



1	Current status of model predictions on volatile organic compounds and impacts
2	on surface ozone predictions during summer in China
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17 Abstract

Volatile organic compounds (VOCs) play a crucial role in the formation of 18 tropospheric ozone (O_3) and secondary organic aerosols. VOC emissions are generally 19 20 considered to have larger uncertainties compared to other pollutants, such as sulphur dioxide and fine particulate matter (PM2.5). Although predictions of O3 and PM2.5 have 21 22 been extensively evaluated in air quality modelling studies, there has been limited reporting on the evaluation of VOCs, mainly due to a lack of routine VOCs 23 measurements at multiple sites. In this study, we utilized VOCs measurements from the 24 25 ATMSYC project at 28 sites across China and assessed the predicted VOCs concentrations using the Community Multiscale Air Quality (CMAQ) model with the 26 widely used Multi-resolution Emission Inventory for China (MEIC). The ratio of 27 28 predicted to observed total VOCs was found to be 0.74 ± 0.40 , with underpredictions ranging from 2.05 to 50.61 ppbv (5.77% to 85.40%) at 24 sites. A greater bias in VOCs 29 predictions was observed in industrial cities in the north and southwest, such as Jinan, 30 31 Shijiazhuang, Lanzhou, Chengdu, and Guiyang. In terms of different VOC components, alkanes, alkenes, non-naphthalene aromatics (ARO2MN), and alkynes were 32 consistently underpredicted, with ratios of predicted to observed of 0.53 ± 0.38 , $0.51 \pm$ 33 $0.48, 0.31 \pm 0.38$, and 0.41 ± 0.47 , respectively. Sensitivity experiments were conducted 34 35 to assess the impact of the VOCs prediction bias on O₃ predictions. While emission adjustments improved the model performance for VOCs, resulting in a ratio of total 36 37 VOCs to 0.86 ± 0.47 , they also exacerbated O₃ overprediction relative to the base case by 0.62% to 6.27% across the sites. This study demonstrates that current modelling 38 setups and emission inventories are likely to underpredict VOCs concentrations, and 39 this underprediction of VOCs contributes to lower O3 predictions in China. 40

41 Keywords: volatile organic compounds, O₃ prediction, model evaluation, emissions





42 1. Introduction

Volatile organic compounds (VOCs) in the ambient atmosphere consist of 43 thousands of gaseous organic trace substances emitted from various anthropogenic and 44 45 biogenic sources (Guenther et al., 2012; Li et al., 2017a; Kelly et al., 2018). These compounds undergo complex chemical reactions that form ozone (O₃) and secondary 46 47 organic aerosols (SOA) (Sillman, 1999; Kroll and Seinfeld, 2008). While biogenic VOCs (BVOCs) are the primary source of VOCs worldwide (Guenther et al., 2006), 48 urban areas are predominantly influenced by anthropogenic sources (Guan et al., 2020; 49 50 Guo et al., 2022; Li et al., 2022a). Anthropogenic VOCs (AVOCs) emission inventories are typically developed by estimating the total VOCs emissions using emission factors 51 (EFs) and activity rates from different sources. The VOCs speciation profiles are then 52 53 utilized to determine the emission rates of various VOCs species (Li et al., 2017a). Due to the complexity of VOCs emission processes and presence of numerous small but 54 dispersed nonpoint sources, notable uncertainties exist while determining EFs, activity 55 56 rates, and speciation profiles. It is estimated that the uncertainties associated with VOCs emissions range from approximately 68% to 76%, which are higher than those of 57 sulphur dioxide (SO₂) (12% to 40%), nitrogen dioxide (NO_x) (31% to 35%), and 58 59 particulate matter (PM) (30% to 94%) (Zhang et al., 2009; Li et al., 2019; Kurokawa 60 and Ohara, 2020; An et al., 2021).

61 Chemical transport models (CTMs), such as the Community Multiscale Air 62 Quality (CMAQ) model, Weather Research and Forecasting model coupled with 63 Chemistry (WRF-Chem), and Goddard Earth Observing System Chemical transport 64 model (GEOS-Chem) have been developed and widely used to investigate the 65 formation processes, source apportionment, and emission control strategies for various 66 air pollution issues (Zhang et al., 2021; Dang et al., 2021; Wang et al., 2021). The





67	emissions of VOCs, along with other species such as SO ₂ , NO _x , ammonia, and PM,
68	serve as essential inputs driving air quality model simulations. Uncertainties in VOCs
69	emissions notably impact air quality modelling for O3, SOA, and total PM2.5. A study
70	conducted in the United States reported substantial underprediction of VOCs emission
71	inventories in urban regions (Mcdonald et al., 2018), particularly for volatile chemical
72	products (VCPs). A simulation study that developed four cases based on the baseline
73	inventory demonstrated that augmented VOCs emission inventories have notable
74	effects on air pollutants, highlighting the need for more detailed VCPs emissions in the
75	inventory to enhance model performance (Zhu et al., 2019). In China, notable
76	discrepancies in aromatics have been observed between CMAQ predictions and
77	measurements (Wang et al., 2020). Wu et al. (2022) reconciled the bottom-up
78	methodology and measurement constraints to improve the city-scale non-methane
79	VOCs (NMVOCs) emission inventory in Nanjing, resulting in improved O ₃ simulation
80	performance with the CMAQ model.

Model evaluation serves as the initial step in establishing confidence in air quality 81 model predictions for further analysis. Numerous studies have conducted evaluations 82 of the predicted O₃ and PM_{2.5} concentrations in China (Hu et al., 2016; Li et al., 2021; 83 Li et al., 2020). Overall, the predictions of O3 and PM2.5 concentrations generally align 84 with the observations (Shi et al., 2017; Wang et al., 2021), although substantial biases 85 have been reported in certain circumstances and for specific species, such as O3 and 86 87 SOA (Gong et al., 2021; Liu et al., 2020; Hu et al., 2017; Qin et al., 2018). Given that VOCs are key precursors of O₃ and SOA, evaluating VOCs predictions can help 88 elucidate the causes of these substantial biases in predictions. However, VOCs 89 90 evaluations in regional modelling studies have been infrequent due to limited measurement data. Ambient VOCs have been measured at different locations in China 91





92	in various studies (Yang et al., 2022; Wang et al., 2022a). Unlike O ₃ and PM _{2.5} , which
93	are routinely monitored across major cities and regions in China, VOCs are often
94	measured over short periods at one or specific sites. Different studies may employ
95	different instruments and the study periods may vary, making it challenging to compile
96	VOCs measurement data from multiple studies for a comprehensive model evaluation.
97	In this study, we conducted VOCs evaluations for the first time in China by utilizing
98	summertime observations from 28 sites located in different regions of the country, as
99	part of the "Towards an Air Toxic Management System in China (ATMSYC)" project
100	(Lyu et al., 2020). This study aimed to: (1) assess the disparities in VOC levels between
101	measured ambient concentrations and predicted concentrations in various regions of
102	China using the widely used CMAQ model, (2) quantify the impacts of VOCs species
103	with substantial biases on O3 predictions through emission adjustments based on
104	observation-prediction differences, and (3) evaluate the sensitivity of O ₃ formation to
105	VOCs in key cities, providing recommendations on the necessity of emission
106	inventories and pollution control measures.

107 2. Materials and Methods

108 2.1. Observation description

109 The ATMSYC project involved a collaborative sampling campaign at 28 sites in 18 cities across China, conducted from 6 June to 24 August, 2018, with speciated VOC 110 measurement as part of the observation task (Lyu et al., 2020). Detailed site information 111 112 and sampling times can be found in Table S1. Measurements were taken at intervals of two or four hours between 8:00 and 16:00. The collection devices, analytical 113 instruments, quality controls, and other measurement methods have been previously 114 described (Lyu et al., 2019; Lyu et al., 2020; Liu et al., 2021; Zhou et al., 2023). From 115 the ATMSYC dataset, we selected 61 representative VOCs species and classified them 116





- into 20 categories, according to the SAPRC07 mechanism (Carter, 2010) to facilitate 117 comparison with model predictions. These species can be categorized into five groups: 118 alkanes, alkenes, aromatics, alkynes, and formaldehydes (HCHO). Further details 119 120 regarding these specific classifications are mentioned in Table S2. Observations of O₃ and nitrogen dioxide (NO₂) were collected from 28 ground 121 122 sites, sourced from the Chinese Ministry of Ecology and Environment 123 (https://www.mee.gov.cn/, last accessed on 20 April 2022), to assess the simulation performance of the modelled O₃ and NO₂. To evaluate the impact of meteorological 124 125 conditions, we also collected observation data of meteorological variables (temperature (T2), relative humidity (RH), wind speed (WS) and wind direction (WD)) from the 126 127 nearest meteorological stations to the 28 sites from the Chinese Meteorological Agency 128 (http://data.cma.cn/en, last accessed on 27 April 2022).
- 129 2.2. Model Configurations

The CMAQ version 5.2 model (Appel et al., 2018), coupled with the 130 131 SAPRC07TIC mechanism and aerosol module AERO6i, was utilized to simulate air quality across China from June to August 2018 (Mao et al., 2022). Meteorological fields 132 were generated using WRF version 4.2.1, employing a $1.0^{\circ} \times 1.0^{\circ}$ resolution FNL 133 reanalysis dataset from the National Centre for Atmospheric Research (NCAR). The 134 specific settings of WRF were consistent with those described by Mao et al. (2022), and 135 the simulation performance of the meteorological fields was verified (Mao et al., 2022). 136 The modelling domain with a horizontal resolution of 36 km is shown in Figure 1, 137 which divides China into seven regions: the North China Plain (NCP), Northwest, 138 Northeast, Yangtze River Delta (YRD), Central China, Southwest, and South China 139 (with a higher concentration of sites in the Pearl River Delta (PRD) region). 140

141 We utilized the Multi-resolution Emission Inventory for China (MEIC) v1.3 with





142	a resolution of $0.25^{\circ} \times 0.25^{\circ}$ in 2017 (http://www.meicmodel.org, last accessed on 25
143	January 2022) for anthropogenic emissions within China. For anthropogenic emissions
144	outside of China, we employed the Regional Emission Inventory in Asia (REAS) v3.2
145	in 2015 (https://www.nies.go.jp/REAS/, last accessed on 25 January 2022). Biogenic
146	emissions were generated using the Model for Emissions of Gases and Aerosols from
147	Nature (MEGAN) v2.1 (Guenther et al., 2012), which were then mapped to 27
148	SAPRC07TIC species, including isoprene (ISOP), α -pinene (APIN), and other BVOCs.
149	Further details on the biogenic emissions can be found in (Li et al., 2022b). Open
150	biomass burning emissions were processed using the Fire Inventory (NCAR FINN,
151	https://www2.acom.ucar.edu/modeling/finn-fire-inventory-ncar, last accessed on 28
152	January 2022).

Most emission inventories commonly employ a lumped mechanism to represent 153 VOCs. Li et al. (2014) introduced a method to allocate individual non-methane VOC 154 (NMVOC) emissions in the MEIC inventory to species groups using multiple chemical 155 156 mechanisms, utilizing mechanism-specific mapping tables from Carter (2013). This method has been widely adopted in CTMs. In this study, we followed this approach and 157 utilized a speciation profile processor called Spec DB, which is available from 158 https://intra.engr.ucr.edu/~carter/emitdb/, provided by Carter, to generate the speciation 159 profiles. The mapping scheme for the SAPRC07TIC mechanism in the MEIC and open 160 biomass burning was updated based on the step-by-step assignment framework of the 161 162 SAPRC07 mechanism provided by the MEIC team.

In this study, we examined the performance of CMAQ simulations during the observation period of the ATMSYC project. The days prior to 6 June were considered as a spin-up period. The simulated VOCs values at each site were matched with the observation time to obtain the average concentration during the same period. This





- 167 duration was defined as the study period.
- 168 2.3. Adjustment of VOCs emissions

169 Emissions were adjusted for several species that exhibited significant deviations 170 in simulations. The adjustment factors for emissions were determined by calculating the median of the ratio between observed and predicted values at 18 urban sites, which 171 172 provided an average measure of the deviation for each species. Sensitivity experiments 173 were conducted to examine the impact of the updated VOCs emissions on both predicted VOCs and O3 levels. To quantify the effect of unit increments in VOCs on O3 174 175 concentrations, the Relative Incremental Reactivity (RIR) was calculated. The RIR is a commonly used metric in observation-based model studies (Cardelino and Chameides, 176 177 1995) to assess the sensitivity of O_3 to individual precursors such as NO_x and various 178 types of VOCs. The calculation of RIR is based on Equation (1):

179
$$RIR(X) = \frac{(N_{O_3}(X) - B_{O_3}(X))/B_{O_3}(X)}{(N_X(X) - B_X(X))/B_X(X)}$$
(1)

In the equation, X represents a specific VOCs species, while B₀₃ and N₀₃ represent the O₃ concentrations in the base and adjusted emission case for X, respectively. The denominator on the right-hand side of the equation represents the relative change in emissions after the adjustment for X.

- 184 **3. Results**
- 185 3.1. Model performance evaluation

186 3.1.1. Evaluation of O₃ and NO₂

Figure 2 displays the performance of the CMAQ model for the maximum daily 8hour average (MDA8) O₃ and NO₂ concentrations at 28 sites. Model performance was assessed using statistical parameters, including the normalized mean bias (NMB), normalized mean error (NME), and correlation coefficient (R). The specific values of these statistical metrics can be found in Table S3. The results indicated that the model





192	predictions complied with the observations at most sites in the NCP, Central China, and
193	Southwest, with only slight underpredictions observed at Lanzhou's urban station (LZ-
194	U; NMB = -0.18) and Shanghai's background station (SH-B; NMB = -0.16), and a
195	slight overprediction at Shanghai's urban station (SH-U; $NMB = 0.20$). However, in the
196	PRD, overpredictions of MDA8 O3 were observed in locations such as Shenzhen's
197	station (SZ; NMB = 0.39) and Foshan's station (FS; NMB = 0.32), despite the
198	correlation coefficients being higher than the performance criteria at most sites. The
199	CMAQ's NO2 predictions exhibited underpredictions for most cities in the Northwest,
200	PRD, and some background sites, but substantial overpredictions were evident in
201	certain urban sites, such as Chengdu's urban station (CD-U; NMB = 0.92) and SZ
202	(NMB = 0.52).

203 3.1.2. Evaluation of VOCs

Figure 3 presents the observed VOCs concentrations and corresponding CMAQ 204 205 simulations across all the sites during the observation period. The proportions of the 206 three categorized VOCs groups, namely alkanes, alkenes, and aromatics, are depicted 207 in detail in Figure S1. The results revealed low predicted VOCs concentrations at most 208 sites, with particularly markable underestimation in certain areas. Table S4 displays the 209 mean values of O₃, NO₂, and total VOCs (TVOCs, encompassing the VOCs considered in this study) concentrations at the 28 sites throughout the study period. As indicated in 210 Table 1, the predicted/observed ratio (referred to as ratio hereafter) of TVOCs is $0.74 \pm$ 211 0.40. The underprediction ranged from 2.05 to 50.61 ppbv (5.77% to 85.40%) at 24 212 sites, while overpredictions occurred at four sites, namely SH-U, CU-U, Wuhan's 213 background station (WH-B), and FS, with values ranging from 0.47 to 29.53 ppbv (1.92% 214 to 89.96%). These findings suggested that the CMAQ model, employing the MEIC 215 inventory, underpredicted TVOCs concentrations. Notably, emission the 216





underprediction of TVOCs was more pronounced at sites located in the cities of
Lanzhou, Jinan, Shijiazhuang, Guiyang, and Zhengzhou, where TVOCs were
underpredicted by factors of two to six.

220 The regional averages of the predicted and observed TVOCs were calculated by averaging the predictions and observations from all the sites in each region (Table S4). 221 222 The observed ratios of TVOCs predictions varied across regions as follows: YRD (1.04) > Southwest (0.92) > PRD (0.83) > Central China (0.71) > NCP (0.42) > 223 Northwest (0.16). In Figure S2, despite having the highest observed TVOCs value 224 225 (44.08 ppbv), the model results showed a lower concentration (7.04 ppbv) in the northwest region (specifically in Lanzhou), making it the region with the lowest 226 predicted value. The predicted TVOCs concentration in the YRD region (Shanghai) was 227 228 the closest to the observed value. However, Figure 3 shows that the VOCs concentrations were notably overpredicted at SH-U and underpredicted at SH-B. The 229 southwest region appeared to have the best performance among all the regions, which 230 231 could be due to the overpredicted TVOCs at CD-U, which offsets the underprediction at other sites. Overall, the predicted and observed TVOCs concentrations exhibited 232 notable discrepancies in most regions and the performance varied across the regions. 233

Regarding the VOC components shown in Figure S2, alkanes consistently 234 235 constituted as the most abundant group of VOCs in both observations (38.3% to 50.6%) and predictions (31.6% to 44.9%). This suggested that the predicted proportion of 236 alkanes in TVOCs closely complied with the actual data. Alkenes typically ranked as 237 the second highest VOC component in observations (14.9% to 31.2%), but they were 238 underrepresented in the model (16.5% to 20.0%). In the case of predicted aromatics and 239 HCHO, their proportion in TVOCs often exceeded the observed results, which differed 240 241 from the alkynes. In terms of absolute concentrations, the underestimation of alkanes





and alkenes was relatively pronounced, particularly in the NCP and Northwest regions.
The model performed better in predicting the proportions of various VOCs species in
the PRD and Southwest regions.

245 Figure 4 illustrates the ratios of O₃, NO₂, and various VOCs species at the 28 sites. The discrepancies in ratios between urban and background sites are presented in Figure 246 247 S3. The ratio of alkanes is 0.53 ± 0.38 (median \pm standard deviation), indicating an 248 underprediction of 5.65 ± 6.81 ppbv from a concentration standpoint (Table 1). Notably, the alkanes whose reaction rate constant with hydroxyl radical (OH) between 5×10^2 249 and 2.5×10^3 ppm⁻¹ min⁻¹ (ALK2) exhibited the most notable underprediction. The 250 251 predictions for aromatics showed minor deviations across different sites, but the median 252 ratio was close to one, except for ARO2MN, which was substantially underpredicted 253 with a ratio of 0.31 ± 0.38 (0.32 ± 0.46 at urban sites), and benzene (BENZ), which was 2.75 ± 1.97 at urban sites (Table S5). Regarding alkenes, the ratios for the seven alkenes 254 were generally high $(0.51 \pm 0.48$ for alkenes), indicating underprediction in most sites. 255 256 Particularly, 1,3-butadiene (BDE13) exhibited a notable low ratio, possibly due to its reallocation from the underpredicted alkenes whose reaction rate constant is greater 257 than 7×10^4 ppm⁻¹ min⁻¹ with OH (OLE2) and the allocation factor may not be 258 universally applicable across regions. Furthermore, the predicted content of acetylene 259 260 (ACYE) was lower at all sites, while the predicted HCHO was slightly overpredicted. Considering that the observed VOCs species primarily originated from anthropogenic 261 emissions and that the majority of emitted VOCs were contributed by the MEIC, the 262 ratios between urban and background sites could verify whether the MEIC emission 263 inventory adequately reflected the differences between urban and background areas. 264 3.2. Adjusting VOCs emissions and their impacts on O₃ predictions 265

266 These findings indicated a bias between the model-predicted VOCs and observed





267	ambient VOCs concentrations. To evaluate the impact of these biases on O ₃ predictions,
268	we modified the VOCs emissions of the MEIC based on the differences between
269	observations and predictions. Previous studies have adjusted emission inventories to
270	match observed constraints for predicting VOCs and O3 in specific cities (Wu et al.,
271	2022; Wang et al., 2020). Considering the temporal and spatial variability of the 28 sites
272	we calculated the median ratio of VOCs for the 18 urban sites. We selected coefficients
273	for six representative AVOCs species with deviations exceeding 2.0 times the median,
274	including ALK2, ARO2MN, BENZ, the alkenes (excluding ethene) whose reaction rate
275	constant is less than 7×10^4 ppm ⁻¹ min ⁻¹ with OH (OLE1), propene (PRPE), and ACYE,
276	and adjusted their emission rates in the MEIC, resulting in six new cases. Additionally,
277	we conducted a case (case_all) that incorporated the aforementioned adjustments and a
278	case in which NOx was adjusted by 1.5 based on observational constraints. The
279	adjustment factors for the eight new cases are provided in Table 2.

The impact of adjusting VOCs emissions on the concentrations of O3 and VOCs is 280 presented in Table S6. The underprediction of simulated VOCs and NO2 values was 281 largely reduced for the new case, as indicated in the six cases with single-species 282 changes and the case all. In Table S7, the ratio of TVOCs in case all was modified to 283 284 0.86 ± 0.47 , demonstrating improved performance in VOCs compared to the base case. However, it was worth noting that even after the emission adjustment, the predicted 285 VOCs concentrations remained lower than the observations (particularly for 286 287 case BENZ). This discrepancy resulted from the varying reactivities of different VOC species and NOx in atmospheric chemical reactions, leading to different levels of 288 289 depletion. Additionally, both measured and modelled concentrations were subject to 290 photochemical losses (Ma et al., 2022b; Shao et al., 2011). The increased VOCs 291 concentrations resulted in higher O₃ concentrations. Based on the data presented in





292	Tables S6 and S8, the constrained species ALK2, ARO2MN, OLE1, and PRPE, guided
293	by observational data, contributed to an increase in O3 concentration, especially in
294	case_all, which led to a more pronounced overpredictions ranging from 0.62% to 6.27%
295	across all the sites. In contrast, increasing NO_x had a positive effect and reduced the O_3
296	concentration.

To illustrate regional pollution levels on a broader scale, Figure 5 displays the average concentrations of O₃, NO₂, and the six previously mentioned VOCs species studied in China during the specified period.

300 High O₃ levels were particularly prominent in most areas of the NCP, the eastern part of the Northwest, and the Sichuan Basin in the Southwest. NO2 concentrations 301 were elevated in the NCP, YRD, and PRD regions, as well as in certain megacities. The 302 303 spatial distribution of various VOCs, derived from TVOCs emissions in the MEIC, 304 exhibited broad consistency, with higher concentrations observed in south-eastern China. Megacities, akin to NO2, displayed elevated VOCs levels. Different cities 305 306 exhibited VOCs originating from various sources. ALK2 demonstrated high concentrations in individual cities but less than 1 ppbv in other regions; thus, displaying 307 stronger geographical characteristics compared to the other five VOCs. ARO2MN 308 exhibited the lowest average concentration but exerted a substantial influence on O3 309 310 due to its higher reactivity. Figure S4 illustrates the effects of altering the emission rates of NO_x and VOCs in seven scenarios across China. The left panel displays the 311 concentrations in the new cases, while the middle and right panels show the 312 concentration differences for corresponding species and O₃ between the new cases and 313 the base case, respectively. Spatial variations in NO2 and VOCs exhibited similarities. 314 The increase in NO2 was more pronounced in the NCP and YRD regions, where NO2 315 316 concentrations was consistently high. Previous studies indicate that the NCP and YRD





317	regions are predominantly limited by VOCs during the summer (Li et al., 2017b; Lyu
318	et al., 2019; Liu et al., 2021), resulting in either no change or a reduction in O ₃ when
319	NO2 increases. Conversely, in other areas with low NO2 concentrations, O3
320	concentrations increased by 0 to 10 ppbv. BENZ was the only compound whose
321	concentration decreased, and its impact on O3 in different regions mirrored that of NO2,
322	albeit at a much lower concentration. The increased emissions of ALK2, ARO2MN,
323	ACYE, OLE1, and PRPE favoured O3 production, with the most notable effects
324	observed in the NCP, YRD, and other metropolitan areas. Among these compounds,
325	OLE1 exhibited the strongest effect, while ACYE had a minimal influence.

The section 2.3 describes the calculation of the RIR values, which were used to 326 demonstrate the sensitivity of the model-simulated O₃ to VOCs constrained by 327 observations in different locations. Figure S5 presents the variations in RIR values for 328 the six VOCs across the 28 sites. OLE1, PRPE, and ARO2MN exhibited a higher RIR 329 values. Urban areas within the same city displayed a higher RIR values compared to 330 331 the background areas. With the exception of Chengdu, Guiyang, Lanzhou's background station (LZ-B), Guangzhou's background station (GZ-B), and Zhaoqing's station (ZQ), 332 where O₃ generation was more sensitive to PRPE, other areas showed a greater impact 333 334 of OLE1 concentration on O₃, indicating that adjusting the emission rate of alkenes in 335 the emission inventory was crucial for simulating changes in O₃ concentrations. For instance, improvements could be made in LZ-U, Huizhou's station (HZ), and 336 Jiangmen's station (JM), where O₃ concentrations were underpredicted in the base case. 337 Special attention should be given to the sites with high RIR values such as SH-U, CD-338 U, SZ, Zhuhai's station (ZH), and others, as O₃ generation in these locations will be 339 highly sensitive to changes in the local VOCs emission inventory. Moreover, ALK2, 340 341 ACYE, and BENZ had minimal effects on O₃, and BENZ even exhibited a negative





342 RIR values at certain sites.

343	These findings indicated a notable improvement in the underprediction of VOCs
344	when adjustments were made based on VOCs observations. However, the elevated
345	VOCs concentrations in the model could lead to increased O ₃ formation, thereby
346	enhancing the model's accuracy in areas where both VOCs and O3 were underpredicted.
347	Nonetheless, this adjustment will unavoidably worsen any existing overprediction of
348	O ₃ in the model.

349 4. Discussions

350 4.1. Large bias in TVOCs predictions at specific sites

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Significant discrepancies between predicted and observed TVOCs were observed 351 in Lanzhou, Jinan, Shijiazhuang, and Zhengzhou. Lanzhou and Shijiazhuang have 352 353 developed petrochemical industries, where high concentrations of VOCs are frequently detected downwind of industrial areas (Guan et al., 2020; Guo et al., 2022). Figure 3 354 illustrates that alkanes, alkenes, and aromatic were substantially underpredicted due to 355 356 inadequate prediction of industrial areas with high VOCs emissions in the MEIC. Jinan and Zhengzhou experienced severe air pollution due to heavy industry and traffic 357 (Zhang et al., 2017; Wang et al., 2022c). The simulated levels of TVOCs were 358 359 substantially lower than the observed levels, with alkenes exhibiting an even greater inaccuracy, being more than 10 times lower in Jinan. At certain sites, the simulated 360 TVOCs exceeded the measurements, including the CD-U, SH-U, WH-B, and FS sites. 361 In CD-U, the predicted TVOCs were almost double the measured values, whereas they 362 were underpredicted in CD-B. In Chengdu, VOCs emissions were dominated by 363 LPG/NG usage and vehicle emissions in summer, with a higher proportion of low-364 carbon alkanes compared to other cities in China (Xiong et al., 2021). Clearly, the MEIC 365 366 overpredicted VOCs emissions in CD-U, particularly for HCHO. In SH-U,





367	characterized by a dense population, the simulation of alkenes, aromatics, and HCHO
368	was approximately twice that of the measurements. This aligns complied with the report
369	by Wang et al. (2020) stating that observation-constrained aromatic emissions were
370	roughly half of the estimates provided by the MEIC in Shanghai, 2015. Peng et al. (2023)
371	also observed inconsistencies between the trend of non-methane hydrocarbon
372	emissions in Shanghai from 2009 to 2015 and the growth trend indicated by the MEIC
373	(Li et al., 2019), suggesting the effectiveness of local pollution control measures.
374	However, SH-B was situated in the easternmost part of Chongming Island, which had
375	the minimal local emissions at the 36 km grid resolution. This likely explains the
376	differences observed between the urban background areas in Shanghai. In the cases of
377	WH-B and FS, which demonstrated excellent model performance for VOCs, only the
378	overprediction of aromatics was more pronounced.

Heavy O₃ pollution events, primarily limited by VOCs, have been frequently 379 380 observed in the PRD region since its rapid development in the last century (Chan et al., 2006; Shao et al., 2009; Li et al., 2014). In the PRD region, slightly lower TVOCs 381 simulations were observed at most sites, primarily due to the underestimation of alkanes 382 and alkenes, while aromatics and HCHO were overestimated. Furthermore, the 383 384 differences in VOCs components among the cities in the PRD region could be attributed to local industry characteristics, and variations in prevention and control policies. For 385 instance, observed ethene (ETHE) in FS accounted for over 50% of the alkenes, 386 387 whereas simulations accounted for only 35%. The predicted ETHE ratio in ZH was higher (50%) than the observed ratio (20%), while other cities exhibited similar ETHE 388 389 percentages. Moreover, the proportion of ISOP in Guangzhou's alkenes was higher than that in other PRD cities, suggesting effective control of local anthropogenic alkene 390 391 emissions, consistent with the findings of Zhao et al. (2022).





392 4.2. Urban-background evaluation

Differences in atmospheric VOCs among urban background areas have been 393 extensively demonstrated (Sillman, 1999; Shao et al., 2020). As depicted in Figure 6, 394 395 we compared the average performance of the model for 18 urban sites and 10 background sites. In urban areas, the predicted TVOCs concentration (23.76 ppbv) was 396 397 lower than the observed concentration (32.46 ppbv), primarily due to the underprediction of alkanes, alkenes, and alkynes. Predicted aromatics and HCHO 398 399 exhibited higher proportions and concentrations compared to the observations. In the 400 background areas, TVOCs were also underpredicted, with concentrations lower than those in urban areas, as indicated by both the observed and predicted values. Each of 401 the five VOCs showed lower predictions, with alkanes exhibiting the most notable 402 403 disparity, with a decrease of 6.91 ppbv compared to the observed values. This suggested 404 that the model underpredicted alkanes in urban areas, which were predominantly derived from the petrochemical industry or fuel evaporation (Wang et al., 2022a). The 405 predicted proportions of alkanes, aromatics, and HCHO exhibited urban-background 406 differences consistent with the observations, reflecting the characteristics of urban and 407 408 background areas in the model. These differences were well represented in our horizontal grid resolution of only 36 km. Overall, the CMAQ model captured the 409 characteristics of different regions and urban background areas but underestimated the 410 concentrations of certain individual VOC species. 411

The ratios distinguished between urban and background areas are presented in Figure S3. The comparison revealed that the alkanes were more prominently underpredicted in the background area than in the urban area. Xylene (XYL), 1,2,4trimethylbenzene (B124), OLE1, OLE2, and PRPE were also underpredicted to a greater extent in the background area. This could be attributed to the scarcity of





417	background sites or the model's underprediction of VOCs emissions in the background
418	area. The model's performance in simulating ISOP, a BVOC, in urban areas was not as
419	satisfactory as in the background areas, which was consistent with the findings of Ma
420	et al. (2021) suggesting that MEGAN could underestimate the emissions from urban
421	green spaces. APIN, a notable monoterpene, including anthropogenic emissions from
422	biomass burning and VCPs, could be either underpredicted or disregarded (Wang et al.,
423	2022b; Mcdonald et al., 2018), resulting in common underprediction with a median
424	ratio of five in urban-background areas. Additionally, the simulated HCHO
425	concentrations were higher in the urban areas. Overall, these results indicated that the
426	model generally performed better for anthropogenic VOCs in the urban areas. However,
427	there were still a few notable outliers and significant deviations for a majority of VOCs,
428	particularly those with high chemical reactivity. These deviations will inevitably impact
429	the model's calculation of photochemical reactions involved in O ₃ generation.

430 4.3. Implications and suggestions

Accurately predicting VOCs is crucial for O₃ modelling. However, due to limited
 measurement data and uncertainties in emission inventories, accurately simulating the
 VOCs across China using CTMs remains challenging.

Considerable efforts have been dedicated to the development of VOCs emission 434 inventories in recent years (Li et al., 2019; An et al., 2021; Chang et al., 2022). However, 435 our findings indicate substantial variation in the model performance of VOCs across 436 437 different regions and species. Therefore, the inclusion of accurate local emission factors, activity data, and source profiles is essential. Sha et al. (2021) compiled an integrated 438 dataset of AVOCs source profiles in China, emphasizing the need for supplementary 439 and timely updates to these profiles in the future. Apart from anthropogenic emissions, 440 441 model resolution, and chemical mechanisms meteorological conditions, and BVOCs





442	emissions also contribute to the uncertainty of VOCs modelling, thereby affecting the
443	performance of O ₃ modelling (Zhang et al., 2021; Wang et al., 2021; Liu et al., 2022).
444	High-resolution models require higher emission inventory resolution (Li et al.,
445	2022; An et al., 2021), which can improve simulation performance to a certain extent.
446	Given the large scope of the model used in this study and the $0.25^{\circ} \times 0.25^{\circ}$ horizontal
447	resolution of the MEIC inventory, a resolution of 36 km was chosen to balance
448	computational efficiency and the preservation of information from the emission
449	inventory, but inevitably results in deviation of the modelled VOCs and other elements.
450	On the one hand, urban and background sites in close proximity may be assigned to the
451	same grid in the model, as shown in Table S3, making it difficult to distinguish the
452	differences in modelled VOCs between urban and background sites in cities such as
453	Shijiazhuang, Jinan, Wuhan, and Guiyang; on the other hand, in real atmospheres, even
454	with close proximity, the observed VOCs may differ greatly in concentration, which is
455	challenging to capture in a coarse-resolution model. When applying coarse-resolution
456	emission inventories, increasing the model resolution can enhance the spatial
457	correlation between observed and predicted concentrations, but does not always
458	improve simulation performance (Zheng et al., 2021). High-resolution models may
459	introduce more emission mapping errors, which can be reduced by using coarse-
460	resolution model grids (Zheng et al., 2021). Therefore, addressing this issue requires
461	not only finer model resolution but also improved emission inventories.

The SAPRC07tic chemical mechanism used in this study has been proven reliable in previous model applications (Qin et al., 2022), reducing the computational effort compared to the explicit MCM mechanism (Li et al., 2015) while retaining the chemical reactivity of various VOCs. However, the lumped VOCs species contain more VOCs species than those in corresponding observations. Therefore, if both the emission





- inventory and model are sufficiently accurate, the predicted values should theoretically
- 468 be higher.

469 Meteorology bias also contributed to some bias of the VOCs predictions. We added 470 evaluation of the meteorology predictions in this study, and the results are shown in Table S9 and S10. The results are consistent with other studies in China (Mao et al., 471 472 2022; Wang et al., 2021). It is observed that temperature is overpredicted at most sites, 473 while RH is mostly underpredicted. The combination of high temperature and low RH facilitates the consumption of VOCs through photochemical reactions, which may 474 475 explain the tendency of our modelled VOCs to be underestimated. But we believe it is insufficient to account for the underestimation of low-reactivity VOC species (mainly 476 alkanes). Furthermore, the modelled wind speeds slightly exceed the observations, 477 which may also contribute to VOCs underprediction (Table S10). While the bias in 478 meteorological conditions contributes to the underestimation of modelled VOCs, the 479 underestimated VOCs emissions is the key factor for the VOCs underprediction across 480 481 most of the cities.

In this study, the adjustment of VOCs emissions resulted in increased predicted 482 emission levels, subsequently leading to higher O₃ predictions. However, these 483 adjustments are simplistic and fail to account for regional variations in VOCs biases. 484 The accuracy of VOCs measurement data is also crucial. Therefore, there is a need to 485 promote the establishment of a national O₃ precursor monitoring network and develop 486 a standardized framework with quality control systems. This would facilitate the 487 comparability of VOCs measurements between regions, thereby supporting related 488 research and the implementation of collaborative regional prevention and control 489 490 measures.

491 5. Conclusion





In this study, we conducted a comprehensive evaluation of the simulation performance of VOCs using the CMAQ model and investigated the influence of predicted VOCs on O₃ formation. The inclusion of summertime-observed VOCs data from the ATMSYC project for 28 sites in China enhanced the spatiotemporal comparability of our model evaluation.

During the study period, TVOCs were found to be underpredicted by 14.1 ± 13.2 497 498 ppbv at 24 sites, except for SH-U, CD-U, WH-B, and FS. Despite some sites exhibiting similar TVOCs concentrations, differences still persisted in their specific components. 499 500 Through considering the uncertainties of the MEIC model and relevant factors, we found several sites with substantial inaccuracies, such as Jinan, Shijiazhuang, Lanzhou, 501 Chengdu, and Guiyang. The model's performance in predicting TVOCs and their 502 503 components varied across regions, with better predictions observed in urban areas compared to background areas. 504

Alkanes, alkenes, ARO2MN, and alkynes are generally underpredicted, with ratios 505 506 of 0.53 ± 0.38 , 0.51 ± 0.48 , 0.31 ± 0.38 , and 0.41 ± 0.47 , respectively. In urban areas, the CMAQ model exhibited underpredictions for OLE1, ALK2, ARO2MN, PRPE, 507 508 ACYE, and NO_x, ranging from 2.0 to 4.6 times, while overpredicting BENZ by 2.75times. For sensitivity experiments, their emissions were adjusted and their impact on 509 O₃ and VOCs was evaluated. These adjustments improved the model's VOCs 510 performance, resulting in a change in the ratio of total VOCs to 0.86 ± 0.47 . However, 511 the increased VOCs contributed to higher reactivity, exacerbating O₃ overpredictions 512 by 0.62% to 6.27% across the sites. Consequently, RIR values were calculated to depict 513 the varying reactivities of VOCs in different regions, with OLE1, PRPE, and ARO2MN 514 contributing the highest RIR values during the study period. 515

516 Due to the inaccuracies present in current VOCs emission inventories, notable





- efforts are needed to enhance the development and updating of emission inventories, particularly in regions characterized by developed industries, evolving energy structures, and relatively underdeveloped conditions. It is only through improving the accuracy of VOCs emission inventories that we can ensure reliable model performance
- 521 in predicting O₃ levels, thereby establishing a solid foundation for addressing the
- 522 escalating issue of O₃ pollution.
- 523

524 Code and data availability

525 The model outputs and observation data are currently available upon request.

526 Author contributions

YS and JH designed research. YS, JH, JL, MQ, XX, KG, FY, and JM contributed to model development, simulations, and data processing. XL and HG provided the observation data. XL and LH contributed to result discussion. YS prepared the manuscript and all coauthors helped improve the manuscript.

531 Competing interests

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

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- 751





752 Table 1. Mean, median, maximum (max), minimum (min), and standard deviation (std)

			- ()		5F ·			
		Alkanes	Alkenes	Aromatics	ARO2MN	Alkyne	НСНО	TVOCs
					(Aromatics)	-		
	mean	0.59	0.60	1.33	0.40	0.55	1.66	0.70
Ratio(pre/obs)	median	0.53	0.51	1.30	0.31	0.41	1.21	0.74
	max	1.87	2.46	3.29	1.96	2.36	8.70	1.90
	min	0.13	0.09	0.10	0.05	0.09	0.25	0.15
	std	0.38	0.48	0.89	0.38	0.47	1.61	0.40
	mean	-6.18	-4.02	0.42	-0.28	-1.16	0.16	-10.78

0.83

6.09

-8.18

3.47

-0.25

0.24

-0.74

0.20

-1.04

0.87

-2.64

0.97

0.49

5.57

-8.90

2.99

-7.57

29.53

-50.61

16.11

753 of the Ratios and differences (Diff) for five VOCs groups and TVOCs at 28 sites

754 755

Diff(pre-obs)

median

max

min

std

-5.65

14.12

-19.40

6.81

-2.56

3.50

-15.50

4.69





Cases in CMAO	Changing species	Adjusted			
Cases in CMAQ	in MEIC	coefficient			
base case					
case_NO _x	NO, NO ₂	1.5			
case_ALK2	ALK2	4.6			
case_ARO2MN	ARO2MN	3.2			
case_BENZ	BENZ	0.4			
case_OLE1	OLE1	2.0			
case_PRPE	PRPE	2.1			
case_ACYE	ACYE	2.8			
case_all	all of the above VOCs				

756 Table 2. New cases of adjusting emission coefficient under observation constraints

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Figure 1. The CMAQ modelling domain cover China and the surrounding countries andregions in this study, including 28 blue dots that represent the positions of VOCs





sampling sites. We divided China into seven regions according to the geographical
location of different provinces, which comprise the following sites: NCP: BJ-B, BJ-U,
SJZ-B, SJZ-U, JN-B, JN-U; Northwest: LZ-B, LZ-U; Northeast (No observation site);
YRD: SH-B, SH-U; Central China: ZZ-B, ZZ-U, WH-B, WH-U; Southwest: CD-B,
CD-U, GY-B, GY-U; South China: Most of the sites are concentrated in PRD region
(shown in the enlarged subgraph in the lower left): GZ-B, GZ-U, SZ, HZ, DG, FS, JM,
ZQ, ZS, ZH.



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Figure 2. Model performance on MDA8 O₃ and NO₂ of 28 sites in different regions from June 6th to August 24th in 2018. The blue and red lines denote performance criteria for MDA8 O₃ suggested by Emery et al. (2017) and the symbols in different colors distinguish different regions of China.













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Figure 4. The predicted/observed ratio (pre/obs) of O₃, NO₂ and different VOCs species at 28 sites (both urban and background). The rectangles with different colors represent the ratio range of 25% to 75% for all sites. The vertical lines with a horizontal bar are called 1.5 Interquartile Range (1.5 IQR). The horizontal lines in rectangles represent the median value and the hollow dots are the mean value. The dots outside the 1.5 IQR are Outliers.

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Figure 5. Prediction concentration of O₃, NO₂ and six VOCs species in the base case
from June 6th to August 24th in 2018.

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791 Figure 6. Observed and predicted values of different VOCs species by sites average.