Secondary organic aerosol formation from photochemical aging of light-duty gasoline vehicle exhausts in a smog chamber

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Abstract. In China, a rapid increase in passenger vehicles has led to the growing concern of vehicle exhaust as an important source of anthropogenic secondary organic aerosol (SOA) in megacities hard hit by haze. In this study, the SOA formation of emissions from two idling light-duty gasoline vehicles (LDGVs) (Euro 1 and Euro 4) operated in China was investigated in a 30 m3 smog chamber. Five photooxidation experiments were carried out at 25°C with relative humidity at around 50%. After aging at an OH exposure of 5 × 10^6 molecules cm^{-3} h, the formed SOA was 12–259 times as high as primary organic aerosol (POA). The SOA production factors (PF) were 0.001–0.044 g kg^{-1} fuel, comparable with those from the previous studies at comparable OH exposure. This quite lower OH exposure than that in typical atmospheric conditions might however lead to the underestimation of the SOA formation potential from LDGVs. Effective SOA yields in this study were well fit by a one-product gas-particle partitioning model but quite lower than those of a previous study investigating SOA formation from three idling passenger vehicles (Euro 2–4). Traditional single-ring aromatic precursors and naphthalene could explain 51–90% of the formed SOA. Unspeciated species such as branched and cyclic alkanes might be the possible precursors for the unexplained SOA. A high-resolution time-of-flight aerosol mass spectrometer was used to characterize the chemical composition of SOA. The relationship between f_{43} (ratio of m/z 43, mostly C_2H_3O^+, to the total signal in mass spectrum) and f_{44} (mostly CO_2) of the gasoline vehicle exhaust SOA is similar to the ambient semi-volatile oxygenated organic aerosol (SV-OOA). We plot the O : C and H : C molar ratios of SOA in a Van Krevelen diagram. The slopes of ΔH : C / ΔO : C ranged from −0.59 to −0.36, suggesting that the oxidation chemistry in these experiments was a combination of carboxylic acid and alcohol/peroxide formation.

1 Introduction

The formation mechanisms, magnitude and chemical composition of airborne fine-particle matter (PM_{2.5}) are important to evaluate its effects on human health and climate (Hallquist et al., 2009). Organic aerosol (OA) contributes roughly ~20–50% of the total fine particle mass at continental midlatitudes (Saxena and Hildemann, 1996; Kanakidou et al., 2005). Atmospheric OA includes primary OA (POA) emitted from sources such as combustion of fossil fuels, biomass burning and volcanic eruptions, and secondary organic aerosol (SOA) formed via gas-particle conversion such as nucleation, condensation, and heterogeneous and multiphase chemistry or the aging of POA (Donahue et al., 2009;
Jimenez et al., 2009). SOA is ubiquitous and dominates the total OA in various atmospheric environments, accounting for approximately two-thirds of the total OA in urban areas to almost 90 % in urban downwind and rural areas in Northern Hemisphere midlatitudes (Zhang et al., 2007). China, for example, has serious air quality problems due to PM2.5 pollution in the recent decades (Chan and Yao, 2008; Q. Zhang et al., 2012) and SOA contributed 30–90 % of OA mass in its megacities (He et al., 2001; Cao et al., 2003; Duan et al., 2005, 2007; Yang et al., 2005; Hagler et al., 2006). However, models generally underestimate the observed OA levels mainly due to the unclear sources and formation processes of SOA (de Gouw et al., 2005; Heald et al., 2005; Johnson et al., 2006; Volkamer et al., 2006).

Vehicle exhausts emit plenty of primary PM and volatile organic compounds (VOCs) containing precursors of SOA, influencing the near-surface atmospheric chemistry and the air quality, especially in urban areas. SOA formation from diesel generators and vehicles has been widely studied in smog chambers, demonstrating that the SOA mass formed from the exhaust of diesel generators and medium- and heavy-duty diesel vehicles (HDDVs) usually exceeds the mass of emitted POA (Robinson et al., 2007; Weitkamp et al., 2007; Chirico et al., 2010; Miracolo et al., 2010; Samy and Zielinska, 2010; Nakao et al., 2011; Kroll et al., 2012). However, there are few studies on the SOA formation from gasoline vehicle exhausts. Nordin et al. (2013) investigated SOA formation from idling gasoline exhausts from three passenger vehicles (Euro 2–4), finding that C6–C9 light aromatics contributed up to 60 % of the formed SOA. While Platt et al. (2013) estimated aromatic precursors including C6–C10 light aromatics and naphthalene were responsible for less than 20 % of the SOA formed from the aging of emissions from a Euro 5 gasoline car operated during a New European Driving Cycle. To exclude the influence of a small sample size, Gordon et al. (2014) studied aging of emissions from 15 light-duty gasoline vehicles with model years ranging from 1987 to 2011, concluding that traditional precursors could fully explain the SOA from older vehicles and unspecified organics were responsible for the majority of the SOA from the newer vehicles. Therefore, chemical compositions of SOA formed from gasoline vehicle exhaust varied a lot among vehicles of different types, model years and operating conditions.

In China, the number of light-duty gasoline vehicles (LDGVs) reached 98.8 million in 2012, increasing at a rate of approximately 20 % per year since 2005 (NBSC, 2013). Furthermore, gasoline fuel in China has a relatively higher mass content of alkenes and aromatic hydrocarbons than in the USA (Schauer et al., 2002; Zhang et al., 2013), and current emission standards of LDGVs in China lag behind European countries and the USA. The emission factors of PM2.5, organic carbon (OC), element carbon (EC), NOx, SO2, NH3 and nonmethane hydrocarbons (NMHCs) for on-road vehicles in China were quite different from those in other countries (Liu et al., 2014; Y. L. Zhang et al., 2015). Therefore, it is urgent to investigate the SOA formation from vehicle exhaust in China to help make suitable policies to mitigate air pollution and also to provide valuable parameters to chemical transport models.

Here, we directly introduced dilute emissions from two idling LDGVs operated in China to a smog chamber to investigate the SOA formation. The magnitude and composition of the SOA formed from gasoline vehicle exhaust and whether traditional SOA precursors can explain the formed SOA was evaluated and discussed in this paper.

2 Materials and methods

2.1 Experimental setup

The photochemical aging experiments were carried out in the smog chamber of the 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 have been described elsewhere (Wang et al., 2014). Briefly, black lamps (1.2 m long, 60 W Philips/10R BL, Royal Dutch Philips Electronics Ltd, the Netherlands) are used as light source, providing a NO2 photolysis rate of 0.49 min−1. Two Teflon-coated fans are installed inside the reactor to guarantee proper mixing of the introduced gas species and particles within 120 s. Temperature can be set in the range from −10 to 40 °C with an accuracy of ±1 °C and is measured by eight temperature sensors inside the enclosure and one just inside the reactor. Relative humidity (RH) inside the reactor is achieved by vaporizing Milli-Q ultrapure water contained in a 0.5 L Florence flask and then flushing the water vapor into the reactor with purified dry air until target RH is reached. In the present study, temperature and RH inside the reactor were all set to 25 °C and 50 %, respectively. During the experiments, the top frame is automatically lowered to maintain a differential positive pressure inside the reactor against the enclosure to avoid the contamination of the enclosure air.

Gasoline vehicle exhausts were injected to the reactor through Teflon lines using two oil-free pumps (Gast Manufacturing, Inc, USA) at a flow rate of 40 L min−1. The injection time varied from a few minutes to more than 1 h based on the primary emissions of different vehicles. A schematic of the smog chamber and the vehicle exhaust injection system is shown in Fig. 1.

2.2 Characterization of gas- and particle-phase chemical compositions and particle sizes

Gas-phase ozone (O3) and NOx were measured online with dedicated monitors (EC9810 and 9841T, Ecotech, Australia). Online monitoring of parent VOCs such as C6–C10 single-ring aromatic hydrocarbons and their oxidation products (Liu et al., 2014; Y. L. Zhang et al., 2015). Therefore, it is urgent to investigate the SOA formation from vehicle exhaust in China to help make suitable policies to mitigate air pollution and also to provide valuable parameters to chemical transport models.

Here, we directly introduced dilute emissions from two idling LDGVs operated in China to a smog chamber to investigate the SOA formation. The magnitude and composition of the SOA formed from gasoline vehicle exhaust and whether traditional SOA precursors can explain the formed SOA was evaluated and discussed in this paper.
products were available with a commercial proton-transfer-reaction time-of-flight mass spectrometer (PTR-TOF-MS; Model 2000, I onicon Analy t i g GmbH, Austria). Detailed descriptions of the PTR-TOF-MS technique can be found elsewhere (Lindinger et al., 1998; Jordan et al., 2009). In this study the decay curve of toluene measured by PTR-TOF-MS were also used to derive the average hydroxyl radical (OH) concentration during each experiment. A wide spectrum of VOCs were also measured offline by drawing 250 ml of air inside the reactor to a Model 7100 Preconcentrator (Entech Instruments Inc., USA) coupled with an Agilent 5973N gas-chromatography mass-selective detector/flame ionization detector (GC-MSD/FID; Agilent Technologies, USA). Detailed descriptions of the method can be found elsewhere (Wang and Wu, 2008; Zhang et al., 2010; Y. Zhang et al., 2012, 2013). C2 and C3 and C4–C12 hydrocarbons were measured by GC-FID and GC-MSD, respectively. In this study, the offline measurement was the standard method to determine the mass concentrations of VOCs. PTR-TOF-MS was used for deriving the time-resolved concentrations of VOCs. The VOC concentrations measured offline were also used as an independent check of that measured online by the PTR-TOF-MS. To determine CO/CO2 concentrations before and after the introduction of exhausts, air samples were also collected into 2 L cleaned Teflon bags. CO was analyzed using gas chromatography (Agilent 6980GC, USA) with a flame ionization detector and a packed column (5A molecular sieve 60/80 mesh, 3 m × 1/8 in.) (Y. Zhang et al., 2012), and CO2 was analyzed with an HP 4890D gas chromatograph (Yi et al., 2007). The detection limits of CO and CO2 were <30 ppb. The relative standard deviations (SDs) were all less than 3 % based on seven duplicates running 1.0 ppm CO and CO2 standards (Spectra Gases Inc, USA).

A high-resolution time-of-flight aerosol mass spectrometer (HR-TOF-MS; Aerodyne Research Incorporated, USA) was used to measure the particle chemical compositions (Jayne et al., 2000; DeCarlo et al., 2006). The instrument was operated in the high sensitivity V mode and high resolution W mode alternatively every 2 min. The toolkit Squirrel 1.51H was used to obtain 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 (Aiken et al., 2007, 2008). 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 of monodisperse ammonium nitrate particles.

Particle number/volume concentrations and size distributions were measured with a scanning mobility particle sizer (SMPS; TSI Incorporated, USA., classifier model 3080, CPC model 3775). Flow rates of sheath and aerosol flow were 3.0 and 0.3 L min−1, respectively, allowing for size distribution scanning ranging from 14 to 700 nm within 255 s. The accuracy of the particle number concentration is ±10 %. An aerosol density of 1.4 g cm−3 was assumed to convert the particle volume concentration into the mass concentration (Zhang et al., 2005). Conductive silicon tubes were used as sampling lines for HR-TOF-MS and SMPS to reduce electrostatic losses of particles.

2.3 Experimental procedure

Two light-duty gasoline-powered vehicles were used in this study, one Euro 1 vehicle and one Euro 4 vehicle. They are both port fuel injected vehicles. More details of the vehicles are listed in Table 1. Both of the vehicles were fueled with Grade 93 gasoline, which complies with the Euro III gasoline fuel standard. Details of the oil compositions can be found in our previous study (Zhang et al., 2013).

Prior to each experiment, the reactor was evacuated and filled with purified dry air at least 5 times, then the reactor was flushed with purified dry air for at least 48 h until no residual hydrocarbons, O3, NOx, or particles were detected in the reactor to avoid carry-over problems from day-to-day experiments. Prior to the introduction of exhaust, the temperature control system and Teflon-coated fans were turned on. The exhaust could be injected when the temperature in the reactor was stable at the set temperature of 25°C.

The LDGV was parked outside the laboratory and tested while idling. Before the injection of exhaust, the cars were idling for at least half an hour to warm up the three-way catalysts, and then the vehicle exhausts were injected into the reactor. During their introduction, the raw exhausts were also sampled in 8 L cleaned aluminum foil bags by a mechanical pump with a flow rate of about 5 L min−1. VOCs and CO2 in these samples were measured offline by the same methods mentioned above to characterize the primary emissions from the exhaust pipe. The exhaust in the reactor was diluted by a factor of 13–30 compared to the tailpipe.

Additional NO was then added to adjust the VOC/NOx ratios to around 10.0 or 2.0 (Table 2), within the range of 0.5–
10 reported in gasoline vehicle exhaust tests and downwind of urban areas (Clairotte et al., 2013). The initial concentrations of NO\textsubscript{x} at the start of the experiments ranged from 134 to 956 ppb. In each experiment CH\textsubscript{3}CN was used as an indicator of dilution in the reactor. After being characterized in the dark for more than 30 min, the exhaust was exposed to black light continuously for 5 h. After the black lamps were switched off, the formed SOA was characterized for another 2–3 h to correct the particle wall loss. Blank experiments with no vehicle exhaust introduced were performed to quantify the reactivity of the matrix gas. After 5 h of irradiation, the number and mass of formed particles were < 5 cm\textsuperscript{-3} and 0.1 µg m\textsuperscript{-3}, respectively.

During the introduction of exhausts, particles and VOCs might deposit to the surface of the transfer lines. Therefore, a flow rate of as high as 20 L min\textsuperscript{-1} and a transfer line of as short as 5 m were used to provide residence time within seconds and thus reduce the losses of particles and VOCs in the transfer lines. Furthermore, before being introduced into the reactor, exhausts were generally pumped through the transfer lines for half an hour to saturate the transfer lines with particles and VOCs while warming the catalytic converter. Losses of particles and VOCs in the introduction lines were determined by comparing the concentrations of total particle number and VOCs in the directly emitted exhausts with the ones after passing through the transfer lines. The loss of total particle number was estimated to be less than 3 %. The penetration efficiency of particles due to diffusion in a cylindrical tube, \( \eta(d_p) \), can also be estimated by a laminar diffusional deposition model (Gormley and Kennedy, 1949). For particles with diameters larger than 10 nm, the penetration efficiency was higher than 95 %, indicating minor losses of particles in the transfer line. The losses of VOCs in the transfer line were estimated to be less than 5 %, which might lead to a small underestimation of SOA production.

### 2.4 Data analysis

#### 2.4.1 Wall-loss corrections

The loss of particles and organic vapors to the reactor walls has to be accounted for to accurately quantify the SOA formation. The loss of particles onto the walls has been well constrained and is treated as a first-order process (McMurry and Grosjean, 1985). The wall-loss rate constant was determined separately for each experiment by fitting the SMPS and aerosol mass spectrometer (AMS) data with first-order kinetics when UV lamps were turned off. By applying this rate to the entire experiment, we use the same method as Pathak et al. (2007), treating the particle wall loss as a first-order process to correct the wall loss of the particles. The wall loss of particles is a size-dependent process; therefore, the influence of nucleation needs to be examined due to the rapid loss of nucleation mode particles. In this study, the impact of the nucleation event on wall-loss estimate is considered to be negligible for only less than 3 % of the particle mass is in the nucleation mode 10 min after nucleation for all the experiments (Fig. S1 in the Supplement). In general, the loss of condensable organic vapors to the walls is estimated for two limiting cases (Weitkamp et al., 2007; Hildebrandt et al., 2009). In the first case (designated \( \omega = 0 \)), no organic vapors is lost to the walls (only to suspended particles). In the second case (designated \( \omega = 1 \)), the particles on the walls are in equilibrium with the organic vapors; therefore, condensation to the particles on the walls is identical to the suspended particles. We use the \( \omega = 0 \) wall-loss correction assuming the organic vapors’ only condensation onto suspended particles. The \( \omega = 1 \) wall-loss correction is not suitable for the experiments here in which nucleation occurred and no seed particles were added (Henry et al., 2012).

#### 2.4.2 AMS data corrections

Theoretically, the sum of the PM mass measured by AMS should be equal to the mass calculated from the SMPS mass size distributions. However, both methods have limitations. One must assume a particle shape and density to convert the volume concentration measured by SMPS to the mass concentration. Here, we assume that particles are spherical with an average density of 1.4 g cm\textsuperscript{-3} (Zhang et al., 2005). Fractal-like particles will cause the overestimate of the spherical equivalent diameter, thus overestimating the particle mass. AMS tends to underestimate the PM mass due to the transmission efficiency (Liu et al., 2007) and the AMS collection efficiency (Gordon et al., 2014), leading to the discrepancy between the AMS data and SMPS data. Figure S2 shows the particle volume distribution measured by SMPS for a typical smog chamber experiment (experiment 2). Most particles were in the range 40–120 nm after SOA formation. Since the transmission window of the standard lens of HR-TOF-AMS is 60–600 nm (aerodynamic diameter) (Liu et al., 2007), particles with diameter smaller than 40 nm (mobility diameter) were cut from the lower edge of the volume distri-

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Table 1. Detailed information of the two light-duty gasoline vehicles.

<table>
<thead>
<tr>
<th>ID</th>
<th>emission standard class</th>
<th>vehicle model year</th>
<th>mileage (km)</th>
<th>displacement (cm\textsuperscript{3})</th>
<th>power (kW)</th>
<th>weight (kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Euro 4</td>
<td>Golf</td>
<td>2011</td>
<td>25 000</td>
<td>1598</td>
<td>77</td>
</tr>
<tr>
<td>II</td>
<td>Euro 1</td>
<td>Accord</td>
<td>2002</td>
<td>237 984</td>
<td>2298</td>
<td>110</td>
</tr>
</tbody>
</table>
To determine the mass of reacted organic precursors, evacuating stainless steel canisters and analyzing by GC-MSD. The AMS measured by HR-TOF-AMS; thus, we assumed that the mass of particles measured by SMPS was comparable with that measured by the AMS. As shown in Fig. S3, the mass of primary SOA measured by AMS was outside the transmission window of HR-TOF-MS, to correct the AMS data.

For all the experiments with discrepancies between the AMS and SMPS data (Fig. S3), we assume that the difference in mass has the same composition as the measured components. We then calculate scaling factors, AMS$_{sf}$, to correct the PM mass measured by AMS and make it accordant with the SMPS measurements. The scaling factor is

$$AMS_{sf} = \frac{C_{SMPS}}{C_{ Org} + C_{SO_4} + C_{NO_3} + C_{NH_4}}.$$  

where $C_{SMPS}$ is the total particle mass concentration derived by the SMPS, $C_{Org}$, $C_{SO_4}$, $C_{NO_3}$ and $C_{NH_4}$ are the mass concentrations of organics, sulfate, nitrate and ammonium measured by the AMS. As shown in Fig. S3, the mass of primary particles measured by SMPS was comparable with that measured by HR-TOF-AMS; thus, we assumed that the mass of black carbon (BC) in the reactor was negligible. The AMS$_{sf}$ for each time step after nucleation is calculated and used to scale the AMS data for the entire experiment.

### 2.4.3 Effective SOA yields

To compare the SOA formation with other studies, we calculated effective SOA yields for all experiments. The effective SOA yield $Y$ was defined as the ratio of the wall-loss-corrected SOA mass to the mass of reacted organic precursors (Odum et al., 1996, 1997; Donahue et al., 2006). In this study, reacted organic precursors included in calculation are only those quantified by GC-MSD, including benzene, toluene, $C_2$-benzene, $C_3$-benzene, $C_4$-benzene and naphthalene. A detailed list of these compounds is presented in Table S2. At the beginning and end of each experiment, air samples in the reactor were collected into 2 L electropolished and evacuated stainless steel canisters and analyzed by GC-MSD to determine the mass of reacted organic precursors.

### 2.4.4 Emission factors

Emission factor (EF) of a pollutant $P$ is calculated on a fuel basis (g kg$^{-1}$):

$$EF = 10^3 \cdot \frac{[\Delta P]}{[\Delta CO_2] + [\Delta CO] + [\Delta C]} \cdot \frac{\omega_C}{MW_C},$$

where $[\Delta P]$, $[\Delta CO_2]$, $[\Delta CO]$, and $[\Delta C]$ are the background corrected concentrations of $P$, CO$_2$, CO and the total hydrocarbons in the reactor (in µg m$^{-3}$); $MW_{CO_2}$, $MW_{CO}$, $MW_{HC}$, and $MW_C$ are the molecular weights of CO$_2$, CO, HC and C. $\omega_C$ (0.85) is the carbon intensity of the gasoline (Kirchstetter et al., 1999).

### 2.4.5 Determination of OH exposure

Decay of toluene measured by PTR-TOF-MS is used to derive the average OH concentration during each experiment. Changes in the toluene concentration over time can be expressed as

$$\frac{d[\text{toluene}]}{dt} = -k \cdot [\text{OH}] \cdot [\text{toluene}],$$

where $k$ is the rate constant for the reaction between toluene and OH radical. Assuming a constant OH concentration during an experiment, we can integrate Eq. (3) to get Eq. (4):

$$\ln \left( \frac{[\text{toluene}]_0}{[\text{toluene}]_t} \right) = k \cdot [\text{OH}] \cdot t.$$  

So, by plotting $\ln([\text{toluene}]_0/[\text{toluene}]_t)$ vs. time $t$, we can obtain a slope that equals $k \cdot [\text{OH}]$. The average OH concentration is therefore calculated as

$$[\text{OH}] = \frac{\text{slope}}{k}.$$  

The OH exposure is then determined through multiplying the average OH concentration by time.

### 3 Results and discussion

#### 3.1 VOC composition

Figure 2 shows the average composition of gasoline vehicle exhausts from vehicle I (Euro 4) and II (Euro 1). For Euro 4...
and 1 vehicles, alkanes contributed about 42.9 and 66.2 % of the total speciated VOCs measured with the GC-FID/MSD by mass, respectively, dominating the speciated VOC emissions in gasoline vehicle exhausts. Due to the high concentrations of isopentane and methylpentane, branched alkanes contributed approximately 44.9 % of the total VOCs for the Euro 1 vehicle, quite higher than that for the Euro 4 vehicle (23.3 %). Aromatic hydrocarbons accounted for about 38.0 and 22.5 % of the total VOCs for Euro 4 and 1 vehicles, respectively, relatively higher than the 10–15 % observed by Nordin et al. (2013) for idling Euro 2, 3 and 4 vehicles. The mass fraction of aromatic hydrocarbons for the Euro 4 vehicle was comparable with 32.2 % for idling private cars in Hong Kong (Guo et al., 2011) and 38.3 % for Euro 3 light-duty gasoline vehicles operated through ECE cycles with an average speed around 18.7 km h$^{-1}$ (Wang et al., 2013). Both Schauer et al. (2002) and Gentner et al. (2013) observed that aromatic hydrocarbons contributed around 27 % of the total VOCs for gasoline-powered automobiles driven through the cold-start Federal Test Procedure urban driving cycle and on-road gasoline vehicles in the Caldecott tunnel, similar to that of the Euro 1 vehicle. Recently, Huang et al. (2015) reported that mass fractions of aromatic hydrocarbons were as high as 46.4 % for Euro 1, 2, and 3 light-duty gasoline vehicles operated through ECE cycles. Therefore, the variations of the composition of LDGV exhausts in this study were within the range of previous studies.

The averaged emission factors of VOCs and aromatic hydrocarbons for the Euro 4 vehicle were 2.1 and 0.8 g kg$^{-1}$, approximately 26.0 and 43.5 % of those for the Euro 1 vehicle, respectively. Compared with a Euro 5 gasoline vehicle operated during a New European Driving Cycle, the emission factor of VOCs for the Euro 4 vehicle was about 1.7 times higher (Platt et al., 2013). Using 12.7 km L$^{-1}$ as the average fuel efficiency (Wagner et al., 2009), we obtained the VOC emission factors based on grams per kilometer for Euro 4 and 1 vehicles to be 0.12 and 0.46 g km$^{-1}$, respectively, comparable with the previous reported values for Euro 1 and 4 gasoline vehicles in China (Huo et al., 2012; Huang et al., 2015). According to previous studies, there is a clear reduction of VOC emissions from gasoline vehicles with stricter emission standards (Huo et al., 2012; Huang et al., 2015). It is worth noting that emissions of HC from gasoline vehicles during idling were observed to be lower than those in the acceleration and deceleration modes (Tong et al., 2000; Yamamoto et al., 2012; Huang et al., 2013), but in a similar level with those in the cruising mode (Tong et al., 2000). It is important to note that the reported data are only based on five chamber experiments with two LDGVs under idling conditions. More tests are needed to assess SOA formation from gasoline vehicle exhausts in China.

3.2 SOA formation

Figure 3 shows the temporal evolution of gas-phase and particle-phase species during a typical smog chamber experiment. During the period $-1.3$ to $-0.85$h, the vehicle exhausts were introduced into the reactor. At time $-0.55$h, the relative humidity was adjusted to approximately 40 % and HR-TOF-AMS was connected to characterize the primary PM. NO was injected to adjust the VOC/NO$_x$ ratio at the approximate time of $-0.25$h. After the black lamps were turned on, NO was fast converted to NO$_2$ in less than 1 h and then O$_3$ was accumulated and OH radicals were formed. When NO concentration decreased to a low level of about 5 ppb, gas-phase light aromatics, especially C$_3$-benzene with higher reactivity, decayed rapidly due to the reaction with the OH radical. SOA was thus rapidly formed and increased to a high level in less than 2 h. As shown in Table 3, at the end of all the experiments, the formed SOA was $12$–$259$ times as high as POA. This enhancement is consistent with $9$–$500$ times increase recently reported by Nordin et al. (2013) when studying SOA formation from idling European gasoline passenger vehicle emissions. As shown in Fig. 3c, the total particle number concentration increased fast from 82 to $116,143$ cm$^{-3}$, indicating dramatic new particle formation, which might be due to the starting surface concentrations of particles being below a critical value (100–2000 µm$^2$ cm$^{-3}$; Table S1) (Wehner et al., 2004). As shown in Table S1, primary particle numbers in the reactor in this study ranged from 82 to 18,948 cm$^{-3}$, 1–2 orders of magnitude higher than that of a Euro 2 car operated at idling with a similar dilution ratio (Nordin et al., 2013), indicating that the small starting particle number concentrations might mainly be due to the idling condition of tested cars rather than the losses in the introduction lines. In addition, upon entering into the chamber, emitted particles would partition due to dilution, as in the atmosphere, regardless of the temperature and concentration in the sampling system, which might lead to the decrease of starting number concentrations. A certain extent of primary particles under the detection limit of 14 nm of SMPS also contributed to the measured small starting number concentration of particles.

Deposition of SOA-forming vapors on the walls might lead to the underestimation of SOA production. The wall-loss rate coefficient of vapors is related to the numbers of carbon and oxygen in the molecule (X. Zhang et al., 2015). Here, we take C$_7$H$_{8}$O$_4$, a product of the photo-oxidation of toluene, as an example. The loss of C$_7$H$_{8}$O$_4$ to walls would be 7 % in an hour before SOA formation when a wall-deposition rate of $2 \times 10^{-5}$ s$^{-1}$ was used (X. Zhang et al., 2015). After SOA formation, the surface concentrations of particles increased fast to as high as 2000 µm$^2$ cm$^{-3}$ in an hour, which would reduce the vapor wall losses.

SOA production factors (PFs) for the LDGVs tested in this study were estimated to vary from 0.001 to 0.044 g kg$^{-1}$ fuel, within the results of Nordin et al. (2013).
Table 3. Summary of the results for the light-duty gasoline vehicle photo-oxidation experiments.

<table>
<thead>
<tr>
<th>Experiment ID</th>
<th>Vehicle ID</th>
<th>OH (× 10^6 molecules cm^{-3})</th>
<th>POA (µg m^{-3})</th>
<th>SOA (µg m^{-3})</th>
<th>SOA/POA effective yield</th>
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<tr>
<td>1</td>
<td>I</td>
<td>1.23</td>
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<td>51.1</td>
<td>46</td>
</tr>
<tr>
<td>2</td>
<td>II</td>
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<td>0.2</td>
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</tr>
<tr>
<td>3</td>
<td>I</td>
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<td>0.3</td>
<td>77.6</td>
<td>259</td>
</tr>
<tr>
<td>4</td>
<td>I</td>
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<td>1.0</td>
<td>125.4</td>
<td>125</td>
</tr>
<tr>
<td>5</td>
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<td>0.79</td>
<td>0.3</td>
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</tbody>
</table>

Figure 2. Concentration–time plots of gas-phase and particle-phase species and particle number concentration distribution as a function of time during a typical smog chamber experiment (experiment 2): (a) NO, NO_2, O_3 (left y axis) and organic aerosol (right y axis); (b) gas-phase light aromatics (measured by PTR-TOF-MS) (benzene characterized by m/z 79; toluene characterized by m/z 93; C_2-benzene characterized by m/z 107; C_3-benzene characterized by m/z 121; C_4-benzene characterized by m/z 135); (c) particle size–number concentration distributions as a function of time. The vehicle exhaust was introduced into the reactor between -1.3 and -0.85 h; the primary emissions were characterized from -0.85 to 0 h; at time 0 h, the black lamps were turned on.

Figure 3. Composition of (a) VOCs and (b) aromatics of gasoline vehicle exhausts from Euro 1 and Euro 4 private cars, presented as weight percentage of speciated VOCs. C_2 and C_3 and C_4–C_12 hydrocarbons were measured by GC-FID and GC-MSD, respectively. The error bars (1σ) represent variability from measurements for each vehicle.

and Gordon et al. (2014) with OH exposure of around 5.0 × 10^6 molecules cm^{-3} h. A recent study investigating SOA formation from in-use vehicle emissions in a highway tunnel in Pittsburgh indicated that the peak SOA production was measured at an OH exposure of 1.9 × 10^8 molecules cm^{-3} h and current smog chamber studies may underestimate the ultimate SOA production by a maximum factor of about 10 due to the limited OH exposure (Tkacik et al., 2014).
The average OH radical concentration (Table 3) was determined to be 0.79–1.23 × 10^6 molecules cm^{-3} during our experiments. This OH level was about 10 times lower than the average OH concentration of 1.5 × 10^7 molecules cm^{-3} around noon in summer in the Pearl River Delta, China (Hofzumahaus et al., 2009). The OH exposure in this study is only 5 × 10^6 molecules cm^{-3} h, equivalent to 0.3 h of atmospheric oxidation. Therefore, the real-world SOA production factor from LDGVs in the atmosphere in China may be even higher than our estimation.

3.3 SOA yield

Effective SOA yield from vehicle exhaust calculated as described in Sect. 2.4.3 ranged from 2.8 to 17.2%. Pankow (1994a, b) and Odum et al. (1996) indicated that Y is a function of \( M_0 \) and the relation is described as

\[
Y = M_0 \left[ \frac{\alpha_i K_{om,i}}{1 + K_{om,i} M_0} \right],
\]

(6)

where \( K_{om,i} \) and \( \alpha_i \) are the mass-based absorption equilibrium partitioning coefficient and stoichiometric coefficient of product \( i \), respectively; \( M_0 \) is the total mass concentration of organic material. As shown in Table 3, SOA yields for the Euro 1 vehicle were around 3%, quite lower than 10–17% for the Euro 4 vehicle. The mass fraction of aromatic hydrocarbons for the Euro 4 vehicle was about 2 times higher than that for the Euro 1 vehicle (Fig. 2a), which would form more semi-volatile organic compounds (SVOCs) partitioning into particle phase under similar OH exposure and thus lead to the relatively higher SOA yields.

Comparison of effective yield data obtained for the LDGV exhaust in this study with those of Nordin et al. (2013) is shown in Fig. 4. Effective yield data of this study are well fit with the one-product model, namely \( Y = M_0 \left[ \frac{\alpha_i K_{om,i}}{1 + K_{om,i} M_0} \right] \). The appropriate values for \( \alpha_i \) and \( K_{om,1} \) when fitting the yields are 0.350 ± 0.114 and 0.007 ± 0.004, respectively. The effective SOA yields in the study of Nordin et al. (2013) were 60–360% higher than those in this study at the same concentrations of \( M_0 \). In their calculation of the reacted SOA precursors, \( C_4 \)-benzene and naphthalene were excluded. The effective SOA yields would increase 7–34% when \( C_4 \)-benzene and naphthalene were excluded in this study, which could explain a small portion of the discrepancy. According to the estimation above, the loss of VOCs in the transfer lines was less than 5%. A little higher than VOCs, if assumed to be 20%, losses of intermediate volatility organic compounds (IVOCs) and SVOCs in the transfer lines would increase the SOA effective yields by a factor of 2–10% when the unexplained SOA discussed later was all attributed to the contribution from IVOCs and SVOCs. The existence of seed particles in the study of Nordin et al. (2013) might reduce the wall loss of semi-volatile organic vapors and thus increase the effective SOA yield (Kroll et al., 2007; Zhang et al., 2014; X. Zhang et al., 2015). However, Cocker et al. (2001) found that SOA formation from \( m \)-xylene and 1,3,5-trimethylbenzene photo-oxidation was unaffected by the presence of ammonium sulfate seed aerosols. The influence of seed particles on SOA yields still needs further investigations. Faster oxidation rates caused by higher OH concentrations in the study of Nordin et al. (2013) would also result in higher SOA yields (Ng et al., 2007). Additionally, the different VOC profiles of exhausts might also influence the SOA yields.

SOA production from the reacted organic precursors can be estimated by the following formula:

\[
\Delta SOA_{predicted} = \sum_j (\Delta X_j \times Y_j),
\]

(7)

where \( \Delta SOA_{predicted} \) is the predicted SOA concentration (in \( \mu g \ m^{-3} \)); \( \Delta X_j \) is the mass of reacted aromatic hydrocarbon \( X_j \) (in \( \mu g \ m^{-3} \)); and \( Y_j \) is the corresponding SOA yield of \( X_j \). In this study, the SOA yield of benzene and other single-ring aromatics were estimated using the two-product model curves taken from Borräs et al. (2012) and Odum et al. (1997), respectively. While the SOA yield of naphthalene was taken from Shakya et al. (2010). SOA yield curves of toluene and \( m \)-xylene from Ng et al. (2007) were also widely used to estimate SOA production (Platt et al., 2013). However, the introduction of seed aerosols and OH precursor made the SOA yield curves in the study of Ng et al. (2007) not suitable for this study. Considering that the study of Odum et al. (1997) provided a systematic estimation of SOA yields from toluene, \( C_2 \)-benzene, \( C_3 \)-benzene and \( C_4 \)-benzene, we mainly used the two-product curves from Odum et al. (1997) to estimate the SOA production. The aerosol yield curves from the literature were converted to the same aerosol density of 1.4 g cm^{-3} in this study. The SOA yield for each precursor was calculated for the measured con-
contributions of the predicted benzene SOA, toluene SOA, C\(_2\)-benzene SOA, C\(_3\)-benzene SOA, C\(_4\)-benzene SOA and naphthalene SOA to the total formed SOA in all experiments.

Figure 5. Contributions of the predicted benzene SOA, toluene SOA, C\(_2\)-benzene SOA, C\(_3\)-benzene SOA, C\(_4\)-benzene SOA and naphthalene SOA and to the total formed SOA in all experiments.

concentration of OA in the reactor. Then, the predicted SOA production from each precursor can be calculated (Table S2).

Figure 5 shows the contributions of the predicted benzene SOA, toluene SOA, C\(_2\)-benzene SOA, C\(_3\)-benzene SOA, C\(_4\)-benzene SOA and naphthalene SOA to the total measured SOA in all experiments. C\(_4\)-benzene contributed negligible SOA because of the very low emissions of C\(_4\)-benzene from light-duty gasoline vehicles (Fig. 2b). Though benzene accounted for a relatively higher percentage of the total VOCs, benzene also accounted for a negligible proportion of the formed SOA due to its low reactivity with OH radicals. Naphthalene was previously estimated to contribute around 5% of the vehicle SOA mass (Nordin et al., 2013). While in this study naphthalene was calculated to contribute 8–52% of the formed SOA. The initial concentrations of naphthalene in this study ranged from 8.5 to 39.5 ppb, much higher than 2.8–4.4 ppb in the study of Nordin et al. (2013). The high contributions of naphthalene are probably attributed to its relatively higher initial concentrations and higher mass yield than single-ring aromatics for similar experimental conditions (Odum et al., 1997; Ng et al., 2007).

In total, single-ring aromatics and naphthalene accounted for 51–90% of the measured SOA, comparable to the estimation that classical C6–C9 light aromatics were responsible for 60% of the formed SOA from gasoline vehicle exhausts (Nordin et al., 2013), indicating that there are other SOA precursors in the LDGV exhausts. Platt et al. (2013) attributed the unexplained SOA formed from the aging of emissions from a Euro 5 gasoline car to highly oxygenated hydrocarbons. In addition, IVOCs such as branched and cyclic alkanes were recognized as important SOA precursors derived from wood burning, diesel engine and aircraft exhaust (Robinson et al., 2007; Weitkamp et al., 2007; Grieshop et al., 2009; Tkacik et al., 2012). Gordon et al. (2014) found that unspeciated species including branched and cyclic alkanes contributed about 30% of the nonmethane organic gas emissions from LDGVs with model years of 1995 or later and are associated with the majority of the SOA formation. Tkacik et al. (2014) also found that unspeciated species were predicted to contribute twice as much SOA from in-use vehicle emissions as traditional precursors. It is worth noting that photooxidation of aromatic hydrocarbons in a complex mixture such as gasoline vehicle exhausts might alter the SOA yield compared to pure precursor experiments, thus, probably influencing the estimation in this study (Song et al., 2007). Wall losses of organic vapors were not considered in this study, which would lead to the underestimation of SOA production. Therefore, the mass closure analysis estimated the maximum amount of SOA that could be explained by aromatics.

Figure 6. (a) The fractions of total organic signal at m/z 43 (f\(_{43}\)) vs. m/z (f\(_{44}\)) at the end of each experiment together with the triangle plot of Ng et al. (2010). The solid square and triangles represent the results of Presto et al. (2014) and Nordin et al. (2013), respectively. The dotted lines define the space where ambient OOA components fall. The ranges of f\(_{44}\) observed for SV-OOA and LV-OOA components are 0.03–0.11 and 0.13–0.21, respectively. (b) Van Krevelen diagram of SOA from light-duty gasoline vehicle exhaust. Dotted lines are to show slopes of 0, −1 and −2. AMS data of the experiment 5 were unavailable.
3.4 SOA composition

Fragmentations derived from the AMS data have been widely used to explore the oxidation degree of the organic aerosols (Zhang et al., 2005; Ng et al., 2010; Heald et al., 2010). The usually used ion fragments include m/z 43, 44 and 57. The dominating organic peaks in gasoline vehicle exhaust SOA are m/z 43 and 44 (Nordin et al., 2013), while m/z 57 is a main hydrocarbon fragment in diesel SOA (Chirico et al., 2010). Here, we use the approach of Ng et al. (2010) by plotting the fractions of total organic signal at m/z 43 (f_{43}) vs. m/z 44 (f_{44}) together with the triangle defined according to the analysis of ambient AMS data. The m/z 43 signal includes C_3H_7^+ and C_2H_5O^+ ions, indicating fresh less-oxidized organic aerosols. The m/z 44 signal, dominated by CO_2^+ and formed from the thermal decarboxylation of organic acids, is an indicator of highly oxygenated organic aerosols (Ng et al., 2010).

Figure 6a shows the f_{43} vs. f_{44} at the end of each experiment and the results of Nordin et al. (2013) and Presto et al. (2014), together with the triangle developed by Ng et al. (2010). The ambient low-volatility oxygenated OA (LV-OOA) and semi-volatile OOA (SV-OOA) factors fall in the upper and lower portions of the triangle, respectively. Our data mainly lie in the SV-OOA region, similar to the results of Nordin et al. (2013) and Presto et al. (2014). However, SOA in one experiment shows relatively lower oxidation degree. This phenomenon reflects the different SOA compositions among different experiments and might be caused by the different VOCs profiles, OH exposure and organic mass loadings (Ng et al., 2010).

The O : C ratio can also be used to characterize the oxidation degree of the organic aerosols. After 5 h of irradiation the H : C ratios varied from 1.22 to 1.37 and the O : C ratios from 0.43 to 0.69 for all the experiments. Almost all the O : C values were lower than 0.6, comparable to the SV-OOA compounds, which typically have O : C ratios between 0.3 and 0.6 (Jimenez et al., 2009). Platt et al. (2013) observed a relatively higher O : C ratio of 0.7 on the aging (OH = 12 × 10^6 molecules cm\(^{-3}\) h) of emissions from a Euro 5 gasoline car operated during a New European Driving Cycle. As a higher OH exposure will lead to a higher O : C ratio, if the gasoline exhaust in this study was irradiated under a similar OH exposure to that of Platt et al. (2013), the O : C ratios might reach the similar level or higher, comparable to the LV-OOA factor (Jimenez et al., 2009).

In Fig. 6b we plot the O : C and H : C molar ratios after SOA was formed during experiments 1, 2 and 3 on a Van Krevelen diagram (Heald et al., 2010). The slopes ranged from −0.59 to −0.36, similar to previous laboratory studies of Tkacik et al. (2012) for cyclic, linear and branched alkanes, Jathar et al. (2013) for unburned fuel and Presto et al. (2014) for light-duty gasoline vehicle exhaust. They are also similar to the ambient data (Ng et al., 2011). A slope of −1, −0.5 and 0 in the Van Krevelen diagram represents the addition of alcohol/peroxide, the addition of carboxylic acid with fragmentation, and the addition of carboxylic acid without fragmentation, respectively (Heald et al., 2010; Ng et al., 2011). Consequently, the slopes in this study indicate that the SOA formation is a combination of the addition of both carboxylic acid and alcohol/peroxide function groups without C–C bond cleavage and/or the addition of carboxylic acid with C–C bond breakage (Heald et al., 2010; Ng et al., 2011).

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