

1           **Oxygenated VOCs as significant but varied contributors**  
2                           **to VOC emissions from vehicles**

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23 **Abstract:**

24 Vehicular emission is an important source for volatile organic compounds (VOCs) in  
25 urban and downwind regions. In this study, we conducted a chassis dynamometer study  
26 to investigate VOC emissions from vehicles using gasoline, diesel, and liquefied  
27 petroleum gas (LPG) as fuel. Time-resolved VOC emissions from vehicles are  
28 chemically characterized by a proton-transfer-reaction time-of-flight mass  
29 spectrometry (PTR-ToF-MS) with high frequency. Our results show that emission  
30 factors of VOCs generally decrease with the improvement of emission standard for  
31 gasoline vehicles, whereas variations of emission factors for diesel vehicles with  
32 emission standards are more diverse. Mass spectra analysis of PTR-ToF-MS suggest  
33 that cold start significantly influence VOCs emission of gasoline vehicles, while the  
34 influences are less important for diesel vehicles. Large differences of VOC emissions  
35 between gasoline and diesel vehicles are observed with emission factors of most VOC  
36 species from diesel vehicles were higher than gasoline vehicles, especially for most  
37 oxygenated volatile organic compounds (OVOCs) and heavier aromatics. These results  
38 indicate quantification of heavier species by PTR-ToF-MS may be important in  
39 characterization of vehicular exhausts. Our results suggest that VOC pairs (e.g. C<sub>14</sub>  
40 aromatics/toluene ratio) could potentially provide good indicators for distinguishing  
41 emissions from gasoline and diesel vehicles. The fractions of OVOCs in total VOC  
42 emissions are determined by combining measurements of hydrocarbons from canisters  
43 and online observations of PTR-ToF-MS. We show that OVOCs contribute 9.4% ±  
44 5.6%7.7% ± 6.2% of gasoline vehicles of the total VOC emissions, while the fractions  
45 are significantly higher for diesel vehicles (52-71%)(40-77%), highlighting the  
46 importance to detect these OVOC species in diesel emissions. Our study demonstrated  
47 that the large number of OVOC species measured by PTR-ToF-MS are important in  
48 characterization of VOC emissions from vehicles.

49

## 50 1. Introduction

51 Volatile organic compounds (VOCs) are important trace components in the  
52 troposphere, as important precursors of ground-level ozone (Shao et al., 2009) and  
53 secondary organic aerosol (SOA) (Seinfeld and Pandis, 2006;Kansal, 2009;Ziemann  
54 and Atkinson, 2012). As the result, it is particularly important to identify emission  
55 sources of VOCs in the atmosphere. Vehicular emission is an important source of VOCs  
56 in cities around the world (Liu et al., 2008;Parrish et al., 2009), contributing  
57 approximately 25% to total VOC emissions in China (Ou et al., 2015;Wu et al.,  
58 2016;Sun et al., 2018). In order to control atmospheric pollution in urban and  
59 surrounding regions, it is necessary to understand source profiles and emission  
60 characteristics of VOCs from vehicles.

61 Emissions of VOCs from vehicles have been investigated extensively from  
62 tunnel studies (Cui et al., 2018;Zhang et al., 2018;Song et al., 2020), on-road mobile  
63 measurements (Li et al., 2017), and chassis dynamometer tests (Guo et al., 2011;Wang  
64 et al., 2013;Yang et al., 2018). Previous studies demonstrated that fuel types of vehicles  
65 strongly impact VOC emissions. Aromatics along with other hydrocarbons are known  
66 as compounds with high emissions in exhausts of gasoline vehicles (Wang et al.,  
67 2013;Ly et al., 2020). Some carbonyl compounds contribute significantly to emissions  
68 of diesel vehicles, at fractions much higher than gasoline vehicles (Tsai et al., 2012;Qiao  
69 et al., 2012;Yao et al., 2015;Mo et al., 2016). Moreover, there are still a large number  
70 of unidentifiable compounds in diesel vehicles (May et al., 2014). Furthermore, VOC  
71 emissions from vehicles significantly decreased in China due to stricter emission  
72 standards~~Furthermore, VOC emissions significantly decreased for stricter emission~~  
73 ~~standards~~ (Liu et al., 2017;Sha et al., 2021). In order to reduce emissions of most  
74 primary pollutants, more stringent emission standards and after-treatment devices have  
75 been implemented. The emission standard of China VI has already been implemented  
76 in July of 2019 in a few key cities in China and in July of 2021 nationwide. The emission  
77 limits for various air pollutants emitted by vehicles are significantly lower under the  
78 China VI emission standard (see details in the Supplement) (Wu et al., 2017). With the

79 continuous development of engine and exhaust after-treatment technologies, emission  
80 characteristics of VOCs from vehicles may change and need to be frequently updated.

81 Oxygenated volatile organic compounds (OVOCs) were found to be an important  
82 class of compounds group in vehicle exhausts, accounting for more than 50% of the  
83 total VOC emissions for diesel vehicles from both chassis dynamometer tests-(Schauer  
84 et al., 1999;Mo et al., 2016) and on-road mobile measurements (Yao et al., 2015).  
85 Traditionally, VOCs are collected in the canister or Tedlar bags, and then analyzed by  
86 gas chromatography-mass spectrometer/flame ionization detector (GC-MS/FID),  
87 mainly reporting emissions of hydrocarbons (Wang et al., 2017;Qi et al., 2019).  
88 Previous work usually collected 2,4-dinitrophenylhydrazine (DNPH) cartridges and  
89 analyzed using high-performance liquid chromatography (HPLC) for carbonyls  
90 (aldehydes and ketones), which are both time-consuming and prone to contaminations  
91 (Mo et al., 2016;Han et al., 2019).

92 The large variability of VOC emissions under different engine activities or  
93 driving conditions require characterization of vehicular emissions at higher time  
94 resolution. Proton-transfer-reaction mass spectrometry (PTR-MS) has been used in a  
95 number of studies for measurements of vehicle emissions. VOCs from vehicle exhausts  
96 under various driving and operational modes were measured by PTR-MS onboard a  
97 mobile laboratory (Zavala et al., 2006;Zavala et al., 2009). Drozd et al. (2016) used a  
98 PTR-MS to emphasize the importance of cold start for vehicles, concluding that VOC  
99 emissions during cold start were equal to a 200 miles distance of driving during hot  
100 stabilized condition. Proton-transfer-reaction time-of-flight mass spectrometry (PTR-  
101 ToF-MS) can provide more powerful detection of various VOCs, thanks to the  
102 measurements of whole mass spectra and high mass resolution (Cappellin et al.,  
103 2012;Yuan et al., 2017). More OVOC species could be quantified from the measured  
104 mass spectra based on parameterization methods for sensitivity of instrument  
105 (Sekimoto et al., 2017;Wu et al., 2020).

106 In this study, we applied a PTR-ToF-MS along with a suite of other instruments  
107 to measure VOCs emitted from gasoline, diesel, and liquefied petroleum gas (LPG)  
108 vehicles. We investigated emission factors from different fuel types and emission

109 standards for representative VOC species exhausted from these vehicles. We used the  
110 dataset to analyze contributions of various VOC groups to total VOC emissions in  
111 different types of vehicles.

## 112 **2. Materials and methods**

### 113 **2.1 Tested vehicles and the chassis dynamometer study methods**

114 In this study, we conducted chassis dynamometer measurements to investigate  
115 VOC emissions from vehicles using gasoline, diesel, LPG as fuel. All gasoline vehicles  
116 are light-duty-gasoline-vehicle (LDGV) with the emission standards from China I to  
117 China VI, whereas diesel vehicles can be classified into light-duty-diesel-truck (LDDT),  
118 middle-duty-diesel-truck (MDDT), heavy-duty-diesel-truck (HDDT), and bus  
119 associated with emission standards of China III to China V. In addition, the test vehicles  
120 using LPG are all taxis, which are under mandatory scrappage after 8 years of driving  
121 in China; as a result only China IV and China V for LPG vehicles were tested. [The](#)  
122 [fractions of gasoline and diesel vehicles with different emission standards in China are](#)  
123 [shown in Table S1 \(MEEPRC, 2019;Li et al., 2021\).](#) Among the 38 vehicles we tested,  
124 a fraction of vehicles was measured several times, with a total of 62 experiments  
125 measured. The detailed information for test vehicles is summarized in [Sect. 1 in the](#)  
126 [Supplement.](#) Table S24 and Table S32.

127 The short transient driving cycle (GB 18285-2018, Figure S1a), as one of the  
128 widely used test methods for vehicle emissions in China (Li et al., 2012;Wang et al.,  
129 2013), was used for measurements of gasoline vehicles and LDDT, each running for  
130 three to five times. The short transient driving cycle methods were initially adapted  
131 based on emission regulations of the Economic Commission for Europe (ECE) cycle  
132 (Yao et al., 2003), which is developed and used in European countries (Laurikko, 1995).  
133 The short transient driving cycle consist of four conditions, namely idling, acceleration,  
134 deceleration and uniform speed, as shown in Fig. S1. For the MDDT and HDDT, we  
135 customized a step-by-step test method, in which the vehicle accelerates to 20 km·h<sup>-1</sup>,  
136 40 km·h<sup>-1</sup> and 60 km·h<sup>-1</sup> in sequence after the engine activates, keeping at 20 km·h<sup>-1</sup>  
137 and 40 km·h<sup>-1</sup> for 2 minutes, and 60 km·h<sup>-1</sup> for 1 minute, respectively (Fig. S1) (Li et

138 al., 2021;Liu et al., 2021;Liao et al., 2021). In addition, the cold start was tested for a  
139 number of vehicles after a cold soak for more than 12 hours at ambient temperature  
140 (20-25 °C) before engine started. The measurements of cold start are compared to  
141 measurements of hot start after a ~10 minutes break for the vehicles after previous  
142 measurement. More details about cold start and hot start in this campaign can be found  
143 in Li et al. (2021).

144 A custom-built sampling and dilution system for vehicles combining online and  
145 offline sampling techniques was used in this study. As shown in Fig. S2, a portable  
146 emission measurement system (PEMS, SEMTECH-DS, Sensors. USA) was employed  
147 to measure emissions of CO, CO<sub>2</sub>, NO<sub>x</sub>, and total hydrocarbon (THC) directly from the  
148 tailpipe of vehicles. A custom-built dilution system (Li et al., 2021;Liao et al., 2021)  
149 was used for dilution of vehicular emissions, achieving dilution ratios of 10-100 for  
150 different vehicles. After dilution, CO<sub>2</sub> and CO were measured using a Li-840A  
151 CO<sub>2</sub>/H<sub>2</sub>O Gas Analyzer (Licor, Inc. USA) and a Thermo 48i-TLE analyzer (Thermo  
152 Fisher Scientific Inc. USA), respectively. Measurements of CO<sub>2</sub> before and after the  
153 dilution system was used to determine the dilution ratio for each test (see details in Fig.  
154 S3).

## 155 **2.2 VOC measurements using PTR-ToF-MS**

156 In this study, a Proton Transfer Reaction Quadrupole interface Time-of-Flight  
157 Mass Spectrometer (PTR-QiToF-MS) (Ionicon Analytik, Innsbruck, Austria) with  
158 H<sub>3</sub>O<sup>+</sup> chemistry was used to measure VOCs (Sulzer et al., 2014). The mass spectra of  
159 PTR-ToF-MS was recorded every 1 s as to capture characteristics of VOC species from  
160 vehicle exhausts in real-time. Background measurements of the instrument were  
161 performed using sampled air through a custom-built platinum catalytical converter  
162 heated to 365 °C for 30 s before vehicle starts in each test. The more detailed setting  
163 parameters for the instrument can be found elsewhere (Wu et al., 2020;Wang et al.,  
164 2020a;He et al., 2022). Data analysis of PTR-ToF-MS was performed using the Tofware  
165 software package (version 3.0.3, Tofwerk AG, Switzerland) (Stark et al., 2015).

166 A 23-component gas standard (Linde Spectra) was used for daily calibration of

167 PTR-ToF-MS during the campaign. VOC sensitivities from automatical calibrations  
168 indicated quite stable instrumental performance for most of the VOC species (Fig. S4).  
169 Another gas standard with 35-component VOCs (Apel Riemer Environmental Inc.) was  
170 used for calibrations during the later period of this campaign to include more VOC  
171 species in the calibration. The Liquid Calibration Unit (LCU, Ionicon Analytik,  
172 Innsbruck, Austria) was used to calibrate a total of 11 organic acids and nitrogen-  
173 containing species (Table S43). The limits of detection for calibrated VOC species are  
174 below 100 ppt for the 1-s measurement, except for ethanol (423 ppt) and formic acid  
175 (166 ppt). Additionally, the humidity dependence for a few VOC species in PTR-ToF-  
176 MS (Yuan et al., 2017;Koss et al., 2018) were corrected using humidity-dependence  
177 curves determined in the laboratory, as previously shown in Wu et al. (2020). To  
178 quantify the ion signals without calibration, we determine the sensitivities based on the  
179 kinetics of proton-transfer reactions of  $\text{H}_3\text{O}^+$  with VOCs (Cappellin et al.,  
180 2012;Sekimoto et al., 2017). The relationship between VOCs sensitivity and kinetic  
181 rate constants for the same instrument has been reported in Wu et al. (2020) and He et  
182 al. (2022). The corrected sensitivities as a function of kinetic rate constants for proton-  
183 transfer reactions of  $\text{H}_3\text{O}^+$  with VOCs during this campaign is shown in Fig. S5. The  
184 fitted line is used to determine sensitivities of uncalibrated species, and the uncertainty  
185 of the concentrations for uncalibrated species are determined to be around 50%.

### 186 **2.3 Other VOC measurements**

187 Whole air samples were collected using canisters after the dilution system for  
188 determination of hydrocarbons emitted from various vehicles. All the canisters were  
189 sent to the laboratory for analysis by an offline GC-MS/FID system, with a total 95  
190 hydrocarbons calibrated by Photochemical Assessment Monitoring Stations (PAMS)  
191 and TO-15 standard mixtures (Table S54). We compared emission factors from PTR-  
192 ToF-MS and the offline canister-GC-MS/FID (Fig. S6c-d), obtaining generally  
193 consistent results, considering the large variation of VOC emissions for driving  
194 conditions and the difficulty to control the fill time for canisters.~~Due to the difference~~  
195 ~~of sampling (e.g., times and dilution ratios), we compared emission factors from PTR-~~

~~ToF-MS and the offline canister GC-MS/FID, obtaining consistent results, except for gasoline vehicles with China I (Fig. S6c).~~

An instrument based on Hantzsch reaction-absorption method was used to measure formaldehyde (Zhu et al., 2020). Good agreement for formaldehyde between PTR-ToF-MS and the Hantzsch instrument was obtained (Fig. S6a). An iodide-adduct time-of-flight chemical ionization mass spectrometer (I<sup>-</sup> ToF-CIMS, Aerodyne Research, Inc.) (Wang et al., 2020c; Ye et al., 2021) was used to measure organic acids, hydrogen cyanide (HCN), and isocyanic acid (HNCO) from vehicles (Li et al., 2021). As shown in Fig. S6b, formic acid measured by PTR-ToF-MS and I<sup>-</sup> ToF-CIMS showed reasonable agreement.

## 2.4 Emission factors and emission ratios calculation

In this study, we determine emission factors of VOC species in two different approaches: the mileage-based emission factors ( $\text{mg}\cdot\text{km}^{-1}$ ) as the mass of these VOCs exhausted per kilometer driving of vehicles, and the fuel-based emission factors ( $\text{mg}\cdot\text{kg}_{\text{fuel}}^{-1}$ ) as the mass of VOCs per kilogram of fuel burned by the vehicles. In addition, emission ratios of VOCs to combustion tracers (usually CO) are widely applied in vehicle emissions in urban regions, as the result we determine emission ratios to CO in  $\text{ppb}\cdot\text{ppm}^{-1}$  as well. More details about the determination of emission factors and emission ratios can be found in Sect. ~~2~~ in the Supplement.

The average emission factors for various types of vehicles are determined from arithmetic means for different emission standards of vehicles. As for diesel vehicles, the average emission factors are obtained from the arithmetic means of LDDT, MDDT, HDDT, and bus. Besides, we also calculate emission factors and emission ratios from weighted means based on the fractions of gasoline and diesel vehicles with different emission standards in China (MEEPRC, 2019; Li et al., 2021) (~~S~~see Sect. ~~4~~~~2~~ in the Supplement for details). In order to evaluate the uncertainties of obtained emission factors, the average limit of detection for VOC species are used to estimate the limit of detection for the determined emission factors (more details can be found in Sect. ~~3~~~~2~~ in the Supplement).



## 225 **3. Results and discussions**

### 226 **3.1 Characteristics of the VOC emissions in the vehicles**

227 Time series of several aromatics and OVOC species measured by PTR-ToF-MS  
228 for a selected gasoline vehicle associated with emission standard of China I and a LDDT  
229 associated with China IV emission standard are shown in Fig. 1. Both tests started with  
230 cold engines for the two vehicles. Benzene and toluene are typical aromatic species  
231 emitted by vehicles~~species exhausted by vehicles~~. As shown in Fig. 1a, high  
232 concentrations of benzene and toluene exhausted by the gasoline vehicle were observed  
233 as the engine started. The concentrations of the two species continued to increase until  
234 ~2 min after the engine started, and then dropped rapidly before a minor increase during  
235 the acceleration condition. These observations are similar to the previous results from  
236 PTR-MS measurements in Drozd et al. (2016). Acetaldehyde and acetone are important  
237 OVOC species emitted from vehicles. They show similar temporal variations as  
238 benzene and toluene. However, concentrations of acetaldehyde and acetone were much  
239 lower than the two aromatics after engine started. Compared to the concentrations at  
240 engine start-up for the gasoline vehicle (the first cycle), concentrations of the VOCs are  
241 3.0 to 40 times lower during the gasoline vehicle running at hot stabilized condition  
242 (the third cycle). As shown in Fig. 1 for the diesel vehicle, enhanced emissions from  
243 cold start are minor, which is different from the gasoline vehicle. The concentration of  
244 these VOCs at engine start-up for the diesel vehicle are only 1.3 to 2.5 times higher  
245 than the periods as the diesel vehicle running at hot stabilized condition. It indicates  
246 that the impact of the engine start-up in diesel vehicles on emissions is much lower than  
247 gasoline vehicles. It might be a combined effect of cold engine and operation  
248 temperature of the after-treatment device (Gentner et al., 2017; George et al., 2015). In  
249 contrast to the gasoline vehicle, we observe higher concentrations of the two OVOC  
250 species than the two aromatics species from the diesel vehicle. These higher OVOC  
251 concentrations in diesel vehicle exhausts are in line with the observations of organic  
252 acids using the I-ToF-CIMS from the same campaign (Li et al., 2021).

253 Based on the high time-resolution measurements of PTR-ToF-MS, we

254 determined emission factors of various VOC species from different vehicles. Fig. 2  
255 shows the determined average mileage-based emission factors of benzene, toluene,  
256 acetaldehyde, and acetone for various types of vehicles (also tabulated in the  
257 Supplement table). In general, we observe a downward trend for emissions factors of  
258 gasoline vehicles from China I to China VI emission standards for the four  
259 representative VOC species. This is consistent with the results in previous studies with  
260 lower emissions for newer emission standards (Wang et al., 2017; Sha et al., 2021). In  
261 addition, the dependence of VOCs emission versus emission standard may also be  
262 attributed to the history of vehicle usage, i.e., the mileage traveled by the vehicles, as  
263 lower mileages of vehicles are usually associated with vehicle with newer emission  
264 standards. As shown in Fig. 3, we observe strong positive relationship between toluene  
265 emission factors and vehicle odometers for both gasoline and diesel vehicles, indicating  
266 the mileages of vehicles can significantly affect VOCs emission factors for vehicles  
267 tested in this study. Intestinally, the emission factors of the representative VOC species  
268 are highest for China II gasoline vehicles rather than China I vehicles, coincidence with  
269 largest mileage of the test vehicles. Emission factors of the four species for China VI  
270 vehicles are 12 to 25 times lower than emissions for China I vehicles, indicating that  
271 newer emission standards successfully reduced VOC emissions of gasoline vehicles.  
272 The decline of emission factors for the four species with newer emission standards for  
273 diesel vehicles are in the range of 1.1 to 7.4 times from China III to China V, compared  
274 to 4.5 to 5.4 times reduction from China III to China V for gasoline vehicles. Emission  
275 factors of benzene and toluene from diesel vehicles are in the range of 0.8 to 7.4 mg·km<sup>-</sup>  
276 <sup>1</sup> and 0.3 to 5.8 mg·km<sup>-1</sup>, which are comparable to emission factors from gasoline  
277 vehicles with China IV to China VI emission standards. This is different from  
278 observations of the two OVOC species (acetaldehyde and acetone), with much higher  
279 emission factors from diesel vehicles (8.0 to 27.9 mg·km<sup>-1</sup> for acetaldehyde and 0.8 to  
280 10.0 mg·km<sup>-1</sup> for acetone) than almost all gasoline vehicles (a maximum of 3.9 mg·km<sup>-</sup>  
281 <sup>1</sup> for acetaldehyde and a maximum of 3.2 mg·km<sup>-1</sup> for acetone). Higher emission factors  
282 from diesel vehicles are also observed for many other common OVOC species, as  
283 shown in Fig. 43. As the largest OVOCs emitted from gasoline vehicles (4.6 ± 5.1

284 mg·km<sup>-1</sup>), methanol is found to be the only common OVOC species, with lower  
285 emission factors from diesel vehicles than gasoline vehicles. The emission factor of  
286 other OVOCs (e.g. formaldehyde, acetone) from diesel vehicles are higher than  
287 gasoline vehicles, which is consistent with previous results (Gentner et al., 2013). The  
288 high emissions of OVOCs from diesel vehicles may be related to combustion processes  
289 in diesel vehicles, with more excess air (i.e., under overall fuel-lean conditions) into  
290 combustion cylinder resulting in higher oxygen contents and more oxidation processes  
291 during fuel combustion (Pang et al., 2008; Qiao et al., 2012; Gentner et al., 2017). Finally,  
292 the determined emission factors of the four VOC species from LPG vehicles are much  
293 lower than both gasoline and diesel vehicles.

### 294 ~~3.2 Analysis VOCs of PTR-ToF-MS mass spectra~~ Analysis of PTR- 295 ToF-MS mass spectra to evaluate VOCs speciation

296 In addition to typical VOC species shown above, PTR-ToF-MS detected  
297 abundant signals for a large number of ions. The determined average mileage-based  
298 emission factors for all detected VOC species are shown as mass spectra in Fig. 4. VOC  
299 species measured by PTR-ToF-MS were divided into groups according to chemical  
300 formula, namely hydrocarbon species only containing C and H atoms (C<sub>x</sub>H<sub>y</sub>), OVOCs  
301 (C<sub>x</sub>H<sub>y</sub>O<sub>z</sub>), species containing nitrogen and/or sulfur atoms (N/S-containing), and some  
302 other ions (others). We observe similar mass spectra of emission factors for gasoline  
303 vehicles with different emission standards (Fig. S7). Highest emission factors from  
304 gasoline vehicles (Fig. 54a) are detected as hydrocarbons, including C<sub>6</sub> to C<sub>10</sub> aromatics.  
305 A few OVOC species, namely methanol, ethanol, formaldehyde, acetaldehyde and  
306 acetone, are also observed as the largest emissions. In contrast to gasoline vehicles, the  
307 largest emissions from diesel vehicles were attributed to a few low-molecular-weight  
308 OVOC species, including formaldehyde, acetaldehyde, formic acid, and acetic acid,  
309 followed by a large number of hydrocarbon species. Comparison between the mass  
310 spectra of gasoline and diesel vehicle emissions suggest that emissions from diesel  
311 vehicles are more evenly distributed among different VOC species, as reflected by 50  
312 and 140 species contributing more than 1% of the total emissions for gasoline and diesel

313 vehicles, respectively. As shown in Fig. 53b, many hydrocarbon ions in the range of  
314  $m/z$  150-200 still account for significant fractions of emissions from diesel vehicles,  
315 whereas only one species in this  $m/z$  range contribute more than 1% of emissions from  
316 gasoline vehicles. These results demonstrate that diesel vehicles emit more heavier  
317 hydrocarbons than those from gasoline vehicles, which is consistent with observations  
318 in previous studies (Gentner et al., 2012; Erickson et al., 2014). It should be noted that  
319 the signals of  $C_{16}H_{22}O_4H$  ( $m/z=279$ ) were higher during the tests based on determined  
320 emission factors. However, we suspect that it may be emitted artifacts from the  
321 sampling or dilution system as it mainly showed higher signals in the latter period of  
322 each test when sampling materials absorb more heat from vehicle exhausts (Fig. S8),  
323 and thus it is not included in Fig. 53 (details in the Sect. 32 in the Supplement).

324 The scatterplot of carbon oxidation states ( $\overline{OS}_C$ ) as a function of carbon number  
325 ( $n_C$ ) provides a framework for describing bulk chemical properties of organics (Kroll  
326 et al., 2011). The details of  $\overline{OS}_C$  calculation is included in Sect. 43 in the Supplement.  
327 The results from gasoline and diesel vehicles are compared in Fig. 65 (LPG vehicles  
328 are shown in Fig. S9). It is apparent that ions with carbon oxidation states between -2.0  
329 to 0 comprise main emissions for each carbon number for both gasoline and diesel  
330 vehicles. It is interesting to observe that averaged  $\overline{OS}_C$  for  $n_C > 6$  increase as the carbon  
331 number decrease for both gasoline and diesel vehicles, whereas the opposite trends are  
332 observed for  $n_C < 5$ . The averaged  $\overline{OS}_C$  in diesel vehicles for  $n_C$  between 1 and 5 are  
333 significantly higher than those in gasoline vehicles, as the result of high emissions of  
334  $C_2$  to  $C_5$  low-molecular-weight OVOCs. Fig. 65c further shows that emission factors of  
335 most VOC species from diesel vehicles were higher than gasoline vehicles, except a  
336 number of species occupying in the right-bottom corner of the two-dimensional space.

337 The determined mass spectra of PTR-ToF-MS in terms of emission factor for  
338 different types of vehicles can be used to explore the dependence of various VOC  
339 emissions to different factors. Fig. 76a-b shows scatterplots of the average mileage-  
340 based emission factors of VOCs between cold start and hot start for gasoline and diesel  
341 vehicles, respectively. We observe strong correlation between emission factors from  
342 cold start and hot start tests ( $R=0.99$  and  $0.92$ ) and generally consistent ratios between

343 cold start and hot start for different types of VOC species for both gasoline and diesel  
344 vehicles, indicating that variation behaviors are similar for different species and thus  
345 chemical compositions of VOC emissions are comparable between different start  
346 conditions. As cold start emissions are richer in unburned fuel than other hot-running  
347 conditions, the observation in Fig. 7a-b also infer that unburned fuel are the major  
348 contributor for vehicle exhaust emissions, which has been previously shown in  
349 California, US (Gentner et al., 2013). It is obvious that emission factors of VOCs  
350 during cold start are significantly higher than those during hot start for gasoline  
351 vehicles (slope=0.40), whereas similar emissions factors between cold start and hot  
352 start are derived for diesel vehicles (slope=0.84). These results suggest that gasoline  
353 vehicles are more significantly influenced by cold start, as the result of compositions  
354 in gasoline fuel are more volatile than diesel fuel (US NRC, 1996). We further explore  
355 the effects of emission standards to VOCs emission factors by comparing determined  
356 emission factors between China I and China V for gasoline vehicle (Fig. 7c, also see  
357 China III versus China V and China V versus China VI in Fig. S10) and between China  
358 III and China V for LDDT (Fig. 7d, also see China III versus China V for MDDT and  
359 HDDT in Fig. S10). Fig. 7c-d show that the chemical compositions of VOC emissions  
360 are comparable between different emission standards for both gasoline and diesel  
361 vehicles (R=0.98 and 0.89), indicating after-treatment devices may not affect the  
362 relative fractions of VOC components. Furthermore, C-comparison of both gasoline  
363 and diesel vehicles demonstrate newer emission standards successfully decreased  
364 VOC emissions. Based on the derived slopes, we obtain VOCs emission factors  
365 reduced by a factor of 10 for gasoline vehicles from China I to China V (a factor of 5  
366 reduction from China III to China V and a factor of 2.5 reduction for China V to China  
367 VI), and a factor of 2 reduction for LDDT from China III to China V (a factor of 1.5  
368 and 8 reduction for MDDT and HDDT from China III to China V). The reduction ratio  
369 for gasoline vehicles from China I to China V are generally similar for most VOC  
370 species, except that some OVOC species with smaller reduction ratios. The reduction  
371 ratios for LDDT vehicles from China III to China V show large variability for different  
372 species. The lowest reduction ratios (a factor of ~2) are observed for the low-molecular

373 weight OVOC species associated with largest emissions, while the reduction ratios for  
374 hydrocarbons and higher-molecular weight OVOCs are in the range of a factor of 10-  
375 100. These results indicate the after-treatment device for diesel vehicles [\(see Sect. 1 in](#)  
376 [the Supplement for details.\)](#) may effectively reduce emissions of some heavier VOC  
377 species, though the after-treatment devices do not aim for VOCs control (Gentner et  
378 al., 2017).

### 379 **3.3 Non-target analysis for comparison between gasoline and diesel** 380 **vehicles**

381 As shown in the previous section, the analysis of PTR-ToF-MS mass spectra  
382 provide rich information on understanding the influences of VOC emissions from  
383 vehicles. This detailed information provided by the PTR-ToF-MS also offer an  
384 opportunity to systematically compare emissions between gasoline and diesel vehicles.  
385 The scatterplot of the determined average emission factors of various VOC species  
386 between gasoline and diesel vehicles is shown in Fig. [87](#). Large difference of VOC  
387 compositions emitted from gasoline and diesel vehicles are observed, as indicated by  
388 the low correlation of the data points ( $R=0.24$ ). A limited number of VOC species,  
389 including C<sub>6</sub>-C<sub>10</sub> aromatics and some N/S-containing species (e.g. C<sub>7</sub>H<sub>5</sub>N) are  
390 associated with higher emission factors from gasoline vehicles, whereas the obtained  
391 emission factors of most VOC species emitted from diesel vehicles are higher,  
392 especially most OVOC species. For example, formic acid is found to be one of the  
393 most significant emission species in diesel vehicles, with emission factors three orders  
394 of magnitude higher than that of gasoline vehicles. In addition, emission factors of  
395 HCN from gasoline vehicles are similar to those from diesel vehicles. These results  
396 are consistent with the measurements using the I ToF-CIMS from the same campaign,  
397 as shown in Li et al. (2021).

398 The scatterplot shown in Fig. [87](#) can also be expressed in terms of the determined  
399 fuel-based emission factors between gasoline and diesel vehicles (Fig. [S11a14](#)).  
400 Generally, similar variability is obtained except the determined slope of the data points,  
401 with higher slopes determined from the scatterplot based on fuel-based emission factor

402 (0.19 versus 0.15). The emission ratios to CO between gasoline and diesel vehicles  
403 (Fig. S11b) show similar results. Furthermore, theThe difference between the slopes  
404 reflects the different average mileage for the same weight of fuel between gasoline  
405 ( $9.7 \text{ km} \cdot \text{kg}_{\text{fuel}}^{-1}$ ) and diesel vehicles ( $7.1 \text{ km} \cdot \text{kg}_{\text{fuel}}^{-1}$ ), as demonstrated for emission  
406 factors of  $\text{CO}_2$  in Table S65.

407 Comparing gasoline and diesel vehicles.~~From the comparison gasoline and diesel~~  
408 ~~vehicles,~~ we can also observe profound differences in relative changes of emission  
409 factors for analogous compounds series. The emission factors of  $\text{C}_6$ - $\text{C}_{10}$  aromatics are  
410 apparently higher for gasoline vehicles than diesel vehicles, whereas emission factors  
411 for larger aromatics ( $n_C > 11$ ) from diesel vehicles start to exceed gasoline vehicles. This  
412 interesting behavior is the result of different variations of emission factors for gasoline  
413 and diesel vehicles as carbon number increases. This may be attributed to the  
414 differences of chemical compositions of gasoline and diesel fuel, such as higher  
415 fractions of polycyclic aromatic hydrocarbons (PAHs) in the diesel fuel (Yue et al.,  
416 2015;Gentner et al., 2017). As shown in Fig. 98, emission factors of aromatics from  
417 gasoline vehicles start to rapidly decrease at  $n_C = 10$  (a factor of 5 for each additional  
418 carbon for  $\text{C}_{10}$ - $\text{C}_{15}$ ), while the emission factors of aromatic for diesel vehicles  
419 demonstrate a relatively flat pattern between  $\text{C}_6$  and  $\text{C}_{15}$ , only with significantly  
420 decrease for  $n_C > 15$ . Based on Fig. 98, we determine that emissions of aromatics with  
421  $n_C \geq 10$  in gasoline and diesel vehicles are account for 14% and 63% of total aromatic  
422 emissions, again suggest the importance of heavier aromatics in emissions from diesel  
423 vehicles. It also highlights that quantification of these heavier species by PTR-ToF-MS  
424 may be important in characterization of vehicular exhausts, especially diesel vehicles.

425 In addition to aromatics, the relative changes of emission factors for carbonyls  
426 with carbon number are apparently different between gasoline and diesel vehicles (Fig.  
427 87 and Fig. 98b). Emission factors of carbonyls tend to decrease as carbon number  
428 increase for both gasoline and diesel vehicles. The decrease magnitudes are observed  
429 to be comparable from  $\text{C}_1$ - $\text{C}_6$  carbonyls for gasoline (97.6%) and diesel vehicles  
430 (97.4%). However, as  $n_C > 6$ , the decrease of carbonyl emissions factors for diesel  
431 vehicles become smaller, result in larger emissions factors than gasoline vehicles for



432 this range of carbon number.

433 The above discussions demonstrate that emission characteristics of aromatics and  
434 OVOCs are significantly different between gasoline and diesel vehicles. As the result,  
435 the ratios of VOC pairs can be identified to distinguish emissions of gasoline and diesel  
436 vehicles. Fig. 910 shows the scatterplots of four representative VOCs (benzene, C<sub>14</sub>  
437 aromatics, formaldehyde, and acetaldehyde) versus toluene based on the determined  
438 emission factors. The data points for each VOCs pair clearly show distinct separation  
439 between gasoline vehicles and diesel vehicles, with apparently higher slopes for diesel  
440 vehicles than gasoline vehicles, as the result of much larger emission factors of toluene  
441 from gasoline vehicles and lower emission factors of the four representative VOCs  
442 from diesel vehicles. The benzene/toluene ratio in gasoline and diesel vehicle are  
443 determined as 0.48 and 1.24 mg·mg<sup>-1</sup> (corresponding to 0.57 and 1.46 ppb·ppb<sup>-1</sup> that  
444 are more widely used in ambient studies). The difference of benzene/toluene ratio  
445 between gasoline and diesel vehicles has been reported in previous studies, and our  
446 results are generally consistent with these previous results (Chan et al., 2002; Barletta  
447 et al., 2005; Qiao et al., 2012; Kumar et al., 2020). Compared to benzene/toluene ratio,  
448 the difference of C<sub>14</sub> aromatics/toluene ratio between gasoline and diesel vehicles are  
449 more substantial (a factor of 3800). The remarkable larger emission factors of C<sub>14</sub>  
450 aromatics from diesel vehicles suggest that diesel vehicles can be a significant or even  
451 predominated source for higher molecular aromatics. The enormous difference of C<sub>14</sub>  
452 aromatics/toluene ratio (and also other higher aromatics/toluene) between gasoline  
453 and diesel vehicles indicate these ratios could potentially provide good indicators for  
454 separation of gasoline and diesel vehicles in ambient or tunnel studies ([see discussion](#)  
455 [in Sect. 5 in the Supplement for details about the feasibility of the ratio using in](#)  
456 [ambient air](#)). Similar discrepancies are observed for formaldehyde/toluene and  
457 acetaldehyde/toluene ratios between gasoline and diesel vehicles. These ratios may  
458 not be able to be used as indicators for distinguish gasoline and diesel vehicles in  
459 ambient studies, since secondary sources may complicate the observed ratios in  
460 ambient air. However, these results strongly suggest that diesel vehicles can be  
461 important in emissions of these OVOC species, though the number of diesel vehicles



462 are smaller than gasoline vehicles in many countries, e.g. China and U.S (Wallington  
463 et al., 2013; Yao et al., 2015; Huang et al., 2021).

### 464 3.4 OVOC fractions in VOC emissions

465 Emission factors of various VOC species measured by PTR-ToF-MS from  
466 different vehicles are summarized in Fig. 1110. As shown in Fig. 1110a, the determined  
467 average mileage-based emission factors of total VOC ions from diesel vehicles were  
468 much higher than gasoline and LPG vehicles. Fig. 1110b-d quantified the proportions  
469 of different categories of ions measured by PTR-ToF-MS. The determined average  
470 mileage-based emission factors of  $C_xH_y$  accounted for the largest fraction in gasoline  
471 vehicles ( $84\% \pm 5.9\%$ ), and lower fractions in diesel ( $47\% \pm 16\%$ ) and LPG vehicles  
472 ( $32\% \pm 0.7\%$ ). OVOCs account for larger fractions in diesel ( $49\% \pm 16\%$ ) and LPG  
473 vehicles ( $58\% \pm 3.7\%$ ), while they only account for  $13\% \pm 6.1\%$  of emissions from  
474 gasoline vehicles. The fractions of different OVOC groups generally demonstrate a  
475 downward trend from  $C_xH_yO_1$  to  $C_xH_yO_{\geq 3}$ , and OVOCs with more than two oxygen  
476 atoms only occupy small percentages (0-7%) in vehicle exhausts, indicating low  
477 emissions of these species.

478 Combined with measurements of other VOCs from canisters measured by GC-  
479 MS/FID, the fractions of OVOCs in total VOC emissions ~~can be determined~~  
480 ~~determined~~ for different vehicles (details in Sect. 64 in the Supplement) (Fig. 1211).  
481 OVOCs account for ~~9.4% ± 5.6%~~~~7.7% ± 6.2%~~ of total VOC emissions for gasoline  
482 vehicles. The OVOC fractions for gasoline vehicles are generally comparable for  
483 different emission standards and cold/hot start, except somewhat higher fractions for  
484 China VI from hot start (Fig. S12). The OVOC fractions obtained in this study for  
485 gasoline vehicles are generally consistent with previous results (Cao et al., 2016; Wang  
486 et al., 2020b) (Fig. 1211). Among these studies, the OVOC fractions determined for  
487 gasoline with 10% ethanol (E10) (Roy et al., 2016) ( $22\% \pm 11\%$ ) are apparently higher.  
488 The fractions of OVOCs in total VOC emissions for diesel vehicles are ~~71% ± 20%,~~  
489 ~~65% ± 22%, 52% ± 18%, and 56% ± 26%~~~~77% ± 15%, 68% ± 15%, 73% ± 14%, and~~  
490 ~~40% ± 10%~~ for LDDT, MDDT, HDDT, and bus, respectively. The variations of OVOC

491 fractions with emission standards are observed to be mixed among different types of  
492 diesel vehicles (Fig. S12). The OVOC fractions from diesel vehicles are obviously  
493 higher than those in gasoline vehicles, indicating the importance of OVOCs in VOC  
494 emissions for diesel vehicles. Compared to previous studies (Tsai et al., 2012; Qiao et  
495 al., 2012; Cao et al., 2016; Mo et al., 2016), determined OVOC fractions for diesel  
496 vehicles in this study are higher. If only considering carbonyls among various types of  
497 OVOCs measured by PTR-ToF-MS, the OVOC fractions determined in this study are  
498 more comparable with previous studies (Fig. ~~124~~), since most previous studies only  
499 detected carbonyls among various types of OVOCs. Finally, we determine that OVOCs  
500 account for ~~41% ± 10%~~  $41% \pm 8.6%$  of total VOC emissions for LPG vehicles, which  
501 is also higher than in one previous study (Wang et al., 2020b) with only carbonyls and  
502 a few esters/alcohols included. These results stress that the large number of OVOCs  
503 measured by PTR-ToF-MS are important in characterization of VOC emissions from  
504 vehicles. It should be noted that the OVOC fractions obtained here only reflect exhaust  
505 emissions. Evaporative emissions may be associated with different fractions of various  
506 VOC groups, which may be more related to fuel compositions (Rubin et al.,  
507 2006; Huang et al., 2021).

#### 508 **4. Conclusions**

509 In this work, we conducted a chassis dynamometer study to measure VOC  
510 emissions from gasoline, diesel, and LPG vehicles using PTR-ToF-MS along with other  
511 offline and online measurement techniques. Using this dataset, we provide emission  
512 factors of many VOCs from these three different types of vehicles associated with  
513 various emission standards in China. Our results show that emission factors of VOCs  
514 generally decrease with the increased stringency of emission standards for gasoline  
515 vehicles, whereas variations of emission factors for diesel vehicles with emission  
516 standards are more diverse. Mass spectra analysis of PTR-ToF-MS suggest that cold  
517 start significantly influence VOCs emission of gasoline vehicles, while the influences  
518 are smaller for diesel vehicles.

519 We observe large differences of VOC emissions between gasoline and diesel

520 vehicles based on PTR-ToF-MS measurements. Emission factors of most VOC species  
521 from diesel vehicles were higher than gasoline vehicles, especially for most OVOCs  
522 and heavier aromatics. The ~~substantially larger~~ ~~substantial larger~~ emission factors of  
523 some OVOCs emission factors for diesel vehicles indicate potentially dominant  
524 emissions of these species from diesel vehicles among vehicular emissions. Our results  
525 suggest that VOC pairs (e.g. C<sub>14</sub> aromatics/toluene ratio) could potentially provide good  
526 indicators for distinguishing emissions between gasoline and diesel vehicles.

527 Based on measurements of PTR-ToF-MS, C<sub>x</sub>H<sub>y</sub> ions account for the largest  
528 fraction in gasoline vehicles (84% ± 5.9%), whereas OVOC ions are the largest  
529 contributor in the mass spectra of emissions from diesel (49% ± 16%) and LPG vehicles  
530 (58% ± 3.7%). In the end, the fractions of OVOCs in total VOC emissions are  
531 determined by combining hydrocarbons measurements from canister results and online  
532 measurements of PTR-ToF-MS. We show that OVOCs contribute ~~9.4% ± 5.6%~~ ~~7.7% ±~~  
533 ~~6.2%~~ of gasoline vehicles of the total VOC emissions, while the fractions are  
534 significantly higher for diesel vehicles (~~52-71%~~ ~~(40-77%)~~), highlighting the importance  
535 to detect these OVOC species in diesel emissions.

536 This study shows significant contributions of OVOCs in VOC emissions from  
537 various vehicles, especially diesel vehicles. As a consequence, vehicular emissions may  
538 account for considerable proportions for primary emissions of these OVOCs in urban  
539 regions. Emissions of many OVOC species are currently not fully represented in  
540 emission inventories of VOCs, which may in turn affect the prediction ability of air  
541 quality models in urban regions. In this study, OVOC species are mainly quantified  
542 from PTR-ToF-MS measurements by taking into account all signals in the mass spectra,  
543 which stress that the large number of OVOC species measured by PTR-ToF-MS are  
544 important in characterization of VOC emissions from vehicles.

## 545 **Data availability**

546 Data are available from the authors upon request.

## 547 **Author contribution**

548 BY designed the research. ZBY, ~~JYZ~~, BY, QES organized vehicle test

549 measurements. SHW, CHW, CMW, TGL, JPQ, QES, and MMZ contributed to data  
550 collection. SHW performed the data analysis, with contributions from TGL, XJH, YBH,  
551 XBL, and QES. SHW and BY prepared the manuscript with contributions from other  
552 authors. All the authors reviewed the manuscript.

### 553 **Competing interests**

554 The authors declare that they have no known competing financial interests or  
555 personal relationships that could have appeared to influence the work reported in this  
556 paper.

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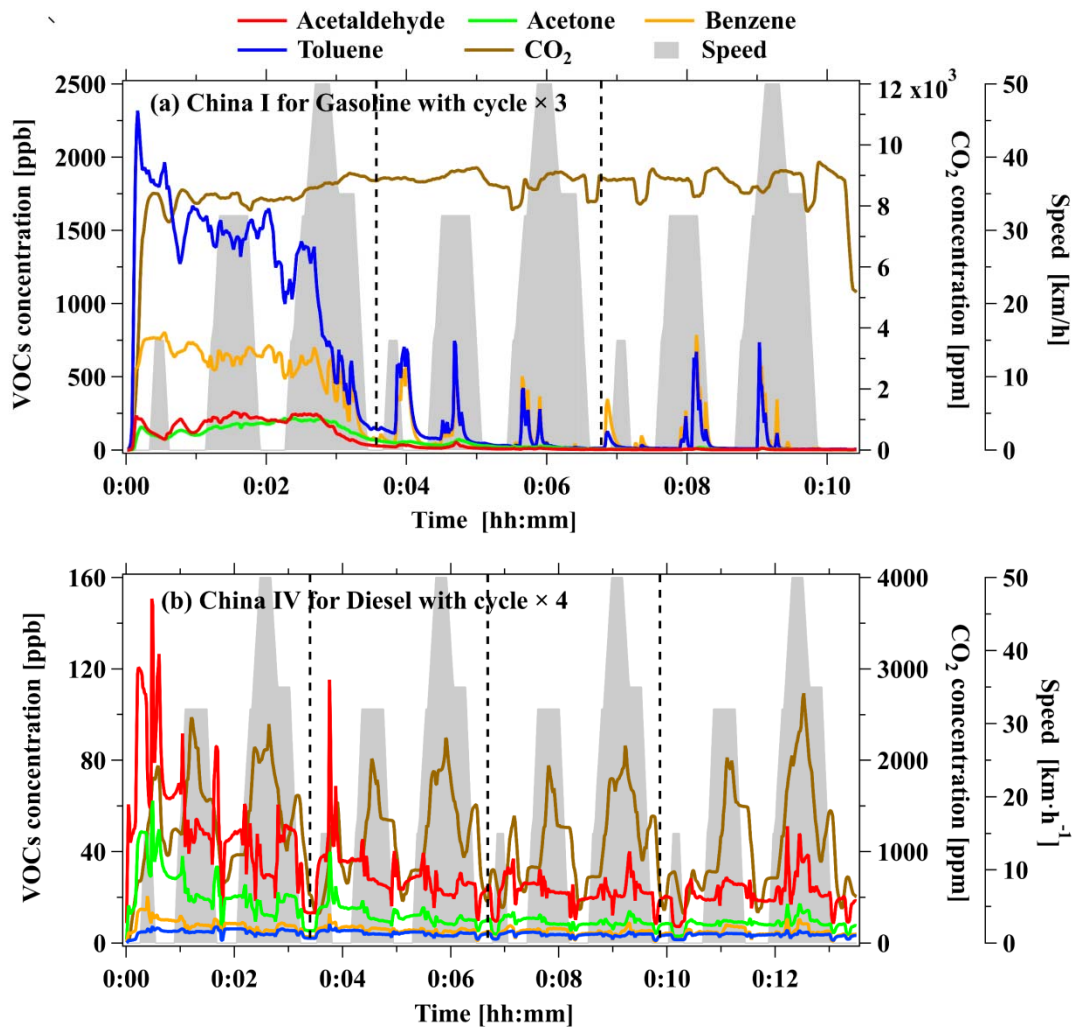
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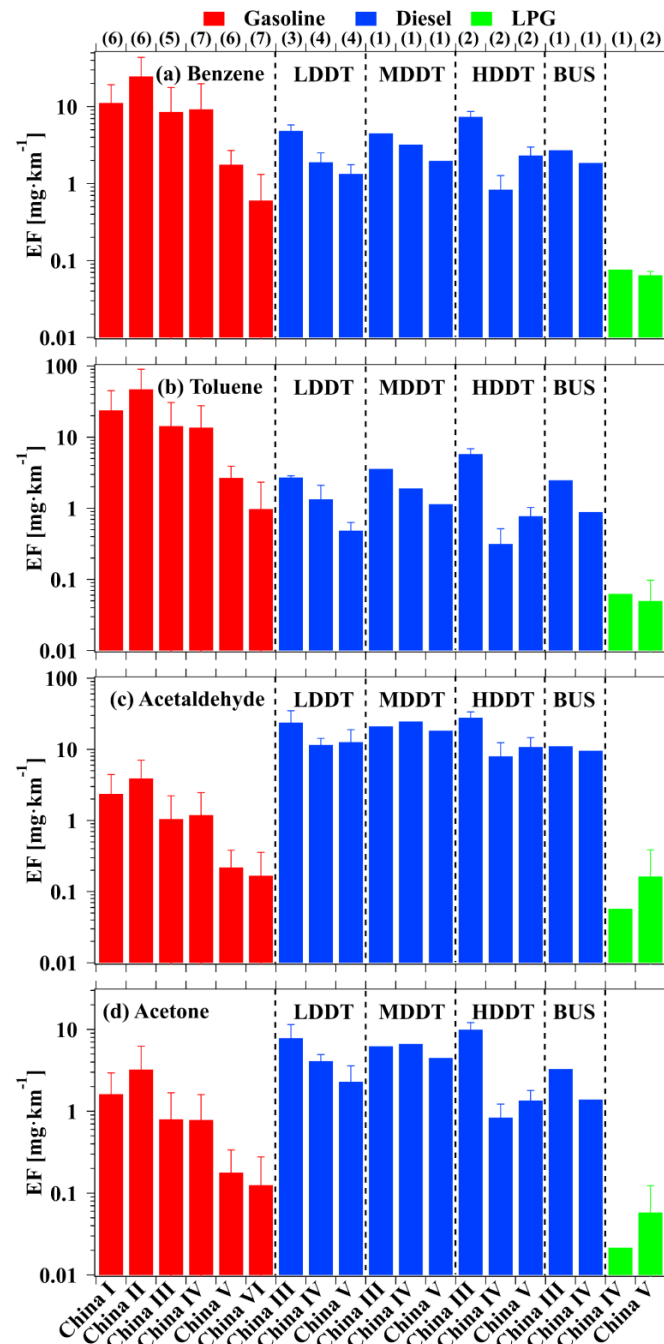
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859 **Figure 1.** Real-time concentrations of acetaldehyde, acetone, benzene, toluene, and  
 860 CO<sub>2</sub> for (a) a gasoline vehicle with emission standard of China I and (b) a light-duty  
 861 diesel vehicle (LDDV) with emission standard of China IV. The two vehicles were both  
 862 cold started. The gray shadows represent the speed of the vehicles on the chassis  
 863 dynamometer.

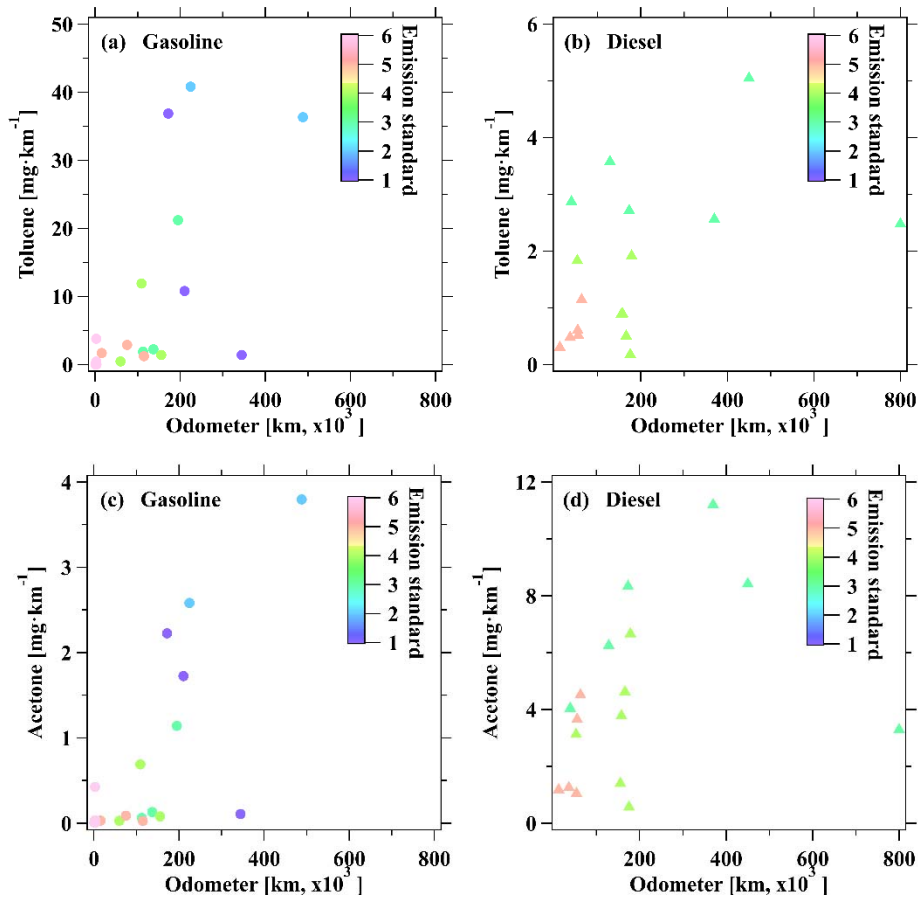
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866 **Figure 2.** The determined average mileage-based emission factors ( $\text{mg}\cdot\text{km}^{-1}$ ) for (a)  
 867 benzene, (b) toluene, (c) acetaldehyde, and (d) acetone for vehicles with different  
 868 emission standards. The numbers above the top axis represent the number of all  
 869 experiments (including multiple measurements for individual test vehicle) for each  
 870 emission standard. LDDT, MDDT, HDDT, and BUS represent light-duty-diesel-truck,  
 871 middle-duty-diesel-truck, heavy-duty-diesel-truck, and bus, respectively. Error bars  
 872 represent standard deviations of emission factors for the specific emission standard.

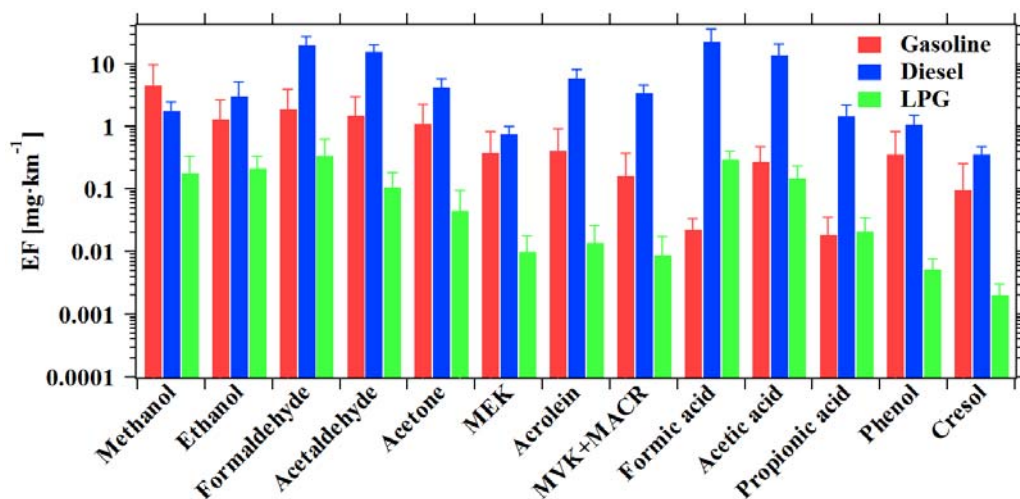
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875 **Figure 3.** Scatterplot of the emission factor of toluene in (a) gasoline and (b) diesel  
 876 vehicles, and acetone in (c) gasoline and (d) diesel vehicles during the hot start based  
 877 on the odometer for each vehicle.

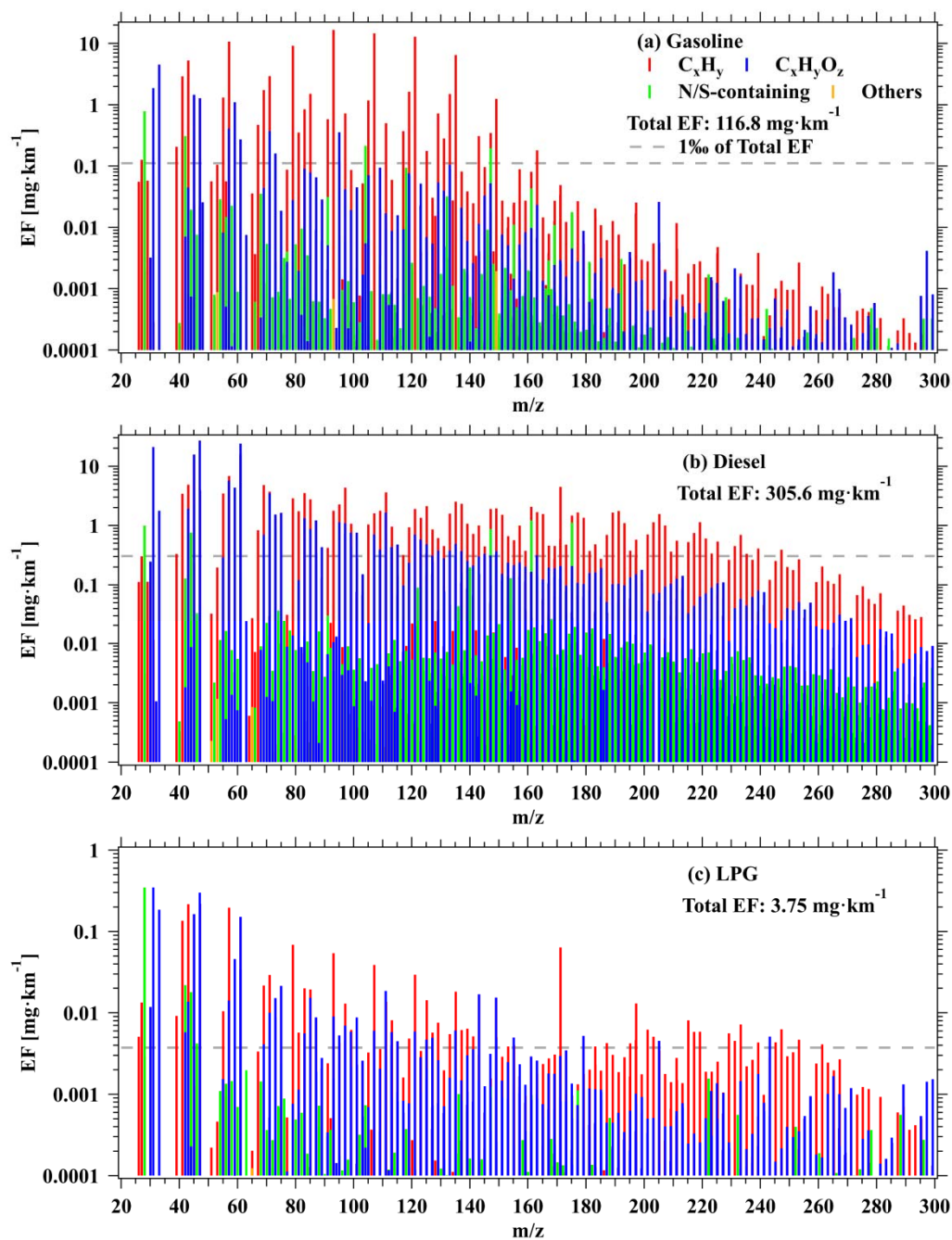
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880 **Figure 43.** The determined emission factors of representative OVOC species from  
 881 different types of vehicles. Error bars represent standard deviations of the emission  
 882 factors for the VOCs.

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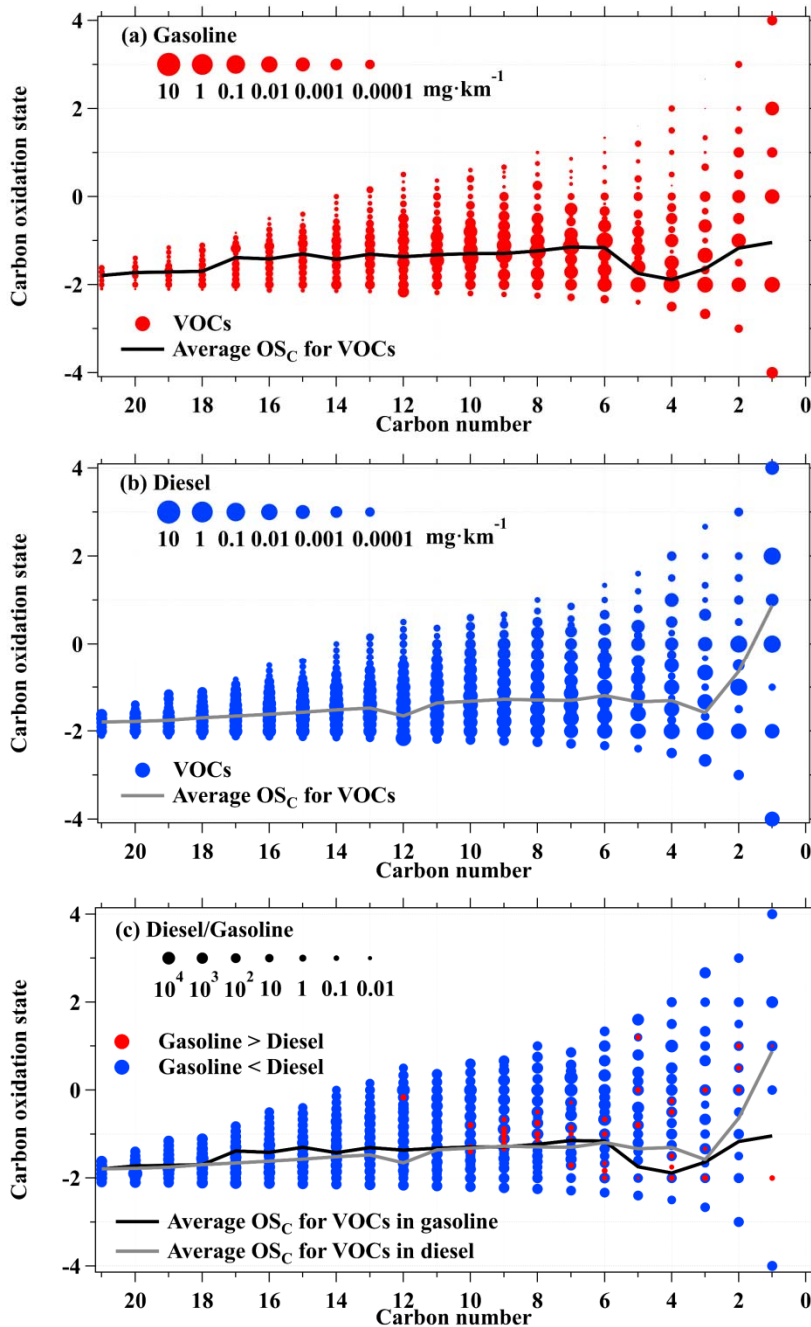


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885 **Figure 54.** The determined average mileage-based emission factors of VOC species  
 886 measured by PTR-ToF-MS from (a) gasoline, (b) diesel, and (c) LPG vehicles. The  
 887 gray dashed lines represent 1% of total VOCs emission factors.

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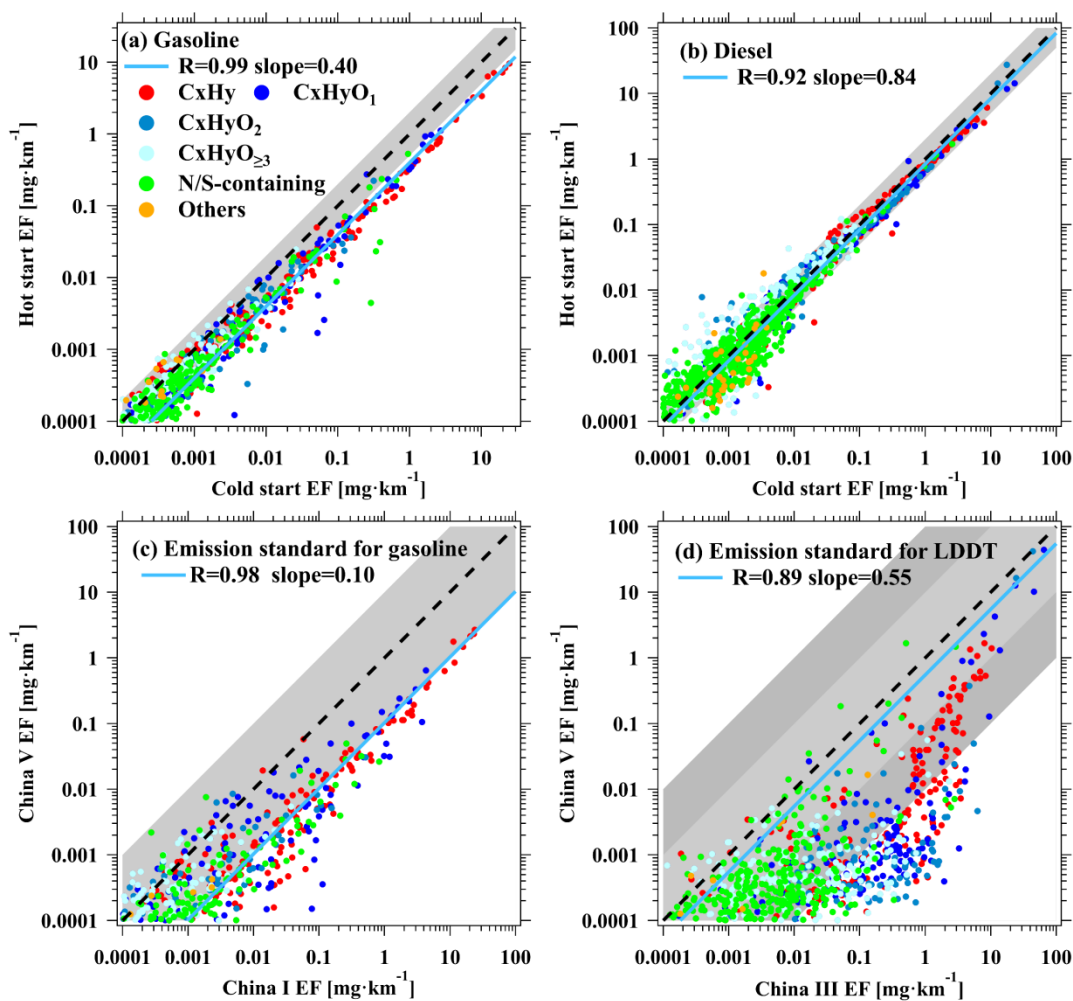




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890 **Figure 65.** The two-dimensional space of  $\overline{OS}_C - n_C$  with data points sized coded using  
 891 emission factors of VOC species from (a) gasoline and (b) diesel vehicles, and (c)  
 892 the ratio of emission factors of diesel vehicle relative to gasoline vehicle. The black and  
 893 gray lines are the average  $\overline{OS}_C$  of each carbon number for VOC species in gasoline and  
 894 diesel vehicles, respectively.

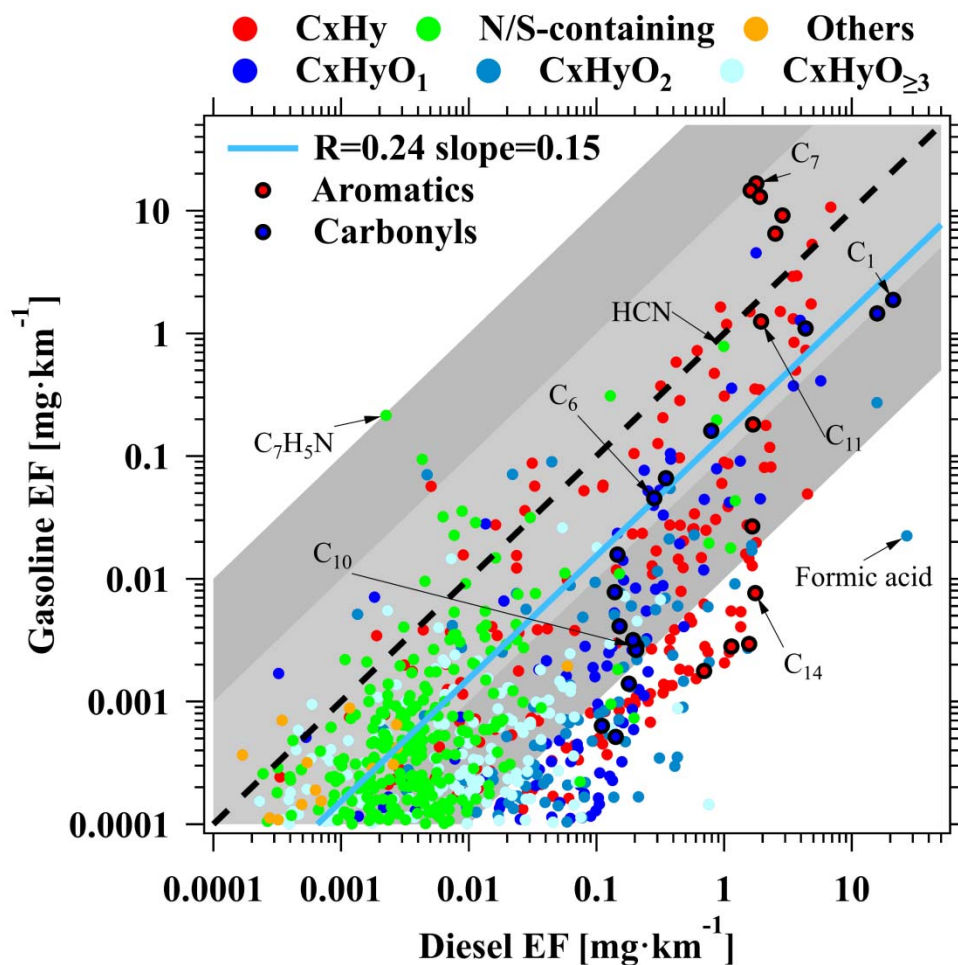
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897 **Figure 76.** Scatterplots of VOCs emission factors between cold start and hot start for  
 898 gasoline (a) and diesel vehicles (b). Scatterplots of VOCs emission factors between  
 899 China I and China V emission standard for gasoline vehicles (c) and between China III  
 900 and China V emission standard for diesel vehicles (d). Each data point indicates a VOC  
 901 species measured by PTR-ToF-MS. The blue lines are the fitted results for all data  
 902 points. The black dashed lines represent 1:1 ratio, and the shaded areas represent ratios  
 903 of a factor of 2 in (a) and (b), and a factor of 10 and 100 in (c) and (d).

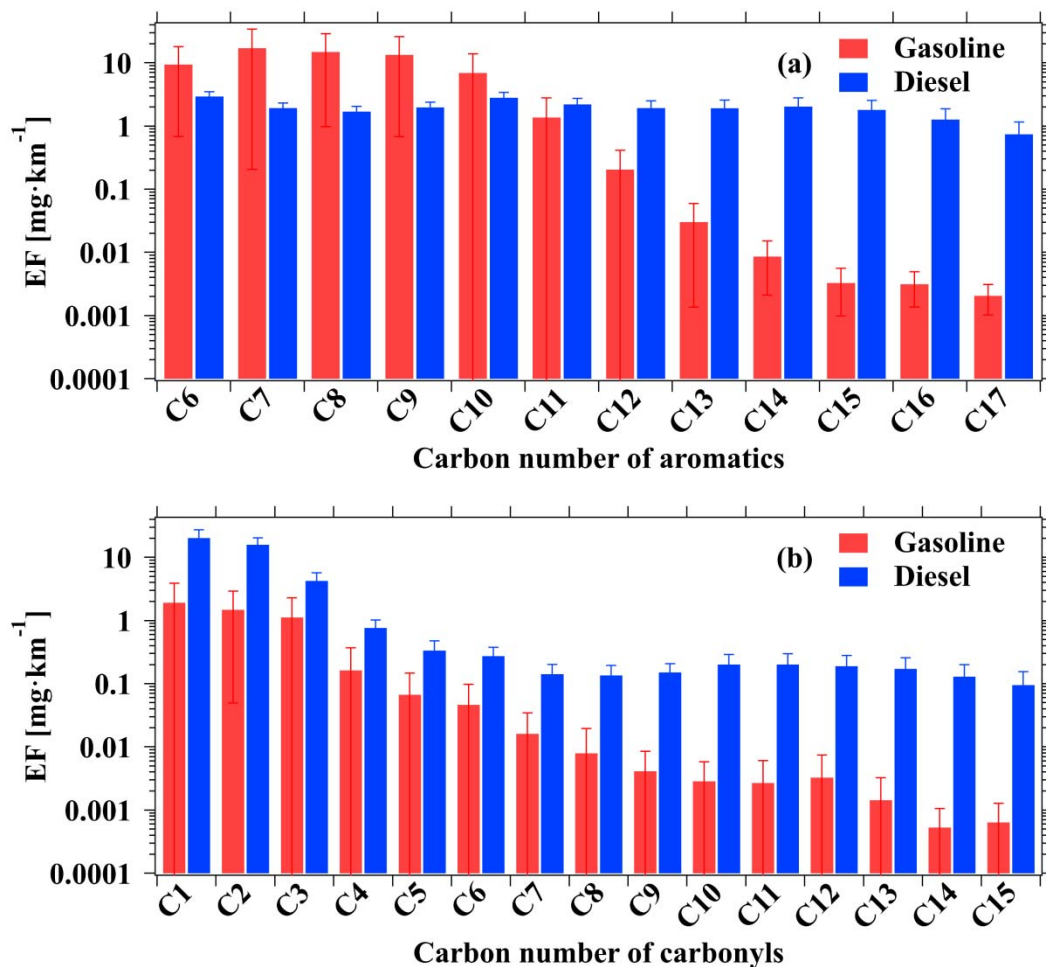
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906 **Figure 87.** Scatterplot of VOCs emission factors between gasoline and diesel vehicles.  
 907 Each data point indicates a VOC species measured by PTR-ToF-MS. The blue line is  
 908 the fitted result for all data points. The black line represents 1:1 ratio, and the shaded  
 909 areas represent ratios of a factor of 10 and 100.

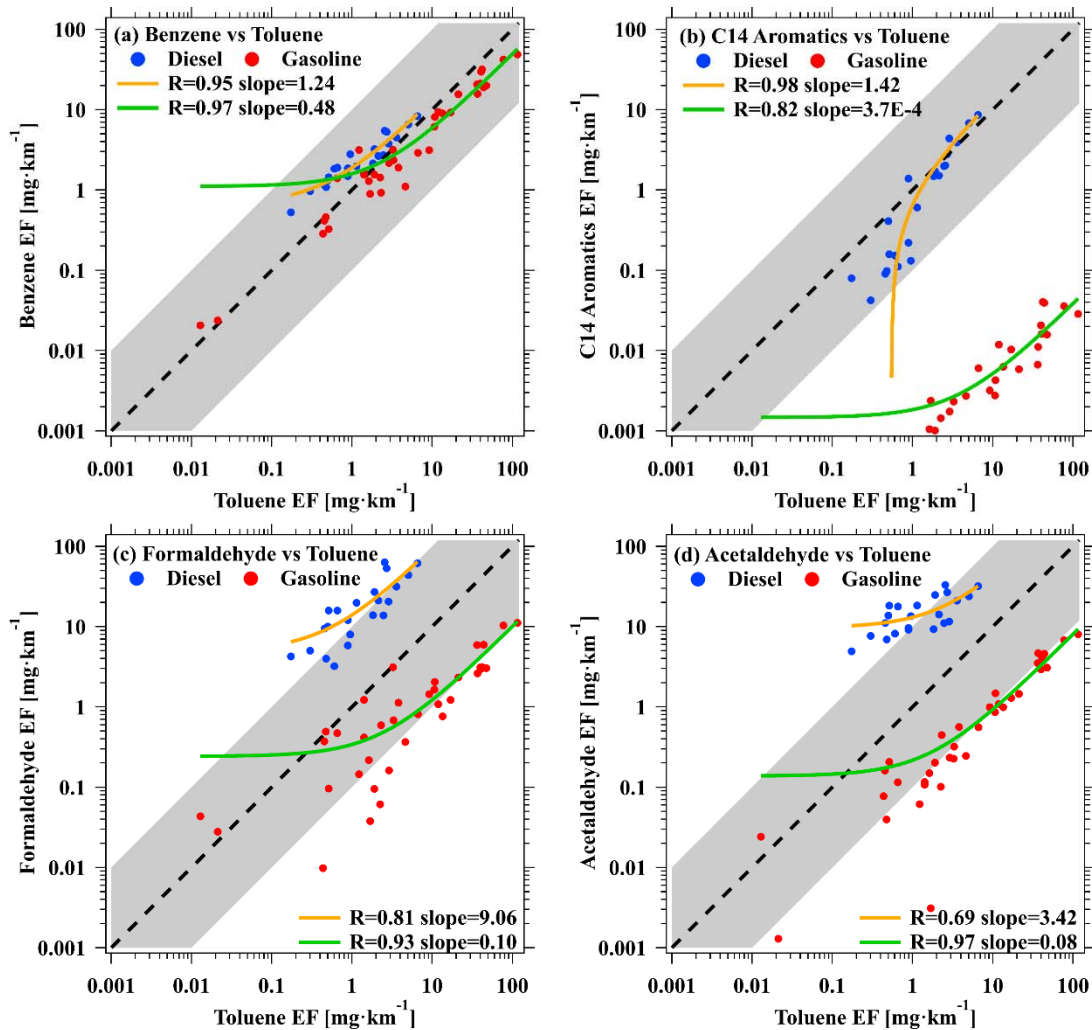
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912 **Figure 98.** The determined emission factors of (a) aromatics and (b) carbonyls for each  
 913 carbon number from gasoline and diesel vehicles. Error bars represent standard  
 914 deviations of the emission factors for the VOCs of different carbon number.

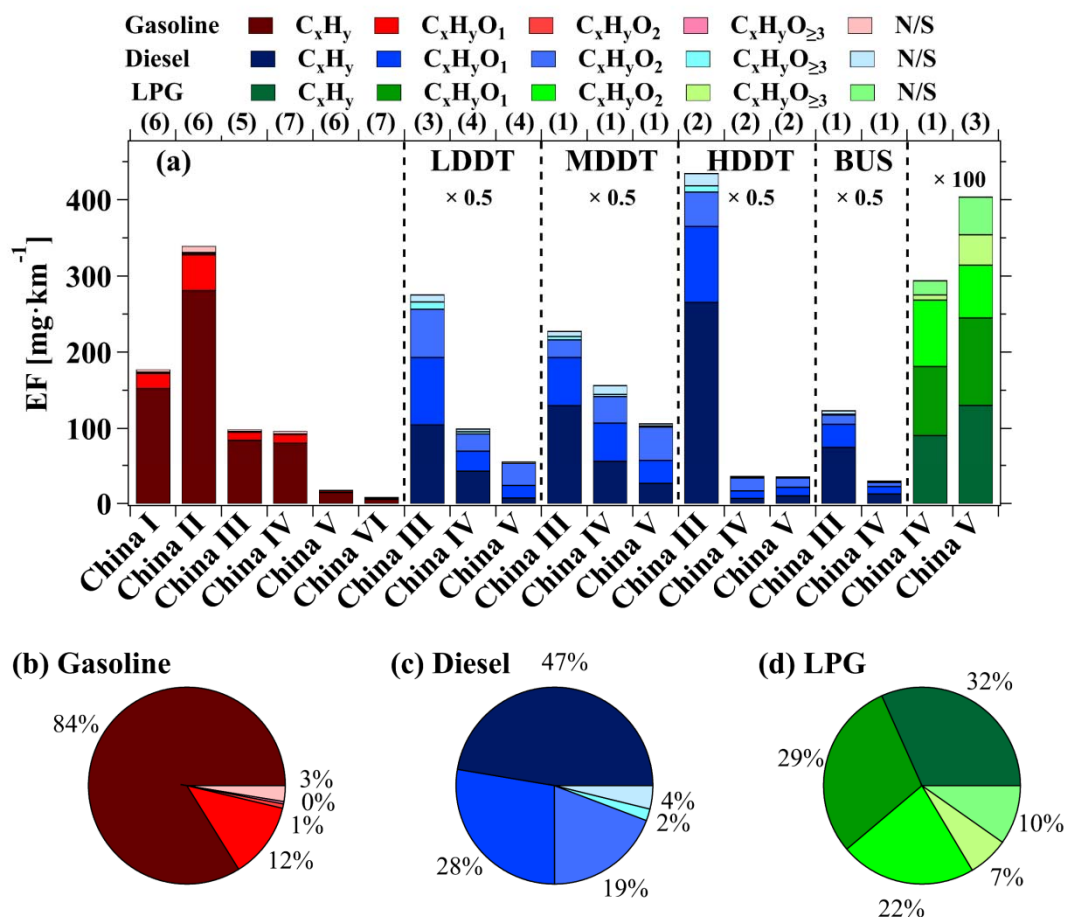
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917 **Figure 109.** Scatterplots of the determined mileage-based emission factors of (a)  
 918 benzene versus toluene, (b) C<sub>14</sub> aromatics versus toluene, (c) formaldehyde versus  
 919 toluene, and (d) acetaldehyde versus toluene for gasoline and diesel vehicles. Each data  
 920 point represents each test vehicle in this study. The green and orange lines are the fitted  
 921 results for gasoline and diesel vehicle. The black line represents 1:1 ratio, and the  
 922 shaded areas represent ratio of a factor of 10. The green and orange line are the fits to  
 923 gasoline and diesel points in each plot. Note that these linear fits are shown in curves  
 924 in log-log space as the result of non-zero y-intercept.

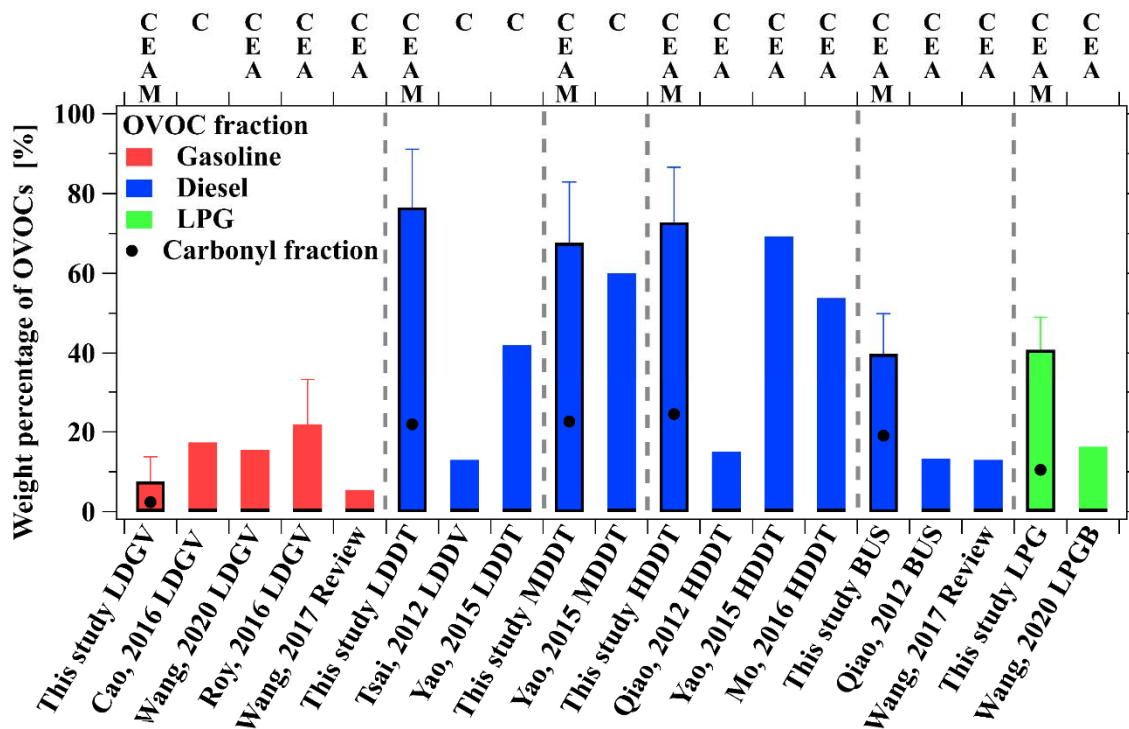
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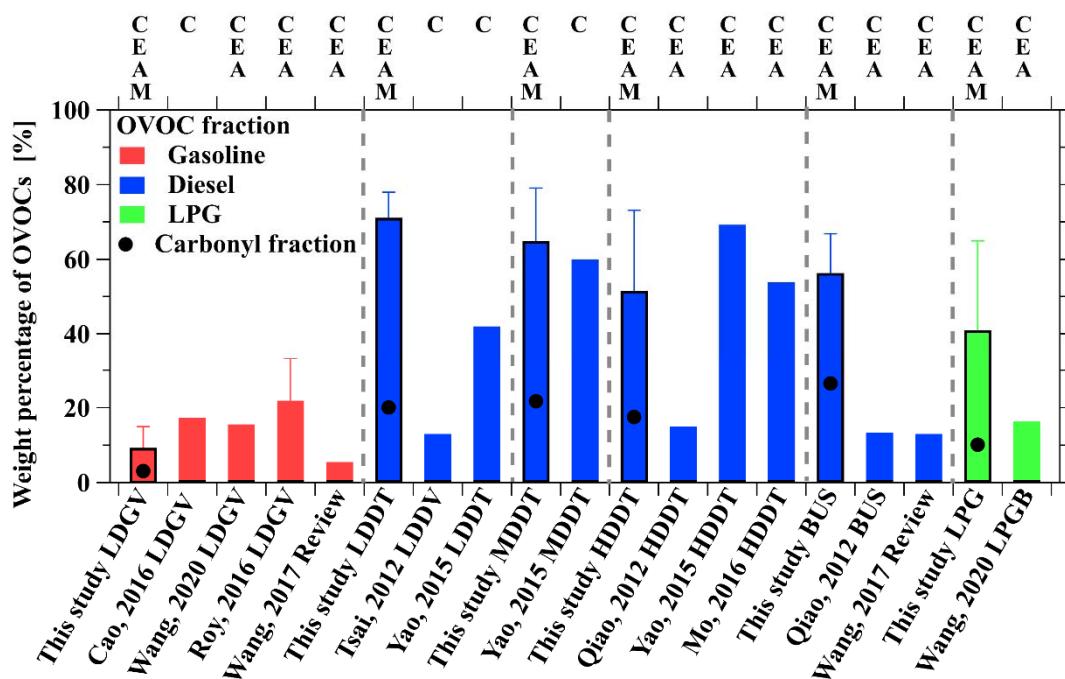
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927 **Figure 1140.** (a) The determined average emission factors for different emission  
 928 standard from gasoline, diesel (×0.5), and LPG (×100) vehicles measured by PTR-ToF-  
 929 MS. The different ion categories are discussed in the manuscript. Fractions of the  
 930 determined average emission factors of VOCs ions in different ion categories from (b)  
 931 gasoline, (c) diesel, and (d) LPG vehicles. The numbers above the top axis represent  
 932 the number of all experiments (including multiple measurements for individual test  
 933 vehicle) for each emission standard.

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937 **Figure 12.1.** Comparison of OVOCs fractions determined in this study and those in  
 938 previous studies. Error bars represent the standard deviations of the weight percentage  
 939 of OVOCs. The C, E, A, M above the top axis represent the four groups of OVOCs  
 940 measured in this study or previous studies, including Carbonyl: C, Ester/Ether: E,  
 941 Alcohol: A, Multiple-functional: M.