Role of ammonia in forming secondary aerosols from gasoline vehicle exhaust

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Ammonia (NH3) plays vital roles in new particle formation and atmospheric chemistry. Although previous studies have revealed that it also influences the formation of secondary organic aerosols (SOA) from ozonolysis of biogenic and anthropogenic volatile organic compounds (VOCs), the influence of NH3 on particle formation from complex mixtures such as vehicle exhausts is still poorly understood. Here we directly introduced gasoline vehicle exhausts (GVE) into a smog chamber with NH3 absorbed by denuders to examine the role of NH3 in particle formation from GVE. We found that removing NH3 from GVE would greatly suppress the formation and growth of particles. Adding NH3 into the reactor after 3 h photo-oxidation of GVE, the particle number concentration and mass concentrations jumped explosively to much higher levels, indicating that the numbers and mass of particles might be enhanced when aged vehicle exhausts are transported to rural areas and mixed with NH3-rich plumes. We also found that the presence of NH3 had no significant influence on SOA formation from GVE. Very similar oxygen to carbon (O:C) and hydrogen to carbon (H:C) ratios resolved by aerosol mass spectrometer with and without NH3 indicated that the presence of NH3 also had no impact on the average carbon oxidation state of SOA from GVE.

ammonia, vehicle exhaust, secondary organic aerosols (SOA), smog chamber, fine particles, nitrogen oxides, volatile organic compounds

1 Introduction

Ammonia (NH3), a ubiquitous gas, is the third abundant nitrogen-containing gas in the atmosphere [1]. Atmospheric NH3 is mainly emitted from animal waste, biological processes in soils and ammonia-based chemical fertilizers, followed by biomass burning, and sewage treatment plants [2,3]. The gas-phase reactions between NH3 and inorganic acids such as nitric and sulfuric acid can form ammonium nitrate and sulfate [4], which are important constituents of airborne fine particles or PM2.5 [5]. In China’s megacities, which are frequently hard-hit by severe haze due to PM2.5 pollution in the recent decade [6,7], ammonium alone has contributed more than 5% of the PM2.5 mass [8–12].

NH3 can enhance the nucleation of sulfuric acid particles and thus influence new particle formation events [13,14]. In addition, recent smog chamber simulations indicate that NH3 can enhance the secondary organic aerosol (SOA) formation from ozonolysis of biogenic precursors by reacting with organic acids [15,16], but reduce SOA formation from the styrene-ozone system [17]. However, the potential
influence of NH₃ on SOA formation is still poorly understood, especially for complex chemical mixtures such as vehicle exhausts brimming with thousands of gaseous and particle-phase components [18]. SOA, which is ubiquitous in various atmospheric environments, respectively accounts for approximately 66% and 90% of the total OA in urban as well as urban downwind and rural sites in Northern Hemisphere mid-latitudes [19]. Current chemical transport models usually underestimate the level of OA mainly due to the unclear formation processes and sources of SOA [20]. A detailed investigation of the potential influence of NH₃ on SOA formation from vehicle exhaust is therefore necessary to better understand the formation pathways and sources of SOA.

In urban areas with high population densities, NH₃ is centralized and significantly emitted from sewage. Recent tunnel studies indicated that emissions from light-duty gasoline vehicles (LDGVs) are also an important source of NH₃ [21–25]. Nordin et al. [26] found that SOA formed from the exhaust of idling gasoline vehicles could be 1–2 orders of magnitude higher than POA. Gordon et al. [18] and Platt et al. [27] also observed an enhancement of SOA/POA from 1 to 15 during the aging of emissions from LDGVs. Significant nitrate and ammonium were observed to be formed synchronously with SOA during the photo-oxidation of LDGV exhaust [18, 26]. However, the formation of secondary aerosols including SOA, nitrate, and ammonium from vehicle exhaust still remain uncertain when NH₃ is removed. Therefore, studying the role of NH₃ in forming secondary aerosols from vehicle exhaust might give a better understanding of the tropospheric chemistry in urban areas, as well as the aging of vehicle exhausts transported into NH₃-rich plumes emitted from livestock waste and fertilizers in rural areas.

Here we introduce exhaust from LDGVs directly into a smog chamber with a 30 m³ Teflon reactor with NH₃ absorbed by denuders, to study the role of NH₃ in particle formation under photo-oxidation. The purpose is to understand the effects of NH₃ on secondary aerosol formation from LDGV exhaust, and to obtain valuable information for the future emission control of NH₃ and vehicle exhaust.

2 Methods

2.1 Vehicles and fuel

Two port fuel injected Euro 1 and Euro 4 LDGVs with model years of 2002 and 2011 were used in this study. More details of each vehicle are listed in Table 1. They were fueled with Grade 93# gasoline, which complies with the Euro III gasoline fuel standard. The gasoline compositions were described in our previous study [28].

| ID | Vehicle Model | Vehicle class | Mileage (km) | Displacement (cm³) | Power (kW) | Weight (kg) |
|----|---------------|--------------|--------------|--------------------|------------|-------------|
| I  | Euro4 Golf    | 2011         | 25000        | 1598               | 77         | 1295        |
| II | Euro1 Accord  | 2002         | 257984       | 2298               | 110        | 1423        |

2.2 Smog chamber experiments

Six photochemical experiments were conducted in the smog chamber in Guangzhou Institute of Geochemistry, Chinese Academy of Sciences (GIG-CAS). The GIG-CAS smog chamber has a 30 m³ fluorinated ethylene propylene (FEP) reactor housed in a temperature-controlled room. Details of setup and facilities about the chamber are described elsewhere [29]. Prior to each experiment, the chamber was evacuated and filled with purified dry air for at least 48 h until no residual hydrocarbons, O₃, NOₓ, or particles were detected. In the present study, temperature and RH inside the reactor were respectively set to 25 °C and 50%.

Before the introduction of vehicle exhaust, the test vehicle was at idling for at least 30 min to warm up the three-way catalysts. Next, vehicle exhaust was directly introduced into the reactor by two oil-free pumps (Gast Manufacturing, Inc., USA). For comparison, ammonia in LDGV exhaust from experiments I-2, II-2, and II-3 (Table 2) was absorbed by denuders coated with a solution of phosphorous acid in 1:9 water and ethanol [30].

After the introduction of exhaust, additional NO was added to adjust the VOC/NOₓ ratios to 2.0–5.0 (Table 2), which was within the range of 0.5–10 in gasoline vehicle exhaust and downwind urban areas [31]. There were experiment-to-experiment differences in initial precursor concentrations due to the technical challenges of introducing the same amounts of exhaust into the reactor. We kept similar non methane hydrocarbons (NMHCs) concentrations for comparison experiments such as I-1 and I-2. After more than 30 min of primary characterization, the exhaust was continuously exposed to black light for 5 h. After the black lamps were switched off, the formed particles were characterized for another 2–3 h to correct the particle wall loss.

A series of instruments was used to characterize the gas- and particle-phase compounds in the reactor. Gas-phase ozone (O₃) and NOₓ were measured online with dedicated monitors (EC9810 and 9841T, Ecotech, Australia). Methane and CO concentrations were determined using a gas chromatography (Agilent 6980GC, USA) with a flame ionization detector and a packed column (5A Molecular Sieve 60/80 mesh, 3 mx1/8 inch) [32], and CO₂ was analyzed with a gas chromatography (HP 4890D, USA) [33]. Gas-phase organic species such as C₆–C₁₀ single-ring aromatic hydrocarbons were measured with a Mode 7100 preconcentrator (Entech Instruments Inc., USA) coupled with an
Table 2  Summary of the initial conditions and results of the light-duty gasoline vehicle photo-oxidation experiments

| Exp # | OH (×10^6 molecules/cm^3) | T (°C) | RH (%) | VOC/NOx | NMHCs (ppbv) | NO (ppbv) | NO2 (ppbv) | SO2 (ppbv) |
|-------|-----------------|-------|--------|---------|-------------|----------|----------|----------|
| I-1   | 1.20            | 24.2  | 52.5   | 2.0     | 1885        | 794.1    | 161.9    | 7.2       |
| I-2   | 1.18            | 24.8  | 54.3   | 3.6     | 1821        | 496.9    | 2.8      | 6.8       |
| II-1  | 0.80            | 24.6  | 55.6   | 3.0     | 5693        | 1833.3   | 45.9     | 14.7      |
| II-2  | 0.79            | 24.4  | 54.6   | 3.9     | 4605        | 1117.8   | 63.3     | 15.0      |
| II-3  | 0.84            | 25.1  | 51.9   | 4.4     | 1352        | 298.1    | 9.7      | 12.1      |
| II-4  | 1.59            | 25.3  | 51.1   | 4.3     | 1303        | 296.2    | 4.4      | 13.3      |

3 Results and discussion

3.1 Effects of NH3 on particle formation and growth

To investigate the role of NH3 in the formation of secondary aerosols from GVE, we conducted two experiments for each vehicle, with and without NH3 absorbed by denuders. The detailed experimental conditions are listed in Table 2. The average concentration of OH radicals during the 5 h of irradiation was similar for each vehicle. Figure 1 shows the temporal evolution of gas-phase species, including NO, NO2, and O3, and the particle number concentrations during the photochemical aging experiments of LDGV exhaust with vehicle II. Between ~2.1 and ~1.5 h, the vehicle exhaust was introduced into the reactor, leading to a slight increase of the particle numbers before irradiation. In experiment II-2, NH3 in the exhaust was absorbed by denuders. Obviously, in both experiments, NO was converted to NO2 after the black lights were turned on. As presented in previous studies, a delay between aerosol formation and the start of irradiation is frequently observed in classic photo-oxidation experiments; specifically, aerosols may not form until the concentration of NO is around zero [45–47]. In the presence of NH3, the number concentration of particles increased from ~1000 to ~4000 cm^-3 when the concentration of NO was still hundreds of ppb (Figure 1(a)); without NH3, no new particles were formed even when the concentration of NO was approximately zero (Figure 1(b)). Ng et al. [48] found that products formed under high NO conditions via RO2+NO reactions partition much less readily into the aerosol phase than products formed via RO2+HO2 reactions under low NO conditions. Wildt et al. [47] observed that high concentration of NO was responsible for the suppression of new particle formation. Accordingly, higher concentration of NO in experiment II-1 than that in experiment II-2 should have suppressed new particle formation. However, new particle formation was observed only in experiment II-1, indicating that the presence of ammonia in LDGV exhaust might play an important role. Kirkby et al. [14] found that the presence of 100 ppt NH3 could increase the nucleation rate of sulfuric acid particles by a factor of 100–1000. According to a recent study by Kulmala et al. [49], NH3 absorbs by denuders coated with a solution of phosphorous acid in 1:9 water and ethanol [30]. Around 100 ppb of NH3 was injected after 4 h (experiment I-2) and 3 h of irradiation (experiment II-2); c) around 100 ppb of NH3 was injected before UV irradiation.

O:C, and N:C [41,42]. The contribution of gas-phase CO2 to determine the average element ratios of organics, like H:C, ammonium, and organics). We used the toolkit Pika 1.1H to the time series of various mass components (sulfate, nitrate, two minutes. The toolkit Squirrel 1.51H was used to obtain the mass concentration [38]. A high-resolution time-of-flight mass spectrometer (HR-TOF-MS, Aerodyne Research Incorporated, USA) was used to measure the particle chemical compositions and nonrefractory PM mass [39,40]. The instrument was operated in the high sensitivity mode and high resolution W-mode alternatively every two minutes. The toolkit Squirrel 1.51H was used to obtain the time series of various mass components (sulfate, nitrate, ammonium, and organics). We used the toolkit Pika 1.1H to determine the average element ratios of organics, like H:C, O:C, and N:C [41,42]. The contribution of gas-phase CO2 to the m/z 44 signal was corrected using the measured CO2 concentrations. The HR-TOF-MS was calibrated using 300 nm monodisperse ammonium nitrate particles. 2.3 Wall loss corrections

The loss of particles and organic vapors to the reactor walls must be accounted for to accurately quantify the SOA formation. Briefly, the loss of particles onto the walls has been well constrained and is treated as a first-order process [43]. The wall-loss rate constant was determined separately for each experiment by fitting the SMPS and AMS data when no new particles were formed. By applying this rate to the entire experiment, we used the same method as Pathak et al. [44] to correct the wall loss of the particles. In addition, the aerosol mass measured by HR-TOF-AMS was corrected with SMPS data with the same method used by Gordon et al. [18].
can stabilize initial sulfuric acid clusters with diameters of about 1.5 nm. These stable clusters are critical for new particle formation. As shown in Table 2, the initial concentration of SO$_2$ for vehicle II was around 15 ppb. NH$_3$ in GVE might stabilize sulfuric acid formed via the oxidation of SO$_2$ and thus facilitate new particle formation. Consequently, removing NH$_3$ in GVE suppressed new particle formation.

The geometric average diameter increased from 45 to 200 nm and from 22 to 220 nm in around 1 h for experiments I-1 and II-1, with respective particle growth rates of 139 and 181 nm h$^{-1}$; these rates were one order of magnitude higher than those observed in urban environments [50–52]. Some studies have shown that particles larger than 80 nm can be activated as cloud condensation nuclei (CCN) at moderate atmospheric supersaturation and that they possibly directly and indirectly impact climate [53,54]. Therefore, large amounts of LDGV exhaust emissions in urban areas via photochemical aging can influence air quality and even the climate.

As shown in Figure 2, a significant amount of ammonium nitrate was formed through the reaction of NH$_3$ with HNO$_3$ oxidized from NO in experiments I-1 and II-1. The maximum ammonium nitrate production was almost 7 times higher than SOA production, which indicated the significant contribution of LDGV exhaust to secondary inorganic aerosols. Nordin et al. [26] and Gordon et al. [18] also observed significant formation of ammonium nitrate during the aging of gasoline vehicle exhaust in smog chambers; in addition, in-use vehicle emissions mainly from LDGVs in a highway tunnel in Pittsburgh were also found to produce substantial ammonium nitrate [55]. Ammonium has frequently been observed to contribute to particle growth in urban environments [56–58]. Zhu et al. [52] hypothesized that ammonium nitrate plays a role in the particle growth in Qingdao and Toronto during spring. In Figure 3, we plot the concentration of ammonium nitrate without wall loss correction against the geometric average diameter derived from the SMPS data in experiments I-1 and II-1. In both experiments, the geometric average diameter showed significant linear correlations ($R^2$>0.84, $P$<0.001) with the concentration of ammonium nitrate, which demonstrated that ammonium nitrate plays an important role in the rapid growth of particles formed from the aging of LDGV exhaust.

A recent study indicated that the NH$_3$ emission factor of LDGVs in China was much higher than those in the United States [23]. However, today’s emission standards for LDGV
exhaust concern only the primary emissions of hydrocarbon, NOx, and CO. As the presence of NH3 strongly enhances the formation and growth of particles from GVE, removing NH3 from GVE might reduce secondary aerosols formation and benefit air pollution control in urban areas.

3.2 Effects of NH3 on SOA formation

To further investigate the effects of NH3 on SOA formation from LDGV exhaust, we conducted two more experiments, II-3 and II-4, with similar initial concentrations of VOCs and NOx (Table 2). NH3 in LDGV exhaust in experiment II-3 was absorbed by denuders, whereas in experiment II-4 around 100 ppb of NH3 was injected before the UV lights were turned on. Variations of the geometric average diameters and concentrations of organic aerosols during the irradiation of LDGV exhaust with and without NH3 are shown in Figure 4. With and without adding NH3, the geometric average diameter respectively increased from around 20 nm to a maximum of 130 and 115 nm (Figure 4(a)). This result further demonstrates that the presence of NH3 can enhance the growth of particles formed from the aging of LDGV exhaust. As shown in Figure 4(b), in the presence of NH3, SOA was formed more slowly and its concentration was relatively lower. Because the average concentration of OH radicals during the whole experiment in the presence of NH3 was higher, according to the gas-particle partitioning model [59], SOA formation under a higher OH concentration and similar concentrations of SOA precursors should be higher. Na et al. [15] observed that NH3 could neutralize organic acids to enhance SOA formation from α-pinene ozonolysis. A recent study [60] demonstrated that NH3 could react with carbonyl groups in SOA molecules to form nitrate-containing compounds. However, in our study the relatively lower concentration of SOA in the presence of NH3 suggested that NH3 might have no significant influences on SOA formation from LDGV exhausts.

The average oxygen to carbon (O:C) and hydrogen to carbon (H:C) molar ratios of SOA in experiments II-3 and II-4 are presented in Table 3. The O:C and H:C ratios in the
presence of NH$_3$ were 0.683±0.042 and 1.352±0.042, respectively, almost same as the values without NH$_3$, indicating that the presence of NH$_3$ had no influence on the O:C and H:C ratios of SOA from LDGV exhaust. An average carbon oxidation state (OS$_c$) can be estimated from the O:C and H:C data [61]. The average OS$_c$ of SOA formed from LDGV exhaust with and without NH$_3$ was 0.013 and 0.016, respectively, which are within or near the range of semi-volatile oxygenated OA (SV-OOA) (−0.5–0) [61] and are consistent with the observation of Nordin et al. [26]. The presence of NH$_3$ had no impact on the average OS$_c$ of SOA from LDGV exhaust.

### 3.3 NH$_3$ with aged gasoline vehicle exhaust

With tremendous nitrogen fertilizer applications and animal feeding, in 2000 China contributed more than 55% of the NH$_3$ emissions in Asia [62]; in 2006, livestock waste and fertilizer in rural areas accounted for more than 85% of China’s NH$_3$ emissions [63]. Obviously, mixing of NH$_3$ with aged GVE transported from urban areas is of concern for the air quality in rural areas. In the present study we added about 100 ppb of NH$_3$ into the reactor after 4 h (experiments I-2) and 3 h of irradiation (experiment II-2) to investigate the aerosol formation from aged GVE mixed with NH$_3$. Due to the unavailability of AMS data, herein we discuss only the influence of added NH$_3$ on the number and volume concentrations of particles. When NH$_3$ was added, the number and volume concentrations jumped immediately to higher levels (Figure 5). For experiment II-2, the number and volume concentrations of particles increased extremely fast to about 1×10$^5$ cm$^{-3}$ and 40 μm cm$^{-3}$, respectively (Figure 5(a)). The addition of high concentrations of NH$_3$ might stabilize sulfuric acid clusters to overcome the nucleation barrier and thereby induce the burst particle formation [49]. If so, this would indicate that the numbers and mass of particles might be greatly enhanced when aged GVE is transported to rural areas and mixed with the substantial amounts of NH$_3$ emitted from livestock wastes and fertilizers. Reducing NH$_3$ emissions in the agricultural sector would therefore help lower both the number and mass contribu-

|         | O:C      | H:C      | OS$_c$  |
|---------|----------|----------|---------|
| II-3    | 0.684±0.042 | 1.352±0.041 | 0.016   |
| II-4    | 0.683±0.042 | 1.353±0.042 | 0.013   |

**Figure 4** Geometric average diameter (a) and concentration of organic aerosols (b) in experiments II-3 (without NH$_3$) and II-4 (with NH$_3$).

**Figure 5** Effects of NH$_3$ on the formation of particles from aging LDGV exhaust in (a) II-2 and (b) I-2.
tions of fine particles in suburban and rural areas.

4 Conclusions

Smog chamber simulations about the formation of secondary aerosols from GVE with and without NH$_3$ absorbed by denuders revealed that the presence of NH$_3$ in GVE could enhance the formation and growth of particles. This indicates that control of NH$_3$ emissions in urban areas, as well as control of NH$_3$ in GVE, would suppress particle formation and growth during the early atmospheric evolution of GVE. Chamber simulations also revealed that particle numbers and mass concentrations increased rapidly to much higher levels when aged GVE was mixed with NH$_3$, suggesting that NH$_3$ from livestock waste and fertilizer applications would promote particle formation and growth when aged urban plumes reach rural areas. The present study therefore indicates NH$_3$ plays vital roles in particle growth and formation in both urban and rural areas. We also found that NH$_3$ had no significant effects on SOA formation from GVE or on the oxidation state of SOA.

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