

# The variations of VOCs based on the policy change of Omicron in traffic-hub city Zhengzhou

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**Abstract:** Online volatile organic compounds (VOCs) were monitored before and after the Omicron policy change at an urban site in polluted Zhengzhou from December 1, 2022, to January 31, 2023. The characteristics and sources of VOCs were explored. The daily average concentration of PM<sub>2.5</sub> and total VOCs (TVOCs) ranged from 53.5 to 239.4  $\mu\text{g}/\text{m}^3$  and from 15.6 to 57.1 ppbv with an average value of  $111.5 \pm 45.1 \mu\text{g}/\text{m}^3$  and  $36.1 \pm 21.0$  ppbv, respectively during the entire period. The values of PM<sub>2.5</sub> and TVOCs in Case 2 (pollution episode after the abolishment of “Nucleic Acid Screening Measures for all staff” policy) were 1.3 and 1.8 times of the values in the Case 1 (pollution episode during “Nucleic Acid Screening Measures for all staff” policy). The concentration of TVOCs in Case 1 and Case 2 were  $48.4 \pm 20.4$  and  $67.6 \pm 19.6$  ppbv, respectively, increased by 63% and 188% compared with values during clean days. Alkanes were found to be the most abundant compounds during the entire period. Equivalent volume contribution of halogenated hydrocarbon and oxygenated VOCs (15%) were found the most in Case 2, followed by alkenes (10%). Though the volume contributions of aromatics were the lowest (6% in Case 1 and 7% in Case 2), the highest increasing ratio was found from clean days to polluted episodes. Positive Matrix Factor model results showed that the main source of VOCs during the observation period was industrial emissions, which accounted for 32% of the TVOCs, followed by vehicular emission (27%) and combustion (21%). In Case 1, industrial emissions constituted the largest contributor, accounting for 32% of the total VOCs. In Case 2, however, the share of vehicular emission source increased to 33%, becoming the primary source of VOCs. Secondary organic aerosol formation potential (SOAP) values were 37.6 and 65.6  $\mu\text{g}/\text{m}^3$  in Case 1 and Case 2, respectively. In Case 1, industrial source accounted for the overwhelming majority (63%, 23.8  $\mu\text{g}/\text{m}^3$ ), while vehicular source, as the second

34 largest source, accounted for only 18%. In Case 2, the distribution of contributions is  
35 more uniform, with solvent usage source and fuel evaporation source accounting for  
36 the majority of SOAP, at 32% (20.9  $\mu\text{g}/\text{m}^3$ ) and 26% (16.8  $\mu\text{g}/\text{m}^3$ ), respectively.  
37 Industrial source and solvent usage continue to be the main contributors to SOAP on  
38 clean days. It is crucial to prioritize the regulation of emissions from industrial and  
39 solvent-using sectors as a means of curbing  $\text{PM}_{2.5}$  pollution in Zhengzhou. Additionally,  
40 it is imperative to consider the impact of rising vehicular emissions on air quality.

41

42 **Keywords: Volatile organic compounds; Pollution episode; Source apportionment;**

## 43 **1. Introduction**

44 Volatile organic compounds (VOCs) in the atmosphere have high reactivity and  
45 can react with nitrogen oxides (NO<sub>x</sub>) to form a series of secondary pollutants such as  
46 ozone (O<sub>3</sub>) and secondary organic aerosol (SOA), resulting in regional air pollution (Li  
47 et al., 2019; Hui et al., 2020). The problem of O<sub>3</sub> pollution has been plaguing major  
48 urban agglomerations in China (Zheng et al., 2010; Li et al., 2014; Wang et al., 2017).  
49 SOA is an important component of fine particulate matter (PM<sub>2.5</sub>) and contributes  
50 significantly to haze pollution (Liu et al., 2019). PM<sub>2.5</sub> remains the most significant air  
51 pollutant in many Chinese cities for years (Shao et al., 2016; Wu et al., 2016). In  
52 addition, VOCs, represented by the benzene homologues, can cause damage to kidneys,  
53 liver, and nervous system of humans when they enter the body (Zhang et al., 2018).

54 Studies have shown that the most common VOC components in China are alkanes,  
55 olefins, aromatic hydrocarbons, oxygenated VOCs (OVOCs), and halogenated  
56 hydrocarbons, among which alkanes are the most abundant species (Liu et al., 2020;  
57 Zhang et al., 2021a). VOCs in the atmosphere have a wide range of sources, and VOCs  
58 in different regions are affected by multiple factors such as local geography, climate,  
59 and human activities (Mu et al., 2023; Zou et al., 2023). The above reasons lead to  
60 significant regional and seasonal differences in the characteristics of VOCs (Song et al.,  
61 2021). For example, the annual average concentration of VOCs in the coastal  
62 background area of the Pearl River Delta is 9.3 ppbv. The seasonal variation trend of  
63 VOCs is high in autumn and winter and low in summer (Yun et al., 2021). In contrast,  
64 the average VOC concentration in autumn and winter in Beijing was  $22.6 \pm 12.6$  ppbv,  
65 and the VOC concentration in the winter heating period was twice that in the autumn  
66 non-heating period (Niu et al., 2022).

67 Moreover, the sources of VOC components in different regions are also related to  
68 the local industrial structure and living habits. In rural areas of North China Plain in  
69 winter, it is found that the SOA formation potential (SOAP) of VOCs under low NO<sub>x</sub>  
70 conditions is significantly higher than that under high NO<sub>x</sub> conditions, and the increase  
71 of aromatic hydrocarbon emissions caused by coal combustion is the main reason for  
72 the higher SOAP in winter (Zhang et al., 2020). Li et al. (2022) found that the average  
73 increased concentration of acetylene was 4.8 times from autumn to winter in the  
74 Guanzhong Plain, indicating that fuel combustion during the heating period in winter  
75 has a significant impact on the composition of VOCs. In contrast, continuous

76 observations conducted by Zhou et al. (2022) in the suburbs of Dongguan in summer  
77 found that industrial solvent usage, liquefied petroleum gas (LPG) and oil and gas  
78 volatilization were the main sources of VOCs. The results highlighted a wide variation  
79 of characteristics, sources and chemical reactions of VOCs in the atmosphere, thus it is  
80 necessary to investigate VOCs in different cities when formulating control measures.

81 Zhengzhou, as the capital of Henan Province, is an important transportation hub  
82 and economic center in the Central Plains region. Zhengzhou is currently facing  
83 significant air pollution problems, with the Air Quality Index at the bottom of the  
84 national ranking of 168 cities for many years. In January 2023, for example, the number  
85 of polluted days with PM<sub>2.5</sub> as the primary pollutant was 17, and the daily average value  
86 of PM<sub>2.5</sub> reached a maximum of 298  $\mu\text{g}/\text{m}^3$   
87 ([https://www.aqistudy.cn/historydata/daydata.php?city=%E9%83%91%E5%B7%9E](https://www.aqistudy.cn/historydata/daydata.php?city=%E9%83%91%E5%B7%9E&month=202301)  
88 &month=202301, Accessed Jan 2024), which is almost 300% higher than the Chinese  
89 daily average standard (grade II, 75  $\mu\text{g}/\text{m}^3$ ). The studies of VOCs were carried out in  
90 Zhengzhou in recent years, which focused on the characteristics and sources of VOCs  
91 during pollution episodes (Lai et al., 2024) or before the coronavirus epidemic outbreak  
92 (Li et al., 2020; Zhang et al., 2021b). While some atmospheric VOCs studies involving  
93 the impact of Covid-19 lockdown have been performed in India (Singh et al., 2023a),  
94 in China (e.g., Pei et al., 2022; Jensen et al., 2023; Zuo et al., 2024), or with respect to  
95 toluene, benzene, m/p-xylene and ethylbenzene only (e.g., Sahu et al., 2022; Singh et  
96 al., 2023b), a gap persisted in the investigation of VOCs due to the impact of  
97 abolishment of China's zero-policy. In addition, there have been some studies  
98 discussing the impact of human factors on air pollution during and after the outbreak  
99 of the Coronavirus disease (e.g., Ma et al., 2022; Jiang et al., 2023; Song et al., 2023),  
100 but as mentioned earlier, only a few studies with in-depth exploration of the changes in  
101 VOCs and none dealing with ending the zero-Covid policy during Omicron variant  
102 infection period.

103 In this study, a continuous online observation of VOCs in polluted winter at an  
104 urban site was carried out, which covered the abolishment of lockdown measures in  
105 Zhengzhou. China lifted the zero-COVID strategies, notably by announcing the '10  
106 measures' about the optimization of COVID-19 rules on 7 December 2022  
107 ([http://www.news.cn/politics/2022-12/07/c\\_1129189285.htm](http://www.news.cn/politics/2022-12/07/c_1129189285.htm), Accessed Jan 2024),  
108 which led to significant changes in social activities. After that, China experiences a  
109 nationwide outbreak of COVID-19. Our research primarily concentrates on the period

110 dominated by COVID-19 Omicron variant, where they demonstrate notable differences  
111 from the early virus strains (i.e., original SARS-CoV-2 virus and Delta) in terms of  
112 geographical transmission, the scale of the infected population, and symptom  
113 manifestation (Petersen et al., 2022; Merino et al., 2023). A two-month-long lockdown  
114 measure was applied to after first Omicron case of student in Zhengzhou University  
115 was confirmed on October 8, 2022. Lockdown measure was abolished from the  
116 beginning of December in 2022, which resulted in a sharp increase of Omicron-infected  
117 people and a decrease in daily social production activities. In fact, the “Nucleic Acid  
118 Screening Measures for all staff” policy was also canceled at 8 October in 2022. People  
119 are basically homebound after the lifting of the lockdown policy due to infection or fear  
120 of infection of Omicron. The resumption of normal production and livelihoods was  
121 based on the assumption of herd immunization. This change is worth exploring in terms  
122 of its impact on transportation and industrial production emissions. Therefore, the  
123 characteristics and variations of VOCs during different periods were investigated to  
124 assess their impact on the formation of SOA and to provide data support for future  
125 pollution control policies in Zhengzhou.  
126

## 127 **2. Materials and methods**

### 128 **2.1 Sample collection and Chemical analysis**

129 The online VOCs observation station is located on the roof of the Zhengzhou  
130 Environmental Protection Monitoring Center, which is in the urban area. The sampling  
131 site is close to main roads on three sides (150 m away from Funiu Road on the east side,  
132 200 m away from Qinling Road on the west side, and connected to Zhongyuan Road  
133 on the south side), and surrounded by residential areas and commercial areas without  
134 other large nearby stationary sources. The sampling period for this study was from  
135 December 1, 2022, to January 31, 2023, which is always the most polluted period in  
136 the entire year. Apart from a brief occurrence of rain and snow on December 25, the  
137 sampling days were either sunny or cloudy. The wind speed (WS), temperature (Temp)  
138 and relative humidity (RH) during this period were  $1.3 \pm 0.9$  m/s,  $5.3 \pm 3.2$  °C and  $38.9$   
139  $\pm 19.0\%$ ), respectively, similar to the values observed in previous years in Zhengzhou.  
140 It is interesting to point out that the sampling period in the present study covered the  
141 entire infection period of Omicron in Zhengzhou, including the phase of surge in  
142 infected population (Infection period, from 2022.12.01 to 2022.12.31) and restoration  
143 of production and livelihood phase (Recovery period, from 2023.1.1 to 2023.1.31 in  
144 2023) (Fig. S1, Chinese Center for Disease Control and Prevention, 2023).

145 The VOCs were measured hourly using a GC-FID/MS (TH-PKU 300 b, Wuhan  
146 Tianhong Instruments Co., China). The instrument TH-PKU300b includes electronic  
147 refrigeration ultra-low temperature pre-concentration sampling system, analysis system  
148 and system control software. The ambient VOCs in the first 5 minutes of each hour  
149 were collected by the sampling system and then entered the concentration system.  
150 Under low temperature conditions, the VOCs samples collected were frozen in the  
151 capillary capture column, and then quickly heated and resolved, so that the compounds  
152 entered the analysis system. After separation by chromatographic column, the  
153 compounds were monitored by FID and MS detectors. During the detection process,  
154 the atmospheric samples collected undergo analysis through two distinct pathways. C2-  
155 C5 hydrocarbons are analyzed using FID, while C5-C12 hydrocarbons, halocarbons,  
156 and OVOCs are analyzed with a MS detector. After excluding species with missing data  
157 exceeding 10%, the detected volatile organic compounds include 29 alkanes, 11 alkenes,  
158 17 aromatics, 35 halocarbons, 12 OVOCs, 1 alkyne (acetylene), and 1 sulfide (CS<sub>2</sub>)  
159 with a total of 106 compounds.

160 The instrument was calibrated per week to ensure the accuracy of VOCs by  
 161 injecting standard gases with a five-point calibration curve. The detection limit of C2-  
 162 C5 hydrocarbons ranges from 0.007 to 0.099 ppbv, other hydrocarbons are 0.004–0.045  
 163 ppbv, halogenated hydrocarbons 0.009-0.099 ppbv, OVOCs and other compounds of  
 164 0.006–0.095 ppbv. Thirty-two of the monitored VOCs had over 90% observed data  
 165 greater than the detection limit, and 34 had more than 50% observed data greater than  
 166 the detection limit.

167 Simultaneous observations at the same site were also carried out for particulate  
 168 matter (PM<sub>2.5</sub>, PM<sub>10</sub>), other trace gases (carbon monoxide (CO), O<sub>3</sub>, nitric oxide (NO),  
 169 nitrogen dioxide (NO<sub>2</sub>)), and meteorological data (Temperature, RH, WS, and wind  
 170 direction (WD)) based on 1 h resolution.

## 171 2.2 Positive Matrix Factorization (PMF) model

172 EPA PMF5.0 model was used for the quantitative source analysis of VOCs (Norris  
 173 et al., 2014). The principles and methods have been described in detail in previous  
 174 studies (Mozaffar et al., 2020; Zhang et al., 2021b). The decomposition of the PMF  
 175 mass balance equations is simplified as follows (Norris et al., 2014):

176

$$177 \quad x_{ij} = \sum_{k=1}^p g_{ik} f_{kj} + e_{ij} \quad (1)$$

178

179 where  $x_{ij}$  is the mass concentration of species  $j$  measured in sample  $i$ ;  $g_{ik}$  is the  
 180 contribution of factor  $k$  to the sample  $i$ ;  $f_{kj}$  represents the content of the  $j$ th species in  
 181 factor  $k$ ;  $e_{ij}$  is the residual of species  $j$  in sample  $i$ ;  $p$  represents the number of factors.  
 182 The fitting objective of the PMF model is to minimize the function  $Q$  to obtain the  
 183 factor contributions and contours. The formula for  $Q$  is given in Eq. (2):

184

$$185 \quad Q = \sum_{i=1}^n \sum_{j=1}^m \left[ \frac{x_{ij} - \sum_{k=1}^p g_{ik} f_{kj}}{u_{ij}} \right]^2 \quad (2)$$

186

187 where  $n$  and  $m$  denote the number of samples and VOC species, respectively.

188 Concentrations and uncertainty data are required for the PMF model. In this study,

189 the median concentration of a given species is used to replace missing values with an  
190 uncertainty of four times of the median values; data less than the Method Detection  
191 Limit (MDL) were replaced with half the MDL, with an uncertainty of 5/6 of the MDL;  
192 and the uncertainty for values greater than the MDL was calculated using Eq. (3). In  
193 Eq. (3),  $EF$  is error fraction, expressed as the precision of VOCs species, and the setting  
194 range can be adjusted from 5 to 20% according to the concentration difference (Buzcu  
195 et al., 2006; Song et al., 2007); and  $c_{ij}$  is the concentration of species  $j$  in sample  $i$ :

$$196 \quad U_{ij} = \sqrt{(EF \times c_{ij})^2 + (0.5 \times MDL)^2} \quad (3)$$

197 when the concentration of VOCs in the species is less than the value of the  
198 detection limit  $U_{ij}$  is calculated using Eq. (4):

$$199 \quad U_{ij} = \left(\frac{5}{6}\right) MDL \quad (4)$$

200 VOC species and concentration input into PMF were carefully selected to ensure  
201 the accuracy of the PMF results. Species were excluded when over 25% of the samples  
202 were missing or concentrations values were below the MDL (Gao et al., 2018); VOCs  
203 with a short lifetime in the atmosphere were also excluded unless they are source-  
204 relative species (Zhang et al., 2014; Shao et al., 2016). After that, retained VOC species  
205 were categorized according to the signal-to-noise ratio (S/N) with  $S/N < 0.2$  species  
206 categorized as bad,  $0.2 < S/N < 2$  species categorized as weak; and  $S/N > 2$  species  
207 categorized as strong (Shao et al., 2016).

208 We used displacement of factor elements (DISP) to assess PMF modelling  
209 uncertainty (for a description, see Paatero et al. (2014)).  $Q$  was less than 1% and no  
210 swaps occurred for the small est  $dQ^{\max}$  in DISP.  $F_{\text{peak}}$  values from -2 to 2 were tested  
211 to explore the rotational stability of the solutions.  $Q_{\text{true}}/Q_{\text{exp}}$  is lowest when  $F_{\text{peak}} = 0$ ,  
212 so we chose the PMF results for that case (Fig. S2a). After examining 3-8 factors, 20  
213 base runs with 5 factors eventually selected to represent final result. We provide an  
214 explanation of factor selection in the supplementary materials. Fig. S2(b) includes  
215  $Q_{\text{true}}/Q_{\text{exp}}$ ,  $Q_{\text{robust}}/Q_{\text{exp}}$  for factors 3-8. The slopes of these two ratios in changed at five  
216 factors, and we found that five factors were more realistic after repeated comparisons  
217 of the results at four, five and six factors.



## 218 **2.3 SOA generation potential**

219 The contributions of VOC species to SOAP were calculated based on the toluene  
220 weighted mass contributions (TMC) method (Derwent et al., 2010). The methodology  
221 for calculating SOAP is as follows:

$$222 \text{SOAPF}_i = \frac{\text{VOCs component } i \text{ to SOA mass concentration increments}}{\text{Toluene to SOA mass concentration increment}} \times 100 \quad (5)$$

224  
225  $\text{SOAPF}_i$  for each VOC is taken from the literature (Derwent et al., 2010). The  
226 SOAP was estimated by multiplying the  $\text{SOAPF}_i$  value by the concentration of  
227 individual VOC species. The SOAP calculations through each VOC are as follows:

$$228 \text{SOAP} = \sum E_i \times \text{SOAPF}_i \quad (6)$$

229  $E_i$  is the concentration of species  $i$ .

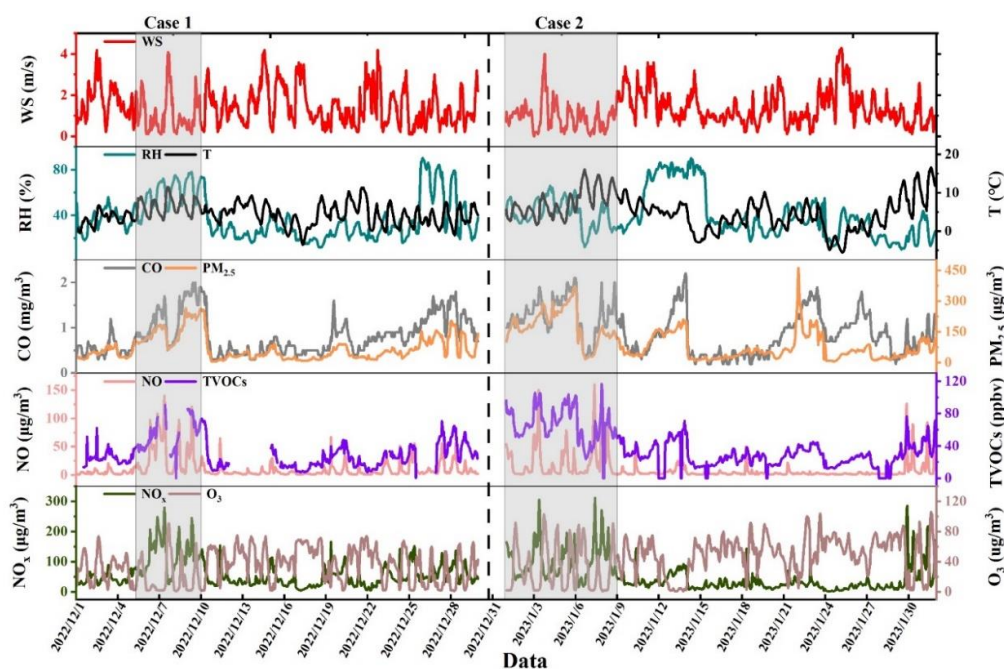
## 230 **3. Results and discussion**

### 231 **3.1 Overview of variation in pollutants and meteorological parameters**

232 Figure 1 shows the time series of meteorological parameters, TVOCs, O<sub>3</sub>, NO<sub>x</sub>,  
233 SO<sub>2</sub>, CO and PM<sub>2.5</sub> during the observed periods. Low WS and Temperature were found  
234 with an average value of  $1.3 \pm 0.6$  m/s and  $5.0 \pm 2.5$  °C, respectively, during the entire  
235 period, comparable with observations at the same site in 2021 (Lai et al., 2024). A total  
236 of 62 days of valid data was acquired with the daily average concentration of PM<sub>2.5</sub>  
237 ranging from 53 to 239  $\mu\text{g}/\text{m}^3$ , with the average value of  $111 \pm 45$   $\mu\text{g}/\text{m}^3$ . The  
238 concentration of TVOCs ranged from 15.6 to 57.1 ppbv with an average of  $36.1 \pm 21.0$   
239 ppbv, higher than the same period in last year ( $27.9 \pm 12.7$  ppbv, Lai et al., 2024).  
240 During the observation period, the average values of T, WS and RH were  $5.0 \pm 2.5$  °C,  
241  $1.3 \pm 0.6$  m/s and  $38.9 \pm 16.7\%$ , respectively.

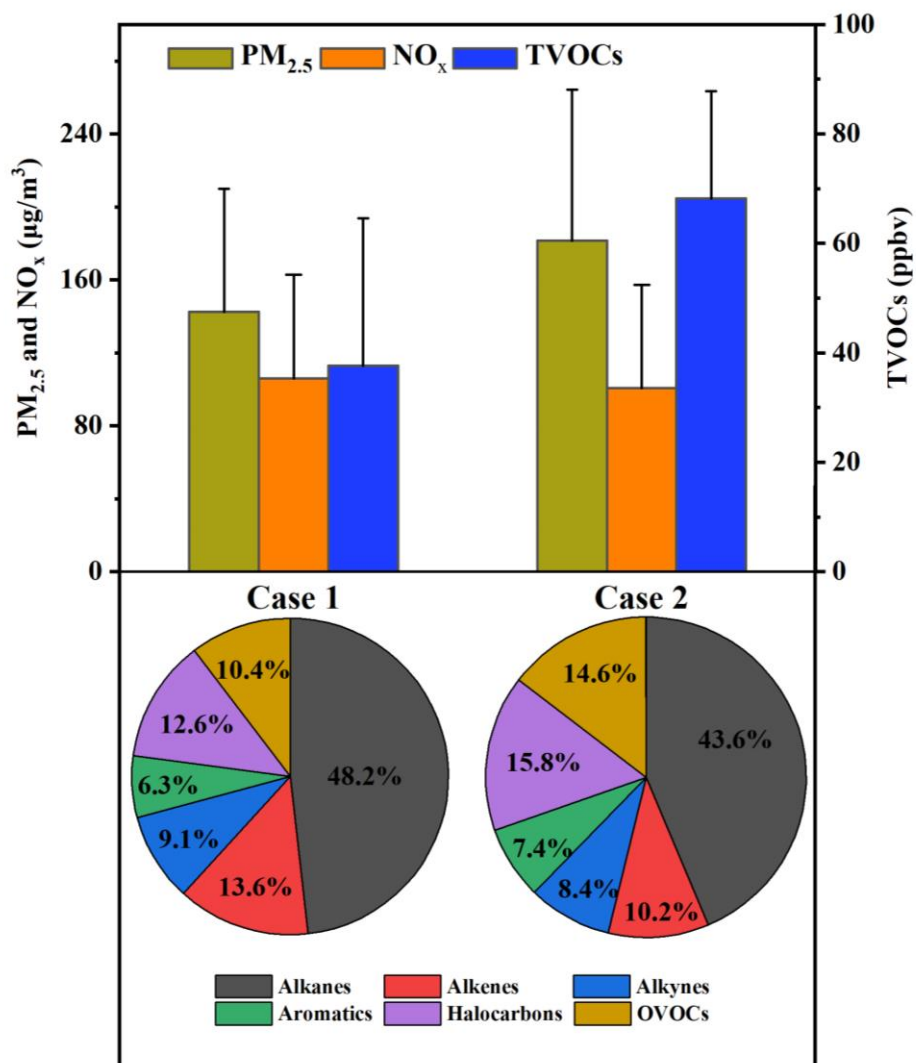
242 The relationship between meteorological parameters and pollutant concentrations  
243 were analyzed and correlations between PM<sub>2.5</sub>, TVOCs and NO<sub>x</sub> and RH were found  
244 (Fig. S3), suggesting that meteorological conditions have an important influence on  
245 pollution formation. The comparisons of average concentrations of different periods

249 between different periods are presented in Tables 1 and 2. WS, Temp and RH  
 250 conditions during infection and recovery periods were generally similar. However, the  
 251 average concentration of PM<sub>2.5</sub> during the recovery period was 1.6 times the value  
 252 during the infection period. Furthermore, the concentrations of other pollutants  
 253 including SO<sub>2</sub>, NO<sub>2</sub>, CO, and O<sub>3</sub> all showed a similar trend between infection and  
 254 recovery periods. The TVOCs concentration during the recovery period was 1.2 times  
 255 the value during the infection period, showing an obvious increase trend after  
 256 resuming production. Decreased trends of air pollutants were found in other studies  
 257 before and after the outbreak of the novel coronavirus (COVID-19) in early 2020 (Qi  
 258 et al., 2021; Wang et al., 2021).



259  
 260 Fig. 1. Time series of WS, T, RH, CO, PM<sub>2.5</sub>, NO, TVOCs, NO<sub>x</sub> and O<sub>3</sub> during the observation  
 261 period.

262 The shadow section in Fig. 1 represents two haze pollution events during the  
 263 monitoring period. A pollution event is determined when the daily average  
 264 concentration of PM<sub>2.5</sub> exceeds 75 µg/m<sup>3</sup> (China's II-level standard) for at least three  
 265 consecutive days. Case 1 (December 5 to December 10 with daily average PM<sub>2.5</sub> =  
 266 142.5 µg/m<sup>3</sup>) and Case 2 (January 1 to January 8 with daily average PM<sub>2.5</sub> = 181.5  
 267 µg/m<sup>3</sup>) were selected as they represent the pollution events in infection and  
 268 recovery periods, respectively, due to their long duration and high pollution levels. Any days with  
 269 a PM<sub>2.5</sub> concentration lower than 35 µg/m<sup>3</sup> (China's I-level standard) is considered as  
 270 Clean days.



271  
 272 Fig. 2. The concentration of PM<sub>2.5</sub>, NO<sub>x</sub>, TVOCs and the composition ratio of VOCs in Case 1 and  
 273 Case 2.

274 As for the two representative pollution processes (Case 1 during the infection  
 275 period and Case 2 during the recovery period), the concentration of TVOCs in Case 1  
 276 and Case 2 were  $48.4 \pm 20.4$  and  $67.6 \pm 19.6$  ppbv (Fig. 2), respectively, increased by  
 277 63% and 188% compared with values during clean days. The average concentrations  
 278 of PM<sub>2.5</sub> and TVOCs during Case 2 were 1.3 and 1.8 times the values in Case 1. The  
 279 highest volume contributions of alkanes were found both in Case 1 (48%) and Case 2  
 280 (44%), consistent with the results in the Yangtze River Delta region (36-43%, Liu et al.,  
 281 2023). While alkenes exhibited higher volume percentages of 13% in Case 1, followed  
 282 by halogenated hydrocarbon (12%) and OVOCs (10%). Higher volume percentages of  
 283 alkanes and alkenes in Case 1 were similar to the results in the gasoline evaporation  
 284 site in winter (Niu et al., 2022). Equivalent volume contribution of halogenated  
 285 hydrocarbon and OVOCs (15%) were found in Case 2, followed by alkenes (10%).

286 Though the volume contributions of aromatics were the lowest (6% in Case 1 and 7%  
 287 in Case 2), the highest increase ratio was found from clean days to polluted episodes.

288 Table 1 The average concentrations of meteorological parameters and pollutants during different  
 289 processes.

Category	Entire process	Infection period	Recovery period	Case 1	Case 2
	N = 62 days	N = 31 days	N = 31 days	N = 6 days	N = 8 days
WS (m/s)	1.3 ± 0.6	1.4 ± 0.6	1.3 ± 0.6	1.2 ± 0.9	0.9 ± 0.7
T (°C)	5.0 ± 2.5	4.7 ± 1.7	5.4 ± 3.1	6.1 ± 2.2	7.4 ± 3.5
RH (%)	38.9 ± 16.7	37.6 ± 15.5	40.2 ± 18.2	55.7 ± 14.7	42.0 ± 12.1
TVOCs (ppbv)	36.1 ± 21.0	31.9 ± 18.1	39.8 ± 22.4	37.6 ± 27.0	68.2 ± 19.6
SO <sub>2</sub> (µg/m <sup>3</sup> )	11.4 ± 2.7	10.2 ± 2.8	12.7 ± 2.3	11.0 ± 3.7	16.2 ± 6.1
NO <sub>2</sub> (µg/m <sup>3</sup> )	47.2 ± 10.0	46.8 ± 8.6	47.8 ± 11.7	62.7 ± 20.5	65.0 ± 21.3
CO (mg/m <sup>3</sup> )	0.9 ± 0.2	0.8 ± 0.2	1.1 ± 0.2	1.2 ± 0.5	1.3 ± 0.4
O <sub>3</sub> (µg/m <sup>3</sup> )	34.9 ± 6.0	31.1 ± 4.5	39.0 ± 4.6	21.8 ± 23.7	32.5 ± 29.6
PM <sub>2.5</sub> (µg/m <sup>3</sup> )	111.5 ± 45.1	86.6 ± 34.6	138.3 ± 39.6	142.5 ± 67.4	181.5 ± 82.7

290 Table 2 Concentration of VOC species during different processes (ppbv).

Category	Entire process	Infection period	Recovery period	Case 1	Case 2	Clean days
TVOCs	36.1 ± 21.0	31.9 ± 18.1	39.8 ± 22.4	48.4 ± 20.4	67.6 ± 19.6	17.5 ± 9.5
alkanes	16.8 ± 9.2	15.0 ± 8.4	18.4 ± 9.5	23.1 ± 10.0	29.5 ± 8.4	9.2 ± 5.6
alkenes	4.1 ± 2.7	3.8 ± 2.6	4.4 ± 2.7	6.5 ± 2.9	7.0 ± 2.6	1.7 ± 1.3
alkynes	3.1 ± 2.0	2.7 ± 1.7	3.4 ± 2.1	4.3 ± 2.0	5.8 ± 1.9	1.3 ± 0.8
aromatics	2.1 ± 2.0	1.8 ± 1.5	2.3 ± 2.2	3.0 ± 1.8	4.9 ± 2.8	0.7 ± 0.5
halogenated hydrocarbon	5.4 ± 3.3	4.4 ± 2.3	6.2 ± 3.8	6.0 ± 1.9	10.7 ± 3.6	2.7 ± 1.4
OVOCs	4.6 ± 3.2	3.5 ± 2.7	5.1 ± 3.5	5.0 ± 2.4	9.7 ± 2.8	1.9 ± 1.1

### 291 3.2 Source Analysis of VOCs

292 Specific VOC ratios can be used for initial source identification of VOCs and  
 293 determination of photochemical ages of air masses (Monod et al., 2001; An et al., 2014;  
 294 Li et al., 2019). In this study, the ratios of toluene/benzene (T/B), isopentane/n-pentane,  
 295 isobutane/n-butane, and m/p-xylene/ethylbenzene (X/E) were selected to initially  
 296 identify the potential sources of VOCs (Fig. 3). Concentrations of selected pollutants  
 297 and ratios used are shown in Table S1.

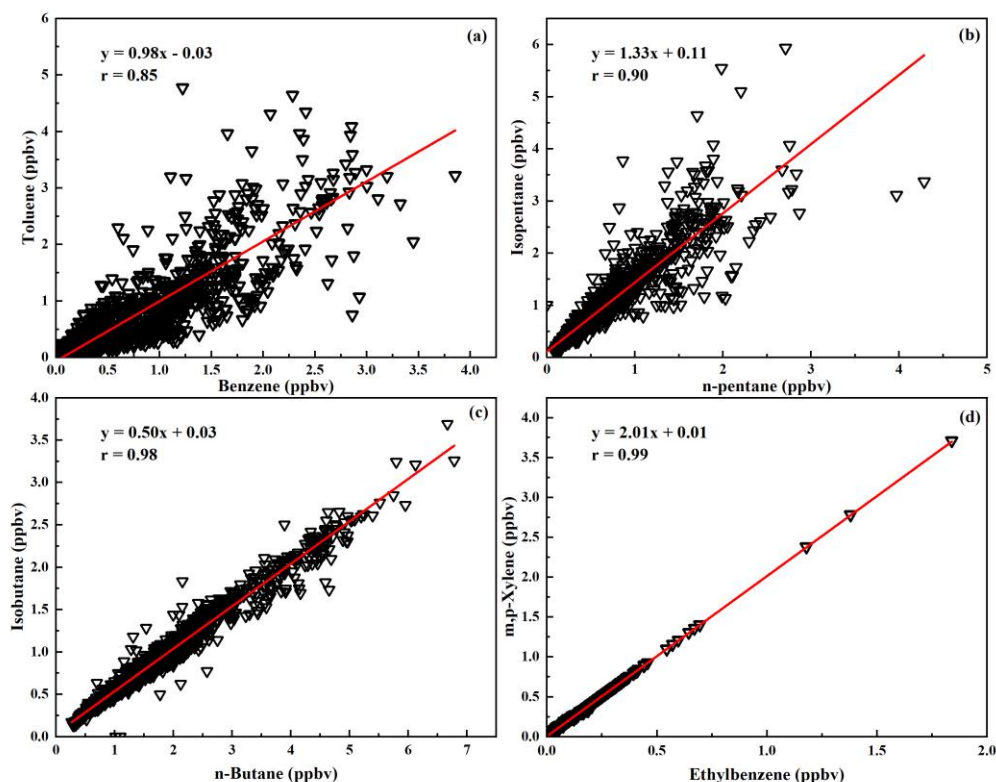
298 Toluene-to-benzene ratio (T/B ratio) was widely used to assess the relative

299 importance of different sources. Specifically, T/B ratio with the value of 1.3–3.0 was  
300 observed in vehicle emissions for vehicles with different fuel types (Schauer et al., 2002;  
301 Wang et al., 2015). The reported T/B ratio for combustion processes was between 0.13  
302 and 0.7 (Li et al., 2011; Wang et al., 2014). The average T/B value for the entire period  
303 was 1.0, indicating that both traffic emissions and combustion are significant sources  
304 of VOCs.

305 The isopentane/n-pentane concentration ratios of 0.6-0.8 represent mainly coal  
306 combustion emissions, ratios of 0.8-0.9 represent liquefied petroleum gas (LPG)  
307 emissions, 2.2-3.8 represent vehicle exhaust emissions, and 1.8-4.6 represent fuel  
308 evaporation (Conner et al., 1995; Liu et al., 2008; Li et al., 2019). The overall ratio of  
309 isopentane/n-pentane is 1.4, indicating that pentane is mainly derived from the  
310 combined effects of liquid petrol and fuel evaporation.

311 Isobutane/n-butane concentration ratios of 0.2-0.3 represent vehicle emissions,  
312 0.4-0.6 represent LPG usage, and 0.6-1.0 represent natural gas emissions (Russo et al.,  
313 2010; Zheng et al., 2018). The ratio of isobutane/n-butane in this study was 0.50, which  
314 suggests that the VOC concentrations at the observation sites are influenced by natural  
315 gas emissions (Shao et al., 2016; Zeng et al., 2023).

316 The ratio of X/E can be used to infer the photochemical age of the air mass. X/E  
317 ratios around 2.5-2.9 are typical of urban areas, indicating that VOCs are mainly from  
318 the urban area (fresh air mass) (Kumar et al., 2018). When this ratio is significantly  
319 lower than 3, it indicates that VOCs are mainly transported from distant sources (aging  
320 air masses) (Kumar et al., 2018). The average X/E value in this study was 2.0 (Fig.  
321 3(d)), indicating low photochemical activity and aging of the air mass at the observation  
322 site. Potential source analyses also indicate that air masses are affected by long-range  
323 transport (Fig. S4).



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Fig. 3. Correlation analysis between specific VOC species.

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Figure 4 shows the chemical profiles of individual VOCs resolved by the PMF model during the entire observation period. These five factors eventually selected as potential sources for the observed VOCs are: (1) Fuel evaporation; (2) Solvent usage; (3) Vehicular emission; (4) Industrial source; and (5) Combustion. These 5 factors have been commonly reported before, e.g., in Shijiazhuang, northern China (Guan et al, 2023) and in Beijing (Cui et al., 2022).

332

Alkanes of C4-C6 substances were predominant in factor 1, including 2-methylpentane, 3-methylpentane, isobutane, n-butane, isopentane and n-pentane from oil and gas (Xiong et al., 2020). Fig. S5 shows that emissions from this source peak at midday, when fuel volatilization is high, The CPF plot shows that south-east is the dominant direction at wind speeds of less than 2 m/s (Fig. 5a). Therefore, factor 1 was identified as the source of oil and gas volatilization.

338

The contribution of benzene, toluene, methylene chloride, 1,2-dichloroethane and ethyl acetate was high in factor 2. It has been shown that benzene, toluene, ethylbenzene, and xylene is an important component in the use of solvents (Li et al., 2015); methylene chloride is often used as a chemical solvent, while esters are mostly used as industrial solvents or adhesives (Li et al., 2015). Factor 2 is determined to be solvent usage source.

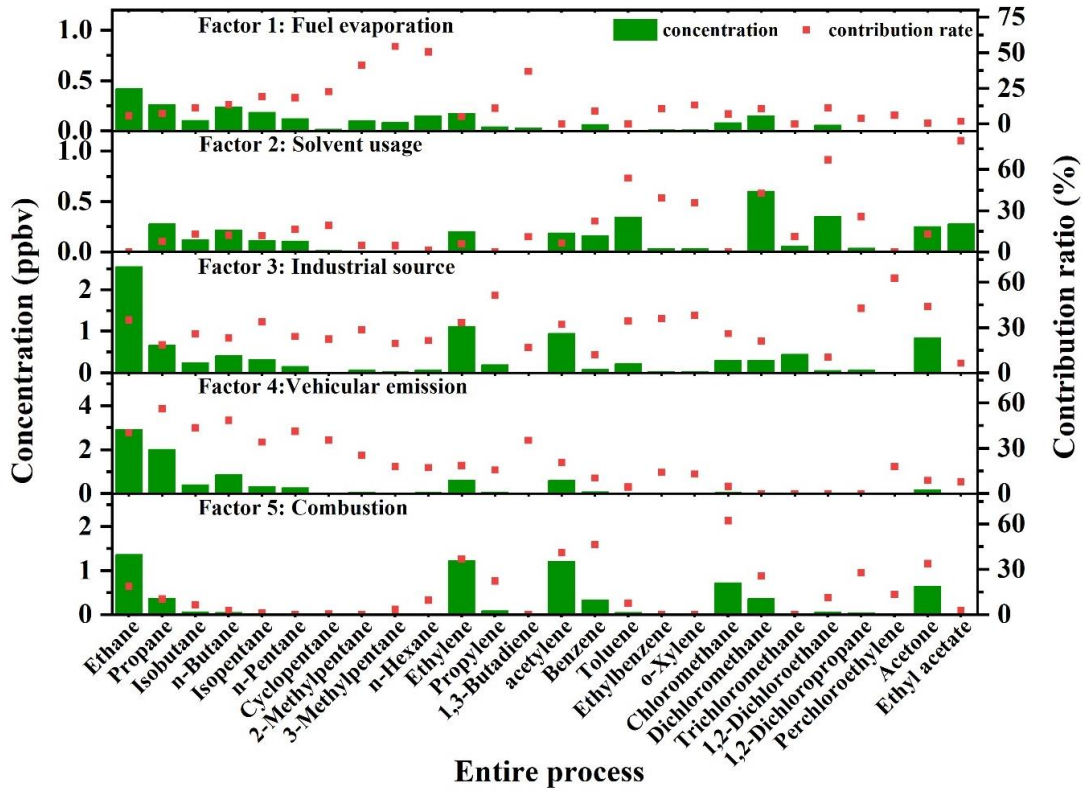
342

343 The CPF plot shows that local sources with wind speeds less than 1 m/s are the main  
344 sources (Fig. 5b).

345 Factor 3 contains predominantly C3-C8 alkanes, olefins and alkynes, and  
346 relatively high concentrations of benzene. These substances are usually emitted by  
347 industrial processes (Shao et al., 2016), so Factor 4 is defined as an industrial source.  
348 The CPF plots indicate that a local source at low wind speeds is the dominant sources  
349 (Fig. 5c).

350 Factor 4 is characterized by relatively high levels of C2-C6 low-carbon alkanes  
351 (ethane, propane, isopentane, n-pentane, isobutane and n-butane), olefins (ethylene and  
352 propylene), and benzene and toluene, which are important automotive exhaust tracers  
353 (Song et al., 2021; Zhang et al., 2021b). Ethylene and propylene are important  
354 components derived from vehicle-related activities. Previous studies of VOCs in  
355 Zhengzhou have shown a high percentage of VOCs emitted from gasoline vehicles,  
356 with the main source of alkanes being on-road mobile sources (Bai et al., 2020). The  
357 daily variation of this source in Fig. S5 shows a bimodal trend, with peaks occurring in  
358 the morning and evening peaks of traffic, consistent with motor vehicle emissions. Fig.  
359 5d shows that this source is mainly from the west where wind speeds are below 2 m/s,  
360 and in this direction, there are a number of urban arterial roads with high traffic volumes.  
361 Therefore, factor 4 was defined as vehicular emission source.

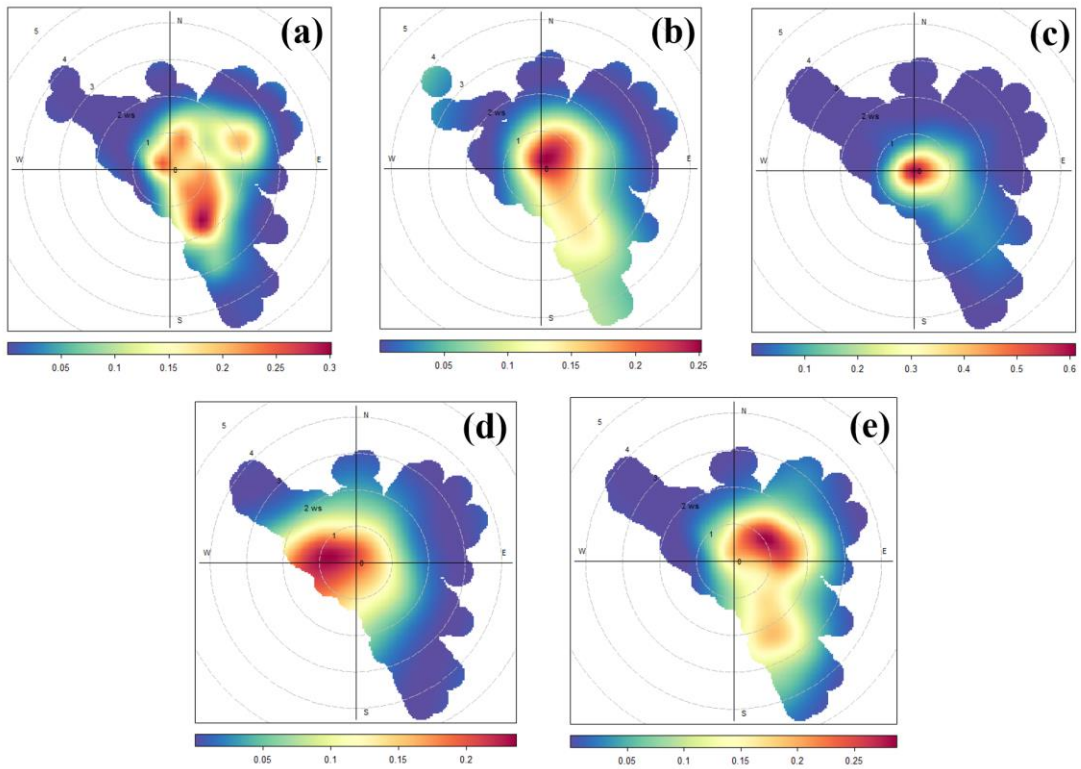
362 The highest contribution to factor 5 is chloromethane (62%). Benzene (46%) and  
363 acetylene (41%) also contribute highly to factor 5. Chloromethane is the key tracer for  
364 biomass combustion and acetylene is the key tracer for coal combustion (Xiong et al.,  
365 2020). Therefore, factor 5 is defined as a combustion source. The CPF plot shows that  
366 at wind speeds below 2 m/s, the north-east direction is the dominant source direction  
367 (Fig. 5e).



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Fig. 4. Concentration of VOC species in each factor and contribution to each source.



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Note: a: Fuel evaporation; b: Solvent usage; c: Industrial source; d: Vehicular emission; e: Combustion.



373

Fig. 5. CPF plots of five VOCs sources obtained using the PMF model.

374 Figure S6 compares the differences in PMF factor/source profiles during the peak  
375 of Omicron infection with those during the recovery phase after the peak, as well as  
376 between contaminated and clean days. We present the concentrations of the five main  
377 VOCs in all five factors in Table S2. Ethane (vehicular emission), 2-methylpentane  
378 (fuel evaporation), benzene (industry source), chloromethane (combustion), and ethyl  
379 acetate (solvent usage) were selected as tracers for five sources. Ethane concentration  
380 in Case 2 (5.9 ppbv) is much higher than in other processes, and ethane concentration  
381 during the recovery period (3.4 ppbv) is also higher than during the infection period  
382 (2.4 ppbv), which may to some extent reflect increased vehicular emissions during the  
383 recovery period.

384 Concentrations of most species were significantly higher during the recovery  
385 period than during the infection period. The representative pollution processes in both  
386 periods showed the same results as well, with a 79% higher concentration of TVOCs in  
387 Case 2 (65.1 ppbv) compared to Case 1 (36.3 ppbv) (Fig. 6). While in Case 1 industry  
388 was the dominant source of VOCs, by Case 2 motorized sources reached a  
389 concentration value of 21.2 ppbv, accounting for 33% of the observed VOCs, and  
390 became the dominant source of emissions. This is consistent with the fact that people's  
391 mobility activities have increased after the epidemic has entered the recovery period.  
392 As a group of VOCs species with the highest concentration share, ethane and propane  
393 contributed more to the clean days motor vehicle source than other processes, which  
394 also resulted in a 34% clean days motor vehicle source share.

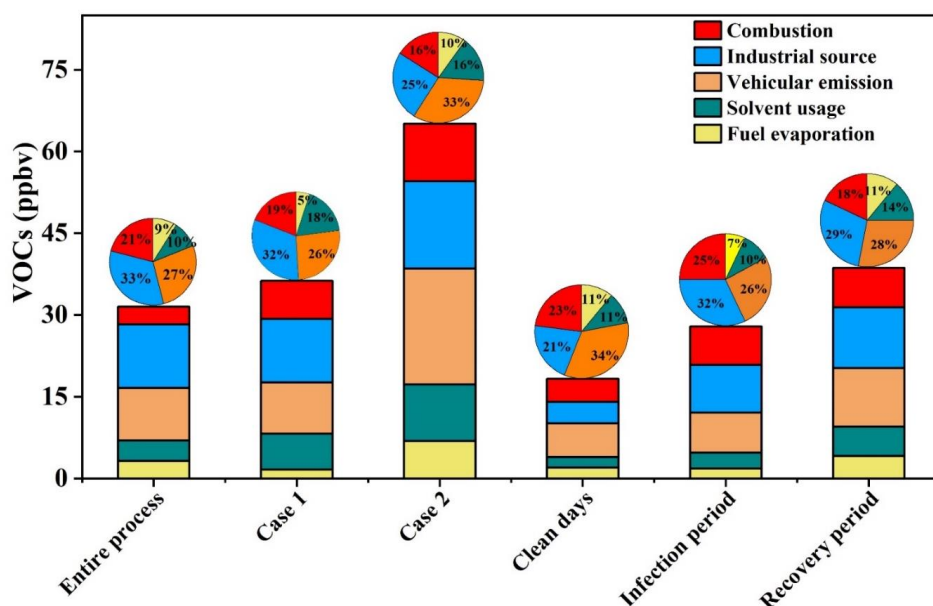


Fig. 6. Contribution of each to VOCs for different processes.

### 3.3 SOAP

VOCs are estimated to contribute about 16–30% or more of  $PM_{2.5}$  by mass through SOA production (Huang et al., 2014). Therefore, by calculating the SOAP value, the influence of different sources on  $PM_{2.5}$  production can be reflected to a certain extent.

We have included quantitative analysis for SOAP as well. Fig. 7 shows the SOAP concentrations and contribution rates of the top ten species throughout the entire process, during two pollution processes, and clean days. The top ten species all reached close to 100% of the total SOAP contribution, with Case 1 reaching 98%. In each process, the composition of the top ten substances is essentially the same. Aromatic hydrocarbons contributed the most, with BTEX always occupying the top five positions and toluene the most. The SOAP values of the top ten contributing species for the two polluting processes are shown in Tables S3 and S4. Toluene, the highest contributing species, reached a SOAP value of  $49.4 \mu\text{g}/\text{m}^3$  in the most polluted Case 2, which was 3.2 times higher than the SOAP sum of all species on the clean day ( $15.5 \mu\text{g}/\text{m}^3$ ). The SOAP value for Case 1, which is also a contaminated process, was  $67 \mu\text{g}/\text{m}^3$ , and the main species (m/xylene:  $9.8 \mu\text{g}/\text{m}^3$ , benzene:  $8.5 \mu\text{g}/\text{m}^3$ ) including toluene ( $34.6 \mu\text{g}/\text{m}^3$ ) were lower than those for Case 2 (m/xylene:  $19.4 \mu\text{g}/\text{m}^3$ , benzene:  $13.4 \mu\text{g}/\text{m}^3$ ).

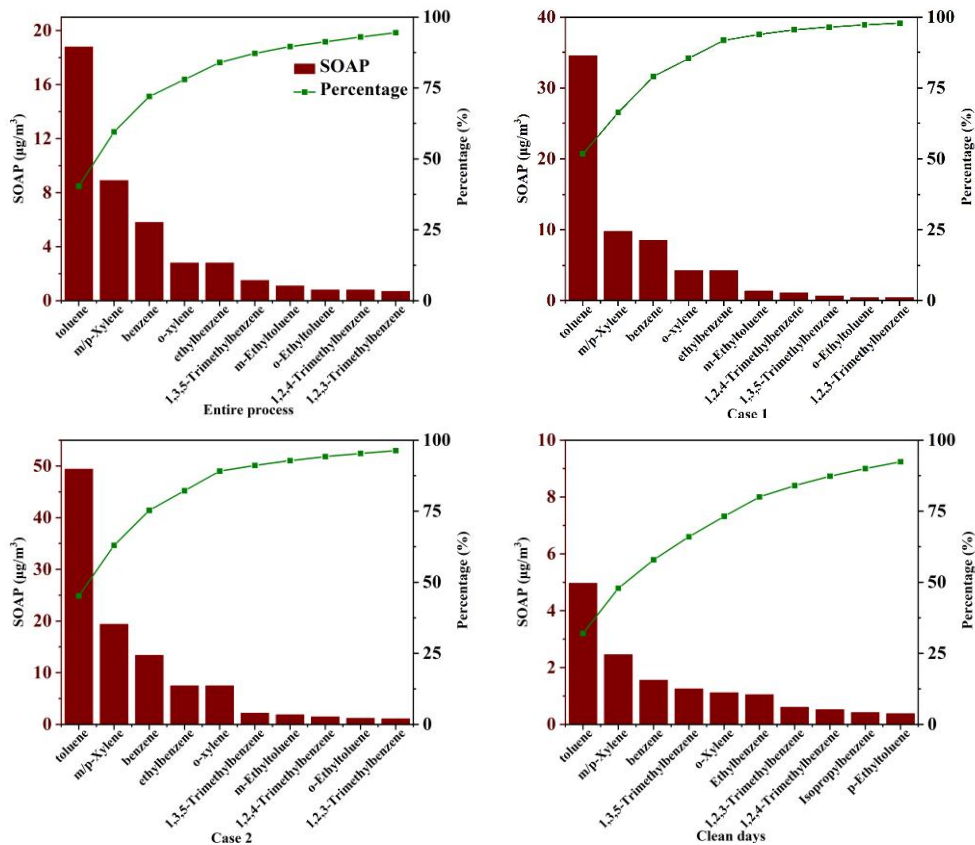


Fig. 7. SOAP dominant species in different processes

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Figure 8 shows the SOAP calculated after source resolution of the two pollution processes by PMF for clean days, respectively. In Case 1, industrial source is the dominant source with a contribution ratio of 63%. In Case 2, the pollution sources exhibit a more evenly distributed contribution, where the solvent usage and fuel evaporation sources emerge as the primary contributors to SOAP, with their respective contribution levels rising to 32% and 26%. The clean day result with a SOAP of 8.8 µg/m³ also indicates that industrial and solvent usage sources are the most dominant SOAP sources. The primary sources of aromatic compounds, which are the most significant contributors to SOAP, are solvent usage and industrial process emissions. This finding aligns with the results of other studies (Wu et al., 2017). Consequently, it is imperative to implement measures to reduce PM<sub>2.5</sub> pollution by regulating emissions from industrial and solvent usage sources.

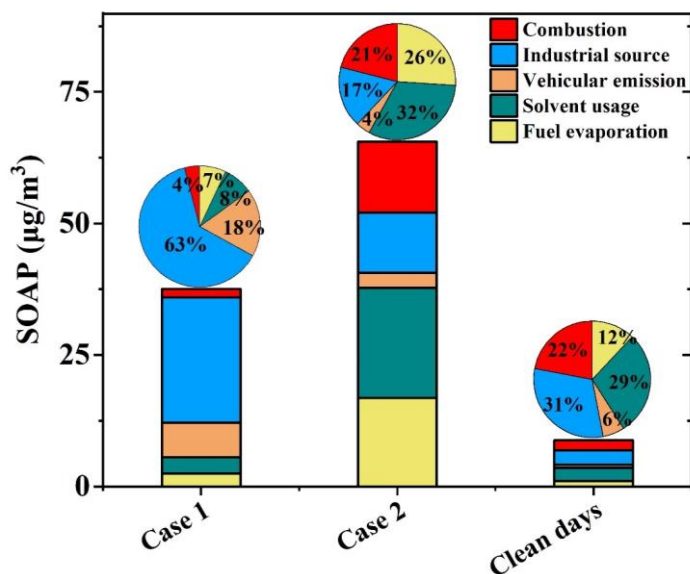


Fig. 8. SOAP value and contribution ratio of each process

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#### 431 4. Conclusions

432 Continuous observation of VOCs during the infection of the Omicron epidemic  
 433 was carried out at an urban site in polluted Zhengzhou from December 1, 2022, to  
 434 January 31, 2023. The daily average concentration of PM<sub>2.5</sub> ranged from 53.5 to 239.4  
 435 µg/m<sup>3</sup> with an average value of 111.5 ± 45.1 µg/m<sup>3</sup> during the whole period. The  
 436 concentration of TVOCs ranged from 15.6 to 57.1 ppbv with an average of 36.1 ± 21.0  
 437 ppbv, higher than the same period in last year (27.9 ± 12.7 ppbv, Lai et al., 2024).

438 Two representative contamination processes were identified (Case 1 during the  
 439 infection period and Case 2 during the recovery period). The concentration of TVOCs  
 440 in Case 1 and Case 2 were 48.4 ± 20.4 and 67.6 ± 19.6 ppbv, respectively, increased by  
 441 63% and 188% compared with values during clean days. The average concentrations  
 442 of PM<sub>2.5</sub> and TVOCs during Case 2 were 1.3 and 1.8 times of the values in Case 1. The  
 443 highest volume contributions of alkanes were found both in Case 1 (48%) and Case 2  
 444 (44%). Though the volume contribution of aromatics were the lowest (6% in Case 1  
 445 and 7% in Case 2), the highest increase ratio was found from clean days to polluted  
 446 episodes.

447 Local sources were initially identified through T/B, isopentane/n-pentane,  
 448 isobutane/n-butane, and X/E ratios. The average X/E value was 2.0, indicating that  
 449 measured levels of airborne VOCs were influenced by emissions from remote sources  
 450 and urban areas. The PMF receptor modeling yielded five major sources of pollution,  
 451 which included industrial emissions (32%), vehicular emissions (27%), combustion

452 (21%), solvent usage (11%), and fuel evaporation (9%). Significant differences were  
453 observed in the sources of VOCs across different pollution periods. In Case 1, industrial  
454 emissions constituted the largest contributor, accounting for 32% of the total VOCs. In  
455 contrast, in Case 2, the proportion of vehicular emissions increased to 33%, becoming  
456 the primary source of VOCs.

457 Aromatic compounds are the main contributors to SOAP, with BTEX being the  
458 main contributor during the entire period. SOAP values reached 37.6 and 65.6  $\mu\text{g}/\text{m}^3$ ,  
459 respectively in Case 1 and Case 2. In Case 1, industrial source accounted for a  
460 substantial majority (63%, 23.8  $\mu\text{g}/\text{m}^3$ ), while vehicular source, as the second most  
461 significant contributor, made up only 18%. In Case 2, the distribution of contribution  
462 rates was more uniform, with solvent usage source and fuel evaporation source  
463 becoming the primary contributors to SOAP, at 32% (20.9  $\mu\text{g}/\text{m}^3$ ) and 26% (16.8  $\mu\text{g}/\text{m}^3$ ),  
464 respectively. The SOAP result for the Clean Day was 8.8  $\mu\text{g}/\text{m}^3$ , with industrial source  
465 and solvent usage remaining the primary contributors. Consequently, it is of paramount  
466 importance to regulate emissions from the industrial and solvent usage sectors with the  
467 objective of reducing PM<sub>2.5</sub> pollution.

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