

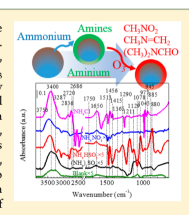


# Ozonolysis of Trimethylamine Exchanged with Typical Ammonium Salts in the Particle Phase

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Supporting Information

ABSTRACT: Alkylamines contribute to both new particle formation and brown carbon. The toxicity of particle-phase amines is of great concern in the atmospheric chemistry community. Degradation of particulate amines may lead to secondary products in the particle phase, which are associated with changes in the adverse health impacts of aerosols. In this study, O<sub>3</sub> oxidation of particulate trimethylamine (TMA) formed via heterogeneous uptake of TMA by (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, NH<sub>4</sub>HSO<sub>4</sub>, NH<sub>4</sub>NO<sub>3</sub> and NH<sub>4</sub>Cl, was investigated with in situ attenuated total reflection Fourier transform infrared spectroscopy (ATR-FTIR) and proton transfer reaction mass spectrometry (PTR-MS). HCOOH, HCHO, CH<sub>3</sub>N=CH<sub>2</sub>, (CH<sub>3</sub>)<sub>2</sub>NCHO, CH<sub>3</sub>NO<sub>2</sub>, CH<sub>3</sub>N(OH)CHO, CH<sub>3</sub>NHOH and H<sub>2</sub>O were identified as products on all the substrates based upon IR (one-dimensional IR and two-dimensional correlation infrared spectroscopy), quantum chemical calculation and PTR-MS results. A reaction mechanism was proposed to explain the observed products. This work demonstrates that oxidation might be a degradation pathway of particulate amines in the atmosphere. This will aid in understanding the fate of particulate amines formed by nucleation and heterogeneous uptake and their potential health impacts during atmospheric aging.



### ■ INTRODUCTION

Amines in the atmosphere are emitted from industrial processes, biomass burning, animal husbandry, automobile emissions, waste treatment facilities and marine emissions. <sup>1</sup> The atmospheric concentration of amines varies with location, and is typically 1–14 nmol Nm<sup>-3</sup>, which is 1–2 orders of magnitude lower than that of ammonia. 1,2 In the atmosphere, amines can interconvert among gas-, liquid- and particle-phases through dissolution,<sup>3</sup> nucleation<sup>4,5</sup> and evaporation.<sup>1</sup> Novakov et al. first reported the presence of amines in particulate matter in 1972, and subsequently more and more field measurements identified amines in atmospheric particles. <sup>2,7-9</sup> It has been suggested that amines may contribute up to 10-20% of the organic content of ambient particles, and frequently over one hundred different amine species containing 1-4 N atoms and ranging in MW from 59-302 Da have been observed in the particle phase.2,10

Amines play an important role in new particle formation (NPF) for the H<sub>2</sub>SO<sub>4</sub>-H<sub>2</sub>O-amines ternary system and their subsequent growth, in particular, in areas of where anthropogenic emission of amines are strong. For example, aliphatic amines contributed 23% to the NPF events at a remote forested site in Hyytiälä, Finland in March, 2006 and

47% to that at an urban site in Tecamac, Mexico in April, 2007. 12 Amines observed in Marshall Field Site, Colorado (2007),13 North Boulder, Colorado (2009), and Atlanta, Georgia (2009)<sup>14</sup> have also been confirmed to play the same role in NPF events. 12 In laboratory studies, NPF was observed when amines were added into a smog chamber or flow tube reactor containing a certain concentration of nitric acid, 15 sulfuric acid, <sup>16</sup> or methanesulfonic acid (MSA). <sup>17,18</sup> Density functional theory (DFT) calculations indicated that amine molecules act as cluster stabilizers for nucleation precursors, <sup>19,20</sup> and therefore promote nuclei formation. <sup>16,21</sup> Heterogeneous uptake of amines by particles also contributes to particle-phase amines. Exchange or substitution reactions between amines and ammonium salts ((NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, NH<sub>4</sub>HSO<sub>4</sub>, NH<sub>4</sub>NO<sub>3</sub> and NH<sub>4</sub>Cl) have been confirmed using both mass spectrometry<sup>22</sup> and Raman spectrometry.<sup>23,24</sup> The uptake coefficient ( $\gamma$ ) of amines on coarse particles of ammonium salts is on the order of  $10^{-3}$ – $10^{-2}$ ,  $2^{3,25}$ – $2^{7}$  while

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it is close to unity with nanoscale ultrafine particles. <sup>22</sup> The  $\gamma$  of amines on organic acids (citric acid and humic acid) via acidbase reaction is  $10^{-5}-10^{-3}$  depending on the type of organic acid. It has been found that stronger acidity of particulate matter,<sup>27</sup> smaller steric effect with fewer substituted groups on the amines are favorable for the heterogeneous uptake of amines.<sup>23</sup> In addition, the uptake coefficient of amines on (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> particles is highly dependent on the metastable (efflorescence or deliquescence) state, which is determined by the relative humidity, of (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>. Heterogeneous uptake of amines on sulfate particles may considerably alter the aerosol properties, such as leading to a transition from the crystalline to an amorphous phase and increased water uptake, considerably enhancing their direct and indirect climate forcing effects.<sup>28</sup>

In the atmosphere, oxidation by OH was considered to be the main degradation pathway of amines. Their atmospheric lifetimes were estimated to be several hours based on the measured second-order rate constants ((2-6)  $\times$  10<sup>-11</sup> cm<sup>-3</sup>. molecule<sup>-1</sup>·s<sup>-1</sup>).<sup>29</sup> Oxidation by O<sub>3</sub> was another degradation path of amines. The lifetime for typical amines reacting with O<sub>3</sub> in the gas phase varies from several hours to around 10 days. Oxidation of amines in the gas phase results in the formation of both volatile and condensed products. 31-33 For example, products including (CH<sub>3</sub>)<sub>2</sub>NCHO, CH<sub>3</sub>N=CH<sub>2</sub>, CH<sub>3</sub>NO<sub>2</sub>, HCHO, HCOOH, and CO<sub>2</sub> have been identified from ozonolysis of gas-phase trimethylamine (TMA).<sup>34</sup> It has also been found that significant nonsalt organics contributed to the secondary organic aerosol (SOA) from ozone oxidation and photooxidation of TMA and TEA.<sup>15</sup> Particulate amines and secondary oxidation products from amines in the particle phase are associated with adverse aerosol health impacts. For example, concentrations of 20 mL·m<sup>-3</sup> trimethylamine were found to found to irritate the mucous membranes and eyes. 35 Alkylamines may also act as endogenous teratogens under certain conditions. 36,37 This implies that heterogeneous oxidation of particle-phase amines should subsequently modify their toxicity. A recent work has found that ozonolysis of particle-phase octadecylamine leads to nitroalkanes, 38 which are carcinogenic.<sup>39</sup> However, the further transformation or fate of particlephase amines formed through heterogeneous uptake of short chain alkylamines is unclear. Therefore, it is important to study the degradation of particle-phase amines formed via nucleation and/or heterogeneous uptake under atmospheric conditions to understand their climatic and health impacts.

In this work, the heterogeneous ozonolysis of particulate TMA exchanged with typical ammonium salts found in atmospheric particles was investigated for the first time using an in situ attenuated total internal reflection infrared (ATR-IR) spectrometer as well as a proton transfer reaction mass spectrometer (PTR-MS). The degradation mechanism of particle-phase TMA by ozone was discussed. The results of this study will aid in understanding the fate of particulate amines formed by nucleation and heterogeneous uptake and their potential impacts during atmospheric aging.

# **■ EXPERIMENTAL SECTION**

Chemicals. (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, NH<sub>4</sub>HSO<sub>4</sub>, NH<sub>4</sub>NO<sub>3</sub> and NH<sub>4</sub>Cl were used to simulate the typical ammonium components in atmospheric particles. (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> (≥99.0%), NH<sub>4</sub>HSO<sub>4</sub>  $(41.5-43.5\% \text{ H}_2\text{SO}_4)$ ,  $\text{NH}_4\text{NO}_3$  ( $\geq 99.0\%$ ), and  $\text{NH}_4\text{Cl}$ (≥99.5%) were used as received and supplied by Sinopharm Chemical Reagent Co. Ltd. TMA (28%) in aqueous solution

was used as received from TCI. Ultrapure water (18.2 M $\Omega$ ) was used to prepare ammonium salt solutions.

Heterogeneous Ozonolysis Experiments. Ozone oxidation of particle-phase TMA was performed in an in situ ATR reactor. The reaction chamber consisted of a ZnSe crystal and a sealed Teflon cover with entry and exit ports. The surface-tovolume ratio was 2 cm<sup>-1</sup> with a volume of 44 cm<sup>3</sup>. More details about the reactor was shown in the Supporting Information (Figure S1). The particles were prepared by depositing small droplets containing ammonium salts onto the ZnSe crystal with an atomizer, followed by purging with 1 L·min<sup>-1</sup> zero air to obtain dry particles. The particle size for the generated ammonium salt particles was 115  $\pm$  76  $\mu$ m measured using a scanning electron microscope (SEM, Hitachi S3000N with an accelerating voltage of 10 kV). Although the particle size is 2-3 orders of magnitude greater than that of atmospheric particles, the reaction path obtained in this work should be credible because particle size has little influence on the reaction mechanism. Amorphous ammonium salt particles (several mg, Table S1) were obtained in this study.

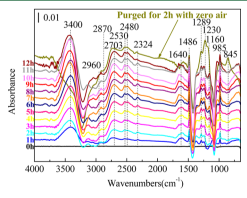
After the particles were dried, which was monitored by observing the IR bands of water, TMA (~385 ppm) vapor was introduced into the reactor by a flow of 200 mL·min<sup>-1</sup> zero air through a water bubbler containing TMA. The RH in the gas flow was 21% measured with a RH sensor (HMP110, HUMICAP). The IR spectra were collected with a time resolution of 1 min using the dried ammonium salts as reference. The exchange reaction was carried out until a dynamic equilibrium was established. Physically adsorbed and gaseous TMA as well as adsorbed water were removed by purging the sample with 1 L·min<sup>-1</sup> zero air until the IR spectrum showed no further changes (around 2 h). Protonated TMA should be the initial form of TMA because the exchange reaction were performed under high TMA concentration for a long time and almost 100% exchange degree was observed in previous work under similar conditions.<sup>2</sup>

100-400 ppbv O<sub>3</sub> was introduced into the reactor to start the heterogeneous ozonolysis of particulate TMA. O<sub>3</sub> was generated by passing zero air through a quartz tube irradiated by a UV lamp (185 nm line dominated, Beijing Lighting Research Institute). The O<sub>3</sub> concentration in the gas flow was measured using an O<sub>3</sub> monitor (model 202, 2B Technologies). Because the mass of ammoniums deposited on the ZnSe crystal (Table S1) varied in each experiment, subsequently, leading to different amount of the particulate TMA exchanged. Therefore, it is reasonable to propose that the amount of exchanged TMA which was determined by the amount of deposited ammoniums on the ZnSe crystal, the equilibrium chemical compositions on different ammonium and the reactivity of particulate TMA toward O<sub>3</sub> should have influence on the concentrations of the surface species. The IR spectra with the best spectral quality were shown and qualitatively analyzed in this work to more confidently identify the surface products and to study the reaction mechanism. To further eliminate the possible influence of the trace TMA in gas phase or adsorbed on the walls of the reactor, the chamber was flushed with O3 balanced in 100 mL· min<sup>-1</sup> zero air for 15 min. Then, the inlet and outlet of the reactor were closed to for the oxidation reaction in a closed system. In situ IR spectra were recorded on a Nicolet 6700 (Thermo Nicolet Instrument Corporation) Fourier transform infrared (FT-IR) spectrometer equipped with an in situ attenuated total reflection chamber and a high sensitivity mercury cadmium telluride (MCT) detector cooled by liquid N<sub>2</sub>. The infrared spectra were collected and analyzed by a computer with OMNIC 6.0 software. All spectra reported here were recorded at a resolution of 4 cm<sup>-1</sup> for 100 scans. A PTR-MS (500 HS, Ionicon Analytik)<sup>40</sup> was used to further identify the products by monitoring the composition of the outflow gas obtained by purging the oxidized sample with 1 L·min<sup>-1</sup> of zero air after ozone oxidation was completed (12 h). Before that, mass spectra were also collected for comparison during purging of the TMA-exchanged ammonium salts with 1 L·min<sup>-1</sup> of zero air before O<sub>3</sub> was introduced into the chamber. PTR-MS measures the molecular ions without substantial fragmentation or adduct formation and provides supplemental information about the products because it is a soft ionization technique. 41,42 Two-dimensional correlation spectroscopy (2DCOS) is an invaluable tool to elucidate the changes that occur at a molecular level when a system is subjected to external perturbation, 43,44 which obtains a higher spectral resolution by resolving the correlations into imaginary and real components and by distributing the peaks over a second dimension. 45,46 Therefore, it is capable of measuring small couplings and can be used as an incisive tool to shed light on the detailed structure and its structural change in time. 47 The 2DCOS was analyzed using SpectraCorr 2DCOS Software (Thermo Fisher Scientific).

**Quantum Chemical Calculations.** Quantum chemical calculations were performed using the Gaussian 09 program which is consistent with the IR assignments for the possible products during ozone oxidation. All geometry optimizations and vibrational frequency calculations were performed at the B3LYP hybrid DFT level of theory with the 6-311++G (d, p) basis set. A scaling factor of 0.975 was used to account for anharmonicity in the calculated vibrational frequencies based upon those products with known IR spectra (NIST).

### RESULTS AND DISCUSSION

Exchange Reaction between TMA and Ammonium. An exchange reaction between TMA and  $(NH_4)_2SO_4$  was conducted first to obtain particulate TMA. As shown in Figure 1, the typical IR bands of TMA, as confirmed with the NIST



**Figure 1.** Time series of IR spectra during reaction between TMA and  $(NH_4)_2SO_4$ .

database, increased as a function of reaction time when the dried  $(NH_4)_2SO_4$  was used as a reference. The peaks at 2960 and 2870 cm<sup>-1</sup> are the asymmetrical stretching vibration of  $CH_3$  ( $\nu_{as}(CH_3)$ ) and the symmetrical stretching vibration of  $CH_3$  ( $\nu_s(CH_3)$ ), respectively, which are accompanied by the deformation vibration of  $CH_3$  ( $\delta(CH_3)$ ) at 1486 and 1446 cm<sup>-1</sup> and the rocking vibration of  $CH_3$  ( $\rho(CH_3)$ ) from 985 to

1290 cm $^{-1.48}$  The band at 845 cm $^{-1}$  is assigned to overtone of asymmetrical deformation vibration of NC $_3$  (2 $\delta_{\rm as}({\rm NC}_3)$ ).  $^{48}$  The peaks in the range of 2300–2700 cm $^{-1}$  are the overtone/combination bands of - CH $_3$  and NC $_3$  groups, which have been well assigned in the literature.  $^{48}$  During the exchange reaction, evidence of the adsorption of water, namely,  $\nu_{\rm s}({\rm OH})$  (3400 cm $^{-1}$ ) and  $\delta({\rm OH})$  (1640 cm $^{-1}$ ) was also observed, because water vapor was present in the gases during bubbling. The absorption bands of water decreased markedly, whereas the peaks attributed to TMA (1289, 1230, 1160, 985, and 845 cm $^{-1}$ ) increased notably after the sample was purged for 2 h with zero air. This can be explained by the symmetry changes from solvated to dry aminium salts.

It should be noted that strong negative peaks at 3250 and 1420 cm<sup>-1</sup> were also observed, as shown in Figure 1, when TMA was introduced into the reactor chamber. They are the typical  $v_s(NH_4^+)$  (3250 cm<sup>-1</sup>) and  $\delta(NH_4^+)$  (1420 cm<sup>-1</sup>) bands of  $(NH_4)_2SO_4$ , 50 respectively. This means that ammonium is replaced by TMA. The reaction is similar to a previous work where NH3 has been identified as a gas-phase product of the reaction between methylamine and ammonium.<sup>51</sup> In Figure 1, the negative peak at 1080 cm<sup>-1</sup> due to  $v_s(SO_4^{2-})^{50}$  can be explained by a part of the sulfate transforming into bisulfate due to TMA or NH3 evaporation from TMAH sulfate or (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> particles, which is similar to the transformation of TEAH sulfate to TEAH bisulfate observed using Raman spectroscopy.<sup>24</sup> On the other hand, a previous work found that the bands at 1484 and 984 cm<sup>-1</sup> are the characteristic IR bands of protonated TMA ((CH<sub>3</sub>)<sub>3</sub>NH<sup>+</sup>), both in solution and on the surface of polyethylene (PE). These frequencies are almost the same as the observed IR bands at 1486 and 985 cm<sup>-1</sup> in this study. This further confirmed that a substitution reaction took place between TMA and (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>. As shown in Figure 1, the intensities for all of these IR bands increased slightly because the exchange rate between TMA and ammoniums gradually slowed down after 3 h of reaction. It should be noted that physisorbed TMA should be negligible compared with the chemically exchanged TMA (TMAH<sup>+</sup>) because the intensity of the IR bands of the surface species, except for water and NH<sub>4</sub><sup>+</sup>, did not decrease after the sample was purged with 1 L·min<sup>-1</sup> of zero air for 2 h as shown in Figure 1.

The exchange reaction was also observed when NH<sub>4</sub>NO<sub>3</sub> and NH<sub>4</sub>Cl were individually exposed to TMA. The IR spectra are shown in Figure 2 after the ammonium salts reacted with TMA had been further purged with dry zero air for 2 h. The peaks attributed to  $-\text{CH}_3$  (1486 cm<sup>-1</sup>,  $\delta(\text{CH}_3)$ ; 1289 cm<sup>-1</sup>, 1230 and 985 cm<sup>-1</sup>,  $\rho(\text{CH}_3)$ ), NC<sub>3</sub> groups (816 cm<sup>-1</sup>,  $2\delta(\text{NC}_3)$ ) and the overtone/combination bands of  $-\text{CH}_3$  and NC<sub>3</sub> groups in the range of 2300–2700 cm<sup>-1</sup> were also observable. At the same time, consumption of NH<sub>4</sub><sup>+</sup> was supported by the negative peaks of  $\nu_s(\text{NH}_4^+)$  (~3200 cm<sup>-1</sup>) and  $\delta(\text{NH}_4^+)$  (1420 cm<sup>-1</sup>) for both NH<sub>4</sub>NO<sub>3</sub> and NH<sub>4</sub>Cl (Figure 2) even though the strong negative peaks at ~3200 cm<sup>-1</sup> make it difficult to discern the  $\nu_{as}(\text{CH}_3)$  and  $\nu_s(\text{CH}_3)$  bands on NH<sub>4</sub>NO<sub>3</sub>.

When  $\mathrm{NH_4HSO_4}$  was exposed to TMA, the negative peaks attributed to  $\mathrm{NH_4^+}$  were very weak, whereas the typical absorbance bands of TMA increased significantly. Additionally, a new peak centered at 1080 cm<sup>-1</sup> due to  $v_\mathrm{s}(\mathrm{SO_4^{2-}})^{50}$  increased, which means TMA had reacted with  $\mathrm{NH_4HSO_4}$  and existed on the crystal. This means an acid—base reaction instead of an exchange/displacement reaction should mainly take place between TMA and  $\mathrm{NH_4HSO_4}$ . At the same time, this

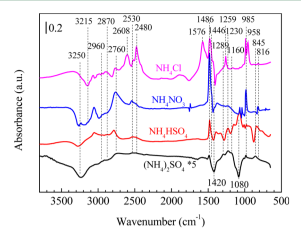


Figure 2. IR spectra after the reacted ammonium with TMA further purged with dry zero air for 2 h.

means that TMA promotes the transformation of HSO<sub>4</sub><sup>-</sup> to SO<sub>4</sub><sup>2-</sup>, which is in agreement with our previous work in which no gas-phase NH3 was detected by a mass spectrometer, and the transformation from  $HSO_4^-$  to  $SO_4^{\ 2-}$  was observed using an in situ Raman spectrometer when NH<sub>4</sub>HSO<sub>4</sub> was exposed to MA.5

It should be noted that protonated TMA ((CH<sub>3</sub>)<sub>3</sub>NH<sup>+</sup>) was formed on all of these ammonium salts as both 1486 and 985 cm<sup>-1</sup> bands were present in the corresponding IR spectra. This implies that aminium salts can be formed on all of these substrates. As shown in Figure 2, frequency shifts were observed for several IR bands of the exchanged TMA on different ammonium substrates. For example, the  $2\delta_{as}$  (NC<sub>3</sub>) band shifted from 845 cm<sup>-148</sup> on (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> and NH<sub>4</sub>HSO<sub>4</sub> to 816 cm<sup>-1</sup> on NH<sub>4</sub>Cl and NH<sub>4</sub>NO<sub>3</sub>. The  $\rho$ (CH<sub>3</sub>) band was observed at 1259 cm<sup>-1</sup> accompanied by new unknown bands at 1576 and 2608 cm<sup>-1</sup> for TMA-exchanged NH<sub>4</sub>Cl. This means that the structure of the formed (HTMA)<sup>+</sup> is sensitive to the surrounding anions.

Ozonolysis of Particulate TMA. To simplify the IR spectral analysis, the IR spectra were collected using the corresponding purged sample (TMA-exchanged ammonium salt) as a reference before O<sub>3</sub> was introduced into the reactor. Figure 3 shows the typical IR spectra during O<sub>3</sub> oxidation of TMA-exchanged (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>. The O<sub>3</sub> concentration was 127 ppbv.

As shown in Figure 3, a number of new bands appeared and increased as a function of reaction time. The band at 3400 cm<sup>-1</sup> with a shoulder at 3287 cm<sup>-1</sup> and the peak 1649 cm<sup>-1</sup> are the characteristic absorption bands of water. 49 Although the TMA exchanged particles were dried with zero air before O<sub>3</sub> oxidation, H2O presenting in the gases (RH 12%) during ozonolysis of particulate TMA should lead to the increase in the IR intensity of water because the aminium salt was highly hygroscopic.<sup>28</sup> On the other hand, OH radical involved in the oxidation of amines.<sup>34</sup> Thus, H<sub>2</sub>O formed through H-abstract reaction, which will be discussed later, might also contribute to the accumulation of water in the particle phase. The bands at 1750 ( $\nu_s(C=O)$ ), 1211 ( $\delta(C-OH)$ ) and 962 cm<sup>-1</sup>  $(w_{oop}(CH), wag of CH out of plane)$  indicate the formation of formic acid. S4 Formaldehyde may also contribute to the absorbance band at 1750 cm<sup>-1</sup> ( $\nu_s(C=O)$ ), which is accompanied by the  $\nu_{\rm as}({\rm CH_2})$  band at 2836 cm<sup>-1</sup> and  $\nu_{\rm s}({\rm CH})$  at 2720 cm<sup>-155</sup> in Figure 3. The bands at 1660–

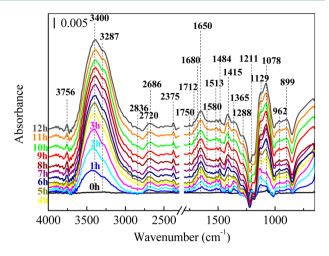
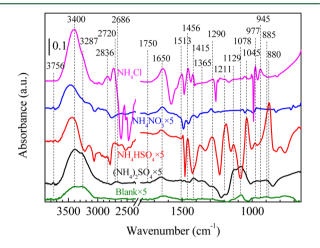


Figure 3. In situ FT-IR spectra during ozone oxidation of TMA exchanged (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>. The initial O<sub>3</sub> concentration was 127 ppb.

1680 cm<sup>-1</sup> can be assigned to the  $\nu_s(C=O)$  of N,Ndimethylformamide (DMF), which is supported by the observation of  $\nu_s(CN)$  at 1288 cm<sup>-1</sup>, the  $r(CH_3)$  (rocking vibration of CH<sub>3</sub>) at 1078 cm<sup>-1</sup> and the mixing of  $\delta$ (CH<sub>3</sub>) and  $v_s(CN)$  at 1513 cm<sup>-1</sup> in DMF.<sup>56</sup> Monomethyleneamine (CH<sub>3</sub>N=CH<sub>2</sub>) was proposed as a possible product because the bands for  $\beta$ (C–H) (bending vibration in plane, 1415 cm<sup>-1</sup>) of the =CH<sub>2</sub> group and  $v_s(C-N)$  (899 cm<sup>-1</sup>) were observable, 57 although it is hard to discern the  $v_s(C=N)$  at around 1660 cm<sup>-157</sup> from bands of other species, including  $\delta(OH)$  of water and  $\nu_s(C=O)$  of formamide. As shown in Figure 3, nitromethane (CH<sub>3</sub>NO<sub>2</sub>) can be confirmed, with  $v_{as}(NO_2)$  and  $v_s(NO_2)$  at 1580 and 1365 cm<sup>-1,58</sup> respectively. The  $v_s(CN)$  band of CH<sub>3</sub>NO<sub>2</sub> at 885 cm<sup>-158</sup> was overlapped with  $v_s(C-N)$  of  $CH_3N=CH_2$  at 899 cm<sup>-1.57</sup> The peaks at 1129 and 1484 cm<sup>-1</sup> in Figure 3 should be related to the  $r(CH_3)$  and  $\delta(CH_3)$  bands in nitromethane, <sup>58</sup> monomethyleneamine<sup>57</sup> and DMF.<sup>56</sup> The band at 2375 cm<sup>-1</sup> is assigned to CO<sub>2</sub>. <sup>59</sup> The peak at 3756 cm<sup>-1</sup> implies that a compound containing OH was formed. In Figure 3, several negative peaks at 1230 cm<sup>-1</sup> ( $\rho$ (CH<sub>3</sub>)) and 845 cm<sup>-1</sup> ( $2\delta_{as}$ (NC<sub>3</sub>))<sup>48</sup> were observed. This is consistent with the consumption of particle-

As discussed above, several possible products including H<sub>2</sub>O, HCHO, HCOOH, CH<sub>3</sub>N=CH<sub>2</sub>, HCON(CH<sub>3</sub>)<sub>2</sub> and CH<sub>3</sub>NO<sub>2</sub> were detected, while some of their IR bands overlapped each other. Therefore, 2DCOS analyses were further performed to confirm the assignments for these species (Figure S3). Besides the bands observed in Figure 3, several new peaks, such as 3340, 3210, 2894, 2854, 2760, 2580, 1660, 1560, 1424, 1140, 921, 877, 820 cm<sup>-1</sup>, have been identified in Figure S3 due to the higher spectral resolution of 2DCOS. As discussed in the SI, (CH<sub>3</sub>)<sub>2</sub>NCHO, CH<sub>3</sub>NO<sub>2</sub>, CH<sub>3</sub>N=CH<sub>2</sub>, HCHO and HCOOH were confirmed by 2DCOS although some peaks were overlapped. On the other hand, the IR frequencies of these products were consistent with DFT calculations and summarized in Table S2. The results further rationalized the assignment for the peaks identified in the IR spectra (Figures 3 and S3). In particular, the band at 3756 cm<sup>-1</sup> was assigned to the  $\nu_s(OH)$  of  $CH_3N(OH)CHO$  or CH<sub>3</sub>NHOH, which is close to the calculated value at 3677 (3771 cm<sup>-1</sup> with unscaled frequency) for CH<sub>3</sub>N(OH)-CHO and 3730 cm<sup>-1</sup> for CH<sub>3</sub>NHOH, and the other IR bands of CH<sub>3</sub>N(OH)CHO and CH<sub>3</sub>NHOH were assigned and summarized in Table S2. This means that CH<sub>3</sub>N(OH)CHO or CH<sub>3</sub>NHOH may be formed during ozonolysis of particulate TMA

Figure 4 summarizes the IR spectra of the TMA-exchanged NH<sub>4</sub>Cl, NH<sub>4</sub>NO<sub>3</sub>, and NH<sub>4</sub>HSO<sub>4</sub> followed by O<sub>3</sub> oxidation for



**Figure 4.** FT-IR spectra of TMA-exchanged  $NH_4Cl$ ,  $NH_4NO_3$ ,  $NH_4HSO_4$  and  $(NH_4)_2SO_4$  after being exposed to 267, 400, 203, 127 ppbv of ozone, respectively, for 12 h. The blank experiment was performed by exposing TMA absorbed on the cleaned ZnSe crystal (without ammonium) to 127 ppbv of ozone.

12 h. For comparison, the IR spectra of TMA-exchanged (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> and the blank experiment have also been included in the same figure. As shown in Figure 4, most of the species observed on TMA-exchanged (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> were present on the other ammonium salts, and the assignments of these species were summarized in Table S2 based on the literature and DFT calculations. Compared with ozone oxidation of the TMAexchanged ammonium salts, oxidation of TMA absorbed on the blank ZnSe crystal (the cleaned ZnSe crystal without ammonium) showed little contribution to the IR signal. On the other hand, the IR signal during ozone oxidation of TMAexchanged NH<sub>4</sub>Cl or NH<sub>4</sub>HSO<sub>4</sub> was much stronger than that of TMA-exchanged NH<sub>4</sub>NO<sub>3</sub> or (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>. This might result from different TMA contents obtained by the exchange reactions due to different reactivities between TMA and the ammonium salts as shown in Figure 2. The properties of anion ions may also have influence on the IR signals. For example, Cl does not have IR absorption in the range of 650-1800 cm<sup>-1</sup>, which leading to a good spectral quality for NH<sub>4</sub>Cl exchanged TMA. In addition, the difference in the IR signals among different substrates implies the different reactivity of particulate TMA toward O<sub>3</sub> due to the role of anion ions. It should be noted that an ATR accessory was used for IR spectra collection. The probe depth of the evanescent wave was less than 1.5  $\mu$ m in the spectral region.<sup>60</sup> This means that particlephase but not gas-phase products mainly contribute to the measured ATR-IR signals. In addition, oxidation of TMA exchanged (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> was further carried out in flow system, from which the species in the gas phase were flushed continuously. As show in Figure S4, the surface products were similar to that observed in the closed system. Therefore, it can be reasonably concluded that the observed products from ozonolysis of TMA were formed via heterogeneous oxidation of particle-phase TMA.

Figure 5 shows the mass spectra before and after  $O_3$  oxidation of TMA-exchanged  $(NH_4)_2SO_4$ . After  $O_3$  oxidation,

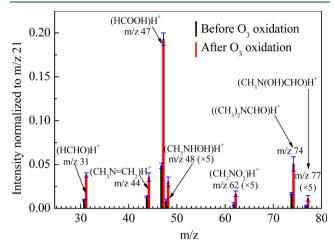


Figure 5. Intensity of selected mass channels before and after  $O_3$  oxidation of TMA-exchanged  $(NH_4)_2SO_4$  for 12 h with 127 ppbv of  $O_3$ .

the mass channels at m/z 31, 44, 47, 48, 62, 74, and 77 corresponding to  $(HCHO)H^+$ ,  $(CH_3N=CH_2)H^+$ ,  $(HCOOH)H^+$ ,  $(CH_3NHOH)H^+$ ,  $(CH_3NO_2)H^+$ ,  $((CH_3)_2NCHO)H^+$  and  $(CH_3N(OH)CHO)H^+$ , respectively, increased significantly. This was consistent with the result of in situ IR spectra and further confirmed HCHO,  $CH_3N=CH_2$ , HCOOH,  $CH_3NO_2$ ,  $(CH_3)_2NCHO$ ,  $CH_3NHOH$  and  $CH_3NI$  to those observed in the gas-phase oxidation of TMA by  $O_3$ . One important observation is that most of these compounds (except for  $CH_3NHOH$ ) could be still detected even when the oxidized sample had been purged with 1 L min<sup>-1</sup> of zero air for 2 h (Figure S5). This indicated that a portion of these compounds were in the particle phase, and confirmed that these products were generated in the particle phase.

Oxidation Paths of Particulate TMA by Ozone. Based on the products observed in this work, the oxidation paths of particulate TMA by O<sub>3</sub> can be summarized in Figure 6. As shown in Figure S6, the symmetry of the highest occupied molecular orbital (HOMO) of TMA matches with that of the lowest unoccupied molecular orbital (LUMO) of O3, while it does not between the HOMO of TMAH+ and the LUMO of O<sub>3</sub>. This means molecular TMA rather than TMAH<sup>+</sup> can be oxidized by O<sub>3</sub>. Although it has been found that aminium sulfate is comparable or more thermostable than ammonium sulfate, 28 an equilibrium between molecular TMA and protonated TMA should be possible on the surface. Then, the N atom in TMA is attacked first by ozone followed by loss of oxygen, forming an energy-rich amine oxide similar to the mechanism described by Slagle et al.<sup>61</sup> for the reaction between O(3P) atoms and alkylamines. Nitromethane, which was identified as a product by both IR and PTR-MS in this study, is directly formed from the attack of amine oxide by a second ozone. Intramolecular H atom migration from the methyl group to the oxygen atom may occur and form an unstable hydroxylamine methyl radical. Thus, N-hydroxylmethylformamide is formed in the presence of O2 by loss of a HO2 radical from peroxide. (CH<sub>3</sub>)<sub>2</sub>NCH<sub>2</sub> radicals can be generated by H abstraction of TMA by OH radical or by loss of OH radical from the hydroxylamine. The corresponding peroxide radical

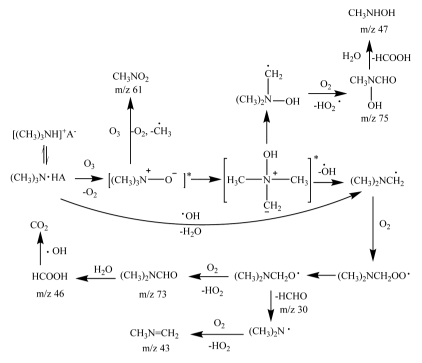


Figure 6. Possible oxidation paths of particulate TMA by ozone. "A" means anion.

((CH<sub>3</sub>)<sub>2</sub>NCH<sub>2</sub>O<sub>2</sub>) and RO radical ((CH<sub>3</sub>)<sub>2</sub>NCH<sub>2</sub>O) are formed in the presence of O<sub>2</sub>. Finally, the CH<sub>3</sub>N=CH<sub>2</sub> and (CH<sub>3</sub>)<sub>2</sub>NCHO can be formed, accompanied by the loss of HCHO and/or HO<sub>2</sub> radical. HCOOH is produced from hydrolysis of (CH<sub>3</sub>)<sub>2</sub>NCHO or CH<sub>3</sub>NCHO and can be finally oxidized to CO<sub>2</sub>. The oxidation mechanism of TMA in the particle phase (Figure 6) is similar to that in the gas phase.<sup>34</sup> However, besides oxidation reactions, hydrolysis in the particle phase has also been observed in this study. For example, a new product (CH<sub>3</sub>NHOH) from hydrolysis of CH<sub>3</sub>NCHO has been identified in the particle phase by both the IR and the PTR-MS.

# ■ ATMOSPHERIC IMPLICATIONS

As pointed out by Chapleski et al.<sup>62</sup> in a recent review paper, adsorption or uptake of volatile organic compounds onto liquid or solid surfaces is one of formation or growth mechanisms of organic particles in the atmosphere. Although the typical concentration of amines is from several pptv to tens of pptv, 63,64 it has been well recognized that amines contribute to both new particle formation <sup>12,16</sup> and brown carbon. <sup>65</sup> For example, nucleation rates with 5 pptv of DMA were enhanced more than 1000-fold compared with 250 pptv ammonia at ambient concentration of sulfuric acid. 16 On the other hand, although it has not been evaluated by modelers, heterogeneous uptake onto fine particles might also contribute to reduced nitrogen in particle phase due to the relative large uptake coefficients as reported previously. <sup>22,53,66,67</sup> Whatever the formation path is, once formed, organic nitrogen containing compounds including particulate amines nearly immediately begin the heterogeneous oxidation by oxidants in the lower atmosphere.62

The lifetimes of alkylamines exposed to ozone in the gas phase usually vary from a few hours to several days. <sup>68</sup> Although oxidized by OH is the dominant degradation path for amines, oxidation by  $O_3$  may also be a degradation path for amines in the atmosphere. For example, the lifetimes of TEA, DEA, TMA,

and DMA are in the range of 4.8-278 h under the typical tropospheric ozone concentration  $(7 \times 10^{11} \text{ molecules cm}^{-3}).^{30}$ Products including HCHO, HCOOH, CH<sub>3</sub>NO<sub>2</sub>, CH<sub>3</sub>N=CH<sub>2</sub>, and (CH<sub>3</sub>)<sub>2</sub>NCHO have been identified from ozonolysis of gasphase amines.<sup>34</sup> As observed in this study, O<sub>3</sub> oxidation of particle-phase amines, which formed through heterogeneous uptake by ammonium, can also take place when exposed to  $O_3$ . HCHO, HCOOH, CH3NO2, CH3N=CH2 and (CH<sub>3</sub>)<sub>2</sub>NCHO were confirmed as the major products by both FTIR and PTR-MS. Heterogeneous oxidation usually leads to increases in oxygen-containing species, <sup>69,70</sup> O/C ratio <sup>71</sup> and structure modifications of aerosol particles, <sup>72</sup> subsequently, may alter its hygroscopicity<sup>73</sup> and optical properties. Although most of these products are volatile from ozone oxidation of particulate TMA, a portion of these compounds are in the particle phase. Thus, heterogeneous oxidation might have influence on the hygroscopic and optical properties of particulate amines, while further studies are required in the future. Because acid-base reactions between amines and inorganic or organic acids contribute to both the nucleation <sup>12,16</sup> and subsequent growth of submicron particles, degradation of particle-phase amines might have a complex influence on air quality through changing their particulate lifetimes. It has been found that heterogeneous oxidation increased the redox activity<sup>75–78</sup> and the cytotoxicity<sup>76,79,80</sup> of black carbon. At the present time, it is poorly understood about the toxicological evolution during atmospheric aging of organic aerosol. However, HCHO, CH<sub>3</sub>NO<sub>2</sub>, and (CH<sub>3</sub>)<sub>2</sub>NCHO from ozone oxidation of particulate TMA are well-recognized toxicants or carcinogens. Therefore, more attention should be paid to the health impacts of the secondary oxidation products from amines in the particle phase.

It should be pointed out that ozone concentration in this work were more representative of highly polluted locations and the particle size was 2–3 orders of magnitude greater than atmospheric particles. Because both the ozone concentration and the particle size should mainly have influence on the

reaction kinetics but not the reaction mechanism, the oxidation path observed in this work should have atmospheric relevance for understanding the fate of amines in the atmosphere. On the other hand, uptake coefficient of ozone on the particulate amines or degradation kinetics of particulate amines are very important to evaluate the atmospheric importance for the possible degradation paths proposed in this work. This requires further investigation of the degradation kinetics of particlephase amines in the future.

### ASSOCIATED CONTENT

# Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.est.6b04375.

Interpretation of 2D correlation spectroscopy (2DCOS) data, reaction conditions in displacement and ozonolysis reactions, assignments of IR spectra, experimental setup, SEM micrographs of ammonium sulfate and aminium sulfate particles, 2DCOS, comparison between the in situ FT-IR spectra during ozone oxidation of TMA exchanged (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> in the flow system and the closed system and concentration changes of the products detected by the PTR-MS (PDF)

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#### **Notes**

The authors declare no competing financial interest.

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