A large role of missing volatile organic compounds reactivity

from anthropogenic emissions in ozone pollution regulation

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Abstract: There are thousands of VOC species in ambient air, while existing techniques 20 can only detect a small part of them (~ several hundred). The large number of 21 unmeasured VOCs prevents us from understanding the photochemistry of ozone and 22 aerosols in the atmosphere. The major sources and photochemical effects of these 23 unmeasured VOCs in urban areas remain unclear. The missing VOC reactivity, which 24 is defined as the total OH reactivity of the unmeasured VOCs, is a good indicator to 25 constrain the photochemical effect of unmeasured VOCs. Here, we identified the 26 27 dominant role of anthropogenic emission sources in the missing VOC reactivity (accounting for up to 70%) by measuring missing VOC reactivity and tracer-based 28 source analysis in a typical megacity in China. Omitting the missing VOC reactivity 29 from anthropogenic emissions in model simulations will remarkably affect the 30 diagnosis of sensitivity regimes for ozone formation, overestimating the degree of 31 VOC-limited regime by up to 46%. Therefore, a thorough quantification of missing 32 VOC reactivity from various anthropogenic emission sources is urgently needed for 33 constraints of air quality models and the development of effective ozone control 34 35 strategies.

36

38 **1 Introduction**

Volatile organic compounds (VOCs) are key precursors of major photochemical 39 40 pollutants, including ozone (O₃) and secondary organic aerosols(Atkinson, 2000; Atkinson and Arey, 2003). Severe O₃ and particle pollution are frequently related 41 to high emissions of VOCs (Atkinson and Arey, 2003; Monks et al., 2015). There exist 42 thousands of VOC species in ambient air that are emitted from either natural processes 43 or anthropogenic activities (Goldstein and Galbally, 2007). No one instrument can 44 capture all VOCs out there and even when they can be measured there is information 45 missing on identification and properties (Yuan et al., 2017; Wang et al., 2014). Gas 46 chromatograph-mass spectrometer/flame ionization detector (GC-MS/FID) can 47 measure C2-C12 non-methane hydrocarbons (NMHCs) and C2-C6 oxygenated VOCs 48 (OVOCs) while cannot measure NMHCs and OVOCs with larger carbon number 49 (Wang et al., 2014). Proton-transfer-reaction time-of-flight mass spectrometer (PTR-50 ToF-MS) is able to measure a huge number of OVOCs and aromatics and several 51 52 alkanes, but cannot measure most alkanes and alkenes, and cannot distinguish isomers (Yuan et al., 2017). The 2,4-dinitrophenylhydrazine (DNPH)/high performance liquid 53 chromatography (HPLC) method can measure several carbonyls but cannot measure 54 non-polar organic species (Wang et al., 2009). The two-dimensional GC is able to 55 56 measure some intermediate-volatile and semi-volatile non-polar organics (Song et al., 2022). A lack of standard gases prevents these technologies from accurate 57 quantification even if these technologies can identify more VOC species. In general, 58 59 many branched alkenes, OVOCs with complex functional groups, intermediate-volatile and semi-volatile organics and complex biogenic VOCs cannot currently be well 60 quantified even if they can be identified by instruments. As a result, the total amount of 61 VOCs in ambient air has generally been underestimated. Currently, emission 62 inventories used in air quality models such as the Community Emissions Data System 63 (CEDS) emission inventory and the multi-resolution Emission Inventory for China 64 (MEIC) only include the VOC species that can be measured such as some C1-C9 65

hydrocarbons and simple-structure OVOCs with small carbon number (<C6). This will
lead to an underestimation of the photochemical effect of total VOCs and thus causes
uncertainties in predicting secondary pollution. The quantification of the unmeasured
VOCs is crucial to assess secondary pollution precisely.

The total OH reactivity (R_{OH}), which can be directly measured, is an index for evaluating the amount of reductive pollutants in terms of ambient OH loss. The total OH reactivity is defined as:

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$$R_{OH} = \sum_{i} k_{OH+Xi} [X_i], \tag{1}$$

where *X* represents a reactive species including carbon monoxide (CO), nitrogen oxides (NOx) and VOCs etc., and k_{OH+Xi} is the reaction rate constant for the oxidation of species *X* by OH. The measured R_{OH} is higher than that calculated based solely on the measured reactive species, and the difference between them is mostly from unmeasured VOCs (Yang et al., 2017). Missing VOC reactivity (missing VOC_R), defined as VOC reactivity (VOC_R) of all unmeasured VOCs, can be obtained by subtracting the calculated R_{OH} from the measured R_{OH}.

$$missing \ VOC_R = measured \ R_{OH} - calculated \ R_{OH}$$
(2)

82

calculated
$$R_{OH} = \sum_{i} k_{OH+reactive \, species_i} [reactive \, species_i]$$
 (3)

where reactive species represents measured VOCs and reactive inorganic species 83 including carbon monoxide (CO), nitric oxide (NO), nitrogen dioxide (NO₂), O₃, sulfur 84 dioxide (SO₂), nitrous acid (HONO), and so on. The missing VOC_R provides a 85 constraint for evaluating the photochemical roles of unmeasured VOCs in the 86 atmosphere (Sadanaga et al., 2005; Yang et al., 2016b). The inclusion of the missing 87 VOC_R can help to improve the performance of box model and air quality models in 88 89 simulating photochemistry processes. Relatively high missing VOC_R have been found in forests (Di Carlo et al., 2004;Hansen et al., 2014;Nakashima et al., 2014;Nölscher et 90 al., 2016; Praplan et al., 2019), urban areas (Shirley et al., 2006; Yoshino et al., 91 2006;Dolgorouky et al., 2012;Yang et al., 2017) and suburban areas (Kovacs et al., 92 2003; Yang et al., 2017; Fuchs et al., 2017; Lou et al., 2010), accouting for 10-75% of 93 total ROH. Given that total VOCR is one part of total ROH, missing VOCR would account 94

95 for a larger percentage of total VOC_R (>10%-75%).

The potential sources of missing VOC_R include anthropogenic emissions, biogenic emissions, soil emissions, and photochemical production, etc (Yang et al., 2016b). Previous studies have reported that the missing VOC_R in forest areas was mainly from either direct emissions or photochemical oxidation of biogenic VOCs (Di Carlo et al., 2004;Hansen et al., 2014;Nakashima et al., 2014;Nölscher et al., 2016;Praplan et al., 2019). Nevertheless, the dominant source of the missing VOC_R in urban and suburban areas remains unclear or under debate.

Surface O₃ pollution has become a major public health concern in cities worldwide 103 (Paoletti et al., 2014;Lefohn et al., 2018). A critical issue in determining an emission 104 control strategy for ozone pollution is to understand the relative benefits of NOx and 105 VOC emission controls. This is generally framed in terms of ozone precursor sensitivity, 106 i.e., whether ozone production is NOx-limited or VOC-limited (Kleinman, 107 1994;Sillman et al., 1990). Nevertheless, the effect of missing VOCs on ozone 108 precursor sensitivity has not been well understood yet. Given that the missing VOC_R 109 110 could potentially account for a large part of total VOCR, clearly clarifying the role of missing VOC_R in determining ozone precursor sensitivity is an urgent need for the 111 diagnosis of ozone sensitivity regimes and formulation of an effective emission 112 reduction roadmap. 113

114 China has become a global hot spot of ground-level ozone pollution in recent years 115 (Lu et al., 2018;Wang et al., 2022). Pearl River Delta (PRD) remains one of the most 116 O₃-polluted regions in China (Li et al., 2022), although many control measures have 117 been attempted. Here, we measured R_{OH} in Guangzhou, a megacity in PRD and 118 quantified the missing VOC_R. The dominant source of the missing VOC_R and its impact 119 on ozone precursor sensitivity were comprehensively investigated.

120 **2 Method**

121 **2.1 Overview of the measurment**

The field campaign was conducted from 25 September to 30 October 2018 continuously at an urban site in downtown Guangzhou (113.2°E, 23°N). The sampling site is located on the ninth floor of a building on the campus of Guangzhou Institute of Geochemistry, Chinese Academy of Sciences, 25 m above the ground level. This site is primarily influenced by industrial and vehicular emissions. ROH, VOCs, NOX, O3, HONO, SO2, CO, photolysis frequencies, and meteorological factors were simultaneously measured during the measurement period.

129 2.2 RoH measurement

Total ROH was measured by the comparative reactivity method (CRM) (Sinha et 130 al., 2008). The CRM system consists of three major components, namely an inlet and 131 132 calibration system, a reactor, and a measuring system. Here, pyrrole (C4H5N) was used 133 as the reference substance in CRM and its concentration was quantified by a quadrupole proton-transfer-reaction mass spectrometer (PTR-MS) (Ionicon Analytik GmbH, 134 Innsbruck, Austria). The CRM system was calibrated by propane, propene, toluene 135 standards and 16 VOC mixed standard (acetaldehyde, methanol, ethanol, isoprene, 136 137 acetone, acetonitrile, methyl vinyl ketone, methyl ethyl ketone, benzene, toluene, oxylene, α -pinene, 1,2,4-trimethylbenzene, phenol, m-cresol, and naphthalene). 138 Measured and calculated RoH agreed well within 15% for all calibrations. The RoH 139 measurement by the CRM method is interfered from ambient nitric oxide (NO), which 140 141 produces additional OH radicals via the reaction of HO₂ radicals with NO (Sinha et al., 2008). To correct this interference, a series of experiments were conducted by 142 introducing different levels of NO (0-160 ppb) and given amounts of VOC into the 143 CRM reactor. A correction curve was acquired from these NO interference experiments, 144 which can be used to correct the R_{OH} thanks to the simultaneous measurement of 145

ambient NO concentrations (Supplementary information S1; Fig. S1). The detection 146 limits of the CRM method were around 2.5 s^{-1} , and the total uncertainty was estimated 147 to be about 15%. The CRM method has been successfully applied to measure OH 148 reactivity in urban areas with high NO_X levels in previous studies (Dolgorouky et al., 149 2012; Yang et al., 2017; Hansen et al., 2015). The intercomparison between the CRM 150 method and pump-probe technique indicates that the CRM method can be used under 151 high-NO_X conditions (NO_X>10 ppb) if a NO_X -dependent correction is applied (Hansen 152 et al., 2015). 153

154 **2.3 VOCs measurements**

Nonmethane hydrocarbons (NMHCs) were measured using a gas chromatograph-155 mass spectrometer/flame ionization detector (GC-MS/FID) system coupled with a 156 157 cryogen-free preconcentration device (Wang et al., 2014). The system contains twochannel sampling and GC column separation, which is able to measure C2-C5 158 hydrocarbons with the FID in one channel and measure C5-C12 hydrocarbons using 159 MS detector in the other channel. After removal of water vapor, VOCs were trapped at 160 -155 °C in a deactivated quartz capillary column (15 cm×0.53 mm ID) and a Porous 161 Layer Open Tubular (PLOT) capillary column (15 cm×0.53 mm ID) for the MS channel 162 and the FID channel, respectively. The system was calibrated weekly by TO-15 (Air 163 Environmental Inc., USA) and PAMS gas standards (Spectra Gases Inc., USA). 164 165 Detection limits for various compounds were in the range of 0.002-0.070 ppbv. A total of 56 NMHCs species were measured (Table S1). The time resolution of the 166 measurement was 1 h. The uncertainties of VOC measurements by GC-MS/FID are in 167 the range of 15 %–20 %. More details of this method can be found in previous studies 168 (Wang et al., 2014; Yuan et al., 2012). 169

An online proton-transfer-reaction time-of-flight mass spectrometer (PTR-ToF-MS) (Ionicon Analytic GmbH, Innsbruck, Austria) with H_3O^+ and NO^+ ion sources was also used to measure VOCs. During the campaign, the PTR-ToF-MS automatically switched between H_3O^+ and NO^+ chemistry every 10–20 min. The H_3O^+ mode was

used to measure OVOCs and aromatics while the NO⁺ model was used to measure 174 alkanes with more carbons (C8-C20). When running in the H₃O⁺ ionization mode, the 175 drift tube was at a temperature of 50 °C, a pressure of 3.8 mbar, and a voltage of 920 176 V, leading to an operating E/N (E is the electric field, and N is the number density of 177 the gas in the drift tube) ratio of 120 Td. When running in the NO⁺ ionization mode, the 178 drift tube was at a temperature of 50 °C, a pressure of 3.8 mbar, and a voltage of 470 179 V, leading to an operating E/N ratio of 60 Td. PTR-ToF-MS technique is capable of 180 measuring oxygenated VOCs (OVOCs) and higher alkanes that GC-MS/FID cannot 181 measure (Wu et al., 2020; Wang et al., 2020a). The time resolution of PTR-ToF-MS 182 measurements was 10 s. A total of 31 VOCs were calibrated using either gas or liquid 183 standards (Table S2). For other measured VOCs, we used the method proposed by 184 Sekimoto et al. (2017) to determine the relationship between VOC sensitivity and 185 kinetic rate constants for proton transfer reactions of H₃O⁺ with VOCs. The fitted line 186 was used to determine the concentrations of those uncalibrated species. The 187 uncertainties of the concentrations for uncalibrated species were about 50 % (Sekimoto 188 189 et al., 2017). By this metod, PTR-ToF-MS can additionally measure 128 VOCs which were included in the analysis of this study. The detailed information for this method can 190 be found in Wu et al. (2020) and all VOC species measured by PTR-ToF-MS were 191 provided in table S4 of that article. The PTR-ToF-MS is capable of measuring 192 additional VOC species that GC-MS/FID cannot measure including alkanes with more 193 carbons (C12-C20) and OVOCs including aldehydes, ketones, carboxylic acids, 194 195 alcohols, and nitrophenols. Formaldehyde (HCHO) was measured by a custom-built 196 instrument based on the Hantzsch reaction and absorption photometry (Xu et al., 2022).

197 **2.4 Other measurements**

Nitrous acid (HONO) was measured by a custom-built LOPAP (Long Path
Absorption Photometer) based on wet chemical sampling and photometric detection
(Yu et al., 2022). The uncertainty of the measurement was 8 %. NO_X, O₃, SO₂, and CO
were measured by NO_X analyzer (Thermo Scientific, Model 42i), O₃ analyzer (Thermo
Scientific, Model 49i), SO₂ analyzer (Thermo Scientific, Model 43i), and CO analyzer

(Thermo Scientific, Model 48i), respectively. The meteorological data, including temperature (T), relative humidity (RH) and wind speed and direction (WS, WD) were recorded by Vantage Pro2 Weather Station (Davis Instruments Inc., Vantage Pro2) with a time resolution of 1 min. Photolysis frequencies of O₃, NO₂, HONO, H₂O₂, HCHO, and NO₃ were measured by a spectrometer (Focused Photonics Inc., PFS-100) (Shetter and Müller, 1999;Wang et al., 2019).

209 2.5 Multiple linear regression

The Multiple Linear Regression (MLR) has been successfully applied to quantify the sources of air pollutants (Li et al., 2019; Yang et al., 2016a). In this study, a tracerbased MLR analysis was used to decouple the individual contributions of anthropogenic emissions, secondary production, biogenic emissions and background level to missing VOC_R, as shown in Eq. (4).

Missing $VOC_R = a\Delta CO + b[O_X] + c[isoprene_{initial}] + C_{background}$ (4) 215 where O_X is defined as O_3+NO_2 . ΔCO , $[O_X]$ and $[isoprene_{initial}]$ are concentrations 216 of tracers for anthropogenic emissions, secondary production and biogenic emissions, 217 respectively. ΔCO is the relative change between ambient CO and background CO of 218 150 ppb (Wang et al., 2020a). [isoprene_{initial}] represents the initial concentration of 219 isoprene from biogenic emissions that has not undergone any photochemical reactions, 220 which is calculated from observed isoprene and its photochemical products methyl 221 vinyl ketone (MVK) and methacrolein (MACR) (Xie et al., 2008). Cbackground 222 223 indicates the background level of missing VOC_R. a, b, c and C_{backgound} are fitted coefficients by the multiple linear regression. 224

225 **2.6 Observation-based box model**

A zero-dimensional box model coupled with the Master Chemical Mechanism (MCM) v3.3.1 chemical mechanism (Jenkin et al., 2003) was used to simulate the photochemical production of RO_X ($RO_X=OH+HO_2+RO_2$) radicals and O_3 during the field campaign. The model was constrained by the observations of meteorological parameters, photolysis frequencies, VOCs, NO, NO₂, O₃, CO, SO₂, and HONO. The
model runs were performed in a time-dependent mode with a time resolution of 1 hour
and a spin-up of four days. A 24-h lifetime was introduced for all simulated species,
including secondary species and radicals, to approximately simulate dry deposition and
other losses of these species (Lu et al., 2013;Wang et al., 2020b). Sensitivity tests show
that this assumed physical loss lifetime has a relatively small influence on ROx radicals
and ozone production rates.

Measured OVOCs such as HCHO, acetaldehyde and acetone were constrained in the model and unmeasured OVOCs were simulated according to the photochemical oxidation of NMHCs by OH radicals. RO_2 , HO_2 and OH radicals were simulated by the box model to calculate the net O₃ production rate (P(O₃)) and O₃ loss rate (L(O₃)) as shown in Equations (5) and (6) as derived by Mihelcic et al. (2003)

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$$P(O_3) = k_{HO_2+NO}[HO_2][NO] + \sum_i (k_{RO_2+NO}^i [RO_2^i][NO]) - k_{OH+NO_2}[OH][NO_2] - L(O_3)$$
243 (5)

244
$$L(O_3) = (\theta j (O^1 D) + k_{OH+O_3} [OH] + k_{HO_2+O_3} [HO_2] + \sum_j (k_{alkene+O_3}^j [alkene^j]) [O_3]$$
245 (6)

where θ is the fraction of O¹D from ozone photolysis that reacts with water vapor, and i and j represent the number of species of RO₂ and alkenes, respectively.

The box model was used to evaluate the impact of missing VOC_R on the O₃ 248 production rate. In the base scenario, the box model was constrained by all measured 249 inorganic and organic gases but the missing VOCR was not considered. To consider the 250 missing VOC_R in the box model, we additionally increased the concentration of 251 NMHCs to exactly compensate for the missing VOC_R by multiplying a factor, on the 252 253 basis of measured NMHC concentrations. We simulated four scenarios by increasing the concentration of: (1) n-pentane, (2) ethylene, (3) toluene, (4) all measured 56 254 NMHCs. For the scenario of increasing all 56 NMHCs, concentrations of 56 NMHC 255 species were increased by multiplying the same factor. Given that the VOC_R of 256 unconstrained secondary products increases with the increase in the concentration of 257 NMHCs, several attempts of different values are needed to determine the increasing 258

259 factor.

260 **3 Results and discussion**

261 **3.1 Quantification of missing VOC**_R during the campaign

Figure 1 shows the time series of measured ROH, calculated ROH according to all 262 measured reactive gases, and missing VOC_R (the gap between measured and calculated 263 R_{OH}) in Guangzhou. By using GC-MS/FID, we measured 56 NMHCs. By using PTR-264 265 ToF-MS, we measured 159 VOCs and 128 of them were difficult to be measured before. Besides the alkanes with carbons less than 12, PTR-ToF-MS can also measure alkanes 266 with more carbons (C12-C20). With regard to OVOCs, not only common OVOC 267 species including formaldehyde and C2-C4 carbonyls but also carbonyls with more 268 269 carbons (C5-C10) and some N-containing OVOC species such as nitrophenol and methyl nitrophenol were measured by PTR-ToF-MS. Thanks to these additional 270 measured VOCs, the measured R_{OH} was close to the calculated R_{OH} within 20% in most 271 272 periods. In some periods the missing VOC_R was negative, which is probably due to the uncertainty in the measurements of ROH and reactive gases. The negative missing VOCR 273 primarily occurred in the afternoon (12:00-17:00) when the photochemistry was most 274 active. Nevertheless, there were still some days exhibiting remarkable missing VOC_R. 275 The days with missing VOC_R of more than 25% of total R_{OH}, namely high missing-276 277 VOC_R days, are indicated by yellow background in Fig. 1a. The largest missing VOC_R occurred on October 15th, 16th, 25th and 26th, with average values of 16 s⁻¹. During the 278 period of October 24th to 26th, the total R_{OH} was highest and the missing VOC_R was also 279 relatively high among all days. Figure 1b shows the contribution of different species 280 classifications to total ROH during high missing-VOCR days. Inorganic species, NMHCs 281 and OVOCs account for 34%, 13% and 14% of total RoH, respectively, with missing 282 VOC_R accounting for 39%. The fraction of missing VOC_R (39%) during the high 283 missing-VOC_R days is comparable to measurements in Los Angeles 2010 (Griffith et 284 al., 2016) and in Seoul 2016 (Sanchez et al., 2021). 285

We evaluated the uncertainty of the missing VOC_R. The uncertainty of the R_{OH} 286 measurement was 15%. In addition, according to reports of Jet Propulsion Laboratory 287 (Burkholder et al., 2020), reaction rate constants used for the calculation of ROH in Eq. 288 (3) have uncertainties of 5%-30%, depending on different species. We took the 289 uncertainties in the reaction rate constants and the measurements of all reactive gases 290 into account when calculating ROH, according to error propagation. As a result, the 291 uncertainties in the missing VOC_R are 3.8 s^{-1} and 5.2 s^{-1} for the whole measurement 292 period and the high missing-VOC_R days, respectively. The average missing VOC_R 293 during the high missing-VOC_R days is 13 s⁻¹, which is significantly higher than the 294 uncertainty of 5.2 s⁻¹, suggesting that the missing VOC_R really exists during the high 295 missing-VOC_R days. 296

297

298 **3.2 The sources of missing VOC**_R

To explore the sources of missing VOC_R during the whole measurement period, 299 we investigated the correlation between missing VOC_R and tracers characterizing 300 primary emissions (CO, NO_X and NMHCs) and secondary production (O_X=O₃+NO₂ 301 and formic acid). The correlation of missing VOC_R with CO, reactivity of NMHCs 302 (NMHC_R) and NO_X is moderate, with correlation coefficient (R) in the range of 0.47– 303 0.56 (Fig. 2a and b, and Fig. S2) while there is no significant correlation of missing 304 305 VOC_R with O_X and formic acid (Fig. 2c and Fig. S2). Furthermore, there is no significant correlation between missing VOC_R and acetonitrile which is a tracer of 306 biomass burning (de Gouw et al., 2003; Wang et al., 2007) (Fig. S2), indicating that 307 biomass burning was not a major contributor to missing VOC_R during this campaign. 308 In terms of the diurnal variation, the missing VOC_R was higher in the morning (7:00– 309 10:00) and evening (18:00-22:00) when the anthropogenic emissions, especially 310 vehicle exhaust were intensive, and was lower in the afternoon when the 311 photochemistry was most active (Fig. 2d). The diurnal profile of missing VOC_R was 312 similar to those of CO, NO_X and NMHC_R. In contrast, the diurnal profiles of secondary 313

species including O_X, formic acid and acetic acid, which peaked in the afternoon, 314 evidently differ from the diurnal profile of missing VOC_R (Fig. S3). Further, we 315 316 investigated the influence of airmass aging on missing VOC_R. The ratio of ethylbenzene to m,p-xylene was used to characterize the degree of air mass aging (De Gouw et al., 317 2005; Yuan et al., 2013). A higher ratio of ethylbenzene to m,p-xylene corresponds to a 318 higher degree of air mass aging as the m,p-xylene has a larger reaction rate constant 319 $(18.9 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1}\text{s}^{-1})$ than ethylbenzene $(7.0 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1}\text{s}^{-1})$ when 320 reacting with the major oxidant - OH radicals. As shown in Fig. 2e, missing VOCR 321 decreases with the ratio of ethylbenzene to m,p-xylene. Given that secondary 322 production generally increased with air mass aging, this result further demonstrates that 323 missing VOC_R was not caused by enhanced secondary production. 324

During the high missing- VOC_R days, the correlation coefficient for missing VOC_R 325 versus CO is 0.76 (Fig. 3a), which is higher than that in the whole measurement period 326 (0.56) shown in Fig. 2a. We then quantify the sources of missing VOC_R during the high 327 missing- VOC_R days by applying MLR. The fitted coefficient a is $0.031 \text{ s}^{-1} \text{ ppb}^{-1}$, b is 328 0.012 s⁻¹ ppb⁻¹, c is 1.8 s⁻¹ ppb⁻¹ and C_{background} is 1.3 s⁻¹. The coefficient of determination 329 (\mathbb{R}^2) for the MLR is 0.68. As shown in **Fig. 3b**, anthropogenic emissions were the largest 330 contributor to missing VOCR, accounting for 70% of missing VOCR. Secondary 331 production, biogenic emissions and background contribution played a minor role in 332 missing VOC_R (13%, 7%, 10%, respectively). The parametric relationship between 333 missing VOC_R and relevant tracers established by MLR provides a valid approach to 334 estimate the missing VOC_R according to readily available gases including CO, O_X and 335 336 isoprene.

Although anthropogenic emissions are identified to be the major source of missing VOC_R, which species dominantly contribute to the missing VOC_R remains unclear. A potential source is the unmeasured branched alkenes for their high reactivity, previously observed from vehicle exhaust (Nakashima et al., 2010) and gasoline evaporation emissions (Wu et al., 2015). Another possible source is emitted OVOCs with a more complex functional group that cannot be accurately measured. In addition, directly emitted semi-volatile and intermediate volatility organic compounds are also possible
sources of missing VOC_R (Stewart et al., 2021).

345 **3.3 The impact of missing VOC**_R on O₃ sensitivity regimes

346 The reaction of OH with VOCs is key to the propagation and amplification of OH radicals, thus determining the ozone production rate (Tonnesen and Dennis, 2000). The 347 box model was used to evaluate the impact of missing VOCR on the O3 production rate 348 during high missing-VOC_R days. The setting of model simulations for different 349 350 scenarios are depicted in Section 2.6. Under the base scenario, on average the measured VOC_R of n-pentane, ethylene, toluene and all 56 NMHCs are 0.14 s⁻¹, 0.53 s⁻¹, 0.60 s⁻¹ 351 and 4.6 s⁻¹ respectively. To consider the missing VOC_R (on average of 13 s⁻¹) in the 352 model, four scenarios were simulated by additionally increasing n-pentane, ethylene, 353 354 toluene and 56 NMHCs by a factor of 70, 16, 13.3 and 1.9, respectively. These increasing factors led to an additional increase in VOCR of both NMHCs and 355 unconstrained secondary products, which exactly compensated for the missing VOC_R. 356 Figure 4 shows the simulated $P(O_3)$ for the base scenario and the scenarios considering 357 missing VOC_R. The daytime average P(O₃) under the scenarios considering missing 358 VOC_R is a factor of 1.5–4.5 for the results under the base scenario. The difference in 359 added species has a large effect on P(O₃). Adding toluene causes a larger increase in 360 P(O₃) than adding n-pentane or ethene, as toluene has a stronger ability to amplify the 361 362 production of radicals.

O₃ precursor sensitivity depends on the dominant loss pathways of RO_X radicals 363 (RO_X=OH+HO₂+RO₂). O₃ production is NO_X-limited if the self-reaction of peroxy 364 radicals (HO₂ and RO₂) dominates the RO_X sink, and VOC-limited if the reaction of 365 NO₂ with OH dominates (Kleinman et al., 1997;Kleinman et al., 2001). Accordingly, 366 the ratio of RO_X sink induced by OH+NO₂ reaction to the total rate of the two RO_X 367 sinks, i.e., L_N/Q , is used to identify O₃ sensitivity regimes. O₃ production is NO_X-368 limited if L_N/Q is lower than 0.5, otherwise, it is VOC-limited (Kleinman et al., 1997). 369 $k_{OH+NO_2}[OH][NO_2]$ 370 $L_{\rm W}/0 = -$

$$k_{HO_2+RO_2}[HO_2][RO_2] + k_{HO_2+HO_2}[HO_2][HO_2] + k_{OH+HO_2}[OH][HO_2] + k_{OH+HO_2}[OH][NO_2]$$

(7)

As shown in **Fig. 5a**, under the base scenario, L_N/Q remained at a stable and high 372 level (>0.9) during the daytime when photochemical production of ozone occurs, 373 indicating O₃ production was VOC-limited. Under the scenarios considering missing 374 VOC_R, L_N/Q decreased significantly regardless of which VOC species was added, 375 compared to the base scenario. Adding toluene caused the largest decrease in L_N/Q, 376 377 followed by adding all measured NMHC species, adding the alkane and adding the alkene. It is worth noting that adding toluene and all measured NMHC species caused 378 the L_N/Q to be close to 0.5 in the afternoon, indicating that the O₃ production shifted to 379 transitional or NO_X-limited regimes in these scenarios. Fig. 5b shows the changes in 380 381 radical sinks before and after considering missing VOC_R. All radical sinks including self-reactions of peroxy radicals and OH+NO2 reaction increased after considering 382 missing VOC_R. Nevertheless, the increased proportion of the self-reactions of peroxy 383 radicals was larger than that of OH+NO₂ reaction, leading to a decrease in L_N/Q and 384 thus a shift toward NO_X-limited regime. 385

386 Figure 5c shows the dependence of daily peak O₃ concentrations on NO_X 387 concentrations, which was calculated by the box model for the base scenario and the scenario considering missing VOC_R. The NO_X concentration level corresponding to the 388 maximum of O3 concentrations was determined. This NOx concentration level reflects 389 the threshold to distinguish between VOC-limited and NOx-limited regimes. The larger 390 threshold of NO_X represents a higher possibility of ozone production in NO_X limited 391 regime. The threshold of NO_X for the scenario considering missing VOC_R is 46% higher 392 than for the base scenario. Note that the uncertainty in missing VOC_R leads to 17% 393 394 uncertainty in the threshold of NO_X for the scenario considering missing VOC_R. Overall, Fig. 5 suggests that omitting the missing VOC_R will overestimate the degree of the 395 VOC-limited regime and thus overestimate the effect of VOCs abatement in reducing 396 ozone pollution, which in turn may mislead ozone control strategy. 397

398 **3.4 Atmospheric implications**

Although many previous studies have reported that photochemical production 399 processes and biogenic emissions are important sources of missing VOC_R (Lou et al., 400 2010;Dolgorouky et al., 2012;Yang et al., 2017;Sanchez et al., 2021;Di Carlo et al., 401 2004), we find that anthropogenic emissions may dominate the missing VOC_R in urban 402 403 regions. In zero-dimensional box models and three-dimensional chemistry-transport models, the input of VOCs emission information mainly contains well-studied simple-404 structure alkanes, alkenes and aromatics, while those unmeasured/unknown VOC 405 species have been neglected. This will lead to biases in quantifying ozone production 406 407 and diagnosing ozone sensitivity regimes. Our study demonstrates that the ambient measurement of R_{OH} at urban sites can provide quantification of missing VOC_R, which 408 can be used in models to account for the missing VOC_R from anthropogenic emissions. 409 In addition, the parametric equation of missing VOC_R derived from MLR method (Eq 410 (4)) here can be used to estimate missing VOC_R according to measurements of CO, O_X 411 and isoprene. Further study should try to parse the specific sources of the missing VOC_R, 412 e.g., whether the missing VOC_R is from intermediate-volatility and semivolatile organic 413 414 compounds emitted from vehicles or whether it is from some other sources. Furthermore, future studies can focus on direct measurements of missing VOC_R for 415 various emission sources to develop a comprehensive emission inventory of missing 416 VOC_R, which will help to improve O₃ pollution mitigation strategies. 417

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Figure 1. The level of missing VOC_R during the measurements in Guangzhou. (a) Time series of measured R_{OH} and calculated R_{OH} from all measured reactive gases in Guangzhou. Yellow background represents the high missing-VOC_R days with missing VOC_R accounting for more than 30% of total R_{OH}. (b) Contributions of different compositions to R_{OH} in high missing-VOC_R days. The error bar represents standard deviation of missing VOC_R.

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Figure 2. Correlation of missing VOC_R with major tracers during the whole measurement period. (a-c) Correlation of missing VOC_R with CO, OH reactivity of NMHCs (NMHC_R) and O_X. Each point represents hourly data. (d) Diurnal variations in missing VOC_R, CO, NO_X and NMHCs. (e) The dependence of missing VOC_R on ethylbenzene to m, p-xylene ratio. The red squares indicate the mean values of missing VOCR in different ranges of ethylbenzene/m,p-xylene with classification width of 0.1, and the error bars represent standard deviation.





Figure 3. The source apportionment of missing VOC_R in high missing-VOC_R days.

657 (a) Correlation of missing VOC_R with CO. Each point represents hourly data. (b)

658 Contributions of different sources to missing VOC_R according to the MLR.

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Figure 4. Simulated daytime mean $P(O_3)$ for the base scenario (without missing VOC_R) and the scenario considering missing VOC_R , respectively, in high-missing VOC_R days. The missing VOC_R is considered by adding individual species (n-pentane, ethene or toluene) or increasing all measured NMHCs to compensate for the missing VOC_R . The error bar represents standard deviation of $P(O_3)$ induced by the uncertainty of missing VOC_R .



Figure 5. The impact of missing VOC_R on O₃ sensitivity for the high-missing VOC_R days. (a) Diurnal variations in L_N/Q for the base scenario and the scenarios considering missing VOC_R. The missing VOC_R is considered by adding individual species (n-pentane, ethene or toluene) or increasing all measured NMHCs to fill the missing VOC_R. The dashed line represents the threshold value of L_N/Q that distinguishes VOC-limited and NO_X-limited regimes. (b) The averages of radical sinks in the afternoon (12:00-18:00) for the base scenario (red bar) and the scenario considering missing VOC_R (blue bar) by increasing all measured NMHCs to fill the missing VOC_R (blue bar) by increasing all measured NMHCs to fill the missing VOC_R. (c) Model-simulated dependence of daily peak O₃ concentrations on daily mean NO_X concentrations for the

base scenario (red curve) and the scenario considering missing VOC_R (blue curve) by increasing all measured NMHCs to fill the missing VOC_R. The dashed lines parallel to Y-axis represent the threshold of NO_X levels to distinguish between VOC-limited and NO_X-limited regimes. The shaded area represents standard deviation induced by the uncertainty in missing VOC_R.