1	Measurement Report: Urban Ammonia and Amines in Houston, Texas
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12 Abstract. Ammonia and amines play critical roles in secondary aerosol formation, especially in 13 urban environments. However, fast measurements of ammonia and amines in the atmosphere are very scarce. We measured ammonia and amines with a chemical ionization mass spectrometer 14 15 (CIMS) at the urban center in Houston, Texas, the fourth most populated urban site in the United 16 States, during October 2022. Ammonia concentrations were on average 4 parts per billion in 17 volume (ppby), while the concentration of an individual amine ranged from several parts per 18 trillion in volume (pptv) to hundreds of pptv. These reduced nitrogen compounds were more 19 abundant during the weekdays than on weekends and correlated with measured CO concentrations, implying they were mostly emitted from pollutant sources. Both ammonia and amines showed a 20 21 distinct diurnal cycle, with higher concentrations in the warmer afternoon, indicating dominant 22 gas-to-particle conversion processes taking place with the changing ambient temperatures. Studies 23 have shown that dimethylamine is critical for new particle formation (NPF) in the polluted 24 boundary layer, but currently, there are no amine emission inventories in global climate models 25 (as opposed to ammonia). Our observations made in very polluted Houston, as well as a less 26 polluted site (Kent, Ohio) from our previous study (You et al., 2014), indicate there is a consistent ratio of dimethylamine over ammonia at these two sites. Thus, our observations can provide a 27 28 relatively constrained proxy of dimethylamine using 0.1% ammonia concentrations at polluted 29 sites in the United States to model NPF processes.

1. Introduction

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31 Atmospheric ammonia and amines are ubiquitous in the atmosphere, and they have been found in the gas phase, aerosol, clouds, and fog droplets (Ge et al., 2011a, b). Ammonia and amines are 32 33 emitted from various natural and anthropogenic sources, such as agricultural activity, animal husbandry, vegetation, soil, waste processing, automobile traffic, power plants, and biomass 34 35 burning (Ge et al., 2011a). Ammonia and amines often share the same emission sources. In general, 36 ambient concentrations of ammonia are at the parts per billion in volume (ppbv) range, and amines 37 are approximately two to three orders of magnitude lower than ammonia concentrations. Ambient 38 concentrations of ammonia and amines vary rapidly due to emission, gas-to-particle conversion, 39 and wet deposition processes (You et al., 2014; Yu and Lee, 2012).

40 Laboratory studies have shown that ammonia and amines play key roles in new particle 41 formation (NPF) as they can stabilize sulfuric acid clusters (Yu et al., 2012; Almeida et al., 2013; Lehtipalo et al., 2018; Xiao et al., 2021; Glasoe et al., 2015; Jen et al., 2016). In particular, 42 43 dimethylamine can have a profound effect on atmospheric processes even at the pptv level 44 (Almeida et al., 2013; Glasoe et al., 2015). Field observations show that ammonia and amines are 45 associated with NPF events in Chinese megacities (Yao et al., 2016; Yan et al., 2021; Cai et al., 2021; Cai et al., 2023), urban areas in the United States (Jen et al., 2016; Smith et al., 2010), 46 47 European cities (Brean et al., 2020), a high altitude site (Bianchi et al., 2016), and the Arctic and Antarctic (Beck et al., 2021; Brean et al., 2021; Jokinen et al.; Köllner et al., 2017). However, 48 49 global models cannot simulate urban NPF processes currently because of the lack of amine

50 emission inventories in models.

Ammonia and amines also contribute to secondary organic aerosol (SOA) formation by condensation of oxidation products formed by reactions with ozone, OH, or NO₃ radicals and produce light-absorbing particles (Erupe et al., 2010; Malloy et al., 2009; Silva et al., 2008; Nielsen, 2016; Nielsen et al., 2012; Qiu and Zhang, 2013). As a result, reducing ammonia emissions has been identified as a cost-effective way to mitigate ambient fine particle concentrations (Gu et al., 2021).

57 Fast-response measurements of ammonia and amines at atmospheric concentrations are very 58 challenging (Lee, 2022), although such measurements are necessary because these reduced 59 nitrogen compounds have relatively short atmospheric lifetimes (Nielsen et al., 2012). Previously, 60 (Schwab et al., 2007) made an intercomparison of six different ammonia detection methods in the 61 laboratory and found a large variance in the measured concentrations and vastly different response 62 times (over several hours) within different instruments. Difficulties in the detection of base 63 compounds also arise because these "sticky" compounds can rapidly adsorb and desorb on/from 64 the surfaces of sampling inlets to cause background signals that vary depending on ambient 65 concentrations, air humidity, and other atmospheric conditions. Thus frequent, in situ 66 measurements of instrument background signals using proper zero gases are required, especially 67 for field observations with rapidly changing ambient concentrations of base compounds.

68 Chemical ionization mass spectrometers (CIMS) using ion reagents such as protonated ethanol, 69 acetone, and water ions can detect ammonia and amines in the atmosphere with fast response 70 (Nowak et al., 2006; Benson et al., 2010; Yu and Lee, 2012; Hanson et al., 2011; Jen et al., 2016; 71 Nowak et al., 2010). As summarized in Table 1, CIMS technique has been used for the detection 72 of ambient ammonia and amines at a polluted site in Ohio (You et al., 2014; Yu and Lee, 2012), 73 a rural Alabama forest (You et al., 2014), and polluted urban sites in China (Zheng et al., 2015; 74 Wang et al., 2020a; Wang et al., 2016; Zhu et al., 2022). As shown in Table 1, there are even fewer 75 studies that simultaneously measured ammonia and amines. The CIMS using ethanol reagent can 76 measure amines at or below single-digit ppty concentrations with a time response of 1 minute and 77 measure simultaneously amines and ammonia (You et al., 2014; Yu and Lee, 2012; Erupe et al., 78 2011; Benson et al., 2010). The CIMS using protonated water ions (i.e., proton-transfer chemical 79 ionization mass spectrometer, PTR-CIMS) can measure mono- and di-amines (Hanson et al., 2011; 80 Jen et al., 2016). Using a high-resolution time-of-flight (HR-TOF) detector coupled to CIMS (HR-81 TOF CIMS) (with ethanol reagent), (Yao et al., 2016) measured various amines and amides in 82 Shanghai. However, isomers of amines were still not resolved in the detection; for example, the 83 measured C2-amines still contained dimethylamine and ethylamine. Thus, a major disadvantage 84 of a mass spectrometer (regardless of mass resolution) is the inability to resolve/identify isomers. 85 To resolve isomers, tandem MS/MS analysis or an additional independent separation method (such 86 as chromatography) coupled to the mass spectrometer is necessary. 87 In situ measurements of ammonia have been made in various atmospheric environments also

with optical techniques such as open-path absorption (Miller et al., 2014), closed-path absorption
(Griffith and Galle, 2000; Ellis et al., 2010; Mcmanus et al., 2010; Leen et al., 2013; Pollack et al.,

90 2019), cavity ring-down spectroscopy (Martin et al., 2016), and photoacoustic spectroscopy

91 (Pushkarsky et al., 2002). These fast-response optical techniques were used for flux and aircraft 92 measurements of ammonia.

93 We measured ammonia and C1-C6 amines with an ethanol CIMS in October 2022 at the urban center in Houston, Texas. Houston is the fourth most populated urban center in the U.S. and 94 95 contains a diverse range of pollutant emissions from urban activity, traffic, ship channels, oil production, marine air masses, and agricultural activity. The primary goal of these measurements 96 is to quantify ammonia and C1-C6 amines in an urban setting and identify the atmospheric 97 conditions that affect their abundance. The study is amongst very few observations of ammonia 98 99 and amines at highly polluted urban sites in the U.S. We also compare observations in Houston 100 with previous measurements taken with the same instrument in Kent, Ohio (less polluted) (You et al., 2014) and establish a quantitative relationship between ammonia and dimethylamine in a 101 102 different range of polluted conditions. This relationship will allow global models to simulate urban 103 NPF processes using the existing ammonia emission inventories.

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2. Methods

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Figure 1. Location of the measurement platform, indicated by a red pin in the center of the map. Nearby commercial, industrial, and residential areas are labeled by yellow, red, and blue shaded sections, respectively. The nearby University of Houston power plant is circled in orange to the southwest of the measurement platform. The map of the greater Houston urban area, as well as the satellite view of the nearby vicinity of the measurement site, are shown in Figure S1.

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The field observation took place in Houston continuously from the 8th to the 27th of October in 2022. Measurements were made at a stationary platform located on the campus of the University of Houston (29.72° N, 95.34° W) ~2.5 km from central downtown Houston. Maps of the measurement site (Figures 1 and S1). The measurement platform was located ~5 m from an active parking lot, ~200 m from a low-traffic road, ~300 m from a high-traffic thoroughfare, and ~500 m from an interstate highway. The immediate vicinity of the site was the University of Houston campus, containing classroom buildings, dormitories, facilities services, and dining halls. Nearby to the southeast of the site were several restaurants as well as an industrial park containing sites of chemical supply companies, construction, machining services, and automobile shops. The site was surrounded by residential areas to the south, northeast, and west. The city center and highest population densities were to the northeast of the measurement site.

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127 The ethanol CIMS instrument used has been described in detail previously (Benson et al., 128 2010; You et al., 2014; Yu and Lee, 2012). The CIMS draws 10 standard liter per minute (slpm) 129 of sample air into a low-pressure ion-molecule region (about 2,000 Pa) where the flow mixes with 130 a pure nitrogen flow with a 2 slpm through a stainless-steel vessel of 200-proof ethanol, followed 131 by a $^{210}P_{o}$ radiation source. Ammonia and amines were detected with the following ion-molecule 132 reactions based on (Erupe et al., 2011), (Yu and Lee, 2012), and (Nowak et al., 2006):

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 $(C_2H_5OH)_nH^+ + NH_3 \rightarrow (C_2H_5OH)_{n-1}NH_4^+ + C_2H_5OH$ $(C_2H_5OH)_nH^+ + B \rightarrow BH^+ + nC_2H_5OH$

Here, "B" refers to amines, and "n" is the number of reagent ions measured by the CIMS (n=1-3). The $(C_2H_5OH)_2H^+$ (m/z = 93) peak was the highest among the three reagent ions (m/z = 47, 93, and 140). As shown in Figure S2, the production ions of amines were protonated ions: C1-amine (m/z = 32), C2 (m/z = 46), C3 (m/z = 60), C4 (m/z = 74), C5 (m/z = 88), and C6 (m/z = 102). Ammonia product ions were NH₄⁺ (m/z = 18, higher peak) and (C₂H₅OH)NH₄⁺ (m/z = 64, lower peak); these two ions were strongly correlated to each other during the ammonia calibration and ambient measurements, indicating they represent ammonia signals.

142 To obtain a background signal, the CIMS is operated with 10 minutes of sampling followed 143 by 10 minutes of background measurements. Figure S2 shows the main reagent and base 144 compound product ions during the switching between ambient and background measurements. 145 Background measurements were taken by switching a 3-way valve to supply the inlet with a flow 146 of zero air through a silicon phosphate medium (Pan Tech, Texas) to scrub ammonia and amines. 147 The reagent signal was taken as the sum of three ethanol reagent ions. Reagent ion signals were 148 typically around 400 kHz with less than 10 % difference between ambient and background 149 measurement modes. Ammonia and amine concentrations were calculated by the difference 150 between the ambient and background signals normalized to 1,000,000 Hz of reagent ion signal 151 multiplied by a calibration factor. Calibration of the instrument was carried out with diluted 152 ammonia in nitrogen and permeation tubes of methylamine, dimethylamine, trimethylamine, 153 diethylamine, and diisopropylamine (Kin-tek, USA). Due to the difficulty of obtaining a 154 calibration standard, C5 amines were assumed to have the same sensitivity as C6 amines. The calibration factors for each compound and detection limits were found to be similar to the results 155 156 from the calibration of the instrument by (You et al., 2014) (Table S1), over a period of nearly 10 157 years, demonstrating an excellent reproducibility in the instrument performance. The time 158 response of the CIMS instrument to ammonia and amines is defined as where the signal stabilizes at its "double e-folded" concentration of 1/e² during the calibration. Average response times for ammonia and amines were smaller than 1 minute. For each 10-minute cycle of background and measurement, the first two minutes of each background/measurement cycle were excluded from the data analysis to allow the instrument to reach a steady concentration.

The uncertainty in the CIMS included error in the permeation sources, which ranged from 2% to 5% depending on the compound. The permeation sources were diluted in two stages using flow controllers that each had uncertainties of 1.5%. Total error in the calibration of the CIMS was 6.7%. Overall uncertainty in the CIMS was 30%, accounting for calibration error, variability of

167 ion signals, and inlet losses.

168 Meteorological data was measured concurrently on the platform by a Vaisala HMP-45c for 169 temperature and relative humidity, and a RM Young 05305 wind speed and direction sensor. Additionally, CO and NO_x (NO+NO₂) were measured with Thermo 48c and Thermo 42c-TL, 170 171 respectively. These measurements were provided by the University of Houston. The uncertainty 172 in trace gas (CO and NO_x) measurements arises from instrumental uncertainty in the Thermo 48c CO analyzer and Thermo 42c-TL NO_x analyzer. Zero correction was performed on this instrument 173 174 daily by switching to a flow of zero air. The typical uncertainty of each of these instruments was 175 5%.

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3. Results and Discussion

Figure 2. Time series of ammonia and C1-C6 amines observed at the urban center in Houston,
Texas, in October 2022.

183 The time series of ammonia and amines during the ambient measurement period is shown in 184 Figure 2. The average ammonia concentration during the measurement campaign was 4 ppby with 185 several short-term spikes above 10 ppbv and one occasion when the concentration exceeded 20 186 ppbv. Concentrations of C1 amine averaged 4 pptv with several spikes up to 15 pptv. Average C2 187 amine concentrations were 6 pptv with frequent but brief periods of concentrations more than 10 188 pptv. Average C3 amine concentrations were 31 pptv with brief increases in concentration above 189 100 ppty. C4 amine was the most abundant amine observed during the measurement period with 190 an average concentration of 79 pptv with spikes in concentration into the hundreds of pptv. 191 Average C5 and C6 amine concentrations were 33 and 12 pptv, respectively. These concentrations 192 in Houston were generally consistent with concentrations measured in other urban sites (Table 1). 193 Previous CIMS ammonia measurements from aircraft flights above Houston observed similar 194 baseline concentrations of ammonia (0.2-3 ppbv) with brief spikes in concentration (up to 80 ppbv) 195 associated with agricultural or industrial activity (Nowak et al., 2010). Additionally, ammonia 196 concentrations of similar magnitude to the high spikes in concentration observed in this study have 197 been reported in Shanghai (Xiao et al., 2015) as well as an urban site in Romania (Petrus et al., 198 2022), with high ammonia concentrations corresponding to high temperatures and high traffic 199 activity. Long-term measurements taken in Nanjing with a cavity ring-down spectrometer also 200 showed an average ammonia concentration of 12 ppbv (Liu et al., 2024). Measurements of amines 201 in Atlanta, Georgia showed <1 to 3 pptv concentrations of C1 and C2 amines, and C3 and C6 202 amines up to 15-25 pptv (Hanson et al., 2011). Yao et al. (Yao et al., 2016) measured amines at 203 the level of pptv or sub-pptv, e.g., C2 amines of 3.9 ± 1.2 pptv, in urban Shanghai during the 204 summer. It is possible that measured concentrations of amines measured here contain some interference from amides formed from oxidation of emitted amines. The CIMS does not have 205 206 sufficient resolving power to separate trimethylamine (m/z 59.11) from acetamide (m/z 59.07), for 207 example. Therefore, these amine concentrations represent an upper limit of amine concentrations 208 (assuming all of the detected signal is due to the presence of amines). However, (Yao et al., 2016) 209 measured amide concentrations in urban Shanghai in the tens to hundreds of pptv, while C1-C2 210 amine concentrations in Shanghai were similar to Houston observations reported here. Considering 211 the consistency between amine measurements at these two urban locations, it is likely that 212 interference from amides in the CIMS was minimal for C1 and C2 amines. The discrepancies 213 between these two urban areas become more pronounced for C3-C6 amines (Table 1), which 214 makes amide interference a possible explanation for elevated concentrations of C3 amines and 215 above.



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Figure 3. Averaged diurnal cycles of (a) ammonia, (b-g) C1-C6 amines, (h) temperature, and RH in Houston, Texas, during the observation period (19 days continuously). Shaded areas indicate 1 standard deviation from the mean values of observation data.

221 Figure 3 shows the averaged diurnal concentrations of ammonia and amines during the 222 observation period. Ammonia and amines had a diurnal cycle with peak concentrations in the 223 afternoon with higher ambient temperatures. Generally, ammonia and amines correlated with one 224 another throughout the measurement campaign, while C1-C3 amines showed the highest 225 correlation with ammonia. Peak concentrations of all compounds corresponded with the high 226 temperature of the day at around 3 pm local time. This was especially pronounced for ammonia, 227 C1 and C3 amines. The relationships between ammonia and amines and temperature are shown in 228 Figure 4. Ammonia had the strongest correlation with temperature, and the relationship fit an 229 exponential parameterization, as the following:

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$$[NH_3] = 2.85 + 9.66 \times 10^{15} e^{-\frac{10619}{T}}$$

Amines generally showed linear relationships with temperature, with C3 and C4 amines displaying 231 the strongest relationships. C3 amines increased by 2.3 pptv per °C ($r^2 = 0.86$) and C4 by 2.9 pptv 232 per °C ($r^2 = 0.65$). C5 and C6 amines were also moderately correlated with temperature, increasing 233 by 1.2 pptv per °C and 0.5 pptv per °C, respectively ($r^2 = 0.60$ for both C5 and C6). On the other 234 hand, the correlation of C1 and C2 amines with temperature were weaker: C1 only increased by 235 0.1 pptv per °C with almost no correlation ($r^2 = 0.22$), and C2 increased by 0.8 pptv per °C ($r^2 =$ 236 0.50). The temperature dependence of ammonia and amines was previously observed in a rural 237 238 forest in Alabama by (You et al., 2014), which attributed this partially to particle-to-gas conversion 239 of ammonia and amine containing particles at elevated temperatures. The temperature dependence 240 could also be due to higher emissions at higher temperatures. The temperature dependence of ammonia and amines has been observed at other urban, suburban and rural locations such as Kent, 241

242 Ohio (You et al., 2014), Atlanta (Hanson et al., 2011), Delaware (Freshour et al., 2014), the

Southern Great Plains (Freshour et al., 2014), and rural central Germany (Kürten et al., 2016).



Figure 4. Temperature dependence of (a) ammonia and (b-g) C1-C6 amines measured in Houston.
Vertical bars indicate 1 standard deviation from the mean values of observation data. Binned
temperatures are shown in colored squares, 1-minute averaged data is shown in gray squares.
Horizontal bars indicate bin width. Black dashed lines indicate exponential fit for ammonia and
linear fits for amines.

252 Anthropogenic pollutants such as CO and NO_x and CO can serve as tracers for industrial and traffic activities. Ammonia and amines in general showed a positive correlation with CO, with the 253 254 exception of C3 amines (Figure 5). As ammonia, amines, and CO can be traced to traffic or 255 industrial emissions, the positive relationship between these compounds implies that these base 256 compounds were emitted from pollutant sources. Unlike with CO, there was a negative correlation 257 with NO_x (Figure S3). This lack of a strong correlation between NO_x and ammonia was previously 258 observed in Nanjing where a strong reduction in NO_x concentration during COVID-19 lockdown 259 periods was not accompanied by an equivalent reduction in ammonia concentrations (Liu et al., 260 2024). This may indicate some unique emission sources for ammonia and amines that do not co-261 emit NO_x.

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Figure 5. Correlation between ammonia (a) and C1-C6 amines (b-g) with the collocated CO concentrations during the measurement campaign. Binned CO concentrations are shown in colored squares, 5-minute averaged data shown in gray squares. Vertical bars indicate 1 standard deviation from the mean values of observation data. Horizontal bars indicate bin widths. Black dashed lines indicate linear fits.

269 Wind speed and direction can help to identify local sources of ammonia and amines near the 270 measurement site. Figures 6 and S4 show the correlation of ammonia and amines with wind speeds 271 and direction throughout the observation period. Consistent between all base compounds is the 272 high concentration coming from the southeast. This is the direction of the interstate highway, 273 industrial areas, and train yards (Figures 1 and S1). Ammonia and most amines also have a 274 pronounced source from the northwest - this is the direction of downtown Houston, where 275 population density is highest. Except for C2 and C4 amines, the observed ammonia and amines in 276 Houston were higher during periods of low wind speeds. The abundant C2 and C4 at high wind 277 speeds may suggest that C2 and C4 amines were transported from more distant sources.

278 Figure S5 shows the average diurnal cycle of ammonia and amines on weekdays as opposed 279 to weekends. Except for C2 and C4 amines, there was a clear decrease in concentrations during 280 weekends during the afternoon peak. Weekends saw much less traffic and activity on the 281 University of Houston campus. During this observation period, ambient temperatures were higher 282 during the weekends, which would increase emissions. Therefore, the differences in weekdays vs. 283 weekends indicate that amines and ammonia were indeed emitted from traffic and industrial 284 activities. Lower average amine concentrations on weekends were also observed during mobile 285 measurements in Yangtze River Delta cities (Chang et al., 2022).

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Figure 6. Wind rose plots of (a) ammonia and (b-g) C1-C6 amines observed in urban Houston.
The color scale indicates concentration, and radial intensity shows wind speed.

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4. Atmospheric Implications

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295 Field observations show that sulfuric acid and amines are responsible for aerosol nucleation 296 (Yao et al., 2016; Yan et al., 2021; Cai et al., 2021; Cai et al., 2023; Jen et al., 2016; Smith et al., 297 2010; Brean et al., 2020), however, currently, global models do not have amine emission 298 inventories. Figure 7 shows the correlation of ammonia with C1-C6 amines measured during this 299 campaign. This figure also includes that data obtained with the same instrument in Kent, Ohio, 300 (You et al., 2014). It is clear from this figure that concentrations of ammonia, C1, C2, C3, C5, and 301 C6 amines were positively correlated with one another throughout the study: r^2 values for the 302 correlation between ammonia and amines were 0.61 for C1, 0.42 for C2, 0.47 for C3, 0.26 for C5

303 and 0.88 for C6. These relationships imply that these compounds are mostly co-emitted from 304 similar sources and undergo similar atmospheric transport. C4 amines showed no correlation with 305 ammonia and lower-mass amines – the r^2 value for C4 vs. NH₃ was 0.048. This indicates a unique 306 source for C4 amines, consistent with both elevated C4 concentrations at high wind speeds and 307 higher weekend C4 concentrations as discussed previously. Correlations of C1-C3 amines 308 concentrations, taken from the linear fits of the plots shown in Figure 7, were approximately equivalent to 1.1×10^{-3} [NH₃], 1.4×10^{-3} [NH₃], and 8.4×10^{-3} [NH₃], respectively. C5 and C6 309 amine concentrations were 1.9×10^{-2} [NH₃] and 3.5×10^{-3} [NH₃], respectively (Table S2). From 310 these results, we propose that global modelers use 0.1 % of the ammonia concentration as a proxy 311 of dimethylamine to simulate urban NPF processes. However, this recommendation comes with 312 313 the caveat that measured C2 amines may include dimethylamine as well as ethylamine due to the 314 inability of mass spectrometry to resolve isomers. Therefore, this correlation represents only the 315 upper bound of dimethylamine concentrations.

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Figure 7. Correlations of C1-C6 amines with ammonia throughout the observation period in Houston (diamonds) and Kent, OH (squares) as reported by (You et al., 2014). Binned concentrations are shown in colored squares, 1-minute averaged data from Houston are shown in

321 gray squares. Vertical bars indicate one standard deviation from the mean values of observation

- 322 data. Horizontal bars indicate bin widths. Black dashed lines indicate linear fits of the combined
- 323 data from Kent and Houston.

From these observations made in very polluted Houston and less polluted Kent, we propose 324 325 that at the polluted sites in the United States, dimethylamine concentrations can be estimated using 326 the proxy, 0.1% ammonia concentrations. The caveat of this proxy is that it is based on only two 327 locations in the United States and did not consider different emission sectors. There has been so 328 far only one attempt to use quantify the aerosol nucleation processes using sulfuric acid and 329 dimethylamine in the global model by Zhao et al. (Zhao et al., 2024) and they concluded that this 330 nucleation process is dominant globally in polluted boundary layer, from China, India, Europe to 331 United States. In this cited study the authors used the proxy of dimethylamine using ammonia 332 concentrations, for example, dimethylamine/ammonia ratio of 0.0070, 0.0018, and 0.0100, for 333 chemical industrial, other industrial, and residential sources; these proxies were derived by Mao 334 et al. (Mao et al., 2018) based on the measurement made in Nanjing (Zheng et al., 2015). Our 335 observations indicate that (Zhao et al., 2024) likely overestimated dimethylamine concentrations 336 for polluted sites in the United States overall, and thus overpredicted nucleation rates as well. Thus, 337 our results can provide more constrained proxy for polluted sites in the United States for future 338 modeling studies.

5. Conclusions

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340 Our observations in urban Houston show that ammonia and amines generally followed a clear 341 diurnal cycle, peaking in the early afternoon when the ambient temperature was highest during the 342 day. We found a correlation between ammonia/amines and ambient temperature. The diurnal 343 cycles and temperature dependence of these compounds are consistent with (You et al., 2014) 344 which showed that the gas-to-particle conversion contributes to the temperature dependence. To 345 verify this process, the chemical composition of particle is needed, but particle measurements were 346 not available during the present study. Additionally, the observed temperature dependence could 347 be due to increased emissions of ammonia and amines from biogenic and anthropogenic sources. 348 On the other hand, photochemical aging that occurs typically during the higher solar flux can also 349 reduce the gas phase amines at the noontime; thus, photochemical aging was unlikely the main 350 driving factor to produce higher concentrations of amines around noon.

351 High concentrations of ammonia and amines were correlated with local air masses from 352 densely populated areas and areas of high traffic, industry, and other human activity. This suggests 353 that most ammonia and amines measured in Houston originated from pollutant sources, consistent 354 with the correlation observed with CO concentrations. There was also a clear increase in ammonia 355 and amines on days with more human activity as shown by the results of concentrations on 356 weekends vs weekdays. We observed a consistent relationship between ammonia and amines 357 during our measurement campaign as well as with observations in less densely populated Kent, 358 Ohio, suggesting that it is reasonable to parameterize amine emission inventories based on existing 359 ammonia inventories to simulate urban NPF processes. However, as the CIMS is incapable of 360 resolving amides or isomers, this parameterization is only capable of representing the upper 361 bounds of amines. Further work involving instrumentation capable of isomer resolution such as 362 tandem MS/MS or chromatographic separation is needed to determine typical isomer ratios of 363 amines for more accurate parameterizations.

- 364 Measuring ammonia and amines in the atmosphere is one of the most challenging areas in the 365 development of atmospheric analytical instruments (Lee, 2022; Lee et al., 2019). The CIMS used in this campaign is currently one of the few instruments in the world that is capable of simultaneous 366 367 measurements of ammonia and amines at atmospherically relevant detection limits and timescales. Very importantly, our CIMS, despite its relatively low mass resolution, has measured ammonia 368 369 and amines at various atmospheric conditions, ranging from the rural forests (You et al., 2014; 370 Kanawade et al., 2014), a relatively less polluted site (Yu and Lee, 2012; You et al., 2014; Erupe 371 et al., 2010), to the extremely polluted urban environment (this study), with consistent instrument 372 sensitivities over the decade; to our knowledge, this is the only instrument that demonstrated such 373 consistency in the performance. Studies have shown that the co-presence of ammonia and amines 374 can enhance sulfuric acid nucleation rates compared to ammonia alone (Yu et al., 2012; Glasoe et 375 al., 2015; Myllys et al., 2019). From this perspective, simultaneous measurements of ammonia and 376 amines will be required for the correct prediction of NPF processes in the atmosphere. 377 Measurements of ammonia and amines with comprehensive calibration as shown in the present 378 study are very even rarer, but such measurements are needed for mitigating urban air quality 379 problems and the health effects of ultrafine particles.
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381 Author Contributions

382 SHL designed the research; LT and SHL performed measurements; JF provided the measurement
 383 platform as well as the trace gas and meteorology data; LT and SHL wrote the manuscript.

384 Competing interests

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

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- **Table 1.** Ammonia and amine measurements with CIMS at various locations reported in the
- 392 literature. DL, detection limit of each instrument.

Location	NH ₃	C1 Amine	C2	C3	C4	C5	C6
	(ppbv)	(pptv)	Amine	Amine	Amine	Amine	Amine
			(pptv)	(pptv)	(pptv)	(pptv)	(pptv)
Rural Alabama	Up to 1-2	< DL	< DL	1 - 10	< DL	< DL	< DL
Forest							
(You et al.,							
2014)*							
Kent, Ohio	Up to 6	1-4	< DL	5 - 10	10 - 50	10 - 100	< DL
(You et al.,							
2014)*							
Kent, Ohio	0.5 ± 0.26	-	8 ± 3	16 ± 7	-	-	-
(Yu and Lee,							
2012)*							
Atlanta,	-	< 1	3	4 – 15	25	-	-
Georgia							
(Hanson et al.,							
2011)†	0.0	-	20	6	1.50	1	
Lewes,	0.8	5	28	6	150	1	2
Delaware							
(Freshour et							
al., 2014)†	0.0	4	1.4	25	150	00	20
Lamont,	0.9	4	14	35	150	98	20
Oklanoma							
(Freshour et $(1, 2014)$							
al., 2014)	1.0	4	42	10	14	20	5
Minnesota	1.0	4	42	19	14	20	5
(Freshour et							
$(110511041) \div$							
Shanghai	_	39 + 12	66 + 12	0.4 + 0.1	36 + 10	0.7 ± 0.3	18 + 08
(Yao et al		5.7 ± 1.2	0.0 ± 1.2	0.1 ± 0.1	5.0 ± 1.0	0.7 ± 0.5	1.0 ± 0.0
(100 et al.) 2016)†							
Naniing	1.7 ± 2.3	7.2 ± 7.4 (C)	+ C2 + C3)	-	-	_
(Zheng et al.,			30				
2015)‡							
Wangdu	-	-	14.6 ±	-	-	-	-
			14.9				
1							

(Wang et al., 2020b)§							
Beijing	2.8 ± 2.0	5.2 ± 4.3	-	-	-		
(Zhu et al.,		(C1 + C2 +					
2022)‡		C3)					
Houston, TX	4 ± 1	4 ± 2	6 ± 2	31 ± 9	79 ± 30	33 ± 12	12 ± 4
(This study)*							

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394 * CIMS with ethanol reagent

395 † Proton-transfer chemical ionization mass spectrometer (PTR-CIMS)

396 ‡ High-resolution time of flight chemical ionization mass spectrometer (HR-TOF CIMS) with

397 ethanol reagent

398 § Vocus proton transfer time-of-flight mass spectrometer (PTR-TOF MS)

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