1	Measurement report: Occurrence of aminiums
2	in PM _{2.5} during winter in China: aminium
3	outbreak during polluted episodes and potential
4	constraints
5	
6	Yu Xu ^{1,2} , Tang Liu ¹ , Yi-Jia Ma ¹ , Qi-Bin Sun ³ , Hong-Wei Xiao ^{1,2} , Hao Xiao ^{1,2} , Hua-
7	Yun Xiao ^{1,2} *, Cong-Qiang Liu ⁴
8	
9	¹ School of Agriculture and Biology, Shanghai Jiao Tong University, Shanghai 200240,
10	China
11	² Shanghai Yangtze River Delta Eco-Environmental Change and Management
12	Observation and Research Station, Ministry of Science and Technology, Ministry of
13	Education, Shanghai 200240, China
14	³ Dongguan Meteorological Bureau, Dongguan, Guangdong, 523086, China
15	⁴ Institute of Surface-Earth System Science, School of Earth System Science, Tianjin
16	University, Tianjin 300072, China
17	
18 19	
	*Company ding outhous
20	*Corresponding authors
21	Hua-Yun Xiao
22	E-mail: xiaohuayun@sjtu.edu.cn
23	
24	

Abstract: Amines and aminiums play an important role in particle formation, liquid-25 phase reactions, and climate change, attracting considerable attention over the years. 26 27 Here, we investigated the concentrations and compositions of aminiums in PM_{2.5} in 11 Chinese cities during the winter, focusing on the characteristics of aminiums 28 during the polluted days and the key factors influencing aminium outbreak. 29 Monomethylaminium was the dominant aminium species in most cities except 30 Taiyuan and Guangzhou, followed by dimethylaminium. Diethylaminium dominated 31 the total aminiums in Taiyuan and Guangzhou. Thus, the main amine sources in 32 33 Taiyuan and Guangzhou were significantly different from those in other cities. The concentrations of the total aminiums (TA) in all cities increased significantly during 34 the polluted days, with weak aminium outbreaks in Xi'an and Beijing. Additionally, 35 36 the concentrations of TA in Xi'an and Beijing were insignificantly correlated with those of PM_{2.5} and the major acidic aerosol components, while the opposite pattern 37 was observed in 9 other cities. Thus, acid-base chemistry was significantly associated 38 39 with the formation of aminiums in PM_{2.5} in all cities except Xi'an and Beijing. Based on the sensitivity analysis of the aminiums/ammonium ratio to ammonium changes as 40 well as excluding the effects of relative humidity and atmospheric oxidation, we 41 proposed the possibility of the competitive uptake of ammonia versus amines on 42 acidic aerosols or the displacement of aminiums by ammonia in Xi'an and Beijing 43 (constraining aminium outbreaks). Overall, this study deepens the understanding of 44 the spatiotemporal differences in aminium characteristic and formation in China. 45 However, the uptake of amines on particles to form aminiums and the relevant 46

47 influencing factors require further mechanistic research.

48

Keywords: Aminiums, PM_{2.5} pollution, Aerosol acidity, Spatiotemporal variations,
Formation mechanism

- 51
- 52

53 **1. Introduction**

54 Low-molecular-weight amines are ubiquitous and important in the gaseous and 55 particulate phases (Nielsen et al., 2012; Ge et al., 2011a; Berta et al., 2023). More than 150 amines have been identified in the atmosphere (Ge et al., 2011a). The most 56 abundant and frequently reported amines in field observations are typically C1-C6 57 58 alkylamines including dimethylamine, monomethylamine, trimethylamine, diethylamine, ethylamine, 1-propanamine, and 1-butanamine (Yang et al., 2023b; Liu 59 et al., 2023). Amines can participate in various chemical and physical processes in the 60 61 atmosphere, promoting the formation and growth of new particles and contributing to the production of secondary organic aerosols (Yao et al., 2018; Tong et al., 2020; 62 Møller et al., 2020). Amines are thus considered to have a direct or indirect impact on 63 air quality (Li et al., 2019; Tao et al., 2016; Shen et al., 2023). Air pollution (e.g., 64 haze) caused by high levels of atmospheric fine particles (PM2.5) has received 65 considerable attention in China over the past decade due to rapid industrialization and 66 urbanization (Liu et al., 2022b; Liu et al., 2022c). Evidently, controlling the emission 67 strength of amine sources and understanding the transformation of atmospheric 68

69 amines can effectively reduce air pollution in cities.

The main sources of atmospheric amines during the air pollution period in cities 70 71 in China are typically fossil fuel combustion and biomass burning rather than agricultural emissions (Feng et al., 2022; Liu et al., 2022c; Wang et al., 2022; Shen et 72 al., 2017; Ho et al., 2016; Chang et al., 2022). Owing to the water solubility and 73 alkalinity of amines, low-molecular-weight amines in PM2.5 during the air pollution 74 period are mainly present in the form of amine salts (i.e., aminiums) via the gas-to-75 particle partitioning of gaseous amines and subsequent acid-base chemistry (Zhang et 76 77 al., 2021; Liu et al., 2022a; Ge et al., 2011a; Xie et al., 2018). It should be noted that organic amines (e.g., dimethylamine and trimethylamine) in nanoparticles (<200 nm) 78 may also be largely present in the organic phase (Xie et al., 2018). In addition, 79 80 oxidative degradation of higher-molecular-weight amines and displacement reactions are also potential formation pathways of aminiums in PM_{2.5} (Tao et al., 2021; Qiu and 81 Zhang, 2013; Tong et al., 2020). Although previous observational studies have 82 83 investigated the compositions, concentrations, sources, and formation processes of low-molecular-weight aminiums in the particle phase in urban areas of Shanghai (Liu 84 et al., 2023), Guangzhou (Shu et al., 2023), Qingdao (Liu et al., 2022c), Xuzhou 85 (Yang et al., 2023b), China, there has been relatively little focus on the association 86 between PM_{2.5} and amine outbreaks. A recent study conducted in Wangdu County, 87 Hebei Province, China has suggested that amines exhibited outbreak characteristics 88 during the haze episode (Feng et al., 2022). Climate and air pollution conditions can 89 vary greatly from city to city due to the vastness of China. However, it is poorly 90

understood how the characteristics and formation processes of low-molecular-weight
aminiums in PM_{2.5} vary between clean and polluted days in different cities in China,
which may hinder the further assessment of the environmental impacts of amines with
regional differences.
In winter in China, air pollution episodes are more frequent compared to other

seasons. Thus, we present the measurements of aminiums in $PM_{2.5}$ collected from 11 different Chinese cities during the winter (2017–2018). The aims of this study are (1) to investigate the spatial differences in the compositions and concentrations of aminiums in $PM_{2.5}$, with a focus on the difference between them on clean days and polluted days, and (2) to understand the key factors controlling the formation of aminiums in $PM_{2.5}$ in different cities.

102

103 **2. Materials and Methods**

104 **2.1. Site Description and Sample Collection**

105 A total of eleven urban sites were selected for aerosol sample collection, including Beijing (BJ; 116.41°E, 40.04°N), Taiyuan (TY; 112.58°E, 37.80°N), Xi'an 106 (XA; 108.98°E, 34.25°N), Lanzhou (LZ; 103.73°E, 36.11°N), Haerbin (HEB, i.e., 107 Harbin; 126.64°E, 45.77°N), Wulumuqi (WLMQ, i.e., Urumqi; 87.75°E, 43.86°N), 108 Chengdu (CD; 104.14°E, 30.68°N), Guiyang (GY; 106.73°E, 26.58°N), Guangzhou 109 (GZ; 113.35°E, 23.18°N), Wuhan (WH; 114.36°E, 30.55°N), and Hangzhou (HZ; 110 120.16°E, 30.30°N) sites (Figure S1). HZ and GZ are megacities situated in the 111 Yangtze River Delta (YRD) and Pearl River Delta (PRD) regions respectively, both of 112

which have developed economies. WH is located in the central region of China. CD and GY are representative cities in southwest China. LZ, XA, TY, BJ, and HEB are cities in northern China. WLMQ, located in northwest China, is the largest inland city farthest from the ocean in the world. Obviously, the varying geographical locations and economic development levels of different cities lead to different air pollution and climate conditions between them.

PM_{2.5} sampling in most cities was conducted on the rooftops of buildings (4-6 119 floors in total) using a high-volume air sampler (Series 2031, Laoying, China) from 120 121 December 1, 2017 to January 21, 2018 (winter). Specifically, the sampling periods in LZ, TY, HEB, BJ, XA, WLMQ, CD, WH, HZ, GZ, and GY were Dec. 2-30, 2017, 122 Dec. 2-30, 2017, Dec. 18, 2017 - Jan. 15, 2018, Dec. 22, 2017 - Jan. 21, 2018, Dec. 123 124 22, 2017 – Jan. 20, 2018, Mar. 3–28, 2018, Dec. 1 – 31, 2017, Dec. 6–29, 2017, Dec. 4-31, 2017, Dec. 1-30, 2017, and Dec. 10, 2017 - Jan. 11, 2018, respectively (Tables 125 S1-S3). At each site, $PM_{2.5}$ was sampled once every one to two days for ~24 hours on 126 127 prebaked quartz fiber filters (500 °C for 8 hours). Moreover, two random blank filters were collected. The total number of PM_{2.5} samples at each sampling site was shown in 128 Tables S1-S3. All samples were stored at -30 °C. Meteorological data such as 129 precipitation, wind speed, temperature, and relative humidity (RH), as well as 130 concentrations of various pollutants were recorded during the sampling campaigns 131 from the adjacent environmental monitoring stations. Sampling periods were 132 classified as either clean or polluted days based on a daily average PM_{2.5} mass 133 concentration of 75 μ g m⁻³ (Zhang and Cao, 2015). 134

136 **2.2. Chemical Analysis**

137 The extraction of low-molecular-weight aminiums in the filter samples was carried out using the method described in our recent publication (Liu et al., 2023) and 138 139 in a previous study (Liu et al., 2017). Briefly, the sample was filtered using a 0.22 µm Teflon syringe filter (CNW Technologies GmbH) after extraction with Milli-Q water 140 (~18.2 M Ω cm). The aminiums in the extracts that underwent pH regulation were 141 derivatized using 0.1 mL of benzenesulfonyl chloride (BSC). The tube containing the 142 143 derivatives was sealed and agitated for 30 minutes. To remove excess derivatization reagents, the extracts were agitated again for 30 minutes at 80°C after adding NaOH 144 solution (0.5 mL of 10 mol L^{-1}). Once the mixed solution had cooled down, it was 145 146 acidified with a solution of HCl to adjust the pH to 5.5. A further extraction of derivatives was carried out by adding dichloromethane. It is important to mention that 147 the organic phase was treated with Na₂CO₃ solution and anhydrous Na₂SO₄ 148 sequentially. A stream of nitrogen gas was used to concentrate the organic extracts. 149 Finally, the sample was analyzed using GC-MS after adding dichloromethane and 150 hexamethylbenzene. Dimethylaminium (DMAH⁺), monomethylaminium (MMAH⁺), 151 diethylaminium $(DEAH^{+}),$ ethylaminium $(EAH^{+}),$ propylaminium 152 $(PAH^{+}),$ butylaminium (BAH⁺), and pyrrolidinium (PYRH⁺) were quantified. Aminium 153 recoveries varied between 73% for DMAH⁺ and 112% for PAH⁺. The detection limits 154 of the aminium measurements ranged from 0.8 ng mL⁻¹ for DEAH⁺ to 2.8 ng mL⁻¹ for 155 MMAH⁺. Aminiums are undetectable in the blank. Detailed data quality controls were 156

described in our recent publication (Liu et al., 2023). It should be noted that we did not consider the impact of continuous aging of aminiums collected on the filter on the measurement results. This is mainly due to the following reasons. The PM_{2.5} samples investigated in this study are all acidic (**Tables S1–S3**), promoting the protonation of amino groups. The protonated amino group is difficult to undergo oxidation by oxidants (e.g., hydroxyl radicals and ozone) (Nielsen et al., 2012).

Another filter cut was extracted with Milli-Q water to measure the concentrations of inorganic ions (e.g., NO_3^- , SO_4^{2-} , NH_4^+ , K^+ , Na^+ , Ca^{2+} , and Mg^{2+}) and organic acids (e.g., acetic acid, formic acid, succinic acid, oxalic acid, glutaric acid, and methanesulfonic acid) (Xu et al., 2022a; Xu et al., 2023; Liu et al., 2023; Lin et al., 2023). These inorganic ions were quantified via an ion chromatograph system (Dionex Aquion, Thermo Scientific, USA).

169

170 **2.3. Parameter calculation**

The thermodynamic model (ISORROPIA-II) was used for the prediction of the mass concentration of aerosol liquid water (ALW) and the pH value, which was detailed in our previous studies (Xu et al., 2022b; Xu et al., 2020; Xu et al., 2023). The ventilation coefficient (VC) can be used as an indicator to assess the state of atmospheric dilution of pollutant concentrations (Gani et al., 2019). It is calculated by multiplying the wind speed by the planetary boundary layer height (PBLH) (Yang et al., 2023a).

179 **3. Results and discussion**

180 **3.1.** Compositions of aminiums in PM_{2.5} in China during winter

181 Figure 1 shows the average percentage distributions of various aminiums in PM_{2.5} collected in different cities in China during winter, with a comparison between 182 183 their mass fractions on clean and polluted days. MMAH⁺ was the predominant species among the aminiums investigated in PM_{2.5} in most cities in northern China, including 184 LZ, XA, HEB, BJ, and WLMQ. MMAH⁺ and DMAH⁺ (as the second most abundant 185 species) constituted over 63% of the total aminium concentrations in those northern 186 187 cities. The relatively minor species, including DEAH⁺, EAH⁺, PAH⁺, BAH⁺, and PYRH⁺, contributed between 1% and 18% of the total aminium concentrations. The 188 predominance of MMAH⁺ was also found in cities in the YRD (HZ), central (WH), 189 190 and southwestern (CD and GY) China, closely followed by DMAH⁺. Previous studies conducted in Xi'an (winter, China) (Ho et al., 2015), Beijing (winter, China) (Wang et 191 al., 2022; Ho et al., 2016), Nanjing (winter, China) (Liu et al., 2023) Shanghai (winter, 192 193 China) (Liu et al., 2023), Xiamen (winter, China) (Ho et al., 2016), Hong Kong (winter, China) (Ho et al., 2016), and Arabian Sea (autumn and winter) (Gibb et al., 194 1999), as well as at mountain (autumn, Nanling, China) (Liu et al., 2018) and 195 background (winter, Puding, China) (Liu et al., 2023) sites have suggested that the 196 mass concentration fraction of MMAH⁺ was highest in the measured aerosol amine 197 salts. The Henry's constants of MMA ($3.65 \times 10^1 \text{ mol kg}^{-1} \text{ atm}^{-1}$), DMA (3.14×10^1 198 mol kg⁻¹ atm⁻¹), and EA (3.55×10^1 mol kg⁻¹ atm⁻¹) are relatively lower than those of 199 the other amines investigated (e.g., 1.32×10^2 mol kg⁻¹ atm⁻¹ for DEA) (Ge et al., 200

201 2011b), implying that the potential of MMA, DMA, and EA to be partitioned into 202 aqueous particles was weaker compared to DEA. Additionally, the gaseous forms of 203 these determined aminiums typically have strong alkalinity (Ge et al., 2011b). The 204 aerosol samples in this study were all acidic (**Tables S1–S3**). Thus, these results imply 205 that the increased emissions of MMA and DMA may partially explain the higher 206 abundance of MMAH⁺ and DMAH⁺ in PM_{2.5} in these investigated cities during winter.



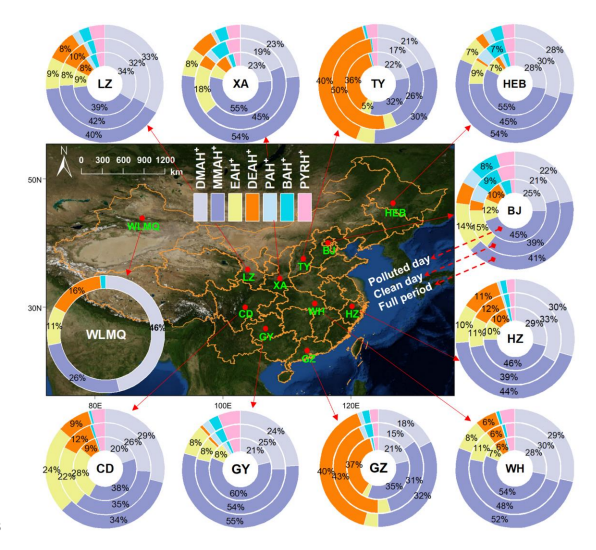




Figure 1. Average percentage distributions of various aminiums in PM_{2.5} collected in different cities in China during winter. The map was obtained from [©]MeteoInfoMap (version 3.3.0) (Chinese Academy of Meteorological Sciences, China).

213	In another northern city (i.e., TY), DEAH ⁺ was the most abundant aminium
214	species (40% of the total aminium concentrations), followed by $MMAH^+$ (30%) and
215	DMAH ⁺ (21%). The composition characteristic of aminiums in the city of GZ (PRD
216	area) was similar to that observed in TY (Figure 1). Anthropogenic emissions,
217	including vehicle exhaust and industrial production are considered to be the main
218	contributors to aerosol DEAH ⁺ in urban areas (Chen et al., 2022b; Chen et al., 2019;
219	Yang et al., 2023b; Chang et al., 2022). A recent study has suggested that ethanol
220	gasoline vehicles can emit a large amount of ethyl-amines, leading to the outbreak of
221	DEAH ⁺ during the haze episodes in Hebei Province (North China) (Feng et al., 2022).
222	Thus, the relative emission strength of anthropogenic DEA in the investigated amines
223	was probably higher in TY (an inland city with application of ethanol gasoline
224	vehicles) than in other cities. In addition, previous studies have suggested that aerosol
225	DEAH ⁺ can also be largely derived from marine emissions (Facchini et al., 2008;
226	Dall'osto et al., 2019). Since GZ is a developed coastal city, local aerosol aminiums
227	may be influenced by large gaseous DEA inputs from both local industrial production
228	and marine sources.

The mass concentration fractions of aminiums on clean and polluted days were also compared (**Figure 1**). The dominant aminium species (i.e., $MMAH^+$, $DMAH^+$, or DEAH⁺) in PM_{2.5} in all cities were not replaced by other aminiums from the clean days to the polluted days. This likely suggests that the main sources of atmospheric gas-phase amines in the cities did not change significantly on the polluted days. In addition, the proportions of MMAH⁺ and DMAH⁺ tended to further increase from the
clean days to the polluted days, while that of DEAH⁺ with relatively low solubility
showed a decreasing trend, especially in TY and GZ (where DEAH⁺ was dominant).
The concentrations of ALW in PM_{2.5} were generally much higher on polluted days
than on clean days, especially in the northern cities (**Tables S1–S3**). Clearly, liquidphase processes likely played an important role in the formation of aminiums on
polluted days.

241

242 **3.2.** Aminium concentrations and their linkage with PM_{2.5} variations

Figure 2 shows the average concentration distributions of various aminiums in 243 PM_{2.5} collected in different cities in China during winter, focusing on the difference 244 245 between their concentrations on clean days and polluted days. The concentrations of total aminiums (TA) in TY ranged from 17.50 to 149.00 ng m⁻³, with an average of 246 56.90 ± 41.81 ng m⁻³. This average TA level was the highest among all the cities 247 248 investigated. The average concentration of TA in WLMQ was found to be the lowest $(4.16 \pm 1.24 \text{ ng m}^{-3})$, with a range of 2.10–6.50 ng m⁻³. As previously mentioned, 249 250 WLMQ is a vast city with a lower population density and less developed industries compared to the more developed northern and coastal cities in China. Additionally, 251 this region is surrounded by barren mountains and sandy land (Ma et al., 2024) 252 (Figure 2). Apparently, the weak amine emission intensity appears to be responsible 253 254 for the low levels of aminiums in the WLMQ.

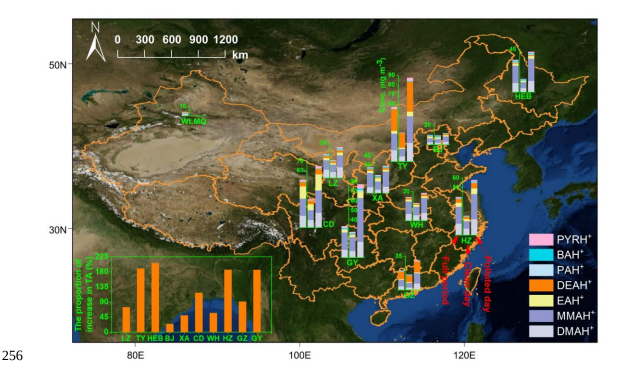


Figure 2. Average concentration distributions of various aminiums in PM_{2.5} collected in different cities in the winter in China. The stacked bar chart from left to right indicates the data for the full sampling period, the clean day, and the polluted day in turn. The column chart in the bottom left corner shows the proportion of the increase in TA concentration from the clean days to the polluted days. The map was obtained from [©]MeteoInfoMap (version 3.3.0) (Chinese Academy of Meteorological Sciences, China).

Table S4 provides an overview of the aminiums detected in atmospheric fine particles detected in different seasons and regions. The ranges of average TA concentrations in the northern cities (i.e., HEB, BJ, TY, XA, LZ, and WLMQ) generally overlapped with those measured in the coastal (GZ and HZ), central (WH), and southwestern (CD and GY) cities in this study (Tables S1–S3). Moreover, the average TA concentrations investigated here (4.16 ng m⁻³ – 56.90 ng m⁻³) were also

271	within the observation ranges reported in previous studies (1.49 ng m ⁻³ – 329.80 ng m ⁻
272	³) (Table S4) (Ho et al., 2016; Liu et al., 2023; Shen et al., 2017; Huang et al., 2016;
273	Choi et al., 2020; Liu et al., 2018; Shu et al., 2023). MMAH ⁺ , as the dominant
274	aminium species in most of cities, showed the highest (18.33 \pm 12.82 ng m^-3) and
275	lowest (1.07 \pm 0.55 ng m ⁻³) average concentrations in HEB and WLMQ, respectively.
276	$\mathrm{DEAH}^{\scriptscriptstyle +}$ was the most abundant aminium species in TY and GZ, with average
277	concentrations of 22.62 \pm 17.62 ng m^{-3} and 8.16 \pm 4.65 ng m^{-3}, respectively (Tables
278	S1 and S3). Two previous studies conducted in the GZ area in winter (2021 and 2015–
279	2016) showed similar average DEAH ⁺ (~ 7 ng m ⁻³) levels to this study (Liu et al.,
280	2022b; Shu et al., 2023). However, $DEAH^+$ was not identified as the dominant
281	aminium component in those two previous studies. Furthermore, lower aminium
282	concentrations (< 8 ng m ⁻³) were generally found in most of the marine and polar
283	regions (Dall'osto et al., 2019; Corral et al., 2022). In general, the concentration and
284	composition of aminiums vary spatially, which may be attributed to spatial differences
285	in amine sources, emission intensities, and the main factors affecting aminium
286	formation.

The average concentrations of TA in all the investigated cities exhibited a similar variation pattern from clean to polluted days, which was characterized by higher levels on polluted days (**Figure 2**). Specifically, the average aminium concentration showed an increase of up to 206% in HEB during the polluted period. TA concentrations in LZ, TY, CD, HZ, and GZ also increased greatly by 91% (in GZ) -190% (in TY). It seems that PM_{2.5} pollution can be accompanied by an outbreak of

aminiums. In contrast, a relatively small percentage increase in TA concentration 293 during the polluted days was found in WH (57%), XA (50%), and BJ (25%). To 294 295 further explore the linkage between changes in PM_{2.5} and fluctuations in aminiums, the temporal variations in the mass concentrations of aminiums and PM2.5 were 296 compared across various cities (Figure 3). The concentrations of total and major 297 aminiums in LZ, TY, HEB, WLMQ, CD, WH, HZ, GZ, and GY showed a temporal 298 variation highly similar to that of PM_{2.5}, as indicated by a significant correlation 299 between TA and PM_{2.5} in these cities (r = 0.61-0.85, P < 0.05). However, high levels 300 of PM_{2.5} can correspond to low levels of aminiums in XA (e.g., Dec. 29 and Jan. 2, 301 14, 15, and 16) and BJ (e.g., Dec. 28, 30). The correlations between TA and PM_{2.5} in 302 these two cities were also insignificant (P > 0.05). These results suggest that the 303 304 formation of aminiums in XA and BJ during the polluted period may be constrained by some special factors, which will be discussed in the following discussion. 305

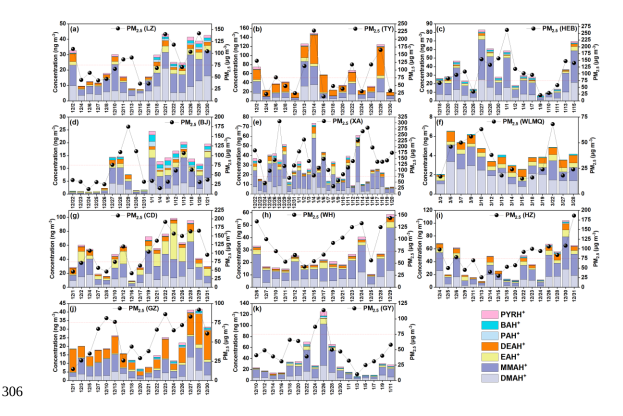


Figure 3. Temporal variations in the mass concentrations of aminiums and PM_{2.5}
observed at the (a) LZ, (b) TY, (c) HEB, (d) BJ, (e) XA, (f) WLMQ, (g) CD, (h) WH,
(i) HZ, (j) GZ, and (k) GY sites.

311 3.3. Formation of aminiums and potential ammonia suppression in aminium 312 outbreaks

It is well documented that aminiums in PM_{2.5} can be formed mainly via the uptake of their gaseous form (i.e., amines) by aqueous particles, followed by acidbase neutralization reactions (Ge et al., 2011b; Xie et al., 2018; Sauerwein and Chan, 2017; Qiu and Zhang, 2013; Liu et al., 2023). Clearly, the formation of particle-phase aminiums was closely associated with the origins of the corresponding gas-phase amines (as precursors of aminiums). We found that TA and major aminiums (e.g., MMAH⁺, DMAH⁺, and DEAH⁺) showed a significant positive correlation (P < 0.05)

320	with either SO ₂ , NO ₂ , or K^+ (as indicators of fuel combustion and biomass burning
321	(Tian et al., 2020; Liu et al., 2023; Kunwar and Kawamura, 2014)) in LZ, TY, HEB,
322	BJ, WLMQ, CD, WH, HZ, GZ, and GY (Figure 4 and Figure S2). Thus, although
323	lacking sufficient indicators (e.g., biogenic source traces) to trace the source of
324	amines, our results can at least indicate that fossil fuel combustion or biomass burning
325	may be important contributors to atmospheric amines in most of the investigated cities
326	during the winter. This consideration was also supported by previous studies about the
327	potential source analysis of aerosol aminiums in Guangzhou, Xuzhou, and Wulumuqi
328	during the winter (Yang et al., 2023b; Shu et al., 2023; Ma et al., 2024). In contrast,
329	the concentrations of TA in XA were weakly correlated ($P > 0.05$) with those of K ⁺ ,
330	SO ₂ , and NO ₂ . Several studies conducted in XA have suggested that aerosol nitrogen-
331	containing organic compounds can be largely derived from fossil fuel combustion and
332	biomass burning (Zhang et al., 2023a; Zhang et al., 2023b; He et al., 2023; Yang et al.,
333	2024). Moreover, the traditional method of identifying amine sources through
334	correlation analysis (Berta et al., 2023; Liu et al., 2022b; Liu et al., 2022a; Huang et
335	al., 2022; Corral et al., 2022) can also have significant uncertainties, as implied by the
336	following two cases. First, the uptake of amines by aerosol particles might be
337	constrained by low ALW concentration, weak particle acidity, or high ammonia levels
338	(Liu et al., 2022b; Chen et al., 2022a; Ge et al., 2011b; Sauerwein and Chan, 2017;
339	Chan and Chan, 2013; Wang et al., 2010). Second, amines might be largely
340	decomposed by atmospheric oxidants (e.g., hydroxyl radical and ozone) (Nielsen et
341	al., 2012; Qiu and Zhang, 2013). Thus, the abovementioned weak correlations

between aminiums and indicators in XA cannot definitely indicate that the contributions of fossil fuel combustion and biomass burning to amines in XA were insignificant. Presumably, the prerequisite for amine source apportionment using the correlation between aminiums and indicators is that the gas-phase amines can be largely converted into aminiums in $PM_{2.5}$ through secondary processes without the influence of constrained factors. To further explore this issue, the following discussion focuses on the main factors affecting the formation of aminiums in particles.

349

350

(a)	PM _{2.5}	NH_4^+	SO4-	NO ₃	Total org. acids	s K⁺	o,	(b)	PM _{2.5}	NH_4^*	SO4-	NO ₃	Total org. acid	s K ⁺	O _x	(C)	PM _{2.5}	NH_4^*	SO4-	NO3	Total org. acids	K+	O _x
TA (LZ)	0.75	0.69	0.78	0.71	Х	0.77	Х	TA (TY)	0.85	0.87	0.89	0.87	0.76	0.80	Х	TA (HEB)	0.70	0.72	0.72	0.59	0.64	0.75	0.49
TA (BJ)	Х	Х	Х	Х	0.59	0.59	-0.75	TA (XA)	Х	Х	Х	Х	Х	Х	-0.47	TA (WLMQ)	0.72	0.70	0.74	0.70	Х	Х	Х
TA (CD)	0.78	0.79	0.80	0.82	0.86	0.84	Х	TA (WH)	0.75	0.58	0.63	0.75	0.74	Х	Х	TA (HZ)	0.61	0.84	0.86	0.86	0.78	0.62	Х
TA (GZ)	0.63	0.64	0.78	Х	0.60	0.54	Х	TA (GY)	0.72	0.80	0.86	0.74	0.76	0.81	Х].	-1 -0.8	-0.6	0.4 -0.:	2 0	0.2 0.4	0.6	0.8 1

Figure 4. Diagrams presenting correlations between the concentrations of TA and other parameters at (a-c) different sites. The colors of the different solid circles indicate different correlation coefficients *r*. The size of the solid circle indicates the significance of the correlation between the two corresponding parameters: the larger circle indicates that the correlation is more significant, while the symbol "×" indicates that the *P*-value is greater than 0.05.

357

The concentrations of TA in LZ, TY, HEB, WLMQ, CD, WH, HZ, GZ, and GY showed significant positive correlations (P < 0.01) with those of the acidic components (e.g., NO₃⁻, SO₄²⁻, organic acids, and acidity (expressed as [(NO₃⁻ + $2SO_4^{2-}) - NH_4^+$] (Feng et al., 2022)), whereas an insignificant correlation (P > 0.05)

was found between them in BJ and XA (Figure 4 and Figure S3). Thus, acid-base 362 chemistry was tightly associated with the formation of aminiums in PM_{2.5} at all sites 363 364 excepting BJ and XA. A recent laboratory study has suggested that amines can be neutralized by H₃O⁺ to form aminiums within picoseconds under conditions of high 365 concentrations of particle sulfuric acid (Zhang et al., 2021). In addition, it has also 366 been found that organic acids (e.g., formic acid) are able to participate in the 367 nucleation of methanesulfonic acid-methylamine through an acid-base reaction 368 (Zhang et al., 2022). The particles are acidic (especially on polluted days) at all study 369 370 sites, with an average pH value ranging from 2.4 to 5.7 (Tables S1-S3). Amines can also partition into the particles by direct dissolution under high RH conditions (Ge et 371 al., 2011b). Significantly increased RH values (i.e., high ALW) (Figure 5a) and acidic 372 373 components (Figure 5b) on polluted days were also observed in XA and BJ. Nevertheless, the insignificant correlation between aminiums and acidic components 374 and ALW concentrations in XA and BJ, together with a relatively small proportional 375 376 increase in aminiums (Figure 2) from clean to polluted days at these two sites suggest that besides acidity and RH, there were other key factors affecting aminium formation 377 in XA and BJ. As we know, the oxidative degradation of amines is one of the main 378 pathways for the removal of atmospheric amines (Qiu and Zhang, 2013; Murphy et 379 al., 2007). Furthermore, for atmospheric oxidants (e.g., hydroxyl radical) reacting 380 with low-molecular-weight alkylamines, a negative temperature dependence of the 381 rate coefficients has been reported (Nielsen et al., 2012). However, the winter air 382 temperature in northern China was relatively low (< 0 °C in XA and BJ) (Tables S1-383

S3); moreover, there was no significant change in the atmospheric oxidation 384 (indicated by O_x levels ($O_x = O_3 + NO_2$)) of polluted and clean days in XA (higher O_x 385 386 level during clean days) and BJ. In particular, the protonated amino group has been suggested to be difficult to undergo oxidation by hydroxyl radicals and ozone 387 (Nielsen et al., 2012). Accordingly, atmospheric oxidation and temperature may not 388 be the main factors affecting changes in aminium concentrations during clean and 389 polluted days. Furthermore, the insignificant correlation between aminiums and acidic 390 components in XA and BJ suggests that other factors affecting aminium formation 391 must be considered. 392

393

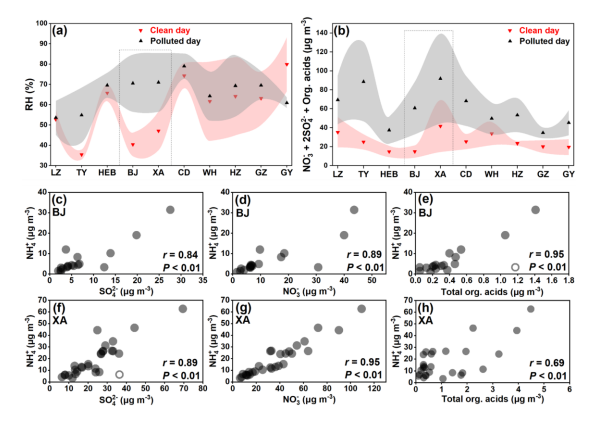


Figure 5. The values of (a) RH and the concentrations of (b) major acidic components (expressed as $NO_3^- + 2SO_4^{2-}$ + total organic acids) on clean and polluted days in different cities. The triangle and the shaded area represent the mean value and

the associated standard deviation, respectively. The correlations of NH_4^+ with the concentrations of NO_3^- , SO_4^{2-} , and total organic acids at (**c**–**e**) BJ and (**f**–**h**) XA. Open circles represent outliers.

401

Furthermore, we found that the concentrations of NH_4^+ were strongly (P < 0.01) 402 correlated with those of acidic components in XA and BJ (Figures 5c-h). This 403 indicates that the acidity of the particles was sufficient for the uptake of ammonia to 404 form ammonium at these two study sites. Typically, the concentration of ammonia in 405 406 the atmosphere is 1 to 3 orders of magnitude higher than that of low-molecular-weight alkylamines (Zheng et al., 2015; You et al., 2014; Yao et al., 2016; Wang et al., 2010). 407 The uptake coefficient of alkylamines on acidic particles is lower than that of 408 409 ammonia (Wang et al., 2010); moreover, Wang et al. (2010) proposed that fresh H₂SO₄ particles can be overwhelmingly neutralized by ammonia when both amines 410 and ammonia are present in the air. In particular, although the strong acidic condition 411 412 was conducive to the formation of aminiums, amines and ammonia may compete for uptake into acidic aerosol particles (Chen et al., 2022a). Thus, the constraint of 413 414 ammonia on amine uptake at much higher ammonia levels than amine levels may be a possible explanation for the insignificant acid-dependent aminium formation in XA 415 416 and BJ (Figures 4a,b).

417 To further explore the role of ammonia (or ammonium) in aminium formation, 418 the average ratios of TA to NH_4^+ on clean and polluted days in different cities were 419 examined (**Figure 6a** and **Table S1–S3**). The average ratios of TA to NH_4^+ were found

420	to be lower in XA and BJ, especially on the polluted days, which was similar to the
421	characteristics of the TA/($NH_3 + NH_4^+$) ratios (Figure S4). The sensitivity analysis of
422	the TA/NH ₄ ⁺ ratio (the lowest in XA and BJ) to NH_4^+ changes (Figures 6b,c and
423	Figure S5) suggests that when TA/NH ₄ ⁺ >2, the NH ₄ ⁺ concentrations in XA and BJ
424	remained at a relatively low level (less than 6 μg m $^{-3}$ and 15 μg m $^{-3}$ in BJ and XA,
425	respectively) with the increase of TA/NH_4^+ ratio, indicating that the formation of
426	aminiums was not limited by ammonia at low amine and ammonium levels (in this
427	case, TA was significantly ($P < 0.01$) correlated with NH ₄ ⁺). When TA/NH ₄ ⁺ < 2, the
428	formation of aminiums may be constrained by higher amine and ammonium levels,
429	which can also be supported by the insignificant $(P > 0.05)$ correlation between TA
430	and $\mathrm{NH_4^+}$ in this case. In contrast, the distributions of the ratios of TA to $\mathrm{NH_4^+}$ in other
431	cities were mainly in regions greater than 2 (Figure S5). The TA concentrations were
432	thus significantly positively correlated with ammonium in these cities (excepting BJ
433	and XA) (Figure 4). A recent study on the uptake of marine aerosol DMA by acidic
434	aerosols has found that the concentrations of particle $DMAH^+$ generally decreased
435	with increasing atmospheric ammonia concentrations (Chen et al., 2022a); moreover,
436	these researchers proposed the possibility that aminiums can be displaced by
437	ammonia in a high ammonia environment. Accordingly, high atmospheric ammonia
438	levels can indeed constrain the conversion of amines to aminiums, even if the aerosol
439	is acidic. In addition, due to the lower VC values (Tables S1-S3) on polluted days
440	compared to clean days, the atmospheric amines were less able to diffuse on polluted
441	days. This may result in an accumulation of aminiums on polluted days via acid-base

chemistry. However, a large decrease in average TA/NH₄⁺ and TA/(NH₃ + NH₄⁺) ratios 442 from clean to polluted days occurred in XA (t-Test, P < 0.05) (Figure 6a, Figure S4, 443 444 and Table S1–S3), followed by BJ. These results indicate that the uptake of amines on acidic particles relative to that of ammonia was significantly reduced from clean to 445 polluted days in XA and BJ. Presumably, the aminiums/ammonium ratio was likely an 446 important indicator to reveal the competitive uptake of ammonia against amines on 447 acidic aerosols, or the displacement of aminiums by ammonia in a high ammonia 448 environment. Thus, this study provides a special field case that emphasizes the 449 potential suppression of ammonia on aminium outbreaks during the polluted days. 450

451

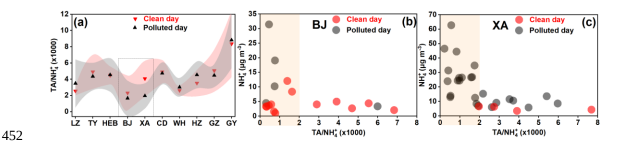


Figure 6. The (a) average ratio of TA to NH_4^+ on clean and polluted days in different cities. The triangle and the shaded area represent the mean value and the associated standard deviation, respectively. Scatterplots of the mass concentration of NH_4^+ with the ratio of TA to NH_4^+ at the (b) BJ and (c) XA sites.

457

458 **4. Conclusions and atmospheric implications**

459 The concentrations, compositions, and temporal and spatial variations of 460 aminiums in $PM_{2.5}$ in 11 different Chinese cities during the winter were systematically 461 investigated to reveal the key factors affecting the aminium outbreak during the

polluted days. Specifically, MMAH⁺ was the dominant species among the aminiums 462 investigated in PM2.5 in most cities, including LZ, XA, HEB, BJ, WLMQ, HZ, WH, 463 CD, and GY, followed by DMAH⁺. In contrast, DEAH⁺ was found to be the most 464 abundant aminium species in TY and GZ, followed by MMAH⁺ and DMAH⁺. This 465 result can be attributed to the fact that the main sources of amines in TY and GZ were 466 significantly different from those in other cities. However, due to the lack of amine 467 emission inventories and sufficient tracers in these investigated cities, this study did 468 not provide a detailed analysis of the specific sources of amines in these investigated 469 470 cities.

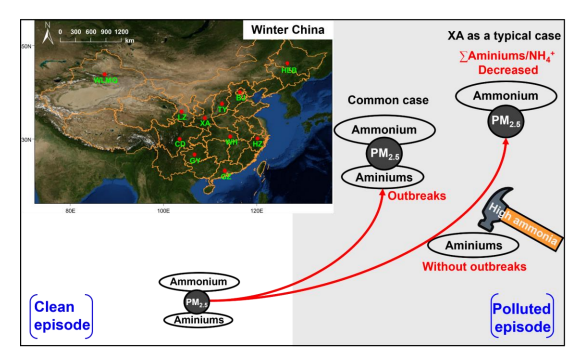
We found that the concentrations of TA and major aminiums in all cities showed 471 a similar pattern of variation from the clean days to the polluted days, which was 472 473 characterized by higher levels on the polluted days. However, the lowest percentage increase in TA concentration during the polluted days was found in XA (50%) and BJ 474 (25%). Moreover, the concentrations of TA in XA and BJ were insignificantly (P >475 476 0.05) correlated with those of PM_{2.5} and the main acidic components in PM_{2.5}. However, the significant correlations of TA with PM2.5 and the main acidic 477 components were observed in other cities. Thus, acid-base chemistry was strongly 478 associated with the formation of aminiums in PM_{2.5} in all cities with the exception of 479 XA and BJ. The concentrations of NH_4^+ were significantly (P < 0.01) correlated with 480 those of the acidic components in XA and BJ, indicating that the acidity of the 481 particles was sufficient for the uptake of ammonia to form ammonium at these two 482 sites. Further, based on the sensitivity analysis of the TA/NH₄⁺ ratio (the lowest in XA 483

and BJ) to NH₄⁺ changes as well as excluding the effects of ALW and atmospheric oxidation, we proposed a possibility about the competitive uptake of ammonia against amines on acidic aerosols in the ambient atmosphere in XA and BJ. This consideration may explain the insignificant acid-dependent aminium formation in XA and BJ. The main finding of this study has been illustrated in a diagram (**Figure 7**).

In general, this study has preliminarily explored the characteristics of aminiums, 489 ammonium, and PM_{2.5} from the clean days to the polluted days according to the 490 observational data from 11 different Chinese cities, highlighting the possibility of the 491 492 competitive uptake of ammonia versus amines on acidic aerosols, or the displacement of aminiums by ammonia under a high ammonia condition. Although a recent study 493 has also demonstrated the possibility of individual aminium being displaced by 494 495 ammonia in an environment of high ammonia level (Chen et al., 2022a), the uptake of amines on particles to form aminiums and the impact mechanisms of relevant 496 influencing factors are still not fully understood. This is because acidity, 497 498 environmental ammonia and amine content, temperature, and liquid-phase reactions all affect the uptake of amines, although acid-base neutralization of amines seems to 499 500 be the most important pathway for amine uptake. Furthermore, if the uptake of amines is significantly constrained by the aforementioned factors, the traditional source 501 apportionment methods using correlation analysis between particle aminiums and 502 tracers will have significant uncertainty due to the weakened partitioning of the 503 amines into the particle phase (i.e., causing insignificant correlations between 504 aminiums and indicators). Further laboratory validation experiments are required to 505

506 substantiate this inference. In addition, it is essential to conduct prolonged 507 observational research in settings with elevated ammonia levels and depleted amine 508 concentrations in the near future.

509



511 **Figure 7.** Conceptual illustration showing the characteristics of aminiums, 512 ammonium, and PM_{2.5} from the clean days to the polluted days. The map was 513 obtained from [©]MeteoInfoMap (version 3.3.0) (Chinese Academy of Meteorological 514 Sciences, China).

515

510

516 **Data availability.** The data in this study are available at 517 https://doi.org/10.5281/zenodo.11102019 (Xu et al., 2024).

518

519	Supplement. Four tables (Tables S1–S4) and five extensive figures (Figures S1–S5)
520	are provided in the Supplement. The supplement related to this article is available
521	online.
522	
523	Author contributions. YX and HYX designed the study. YX, YJM, QBS, HWX, and
524	HX performed field measurements and sample collection; TL performed chemical
525	analysis; YX performed data analysis; YX wrote the original manuscript; and YX,
526	HYX, and CQL reviewed and edited the manuscript.
527	
528	Competing interests. The contact author has declared that none of the authors has
529	any competing interests.
530	
531	Acknowledgements. The authors are very grateful to the editor and the anonymous
532	referees for the kind and valuable comments that improved the paper.
533	
534	Financial support. This study has been kindly supported by the National Natural
535	Science Foundation of China (grant no. 42303081) (Yu Xu) and the Shanghai Sailing
536	Program of Shanghai Science and Technology Commission (grant no. 22YF1418700)
537	(Yu Xu).
538	
539	Review statement. This paper was edited by Roya Bahreini and reviewed by three
540	anonymous referees.

542 **References**

Berta, V. Z., Russell, L. M., Price, D. J., Chen, C. L., Lee, A. K. Y., Quinn, P. K.,
Bates, T. S., Bell, T. G., and Behrenfeld, M. J.: Non-volatile marine and nonrefractory continental sources of particle-phase amine during the North Atlantic
Aerosols and Marine Ecosystems Study (NAAMES), Atmos. Chem. Phys., 23, 27652787, 10.5194/acp-23-2765-2023, 2023.

548 Chan, L. P. and Chan, C. K.: Role of the Aerosol Phase State in 549 Ammonia/Amines Exchange Reactions, Environmental Science & Technology, 47, 550 5755-5762, 10.1021/es4004685, 2013.

551 Chang, Y., Wang, H., Gao, Y., Jing, S. a., Lu, Y., Lou, S., Kuang, Y., Cheng, K.,

552 Ling, Q., Zhu, L., Tan, W., and Huang, R.-J.: Nonagricultural Emissions Dominate

553 Urban Atmospheric Amines as Revealed by Mobile Measurements, Geophysical

554 Research Letters, 49, e2021GL097640, https://doi.org/10.1029/2021GL097640, 2022.

Chen, D., Yao, X., Chan, C. K., Tian, X., Chu, Y., Clegg, S. L., Shen, Y., Gao, Y.,
and Gao, H.: Competitive Uptake of Dimethylamine and Trimethylamine against
Ammonia on Acidic Particles in Marine Atmospheres, Environmental Science &
Technology, 56, 5430-5439, 10.1021/acs.est.1c08713, 2022a.

559 Chen, Y., Lin, Q., Li, G., and An, T.: A new method of simultaneous 560 determination of atmospheric amines in gaseous and particulate phases by gas 561 chromatography-mass spectrometry, Journal of Environmental Sciences, 114, 401-562 411, https://doi.org/10.1016/j.jes.2021.09.027, 2022b.

563	Chen, Y., Tian, M., Huang, R. J., Shi, G., Wang, H., Peng, C., Cao, J., Wang, Q.,
564	Zhang, S., Guo, D., Zhang, L., and Yang, F.: Characterization of urban amine-
565	containing particles in southwestern China: seasonal variation, source, and processing,
566	Atmos. Chem. Phys., 19, 3245-3255, 10.5194/acp-19-3245-2019, 2019.
567	Choi, N. R., Lee, J. Y., Ahn, Y. G., and Kim, Y. P.: Determination of atmospheric
568	amines at Seoul, South Korea via gas chromatography/tandem mass spectrometry,
569	Chemosphere, 258, 127367, 10.1016/j.chemosphere.2020.127367, 2020.
570	Corral, A. F., Choi, Y., Collister, B. L., Crosbie, E., Dadashazar, H., DiGangi, J.
571	P., Diskin, G. S., Fenn, M., Kirschler, S., Moore, R. H., Nowak, J. B., Shook, M. A.,
572	Stahl, C. T., Shingler, T., Thornhill, K. L., Voigt, C., Ziemba, L. D., and Sorooshian,
573	A.: Dimethylamine in cloud water: a case study over the northwest Atlantic Ocean,
574	Environmental Science: Atmospheres, 2, 1534-1550, 10.1039/D2EA00117A, 2022.
575	Dall'Osto, M., Airs, R. L., Beale, R., Cree, C., Fitzsimons, M. F., Beddows, D.,
576	Harrison, R. M., Ceburnis, D., O'Dowd, C., Rinaldi, M., Paglione, M., Nenes, A.,
577	Decesari, S., and Simó, R.: Simultaneous Detection of Alkylamines in the Surface
578	Ocean and Atmosphere of the Antarctic Sympagic Environment, ACS Earth and
579	Space Chemistry, 3, 854-862, 10.1021/acsearthspacechem.9b00028, 2019.
580	Facchini, M. C., Decesari, S., Rinaldi, M., Carbone, C., Finessi, E., Mircea, M.,
581	Fuzzi, S., Moretti, F., Tagliavini, E., Ceburnis, D., and O'Dowd, C. D.: Important
582	Source of Marine Secondary Organic Aerosol from Biogenic Amines, Environmental
583	Science & Technology, 42, 9116-9121, 10.1021/es8018385, 2008.
584	Feng, X., Wang, C., Feng, Y., Cai, J., Zhang, Y., Qi, X., Li, Q., Li, J., and Chen,

Potential Source of Amines from Ethanol Gasoline Vehicle Emission, Environmental 586 587 Science & Technology Letters, 9, 306-311, 10.1021/acs.estlett.2c00145, 2022. Gani, S., Bhandari, S., Seraj, S., Wang, D. S., Patel, K., Soni, P., Arub, Z., Habib, 588 G., Hildebrandt Ruiz, L., and Apte, J. S.: Submicron aerosol composition in the 589 world's most polluted megacity: the Delhi Aerosol Supersite study, Atmos. Chem. 590 Phys., 19, 6843-6859, 10.5194/acp-19-6843-2019, 2019. 591 Ge, X., Wexler, A. S., and Clegg, S. L.: Atmospheric amines - Part I. A review, 592 593 Atmospheric Environment, 45. 524-546, https://doi.org/10.1016/j.atmosenv.2010.10.012, 2011a. 594 Ge, X., Wexler, A. S., and Clegg, S. L.: Atmospheric amines - Part II. 595 596 Thermodynamic properties and gas/particle partitioning, Atmospheric Environment,

Y.: Outbreaks of Ethyl-Amines during Haze Episodes in North China Plain: A

- 45, 561-577, https://doi.org/10.1016/j.atmosenv.2010.10.013, 2011b. 597
- Gibb, S. W., Mantoura, R. F. C., and Liss, P. S.: Ocean-atmosphere exchange and 598 599 atmospheric speciation of ammonia and methylamines in the region of the NW Arabian Global 600 Sea. Biogeochemical Cycles, 13, 161-178, https://doi.org/10.1029/98GB00743, 1999. 601
- He, K., Fu, T., Zhang, B., Xu, H., Sun, J., Zou, H., Zhang, Z., Hang Ho, S. S., 602 Cao, J., and Shen, Z.: Examination of long-time aging process on volatile organic 603 compounds emitted from solid fuel combustion in a rural area of China, 604 Chemosphere, 333, 138957, https://doi.org/10.1016/j.chemosphere.2023.138957, 605 2023. 606

Ho, K.-F., Ho, S. S. H., Huang, R.-J., Chuang, H.-C., Cao, J.-J., Han, Y., Lui, K.-

607

- H., Ning, Z., Chuang, K.-J., Cheng, T.-J., Lee, S.-C., Hu, D., Wang, B., and Zhang, 608 R.: Chemical composition and bioreactivity of PM2.5 during 2013 haze events in 609 China, Atmospheric Environment, 126. 162-170, 610 https://doi.org/10.1016/j.atmosenv.2015.11.055, 2016. 611 Ho, K. F., Ho, S. S. H., Huang, R.-J., Liu, S. X., Cao, J.-J., Zhang, T., Chuang, 612 H.-C., Chan, C. S., Hu, D., and Tian, L.: Characteristics of water-soluble organic 613 nitrogen in fine particulate matter in the continental area of China, Atmospheric 614 615 Environment, 106, 252-261, https://doi.org/10.1016/j.atmosenv.2015.02.010, 2015. Huang, S., Song, Q., Hu, W., Yuan, B., Liu, J., Jiang, B., Li, W., Wu, C., Jiang, 616 F., Chen, W., Wang, X., and Shao, M.: Chemical composition and sources of amines 617 618 in PM2.5 in an urban site of PRD, China, Environmental Research, 212, 113261, https://doi.org/10.1016/j.envres.2022.113261, 2022. 619 Huang, X., Deng, C., Zhuang, G., Lin, J., and Xiao, M.: Quantitative analysis of 620
- aliphatic amines in urban aerosols based on online derivatization and high
 performance liquid chromatography, Environmental Science: Processes & Impacts,
 18, 796-801, 10.1039/C6EM00197A, 2016.
- Kunwar, B. and Kawamura, K.: One-year observations of carbonaceous and
 nitrogenous components and major ions in the aerosols from subtropical Okinawa
 Island, an outflow region of Asian dusts, Atmos. Chem. Phys., 14, 1819-1836.
 https://doi.org/1810.5194/acp-1814-1819-2014, 2014.
- 628 Li, G., Liao, Y., Hu, J., Lu, L., Zhang, Y., Li, B., and An, T.: Activation of NF-κB

pathways mediating the inflammation and pulmonary diseases associated with
atmospheric methylamine exposure, Environmental pollution, 252, 1216-1224,
https://doi.org/10.1016/j.envpol.2019.06.059, 2019.

Lin, X., Xu, Y., Zhu, R.-G., Xiao, H.-W., and Xiao, H.-Y.: Proteinaceous Matter
in PM2.5 in Suburban Guiyang, Southwestern China: Decreased Importance in LongRange Transport and Atmospheric Degradation, J. Geophys. Res.: Atmos., 128,

e2023JD038516, https://doi.org/10.1029/2023JD038516, 2023.

- 636 Liu, C., Li, H., Zheng, H., Wang, G., Qin, X., Chen, J., Zhou, S., Lu, D., Liang,
- 637 G., Song, X., Duan, Y., Liu, J., Huang, K., and Deng, C.: Ocean Emission Pathway
- and Secondary Formation Mechanism of Aminiums Over the Chinese Marginal Sea,
- 639 Journal of Geophysical Research: Atmospheres, 127, e2022JD037805,
 640 https://doi.org/10.1029/2022JD037805, 2022a.
- Liu, F., Bi, X., Zhang, G., Peng, L., Lian, X., Lu, H., Fu, Y., Wang, X., Peng, P.
 a., and Sheng, G.: Concentration, size distribution and dry deposition of amines in
 atmospheric particles of urban Guangzhou, China, Atmospheric Environment, 171,
 279-288, https://doi.org/10.1016/j.atmosenv.2017.10.016, 2017.
- Liu, F., Zhang, G., Lian, X., Fu, Y., Lin, Q., Yang, Y., Bi, X., Wang, X., Peng, P. a., and Sheng, G.: Influence of meteorological parameters and oxidizing capacity on characteristics of airborne particulate amines in an urban area of the Pearl River Delta, China, Environmental Research, 212, 113212, https://doi.org/10.1016/j.envres.2022.113212, 2022b.
- Liu, F., Bi, X., Zhang, G., Lian, X., Fu, Y., Yang, Y., Lin, Q., Jiang, F., Wang, X.,

651	Peng, P. a., and Sheng, G.: Gas-to-particle partitioning of atmospheric amines
652	observed at a mountain site in southern China, Atmospheric Environment, 195, 1-11,
653	https://doi.org/10.1016/j.atmosenv.2018.09.038, 2018.
654	Liu, T., Xu, Y., Sun, QB., Xiao, HW., Zhu, RG., Li, CX., Li, ZY., Zhang,
655	KQ., Sun, CX., and Xiao, HY.: Characteristics, Origins, and Atmospheric
656	Processes of Amines in Fine Aerosol Particles in Winter in China, J. Geophys. Res.:
657	Atmos., 128, e2023JD038974, https://doi.org/10.1029/2023JD038974, 2023.
658	Liu, Z., Li, M., Wang, X., Liang, Y., Jiang, Y., Chen, J., Mu, J., Zhu, Y., Meng,
659	H., Yang, L., Hou, K., Wang, Y., and Xue, L.: Large contributions of anthropogenic
660	sources to amines in fine particles at a coastal area in northern China in winter,
661	Science of The Total Environment, 839, 156281,
662	https://doi.org/10.1016/j.scitotenv.2022.156281, 2022c.
663	Ma, Y. J., Xu, Y., Yang, T., Xiao, H. W., and Xiao, H. Y.: Measurement report:
664	Characteristics of nitrogen-containing organics in PM2.5 in Ürümqi, northwestern
665	China - differential impacts of combustion of fresh and aged biomass materials,
666	Atmos. Chem. Phys., 24, 4331-4346, 10.5194/acp-24-4331-2024, 2024.
667	Møller, K. H., Berndt, T., and Kjaergaard, H. G.: Atmospheric Autoxidation of

- 668 Amines, Environmental Science & Technology, 54, 11087-11099,
 669 10.1021/acs.est.0c03937, 2020.
- 670 Murphy, S. M., Sorooshian, A., Kroll, J. H., Ng, N. L., Chhabra, P., Tong, C.,
- 671 Surratt, J. D., Knipping, E., Flagan, R. C., and Seinfeld, J. H.: Secondary aerosol
- 672 formation from atmospheric reactions of aliphatic amines, Atmos. Chem. Phys., 7,

673 2313-2337, 10.5194/acp-7-2313-2007, 2007.

674	Nielsen, C. J., Herrmann, H., and Weller, C.: Atmospheric chemistry and
675	environmental impact of the use of amines in carbon capture and storage (CCS),
676	Chemical Society Reviews, 41, 6684-6704, 10.1039/C2CS35059A, 2012.
677	Qiu, C. and Zhang, R.: Multiphase chemistry of atmospheric amines, Physical
678	Chemistry Chemical Physics, 15, 5738-5752, 10.1039/C3CP43446J, 2013.
679	Sauerwein, M. and Chan, C. K.: Heterogeneous uptake of ammonia and
680	dimethylamine into sulfuric and oxalic acid particles, Atmos. Chem. Phys., 17, 6323-
681	6339, 10.5194/acp-17-6323-2017, 2017.
682	Shen, W., Ren, L., Zhao, Y., Zhou, L., Dai, L., Ge, X., Kong, S., Yan, Q., Xu, H.,
683	Jiang, Y., He, J., Chen, M., and Yu, H.: C1-C2 alkyl aminiums in urban aerosols:
684	Insights from ambient and fuel combustion emission measurements in the Yangtze
685	River Delta region of China, Environmental pollution, 230, 12-21,
686	https://doi.org/10.1016/j.envpol.2017.06.034, 2017.
687	Shen, X., Chen, J., Li, G., and An, T.: A new advance in the pollution profile,
688	transformation process, and contribution to aerosol formation and aging of
689	atmospheric amines, Environmental Science: Atmospheres, 3, 444-473,
690	10.1039/D2EA00167E, 2023.

Shu, Q., Pei, C., Lin, X., Hong, D., Lai, S., and Zhang, Y.: Variations of
aminiums in fine particles at a suburban site in Guangzhou, China: Importance of
anthropogenic and natural emissions, Particuology, 80, 140-147,
https://doi.org/10.1016/j.partic.2022.11.019, 2023.

695	Tao, Y., Liu, T., Yang, X., and Murphy, J. G.: Kinetics and Products of the
696	Aqueous Phase Oxidation of Triethylamine by OH, ACS Earth and Space Chemistry,
697	5, 1889-1895, 10.1021/acsearthspacechem.1c00162, 2021.

Tao, Y., Ye, X., Jiang, S., Yang, X., Chen, J., Xie, Y., and Wang, R.: Effects of
amines on particle growth observed in new particle formation events, Journal of
Geophysical Research: Atmospheres, 121, 324-335,
https://doi.org/10.1002/2015JD024245, 2016.

Tian, D., Fan, J., Jin, H., Mao, H., Geng, D., Hou, S., Zhang, P., and Zhang, Y.:
Characteristic and Spatiotemporal Variation of Air Pollution in Northern China Based
on Correlation Analysis and Clustering Analysis of Five Air Pollutants, Journal of
Geophysical Research: Atmospheres, 125, e2019JD031931,
https://doi.org/10.1029/2019JD031931, 2020.

Tong, D., Chen, J., Qin, D., Ji, Y., Li, G., and An, T.: Mechanism of atmospheric 707 organic amines reacted with ozone and implications for the formation of secondary 708 709 organic aerosols, Science of The Total Environment, 737, 139830, https://doi.org/10.1016/j.scitotenv.2020.139830, 2020. 710

Wang, L., Lal, V., Khalizov, A. F., and Zhang, R.: Heterogeneous Chemistry of
Alkylamines with Sulfuric Acid: Implications for Atmospheric Formation of
Alkylaminium Sulfates, Environmental Science & Technology, 44, 2461-2465,
10.1021/es9036868, 2010.

Wang, M., Wang, Q., Ho, S. S. H., Li, H., Zhang, R., Ran, W., Qu, L., Lee, S.-c.,
and Cao, J.: Chemical characteristics and sources of nitrogen-containing organic

compounds at a regional site in the North China Plain during the transition period of
autumn and winter, Science of The Total Environment, 812, 151451,
https://doi.org/10.1016/j.scitotenv.2021.151451, 2022.

Xie, H., Feng, L., Hu, Q., Zhu, Y., Gao, H., Gao, Y., and Yao, X.: Concentration
and size distribution of water-extracted dimethylaminium and trimethylaminium in
atmospheric particles during nine campaigns - Implications for sources, phase states
and formation pathways, Science of The Total Environment, 631-632, 130-141,
https://doi.org/10.1016/j.scitotenv.2018.02.303, 2018.

Xu, Y., Dong, X.-N., Xiao, H.-Y., He, C., and Wu, D.-S.: Water-Insoluble
Components in Rainwater in Suburban Guiyang, Southwestern China: A Potential
Contributor to Dissolved Organic Carbon, Journal of Geophysical Research:
Atmospheres, 127, e2022JD037721, https://doi.org/10.1029/2022JD037721, 2022a.

Xu, Y., Dong, X.-N., Xiao, H.-Y., Zhou, J.-X., and Wu, D.-S.: Proteinaceous
Matter and Liquid Water in Fine Aerosols in Nanchang, Eastern China: Seasonal
Variations, Sources, and Potential Connections, J. Geophys. Res.: Atmos., 127,
e2022JD036589. https://doi.org/036510.031029/032022JD036589, 2022b.

Xu, Y., Dong, X. N., He, C., Wu, D. S., Xiao, H. W., and Xiao, H. Y.: Mist
cannon trucks can exacerbate the formation of water-soluble organic aerosol and
PM2.5 pollution in the road environment, Atmos. Chem. Phys., 23, 6775-6788,
10.5194/acp-23-6775-2023, 2023.

Xu, Y., Miyazaki, Y., Tachibana, E., Sato, K., Ramasamy, S., Mochizuki, T.,
Sadanaga, Y., Nakashima, Y., Sakamoto, Y., Matsuda, K., and Kajii, Y.: Aerosol

Liquid Water Promotes the Formation of Water-Soluble Organic Nitrogen in
Submicrometer Aerosols in a Suburban Forest, Environ. Sci. Technol., 54, 1406-1414.

https://doi.org/1410.1021/acs.est.1409b05849, 2020.

- Yang, T., Xu, Y., Ye, Q., Ma, Y. J., Wang, Y. C., Yu, J. Z., Duan, Y. S., Li, C. X.,
 Xiao, H. W., Li, Z. Y., Zhao, Y., and Xiao, H. Y.: Spatial and diurnal variations of
 aerosol organosulfates in summertime Shanghai, China: potential influence of
 photochemical processes and anthropogenic sulfate pollution, Atmos. Chem. Phys.,
 23, 13433-13450, 10.5194/acp-23-13433-2023, 2023a.
- Yang, X.-Y., Cao, F., Fan, M.-Y., Lin, Y.-C., Xie, F., and Zhang, Y.-L.: Seasonal
 variations of low molecular alkyl amines in PM2.5 in a North China Plain industrial
 city: Importance of secondary formation and combustion emissions, Science of The
 Total Environment, 857, 159371, https://doi.org/10.1016/j.scitotenv.2022.159371,
 2023b.
- Yang, X., Huang, S., Li, D., Xu, H., Zeng, Y., Yang, L., Wang, D., Zhang, N.,
 Cao, J., and Shen, Z.: Water-soluble organic matter with various polarities in PM2.5
 over Xi'an, China: Abundance, functional groups, and light absorption, Particuology,
 84, 281-289, https://doi.org/10.1016/j.partic.2023.07.005, 2024.
- Yao, L., Wang, M. Y., Wang, X. K., Liu, Y. J., Chen, H. F., Zheng, J., Nie, W.,
 Ding, A. J., Geng, F. H., Wang, D. F., Chen, J. M., Worsnop, D. R., and Wang, L.:
 Detection of atmospheric gaseous amines and amides by a high-resolution time-offlight chemical ionization mass spectrometer with protonated ethanol reagent ions,
 Atmos. Chem. Phys., 16, 14527-14543, 10.5194/acp-16-14527-2016, 2016.

761	Yao, L., Garmash, O., Bianchi, F., Zheng, J., Yan, C., Kontkanen, J., Junninen,
762	H., Mazon, S. B., Ehn, M., Paasonen, P., Sipilä, M., Wang, M., Wang, X., Xiao, S.,
763	Chen, H., Lu, Y., Zhang, B., Wang, D., Fu, Q., Geng, F., Li, L., Wang, H., Qiao, L.,
764	Yang, X., Chen, J., Kerminen, VM., Petäjä, T., Worsnop, D. R., Kulmala, M., and
765	Wang, L.: Atmospheric new particle formation from sulfuric acid and amines in a
766	Chinese megacity, Science, 361, 278-281, doi:10.1126/science.aao4839, 2018.
767	You, Y., Kanawade, V. P., de Gouw, J. A., Guenther, A. B., Madronich, S., Sierra-
768	Hernández, M. R., Lawler, M., Smith, J. N., Takahama, S., Ruggeri, G., Koss, A.,
769	Olson, K., Baumann, K., Weber, R. J., Nenes, A., Guo, H., Edgerton, E. S., Porcelli,
770	L., Brune, W. H., Goldstein, A. H., and Lee, S. H.: Atmospheric amines and ammonia
771	measured with a chemical ionization mass spectrometer (CIMS), Atmos. Chem.
772	Phys., 14, 12181-12194, 10.5194/acp-14-12181-2014, 2014.
773	Zhang, B., Shen, Z., He, K., Sun, J., Huang, S., Xu, H., Li, J., Ho, S. S. H., and
774	Cao, Jj.: Insight into the Primary and Secondary Particle-Bound Methoxyphenols
775	and Nitroaromatic Compound Emissions from Solid Fuel Combustion and the
776	Updated Source Tracers, Environmental Science & Technology, 57, 14280-14288,
777	10.1021/acs.est.3c04370, 2023a.
778	Zhang, B., Shen, Z., He, K., Zhang, L., Huang, S., Sun, J., Xu, H., Li, J., Yang,
779	L., and Cao, J.: Source Profiles of Particle-Bound Phenolic Compounds and Aromatic

Acids From Fresh and Aged Solid Fuel Combustion: Implication for the Aging

- Mechanism and Newly Proposed Source Tracers, Journal of Geophysical Research:
- Atmospheres, 128, e2023JD039758, https://doi.org/10.1029/2023JD039758, 2023b.

783	Zhang, R., Shen, J., Xie, H. B., Chen, J., and Elm, J.: The role of organic acids in
784	new particle formation from methanesulfonic acid and methylamine, Atmos. Chem.
785	Phys., 22, 2639-2650, 10.5194/acp-22-2639-2022, 2022.
786	Zhang, W., Zhong, J., Shi, Q., Gao, L., Ji, Y., Li, G., An, T., and Francisco, J. S.:
787	Mechanism for Rapid Conversion of Amines to Ammonium Salts at the Air-Particle
788	Interface, Journal of the American Chemical Society, 143, 1171-1178,
789	10.1021/jacs.0c12207, 2021.
790	Zhang, YL. and Cao, F.: Fine particulate matter (PM2.5) in China at a city level,
791	Scientific Reports, 5, 14884, 10.1038/srep14884, 2015.
792	Zheng, J., Ma, Y., Chen, M., Zhang, Q., Wang, L., Khalizov, A. F., Yao, L.,
793	Wang, Z., Wang, X., and Chen, L.: Measurement of atmospheric amines and ammonia
794	using the high resolution time-of-flight chemical ionization mass spectrometry,
795	Atmospheric Environment, 102, 249-259,
796	https://doi.org/10.1016/j.atmosenv.2014.12.002, 2015.
797	