1 Vertical changes in volatile organic compounds (VOCs) and

2 impacts on photochemical ozone formation

3 Xiao-Bing Li¹, Bin Yuan^{1,*}, Yibo Huangfu¹, Suxia Yang², Xin Song¹, Jipeng Qi¹,

4 Xianjun He¹, Sihang Wang¹, Yubin Chen¹, Qing Yang¹, Yongxin Song¹, Yuwen Peng¹,

5 Guiqian Tang^{3, 4}, Jian Gao⁵, Dasa Gu⁶, Min Shao¹

6 ¹ College of Environment and Climate, Institute for Environmental and Climate

7 Research, Guangdong-Hongkong-Macau Joint Laboratory of Collaborative Innovation

8 for Environmental Quality, Jinan University, Guangzhou 511443, China

² Guangzhou Research Institute of Environment Protection Co., Ltd., Guangzhou
510620, China

³ State Key Laboratory of Atmospheric Environment and Extreme Meteorology,

12 Institute of Atmospheric Physics, Chinese Academy of Sciences, Beijing 100029, China

⁴ University of Chinese Academy of Sciences, Beijing, 100049, China

⁵ State Key Laboratory of Environmental Criteria and Risk Assessment, Chinese
 Research Academy of Environmental Sciences, Beijing 100012, China

16 ⁶ Guangdong-Hongkong-Macau Joint Laboratory of Collaborative Innovation for

17 Environmental Quality and Division of Environment and Sustainability, The Hong

18 Kong University of Science and Technology, Hong Kong 999077, China

^{*}Corresponding author: Bin Yuan (byuan@jnu.edu.cn)

20 Abstract

21 Volatile organic compounds (VOCs) play crucial roles in regulating the formation 22 of tropospheric ozone. However, limited knowledge on the interactions between 23 vertical VOC variations and photochemical ozone formation in the planetary boundary 24 layer (PBL) has hindered effective ozone control strategies, especially in large cities. 25 In this study, we investigated the vertical changes in concentrations, compositions, and 26 key driving factors of a large suite of VOCs using online gradient measurements taken 27 from a 325 m tall tower in urban Beijing. We also analyzed tThe impact of these vertical 28 VOC variations on photochemical ozone formation were also analyzed using box 29 model simulations. Our results indicate that VOCs exhibited distinct vertical variation 30 patterns due to their differences in sources and chemical reactivities, the vertical 31 variations of various VOC species are strictly regulated by along with the diurnal 32 evolution of the planetary boundary layerPBL. During daytime, reactive VOCs (e.g., hydrocarbons) are rapidly oxidized as they mix upwards, leading accompanied byto the 33 34 formation and accumulation of oxygenated VOCs (OVOCs) in the middle and upper 35 layers.- In additionThis process plays a more significant role in regulating 36 photochemical ozone formation with increasing height. In the lower layer, the 37 photochemical formation of ozone responds positively to changes in both NOx and 38 VOCs. As a result, the production rate of ozone decreases declines with height due to 39 the simultaneous decreases in concentrations of reactive VOCs and NOx-due to 40 significant reductions in the concentrations of both NOx and VOCs, but remains high 41 in the middle and upper layers. The strong production of ozone aloft is primarily driven 42 by the presence of high OVOCs concentrations of OVOCs and hydroxyl radicals, which 43 can act as an important source of ozone at ground level. Therefore, careful consideration 44 should be given to the vertical variations in both photochemical ozone production rates 45 and formation regimes in the whole PBL boundary layer when developing regional 46 ozone control strategies.

47 **1** Introduction

48 Volatile organic compounds (VOCs) are crucial constituents of atmospheric 49 chemicals (Li et al., 2022c) and play important roles in regulating the atmospheric 50 oxidation capacity and contributing to the photochemical formation of tropospheric 51 ozone (Zhao et al., 2022; Yang et al., 2024b). Ozone is a major air pollutant in urban 52 environments, with increasing trends reported globally over recent decades (Fleming et 53 al., 2018; Cooper et al., 2020), despite stringent measures to control its precursor 54 emissions (Wang et al., 2020b; Yeo and Kim, 2021; Li et al., 2022b; Perdigones et al., 55 2022). As highlighted in previous studies, reducing emissions of reactive VOCs is key 56 to controlling ozone pollution at present and in the foreseeable future (Zhao et al., 2022; 57 Wang et al., 2024).

58 The primary prerequisite for effective regional ozone pollution control is the 59 determination of the photochemical ozone formation regime (Souri et al., 2020; Zhao 60 et al., 2022), which facilitates the development of reduction schemes for key precursor 61 emissions (Ou et al., 2016; Wang et al., 2019). The main challenges in controlling 62 ozone pollution stem from the complex compositions of its precursors (e.g., VOCs and 63 NOx) in ambient air (Guo et al., 2017; Wu et al., 2020; Li et al., 2022c), as well as the 64 complicated responses of photochemical ozone formation to changes in these 65 precursors (Shao et al., 2021; Perdigones et al., 2022). Furthermore, the interactions 66 between vertical variations of ozone precursors and ozone formation remain unclear 67 (Tang et al., 2017; Sun et al., 2018; Li et al., 2024), adding to the complexity of ozone 68 pollution control.

In most cases, the identification of key ozone precursors has been conducted using ground-level observations (*Qi et al., 2021; Lu et al., 2022*) or compiled source emission inventories (*Ou et al., 2015; An et al., 2021; Wang et al., 2022b*). While these methods are undoubtedly helpful in determining key ozone precursors and corresponding reduction strategies, they often encounter unexpected uncertainties in urban regions. (*Mo et al., 2018; Mo et al., 2020*). Consequently, ground-level measurements of ozone precursors have been favored to constrain model calculations (*Lu et al., 2012; Wang et al., 2022a; Yang et al., 2022*) or provide empirical evidence for hypothesized theories (*Hofzumahaus et al., 2009; Wang et al., 2022c*). However, these ground-level measurements cannot fully characterize atmospheric chemical processes in the entire planetary boundary layer (PBL) due to strong vertical variations in precursor concentrations (*Velasco et al., 2008; Li et al., 2018; Sun et al., 2018*).

81 Ambient VOCs, as crucial ozone precursors, are composed of myriad species (Wu 82 et al., 2020; Gkatzelis et al., 2021; Ye et al., 2021; He et al., 2022) and serve diverse 83 functions in photochemical ozone formation (Vo et al., 2018; Li et al., 2022a; Zhang et 84 al., 2022). Owing to the impact of variations in emission sources, chemical removal, advection and convection transport, and secondary formation, the concentration and 85 86 composition of VOCs typically display notable vertical variability within the PBL, 87 especially in urban areas (Li et al., 2022c). The ozone formation regime likely 88 undergoes significant transitions from the ground to the upper boundary layer (Li et al., 89 2024; Liu et al., 2024a). Ozone generated throughout the PBL can influence surface 90 ozone levels due to enhanced atmospheric vertical mixing during the day. Consequently, 91 it is imperative to comprehend the vertical variations and principal determinants of 92 VOCs, as well as their effects on photochemical ozone formation within the PBL.

93 With the rapid development of cities in the recent two decades in China, a large 94 number of pollution-emitting industries and factories have been relocated from city 95 centers to alleviate air pollution. Concurrently, there's been a swift increase in the 96 ownership of electric vehicles (Guo et al., 2021). These shifts in energy consumption 97 have driven the change in concentrations and compositions of VOCs in major cities like 98 Beijing (Liu et al., 2024b), subsequently affecting photochemical ozone formation 99 (Wang et al., 2024). However, the vertical variations and key drivers of VOCs and their 100 impacts on photochemical ozone formation in the urban PBL remain elusive. A primary 101 hurdle in studying these vertical changes in photochemical ozone formation is the 102 scarcity of reliable vertical VOC data (Dieu Hien et al., 2019; Li et al., 2022c). 103 Engaging in vertical profiling of VOCs, ensuring all necessary species represented and 104 obtaining sufficient sample size, is especially challenging in the lower PBL where 105 atmospheric chemical reactions are most intense *(Benish et al., 2020; Kim et al., 2021)*.

106 Previous studies on vertical distributions of photochemical ozone formation in the 107 PBL have been conducted using measurements of a limited number of VOC species and samples (Zhang et al., 2018; Benish et al., 2020; Geng et al., 2020). In this study, 108 109 online gradient measurements of ozone, NOx, and a large suit of VOCs were made on 110 a 325 m tall tower in urban Beijing during the summer of 2021. Additionally, box model 111 simulations constrained by the gradient measurements were performed to analyze the 112 vertical variations and key drivers of VOCs as well as their impacts on photochemical 113 ozone formation.

114 **2** Methods and materials

115 **2.1 Description of the site, instrument, and field campaign**

116 The data utilized in this study was derived from an intensive field campaign 117 conducted at the Beijing Meteorological Tower (BMT: 39°58' N, 116°23' E) between 118 July 6 and August 4, 2021. The BMT has a height of 325 m and is located in the northern 119 part of downtown Beijing, positioned between the third and fourth ring roads (Fig. S1). 120 A vertical observation system, established using long perfluoroalkoxy alkane (PFA) 121 Teflon tubes (OD: 1/2 in.), was used to make online gradient measurements of ozone, 122 NOx, and a set of VOCs on the BMT. Five specific heights, namely 15, 47, 102, 200, 123 and 320 m above ground level, were selected to mount the tube inlets, as depicted in 124 Fig. S2S3. An additional inlet, situated approximately 5 m above ground level, was 125 mounted on the rooftop of the observation room that was adjacent to the tower. 126 Consequently, the vertical observation system totally included a total of six sampling 127 inlets. The sampling inlet at the 15 m height was not utilized during this field campaign.

Filters were installed downstream of the tubing inlets on the tower to remove fine particles. A rotary vane vacuum pump was used to simultaneously and continuously draw sample air from the five tubes, ensuring that all tubes were flushed by ambient air 131 to reduce tubing delays of sticky organic compounds (Pagonis et al., 2017; Liu et al., 132 2019). Five critical orifices were employed to control the flow rate of the air stream in 133 each tubing, resulting in flow rates ranging between 15 and 20 standard liter per minute 134 (SLPM). Instruments drew sample air from the five tubes sequentially through a Teflon 135 solenoid valve group at designated time intervals. The switching time intervals of the 136 Teflon solenoid valve group were set as 4 minutes during this field campaign. The 137 measurements of trace gases in the first and last 1 minute of a four-minute period were 138 discarded to eliminate cross interferences between different inlet heights. Detailed 139 information on the vertical observation system and the assessment of trace gas 140 measurements through hundreds of meters long PFA tubes has been provided in our 141 previous works (Li et al., 2023; Song et al., 2024; Yang et al., 2024a).

142 Ozone was measured using the ultraviolet photometry method (49i, Thermo Fisher 143 Scientific Inc., USA). NO, NO₂, and NOx were measured using the chemiluminescence 144 method (42i, Thermo Fisher Scientific Inc., USA). Gradient measurements of ozone 145 and NOx were conducted at a time resolution of 10 seconds. The photolysis frequencies 146 of NO₂, represented by $i(NO_2)$, were measured by a spectrometer (PFS-100, Focused 147 Photonics Inc., China) situated on the rooftop of the observation room and have a time 148 resolution of 8 seconds. In situ measurements of meteorological parameters including 149 wind speed, air temperature, and relative humidity were made at 15 heights between 8 150 m and 320 m on the BMT with a time resolution of 20 seconds. Planetary boundary 151 layer height (PBLH) was obtained from the Air Resources Laboratory 152 (https://ready.arl.noaa.gov/READYamet.php, last access: 10 June 2024) and was 153 linearly interpolated to hourly values based on the initial time resolutions of three hours 154 (Li and Fan, 2022).

155 A high-resolution proton-transfer-reaction quadrupole interface time-of-flight 156 mass spectrometer (PTR-ToF-MS, Ionicon Analytik, Austria) was employed to measure 157 VOCs at a time resolution of 10 seconds. The PTR-ToF-MS used both hydronium ion 158 (H_3O^+) (Yuan et al., 2017; Wu et al., 2020; Li et al., 2022c) and nitric oxide ion (NO⁺) 159 (Wang et al., 2020a) as reagent ions. These two reagent ions were automatically

160 switched every 60 min for H_3O^+ and every 22 min for NO^+ throughout the campaign. 161 The PTR-ToF-MS operated at an E/N value of approximately 120 Td in H_3O^+ mode and an E/N value of around 60 Td in NO⁺ mode. Instrument backgrounds were 162 163 automatically measured during the last two minutes of each operation mode by passing 164 ambient air through a platinum catalyst heated to 365 °C. A gas standard containing 39 165 VOC species was used to calibrate the PTR-ToF-MS daily. Sensitivities for the 166 remaining species were determined based on reaction kinetics of the PTR-ToF-MS (Wu 167 et al., 2020). Impacts of ambient humidity on the PTR-ToF-MS measurements were 168 corrected by using humidity-dependence curves of VOCs obtained in our laboratory 169 (Wang et al., 2020a; Wu et al., 2020). Carbon dioxide (CO₂ in dry air) and humidity 170 were measured using a CO2 and H2O gas analyzer (Li-840A, Licor Inc., USA) at a time 171 resolution of 10 seconds.

172 Gradient measurements of the total OH reactivity (OHR) of atmospheric trace 173 gases were made using the improved comparative reactivity method (ICRM) developed 174 by our team (Wang et al., 2021a) from July 28 to 31. In addition, gradient measurements 175 of carbon monoxide (CO), methane (CH₄), CO₂, and H₂O were simultaneously 176 measured using the cavity ring-down spectroscopy (CRDS) method (G-2401, Picarro 177 Inc., USA) at a time resolution of 10 seconds from May 15 to June 25. Sulfur dioxide 178 (SO₂) was measured using the ultraviolet fluorescence method (43i, Thermo Fisher 179 Scientific Inc., USA) at a time resolution of 10 seconds from June 25 to August 3. The 180 total OHR of VOCs, denoted by OHR_{VOCs}, can be estimated by excluding those of the 181 inorganic species (namely ozone, NOx, CO, SO₂, and CH₄). It should be noted that 182 gradient measurements of CH₄ and CO were not made during July 28-31, and their 183 average concentrations in daytime (11:00-16:00 LT) between May 15 and June 25 at 5 184 m were used for all altitudes to calculate OHR_{VOCs}. This method will bring minor 185 uncertainties due to the minor vertical differences in concentrations of CH₄ and CO in 186 daytime (Fig. <u>\$3</u>\$4). The OHR of VOCs can also be calculated by summing the 187 products of their measured concentrations and their reaction rate coefficients with OH 188 radicals, as formulated in Eq. (1):

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$$OHR = \sum k^{i}{}_{OH-R}[VOC_{i}]$$
 Eq. (1)

189 where k^i_{OH-R} is the reaction rate coefficient of the *i*th VOC species with OH radical 190 and $[VOC_i]$ is the concentration of the *i*th VOC species.

191 **2.2 Estimation of NMHC concentrations at the BMT site**

192 The PTR-ToF-MS is limited in its ability to measure VOC species with proton affinities higher than H₂O (691 kJ mol⁻¹) when operating in the H₃O⁺ mode (Yuan et 193 194 al., 2017). This limitation results in the absence of certain nonmethane hydrocarbons 195 (NMHCs), such as alkanes and many alkene species, which play important roles in 196 photochemical ozone formation. To obtain a comprehensive understanding of vertical 197 variations in concentrations, compositions, and environmental impacts of VOCs, this 198 study estimated the vertical profiles of those unmeasured NMHC species based on the 199 concentrations of measured VOCs using the PTR-ToF-MS. Detailed information on 200 estimation of NMHC concentrations is provided in SI.

201 2.3 Box model setup

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A zero-dimension box model (F0AM) coupled with the Master Chemical Mechanism (v3.3.1) *(Wolfe et al., 2016; Yang et al., 2022)* was used to compute the production rate of ozone, denoted by P(O₃) as formulated in Eq. (2):

$$P(O_3) = k_{HO_2+NO}[HO_2][NO] + \sum k^i_{RO_2+NO}[R^iO_2][NO]$$
Eq. (2)
$$k_{RO_2+NO}[RO_2][NO]$$

where $[HO_2]$ and [NO] is the concentrations of HO_2 and NO, $[R^iO_2]$ is the concentration of the *i*th organic peroxyl radical. The relative incremental reactivity (RIR) of photochemical ozone production to changes in different precursors was determined using Eq. (3):

$$RIR(X) = \frac{\left[P_{O_{X}}^{S}(X) - P_{O_{X}}^{S}(X - \Delta X)\right] / P_{O_{X}}^{S}(X)}{\Delta S(X) / S(X)}$$
Eq. (3)

where X represents ozone precursors, $P_{O_X}^S(X)$ is the contribution of X to the production rate of Ox, ΔX is the amount of change in ozone precursors, S(X) is the initial concentration of X. RIR values were used to discern sensitivities of photochemical ozone formation to changes in precursor gases. A positive RIR(X) value suggests that an increase in X enhances ozone formation, while a negative RIR value indicates that an increase in X inhibits ozone formation.

216 Model calculations were constrained by measurements of ozone, NOx, CO, a suit 217 of VOCs, air temperature, and relative humidity. In addition to the measured or 218 estimated concentrations of NMHCs, nine oxygenated VOC (OVOC) species (Table 219 S1) measured by PTR-ToF-MS were used to constrain the model calculation. The 220 model was run in a time-dependent mode with a time resolution of 5 minutes and a 221 spin-up period of 2 days (Lu et al., 2012; Wang et al., 2022c). The dry deposition 222 velocity of ozone was set as 0.27 cm s^{-1} when calculating P(O₃) 5 m and was zeroed 223 out when calculating $P(O_3)$ at other heights.

224 **3 Results and discussions**

225 **3.1** Temporal and vertical variations in concentrations of trace gases

226 As shown in Fig. 1, the meteorology in Beijing was characterized by high air 227 temperature (27.3±2.9 °C), high humidity (83.9%±16.2%), and gentle winds (1.1±0.4 m s⁻¹) throughout the campaign. The intense solar radiation, elevated air temperature, 228 229 and mild winds favored the photochemical formation and accumulation of ozone, 230 leading to frequent occurrences of ozone pollution episodes. Fig. 1 also presents time 231 series of mixing ratios of ozone and its selected precursors (namely isoprene, toluene, 232 monoterpenes, and NOx) along with $i(NO_2)$ measured at 5 m. The campaign mean ozone mixing ratio was 45.6±25.3 ppb, but the maximum hourly mean ozone mixing 233 ratio reached 129.3 ppb, indicating strong photochemical reactions in urban Beijing 234 235 during the campaign. Surface ozone concentrations exhibited a typical diurnal variation pattern with the maximum occurring at 16:00 LT (Fig. <u>\$5</u><u>\$6</u>), implying its predominant
source from local photochemical production.

238 Isoprene is a typical tracer of biogenic emissions and is also a highly reactive VOC 239 species (Atkinson and Arey, 2003). Isoprene had a campaign mean mixing ratio of 240 0.7±0.6 ppb. The average diurnal profile of isoprene at 5 m has a unimodal pattern with 241 the maximum occurring at 14:00 LT (Fig. <u>\$556</u>), exhibiting strong dependence on solar 242 radiation. Monoterpenes were also generally recognized as typical tracers of biogenic 243 emissions (*Gómez et al., 2020*) and have a campaign mean mixing ratio of 0.3 ± 0.3 ppb. 244 The average diurnal profile of monoterpenes was characterized by low mixing ratios in 245 daytime with two peaks occurring at 05:00 and 20:00 LT, respectively.

246 Toluene and NOx are recognized as typical tracers of anthropogenic emissions in 247 urban regions (Niu et al., 2017; Li et al., 2022c), with campaign mean mixing ratios of 248 0.7±0.7 and 8.1±4.8 ppb, respectively. The average diurnal profiles of toluene and NOx 249 at 5 m exhibited similar variations with larger values at night than during the day. Based 250 on the measured concentrations and diurnal variations of ozone and its key precursors 251 at ground level, it can be inferred that urban Beijing is experiencing severe ozone 252 pollution, which is predominantly contributed by local photochemical production. As 253 key ozone precursors, ambient concentrations of VOCs are contributed by the mixture of anthropogenic and biogenic sources. 254

255 Fig. 2 shows the average diurnal and vertical variations in mixing ratios of ozone, 256 NOx, Ox (O₃+NO₂), and six selected VOC species (three hydrocarbons and three 257 OVOCs) within the measurement height range of 5-320 m. High mixing ratios of ozone 258 were observed in the afternoon following the enhancement of solar radiation, which 259 was consistent with the diurnal change pattern of ozone concentrations at the ground 260 level. The vertical gradients of ozone mixing ratios were positive throughout the day 261 but substantially enhanced at night (Fig. 3). The lower ozone mixing ratios near the 262 surface than aloft were mainly caused by the enhancement of dry deposition and NO 263 titration (Brown et al., 2007; Ma et al., 2013; Li et al., 2022b).

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264 NOx is a primary pollutant and mainly contributed by vehicular exhausts in urban 265 regions. In contrast to ozone, NOx mixing ratios were low in daytime and exhibited 266 negative vertical gradients throughout the day, as shown in Figs. 2B and 3A-B. In 267 nighttime, large amounts of local NOx emissions were trapped and accumulated in a 268 shallow boundary layer (<100 m). NOx concentrations rapidly decreased with height 269 even in the overlying residual layer due to the suppression of turbulence vertical mixing. 270 With the onset of sunlight, the PBL rapidly expanded due to the surface heating effect. 271 The accumulated high concentrations of NOx in the shallow nocturnal boundary layer 272 were thereupon diluted and removed by photochemical reactions.

273 Ox is frequently used as a conserved metric to investigate temporal and spatial 274 variability of ozone by eliminating the NO titration effect. As shown in Fig. 2C, the 275 mixing ratios of Ox had similar diurnal and vertical variation patterns to those of ozone, 276 but the vertical gradients of Ox were weaker than those of ozone. This result suggests 277 that the vertical distribution of NO concentrations played an important role in regulating 278 the vertical change of ozone concentrations. The enhanced positive gradients of ozone 279 mixing ratios at night were predominantly due to the strict suppression of turbulence 280 vertical mixing (Geyer and Stutz, 2004). The higher concentrations of ozone aloft are 281 considered as the residual of the ozone produced in the daytime PBL and have been 282 recognized as an important reservoir for the enhancement of surface ozone in morning 283 periods (Kaser et al., 2017; Li and Fan, 2022; He et al., 2023).

284 Benzene and toluene demonstrated similar diurnal and vertical variations to NOx, 285 with low concentrations in daytime and high concentrations at night, as shown in Figs. 286 2D-F and 3A-B. The concentrations of both benzene and toluene decreased with height 287 throughout the day, confirming their primary emissions from ground-level sources. 288 However, unlike benzene, the diurnal and vertical variations of toluene were more 289 pronounced. Isoprene emissions are highly dependent on solar radiation, resulting in its 290 higher concentrations in the early afternoon compared to other times of the day. 291 Isoprene mixing ratios also exhibited strong negative vertical gradients below 320 m 292 throughout the day. In contrast to toluene, isoprene concentrations decreased more

rapidly with height in the daytime. For instance, the mixing ratios of isoprene decreased
by approximately 70% from 5 to 320 m in the daytime, while it was only 30% for
toluene.

296 Fig. 3A-B show the average vertical profiles of the NMHCs, normalized to their 297 respective ground-level concentrations measured by the PTR-ToF-MS in daytime and 298 nighttime. The normalized mixing ratios of the NMHCs exhibited significantly 299 differentiated gradients in daytime. In contrast, apart from monoterpenes, the 300 differences in vertical gradients of the normalized vertical profiles for other NMHCs 301 were relatively small at night. The differentiated vertical gradients of the NMHCs in 302 daytime were primarily caused by their intrinsic chemical reactivities, such as reactions 303 with OH radicals. As shown in Fig. 4, concentration ratios of the NMHC species between 320 m and 5 m with k_{OH} values lower than $2.5 \times 10^{-11} \text{ cm}^{-3}$ molecule⁻¹ s⁻¹ 304 305 exhibited slight variability and rapidly declined with the further increases in k_{OH} . The 306 lower NMHC concentrations at higher altitudes were predominantly caused by the 307 combined effects of atmospheric diffusion and chemical removal (Sangiorgi et al., 308 2011).

309 Considering the effects of atmospheric diffusion and chemical removal by 310 reactions with OH radicals, concentration ratios of NMHC species between 320 m and 311 5 m in daytime can be estimated using Eq. (4):

$$y = A \times \exp(-k_{OH}[OH]\Delta t) \qquad \qquad \text{Eq. (4)}$$

312 where y represents concentration ratios of the NMHC species between two altitudes, A 313 represents the effect of atmospheric dilution, k_{OH} is the reaction rate coefficient of 314 NMHCs with OH radicals, [OH] is the concentration of OH radical, Δt is the 315 turbulence mixing time scale between the two altitudes. The term $[OH]\Delta t$ thus refers 316 to the exposure of NHMCs to OH radicals between the two altitudes. As shown in Fig. 317 4, the average concentration ratios of NMHCs between 320 m and 5 m in daytime 318 during the campaign can be well reproduced using Eq. (4) with the coefficients A of 0.88 and $[OH]\Delta t$ of 1.0×10^{10} molecules cm⁻³ s. Atmospheric diffusion processes have 319 320 same impact on the vertical distributions of all trace gases. The differences in vertical

gradients of NMHCs were mainly determined by the differences in their chemicalremoval rates without considering influences from advection transport.

323 Methanol, as one of the most abundant OVOC species in the atmosphere, had its 324 lowest concentrations during daytime and displayed negative vertical gradients 325 throughout the day, as shown in Fig. 2G. The vertical and diurnal variations of methanol 326 suggest that its ambient concentrations in urban Beijing were mainly contributed by 327 local primary emissions. Conversely, formaldehyde and MVK+MACR (the firstgeneration oxidation products of isoprene), as the photochemical oxidation products of 328 329 NMHCs, had higher concentrations during daytime than at night and exhibited 330 relatively weak vertical concentration gradients (Fig. 2H-I). This is mainly because 331 these OVOCs are produced from the oxidation of NMHCs during turbulence vertical 332 mixing and will accumulate in high altitudes. These phenomena were also observed for 333 other OVOC species, as shown in Fig. 3.

334 The vertical and diurnal variations in concentrations of ozone, NOx, and VOCs are intricately governed by their sources, chemical reactivities, and the evolution of the 335 336 PBL (namely the vertical dilution conditions). A significant accumulation of VOCs in 337 the shallow nocturnal PBL is subsequently vertically diluted and chemically removed 338 during daytime, thereby impacting the photochemical formation of ozone within the daytime PBL. In addition, the observed vertical changes in concentrations of VOCs 339 340 imply that they will play distinct roles in contributing to photochemical ozone formation. 341

342 **3.2** Vertical variations in contributions of VOCs to OHR

During the daytime, VOCs are primarily oxidized by OH radicals and contribute to the photochemical formation of ozone. To provide an overview on the vertical variations in contributions of different VOCs to OHR, another 1204 ions measured by the PTR-ToF-MS and can be quantified were used for analysis. All the VOCs were classified into three large categories, namely C_xH_y (including alkanes, alkenes, aromatics, and other hydrocarbons; 121 species), OVOCs ($C_xH_yO_1$, 121 species; 349 $C_xH_yO_2$, 120 species; $C_xH_yO_{\geq 3}$, 256 species), and N/S-containing (653 species), as 350 shown in Fig.5. Acetylene is included in alkenes.

351 Fig. 5A illustrates that the total mixing ratios of VOCs in daytime exhibited a slight 352 downward trend from 5 m to 320 m, primarily due to the rapid decrease in mixing ratios 353 of the C_xH_y category. The total mixing ratios of the C_xH_y category decreased from 16.8 354 to 10.6 ppb from 5 m to 320 m, with alkanes making the largest contribution, followed 355 by alkenes, aromatics, and other C_xH_y. Alkanes constituted 58% of the total mixing ratios of C_xH_y at 5 m, but this proportion increased to 65% at 320 m. The fractional 356 357 contributions of alkenes and aromatics in the total mixing ratios of C_xH_y slightly 358 declined from 28% to 22% and from 12% to 10%, respectively, between these two 359 altitudes. As for OVOCs, the C_xH_vO₁ category was the most abundant among the 360 measurements, contributing to 52%-58% of the total mixing ratios at the five heights, 361 followed by the $C_xH_yO_2$ (8%-10%), and $C_xH_yO_{\geq 3}$ (2%) categories. The mixing ratios of the N/S-containing category slightly varied around 2.8 ppb between 5-320 m, 362 363 contributed to approximately 6% of the total VOC concentrations.

364 Similar to the vertical variations in concentrations, OHRs of the C_xH_y category, denoted by OHR_{CH}, also rapidly decreased from 6.9 s⁻¹ to 2.5 s⁻¹ between 5 and 320 m, 365 366 accounting for 52%-31% in the total OHRs of VOCs (Fig. 5B). Fractional contributions 367 of alkenes (40-18%), alkanes (5%), and aromatics (5%-4%) to the total OHRs of VOCs 368 all exhibited decreasing tendencies from 5 m to 320 m. The total OHRs of alkenes 369 decreased more quickly from 5 to 320 m than those of alkanes and aromatics. OHRs of the other C_xH_y category stabilized at approximately 0.3 s⁻¹ below 320 m, exhibiting an 370 371 increasing contribution (2%-4%) to the total OHRs of VOCs with the increase in height. 372 The OHRs of other VOC categories only slightly varied without exhibiting a clear 373 variation trend from 5 to 320 m during the day. As a result, fractional contributions of the $C_xH_vO_1$ (27%-42%), $C_xH_vO_2$ (12%-18%), and $C_xH_vO_{>3}$ (5%-7%), and N/S-374 375 containing (2%-4%) categories in the total OHRs of VOCs all increased with height. The increased contributions of OVOCs and N/S-containing species to the total 376

377 concentrations and OHRs of VOCs implied that air masses became more aged with the378 increase in height.

379 As depicted in Fig. 6A-B, high OHR_{CH} values were mainly constrained in the PBL 380 and is mainly contributed by biogenic hydrocarbons, specifically isoprene, during 381 daytime due to their high OH reactivities and enhanced emissions. The fractional 382 contributions of isoprene in OHR_{CH} decreased rapidly with increasing height (Fig. 7A). 383 For instance, isoprene accounted for a campaign median fraction of 58% in OHR_{CH} at 384 5 m in daytime, making it a frequent contributor to photochemical ozone formation in 385 urban regions. However, this faction decreased to 38% at 320 m. Therefore, it can be 386 speculated that the total contributions of hydrocarbons to the total OHRs of VOCs will 387 also rapidly decline from 320 m to the top of the PBL, which typically ranges between 388 several hundreds of meters to approximately 2~3 km in daytime (Fig. <u>\$6</u>\$7).

389 The total concentrations and OHRs of OVOCs only slightly decreased with the 390 increase in height below 320 m in daytime, as shown in Fig. 6C-D. This is consistent 391 with the results of (Wang et al., 2021b), which observed high concentrations of OVOCs 392 in the upper PBL. Consequently, the ratio of OHROVOC to OHRCH, denoted by 393 OHR_{OVOC}/OHR_{CH}, rapidly increased from 0.87 at 5 m to 2.6 at 320 m (Fig. 7A). This 394 suggests that OVOCs may play more important roles in regulating the photochemical 395 ozone formation in the middle and upper layers. To assess their potential roles in 396 contributing to the photochemical ozone formation throughout the PBL, we calculated 397 the mean OHRs (MOHR) of different VOC categories in daytime using Eq. (5):

$$MOHR(X) = \left(\sum ([X]_i + [X]_{i-1})(h_i - h_{i-1})/2\right)/(320 - 5)$$
 Eq. (5)

398 where MOHR(X) is the MOHR of the VOC category X, $[X]_i$ is the concentration of 399 X at the *i*th altitude (namely 5, 47, 102, 200, and 320 m for h_i) above ground level.

400 As shown in Fig. 7B, the campaign median MOHR for isoprene was 1.7 s^{-1} and 401 accounted for 48% of the campaign median MOHR of the C_xH_y category. This fraction 402 was significantly lower than that of isoprene (57%) in OHR_{CH} at 5 m. In addition, the 403 campaign median MOHR of the C_xH_y category (3.5 s⁻¹) was also significantly lower 404 than the OHR_{CH} (6.0 s⁻¹) at 5 m. By contrast, the campaign median MOHR of OVOCs (4.8 s⁻¹) was comparable to that of OHR_{OVOC} (4.9 s⁻¹) at 5 m. As unsaturated 405 hydrocarbons, most alkene species are more reactive than alkanes and aromatics 406 407 (Atkinson and Arey, 2003). As a result, alkenes had dominant contributions to the 408 MOHR of the C_xH_y category and the OHR_{CH} at 5 m in daytime. As shown in Fig. 7C, 409 the campaign mean OHRs of alkanes, alkenes, and aromatics at 5 m in daytime were 410 0.7, 5.2, and 0.7 s⁻¹, respectively, accounting for 10%, 75%, and 10% of the OHR_{CH}. However, the campaign mean MOHRs of alkanes, alkenes, and aromatics were 0.5, 2.7, 411 412 and 0.5 s⁻¹, respectively, accounting for 12%, 68%, and 12% of the MOHR of NMHC. We can also expect that the total contributions of alkenes to the MOHR of the C_xH_y 413 414 category in daytime will significantly decrease if their vertical distributions in the whole 415 PBL are considered.

416 This study investigated and compared the vertical profiles of measured OHR_{VOCs} and calculated OHR_{CH} during daytime over the period of July 28-31, as shown in Fig. 417 7D. The campaign median of the measured OHR_{VOCs} exhibited a slow decrease from 418 38.4 s⁻¹ at 5 m to 25.4 s⁻¹ at 320 m. As anticipated, the OHR_{CH}/OHR_{VOCs} ratio declined 419 420 rapidly from 16% to 7% from 5 to 320 m. It is important to note that the small 421 OHR_{CH}/OHR_{VOCs} ratio and its declining trend with the increasing height do not imply 422 the insignificant roles of hydrocarbons in regulating the secondary pollutant formation 423 in higher altitudes. The measured concentrations of hydrocarbons are merely the 424 remnants of chemical reactions. The oxidation products of NMHCs, such as OVOCs 425 and organic nitrates, formed during vertical mixing in daytime, will continue to 426 participate in atmospheric chemical reactions.

427 **3.3** Vertical variations in photochemical ozone formation

428 The surface ozone budget is intimately linked to the vertical variations of 429 photochemical ozone formation throughout the PBL. Previous studies have consistently 430 reported that the photochemical formation of ozone, encompassing both $P(O_3)$ and 431 ozone formation regimes (namely the NOx-limited, VOCs-limited, and transition regimes), are highly dependent on the change in its precursors (*Shao et al., 2021; Yang et al., 2022*). Consequently, any changes in the concentrations and compositions of
VOCs and NOx within the PBL will inevitably lead to alternations in the vertical
distribution of P(O₃) and ozone formation regimes (*Tang et al., 2017; Li et al., 2024*).

436 Fig. 8A illustrates the average dependence of $P(O_3)$ on NOx concentrations along 437 with the normalized probability density (NPD) distribution of NOx concentrations at 5 438 m, 200 m, and 320 m in daytime during the field campaign. At different heights, P(O₃) 439 all rapidly increased with the rise in NOx until a critical NOx mixing ratio was reached, 440 after which $P(O_3)$ decreased slowly. The critical NOx mixing ratios decreased from 441 approximately 9.5 ppb at 5 m to 5.0 m ppb at 320 m, primarily caused by the decreases in both NOx concentrations and the OHRs of VOCs. As also shown in Fig. 8A, the 442 443 majority of the measured NOx mixing ratios fall into the transition zone of the $P(O_3)$ -444 NOx curves, suggesting that the photochemical ozone formation in Beijing belonged to 445 the transition regime below 320 m.

446 RIR values were also calculated using the box model results to further elucidate 447 the sensitivities of photochemical ozone formation to changes in multiple precursors at 448 different altitudes. As shown in Fig. 8B, positive RIR values were observed for both 449 NOx and various VOC groups at the five heights, further confirming that the 450 photochemical ozone formation belonged to the transition regime in the lower layer. 451 RIR values for NOx rapidly declined from 5 to 320 m, implying that the photochemical 452 ozone formation in higher altitudes were was more prone to be controlled by the 453 abundance of VOCs. This is also manifested by the increasing RIR values for both 454 AVOCs and OVOCs from 5 m to 320 m. RIR values for BVOCs significantly decreased 455 with height due to their rapid removal by reactions with OH radicals when being 456 vertically mixed. These results are consistent with the results in section 3.3 that the less 457 reactive AVOCs and OVOCs are the dominant species in regulating the photochemical 458 formation of ozone in urban regions aloft.

459 According to the vertical distribution patterns of the photochemical ozone 460 formation regime, $P(O_3)$ decreases with increasing height alongside simultaneous 461 declines in concentrations of both NOx and VOCs. Fig. 8C presents the average diurnal 462 and vertical variations in $P(O_3)$ calculated by the box model during the campaign. The 463 $P(O_3)$ values were higher in daytime and correlated well with $j(NO_2)$. As shown in Fig. 464 S7, OH radical concentrations and $P(O_3)$ exhibited contrasting vertical distribution 465 patterns in daytime. $P(O_3)$ decreased from the ground to 320 m, where it still maintained a relatively high value of approximately 10 ppb h⁻¹ at noon. These results highlight that 466 467 the photochemical formation of ozone aloft also remained strong compared to those at 468 ground level. Consequently, the downward transport of ozone from high altitudes, 469 driven by turbulence mixing, can become significant sources of surface ozone during 470 the day (Karl et al., 2023).

471 Due to the measurement height limitation, the vertical distributions of $P(O_3)$ in the 472 middle and upper parts of the PBL were not determined in this study. As reported by 473 the work in (Benish et al., 2020), P(O₃) typically exhibited weak and nearly linear 474 decline tendencies from 300 m to the top of the PBL during daytime. $P(O_3)$ at the PBL top was approximately half of that at 300 m. Consequently, we can assume that $P(O_3)$ 475 476 decreased linearly from 320 m to the top of the PBL. The integral of $P(O_3)$ at different 477 heights within the PBL can then be estimated using a similar method as described in 478 Eq. (5).

479 As shown in Fig. 8D, the total amount of ozone photochemically produced below 480 47 m constituted a mere 6% of the entire PBL. This fractional contribution increased to 481 approximately 35% at 320 m, further corroborating that the majority of the boundary-482 layer ozone was produced in the middle and upper layers. Given the enhancement of 483 turbulence vertical mixing in daytime, ozone produced at high altitudes becomes a 484 significant source of surface ozone. This is substantiated by the widespread reports of 485 strong downward ozone fluxes in the bottom part of the PBL (tens of meters above ground level) (Fares et al., 2010; Liu et al., 2021; Karl et al., 2023). Consequently, 486 487 when devising ozone control strategies, particularly in urban regions with intricate precursor emissions, careful considerations should be given to the vertical variations in 488 489 the formation regimes of ozone in the PBL.

490 **4** Conclusions

491 In this study, we investigated the vertical variations, key drivers, and 492 environmental impacts of VOCs in the PBL using tower-based online gradient 493 measurements in urban Beijing during the summer of 2021. The diurnal and vertical 494 variations of various VOC species were strictly regulated by the diurnal evolution of 495 the PBL. In daytime, reactive NMHC species were rapidly oxidized when they were 496 mixed upward along with the formation of OVOCs. As a result, OVOC species played 497 more significant roles in regulating the photochemical ozone formation in urban regions 498 aloft. The photochemical formation of ozone belongs to the transition regime in the 499 lower part of the PBL and becomes more sensitive to changes in the concentrations of 500 AVOCs and OVOCs with increasing height. $P(O_3)$ exhibited decreasing tendencies 501 with height but remained very large in high altitudes, likely driven by the high 502 concentrations of OVOCs and OH radicals. Therefore, careful consideration should be 503 given to the vertical variations in both $P(O_3)$ and photochemical ozone formation 504 regimes in the whole PBL when making regional ozone control strategies.

505 The vertical variations in concentrations and compositions of VOCs significantly 506 influence the formation of secondary pollutants. Furthermore, vertical changes in chemical reaction environments (e.g., temperature, humidity, and solar radiation) and 507 508 concentrations of other chemicals (e.g., particulate matters, NOx, ozone) can also 509 impact the degradation pathways of VOCs. These factors also affect the formation 510 pathways and production yields of secondary pollutants. This is particularly crucial for 511 the highly reactive NMHCs in urban areas with complex anthropogenic emissions and 512 is expected to be thoroughly elucidated in future studies.

513 Data availability

514 The observational data used in this study are available from corresponding authors515 upon request.

516 Author contributions

- 517 BY, XBL, and YH designed the research. XBL, BY, YH, XS, JQ, XH, SW, YC,
- 518 QY, YS, YP, GT, JG, and MS contributed to the data collection and data analysis. XBL,
- 519 SY, and BY designed and performed the box model simulations. XBL and BY wrote
- 520 the paper with contributions from all coauthors. All the coauthors discussed the results
- 521 and reviewed the paper.

522 Competing interests

523 The authors declare that they have no conflict of interest.

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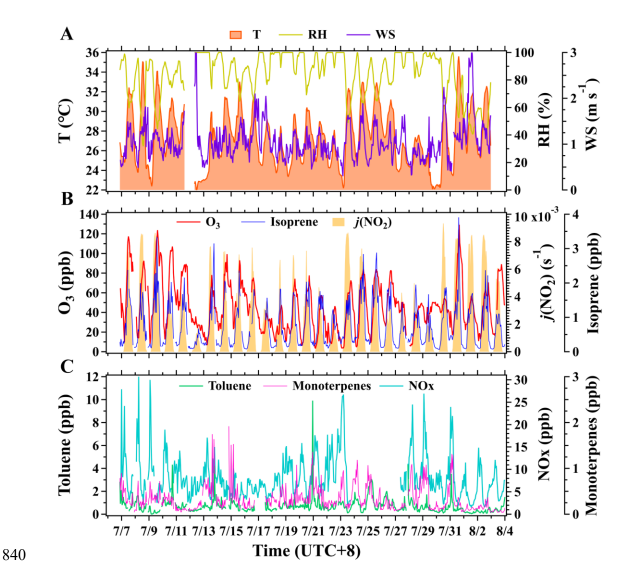


Figure 1. Time series of hourly mean air temperature (T), relative humidity (RH), wind speed (WS), and mixing ratios of surface ozone, NOx, and VOC species along with $j(NO_2)$ at the BMT site during the campaign. Meteorological parameters were measured at 8 m above ground level and mixing ratios of ozone and its selected precursors were measured at 5 m above ground level.

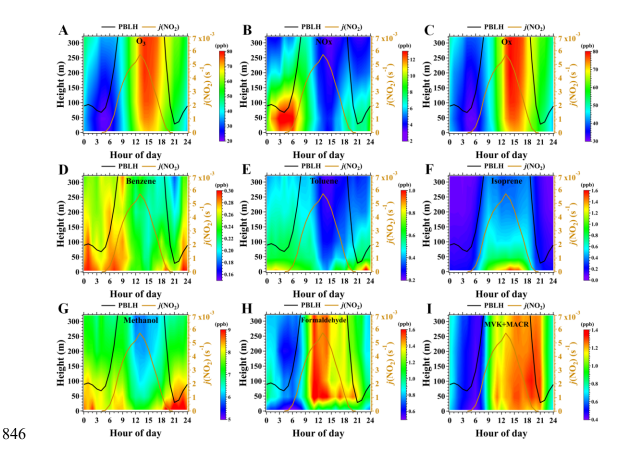


Figure 2. Average diurnal and vertical variations in mixing ratios ozone, NOx, Ox (O_3+NO_2) , and six selected VOC species along with the average diurnal profiles of PBLH and $j(NO_2)$ during the campaign. The figures were obtained by linearly interpolating the data at the five inlet heights on both altitude and temporal scales.

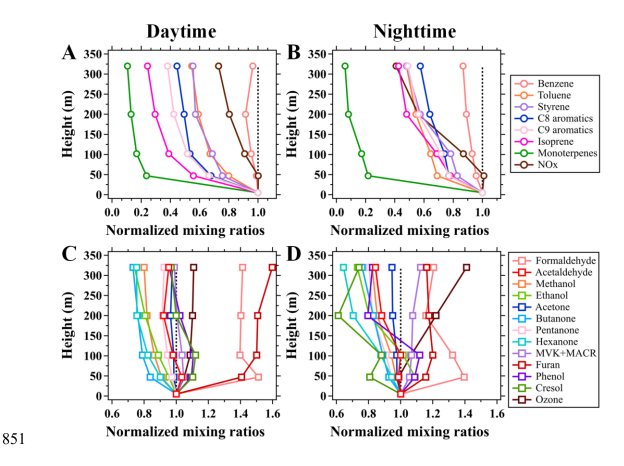
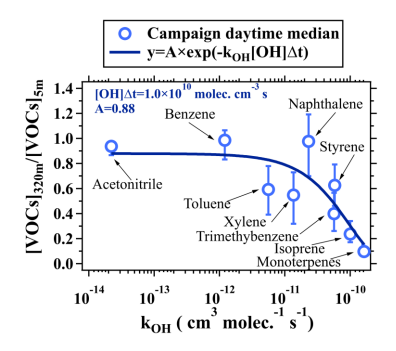
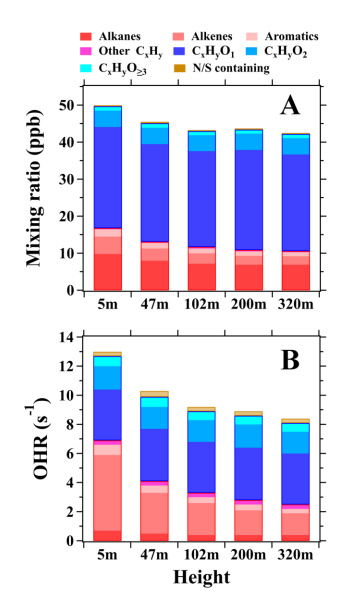


Figure 3. Average vertical profiles of (A-B) NMHCs and NOx, (C-D) OVOCs and O₃
during the daytime (11:00-16:00 LT) and nighttime (23:00-04:00 LT) of the campaign.
The mixing ratios of the chemical species measured above 5 m are normalized to those
at 5 m.



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Figure 4. The change in ratios of NMHC concentrations (including acetonitrile) between 320 m and 5 m as a function of k_{OH} . The vertically-resolved measurements of VOCs made on the BMT in daytime during the campaign were used for analysis. Hollow markers represent median values and error bars indicate the range between 25th and 75th percentiles.



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863 Figure 5. (A) Mean mixing ratios and (B) OHRs of different VOC categories at the five

864 inlet heights in daytime during the campaign.

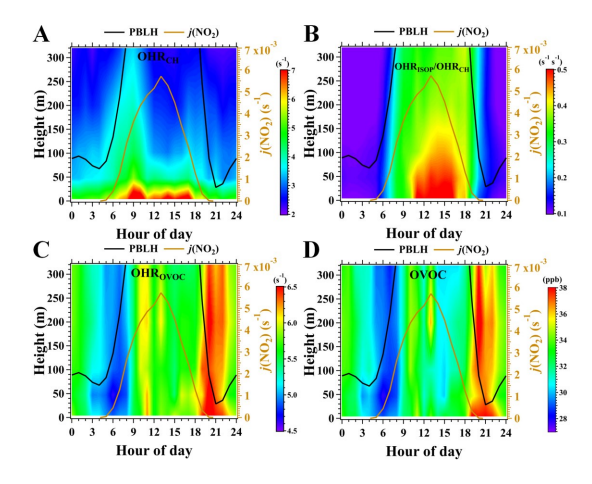
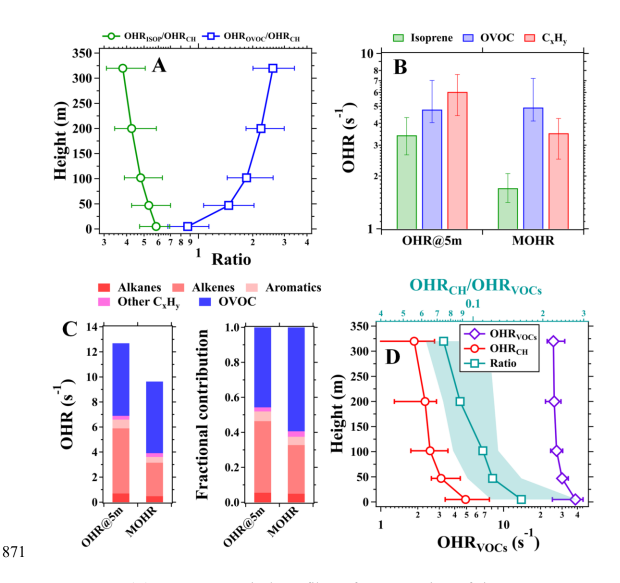
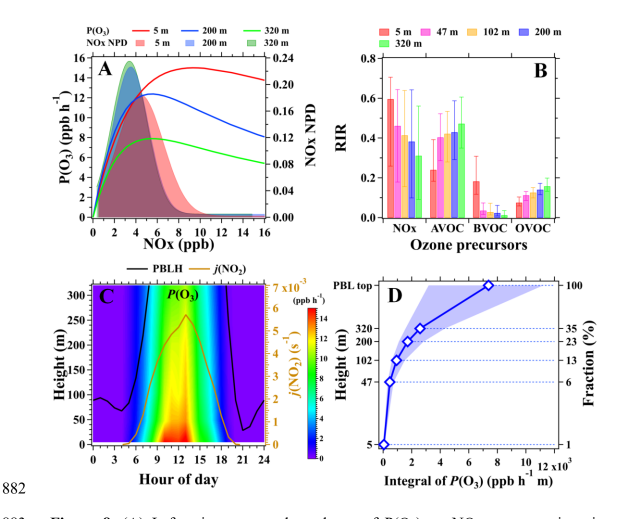


Figure 6. (A-B) Average diurnal and vertical variations in OHRs of C_xH_y and the OHR ratios of isoprene to C_xH_y (OHR_{ISOP}/OHR_{CH}) during the campaign. (C-D) Average diurnal and vertical variations in mixing ratios and OHRs of OVOC. ISOP refers to isoprene. The figures were obtained by linearly interpolating the data at the five measurement heights on both altitude and temporal scales.

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872 Figure 7. (A) Average vertical profiles of OHR ratios of isoprene to C_xH_v (OHR_{ISOP}/OHR_{CH}) and OVOC to NMHC (OHR_{OVOC}/OHR_{CH}). (B) Median values of 873 874 the OHR at 5 m and the mean OHR (MOHR) between 5 m and 320 m for isoprene, OVOC, and C_xH_v. (C) Mean contributions of different VOC categories to the MOHR 875 876 below 320 m and the OHR at 5 m. (D) Vertical profiles of the measured OHR_{VOCs} and the calculated OHR_{CH} (bottom axis) and the OHR_{CH}/OHR_{VOCs} ratios (top axis) during 877 878 July 28-31, 2021. The data used for analysis in panels A-D was within the time window 879 from 11:00 to 16:00 LT during the campaign. Markers in panels A and D represent 880 median values. Shaded areas and error bars in panels A, B, and D indicate the range between 25th and 75th percentiles. 881



883 Figure 8. (A) Left axis: average dependence of P(O₃) on NOx concentrations in 884 daytime during the campaign; Right axis: normalized probability density (NPD) of 885 NOx mixing ratios in daytime at the three inlet heights. (B) Median RIR values of 886 photochemical ozone formation to changes in NOx, AVOC (NMHCs excluding BVOC), 887 BVOC (isoprene), and OVOC (nine OVOC species in Table S1) at the five inlet heights; Error bars indicate the range between 25th and 75th percentiles. (C) Average diurnal and 888 889 vertical variations in $P(O_3)$ during the campaign; The figure was obtained by linearly 890 interpolating the data at the five measurement heights on both altitude and temporal 891 scales. (D) The vertical profile of the integral of $P(O_3)$ in daytime during the campaign; Markers indicate median values and Shaded areas indicate the range between 25th and 892 75th percentiles. 893