Measurement report: Emissions of intermediate-volatility organic compounds from vehicles under real-world driving conditions in an urban tunnel

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Abstract

Intermediate-volatility organic compounds (IVOCs) emitted from vehicles are important precursors to secondary organic aerosols (SOA) in urban areas, yet vehicular emission of IVOCs, particularly from on-road fleets, is poorly understood. Here we initiated a field campaign to collect IVOCs with sorption tubes at both the inlet and the outlet in a busy urban tunnel (>30,000 vehicles per day) in south China for characterizing emissions of IVOCs from on-road vehicles. The average emission factor of IVOCs (EF_{IVOCs}) was measured to be 16.77 ± 0.89 mg km\(^{-1}\) (Average ± 95% C.I.) for diesel and gasoline vehicles in the fleets, and based on linear regression the average EF_{IVOCs} was derived to be 62.79 ± 18.37 mg km\(^{-1}\) for diesel vehicles and 13.95 ± 1.13 mg km\(^{-1}\) for gasoline vehicles. The EF_{IVOCs} for diesel vehicles from this study was comparable to that reported previously for non-road engines without after-treatment facilities, while the EF_{IVOCs} for gasoline vehicles from this study was much higher than that recently tested for a China V gasoline vehicle. IVOCs from the on-road fleets did not show significant correlation with the primary organic aerosol (POA) or total non-methane hydrocarbons (NMHCs) as results from previous chassis dynamometer tests. Estimated SOA production from the vehicular IVOCs and VOCs surpassed the POA by a factor of ~ 2.4, and IVOCs dominated over VOCs in estimated SOA production by a factor of ~ 7, suggesting that controlling IVOCs is of greater importance to modulate traffic-related OA in urban areas. The results demonstrated that although on-road gasoline vehicles have much lower EF_{IVOCs}, they contribute more IVOCs than on-road diesel vehicles due to its dominance in the on-road fleets. However, due to greater diesel than gasoline fuel consumption in China, emission of IVOCs
from diesel engines would be much larger than that from gasoline engines, signaling the

overwhelming contribution of IVOC emissions by non-road diesel engines in China.
1 Introduction

Intermediate-volatility organic compounds (IVOCs) refer to organics with effective saturated concentrations ranging from $10^3$ to $10^6$ μg m$^{-3}$, roughly corresponding to the volatility range of C$_{12}$-C$_{22}$ normal alkanes (n-alkanes) (Donahue et al., 2006; Zhao et al., 2014). Robinson et al. (2007) have demonstrated that IVOCs, as the missing secondary organic aerosol (SOA) precursors in many model studies, could efficiently narrow the gap between model predicted and field observed SOA. Smog chamber studies involving individual IVOCs species, like higher n-alkanes and 2-ring aromatics, have confirmed their significantly higher SOA formation potentials (Chan et al., 2009; Presto et al., 2010; Liu et al., 2015). In addition, recent model simulations including IVOCs as SOA precursors revealed that 30% ~ 80% of ambient SOA could be explained by IVOCs (Ots et al., 2016; Zhao et al., 2016; Yang et al., 2019; Lu et al., 2020; Huang et al., 2020). However, due to lack of direct measurements, these model simulations used the ratios of IVOCs to other species like primary organic aerosol (POA) or non-methane hydrocarbons (NMHCs) to estimate IVOCs emissions.

Vehicular emission is an important anthropogenic source of IVOCs especially in urban environments (Tkacik et al., 2014; Jathar et al., 2014; Cross et al., 2015; Zhao et al., 2015, 2016; Ots et al., 2016). IVOCs could account for ~ 60% of non-methane hydrocarbons (NMHCs) from diesel vehicles and 4 ~ 17% from gasoline vehicles, explaining a dominant portion of estimated SOA mass from diesel and gasoline exhaust (Zhao et al., 2015, 2016). Previous chamber simulations on SOA formation from vehicle exhaust revealed that traditional volatile organic compounds (VOCs) could not explain the formed SOA, and IVOCs instead might dominate the SOA productions (Deng et al., 2020; Zhang et al., 2020). In megacities like London, diesel-emitted IVOCs alone could contribute ~ 30% SOA formed in ambient air (Ots 2016).
et al., 2016). Therefore, for the control of fine particle pollution in urban areas, it is necessary to compile and upgrade emission inventories for IVOCs, and more works are needed to characterize their emissions from on-road vehicles.

Although previous chassis dynamometer tests used limited numbers of vehicles to characterize IVOCs emission (Zhao et al., 2015, 2016; Tang et al., 2021), the results obtained from the tests were widely applied to recent models and emission inventories (Liu et al., 2017; Lu et al., 2018; Wu et al., 2019; Huang et al., 2020). However, driving conditions were recently found to significantly influence vehicular IVOCs emissions (Drozd et al., 2018; Tang et al., 2021), highlighting the importance of conducting on-road measurements of vehicle-emitted IVOCs under real-world driving condition, which could further narrow the uncertainty of vehicular IVOCs estimates in models and emission inventories. Tunnel test is a widely used method to characterize vehicle emissions in light of its advantage in capturing real-world emissions with a large number of driving vehicles. The emissions of PM$_{2.5}$, carbonaceous aerosols, VOCs, NOx, and NH$_3$ from on-road vehicles have been widely studied based on tunnel tests (Liu et al., 2014; Zhang et al., 2016, 2017, 2018). However, to the best of knowledge, till present no reports are available about vehicular emission factors of IVOCs through tunnel tests.

In China, the number of on-road vehicles reached 348 million in 2019, more than double that in 2009 (http://www.mee.gov.cn/hjzl/sthjzkydyhjgl/). However, emissions of IVOCs from mobile sources in China are much understudied. Only very recently, Tang et al. (2021) tested emission of IVOCs from a China V light-duty gasoline vehicle. For this reason, IVOC emission factors derived from vehicle tests in the US have been used to update China’s emission inventories with the inclusion of IVOCs (Liu et al., 2017). It is unknown whether the borrowed
emission factors could well reflect the vehicular emissions of IVOCs in China. On the other hand, although China has made great achievements in combating air pollution in recent years, fine particle pollution is still an air quality problem in many of China’s cities (Wang et al., 2020).

As organic matters are often the most abundant components in PM$_{2.5}$ and SOA pollution is increasingly standing out with the intensified primary emission control (Guo et al., 2020), understanding IVOC emissions from on-road vehicles is of great importance given that vehicle-emitted IVOCs contribute greatly to urban SOA formation (Gentner et al., 2012; Wu et al., 2019; Huang et al., 2020).

In this study, the emissions of IVOCs from on-road vehicles under real-world driving conditions were characterized through tests in an urban tunnel in Guangzhou, a megacity in south China. The study aims to: 1) investigate chemical compositions and volatility of IVOCs from on-road driving vehicles; 2) obtain average IVOC emission factors for on-road fleet based on tests in the tunnel; 3) retrieve average IVOC emission factors for gasoline- and diesel-fueled vehicles by regression analysis, taking advantage of a large number of vehicles (>30,000 per day) passing the tunnel; 4) compare the SOA formation potential of vehicle-emitted IVOCs to that of vehicle-emitted VOCs measured in the same campaign.

2. Methodology

2.1 Field sampling

Sampling campaign was concurrently conducted both at the inlet and at the outlet of the Zhujiang tunnel (23° 6’ N, 113° 14’ E), which is located in urban Guangzhou, South China (Fig. S1), on three weekdays (October 14th-16th, 2019) and two weekend days (October 13th and...
October 19th, 2019). Detailed description of the Zhujiang tunnel could be found in our previous studies (Liu et al., 2014; Zhang et al., 2016, 2017, 2018). IVOCs were collected by a sorption tube (Tenax TA/ Carbograph 5TD, Marks International Ltd, UK) using an automatic sampler (JEC921, Jectec Science and Technology, Co., Ltd, Beijing, China). A Teflon filter was installed before the tube to remove particles in the air flow. The sampling flow rate was set at 0.6 L min⁻¹ and hourly samples were collected from 5:00 am to 24:00 pm on each sampling day. In order to compare SOA productions from IVOCs and VOCs, hourly VOCs samples were collected on 13th October 2019 with stainless-steel canisters at a flow rate of 66.7 mL min⁻¹ using a Model 910 Pressurized Canister Sampler (Xonteck, Inc., CA, USA). 2-hour quartz filter samples were also collected by a high-volume PM₂.₅ sampler (Thermo Electron, Inc., USA) at the outlet and inlet sampling sites from 13th October to 19th October. Trace gases were measured by online analyzers (CO, Model 48i, Thermo Electron Inc., USA; NOx, Model 42i, Thermo Electron Inc., USA). A video camera was installed at the inlet to record the vehicle flow during the campaign. After sampling, the videotapes were used to count the passing vehicles and classify the vehicle types.

### 2.2 Laboratory analysis

Sampled sorption tubes were analyzed by a thermal desorption (TD) system (TD-100, Markes International Ltd, UK) coupled to a gas chromatography / mass selective detector (GC/MSD; Agilent, 7890 GC/5975 MSD, USA) with a capillary column (Agilent, HP-5MS, 30 m × 0.25 mm × 0.25 μm). Deuterated standards (C₁₂-d₂₆, C₁₆-d₃₄, C₂₀-d₄₂, naphthalene-d₈, acenaphthene-d₁₀ and phenanthrene-d₁₀) were injected into the sorption tubes to determine their recoveries before analysis. The sampled sorption tubes and field blanks were thermally desorbed at 320 °C.
for 20 min, and the desorbed compounds were carried by high purity helium into a cryogenic trap at -10 °C, and then the trap was rapidly heated to transfer them into the GC/MSD system. The initial temperature of GC oven was set at 65 °C, held for 2 min, then increased to 290 °C at 5 °C min\(^{-1}\) and kept at 290 °C for 20 min. The MSD was used in the SCAN mode with an electron impacting ionization at 70 eV. Individual speciated IVOCs were quantified with the calibration curves by using authentic standards. The total IVOCs mass was determined using the approach developed by Zhao et al. (2014, 2015, 2016) and the detailed description was provided in the supporting information (Text S1). Briefly, the total ion chromatogram (TIC) of IVOCs was divided into 11 bins based on the retention time of C\(_{12}-C_{22}\) n-alkanes. Each bin centered on the retention time of n-alkane. The start time and end time of the bin was determined by the average retention time of two successive n-alkanes. For example, the start time of Bin16 (B16) was calculated as the average retention time of n-C\(_{15}\) and n-C\(_{16}\), and the end time of B16 as the average retention time of n-C\(_{16}\) and n-C\(_{17}\). The IVOCs mass in each bin was quantified by the response factor of n-alkane in the same bin. The total IVOCs mass was the sum of IVOCs mass determined in 11 bins. The mass of unresolved complex mixtures of IVOCs (UCM-IVOCs) was determined by the difference between the total IVOCs and speciated IVOCs in each bin. The UCM-IVOCs were further classified into unspeciated branch alkanes (b-alkanes) and cyclic compounds (Zhao et al., 2014) (Text S1). The analysis of VOCs can be found elsewhere (Zhang et al., 2018). The POA emission was estimated as 1.2 times of organic carbon that measured in quartz filter samples (Zhao et al., 2015), which were analyzed by an OC/EC analyzer (DRI Model 2015, Nevada, USA) (Li et al., 2018).
2.3 Quality assurance and quality control (QA/QC)

Before their use for field sampling, sorption tubes were conditioned at 320 °C for 2 hours at oxygen-free nitrogen flow and then sealed at both ends with brass storage caps fitted with PTFE ferrules. About 15% of conditioned tubes were selected randomly to be analyzed in the same way as normal samples to check if any targeted species existed in the tubes. The batch of sorption tubes were certified as clean if speciated IVOCs were not found or presented in levels below the method detection limits (MDLs). Before and after sampling, the flow rates of samplers were calibrated by a soap-membrane flowmeter (Gilian Gilibrator-2, Sensidyne, USA). During the sampling, ten field blanks (five at the inlet and five at the outlet) were collected by installing a sorption tube onto the sampler each day but with pump off at both the inlet and the outlet. The speciated IVOCs were not detected or presented in levels below their MDLs in the blanks. MDLs for all the speciated IVOCs, including n-alkanes and polycyclic aromatic hydrocarbons (PAHs), were below 8 ng m⁻³, such as 5.8 ng m⁻³ for n-C₁₂, 5.9 ng m⁻³ for n-C₁₆ and 4.7 ng m⁻³ for n-C₂₂. To check if any breakthrough occurs during the sampling, prior to the field campaign two sorption tubes were connected in series to sample at the tunnel outlet station in the same way. IVOCs detected in the second tube only accounted for 2.6 ± 1.4% of the total in the two tubes, indicating no breakthrough during the sampling. To check the recoveries during thermo-desorption, selected sampled sorption tubes were analyzed twice by the TD-GC/MS system, and the desorption recoveries, calculated as the percentage of IVOCs in first analysis, were 96.7 ± 3.2% on average. Duplicated samples revealed less than 15% differences for all the speciated IVOCs.

2.4 Calculation of IVOCs emission factor
The vehicular EF of IVOCs can be calculated by following equation (Pierson et al., 1983; Zhang et al., 2016, 2017, 2018):

\[
EF = \frac{\Delta C \times V_{air} \times T \times A}{N \times l}
\]  

(1)

where EF (mg km\(^{-1}\) veh\(^{-1}\)) is the fleet-average emission factor of given species during the time interval T (1 h in this study); \(\Delta C\) (mg m\(^{-3}\)) is the inlet-outlet incremental concentration of IVOCs; \(V_{air}\) (m s\(^{-1}\)) is wind speed parallel to the tunnel measured by a 3-D sonic anemometer (Campbell, Inc.); A (m\(^2\)) is the cross-sectional area of the tunnel; N is the number of vehicles travelling through the tunnel during the time interval T; l (km) is the distance between the outlet and the inlet.

3. Results and discussions

3.1 Emission factors and compositions of IVOCs

Fig. 1 shows diurnal variations of vehicle numbers and vehicular IVOCs emission factors (EF\(_{IVOCs}\)) during the campaign. Traffic flow in the tunnel varied 571-2263 vehicles per hour during the campaign, and gasoline vehicles (GVs) dominated the vehicle fleets with a share of 76.3%, diesel vehicles (DVs) only accounted for 4.0%, and other types of vehicles, including liquefied petroleum gas vehicles (LPGVs) and electrical vehicles (EVs), had a percentage of 18.7% (Fig. S2). As LPGVs and EVs are considered to have no IVOCs emissions (Stewart et al., 2021), only GVs and DVs are responsible for the inlet-outlet incremental concentrations of IVOCs. Based on above equation (1), average EF\(_{IVOCs}\) for GVs and DVs in the vehicle fleets ranged from 13.29 ± 5.08 mg km\(^{-1}\) veh\(^{-1}\) to 21.40 ± 5.01 mg km\(^{-1}\) veh\(^{-1}\), with an average of 16.77 ± 0.89 mg km\(^{-1}\) veh\(^{-1}\) (Average ± 95% C.I.) (Fig. 1). The average EF\(_{IVOCs}\) for DVs and GVs could be further derived through linear regression as below (Ho et al., 2007; Kramer et al.,
$EF_{IVOCs} = EF_{DV} \times \alpha + EF_{GV} \times (1 - \alpha)$  \hspace{1cm} (2)

where $EF_{IVOCs}$ represents the fleet-average emission factor measured during a time interval; $EF_{DV}$ and $EF_{GV}$ are the average $EF_{IVOCs}$ for DVs and GVs, respectively; $\alpha$ is the fraction of DVs in the total IVOCs-emitting diesel and GV vehicles traveling through the tunnel. Based on the regression results (Fig. S3), the average $EF_{IVOCs}$ for DVs ($62.79 \pm 18.37$ mg km$^{-1}$veh$^{-1}$) was ~4.5 times that for GV vehicles ($13.95 \pm 1.13$ mg km$^{-1}$veh$^{-1}$).

The mileage-based EF can be converted to fuel-based EF with the fuel density and fuel efficiency (Text S2) (Zhang et al., 2016). Thus, we could obtain an average fuel-based $EF_{IVOCs}$ of $239.5 \pm 19.5$ mg kg$^{-1}$ for GV vehicles and $984.9 \pm 288.2$ mg kg$^{-1}$ for DV vehicles. Zhao et al. (2015, 2016) measured IVOCs emissions from DVs and GV vehicles in the US by the dynamometer tests. As shown in Fig. 2, the average $EF_{IVOCs}$ for DVs measured in our study was significantly lower than that for DVs without any diesel particulate filter (DPF) in the US, but over 4 times higher than that with DPF. It is worth noting that the $EF_{IVOCs}$ for DVs from this study was comparable to that for ships and non-road construction machineries (NRCMs) with diesel-fueled engines in China (Fig. 2) (Huang et al., 2018; Qi et al., 2019). As a matter of fact, China III or lower emission standard DVs accounted for ~40% of China’s total in-use DVs in 2019 (http://www.mee.gov.cn/), and like the non-road engines, they are not equipped with any after-treatment facilities. Although the after-treatment systems are installed in the China IV and China V DVs, their working performance might be not so satisfactory (Wu et al., 2017). This may explain why the DVs in this study had IVOCs-EFs comparable to non-road engines. The $EF_{IVOCs}$ for GV vehicles from this study fell into the ranges of that for GV vehicles in the US, but was at the
high-end of the tested values (Fig. 2). A recent study revealed a significantly lower EF\textsubscript{IVOCs} of 83.7 mg kg\textsuperscript{-1} for a China V gasoline vehicle (Tang et al., 2021), implying that upgrading the emission standard could help reduce emissions of IVOCs from GVs, as China IV and China III GVs still share a much larger portion than the China V and VI ones in the on-road fleets (http://www.mee.gov.cn/).

Fig. 3 shows the EFs and compositions of the vehicular IVOCs in each retention-time based bin (Table S1). Similar to previous studies (Zhao et al., 2015, 2016; Huang et al., 2018; Qi et al., 2019; Tang et al., 2021), the unspeciated cyclic compounds dominated the IVOCs, accounting for 59.07 ± 1.06%, followed by unspeciated b-alkanes (25.27 ± 0.75%) and speciated IVOCs (15.66 ± 0.60%). The speciated IVOCs consist of n-alkanes, b-alkanes and PAHs. Naphthalene dominated the quantified PAHs, accounting for 56.82 ± 1.21% of total PAHs emissions. The distribution of IVOCs in retention-time based bins presented a significant decreasing trend with bin numbers. Previous studies have reported that more than 50% of IVOCs concentrated in higher-volatility bins like B12, B13 and B14 in gasoline exhaust while much broader volatility distributions were found in diesel exhaust (Zhao et al., 2015, 2016; Tang et al., 2021). The IVOCs in B12 measured in this study was also the most abundant as the GVs previously tested in the US (Fig. S4). This was reasonable since the GVs dominated the vehicle fleets during our tunnel experiments (Fig. S2). As shown in Fig. S5, the IVOCs determined in each volatility bin well correlated with those in the volatility bins close to them, and the total IVOCs have stronger correlations with IVOCs in the higher-volatility bins like B12, B13 and B14. In addition, the n-alkanes, as displayed in Table S2, were found to be significantly correlated to the total IVOCs that determined in the same volatility bin except for
B20 and n-C20. The mass ratios of IVOCs in each bin to the n-alkane in the same bin ranges 9.0-15.8 (Table S2). As n-alkanes are more easily and routinely quantified, the relationships of IVOCs and n-alkanes in each volatility bin might be used to estimate total IVOCs from on-road vehicles. However, vehicles types should be taken into consideration when using these ratios, as the results obtained here were based on a fleet dominated by GVs.

3.2 Relationships of IVOCs with other species

Emissions of IVOCs from vehicles are often estimated by assuming a ratio of IVOCs to other species such as POA or NMHCs (Shrivastava et al., 2008; Pye et al., 2010; Gentner et al., 2012; Murphy et al., 2017; Wu et al., 2019). However, these ratios might be highly variable with fuel types, operation conditions and engine performance (Lu et al., 2018). As demonstrated in Fig. S6 (a) and (b), IVOCs correlated well with NOx (R = 0.63, p < 0.05) and CO (R = 0.58, p < 0.05), with an average IVOCs-to-NOx ratio of 0.039 ± 0.004 and an average IVOCs-to-CO ratio of 0.033 ± 0.015. The measured IVOCs-to-POA ratio was 3.35 ± 1.79 (Fig. S6 (c)), comparable to that of 3.0 ± 0.9 for GVs previously measured in dynamometer tests simulating arterial and freeway cycles, but much higher than that of 1.5 previously used for estimating vehicle emissions in models (Robinson et al., 2007; Hodzic et al., 2010). As shown in Fig. S6 (d), the average IVOCs-to-NMHCs ratio measured in this study was 0.36 ± 0.09, lower than that previously measured for diesel vehicle exhaust (0.6 ± 0.1) (Zhao et al., 2015), but higher than that previously measured for gasoline vehicle exhaust (< 0.2) (Zhao et al., 2016; Tang et al., 2021). It is worth noting that the IVOCs did not present significant correlations with POA or NMHCs from this study for on-road vehicle fleets (Fig. S6 (c) and (d)). This would cast uncertainty over the emission estimates of IVOCs based on their ratios to POA or NMHCs.
3.3 Estimated SOA production from IVOCs

SOA formation potentials of IVOCs from on-road vehicle fleet as measured in this tunnel study can be estimated as:

\[ \text{SOA}_{FP} = \sum EF_{i} \times Y_{i} \]  

(3)

where \( \text{SOA}_{FP} \) is the SOA formation potential from the gaseous precursors; \( EF_{i} \) represents the emission factor of precursor \( i \) and \( Y_{i} \) is the SOA yield of precursor \( i \) under high-NOx at OA concentration of 20 \( \mu g \) m\(^{-3}\) (Zhao et al., 2015; Huang et al., 2018; Qi et al., 2019; Tang et al., 2021) (Table S3). As shown in Fig. 4, the SOA formation potentials from vehicular VOCs and IVOCs totalled 8.24 ± 0.68 mg km\(^{-1}\). The SOA-to-POA ratio was 2.41 ± 1.45, which was comparable to that of GVs tested in China (1.8-4.4) (Tang et al., 2021), and that of GVs (3.6) (Zhao et al., 2016) and high-speed DVs (3.2 ± 1.7) without DPF in the US (Zhao et al., 2015). Our previous chamber studies simulating SOA formation from vehicles exhaust revealed the SOA-to-POA ratios of 2.0 for DVs and 3.8 for GVs when cruising at 40 km h\(^{-1}\) (Deng et al., 2020; Zhang et al., 2020), which is near the average driving speed of vehicles in the tunnel. Among the vehicle-emitted SOA precursors, similar to previous studies (Zhao et al., 2015, 2016; Huang et al., 2018; Qi et al., 2019; Tang et al., 2021), IVOCs produced significantly higher SOA (7.19 ± 0.62 mg km\(^{-1}\)), ~7 times that from traditional VOCs (1.04 ± 0.30 mg km\(^{-1}\)). Previous smog chamber studies found that SOA formed during photoaging of vehicle exhaust could not be explained by traditional VOCs especially for vehicles cruising at higher speeds (Robinson et al., 2007; Deng et al., 2020; Zhang et al., 2020). If this \( \text{SOA}_{\text{IVOC}} \)-to-\( \text{SOA}_{\text{VOC}} \) ratio of 7 from this study is used to re-estimate the SOA formation from exhaust for vehicles cruising at 40 km h\(^{-1}\) in our previous chamber studies (Deng et al., 2020; Zhang et al., 2020), the VOCs...
plus IVOCs precursors could explain 91% − 98% SOA formed for GVs and 31.2% − 48.2%
SOA formed for DVs. Zhao et al. (2015, 2016) reported significant higher \( \text{SOA}_{\text{IVOC-to-SOAVOC}} \) ratio for diesel vehicle exhaust than gasoline vehicle exhaust. Thus, \( \text{SOA}_{\text{IVOC-to-SOAVOC}} \) ratio of 7 obtained in a tunnel dominated by GVs would underestimate \( \text{SOA}_{\text{IVOCs}} \) from
DV, consistent with higher NMHCs to IVOCs ratios in gasoline exhaust than in diesel exhaust
(Zhao et al., 2015, 2016; Huang et al., 2018; Qi et al., 2019; Tang et al., 2021). Overall, the
observed vehicular IVOCs as SOA precursors can help achieve mass closure between predicted
and measured SOA.

4. Conclusions and implications

Organic aerosol (OA), primary or secondary, accounts for a large fraction of particle matters
(Zhang et al., 2007; Jimenez et al., 2009). On-road vehicles could be an important source of
OA especially in urban environment (Gentner et al., 2017). Similar to previous smog chamber
simulation results about SOA formed from photochemical aging of vehicle exhaust (Deng et
al., 2020; Zhang et al., 2020), our tunnel test also demonstrated that estimated SOA surpassed
the POA emission. In addition, IVOCs was found to dominate over traditional VOCs in SOA
formation potentials by a factor of \(~7\), implying that reducing vehicle-emitted IVOCs is of
greater importance to modulate SOA for further reducing fine particle pollution particularly in
urban areas. As for the ratios of IVOCs to other primary species, our tunnel tests for on-road
fleet revealed complex and different results when compared to that from previous chassis
dynamometer tests, implying that cautions should be taken when applying the ratios from
chassis dynamometer tests to estimate real-world traffic emissions, or applying the ratios in the
US to estimate the emissions in China or other regions. As IVOCs is not considered in normal
vehicle emission tests, more field works characterizing real-world vehicular emissions of IVOCs are needed to further constrain these ratios.

$EF_{IVOC}$ for the GV-dominated fleets from our tunnel test, or $EF_{IVOC}$ for GV$s$ derived from regression, was much higher than that from a recent chassis test for a China V gasoline vehicle (Tang et al., 2021), suggesting that stricter emission standards might help reduce emissions of IVOC$s$ from GV$s$. Meanwhile, the $EF_{IVOC}$ for on-road DV$s$ was comparable to that for non-road engines without any after-treatments (Huang et al., 2018; Qi et al., 2019), suggesting that facilitating the installation of after-treatment devices with stricter emission standards or improving the performance of existing after-treatment devices are crucial to lower IVOC emissions from DV$s$, which have much bigger $EF_{IVOC}$ than GV$s$.

Based on the regression-derived average $EF_{IVOC}$ for GV$s$ and DV$s$ and the camera-recorded fleet compositions, we could estimate that ~81% of IVOC$s$ by vehicles travelling through the tunnel were coming from GV$s$ and only ~19% were from DV$s$. This is reasonable since DV$s$ have bigger $EF_{IVOC}$ and however much lower proportions in the fleets. These percentages may underestimate the contribution to IVOC$s$ by on-road DV$s$ in regional or national scales since DV$s$ travel less in core urban areas due to traffic restriction rules in China. Differently, in an updated emission inventory of vehicular IVOC$s$ in China (Liu et al., 2017) based on $EF_{IVOC}$ tested in the US, emission of IVOC$s$ from DV$s$ (145.07 Gg) was about 2.6 times that from GV$s$ (55.30 Gg) in China in 2015. However, the ratio of DV-$EF_{IVOC}$ to GV-$EF_{IVOC}$ used in the study (Liu et al., 2017) on average was much higher than that of ~4.5 from this study for on-road vehicles. Using the $EF_{IVOC}$ from tests in the US might underestimate IVOC$s$ emissions from GV$s$ but overestimate that from DV$s$ in China. As an example, $EF_{IVOC}$ of 83.7 mg kg$^{-1}$ reported
very recently for a China V gasoline vehicle (Tang et al., 2021) was still much higher than the
maximum EF$_{IVOC}$ (47.15 mg kg$^{-1}$) they adopted for China V GVs, and the EF$_{IVOC}$ used for
China III and China IV DVs were however significantly larger than that from our tunnel tests
(Fig. 2) for on-road DVs (mostly China III and China IV) (Liu et al., 2017). In 2019 the gasoline
and diesel fuel consumptions in China were $1.20 \times 10^2$ Tg and $1.50 \times 10^2$ Tg, respectively
(http://www.mee.gov.cn/hjzl/sthjzl/ydyhjgl/). Since that gasoline is mostly used for on-road
vehicles while diesel may be used for both on-road and non-road engines, and that EF$_{IVOC}$ for
on-road diesels are comparable to the non-road diesel engines (Huang et al., 2018; Qi et al.,
2019), we could use the fuel-based EF$_{IVOC}$ converted from our study to roughly estimate IVOCs
from diesel and gasoline combustion. This way estimated emission of IVOCs from diesel
engines (147.74 Gg) was about 5 times that from gasoline engines (28.74 Gg) in China in 2019.
In comparison of previous study (Liu et al., 2017), this result implies large uncertainties or even
inconsistencies about China’s vehicular IVOC emission estimates. Moreover, as diesel vehicle
shares less than 10% among China’s motor vehicles and a substantial part of diesel fuel is
consumed by non-road engines, the diesel-related IVOCs may largely come overwhelmingly
from non-road engines instead of on-road DVs, signaling the increasingly important role of
non-road engines as sources of IVOCs with the progress in on-road vehicles emission control.
Data availability. The dataset for this paper is available upon request from the corresponding author (wangxm@gig.ac.cn).

Competing interests. The authors declare no competing financial interest.

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Figure 1. Diurnal variations of vehicle fleets and fleet-average $E_{IVOC}$ during the sampling period. Error bars represent 95% confidence intervals.

Figure 2. Comparison of the $E_{IVOC}$ measured in this study with that previously measured for fossil fuel combustion sources. The error bars in (a) represent 95% confidence interval. In (b), the boxes represent the 75th and 25th percentiles, the centerlines are the medians and squares are the averages. The whiskers represent 10th and 90th percentiles. SORMs refer to small off-road engines fueled with gasoline. NRCMs represent non-road construction machineries fueled with diesel.

Figure 3. The average emission factor of vehicular IVOCs in different bins measured during the campaign.

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