1	The variations of VOCs based on the policy change of
2	<b>Omicron in traffic-hub city Zhengzhou</b>
3	Bowen Zhang <sup>1, 3</sup> , Dong Zhang <sup>2, 3</sup> , Zhe Dong <sup>2, 3</sup> , Xinshuai Song <sup>1, 3</sup> , Ruiqin Zhang <sup>1, 3</sup> ,
4	Xiao Li <sup>1, 3,</sup> *
5	<sup>1</sup> School of Ecology and Environment, Zhengzhou University, Zhengzhou 450001,
6	China
7	<sup>2</sup> College of Chemistry, Zhengzhou University, Zhengzhou 450001, China
8	<sup>3</sup> Institute of Environmental Sciences, Zhengzhou University, Zhengzhou 450001,
9	China
10	Correspondence to: Xiao Li, E-mail address: lixiao9060@zzu.edu.cn
11	Abstract: Online volatile organic compounds (VOCs) were monitored before and
12	after the Omicron policy change at an urban site in polluted Zhengzhou from December
13	1, 2022, to January 31, 2023. The characteristics and sources of VOCs were
14	investigated. The daily mean concentrations of PM2.5 and total VOCs (TVOCs) ranged
15	from 53.5 to 239.4 $\mu g/m^3$ and 15.6 to 57.1 ppbv, respectively, with mean values of 111.5
16	$\pm$ 45.1 $\mu g/m^{3}$ and 36.1 $\pm$ 21.0 ppbv, respectively, throughout the period. Two severe
17	pollution events (designated as Case 1 and Case 2) were identified in accordance with
18	the National Ambient Air Quality Standards (NAAQS) (China's National Ambient Air
19	Quality Standards (NAAQS) from 2012). Case 1 (December 5 to December 10, PM <sub>2.5</sub>
20	daily mean = 142.5 $\mu$ g/m <sup>3</sup> ) and Case 2 (January 1 to January 8, PM <sub>2.5</sub> daily mean =
21	181.5 $\mu$ g/m <sup>3</sup> ) occurred during the infection period (when the policy of "full nucleic acid
22	screening measures" was in effect) and the recovery period (after the policy was
23	cancelled), respectively. The $PM_{2.5}$ and TVOCs values for Case 2 are, respectively, 1.3
24	and 1.8 times higher than those for Case 1. The results of the positive matrix factor
25	modeling demonstrated that the primary source of volatile organic compounds (VOCs)
26	during the observation period was industrial emissions, which constituted 32% of the
27	total VOCs, followed by vehicle emissions (27%) and combustion (21%). In Case 1,
28	industrial emissions constituted the primary source of VOCs, accounting for 32% of the
29	total VOCs. In contrast, in Case 2, the contribution of vehicular emission sources
30	increased to 33% and became the primary source of VOCs. The secondary organic
31	aerosol formation potential for Case 1 and Case 2 were found to be 37.6 $\mu g/m^3$ and 65.6
32	$\mu$ g/m <sup>3</sup> , respectively. In Case 1, the largest contribution of SOAP from industrial sources
33	accounted for the majority (63%, 23.8 $\mu$ g/m <sup>3</sup> ), followed by vehicular sources (18%).

After the end of the epidemic and the resumption of productive activities in the society, the difference in the proportion of SOA generated from various sources decreased. Most of the SOAP came from solvent use and fuel evaporation sources, accounting for 32% (20.9 μg/m<sup>3</sup>) and 26% (16.8 go/m<sup>3</sup>), respectively. On days with minimal pollution, industrial sources and solvent use remain the main contributors to SOA formation. Therefore, regulation of emissions from industry, solvent-using industries and motor vehicles need to be prioritized to control the PM<sub>2.5</sub> pollution problem.

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

43 Matrix Factorization model; Secondary organic aerosol formation potential;

## 44 **1. Introduction**

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 47 ozone (O<sub>3</sub>) and secondary organic aerosol (SOA), resulting in regional air pollution (Li 48 et al., 2019; Hui et al., 2020). The problem of O<sub>3</sub> pollution has been plaguing major 49 urban agglomerations in China (Zheng et al., 2010; Li et al., 2014; Wang et al., 2017). 50 SOA is an important component of fine particulate matter  $(PM_{2.5})$  and contributes 51 significantly to haze pollution (Liu et al., 2019). PM<sub>2.5</sub> remains the most significant air 52 pollutant in many Chinese cities for years (Shao et al., 2016; Wu et al., 2016). In 53 addition, VOCs, represented by the benzene homologues, can cause damage to kidneys, 54 liver, and nervous system of humans when they enter the body (Zhang et al., 2018).

55 Studies have shown that the most common VOC components in China are alkanes, olefins, aromatic hydrocarbons, oxygenated VOCs (OVOCs), and halogenated 56 57 hydrocarbons, among which alkanes are the most abundant species (Liu et al., 2020; Zhang et al., 2021a). VOCs in the atmosphere have a wide range of sources, and VOCs 58 59 in different regions are affected by multiple factors such as local geography, climate, 60 and human activities (Mu et al., 2023; Zou et al., 2023). The above reasons lead to significant regional and seasonal differences in the characteristics of VOCs (Song et al., 61 62 2021). For example, the annual average concentration of VOCs in the coastal 63 background area of the Pearl River Delta is 9.3 ppby. The seasonal variation trend of 64 VOCs is high in autumn and winter and low in summer (Yun et al., 2021). In contrast, the average VOC concentration in autumn and winter in Beijing was  $22.6 \pm 12.6$  ppbv, 65 66 and the VOC concentration in the winter heating period was twice that in the autumn 67 non-heating period (Niu et al., 2022).

68 Moreover, the sources of VOC components in different regions are also related to 69 the local industrial structure and living habits. In rural areas of North China Plain in 70 winter, it is found that the SOA formation potential (SOAP) of VOCs under low  $NO_x$ 71 conditions is significantly higher than that under high NO<sub>x</sub> conditions, and the increase 72 of aromatic hydrocarbon emissions caused by coal combustion is the main reason for 73 the higher SOAP in winter (Zhang et al., 2020). Li et al. (2022) found that the average 74 increased concentration of acetylene was 4.8 times from autumn to winter in the 75 Guanzhong Plain, indicating that fuel combustion during the heating period in winter 76 has a significant impact on the composition of VOCs. In contrast, continuous observations conducted by Zhou et al. (2022) in the suburbs of Dongguan in summer found that industrial solvent usage, liquefied petroleum gas (LPG) and oil and gas volatilization were the main sources of VOCs. The results highlighted a wide variation of characteristics, sources and chemical reactions of VOCs in the atmosphere thus it is necessary to investigate VOCs in different cities when formulating control measures.

82 Zhengzhou, as the capital of Henan Province, is an important transportation hub 83 and economic center in the Central Plains region. Zhengzhou is currently facing 84 significant air pollution problems, with the Air Quality Index at the bottom of the 85 national ranking of 168 cities for many years. In January 2023, for example, the number of polluted days with PM<sub>2.5</sub> as the primary pollutant was 17, and the daily average value 86 298  $\mu g/m^3$ 87 of PM<sub>2.5</sub> reached а maximum of (https://www.aqistudy.cn/historydata/daydata.php?city=%E9%83%91%E5%B7%9E 88 &month=202301, Accessed Jan 2024), which is almost 300% higher than the Chinese 89 90 daily average standard (grade II, 75  $\mu$ g/m<sup>3</sup>). The studies of VOCs were carried out in 91 Zhengzhou in recent years, which focused on the characteristics and sources of VOCs 92 during pollution episodes (Lai et al., 2024) or before the coronavirus epidemic outbreak 93 (Li et al., 2020; Zhang et al., 2021b). While some atmospheric VOCs studies involving 94 the impact of Covid-19 lockdown have been performed in India (Singh et al., 2023a), in China (e.g., Pei et al., 2022; Jensen et al., 2023; Zuo et al., 2024), or with respect to 95 96 toluene, benzene, m/p-xylene and ethylbenzene only (e.g., Sahu et al., 2022; Singh et 97 al., 2023b), a gap persisted in the investigation of VOCs due to the impact of 98 abolishment of China's zero-policy. Furthermore, some studies have discussed the 99 impact of changes in human production activities on air pollution during and after the 100 outbreak of the coronavirus disease (e.g., Ma et al., 2022; Jiang et al., 2023; Song et al., 101 2023), but as mentioned earlier, only a few studies with in-depth exploration of the 102 changes in VOCs and none dealing with ending the zero-Covid policy during Omicron 103 variant infection period.

In this study, we conducted continuous online observations of VOCs during the polluted winter season at an urban site in Zhengzhou. The study covered the period following the removal of lockdown measures. We focused on pollution events when the daily average  $PM_{2.5}$  concentration exceeded 75 µg/m<sup>3</sup> (China's Class II standard) for more than three consecutive days. Days with  $PM_{2.5}$  concentrations below 35 µg/m<sup>3</sup> (China's Class I standard) were classified as clean days. During this period, China lifted zero-COVID strategies, announcing the '10 measures' for optimizing COVID-19 rules 111 on December 7, 2022 (http://www.news.cn/politics/2022-12/07/c 1129189285.htm, 112 Accessed Jan 2024). Zhengzhou's epidemic prevention and control measures changed 113 with the issuance of Circular No. 163 on December 4, 2022, which allowed the 114 reopening of closed public places. As a result, movement within Zhengzhou increased 115 and social production resumed. Our research specifically examines the period dominated by the COVID-19 Omicron variant. where they demonstrate notable 116 117 differences from the early virus strains (i.e., original SARS-CoV-2 virus and Delta) in 118 terms of geographical transmission, the scale of the infected population, and symptom 119 manifestation (Petersen et al., 2022; Merino et al., 2023).

120 After the quarantine policy was lifted, people basically rested at home due to infection or fear of infection with Omicron. The resumption of normal production and 121 122 life depends on herd immunization. This outbreak event is the longest in duration and 123 the largest in number of infections since the 2020 outbreak of the novel coronavirus in Zhengzhou. It would be beneficial to investigate the impact of this event on emissions 124 125 related to transportation and industrial production. This change is worth exploring in 126 terms of its impact on transportation and industrial production emissions. Therefore, 127 the characteristics and variations of VOCs during different periods were investigated to 128 assess their impact on the formation of SOA and to provide data support for future 129 pollution control policies in Zhengzhou.

# 131 **2. Materials and methods**

## 132 **2.1 Sample collection and Chemical analysis**

133 The online VOCs observation station is located on the roof of the Zhengzhou 134 Environmental Protection Monitoring Center, which is in the urban area. The sampling 135 site is close to main roads on three sides (150 m away from Funiu Road on the east side, 136 200 m away from Qinling Road on the west side, and connected to Zhongyuan Road 137 on the south side), and surrounded by residential areas and commercial areas without 138 other large nearby stationary sources. The sampling period for this study was from 139 December 1, 2022, to January 31, 2023, and serious PM<sub>2.5</sub> pollution in Zhengzhou was 140 of frequent during December occurrence and January. 141 (https://www.aqistudy.cn/historydata/monthdata.php?city=%e9%83%91%e5%b7%9e 142 #:~:text=%E7%94%9F%E5%91%BD%E6%9D%A5%E6%BA%90%E8%87%AA% 143 E7%84%B6%EF%BC%8C%E5%81%A5). Apart from a brief occurrence of rain and 144 snow on December 25, the sampling days were either sunny or cloudy. The wind speed 145 (WS), temperature (Temp) and relative humidity (RH) during this period were  $1.3 \pm 0.9$ 146 m/s,  $5.3 \pm 3.2$  °C and  $38.9 \pm 19.0\%$ ), respectively, similar to the values observed in 147 previous years in Zhengzhou. It is interesting to point out that the sampling period in 148 the present study covered the entire infection period of Omicron in Zhengzhou, 149 including the phase of surge in infected population (Infection period, from 2022.12.01 150 to 2022.12.31) and restoration of production and livelihood phase (Recovery period, 151 from 2023.1.1 to 2023.1.31 in 2023) (Fig. S1, Chinese Center for Disease Control and 152 Prevention, 2023).

153 The VOCs were measured hourly using a GC-FID/MS (TH-PKU 300 b, Wuhan 154 Tianhong Instruments Co., China). The instrument TH-PKU300b includes electronic 155 refrigeration ultra-low temperature pre-concentration sampling system, analysis system 156 and system control software. The ambient VOCs in the first 5 minutes of each hour 157 were collected by the sampling system and then entered the concentration system. Under low temperature conditions, the VOCs samples collected were frozen in the 158 159 capillary capture column, and then quickly heated and resolved, so that the compounds entered the analysis system. After separation by chromatographic column, the 160 161 compounds were monitored by FID and MS detectors. During the detection process, 162 the atmospheric samples collected undergo analysis through two distinct pathways. C2-163 C5 hydrocarbons are analyzed using FID, while C5-C12 hydrocarbons, halocarbons, and OVOCs are analyzed with a MS detector. After excluding species with missing data exceeding 10%, the detected volatile organic compounds include 29 alkanes, 11 alkenes, 17 aromatics, 35 halocarbons, 12 OVOCs, 1 alkyne (acetylene), and 1 sulfide ( $CS_2$ ) with a total of 106 compounds. A detailed description of the instrumentation can be found in our previous study (Zhang et al., 2021b; Shi et al., 2022; Zhang et al., 2024).

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

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

## 180 **2.2 Positive Matrix Factorization (PMF) model**

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

185

186 
$$x_{ij} = \sum_{k=1}^{p} g_{ik} f_{kj} + e_{ij}$$
(1)

187

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

194 
$$Q = \sum_{i=1}^{n} \sum_{j=1}^{m} \left[ \frac{x_{ij} - \sum_{k=1}^{p} g_{ik} f_{kj}}{u_{ij}} \right]^2$$
(2)

195

196 where *n* and *m* denote the number of samples and VOC species, respectively.

197 Concentrations and uncertainty data are required for the PMF model. In this study, the median concentration of a given species is used to replace missing values with an 198 199 uncertainty of four times of the median values; data less than the Method Detection 200 Limit (MDL) were replaced with half the MDL, with an uncertainty of 5/6 of the MDL; 201 and the uncertainty for values greater than the MDL was calculated using Eq. (3). In 202 Eq. (3), *EF* is error fraction, expressed as the precision of VOCs species, and the setting 203 range can be adjusted from 5 to 20% according to the concentration difference (Buzcu 204 et al., 2006; Song et al., 2007); and  $c_{ij}$  is the concentration of species j in sample i:

205 
$$U_{ij} = \sqrt{\left(EF \times c_{ij}\right)^2 + (0.5 \times MDL)^2}$$
(3)

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

208 
$$U_{ij} = \left(\frac{5}{6}\right) MDL \tag{4}$$

209 VOC species and concentration input into PMF were carefully selected to ensure 210 the accuracy of the PMF results. Species were excluded when over 25% of the samples 211 were missing or concentrations values were below the MDL (Gao et al., 2018); VOCs 212 with a short lifetime in the atmosphere were also excluded unless they are source-213 relative species (Zhang et al., 2014; Shao et al., 2016). After that, retained VOC species 214 were categorized according to the signal-to-noise ratio (S/N) with S/N < 0.2 species 215 categorized as bad, 0.2 < S/N < 2 species categorized as weak; and S/N > 2 species 216 categorized as strong (Shao et al., 2016).

We used displacement of factor elements (DISP) to assess PMF modelling 217 218 uncertainty (for a description, see Paatero et al. (2014)). Q was less than 1% and no swaps occurred for the small est dQ<sup>max</sup> in DISP. Fpeak values from -2 to 2 were tested 219 220 to explore the rotational stability of the solutions.  $Q_{true}/Q_{exp}$  is lowest when Fpeak = 0, 221 so we chose the PMF results for that case (Fig. S2a). After examining 3-8 factors, 20 222 base runs with 5 factors eventually selected to represent final result. We provide an explanation of factor selection in the supplementary materials. Fig. S2(b) includes 223 224 Q<sub>true</sub>/Q<sub>exp</sub>, Q<sub>robust</sub>/Q<sub>exp</sub> for factors 3-8. The slopes of these two ratios in changed at five factors, and we found that five factors were more realistic after repeated comparisonsof the results at four, five and six factors.

# 227 **2.3 SOA generation potential**

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

231

232 
$$SOAPF_i = \frac{VOCs \ component \ i \ to \ SOA \ mass \ concentration \ increments}{Toluene \ to \ SOA \ mass \ concentration \ increment} \times 100$$
 (5)

233

SOAPF<sub>i</sub> for each VOC is taken from the literature (Derwent et al., 2010). The
 SOAP was estimated by multiplying the SOAPF<sub>i</sub> value by the concentration of
 individual VOC species. The SOAP calculations through each VOC are as follows:

237

 $SOAP = \sum E_i \times SOAPF_i \tag{6}$ 

239  $E_i$  is the concentration of species *i*.

240

# 241 **3. Results and discussion**

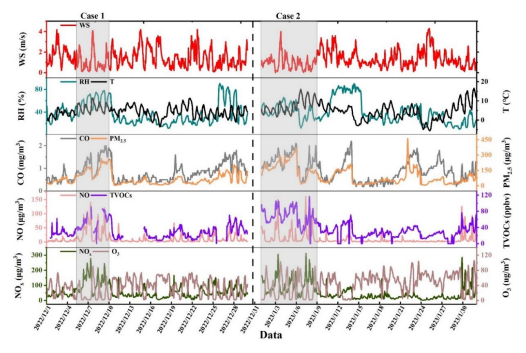
# 242 **3.1 Overview of variation in pollutants and meteorological**

## 243 parameters

244 Figure 1 shows the time series of meteorological parameters, TVOCs, O<sub>3</sub>, NO<sub>x</sub>, SO<sub>2</sub>, CO and PM<sub>2.5</sub> during the observed periods. Low WS and Temperature were found 245 246 with an average value of  $1.3 \pm 0.6$  m/s and  $5.0 \pm 2.5$  °C, respectively, during the entire period, comparable with observations at the same site in 2021 (Lai et al., 2024). A total 247 248 of 62 days of valid data was acquired with the daily average concentration of PM<sub>2.5</sub> ranging from 53 to 239  $\mu$ g/m<sup>3</sup>, with the average value of 111 ± 45  $\mu$ g/m<sup>3</sup>. The 249 250 concentration of TVOCs ranged from 15.6 to 57.1 ppbv with an average of  $36.1 \pm 21.0$ ppbv, higher than the same period in last year (27.9  $\pm$  12.7 ppbv, Lai et al., 2024). 251 During the observation period, the average values of T, WS and RH were  $5.0 \pm 2.5$  °C, 252 253  $1.3 \pm 0.6$  m/s and  $38.9 \pm 16.7\%$ , respectively.

254 Previous studies have shown that meteorological factors such as low WS, high RH,

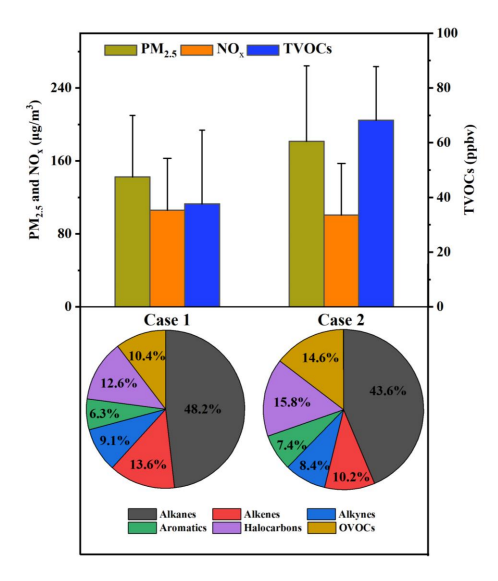
255 and low precipitation are responsible for the increase in PM<sub>2.5</sub> pollution in Zhengzhou in winter (Duan et al., 2019). Our analysis of the correlation between different 256 257 pollutants and meteorological conditions during the pollution period showed that 258  $PM_{2.5}$ , TVOCs and NO<sub>x</sub> were positively correlated with relative humidity (Fig. S3), 259 which is consistent with the results of some previous studies (Wang et al., 2019). Yu et al. (2018) identified RH and WS as the most influential meteorological conditions 260 261 of PM<sub>2.5</sub> during winter. Their findings revealed a positive correlation between hourly  $PM_{2.5}$  concentrations and RH (r = 0.84, p < 0.01) and a negative correlation between 262  $PM_{2.5}$  concentrations and WS (r = -0.62, p < 0.01). In this study, the WS on clean days 263  $(1.4 \pm 0.8 \text{ m/s})$  was higher than in Case 1  $(1.2 \pm 0.9 \text{ m/s})$  and Case 2  $(0.9 \pm 0.7 \text{ m/s})$ , 264 265 while the RH was lower by 26.2% and 12.5% compared to Case 1 and Case 2, respectively. These findings indicate that high RH and low WS significantly influence 266 the occurrence of pollution during the observation period. However, the mean PM<sub>2.5</sub> 267 concentration during the recovery period was found to be 1.6 times higher than that 268 269 observed during the infection period. Furthermore, the concentrations of other 270 pollutants (including SO<sub>2</sub>, NO<sub>2</sub>, CO, and O<sub>3</sub>) exhibited analogous trends during the 271 infection and recovery periods. The concentration of TVOCs during the recovery 272 period was 1.2 times higher than that during the infection period, exhibiting a 273 significant upward trend following the resumption of production. It is notable that WS, 274 which is only 0.3 m/s higher in Case 1 than in Case 2, and RH, which is 13% higher in Case 1 than in Case 2, were relatively stable, while the concentration of pollutants 275 276 is significantly higher in Case 2 than in Case 1. This is presumably attributable to the 277 resumption of production activities in Case 2, which resulted in a notable increase in 278 emissions. Decreased trends of air pollutants were found in other studies before and 279 after the outbreak of the novel coronavirus (COVID-19) in early 2020 (Qi et al., 2021; 280 Wang et al., 2021).



281

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
 period.

The shadow section in Fig. 1 represents two haze pollution events during the 284 monitoring period. A pollution event is determined when the daily average 285 concentration of PM<sub>2.5</sub> exceeds 75  $\mu$ g/m<sup>3</sup> (China's II-level standard) for at least three 286 consecutive days. Case 1 (December 5 to December 10 with daily average  $PM_{2.5}$  = 287 288 142.5  $\mu$ g/m<sup>3</sup>) and Case 2 (January 1 to January 8 with daily average PM<sub>2.5</sub> = 181.5  $\mu g/m^3$ ) were selected as they represent the pollution events in infection and recovery 289 290 periods, respectively, due to their long duration and high pollution levels. Any day with a PM<sub>2.5</sub> concentration lower than 35 µg/m<sup>3</sup> (China's I-level standard) is considered as 291 292 Clean day.



293

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
 Case 2.

296 As for the two representative pollution processes (Case 1 during the infection 297 period and Case 2 during the recovery period), the concentration of TVOCs in Case 1 298 and Case 2 were  $48.4 \pm 20.4$  and  $67.6 \pm 19.6$  ppbv (Fig. 2), respectively, increased by 299 63% and 188% compared with values during clean days. The average concentrations 300 of PM<sub>2.5</sub> and TVOCs during Case 2 were 1.3 and 1.8 times the values in Case 1. The 301 highest volume contributions of alkanes were found both in Case 1 (48%) and Case 2 302 (44%), consistent with the results in the Yangtze River Delta region (36-43%, Liu et al., 2023). While alkenes exhibited higher volume percentages of 13% in Case 1, followed 303 304 by halogenated hydrocarbon (12%) and OVOCs (10%). Higher volume percentages of 305 alkanes and alkenes in Case 1 were similar to the results in the gasoline evaporation 306 site in winter (Niu et al., 2022). Equivalent volume contribution of halogenated 307 hydrocarbon and OVOCs (15%) were found in Case 2, followed by alkenes (10%).

308 Although aromatic hydrocarbons have the lowest volumetric contribution (6% in Case

309 1 and 7% in Case 2), they show the largest increase from clean days to pollution.

310 Table 1 The average concentrations of meteorological parameters and pollutants during different

311 processes.

	Entire process	Infection period	Recovery period	Case 1	Case 2	Clean
C (	(2022.12.1-	(2022.12.1-	(2023.1.1-	(2022.12.5-	(2023.1.1-	Days
Category	2023.1.31)	2022.12.31)	2023.1.31)	2022.12.10)	2023.1.8)	
	N = 62  days	N = 31  days	N = 31  days	N = 6 days	N = 8 days	N = 8 days
WS (m/s)	$1.3\pm0.6$	$1.4\pm0.6$	$1.3\pm0.6$	$1.2\pm0.9$	$0.9\pm0.7$	$1.4\pm0.8$
T (°C)	$5.0 \pm 2.5$	$4.7\pm1.7$	$5.4 \pm 3.1$	$6.1\pm2.2$	$7.4\pm3.5$	$4.1\pm3.0$
RH (%)	$38.9 \pm 16.7$	$37.6\pm15.5$	$40.2\pm18.2$	$55.7\pm14.7$	$42.0\pm12.1$	$29.5\pm18.1$
TVOCs (ppbv)	$36.1\pm21.0$	$31.9\pm18.1$	$39.8\pm22.4$	$37.6\pm27.0$	$68.2\pm19.6$	$22.7\pm11.1$
$SO_2(\mu g/m^3)$	$11.4 \pm 2.7$	$10.2\pm2.8$	$12.7\pm2.3$	$11.0\pm3.7$	$16.2\pm6.1$	$6.5\pm2.5$
$NO_2(\mu g/m^3)$	$47.2\pm10.0$	$46.8\pm8.6$	$47.8 \pm 11.7$	$62.7\pm20.5$	$65.0\pm21.3$	$20.8\pm15.9$
CO (mg/m <sup>3</sup> )	$0.9\pm0.2$	$0.8\pm0.2$	$1.1\pm0.2$	$1.2\pm0.5$	$1.3\pm0.4$	$0.5\pm0.2$
$O_3(\mu g/m^3)$	$34.9\pm6.0$	$31.1\pm4.5$	$39.0\pm4.6$	$21.8\pm23.7$	$32.5\pm29.6$	$52.6\pm18.4$
$PM_{2.5} (\mu g/m^3)$	$111.5\pm45.1$	$86.6\pm34.6$	$138.3\pm39.6$	$142.5\pm67.4$	$181.5\pm82.7$	$23.8\pm16.8$

312

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

Category	Entire process	Infection period	Recovery period	Case 1	Case 2	Clean days
TVOCs	$36.1\pm21.0$	$31.9 \pm 18.1$	$39.8\pm22.4$	$48.4\pm20.4$	$67.6 \pm 19.6$	$17.5\pm9.5$
alkanes	$16.8\pm9.2$	$15.0\pm8.4$	$18.4\pm9.5$	$23.1\pm10.0$	$29.5\pm8.4$	$9.2\pm5.6$
alkenes	$4.1\pm2.7$	$3.8 \pm 2.6$	$4.4\pm2.7$	$6.5\pm2.9$	$7.0\pm2.6$	$1.7 \pm 1.3$
alkynes	$3.1\pm2.0$	$2.7 \pm 1.7$	$3.4\pm2.1$	$4.3\pm2.0$	$5.8 \pm 1.9$	$1.3\pm0.8$
aromatics	$2.1\pm2.0$	$1.8\pm1.5$	$2.3\pm2.2$	$3.0\pm1.8$	$4.9\pm2.8$	$0.7\pm0.5$
halogenated hydrocarbon	$5.4\pm3.3$	$4.4\pm2.3$	$6.2\pm3.8$	$6.0\pm1.9$	$10.7\pm3.6$	$2.7\pm1.4$
OVOCs	$4.6\pm3.2$	$3.5\pm2.7$	$5.1\pm3.5$	$5.0 \pm 2.4$	$9.7\pm2.8$	$1.9 \pm 1.1$

# 313 3.2 Source Analysis of VOCs

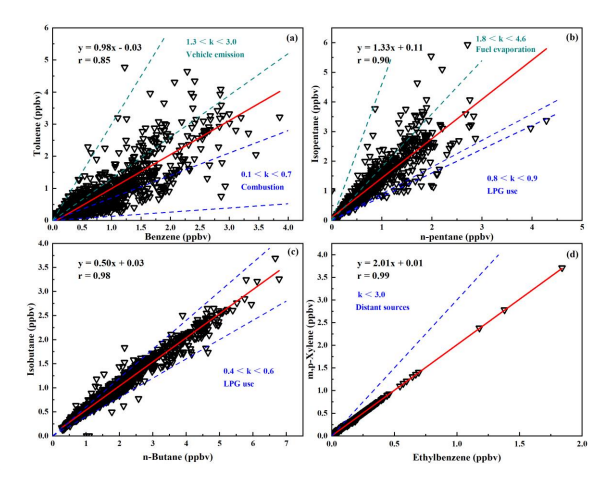
314	Specific VOC ratios can be used for initial source identification of VOCs and
315	determination of photochemical ages of air masses (Monod et al., 2001; An et al., 2014;
316	Li et al., 2019). In this study, the ratios of toluene/benzene (T/B), isopentane/n-pentane,
317	isobutane/n-butane, and m/p-xylene/ethylbenzene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

320 The toluene-to-benzene ratio (T/B ratio) was widely used to assess the relative 321 importance of different sources. Specifically, T/B ratio with a value of 1.3-3.0 was 322 observed in vehicle emissions for vehicles with different fuel types (Schauer et al., 2002; 323 Wang et al., 2015). The reported T/B ratio for combustion processes was between 0.13 324 and 0.7 (Li et al., 2011; Wang et al., 2014). The mean value of T/B ratio for the entire 325 period was 1.0, with the majority of the data (99%) falling between 0.1 and 3.0 and 326 concentrated within the 0.7-1.3 range (49%). This suggests that both traffic emissions 327 and combustion may be significant sources of VOCs.

328 The isopentane/n-pentane concentration ratios of 0.6-0.8 represent mainly coal 329 combustion emissions, ratios of 0.8-0.9 represent LPG emissions, 2.2-3.8 represent 330 vehicle exhaust emissions, and 1.8-4.6 represent fuel evaporation (Conner et al., 1995; 331 Liu et al., 2008; Li et al., 2019). The sources of isopentane and n-pentane in this study 332 were intricate and multifaceted. The mean isopentane/n-pentane ratio was 1.4, with the majority of data points (99%) falling within the range of 0.1-4.6, with a notable 333 334 concentration in the 0.8 to 1.8 interval. This indicates that pentane is susceptible to a 335 combination of LPG emissions and fuel evaporation. It should be noted that this 336 analytical approach is not without limitations. For instance, the proportionality of 337 pentane may be influenced by a combination of LPG emissions and fuel evaporation. 338 Consequently, an in-depth examination of the sources of VOCs was conducted using 339 the PMF model in the next section.

Isobutane/n-butane concentration ratios of 0.2-0.3 represent vehicle emissions, 0.4-0.6 represent LPG usage, and 0.6-1.0 represent natural gas emissions (Russo et al., 2010; Zheng et al., 2018). The mean isobutane/n-butane ratio in this study was 0.5, with the majority of data points (99%) falling within the 0.4-0.6 range, indicating that VOCs at the observation sites were significantly influenced by the use of LPG. (Shao et al., 2016; Zeng et al., 2023).

346 The ratio of X/E can be used to infer the photochemical age of the air mass. X/E ratios around 2.5-2.9 are typical of urban areas, indicating that VOCs are mainly from 347 348 the urban area (fresh air mass) (Kumar et al., 2018). When this ratio is significantly 349 lower than 3.0, it indicates that VOCs are mainly transported from distant sources 350 (aging air masses) (Kumar et al., 2018). The average X/E value in this study was 2.0 351 (Fig. 3(d)), indicating low photochemical activity and aging of the air mass at the 352 observation site. Potential source analyses also indicate that air masses are affected by 353 long-range transport (Fig. S4).



354

355

## Fig. 3. Correlation analysis between specific VOC species.

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).

Alkanes of C4-C6 substances were predominant in factor 1, including 2methylpentane, 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.

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.
The CPF plot shows that due east is the main emission direction at wind speeds less

than 2 m/s and southeast is the main source at wind speeds greater than 2 m/s (Fig. 5b).

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

380 Factor 4 is characterized by relatively high levels of C2-C6 low-carbon alkanes 381 (ethane, propane, isopentane, n-pentane, isobutane and n-butane), olefins (ethylene and 382 propylene), and benzene and toluene, which are important automotive exhaust tracers 383 (Song et al., 2021; Zhang et al., 2021b). Ethylene and propylene are important components derived from vehicle-related activities. Previous studies of VOCs in 384 385 Zhengzhou have shown a high percentage of VOCs emitted from gasoline vehicles, 386 with the main source of alkanes being on-road mobile sources (Bai et al., 2020). The 387 daily variation of this source in Fig. S5 shows a bimodal trend, with peaks occurring in 388 the morning and evening peaks of traffic, consistent with motor vehicle emissions. Fig. 389 5d shows that this source is mainly from the west where wind speeds are below 2 m/s, 390 and in this direction, there are a number of urban arterial roads with high traffic volumes. 391 Therefore, factor 4 was defined as vehicular emission source.

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

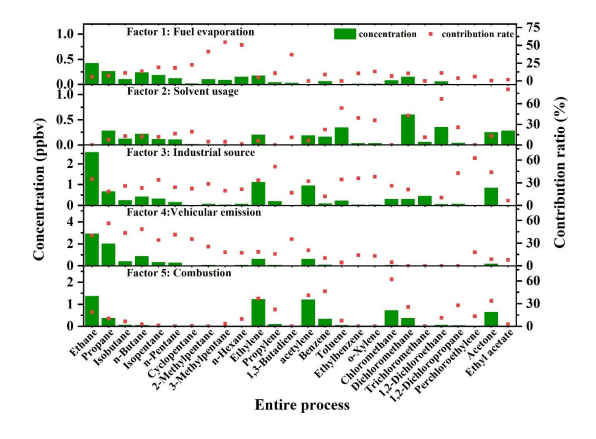




Fig. 4. Concentration of VOC species in each factor and contribution to each source.

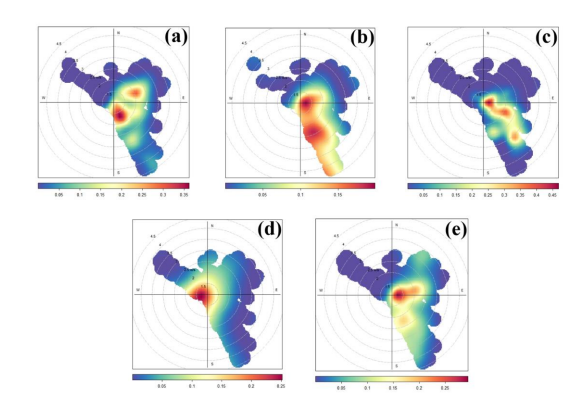




Fig. 5. CPF plots of five VOCs sources obtained using the PMF model.

402 Note: a: Fuel evaporation; b: Solvent usage; c: Industrial source; d: Vehicular emission; e: 403 Combustion.

404 Fig. S6 compares the differences in PMF source profiles between the Omicron 405 infection period and the recovery period, as well as between the pollution day and the 406 clean day. We present the concentrations of the five main VOCs in all five factors in 407 Table S2. Ethane (vehicular emission), 2-methylpentane (fuel evaporation), benzene 408 (industry source), chloromethane (combustion), and ethyl acetate (solvent usage) were 409 selected as tracers for five sources. Ethane concentration in Case 2 (5.9 ppbv) is much 410 higher than in other processes, and ethane concentration during the recovery period (3.4 411 ppbv) is also higher than during the infection period (2.4 ppbv), which may to some 412 extent reflect increased vehicular emissions during the recovery period.

413 Concentrations of most species were significantly higher during the recovery 414 period than during the infection period. The representative pollution processes in both 415 periods showed the same results as well, with a 79% higher concentration of TVOCs in 416 Case 2 (65.1 ppbv) compared to Case 1 (36.3 ppbv) (Fig. 6). While in Case 1 industry was the dominant source of VOCs, by Case 2 motorized sources reached a 417 418 concentration value of 21.2 ppbv, accounting for 33% of the observed VOCs, and 419 became the dominant source of emissions. This is consistent with the fact that people's 420 mobility activities have increased after the epidemic has entered the recovery period. 421 As a group of VOCs species with the highest concentration share, ethane and propane 422 contributed more to the clean days motor vehicle source than other processes, which 423 also resulted in a 34% clean days motor vehicle source share.

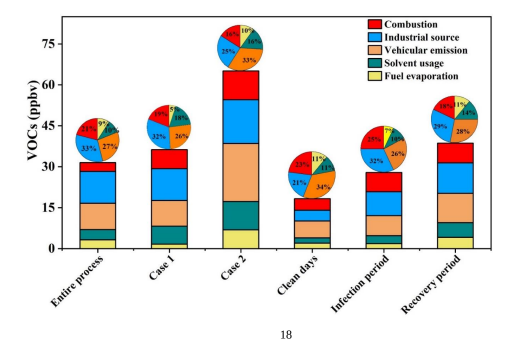


Fig. 6. Contribution of each to VOCs for different processes.

### 426 **3.3 SOAP**

425

427 VOCs are estimated to contribute about 16-30% or more of PM<sub>2.5</sub> by mass through 428 SOA production (Huang et al., 2014). Therefore, by calculating the SOAP value, the 429 influence of different sources on PM<sub>2.5</sub> production can be reflected to a certain extent.

430 We have included quantitative analysis for SOAP as well. Fig. 7 shows the SOAP 431 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 432 433 close to 100% of the total SOAP contribution, with Case 1 reaching 98%. In each 434 process, the composition of the top ten substances is essentially the same. Aromatic 435 hydrocarbons contributed the most, with BTEX always occupying the top five positions 436 and toluene the most. The SOAP values of the top ten contributing species for the two 437 polluting processes are shown in Tables S3 and S4. Toluene, the highest contributing species, reached a SOAP value of 49.4  $\mu$ g/m<sup>3</sup> in the most polluted Case 2, which was 438 3.2 times higher than the SOAP sum of all species on the clean day (15.5  $\mu$ g/m<sup>3</sup>). The 439 SOAP value for Case 1, which is also a contaminated process, was 67  $\mu$ g/m<sup>3</sup>, and the 440 main species (m/xylene: 9.8  $\mu$ g/m<sup>3</sup>, benzene: 8.5  $\mu$ g/m<sup>3</sup>) including toluene (34.6  $\mu$ g/m<sup>3</sup>) 441 442 were lower than those for Case 2 (m/xylene: 19.4  $\mu$ g/m<sup>3</sup>, benzene: 13.4  $\mu$ g/m<sup>3</sup>).

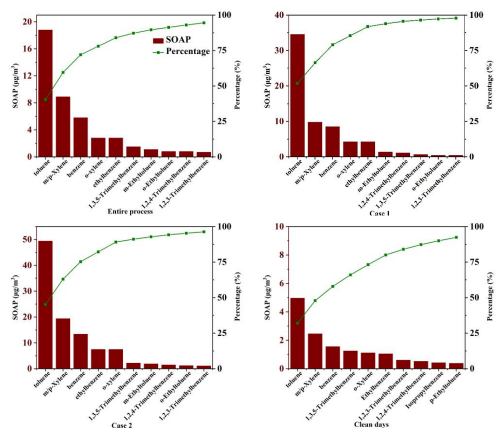


Fig. 7. SOAP dominant species in different processes

445 Figure 8 shows the SOAP calculated after source resolution of the two pollution 446 processes by PMF for clean days, respectively. In Case 1, industrial source is the 447 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 448 449 evaporation sources emerge as the primary contributors to SOAP, with their respective 450 contribution levels rising to 32% and 26%. Case 1 was during the infection period, 451 when social activities had not yet returned to normal. In Case 2, when society had 452 basically returned to normal, the increase in emissions from various sources resulted in 453 a more balanced distribution of SOAP contributions and caused more severe PM2.5 454 pollution. In addition, a few days before Case 2, the Zhengzhou Municipal People's 455 Government initiated the Heavy Pollution Weather Level II response 456 (https://sthij.zhengzhou.gov.cn/tzgg/7037130.jhtml) and introduced control measures 457 for emissions from industrial and mobile sources, which resulted in a significant 458 reduction of SOAP levels from industrial and motorized sources in Case 2. The clean day result with a SOAP of 8.8  $\mu$ g/m<sup>3</sup> also indicates that industrial and solvent usage 459 460 sources are the most dominant SOAP sources. The primary sources of aromatic 461 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 462 et al., 2017). Consequently, it is imperative to implement measures to reduce PM<sub>2.5</sub> 463 464 pollution by regulating emissions from industrial and solvent usage sources.

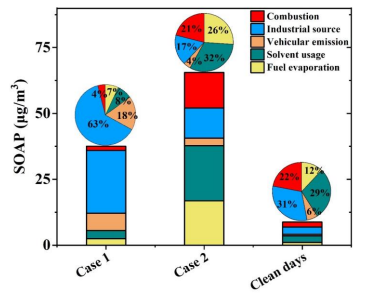


Fig. 8. SOAP value and contribution ratio of each process

466 467

465

## 468 **4.** Conclusions

Continuous observation of VOCs during the infection of the Omicron epidemic 469 470 was carried out at an urban site in polluted Zhengzhou from December 1, 2022, to January 31, 2023. The daily average concentration of PM<sub>2.5</sub> ranged from 53.5 to 239.4 471  $\mu g/m^3$  with an average value of 111.5 ± 45.1  $\mu g/m^3$  during the whole period. The 472 473 concentration of TVOCs ranged from 15.6 to 57.1 ppbv with an average of  $36.1 \pm 21.0$ 474 ppbv, higher than the same period in last year  $(27.9 \pm 12.7 \text{ ppbv}, \text{Lai et al.}, 2024)$ . Two 475 representative contamination processes were identified (Case 1 during the infection 476 period and Case 2 during the recovery period). The concentration of TVOCs in Case 1 477 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. The average concentrations of PM<sub>2.5</sub> 478 479 and TVOCs during Case 2 were 1.3 and 1.8 times of the values in Case 1. This is 480 consistent with the observed increase in pollutant emissions following the return to 481 normal social life from the period of Omicron infection. The highest volume 482 contributions of alkanes were found both in Case 1 (48%) and Case 2 (44%). Though the volume contribution of aromatics were the lowest (6% in Case 1 and 7% in Case 2), 483 484 the highest increase ratio was found from clean days to polluted episodes. Low wind 485 speed and high humidity were the main meteorological reasons for the occurrence of 486 pollution. Analyzing the sources of VOCs revealed that VOCs were found to be affected 487 by a combination of local emissions and regional transport. The primary sources of 488 atmospheric VOCs in Zhengzhou were identified as industrial emissions (32%), vehicle 489 emissions (27%), and combustion (21%). Significant discrepancies were observed in 490 the sources of VOCs between the two pollution processes. In Case 1, industrial emissions constituted the primary source of VOCs, accounting for 32% of the total 491 492 VOC concentration. In contrast, in Case 2, the proportion of vehicle emissions 493 increased to 33%, representing the primary source of VOCs.

A further analysis of the effect of VOCs on SOA generation reveals that aromatic compounds are the primary contributors to SOAP, with BTEX being the predominant contributor throughout the period. The SOAP values reached 37.6 and 65.6  $\mu$ g/m<sup>3</sup> in Case 1 and Case 2, respectively. In Case 1, the greatest contribution to SOAP was made by industrial sources (63%, 23.8  $\mu$ g/m<sup>3</sup>), while vehicular sources, which constituted the second most important source, accounted for only 18%. In Case 2, the contribution of each VOC source was more evenly distributed, with solvent use sources and fuel

- 501 evaporation sources representing the primary contributors to SOAP, accounting for 32%
- 502 (20.9  $\mu$ g/m<sup>3</sup>) and 26% (16.8  $\mu$ g/m<sup>3</sup>), respectively. The SOAP result for the clean day

503 was 8.8  $\mu$ g/m<sup>3</sup>, with industrial sources and solvent use still being the primary

504 contributors. Therefore, the industrial and solvent use sectors are the predominant

505 sources of pollutants during this observation. The aforementioned results substantiate

- 506 the considerable impact of elevated emissions from all sources on the exacerbation of
- 507 pollution following the conclusion of the Omicron infection.

### 508 Author contribution:

- 509 Bowen Zhang: Data curation, Methodology, Formal analysis, Writing Original Draft.
- 510 Dong Zhang: Data curation, Formal analysis, Review & Editing.
- 511 Zhe Dong: Data curation, Formal analysis, Review & Editing.
- 512 Xinshuai Song: Data curation, Formal analysis.
- 513 Ruiqin Zhang: Supervision, Writing-Review & Editing, Funding acquisition.
- 514 Xiao Li: Formal analysis, Investigation, Supervision, Writing-Review & Editing.

### 515 **Competing interests:**

516 The contact author has declared that none of the authors has any competing interests.

### 517 Acknowledgments:

518This research was supported by the Natural Science Foundation of Henan Province519(232300421395) and the National Key Research and Development Program of China

520 (2017YFC0212400).

### 521 **References**

522An, J., Zhu, B., Wang, H., Li, Y., Lin, X., and Yang, H.: Characteristics and source523apportionment of VOCs measured in an industrial area of Nanjing, Yangtze River Delta,524China,Atmospheric525https://doi.org/10.1016/j.atmosenv.2014.08.021, 2014.

Bai, L., Lu, X., Yin, S., Zhang, H., Ma, S., Wang, C., Li, Y., and Zhang, R.: A
recent emission inventory of multiple air pollutant, PM<sub>2.5</sub> chemical species and its
spatial-temporal characteristics in central China, Journal of Cleaner Production, 269,
122114, https://doi.org/10.1016/j.jclepro.2020.122114, 2020.

530 Buzcu, B. and Fraser, M. P.: Source identification and apportionment of volatile

organic compounds in Houston, TX, Atmospheric Environment, 40, 2385-2400,
https://doi.org/10.1016/j.atmosenv.2005.12.020, 2006.

533Conner, T. L., Lonneman, W. A., Seila, R.L.: Transportation-related volatile534hydrocarbon source profiles measured in atlanta, Journal of the Air & Waste535Management536https://doi.org/10.1080/10473289.1995.10467370, 1995.

Cui, L., Wu, D., Wang, S., Xu, Q., Hu, R., and Hao, J.: Measurement report:
Ambient volatile organic compound (VOC) pollution in urban Beijing: characteristics,
sources, and implications for pollution control, Atmospheric Chemistry and Physics,
22, 11931-11944, https://doi.org/10.5194/acp-22-11931-2022, 2022.

541 Derwent, R. G., Jenkin, M. E., Utembe, S. R., Shallcross, D. E., Murrells, T. P., 542 and Passant, N. R.: Secondary organic aerosol formation from a large number of 543 reactive man-made organic compounds, Science of the Total Environment, 408, 3374-544 3381, https://doi.org/10.1016/j.scitotenv.2010.04.013, 2010.

545 Duan, S., Jiang, N., Yang, L., Zhang, R.: Transport Pathways and Potential Sources
546 of PM<sub>2.5</sub> During the Winter in Zhengzhou, Environmental Science, Jan 8;40(1):86-93,
547 https://doi.org/10.13227/j.hjkx.201805187, 2019.

548 Gao, J., Zhang, J., Li, H., Li, L., Xu, L., Zhang, Y., Wang, Z., Wang, X., Zhang, W., Chen, Y., Cheng, X., Zhang, H., Peng, L., Chai, F., and Wei, Y.: Comparative study 549 550 of volatile organic compounds in ambient air using observed mixing ratios and initial 551 mixing ratios taking chemical loss into account – A case study in a typical urban area 552 in Beijing, Science of the Total Environment, 628-629, 791-804. 553 https://doi.org/10.1016/j.scitotenv.2018.01.175, 2018.

554 Guan, Y., Liu, X., Zheng, Z., Dai, Y., Du, G., Han, J., Hou, L. a., and Duan, E.: 555 Summer O<sub>3</sub> pollution cycle characteristics and VOCs sources in a central city of 556 Beijing-Tianjin-Hebei area, China, Environmental Pollution, 323, 121293, 557 https://doi.org/10.1016/j.envpol.2023.121293, 2023.

Huang, R., Zhang, Y., Bozzetti, C. et al.: High secondary aerosol contribution to p
articulate pollution during haze events in China, Nature, 514 (7521), 218–22, https://d
oi.org/10.1038/nature13774, 2014.

Hui, L., Liu, X., Tan, Q., Feng, M., An, J., Qu, Y., Zhang, Y., Deng, Y., Zhai, R., a
nd Wang, Z.: VOC characteristics, chemical reactivity and sources in urban Wuhan, ce
ntral China, Atmospheric Environment, 224, 117340, https://doi.org/10.1016/j.atmose
nv.2020.117340, 2020.

565 Jensen, A., Liu, Z., Tan, W., Dix, B., Chen, T., Koss, A., Zhu, L., Li, L., de Gouw, J.: Measurements of volatile organic compounds during the COVID-19 lockdown in 566 567 Changzhou, China, Geophysical research letters, 48(20), https://doi.org/10.1029/2021 568 GL095560, 2021.

569 Jiang, N., Hao, X., Hao, Q., Wei, Y., Zhang, Y., Lyu, Z., Zhang, R.: Changes in se condary inorganic ions in PM2.5 at different pollution stages before and after COVID-1 570 571 9 control, Environmental Science, 44(5), 2430-2440, https://doi.org/10.13227/j.hjkx.2 572 02206170, 2023.

573 Kumar, A., Singh, D., Kumar, K., Singh, B. B., and Jain, V. K.: Distribution of 574 VOCs in urban and rural atmospheres of subtropical India: Temporal variation, source 575 attribution, ratios, OFP and risk assessment, Science of the Total Environment, 613-614, 576 492-501, https://doi.org/10.1016/j.scitotenv.2017.09.096, 2018.

577 Lai, M., Zhang, D., Yin, S., Song, X., and Zhang, R.: Pollution characteristics, source apportionment and activity analysis of atmospheric VOCs during winter and 578 summer pollution in Zhengzhou City, Environmental Science, 4108, 3500-3510, 579 580 https://doi.org/10.13227/j.hjkx.202001133, 2024.

581 Li, B., Ho, S. S. H., Gong, S., Ni, J., Li, H., Han, L., Yang, Y., Qi, Y., and Zhao, 582 D.: Characterization of VOCs and their related atmospheric processes in a central 583 Chinese city during severe ozone pollution periods, Atmospheric Chemistry and 584 Physics, 19, 617-638, https://doi.org/10.5194/acp-19-617-2019, 2019.

Li, J., Deng, S., Tohti, A., Li, G., Yi, X., Lu, Z., Liu, J., and Zhang, S.: Spatial 585 586 characteristics of VOCs and their ozone and secondary organic aerosol formation 587 potentials in autumn and winter in the Guanzhong Plain, China, Environmental 588 Research, 211, 113036, https://doi.org/10.1016/j.envres.2022.113036, 2022.

589 Li, J., Xie, S. D., Zeng, L. M., Li, L. Y., Li, Y. Q., and Wu, R. R.: Characterization 590 of ambient volatile organic compounds and their sources in Beijing, before, during, and 591 after Asia-Pacific Economic Cooperation China 2014, Atmospheric Chemistry and 592 Physics, 15, 7945-7959, https://doi.org/10.5194/acp-15-7945-2015, 2015.

593 Li, J., Lu, K., Lv, W., Li, J., Zhong, L., Ou, Y., Chen, D., Huang, X., and Zhang, 594 Y.: Fast increasing of surface ozone concentrations in Pearl River Delta characterized by a regional air quality monitoring network during 2006-2011, Journal of 595 596 Environmental Sciences, 26, 23-36, https://doi.org/10.1016/S1001-0742(13)60377-0, 597 2014.

598

Liu, Y., Li, X., Tang, G., Wang, L., Lv, B., Guo, X., and Wang, Y.: Secondary

organic aerosols in Jinan, an urban site in North China: Significant anthropogenic
contributions to heavy pollution, Journal of Environmental Sciences, 80, 107-115,
https://doi.org/10.1016/j.jes.2018.11.009, 2019.

Liu, Y., Shao, M., Fu, L., Lu, S., Zeng, L., and Tang, D.: Source profiles of volatile
organic compounds (VOCs) measured in China: Part I, Atmospheric Environment, 42,
6247-6260, https://doi.org/10.1016/j.atmosenv.2008.01.070, 2008.

605 Liu, Y., Song, M., Liu, X., Zhang, Y., Hui, L., Kong, L., Zhang, Y., Zhang, C., Qu, Y., An, J., Ma, D., Tan, Q., and Feng, M.: Characterization and sources of volatile 606 607 organic compounds (VOCs) and their related changes during ozone pollution days in 257, 608 2016 Environmental Pollution, 113599, in Beijing, China, 609 https://doi.org/10.1016/j.envpol.2019.113599, 2020.

Liu, Z., Hu, K., Zhang, K., Zhu, S., Wang, M., and Li, L.: VOCs sources and roles
in O<sub>3</sub> formation in the central Yangtze River Delta region of China, Atmospheric
Environment, 302, https://doi.org/10.1016/j.atmosenv.2023.119755, 2023.

Li, X., Wang, S., Hao, J.: Characteristics of volatile organic compounds (VOCs)
emitted from biofuel combustion in China, Environmental Science, 32, 3515-3521,
2011.

Li, Y., Yin, S., Zhang R., Yu, S., Yang, J., and Zhang, D.: Characteristics and source
apportionment of atmospheric VOCs at different pollution levels in winter in an urban
area in Zhengzhou, Environmental Science, 4108, 3500-3510,
https://doi.org/10.13227/j.hjkx.202001133, 2020.

Ma, Q., Wang, W., Wu, Y., Wang, F., Jin, L., Song, Y., Han, Y., Zhang, R., Zhang,
D.: Haze caused by NO<sub>x</sub> oxidation under restricted residential and industrial activities
in a mega city in the south of North China Plain, Chemosphere, Volume 305, 135489,
https://doi.org/10.1016/j.chemosphere.2022.135489, 2022.

Merino, M., Marinescu, M., Cascajo, A., Carretero, J., Singh, D.: Evaluating the
spread of Omicron COVID-19 variant in Spain, Future Generation Computer Systems,
149, 547-561, https://doi.org/10.1016/j.future.2023.07.025, 2023.

Monod, A., Sive, B. C., Avino, P., Chen, T., Blake, D. R., and Sherwood Rowland,
F.: Monoaromatic compounds in ambient air of various cities: a focus on correlations
between the xylenes and ethylbenzene, Atmospheric Environment, 35, 135-149,
https://doi.org/10.1016/S1352-2310(00)00274-0, 2001.

631 Mozaffar, A., Zhang, Y.-L., Fan, M., Cao, F., and Lin, Y.-C.: Characteristics of 632 summertime ambient VOCs and their contributions to O<sub>3</sub> and SOA formation in a suburban area of Nanjing, China, Atmospheric Research, 240, 104923,
https://doi.org/10.1016/j.atmosres.2020.104923, 2020.

Mu, L., Feng, C., Li, Y., Li, X., Liu, T., Jiang, X., Liu, Z., Bai, H., and Liu, X.:
Emission factors and source profiles of VOCs emitted from coke production in Shanxi,
China, Environmental Pollution, 335, 122373,
https://doi.org/10.1016/j.envpol.2023.122373, 2023.

639 Niu, Y., Yan, Y., Chai, J., Zhang, X., Xu, Y., Duan, X., Wu, J., and Peng, L.: Effects of regional transport from different potential pollution areas on volatile organic 640 641 compounds (VOCs) in Northern Beijing during non-heating and heating periods, 642 of Total 836, Science the Environment, 155465, 643 https://doi.org/10.1016/j.scitotenv.2022.155465, 2022.

Norris, G., Duvall, R., Brown, S., Bai. S. EPA Positive Matrix Factorization (PMF)
5.0 Fundamentals and User Guide. U.S. Environmental Protection Agency, Washington,
DC, EPA/600/R-14/108 (NTIS PB2015-105147), 2014.

Paatero, P., Eberly, S., Brown, S. G., Norris, G. A.: Methods for estimating
uncertainty in factor analytic solutions, Atmospheric Measurement Techniques, Volume
7, 781-797, https://doi.org/10.5194/amt-7-781-2014, 2014.

650 Pei, C., Yang, W., Zhang, Y., Song, W., Xiao, S., Wang, J., Zhang, J., Zhang, T., Chen, D., Wang, Y., Chen, Y., Wang, X.: Decrease in ambient volatile organic 651 652 compounds during the COVID-19 lockdown period in the Pearl River Delta region, 653 China, of The Total 823, 153720, south Science Environment, 654 https://doi.org/10.1016/j.scitotenv.2022.153720, 2022.

Petersen, M. S., Í Kongsstovu, S., Eliasen, E. H., Larsen, S., Hansen, J. L., Vest,
N., Dahl, M. M., Christiansen, D. H., Møller, L. F., & Kristiansen, M. F.: Clinical
characteristics of the Omicron variant - results from a Nationwide Symptoms Survey
in the Faroe Islands, International Journal of Infectious Diseases, 122, 636–643,
https://doi.org/10.1016/j.ijid.2022.07.005, 2022.

- 660Qi, J., Mo, Z., Yuan, B., Huang, S., Huangfu, Y., Wang, Z., Li, X., Yang, S., Wang,661W., Zhao, Y., Wang, X., Wang, W., Liu, K., and Shao, M.: An observation approach in662evaluation of ozone production to precursor changes during the COVID-19 lockdown,663AtmosphericEnvironment,262,118618,
- 664 https://doi.org/10.1016/j.atmosenv.2021.118618, 2021.

Russo, R. S., Zhou, Y., White, M. L., Mao, H., Talbot, R., and Sive, B. C.: Multiyear (2004–2008) record of nonmethane hydrocarbons and halocarbons in New England: seasonal variations and regional sources, Atmospheric Chemistry and Physics,
10, 4909-4929, https://doi.org/10.5194/acp-10-4909-2010, 2010.

Sahu, L. K., Tripathi, N., Gupta, M., Singh, V., Yadav, R., Patel, K.: Impact of
COVID-19 Pandemic lockdown in ambient concentrations of aromatic volatile organic
compounds in a metropolitan city of western India, Journal of geophysical research,
Atmospheres : JGR, 127(6), https://doi.org/10.1029/2022JD036628, 2022.

673 Schauer, J., Kleeman, M., Cass, G., Simoneit, B.: Measurement of emissions from 674 air pollution sources.5.  $C_1$ - $C_{32}$  organic compounds from gasoline-powered motor 675 vehicles, Environmental Science & Technology, 36, 1169-1180, 676 https://doi.org/10.1021/es0108077, 2002.

Shao, P., An, J., Xin, J., Wu, F., Wang, J., Ji, D., and Wang, Y.: Source
apportionment of VOCs and the contribution to photochemical ozone formation during
summer in the typical industrial area in the Yangtze River Delta, China, Atmospheric
Research, 176-177, 64-74, https://doi.org/10.1016/j.atmosres.2016.02.015, 2016.

Shi, Y., Liu, C., Zhang, B., Simayi, M., Xi, Z., Ren, J., and Xie, S.: Accurate
identification of key VOCs sources contributing to O<sub>3</sub> formation along the Liaodong
Bay based on emission inventories and ambient observations, Science of the Total
Environment, 844, 156998, 10.1016/j.scitotenv.2022.156998, 2022.

685 Singh, B., Sohrab, S., Athar, M., Alandijany, T., Kumari, S., Nair, A., Kumari, S., 686 Mehra, K., Chowdhary, K., Rahman, S., Azhar, E.: Substantial changes in selected volatile organic compounds (VOCs) and associations with health risk assessments in 687 688 industrial areas during the COVID-19 Pandemic, Toxics, 11, 165, 689 https://doi.org/10.3390/toxics11020165, 2023a.

Singh, B., Singh, M., Ulman, Y., Sharma, U., Pradhan, R., Sahoo, J., Padhi, S.,
Chandra, P., Koul, M., Tripathi, P., Kumar, D., Masih, J.: Distribution and temporal
variation of total volatile organic compounds concentrations associated with health risk
in Punjab, India, Case Studies in Chemical and Environmental Engineering, 8, 100417,
https://doi.org/10.1016/j.cscee.2023.100417, 2023b.

Song, M., Li, X., Yang, S., Yu, X., Zhou, S., Yang, Y., Chen, S., Dong, H., Liao,
K., Chen, Q., Lu, K., Zhang, N., Cao, J., Zeng, L., and Zhang, Y.: Spatiotemporal
variation, sources, and secondary transformation potential of volatile organic
compounds in Xi'an, China, Atmospheric Chemistry and Physics, 21, 4939-4958,
https://doi.org/10.5194/acp-21-4939-2021, 2021.

700 Song, X., Zhang, D., Li, X., Lu, X., Wang, M., Zhang, B., Zhang, R.: Simultaneous

observations of peroxyacetyl nitrate and ozone in Central China during static
management of COVID-19: Regional transport and thermal decomposition,
Atmospheric Research, Volume 294, 106958,
https://doi.org/10.1016/j.atmosres.2023.106958, 2023.

Song, Y., Shao, M., Liu, Y., Lu, S., Kuster, W., Goldan, P., and Xie, S.: Source
apportionment of ambient volatile organic compounds in Beijing, Environmental
Science & Technology, 41, 4348-4353, https://doi.org/10.1021/es0625982, 2007.

Wang, H., Li, J., Peng, Y., Zhang, M., Che, H., Zhang, X.: The impacts of the
meteorology features on PM<sub>2.5</sub> levels during a severe haze episode in central-east China,
Atmospheric Environment, Volume 197, Pages 177-189, ISSN 1352-2310,
https://doi.org/10.1016/j.atmosenv.2018.10.001, 2019.

Wang, H., Wang, Q., Chen, J. Chen, C., Huang, C., Qiao, L. Lou, S., Lu, J.: Do
vehicular emissions dominate the source of C6–C8 aromatics in the megacity Shanghai
of eastern China?, Environmental Science, 27, 290-297, https://doi.org/10.
1016/j.jes.2014.05.033, 2015.

Wang, M., Lu, S., Shao, M., Zeng, L., Zheng, J., Xie, F., Lin, H., Hu, K., and Lu,
X.: Impact of COVID-19 lockdown on ambient levels and sources of volatile organic
compounds (VOCs) in Nanjing, China, Science of the Total Environment, 757, 143823,
https://doi.org/10.1016/j.scitotenv.2020.143823, 2021.

Wang, M., Zeng, L., Lu, S., Shao, M., Liu, X., Yu, X., Chen, W., Yuan, B., Zhang,
Q., Hu, M., & Zhang, Z.: Development and validation of a cryogen-free automatic gas
chromatograph system (GC-MS/FID) for online measurements of volatile organic
compounds, Analytical Methods, 6, 9424, https://doi.org/10.1039/C4AY01855A, 2014.

Wang, T., Xue, L., Brimblecombe, P., Lam, Y. F., Li, L., and Zhang, L.: Ozone
pollution in China: A review of concentrations, meteorological influences, chemical
precursors, and effects, Science of the Total Environment, 575, 1582-1596,
https://doi.org/10.1016/j.scitotenv.2016.10.081, 2017.

- Wu, R., Li, J., Hao, Y., Li, Y., Zeng, L., and Xie, S.: Evolution process and sources
  of ambient volatile organic compounds during a severe haze event in Beijing, China,
  Science of the Total Environment, 560-561, 62-72,
  https://doi.org/10.1016/j.scitotenv.2016.04.030, 2016.
- 732Wu, W., Zhao, B., Wang, S., and Hao, J.: Ozone and secondary organic aerosol733formation potential from anthropogenic volatile organic compounds emissions in China,734JournalofEnvironmentalSciences,53,224-237,

735 https://doi.org/10.1016/j.jes.2016.03.025, 2017.

- Xiong, Y., Zhou, J., Xing, Z., and Du, K.: Optimization of a volatile organic
  compound control strategy in an oil industry center in Canada by evaluating ozone and
  secondary organic aerosol formation potential, Environmental Research, 191, 110217,
  https://doi.org/10.1016/j.envres.2020.110217, 2020.
- Yu, F., Wang, Q., Yan, Q., Jiang, N., Wei, J., Wei, Z., Yin, S.: Particle size
  distribution, chemical composition and meteorological factor analysis: A case study
  during wintertime snow cover in Zhengzhou, China, Atmospheric Research, 202, 140147, 0169-8095, https://doi.org/10.1016/j.atmosres.2017.11.016, 2018.
- Yun, L., Li, C., Zhang, M., He, L. and Guo, J.: Pollution characteristics and sources
  of atmospheric VOCs in the coastal background area of the Pearl River Delta,
  Environmental Science, 4191-4201, https://doi.org/10.13227/j.hjkx.202101155, 2021.
- Zeng, X., Han, M., Ren, G., Liu, G., Wang, X., Du, K., Zhang, X., and Lin, H.: A
  comprehensive investigation on source apportionment and multi-directional regional
  transport of volatile organic compounds and ozone in urban Zhengzhou, Chemosphere,
  334, 139001, https://doi.org/10.1016/j.chemosphere.2023.139001, 2023.
- Zhang, C., Liu, X., Zhang, Y., Tan, Q., Feng, M., Qu, Y., An, J., Deng, Y., Zhai, R.,
  Wang, Z., Cheng, N., and Zha, S.: Characteristics, source apportionment and chemical
  conversions of VOCs based on a comprehensive summer observation experiment in
  Beijing, Atmospheric Pollution Research, 12, 230-241,
  https://doi.org/10.1016/j.apr.2020.12.010, 2021a.
- Zhang, D., He, B., Yuan, M., Yu, S., Yin, S., and Zhang, R.: Characteristics,
  sources and health risks assessment of VOCs in Zhengzhou, China during haze
  pollution season, Journal of Environmental Sciences, 108, 44-57,
  https://doi.org/10.1016/j.jes.2021.01.035, 2021b.
- Zhang, D., Li, X., Yuan, M., Xu, Y., Xu, Q., Su, F., Wang, S., Zhang, R.:
  Characteristics and sources of nonmethane volatile organic compounds (NMVOCs) and
  O<sub>3</sub>-NO<sub>x</sub>-NMVOC relationships in Zhengzhou, China, Atmosphere Chemistry and
  Physics, 24, 8549-8567, https://doi.org/10.5194/acp-24-8549-2024, 2024.
- 764 Zhang, F., Shang, X., Chen, H., Xie, G., Fu, Y., Wu, D., Sun, W., Liu, P., Zhang, C., Mu, Y., Zeng, L., Wan, M., Wang, Y., Xiao, H., Wang, G., and Chen, J.: Significant 765 impact of coal combustion on VOCs emissions in winter in a North China rural site, 766 767 Science of the Total Environment, 720. 137617, 768 https://doi.org/10.1016/j.scitotenv.2020.137617, 2020.

- Zhang, J., Sun, Y., Wu, F., Sun, J., and Wang, Y.: The characteristics, seasonal
  variation and source apportionment of VOCs at Gongga Mountain, China, Atmospheric
  Environment, 88, 297-305, https://doi.org/10.1016/j.atmosenv.2013.03.036, 2014.
- Zhang, Z., Yan, X., Gao, F., Thai, P., Wang, H., Chen, D., Zhou, L., Gong, D., Li,
  Q., Morawska, L., and Wang, B.: Emission and health risk assessment of volatile
  organic compounds in various processes of a petroleum refinery in the Pearl River Delta,
  China, Environmental Pollution, 238, 452-461,
  https://doi.org/10.1016/j.envpol.2018.03.054, 2018.
- Zheng, H., Kong, S., Xing, X., Mao, Y., Hu, T., Ding, Y., Li, G., Liu, D., Li, S.,
  and Qi, S.: Monitoring of volatile organic compounds (VOCs) from an oil and gas
  station in northwest China for 1 year, Atmospheric Chemistry and Physics, 18, 45674595, https://doi.org/10.5194/acp-18-4567-2018, 2018.
- Zheng, J., Zhong, L., Wang, T., Louie, P. K. K., and Li, Z.: Ground-level ozone in
  the Pearl River Delta region: Analysis of data from a recently established regional air
  quality monitoring network, Atmospheric Environment, 44, 814-823,
  https://doi.org/10.1016/j.atmosenv.2009.11.032, 2010.
- Zhou, Z., Xiao, L., Fei, L., Yu, W., Lin M., Huang, T., Zhang, Z. and Tao J.:
  Characteristics and sources of VOCs during ozone pollution and non-pollution periods
  in summer in Dongguan industrial concentration area, Environmental Science, 44974505, https://doi.org/10.13227/j.hjkx.202111285, 2022.
- Zou, Y., Yan, X. L., Flores, R. M., Zhang, L. Y., Yang, S. P., Fan, L. Y., Deng, T.,
  Deng, X. J., and Ye, D. Q.: Source apportionment and ozone formation mechanism of
  VOCs considering photochemical loss in Guangzhou, China, Science of the Total
  Environment, 903, 166191, https://doi.org/10.1016/j.scitotenv.2023.166191, 2023.
- Zuo, H., Jiang, Y., Yuan, J., Wang, Z., Zhang, P., Guo, C., Wang, Z., Chen, Y., Wen,
  Q., Wei, Y., Li, X.: Pollution characteristics and source differences of VOCs before and
  after COVID-19 in Beijing, Science of The Total Environment, 907, 167694,
  https://doi.org/10.1016/j.scitotenv.2023.167694, 2024.