

1 **Review of source analyses of ambient VOCs considering reactive losses:**  
2 **Methods of reducing loss effects, impacts of losses, and sources**

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4 Baoshuang Liu<sup>1,2\*</sup>, Yao Gu<sup>1,2</sup>, Yutong Wu<sup>1,2</sup>, Qili Dai<sup>1,2</sup>, Shaojie Song<sup>1,2</sup>, Yinchang Feng<sup>1,2\*</sup>, and  
5 Philip K. Hopke<sup>3,4</sup>

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7 <sup>1</sup>State Environmental Protection Key Laboratory of Urban Ambient Air Particulate Matter Pollution  
8 Prevention and Control & Tianjin Key Laboratory of Urban Transport Emission Research, College  
9 of Environmental Science and Engineering, Nankai University, Tianjin 300350, China

10 <sup>2</sup>CMA-NKU Cooperative Laboratory for Atmospheric Environment-Health Research, Tianjin  
11 300350, China

12 <sup>3</sup>Department of Public Health Sciences, University of Rochester School of Medicine and Dentistry,  
13 Rochester, NY 14642, USA

14 <sup>4</sup>Institute for a Sustainable Environment, Clarkson University, Potsdam, NY 13699, USA

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16 ***Correspondence:*** Baoshuang Liu (lbsnankai@foxmail.com) and Yinchang Feng  
17 (fengyc@nankai.edu.cn)

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21 **Abstract**

22 Chemical losses of ambient reactive volatile organic compounds (VOCs) is a long-term issue yet to  
23 be resolved in VOC source apportionments. These losses substantially reduce the concentrations of  
24 highly reactive species in the apportioned factor profiles and result in the underestimation of source  
25 contributions. This review assesses the common methods and existing issues in ways to reduce losses  
26 and loss impacts in source analyses and suggest research directions for improved VOC source  
27 apportionments. Positive Matrix Factorization (PMF) is now the main VOC source analysis method  
28 compared to other mathematical models. The issue in using any apportionment tool is the processing  
29 of the data to be analyzed to reduce the impacts of reactive losses. Estimating the initial  
30 concentrations of ambient VOCs based on photochemical age has become the primary approach to  
31 reduce reactive loss effects in PMF except for selecting low reactivity species or nighttime data into  
32 the analysis. Currently, the initial concentration method only considers daytime reactions with  
33 hydroxyl ( $\bullet\text{OH}$ ) radicals. However, the  $\bullet\text{OH}$  rate constants vary with temperature and that has not  
34 been considered. Losses from reactions with  $\text{O}_3$  and  $\text{NO}_3$  radicals especially for alkene species  
35 remain to be included. Thus, the accuracy of the photochemical-age estimation is uncertain. Beyond  
36 developing accurate quantitative approaches for reactive losses, source analyses methods for the  
37 consumed VOCs and the accurate quantification of different source contributions to  $\text{O}_3$  and  
38 secondary organic aerosols are important additional directions for future research.

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42 **Keywords:** Reactive loss; Consumed VOCs; Source analyses; PMF; Initial concentrations

## 43 **1 Introduction**

44 Ambient VOCs are key precursors of ozone (O<sub>3</sub>) and secondary organic aerosols (SOAs)  
45 formation (Li et al., 2016; Li et al., 2018a; Wu and Xie, 2018). Accurate apportionment of their  
46 sources can be important in developing effective prevention and control measures for atmospheric O<sub>3</sub>  
47 and secondary organic particulate matter pollution (Carrillo-Torres et al., 2017; Meng et al., 1997;  
48 Wang et al., 2022a). Current research on source analyses of atmospheric VOCs has been primarily  
49 conducted utilizing the species ratio method (Che et al., 2019; Zhang et al., 2021), the photochemical  
50 age-based parameterization method (Huang et al., 2020; Zhu et al., 2021), and receptor models (e.g.,  
51 positive matrix factorization (PMF) (Gu et al., 2020; Liu et al., 2016), chemical mass balance (CMB)  
52 (Song et al., 2019), and principal component analysis/multiple linear regression (PCA/MLR) (Jia et  
53 al., 2016; Sanchez et al., 2008), etc.) based on the measured concentration data. Receptor models,  
54 especially PMF, are the most widely used source apportionment methods (Song et al., 2008; Vega et  
55 al., 2022; Yang et al., 2022b). However, compared to particulate matter, ambient VOC species can  
56 undergo rapid, complex chemical reactions with •OH radicals, NO<sub>3</sub> radicals, O<sub>3</sub>, etc. (Atkinson and  
57 Arey, 2003), resulting in substantial chemical losses during their transport from their sources to the  
58 receptor site (Yang et al., 2022b; Yuan et al., 2012a). Therefore, the source apportioned results based  
59 on the measured VOC data have difficulty to fully reflect the actual impacts of emission sources on  
60 air quality (Wu et al., 2023b; Yang et al., 2022a).

61 To reduce the impact of reactive losses, many studies have selected low-reactivity VOC species  
62 to conduct source analyses when using PMF source analyses (Guan et al., 2020; Yang et al., 2022a).  
63 However, this method cannot fully solve the issues related to reactive losses and provide complete  
64 source apportionments since some highly active marker species such as isoprene (Tan et al., 2020)  
65 cannot be excluded from the PMF input species without a substantial loss of information (Liu et al.,  
66 2023a). Therefore, recent studies estimated the initial concentrations of ambient VOCs (i.e., the VOC  
67 concentrations in the fresh emissions before they can undergo chemical reactions) utilizing the  
68 photochemical age-based parameterization method and then performed source analyses with PMF  
69 (He et al., 2019; Zou et al., 2023). However, there could still be high uncertainties in the estimated  
70 photochemical age of VOC species (Parrish et al., 2007; Yuan et al., 2012b).

71 There were studies that only used nighttime data for source analyses to reduce the loss impacts

72 (Kim et al., 2005), but the representativeness of the apportioned results was likely limited since there  
73 could be daytime only sources that would not be observed at night (Buzcu and Fraser, 2006). Some  
74 studies applied decay factors to correct the impact of reactive losses in using CMB for VOC source  
75 analyses (Friedlander, 1981; Lin and Milford, 1994; Na and Pyo Kim, 2007). However, there were  
76 relatively few studies and the effectiveness of this method still needs to be assessed. In 2023, Liu et  
77 al. (2023a) systematically investigated the impact of VOCs photochemical losses on the PMF source  
78 apportioned results, and found that photochemical losses reduced the concentrations of highly  
79 reactive species in factor profiles resulting in the contributions of biogenic emissions and polymer  
80 production-related industrial sources being substantially underestimated. However, there has been  
81 little related research to assess the limitations of VOC apportionments.

82 With the substantial increase of O<sub>3</sub> concentrations in many locations worldwide in recent years  
83 (Li et al., 2020a; Zhang et al., 2018; Zhao et al., 2021), accurate source apportionment approaches of  
84 the key precursor VOCs have been acquired increasing attention (Gu et al., 2022). Thus, a related  
85 issue is the determination of the VOCs consumed in the formation of the observed O<sub>3</sub> and SOAs (Gu  
86 et al., 2023). Although some studies have investigated the underestimated contributions of emission  
87 sources by comparing the apportioned results based on the initial and measured data (He et al., 2019;  
88 Wu et al., 2023a), there are few publications that conducted source analyses for the consumed VOCs  
89 and apportioned the contributions of different sources to the formation of O<sub>3</sub> and SOA. In 2023 and  
90 2024, Gu et al. (2023) and Cui et al. (2024) attempted to develop a method for apportioning the  
91 primary and oxidative sources of the consumed VOCs, and Wang et al. (2022b) conducted a similar  
92 study. Currently, although some studies have been conducted on these issues of VOC reactive losses  
93 and achieved some important results (Gu et al., 2023; Watson et al., 2001; Wu et al., 2023b), there  
94 remain unresolved issues. To better promote progress on the development of better VOCs source  
95 apportioned methods, the present work systematically investigated the main methods and  
96 shortcomings of those methods that are currently applied to resolve these issues by reviewing  
97 relevant papers, with the aim of identifying directions for the future developments and improvement  
98 of VOCs source apportionment methods.

99

## 100 **2 Materials and methods**

101 Relevant papers were collected by exhaustively searching Science Direct (Elsevier), the Web of  
102 Science, Scopus, Springer, Wiley, and China National Knowledge Infrastructure (CNKI), etc. with  
103 the keywords: volatile organic compounds (VOCs), oxygenated VOCs (OVOCs), initial  
104 concentrations/mixing ratios, chemical/photochemical losses, source apportionment/analysis,  
105 positive matrix factorization (PMF), chemical mass balance (CMB), receptor model, and  
106 photochemical age-based parameterization method. The information extracted from each publication  
107 included methods for reducing the impacts of photochemical losses, the impacts of VOC  
108 photochemical losses on source analyses, the source apportioned methods and the results for the  
109 consumed VOCs in the atmosphere. This study identified 169 papers, of which 68 papers were  
110 published since 2020, accounting for ~40% of the total publications; a total of 36 papers were  
111 published from 2015 to 2019, accounting for ~21%. There were 102 research papers reporting results  
112 in China, accounting for ~60% of the total papers. There were 23 papers located in the United States,  
113 accounting for ~14%. The numbers of papers located in India, South Korea, Canada, and Japan were  
114 only 4, 3, 3, and 3, respectively, and there were a few papers from other countries. Additionally, there  
115 were 19 research papers conducted in Beijing, 9 papers in Guangzhou, and 8 papers in Tianjin,  
116 accounting for ~11%, ~5%, and ~5% of the total papers, respectively. There were fewer reports from  
117 other cities.

118

## 119 **3 Results and discussion**

### 120 **3.1 Methods of reducing the impacts of reactive losses**

121 Ambient VOCs can be substantially oxidized by O<sub>3</sub>, hydroxyl (•OH), and nitrate (NO<sub>3</sub>) radicals  
122 (Atkinson and Arey, 2003; Bey et al., 2001; Finlayson-Pitts and Pitts, 1997), especially oxidation by  
123 •OH radicals primarily during the daytime (Wang et al., 2013). Therefore, reducing the impacts of  
124 VOC reactive losses on source apportionment has long been an important but not easy issue to  
125 resolve in VOC source apportionments (Liu et al., 2023a). Ambient VOCs primarily include alkanes,  
126 alkenes, aromatic hydrocarbons, alkynes, OVOCs, and halogenated hydrocarbons, etc. Studies  
127 suggested that the concentration of Photochemical Assessment Monitoring Stations (PAMS) VOCs  
128 (i.e., alkanes, alkenes, aromatic hydrocarbons, and alkynes) accounted for ~63% of the total VOCs

129 (TVOCs) concentration (i.e., average value of proportions from different literature), while OVOCs  
130 and halogenated hydrocarbons contributed ~22% and ~14% to the TVOCs, respectively (as shown in  
131 Table S1). The reported number of measured species ranged from 13 to 124, including 0-32 alkanes,  
132 2-16 alkenes, 1 alkyne (only acetylene), 3-19 aromatic hydrocarbons, 4-28 OVOCs, and 28-38  
133 halogenated hydrocarbons. There were substantial differences in the identified emission sources for  
134 the different types of VOCs (Mo et al., 2016). For example, PAMS VOCs mainly originate from  
135 primary anthropogenic sources (Chen et al., 2010), while OVOCs can also be formed by oxidation of  
136 PAMS VOCs in addition to primary source emissions (Chen et al., 2014; Seinfeld and Pandis, 1986).  
137 The contributions of secondary formation to some OVOC species (e.g., acetaldehyde and  
138 propionaldehyde) can exceed 50% (de Gouw et al., 2005). In addition to local emissions, the ambient  
139 concentrations of halogenated hydrocarbons can also be affected by the long-distance transport  
140 (Mintz and McWhinney, 2008). Therefore, utilizing only the same source analyses approach for  
141 multiple VOC species from different sources might produce results with high uncertainties in the  
142 apportionments.

143 Source analyses methods for ambient VOCs considering reactive losses mainly included PMF,  
144 CMB, and photochemical age-based parameterization method (Table S2). PMF was the most  
145 commonly method used for source analyses (Yang et al., 2022b). Approximately 52% of the  
146 reviewed publications focused on source analyses of PAMS species using PMF, while OVOCs  
147 primarily used photochemical age-based parameterization method for source apportionment but with  
148 only relatively few studies (Tables S2-S3). Additionally, the studies on data from simultaneous  
149 measurement of both PAMS and OVOC species utilized two methods to conduct source analyses:  
150 one method was to simultaneously input PAMS and some OVOC species into the receptor model  
151 (e.g., PMF) for source analyses and only separated primary and secondary source contributions of  
152 ambient VOCs (Han et al., 2023; Li et al., 2023; Liu et al., 2023b; Tan et al., 2021) (Table S3).  
153 Another approach was to use the PMF and photochemical age-based parameterization to obtain the  
154 source apportioned results for ambient PAMS and OVOCs, respectively, and then combined the two  
155 apportioned results to obtain finally source resolution results for the ambient VOCs (Zhu et al., 2021).  
156 However, the utilization of this method has been limited.

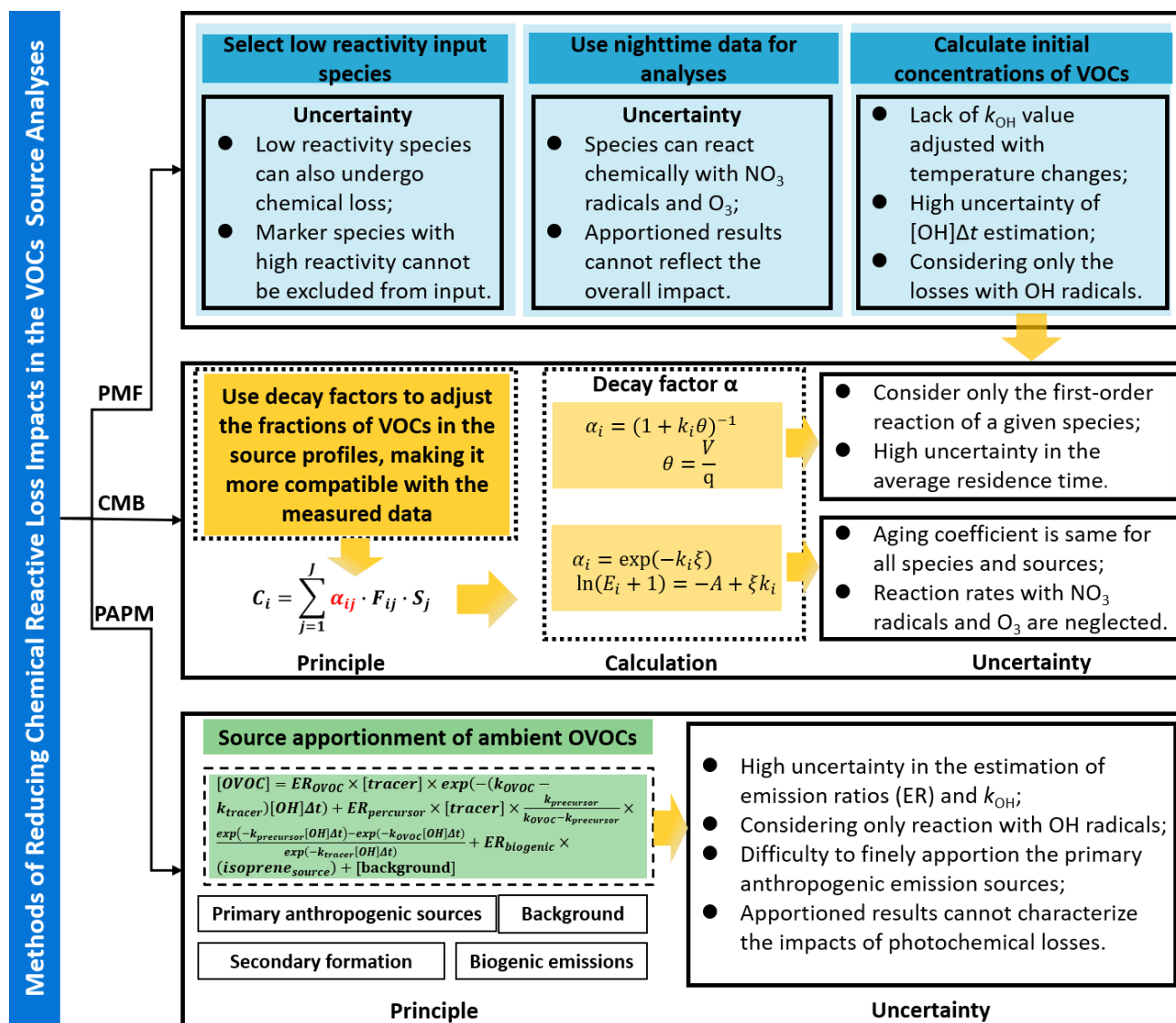
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### 158 **3.1.1 Methods for reducing reactive loss in PMF source analyses**

#### 159 **3.1.1.1 Selecting low activity species or incorporating night only data into PMF**

160 The methods of reducing effects of reactive losses for source apportionments utilizing different  
161 models are shown in Fig. 1. At present, selecting the VOC species with lower reactivity to be input in  
162 PMF for apportionment is the most commonly used approach (Chen et al., 2019; Tan et al., 2020;  
163 Yang et al., 2022b). Many highly active alkene and aromatic hydrocarbon species were not included  
164 in the PMF calculations (Gu et al., 2023; Liu et al., 2023a). For example, Liu et al. (2023a) excluded  
165 highly active species such as 1-hexene, trans-2-butene, trans-2-pentene, cis-2-pentene, and cis-2-  
166 butene. In addition, current approach of incorporating daytime and nighttime VOC data into PMF for  
167 source analyses (e.g., Gu et al., 2020; Li et al., 2020b; Jain et al., 2022) assumes that the daytime and  
168 nighttime factor profiles are consistent. However, the daytime factor profiles can be substantially  
169 influenced by photochemistry (Liu et al., 2025). Therefore, the source contributions obtained by this  
170 method had relatively higher uncertainty. To reduce the impacts of reactive losses on the PMF  
171 apportioned factor profiles and the corresponding contributions, some studies utilized only nighttime  
172 data when reactive losses would be lower for source analyses to obtain more accurate nighttime  
173 contributions of emission sources (Buzcu-Guven and Fraser, 2008; Buzcu and Fraser, 2006; Kim et  
174 al., 2005; Xie and Berkowitz, 2006).

175 However, the volatile emission sources commonly contribute substantially to VOCs during the  
176 daytime (Gu et al., 2023). For example, Buzcu and Fraser (2006) used nighttime and all-day data to  
177 conduct VOC source apportionment in Houston, respectively. They found that the night only data did  
178 not identify biogenic or evaporative gasoline sources due to minimal emissions at night. Thus, the  
179 contributions of the nighttime emission sources substantially increased compared to the all-day  
180 results because of the missing source types. Meanwhile, the all-day VOC source analysis conducted  
181 by Zhao et al. (2004) in Houston during the same period also showed substantial differences from the  
182 results reported by Buzcu and Fraser (2006) utilizing night-only data. An additional consideration is  
183 that VOCs can react with NO<sub>3</sub> radicals and O<sub>3</sub> (especially for highly reactive alkenes) at night  
184 causing reactive losses (de Gouw et al., 2017). Thus, this approach does not eliminate the effects of  
185 all possible loss mechanisms. Although Wang et al. (2024a) also considered reactions with O<sub>3</sub> at  
186 night when estimating VOC chemical reactive losses, the number of related studies remains limited.



188

189 **Figure 1.** Methods for reducing the impacts of reactive losses for different source analyses models.  
 190 PMF represents positive matrix factorization, CMB represents chemical mass balance, and PAPM  
 191 represents photochemical age-based parameterization method.

192

### 193 3.1.1.2 Incorporating VOC initial concentration estimated by photochemical age-based 194 parameterization method into PMF

195 The photochemical age-based parameterization method is an approach to estimate the initial  
 196 concentrations of ambient VOCs (Shao et al., 2011; Wang et al., 2013; Yuan et al., 2012b). This  
 197 method assumes that the chemical loss of ambient VOCs mainly originates only from reactions of  
 198 the VOCs with  $\bullet\text{OH}$  radicals (Parrish et al., 1992; Sun et al., 2016; Wang et al., 2013). The  
 199 photochemical ages ( $\Delta t$ ) of VOC species are estimated, and then combined with the rate constants



200 for reaction for •OH radicals with the VOC species ( $k_{\bullet\text{OH}}$ ) to calculate their initial concentrations and  
201 quantified the VOC photochemical losses (Shao et al., 2011; Wang et al., 2013; Zhan et al., 2021).  
202 The equation for the initial concentration calculation is shown in Eq. (1), where  $k_{\bullet\text{OH}}$  and  $\Delta t$  are the  
203 key parameters in this approach.

$$204 \quad [\text{VOC}]_t = [\text{VOC}]_0 \times \exp(-k_{\bullet\text{OH}} \times [\bullet\text{OH}]\Delta t) \quad (1)$$

205 where  $[\text{VOC}]_0$  and  $[\text{VOC}]_t$  are the initial and measured concentrations of VOC at time  $t$ ,  $[\bullet\text{OH}]$   
206 represents the concentration of •OH radical. However, many studies estimated  $[\bullet\text{OH}] \Delta t$  because of  
207 the lack of •OH radical concentration data and defined the product as the •OH exposure (Shao et al.,  
208 2011; Wei et al., 2022; Yuan et al., 2012b).

209 The  $k_{\bullet\text{OH}}$  of VOC species are substantially affected by atmospheric temperature (Atkinson and  
210 Arey, 2003), which could be commonly measured in laboratory (or chamber) experiments or through  
211 the use of detailed chemical computer models (Atkinson and Arey, 2003). According to the reviewed  
212 papers regarding the estimation of photochemical age, it was found that approximately 47% of the  
213 publications used the Atkinson and Arey's 2003 summary of  $k_{\bullet\text{OH}}$  values at 298° K (Atkinson and  
214 Arey, 2003), and approximately 8% used Carter's 2010 summary of  $k_{\bullet\text{OH}}$  values at 300° K (Carter,  
215 2010). The detailed  $k_{\bullet\text{OH}}$  values for different VOC species summarized in these two publications are  
216 provided in Table S4. However, other relevant studies cited  $k_{\bullet\text{OH}}$  values from other papers by  
217 Atkinson and/or Carter (Atkinson, 1991, 2007; Atkinson et al., 2006; Talukdar et al., 1994). The rate  
218 constant for a bimolecular reaction between a hydroxyl radical and a VOC molecule has a roughly  
219 exponential dependence on temperature in the Arrhenius equation (Liu et al., 2023a). Therefore,  
220 these published papers only used  $k_{\bullet\text{OH}}$  values at specific temperatures, which made it difficult to  
221 characterize the actual reaction rates at the actual temperatures in the atmosphere. To address this  
222 issue, Wang et al. (2022b) and Wang et al. (2023) utilized the Arrhenius expression to locally  
223 optimize the  $k_{\bullet\text{OH}}$  values of VOC species based on the actual temperature conditions to further  
224 improve the accuracy of estimating the VOC initial data. In addition, some studies used the  $k_{\bullet\text{OH}}$  of  
225 *m,p*-xylene instead of values of more reactive species to reduce high-outliers when estimating the  
226 initial concentration (Wang et al., 2013). However, the validity of this method was difficult to assess.

227 There are two main methods (as shown in Table S5) for estimating  $\Delta t$ , namely, the species ratio  
228 method (Roberts et al., 1984; Shao et al., 2011; Wan et al., 2022) and the sequential reaction model

229 (Gong et al., 2018; Shao et al., 2011; Xie et al., 2008). The specific details are as follows:

230 **(1) Estimation of  $\Delta t$  based on the species ratio method**

231 
$$\Delta t = \frac{1}{(k_A - k_B) \times [OH]} \times \left[ \ln \left( \frac{[A]_0}{[B]_0} \right) - \ln \left( \frac{[A]_t}{[B]_t} \right) \right] \quad (2)$$

232 where  $k_A$  and  $k_B$  are the reaction rate constants with  $\bullet OH$  radicals for species A and B of highly  
233 homologous species with substantial differences in reaction activity, respectively,  $\frac{[A]_0}{[B]_0}$  is the initial  
234 concentration ratio of A to B in fresh emissions,  $\frac{[A]_t}{[B]_t}$  is the measured concentration ratio of A to B at  
235 time  $t$ . The determination of initial concentration ratio of A to B was extremely critical for estimating  
236 the  $\Delta t$ . The  $\Delta t$  is used to then calculate the initial concentrations of the VOCs of interest using Eq. (1).  
237 The initial concentrations can then be the input variables to PMF (Liu et al., 2023a).

238 This ratio method was first proposed by Roberts et al. (1984). The method has been  
239 commonly applied to PAMS species emitted by anthropogenic sources. There have been differences  
240 in the species used to calculate the initial ratio in different papers (Table S6). Generally aromatic  
241 hydrocarbons (e.g., benzene, toluene, ethylbenzene, or m,p-xylene) were used as the selected species.  
242 The substantial differences in activity and high homology are generally the main basis for selecting  
243 species-ratio types (Lin et al., 2011; Shao et al., 2009; Zou et al., 2021; Zou et al., 2023). Most  
244 publications used the initial ratios of ethylbenzene/m,p-xylene (E/X) or m,p-xylene/ethylbenzene  
245 (X/E) to calculate  $\Delta t$  (Table S6). These two species ratios (E/X and X/E) accounted for ~33% of the  
246 total reviewed publications of estimating initial ratios, respectively. Approximately 7% of the papers  
247 used toluene/benzene (T/B), while m,p-xylene/benzene (X/B), ethylbenzene/o-xylene (E/O), or O/E  
248 had limited use, accounting for ~2% and ~12% of the total reviewed publications of initial ratio  
249 estimation.

250 The methods for obtaining the initial species ratio in the reviewed papers mainly referred to  
251 prior similar studies (Wang et al., 2016), source emission inventory values (Wang et al., 2013), and  
252 estimations based on the observed concentration data at the receptor sites at times when low  
253 reactivity was expected (Fang et al., 2021; Han et al., 2019; Wu et al., 2023b). There were several  
254 papers using the first two methods to obtain the initial species ratios (Table S6). The initial species  
255 ratios obtained by these methods had difficulty to accurately characterize the comprehensive impacts  
256 of multiple sources in the study area. Most studies were based on the observed data of low

257 photochemical reaction periods at night (~ 19:00 to 08:00 LT) to calculate the initial species ratios  
258 (Table S6). This method assumes that VOC species emitted by different sources during the certain  
259 periods of night would not undergo oxidative reaction, and the concentration ratios of two  
260 homologous VOC species remained unchanged during the transport process from source emissions  
261 to the receptor measured sites (Liu et al., 2023a; Sun et al., 2016; Yuan et al., 2012b). Since there  
262 could be residual reactions with nitrate radical and multiple sources of the indicator species, there  
263 remains uncertainty in the results. For example, McKeen and Liu (1993) and McKeen et al. (1996)  
264 found that the species ratios can be substantially influenced by the oxidative chemistry and  
265 atmospheric mixing. Comparing one species versus another, both the model results and the  
266 observations were consistent with an average rate of dilution roughly equivalent to n-butane  
267 oxidation (McKeen et al., 1996). This result has negative implications for the use of hydrocarbon  
268 ratios as chemical reaction clocks (McKeen et al., 1996). Parrish et al. (2007) also suggested that  
269 there were uncertainties in the determination of  $\Delta t$  by the initial species ratios because of influence of  
270 fresh emissions along the transport path that perturb the results. However, this approach can still  
271 provide a useful measure of chemical processing in the atmosphere (Parrish et al., 2007).

272 There were differences in the estimation details of the initial species ratio in different reports  
273 (Liu et al., 2023a; Shao et al., 2011; Yuan et al., 2012b). For example, Yuan et al. (2012b) utilized the  
274 m,p-xylene and benzene data measured between 0:00 and 5:00 LT to conduct a linear fit. The  
275 regression line was then extrapolated to the highest benzene concentration during this period so that  
276 the initial ratio could be estimated from the m,p-xylene to benzene concentration ratios at this point.  
277 Wu et al. (2023b) and Yang et al. (2022a) used time periods (00:00-04:00 LT and 20:00-05:00 LT,  
278 respectively) with the lowest and the highest E/X and T/B ratios at night to calculate the average  
279 values of corresponding E/X ratios less than various percentiles or the corresponding T/B values  
280 above various percentiles, and then determined the initial ratio by the minimum average ratio (i.e.,  
281 E/X) or the maximum average ratio (i.e., T/B) of the two species with the highest homology (i.e.,  
282 highest correlation) below different percentiles. Wu et al. (2023a) reported that the maximum value  
283 of X/E at 01:00-06:00 LT was considered as the initial species ratio. In addition, Liu et al. (2023a)  
284 utilized the slope of the linear fit of the scatterplots of corresponding ethylbenzene and m,p-xylene  
285 below the 10<sup>th</sup> percentile of E/X ratios during 00:00-05:00 LT to determine the initial ratio. Overall,

286 the E/X in the reviewed papers ranged between 0.22 and 0.75, T/B was between 3.14 and 4.48, X/E  
287 was between 0.39 and 4.42, E/O was between 1.30 and 1.32, O/E was between 1.19 and 3.14, and  
288 X/B was 2.2 (as shown in Table S6).

289 Zhang et al. (2020) attempted to confirm the feasibility and rationality of estimating  $\Delta t$  based on  
290 T/B utilizing the regressions of benzene and toluene versus CO and  $\Delta t$ . Zou et al. (2023)  
291 demonstrated the rationality of selecting the ratio of E and X to estimate the  $\Delta t$  through high  
292 correlation of these two species having substantial activity differences. However, there were few  
293 studies to systematically assess how to choose the most suitable species ratios and the calculated  
294 method.

295 Multiple types of initial species ratios were used for estimating  $\Delta t$  of VOCs (Table S6), and the  
296 results estimated by the different ratios could vary substantially. Shao et al. (2011) suggested that the  
297 differences between the initial concentrations calculated using the ratios of three pairs of compound  
298 ratios (i.e., T/B, X/E, and i-butene/propene (iB/P)) were generally within 50%, and the X/E and iB/P  
299 results were in good agreement. Zou et al. (2021) examined the sensitivity of estimating initial  
300 concentration based on E/X and E/O initial ratios. They found that the relative variation range of the  
301 initial concentrations of PAMS species was between 0.41% and 68.06% for an initial E/O of 1.3  
302 when compared with an initial E/X of 0.5.

303 Additionally, in the ratio method, the same  $\Delta t$  was calculated for different VOCs for each  
304 sample, and the paired species for estimating the ratio were aromatic hydrocarbons (Table S6), which  
305 are mainly emitted from anthropogenic emission sources such as solvent use and petrochemical  
306 enterprises (Mo et al., 2015; Na et al., 2004; Yuan et al., 2010; Zhang et al., 2016). Therefore, the  $\Delta t$   
307 obtained based on these ratios may only reflect the chemical aging of VOC species emitted from  
308 these specific sources. It is difficult to accurately characterize the  $\Delta t$  for these species from other  
309 sources. In the future, different types of species ratios (not just aromatic hydrocarbons) need to be  
310 selected based on VOC species from different sources. The  $\Delta t$  derived from different types of VOC  
311 species should be calculated to improve traditional ratio methods.

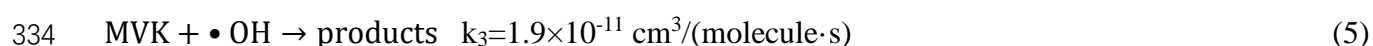
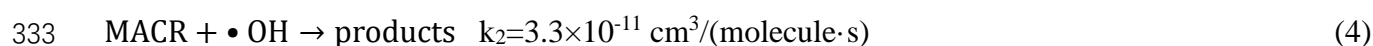
312 An additional issue is that this method only considers the reactions between VOCs and  $\bullet\text{OH}$   
313 radicals during the day, while chemical reactions with  $\text{NO}_3$  radicals at night and/or  $\text{O}_3$  were  
314 commonly excluded. Liu et al. (2023a) attempted to estimate the reactive losses between 19:00 and

315 23:00 LT, but they considered the gradual loss of •OH radicals after sunset and did not consider the  
 316 losses caused by NO<sub>3</sub> radical reactions that may be present at night since they are not photolyzed as  
 317 occurs in the daylight. Additionally, alkenes can directly react with O<sub>3</sub> in the atmosphere causing  
 318 losses. de Gouw et al. (2017) suggested that nighttime removal of highly reactive alkenes by O<sub>3</sub> and  
 319 NO<sub>3</sub> radicals was also substantial. However, there are currently no reports on estimation methods for  
 320 these reactive losses. In future studies, the ratio of two alkene species with substantial differences in  
 321 reaction rates with O<sub>3</sub> could be used as the base ratio as in Eq. (2), and combined with measured O<sub>3</sub>  
 322 reaction rate constants (Atkinson and Arey, 2003) to estimate the chemical reaction time ( $\Delta t$ ) of  
 323 alkene species with O<sub>3</sub>. Then, chemical losses of alkene species in the reaction with O<sub>3</sub> could be  
 324 estimated using Eqs. (1) and (9) and the combined effects of O<sub>3</sub> and •OH could be estimated. de  
 325 Gouw et al. (2017) proposed the calculated method of O<sub>3</sub> exposure (i.e., [O<sub>3</sub>] $\Delta t$ ) at nighttime, which  
 326 can be calculated from the measured ratio of benzene over cis-2-butene.

327

## 328 (2) Estimation of $\Delta t$ based on the sequential reaction model

329 It has been found that using the ratio method to calculate the  $\Delta t$  of isoprene could result in  
 330 overestimation (Wu et al., 2023a; Yang et al., 2022b). An alternative is the sequential reaction  
 331 method. It is based on measurements of the reaction products of these reactions.



335 
$$\frac{[\text{MACR}]_t}{[\text{Isoprene}]_t} = \frac{0.23k_1}{k_2-k_1} (1 - e^{(k_1-k_2)[\bullet\text{OH}]\Delta t})$$
 (6)

336 
$$\frac{[\text{MVK}]_t}{[\text{Isoprene}]_t} = \frac{0.32k_1}{k_3-k_1} (1 - e^{(k_1-k_3)[\bullet\text{OH}]\Delta t})$$
 (7)

337 where MVK is methyl vinyl ketone, MACR is methacrolein;  $k_1$ ,  $k_2$ , and  $k_3$  are the rate constants  
 338 reacting with •OH of isoprene, MACR, and MVK, respectively; [MVK]<sub>t</sub>, [MACR]<sub>t</sub>, and [isoprene]<sub>t</sub>  
 339 are the measured ambient concentrations at time  $t$ .

340 This sequential reaction method was first proposed by Bertman et al. (1995) and was commonly  
 341 used to estimate the  $\Delta t$  of isoprene emitted primarily from biogenic emissions (Gong et al., 2018;  
 342 Roberts et al., 2006; Wu et al., 2023a). Assuming the •OH-driven isoprene oxidation mechanism  
 343 (Eqs. (3)-(5)) from the laboratory chamber studies of Carter and Atkinson (1996), Stroud et al. (2001)

344 quantified the  $\Delta t$  of isoprene by the reaction relationship between isoprene and its oxidation products  
 345 (Eqs. (6)-(7)) (de Gouw et al., 2005; Wu et al., 2023a; Xie et al., 2008). The synchronous  
 346 measurement of MVK and MACR is critical to obtain  $\Delta t$  utilizing this method. Numerous studies  
 347 failed in estimating the isoprene  $\Delta t$  due to the lack of MVK and MACR (Gu et al., 2023; Liu et al.,  
 348 2023a; Wang et al., 2023). Although some studies have used the average of  $\Delta t$  obtained from both  
 349 MVK and MACR as the final  $\Delta t$  (Xie et al., 2008), there has been little assessment of which method  
 350 was more suitable.

351 Additionally, there were studies using the isotopic hydrocarbon clock method (Kornilova et al.,  
 352 2016; Rudolph and Czuba, 2000; Saito et al., 2009) to estimate the species  $\Delta t$ , that was calculated by  
 353 the decay of isotopes in the emissions to the ambient receptor site (Table S5). However, the studies  
 354 only calculated the  $\Delta t$  without estimating the species initial concentrations. Our study found that  
 355 approximately 74% of the publications that calculated the  $\Delta t$  used the species ratio method,  
 356 approximately 11% used both the ratio method and the sequential reaction model, while a few used  
 357 other methods (Table S5).

358

### 359 **3.1.2 Reducing the losses impacts in OVOC source analyses based on photochemical age**

360 Ambient OVOCs are an important fraction of VOCs, which primarily included aldehydes,  
 361 ketones, ethers, and alcohols (Mellouki et al., 2015) since OVOC photolysis is one of the main  
 362 sources of  $\bullet\text{OH}$  radicals (Li et al., 2018b). The OVOCs can be directly emitted from biogenic and  
 363 anthropogenic sources (Huang et al., 2019; Huang et al., 2020; Tanimoto et al., 2014) and can also be  
 364 formed by oxidation of precursors (e.g., PAMS species) with  $\bullet\text{OH}$  radicals,  $\text{O}_3$ , and  $\text{NO}_3$  radicals  
 365 (Legreid et al., 2008; Sahu et al., 2016; Tanimoto et al., 2014). OVOCs were commonly apportioned  
 366 by the photochemical age-based parameterization method as shown in Eq. (8).

$$\begin{aligned}
 367 \quad [\text{OVOC}] &= ER_{\text{OVOC}} \times [\text{tracer}] \times \exp(-(k_{\text{OVOC}} - k_{\text{tracer}})[\bullet\text{OH}]\Delta t) + ER_{\text{precursor}} \times \\
 368 \quad [\text{tracer}] &\times \frac{k_{\text{precursor}}}{k_{\text{OVOC}} - k_{\text{precursor}}} \times \frac{\exp(-k_{\text{precursor}}[\bullet\text{OH}]\Delta t) - \exp(-k_{\text{OVOC}}[\bullet\text{OH}]\Delta t)}{\exp(-k_{\text{tracer}}[\bullet\text{OH}]\Delta t)} + ER_{\text{biogenic}} \times \\
 369 \quad &(\text{isoprene}_{\text{source}}) + [\text{background}] \tag{8}
 \end{aligned}$$

370 where [OVOC], [tracer], and [background] are the concentrations of measured ambient OVOCs, the  
 371 tracer (e.g., benzene, acetylene:  $\text{C}_2\text{H}_2$ , or carbon monoxide:  $\text{CO}$ , Table S7) from primary  
 372 anthropogenic sources, and background OVOC concentration, respectively;  $k_{\text{OVOC}}$ ,  $k_{\text{tracer}}$ , and  $k_{\text{precursor}}$

373 are the  $\bullet\text{OH}$  rate constants of the OVOCs, tracer, and precursor, respectively, and  $k_{\text{OVOC}}$  and  $k_{\text{tracer}}$  are  
374 commonly obtained from the related publications (Atkinson and Arey, 2003);  $ER_{\text{OVOC}}$  and  $ER_{\text{precursor}}$   
375 are the emission ratios of OVOCs and precursors relative to the tracer, respectively,  $ER_{\text{biogenic}}$  is the  
376 emission ratio of OVOCs to the isoprene concentration emitted from biogenic sources (i.e.,  
377  $isoprene_{\text{source}}$ ), and  $ER_{\text{OVOC}}$ ,  $ER_{\text{precursor}}$ ,  $ER_{\text{biogenic}}$ ,  $k_{\text{precursor}}$ , and [background] can be determined from a  
378 linear least-squares fit that minimizes the difference between the measured [OVOC] and those  
379 calculated from Eq. (8), ER values in the reviewed papers are listed in Table S8;  $[\bullet\text{OH}]\Delta t$  represents  
380 the exposure of  $\bullet\text{OH}$  radicals, which can be estimated by Eq. (2); and  $isoprene_{\text{source}}$  can be estimated  
381 by Eqs. (6)-(7) and (1) based on the measured concentrations of ambient isoprene and its  
382 photochemical products (i.e., MVK and MACR). The chemical removal of isoprene in the  
383 atmosphere is so rapid that it is impossible to differentiate between primary and secondary OVOC  
384 sources based on the measured data. Therefore,  $isoprene_{\text{source}}$  represents both primary and secondary  
385 biogenic sources (de Gouw et al., 2005).

386 However, this method remains highly uncertain because of the source complexities of OVOCs  
387 (Mo et al., 2016; Schlundt et al., 2017). This method assumes that (1) anthropogenic emissions of  
388 OVOCs and their precursors are proportional to the selected primary tracer, (2) the removal process  
389 of OVOCs is dominated by reactions with  $\bullet\text{OH}$  radicals, (3) biogenic sources of OVOCs are  
390 proportional to the emission of isoprene, and (4) the photochemical age for a sampled air mass can  
391 be determined (de Gouw et al., 2005).

392 This method estimates the source contributions of primary and secondary anthropogenic sources,  
393 biogenic emissions, and background to different OVOC species, and then further obtained the  
394 contributions of the four types of sources to the ambient OVOCs (de Gouw et al., 2005; Yuan et al.,  
395 2012b; Zhu et al., 2021). This approach was difficult to finely apportion the contributions of primary  
396 anthropogenic sources and to obtain the contributions of primary and secondary biogenic emissions.  
397 Therefore, to obtain refined source apportioned results for OVOCs, many studies incorporated both  
398 OVOC and PAMS species into the receptor model for source analyses (Guan et al., 2020; Yang et al.,  
399 2019; Zhou et al., 2022).

400 This method defines the measured concentrations of OVOCs as the sum of the concentrations  
401 after photochemical losses (i.e., losses caused by the formation of  $\text{O}_3$  and SOAs via photochemical

402 reactions) of OVOCs directly emitted by anthropogenic sources (i.e., the first term in Eq. (8)), the  
403 concentrations after photochemical losses of OVOCs formed by the conversion of precursors emitted  
404 from anthropogenic sources (i.e., the second term in Eq. (8)), the concentration from biogenic  
405 emissions (without considering losses because of small contributions of biogenic emissions) (i.e., the  
406 third term in Eq. (8)), and the background concentration (i.e., the fourth term in Eq. (8)). Therefore,  
407 although this approach considered the influences of photochemical losses in the calculation process,  
408 the final results only reflected the contributions of four types of sources to the measured OVOCs and  
409 could not characterize the impacts of photochemical losses. In addition, to compensate for the  
410 photolytic losses in OVOCs in Eq. (8), which was not considered by de Gouw et al. (2005), Wang et  
411 al. (2017) introduced a modification coefficient ( $m$ ) before the  $k_{\text{OVOC}}$  to modify it, assuming that the  
412 photolysis rate is proportional to the  $\bullet\text{OH}$  reaction rate (de Gouw et al., 2018; Wang et al., 2017).  
413 Meanwhile, Huang et al. (2020) and Zhu et al. (2021) also conducted relevant studies using this  
414 coefficient. The  $m$  value depends on the relative rate of photolysis versus the  $\bullet\text{OH}$  reaction for an  
415 OVOC species (Huang et al., 2020). However, related studies remain limited at present.

416 Additionally, some studies have attempted to estimate the initial concentrations of OVOCs  
417 using traditional photochemical age-based parameterization method (i.e., Eqs. (1)-(2)) to correct their  
418 reactive losses, and then incorporate initial data into PMF for source analyses (Cui et al., 2024; Li et  
419 al., 2023; Ren et al., 2024; Zhang et al., 2024b). However, due to the complexity of the OVOC  
420 sources and the substantial differences in emission sources from PAMS species (de Gouw et al., 2018;  
421 Huang et al., 2020; Zhu et al., 2021), this method for correcting OVOC losses had high uncertainty.  
422 The rational estimation approaches remain to be studied.

423

### 424 **3.1.3 Methods for reducing loss effects in CMB source apportionments**

425 The chemical mass balance method uses known profiles measured at the source. Thus, any  
426 reactions in transit will result in a change in that source's profile and difficulty in fitting the data at  
427 the receptor site. Thus, corrections to the receptor site data are needed to make them comparable to  
428 the measured source profiles. As early as the 1980s and 1990s, studies had been conducted to reduce  
429 the impacts of different VOC species reactions on CMB estimation results, including minimization  
430 of reaction effects by limiting source profiles to VOCs with similar reaction rates (Harley et al., 1992;



431 Lewis et al., 1993; Nelson and Quigley, 1983; Wadden et al., 1986) and VOC samples obtained  
432 during winter (Aronian et al., 1989) or early in the morning (Scheff and Klevs, 1987). In 1983,  
433 Nelson and Quigley (1983) estimated the reactions extent by the changes in the ratios of xylene to  
434 ethylbenzene at the receptor and the release site and then obtained decay factors of other VOC  
435 species. However, it was found that the concentrations adjusted by decay factors at the receptor site  
436 had little impact on their estimated source contributions (Nelson and Quigley, 1983). Since the  
437 emissions varied with time, especially during periods of intense photochemical reactions (Lin and  
438 Milford, 1994), some studies tried to use decay factors to adjust the fractions of VOC species in the  
439 source profiles at different times, making them more compatible with the data measured at the  
440 receptor site (Lin and Milford, 1994; Na and Pyo Kim, 2007). However, this approach has not been  
441 used other than in these two instances.

442 There are two methods to obtain the decay factors: one method was to considering an urban  
443 airshed as a continuous stirred tank reactor and relating the decay factor for a given species to its  
444 first-order reaction rate constant (Friedlander, 1981). The other method was to conduct estimation  
445 based on the reaction rate constants of specific VOC species and “aging coefficients” (Junninen et al.,  
446 2006; Lin and Milford, 1994). The details and limitations of the two methods are provided in Text S1.  
447 However, this method of adjusting source profiles cannot truly address the issue of reactive losses  
448 affecting the CMB apportioned results. Since the receptor measured data and adjusted source profiles  
449 input to CMB were both data after reactive losses. With the progress of VOC source analyses studies  
450 in recent years, this method has not been widely applied and further developed in CMB source  
451 analyses. This change might be because CMB itself required input from VOC source profiles, but  
452 there were relatively few locally measured VOC source profiles due to high costs and difficult to  
453 obtain access to the source facilities (Yang et al., 2022b).

454 Alternatively, with the monitoring of highly time resolved VOCs data in recent years, it became  
455 infeasible to use CMB to conduct source analyses, because CMB requires analysis of one sample  
456 (one hour) at a time resulting in needing thousands of separate analyses (Yang et al., 2022b).  
457 Additionally, the weights in the analyses are generally based only on the measurement uncertainties  
458 of the measurement samples from specific sources at a few specific times. Thus, CMB does not  
459 account for variability in the source profiles that is included in the uncertainty values used to weight

460 the data in PMF (Yang et al., 2022b). To reduce the impact of reactive losses, the initial concentration  
461 data estimated by photochemical age-based parameterization method has been used as input into  
462 CMB for source analyses (Shao et al., 2011). Additionally, the methods for reducing the impacts of  
463 reactive losses in PMF and CMB primarily focused on the PAMS species, and few considerations for  
464 OVOCs.

465

### 466 **3.2 Effects of reactive losses on source analyses**

467 According to the review of relevant publications, it was found that most of the current studies  
468 analyzed the impacts of VOC photochemical losses on the source analyses by comparing the PMF  
469 apportioned results based on measured and initial concentrations estimated by a photochemical age-  
470 based parameterization method (Gao et al., 2018; Gu et al., 2023; Kong et al., 2023; Li et al., 2023;  
471 Liu et al., 2023c; Zou et al., 2023) or comparing the apportioned results based on the daytime and  
472 nighttime VOCs data (Liu et al., 2025). Reactive losses substantially reduced the concentrations  
473 (ppbv/ppbv) of highly reactive VOC species in PMF resolved factor profiles based on the measured  
474 data. For example, Liu et al. (2023a) investigated the impacts of photochemical losses of ambient  
475 VOCs on the PMF resolved profiles by comparing the initial and measured data results. They found  
476 that the concentrations of VOC species with relatively low reactivities (e.g., ethane, propane, n-  
477 butane, and i-butane) were higher in the factor profiles apportioned from the measured data, while  
478 those of VOC species with relatively high reactivities (e.g., m,p-xylene, isoprene, and propene) were  
479 lower in the measured data resolved profiles. Gu et al. (2023) also reported the similar results.  
480 Meanwhile, Liu et al. (2025) also reported that reactive losses clearly reduced the concentrations of  
481 dominant VOC species with high reactivities in the profiles of solvent use, petrochemical industry  
482 emissions, and combustion sources by comparing the daytime and nighttime resolved profiles.

483 Additionally, VOC reactive losses can result in the substantial underestimation of the PMF  
484 apportioned contributions of sources that emitted highly reactive species, and emission sources with  
485 substantially underestimated contributions varied in different cities. For example, Wang et al. (2013)  
486 found that the contributions of biogenic and industrial emissions in Shanghai were underestimated by  
487 30% and 10%, respectively, due to photochemical reactive losses. He et al. (2024) found that the  
488 underestimations of industrial source contributions in Guangzhou were markedly higher than those

489 of other sources. Liu et al. (2023a) suggested that biogenic emissions and polymer production-related  
 490 industrial sources in Tianjin were underestimated by 73% and 50%, respectively. In addition, Wu et  
 491 al. (2023a), Zhang et al. (2024a), and Gu et al. (2023) also suggested that the underestimations of  
 492 contributions of biogenic emissions in Beijing, Langfang, and Qingdao were substantially higher  
 493 than those of any other sources. However, Wang et al. (2024a) found that the contributions of solvent  
 494 usage and biomass burning in Zhengzhou were underestimated by 31.5% and 15.4%, higher than  
 495 other sources. Cui et al. (2024) suggested that the contributions of petrochemical industries, diesel  
 496 vehicle emissions, biogenic emissions, and oxidation formation in Shijiazhuang were underestimated  
 497 by 72.0 %, 71.0 %, 64.5 %, and 44.0 %, respectively. However, due to the uncertainty of initial  
 498 concentration estimation, the reliability of these results needs further validation and evaluation. Thus,  
 499 further research is required.

500

### 501 3.3 Estimation methods of VOC reactive losses

502 There were two main methods for estimating the reactive losses of VOCs in the atmosphere.  
 503 The first method was based on the differences between the initial and measured VOC concentrations  
 504 (Table S9) (as shown in Eq. (9)) (Wang et al., 2023; Wu et al., 2023b). The initial concentration was  
 505 generally estimated using the photochemical age-based parameterization method mentioned in  
 506 Section 3.1.1.2 (as shown in Eq. (1)) (Liu et al., 2023a; Wu et al., 2023b). Due to the uncertainty in  
 507 the initial concentration estimation, there were also uncertainties in the estimated photochemical  
 508 losses.

$$509 [VOC]_{C,t} = [VOC]_t \times (\exp(k_{VOC}[\bullet OH]\Delta t) - 1) \quad (9)$$

510 where  $[VOC]_{C,t}$  represents the photochemical loss of VOC at time  $t$ ,  $K_{voc}$  represents the reaction rate  
 511 constants with  $\bullet OH$  radicals of VOC, and  $[\bullet OH]\Delta t$  represents the exposure of  $\bullet OH$  radicals.

512 The second method applies to other VOC species without isoprene in which the losses could be  
 513 estimated utilizing isoprene conversion (Wiedinmyer et al., 2001). In this study, it was defined as the  
 514 isoprene loss reference method (Table S9). Its principle is to first use the photochemical-age  
 515 parameter method based on the sequential reaction model as shown in Eqs. (1) and (6)-(7), to  
 516 estimate the photochemical loss of isoprene (i.e.,  $\Delta isoprene$ ), and then the photochemical losses of  
 517 other VOC species can be estimated by proportion relationship  $(\frac{k_{VOC}}{k_{isoprene}} \times \frac{[VOC]_t}{[isoprene]_t})$  between  $\Delta$

518 *isoprene* and other species losses (Wiedinmyer et al., 2001; Xie et al., 2008) (as shown in Eq. (10)).  
519 This method assumes that the relative source strengths of VOCs are constant in an immediate area  
520 surrounding the site, that atmospheric transport and dispersion are non-limiting factors compared  
521 with chemistry (Xie et al., 2008), and the photochemical losses of other VOC species can be  
522 calculated by Eq. (10).

$$523 \quad [VOC]_{C,t} = \Delta isoprene \times \frac{k_{VOC}}{k_{isoprene}} \times \frac{[VOC]_t}{[isoprene]_t} \quad (10)$$

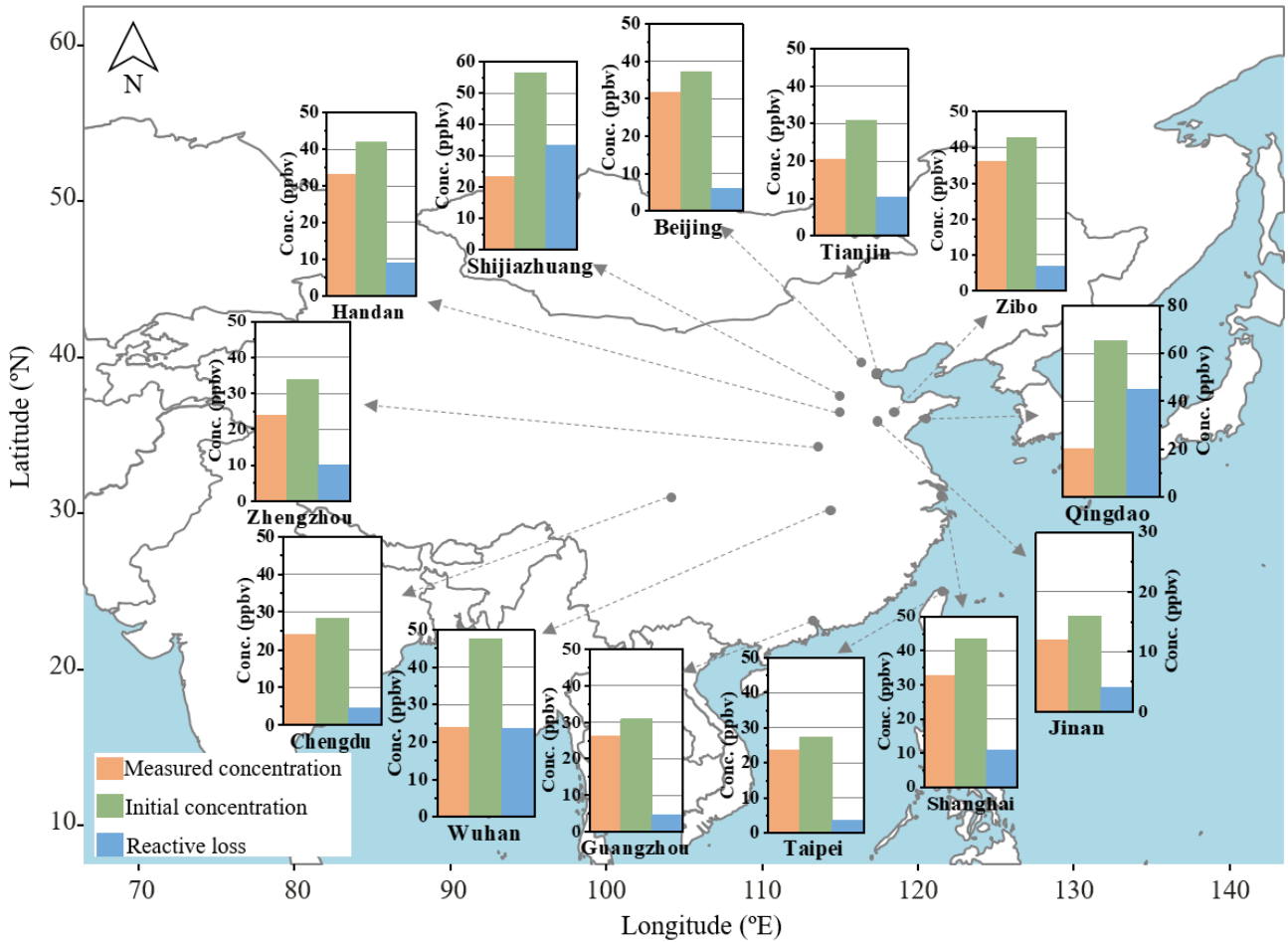
524 where  $k_{isoprene}$  represents the reaction rate constants with  $\bullet$ OH radicals of isoprene, and  $[isoprene]_t$   
525 represents the measured concentration of isoprene at time  $t$ .

526 Because isoprene mainly originates from natural sources (Fu et al., 2008; Kuhn et al., 2004; Lu  
527 et al., 2019), while many other VOC species are primarily emitted from anthropogenic sources (Li et  
528 al., 2021; Seinfeld and Pandis, 1986; Wei et al., 2011). Therefore, the assumption of this method  
529 itself has an obvious issue, which resulted in high uncertainties in the estimations. This defect  
530 substantially limits its application, only 2 publications have utilized this method to quantify chemical  
531 losses of VOCs in this study (Wiedinmyer et al., 2001; Xie et al., 2008), and most studies still used  
532 the first method (Table S9).

533

### 534 **3.4 Spatiotemporal variation of VOC reactive losses**

535 According to the papers reviewed in this study, we found that studies on estimation of VOC  
536 photochemical losses were mainly conducted in Chinese cities. The VOC reactive losses in different  
537 cities are provided in Figs. 2-3 and S1 and Tables S10-S11. Due to relatively limited number of  
538 studies, the representativeness of the analyses of the spatiotemporal distributions of photochemical  
539 losses of VOCs might also be limited. In addition, this study converted the unit of ppbC in some  
540 papers (Chen et al., 2023) to ppbv for better comparative analyses.



541  
 542 **Figure 2.** The measured, initial, and reactive loss concentrations of ambient VOCs in Beijing (Gao et al., 2018; Ma et al., 2022; Zhan et al., 2021), Tianjin (Liu et al., 2023a; Wang et al., 2023), Shanghai  
 543 (Ren et al., 2024; Wang et al., 2013), Chengdu (Kong et al., 2023), Guangzhou (He et al., 2024;  
 544 Wang et al., 2023), Qingdao (Gu et al., 2023), Shijiazhuang (Cui et al., 2024), Jinan (Liu et al.,  
 545 2023c), Zhengzhou (Wang et al., 2024a), Wuhan (Xu et al., 2023), Handan (Wei et al., 2022), Zibo  
 546 (Wang et al., 2024b), and Taipei (Chen et al., 2023). The data in Beijing, Tianjin, Shanghai, and  
 547 Guangzhou was the average from all published papers data. The base map is from Natural Earth.  
 548

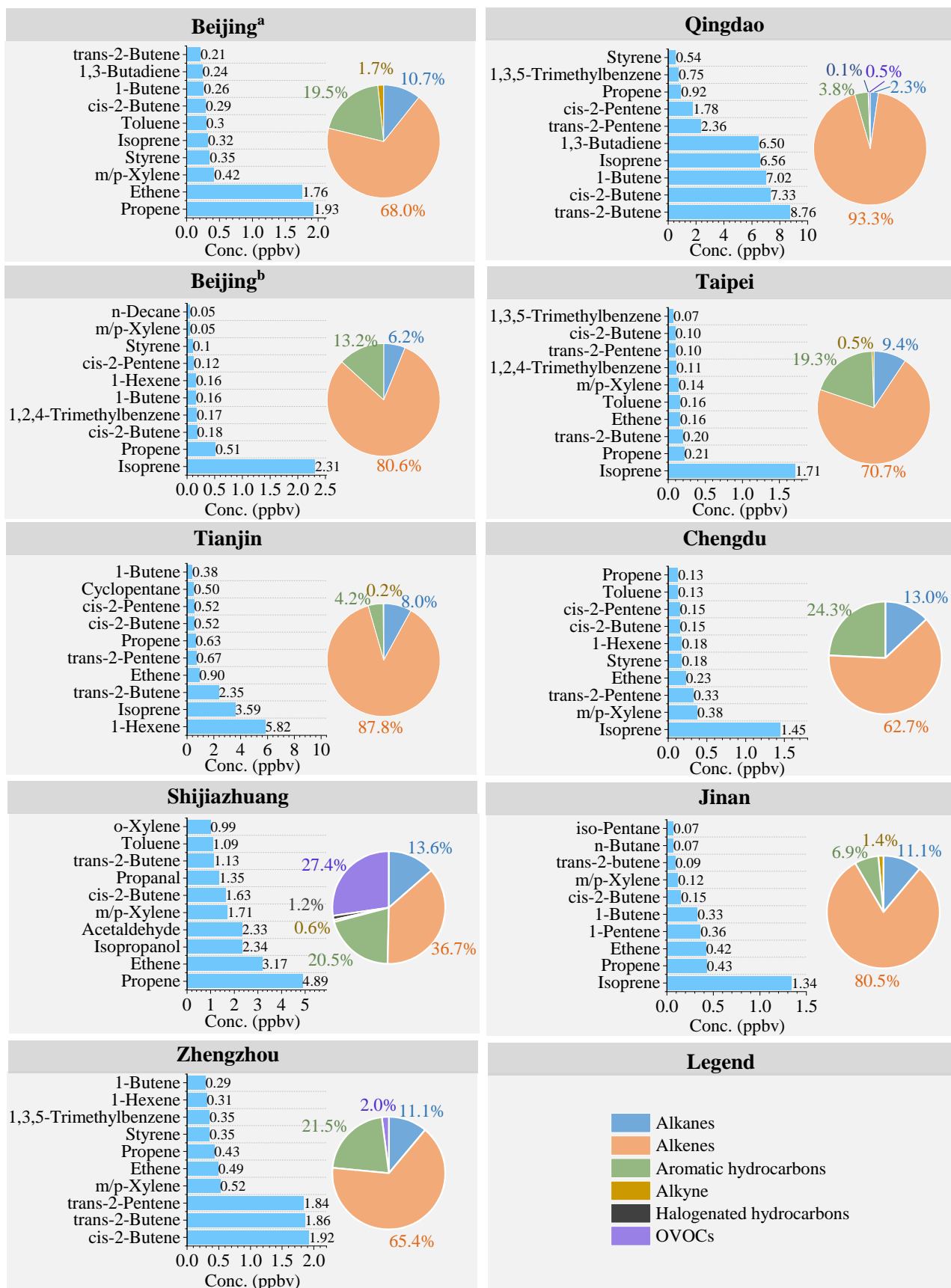
549  
 550 The photochemical reactive loss of ambient VOCs in Qingdao was the highest (45.1 ppbv),  
 551 followed by Shijiazhuang (33.2 ppbv), Wuhan (23.7 ppbv), Shanghai (10.9 ppbv), Tianjin (10.4  
 552 ppbv), Zhengzhou (10.2 ppbv), and Handan (8.90 ppbv) (Fig. 2). VOC reactive losses were relatively  
 553 lower in Zibo (6.8 ppbv), Beijing (6.00 ppbv), Guangzhou (4.65 ppbv), Chengdu (4.48 ppbv), Jinan  
 554 (4.00 ppbv), and Taipei (3.69 ppbv). The chemical loss rates (i.e., the proportion of chemical loss in  
 555 the initial concentration, %) in Qingdao (69.1%) and Shijiazhuang (58.9%) were the highest,  
 556 followed by Wuhan (49.8%) and Tianjin (33.8%). In contrast, chemical loss rates in Zhengzhou  
 557 (29.9%), Shanghai (25.1%), Jinan (25.0%), Handan (21.1%), Beijing (16.1%), Zibo (15.9%),

558 Chengdu (15.8%), Guangzhou (15.1%), and Taipei (13.4%) were relatively lower. However, due to  
559 differences in observation periods and measured VOC species, the comparability of chemical  
560 reactive losses and loss rates between different cities is limited and differences uncertain.

561 Compared to other VOC groups, alkenes had the highest reactive loss (Figs. 3 and S1),  
562 accounting for 36.7%-93.3% of the total losses, followed by aromatic hydrocarbons (3.81%-24.3%),  
563 and alkanes (2.33%-13.6%) (Fig. 3 and Table S11). There were substantial differences in VOC  
564 species with high losses in different cities (Fig. 3). The losses of ethene, propene, and isoprene in  
565 most cities were relatively higher than those of other species (Fig. 3), likely closely related to their  
566 high reactivities (Table S4). The isoprene losses in Beijing, Chengdu, Jinan, and Taipei were all the  
567 highest compared to other species (Fig. 3). However, the reactive losses of trans-2-butene and cis-2-  
568 butene in Qingdao and Zhengzhou were substantially higher than other VOC species. The reactive  
569 loss of 1-hexene in Tianjin was remarkably higher compared to other species. Meanwhile, the trans-  
570 2-butene loss in Tianjin was also relatively higher (Fig. 3).

571 In addition, the VOC reactive losses in spring and summer were substantially higher than those  
572 in autumn and winter. Atmospheric oxidation and meteorological factors can have impacts on the  
573 VOC losses. The trend of the reactive losses of VOCs was often consistent with the changes in the  
574 concentrations of  $O_x$  ( $O_3+NO_2$ ) in the atmosphere (Wang et al., 2013). In addition, studies also found  
575 that both solar radiation and temperature can have a substantial impact on the chemical reactions of  
576 VOCs (Yadav et al., 2016). However, compared to the solar radiation, temperature directly affects  
577 the reaction rates of VOC species (Atkinson and Arey, 2003). Liu et al. (2023a) suggested that the  
578 reactive losses of ambient VOCs were highly correlated with temperature; when the temperature was  
579 above 25 °C, the losses of VOCs increased most substantially.

580



581

582

583

**Figure 3.** The photochemical losses and percentages of the main VOCs in Beijing (a: cited from publication (Gao et al., 2018) and b: cited from publication (Zhan et al., 2021)), Qingdao (Gu et al.,

584 2023), Taipei (Chen et al., 2023), Tianjin (Liu et al., 2023a), Chengdu (Kong et al., 2023),  
585 Shijiazhuang (Cui et al., 2024), Jinan (Liu et al., 2023c), and Zhengzhou (Wang et al., 2024a).

586

### 587 **3.5 Source analyses of VOC reactive losses in the atmosphere**

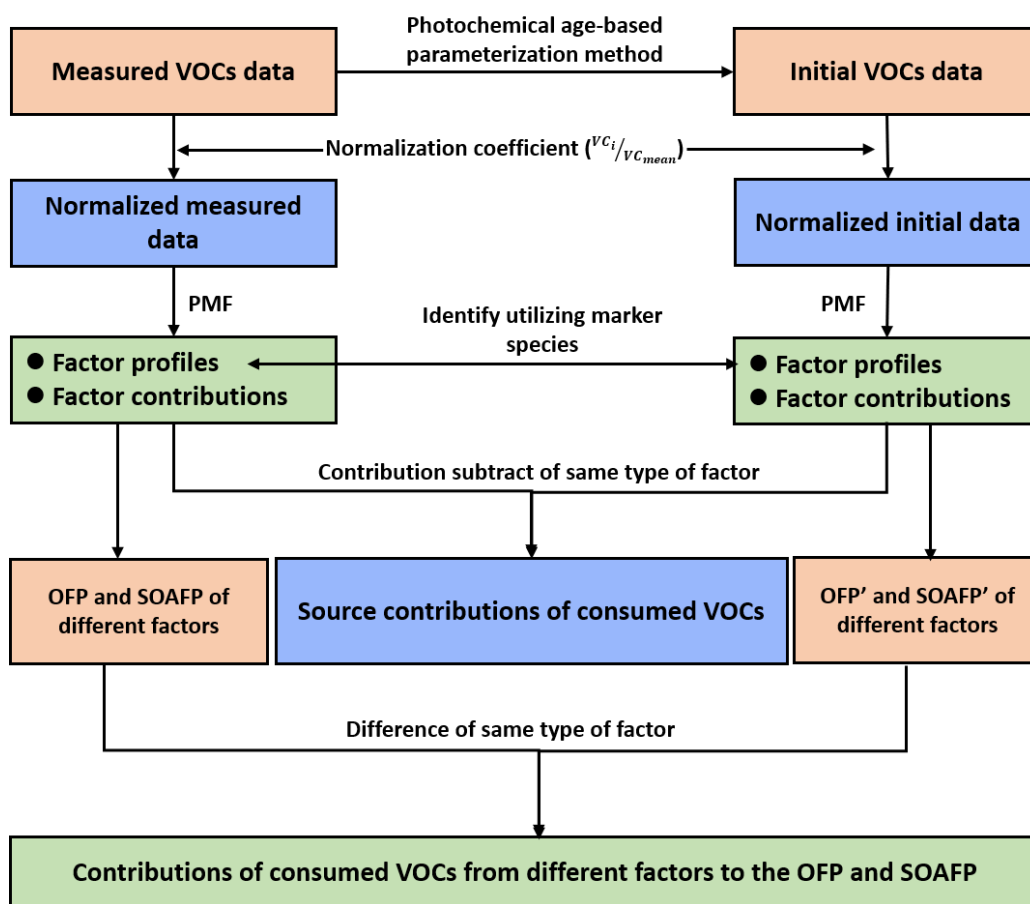
588 At present, source apportionments utilizing the measured and initial VOC concentrations do not  
589 consider the complementary issues of O<sub>3</sub> and SOA formation (Cui et al., 2024; Gu et al., 2023; Wang  
590 et al., 2022b). The chemical losses of VOCs by reaction (i.e., reactive VOCs forming SOA and O<sub>3</sub>)  
591 were real contributors to these pollutants (Ma et al., 2022; Wang et al., 2013). Thus, it is important to  
592 apportion the reactive losses of VOCs to provide the input needed to identify the responsible source  
593 types and thereby allow effective control of O<sub>3</sub> and secondary aerosol pollution. In 2023, Gu et al.  
594 (2023) developed a source analyses method for consumed VOCs (i.e., the reactive losses of VOCs)  
595 in the atmosphere and conducted an applied study. This method was that the measured and initial  
596 data were first used to conduct source analyses by PMF, and then the difference in the contributions  
597 of the same factors from the paired analyses were considered to be the source contribution of  
598 consumed VOCs in the atmosphere. The ozone formation potential (OFP) and SOA formation  
599 potential (SOAFP) of the consumed VOCs from the different sources were estimated. To reduce the  
600 impacts of atmospheric dispersion on the apportioned results, dispersion-normalized PMF (DN-PMF)  
601 was applied for source analyses in this method, and its principle is provided by Dai et al. (2020) and  
602 Gu et al. (2022). The method flowchart is shown in Fig. 4. Similar studies were reported by Wang et  
603 al. (2022b) and Wang et al. (2023). In 2024, based on the studies of Gu et al. (2023), Cui et al. (2024)  
604 developed a primary and oxidative source analyses method of consumed VOCs in the atmosphere.  
605 However, the current methods for apportioning the primary or oxidative sources of consumed VOCs  
606 in the atmosphere are still imperfect. The shortcomings included: the uncertainty in estimating the  
607 initial concentrations of VOCs (as shown in Section 3.1.1.2); the issue of factor identification, even  
608 if identified as the same type of factor, there were substantial differences in their factor profiles (Liu  
609 et al., 2023a). The current studies primarily focused on the PAMS species, and reported  
610 apportionments of OVOC losses are still limited. There were numerous technical issues, for example,  
611 there is still a lack of rational methods for estimating the OVOC initial concentrations.

612 The source analyses of the consumed VOCs conducted by Gu et al. (2023) in Qingdao found  
613 that biogenic emissions (56.3%), vehicle emissions (17.2%), and gasoline evaporation (9.37%) were



614 the main sources of the consumed VOCs. However, the apportioned results from Wang et al. (2023)  
 615 in Tianjin and Guangzhou suggested that biogenic emissions (43% and 35%, respectively), solvent  
 616 usage (14% and 18%, respectively), and industrial sources (14% and 22%, respectively) were the  
 617 major contributors of the consumed VOCs. Therefore, biogenic emissions might be an important  
 618 source of reactive losses of VOCs. However, Cui et al. (2024) suggested that the petrochemical  
 619 industries (36.9% and 51.7%) and oxidation formation (20.6% and 35.6%) were the largest  
 620 contributions to the consumed VOCs and OVOCs in Shijiazhuang during the study period. The  
 621 natural gas (5.0% and 7.6%) and the mixed source of liquefied petroleum gas and solvent use (3.1%  
 622 and 4.2%) had the relatively low contributions (Cui et al., 2024). However, due to the limited  
 623 research currently available, the representativeness of the results is still insufficient at present.

624



625

626

**Figure 4.** The method flowchart of source analyses for consumed VOCs in the atmosphere.

627

## 628 4 Conclusions

629 This study systematically reviewed the major methods of reducing the impacts of reactive loss

630 and the existing critical issues in the current VOC source analyses research. The purpose was to  
631 clarify the future research directions needed to improve the accuracy of VOC source apportionments,  
632 and provided a potential supporting role in completely solving the issue of VOC chemical conversion.  
633 PMF is currently the most useful tool in treating highly time-resolved data compared to other  
634 receptor models. Estimating the initial concentrations of ambient VOCs based on photochemical age  
635 is the primary approach to reduce reactive loss effects in PMF. However, due to the shortcomings  
636 existing in the photochemical-age estimation method, such as only considering the photochemical  
637 reactions with •OH radicals during the day and not considering the reactions with O<sub>3</sub> (especially for  
638 alkene species) and NO<sub>3</sub> radicals, difficulty in dynamically adjusting the required •OH-radical  
639 reaction rate constants with temperature changes, high uncertainty in the estimation of  
640 photochemical age, and difficulty in quantifying the initial concentrations of OVOCs, etc., resulting  
641 in substantial uncertainty in the PMF apportioned results based on the initial concentration data.  
642 Furthermore, both the measured and initial VOCs data do not match the ambient O<sub>3</sub> or SOAs  
643 measured synchronously, the VOCs consumed by reactions are real contributors to these pollutants.  
644 Source analyses of the consumed VOCs can effectively guide the prevention and control of O<sub>3</sub> and  
645 SOAs pollution in the atmosphere. Therefore, in addition to the research into more accurate  
646 quantitative approaches for ambient VOC reactive losses, the source analyses of consumed VOCs  
647 and the accurate quantification of their separate contributions to O<sub>3</sub> and SOAs should also be  
648 important directions for future research.

649

#### 650 **Author contributions**

651 BSL: Writing—original draft, Writing—review & editing. YG: Data curation, Writing—original draft.  
652 YTW: Data curation. QLD and SJS: Investigation and Supervision. YCF: Supervision. PKH:  
653 Supervision—review & editing.

654

#### 655 **Competing interests**

656 The author declares that they have no competing interests.

657

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661

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