

Measurement report: VOC characteristics at different land-use types in Shanghai: spatio-temporal variation, source apportionment, and impact on secondary formations of ozone and aerosol

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Abstract. Volatile organic compounds (VOCs) have important impacts on air quality, climate change and human health. In order to identify the spatio-temporal variations, sources and ozone and secondary organic aerosol formation potentials of the atmospheric VOCs, a concurrent multi-site observation campaign was performed at the supersites of Shanghai, East China, in the first three months of 2019. The sampling sites are located at the different land-use types including industrial district (the Jinshan site: JS), residential and commercial mixed districts (the Pudong site: PD) and background district (the Qingpu site: QP) of Shanghai. During the observation period, the average VOC concentrations were sensitive to the land-use types in the order of JS (21.88 ± 12.58 ppb) > PD (21.36 ± 8.58 ppb) > QP (11.93 ± 6.33 ppb) sites. The predominant VOC category was alkanes (49.32-71.48 %), followed by aromatics (10.70-21.00 %), alkenes (10.13-15.30 %) and alkyne (7.69-14.80 %) at the studied sites. There were distinct diurnal variations and “weekend effects” of VOCs at the sampling sites. The VOC concentrations increased by 27.15, 32.85 and 22.42 % during the haze events relative to the clean days. Vehicle exhaust was determined as the predominant VOC source. The second largest VOC contributor was identified as industrial production at the JS and PD sites, while it proved to be fuel evaporation at the QP site. The industrial emission and biogenic source showed slight contributions to VOC concentrations at the QP and JS/PD sites, respectively. It was consistent with the regional characteristics of anthropogenic activities dominated by land-use types. High potential source contribution function (PSCF) values primarily appeared in the northeastern and northern areas surrounding sampling sites, suggesting the strong local emissions. The ozone formation potentials (OFP) values of each land-use types were in order of JS (50.89 ± 2.63 ppb) > PD (33.94 ± 1.52 ppb) > QP (24.26 ± 1.43 ppb) sites, with alkenes and aromatics being the predominate contributors. Secondary organic aerosol formation potential (SOAFP), mainly contributed by the aromatics, was the highest at the JS site (1.00 ± 2.03 $\mu\text{g m}^{-3}$), followed by the PD (0.46 ± 0.88 $\mu\text{g m}^{-3}$) and QP (0.41 ± 0.58 $\mu\text{g m}^{-3}$) sites. The VOCs-PM_{2.5} sensitivity analysis showed that the VOCs at the QP site could be more sensitive to the variation of PM_{2.5} concentration relative to the

35 other two sites. Alkenes and aromatics are both the key concerns in controlling the VOC-related pollution of O₃ and secondary organic aerosol (SOA) in Shanghai. These findings provide more information on the accurate air-quality control at a city level in China. The results shown herein highlight that the simultaneous multiple-site measurements with the different land-use type in the megacity or city cluster could be more appropriate to fully understand the VOC characteristics relative to a single-site measurement performed normally.

1. Introduction

40 Serious air pollution in China is currently characterized by the high levels of ozone (O₃) and fine particulate matters (PM) especially PM_{2.5} (PM with an aerodynamic diameter less than 2.5 μm). The atmospheric volatile organic compounds (VOCs) greatly influence the O₃ and PM_{2.5} formations, and function as the important precursors (Carter, 1994; Liu et al., 2008; Yuan et al., 2013; Lu et al., 2018; Ma et al., 2019; Yu et al., 2021). In brief, the primary VOCs can be oxidized and/or photolyzed by OH, O₃ and NO₃ to produce secondary VOCs, e.g., non-/semi-volatile organic compounds: NVOCs/SVOCs, which both
45 further transforms into secondary organic aerosol (SOA) *via* a series of atmospheric processes (Odum et al., 1997; Ng et al., 2007; Heald et al., 2020). It was well known that SOA has strong influences on the radiative forcing and climate change (Sadeghi et al., 2021). On the other hand, active VOCs react with NO_x and radicals *via* an array of photochemical pathways to produce O₃. Photochemical Assessment Monitoring Stations (PAMS) have confirmed that totally 57 VOCs, including C₂-C₁₀ alkanes, alkenes, alkynes and aromatics are extremely contributed to the formation of O₃ (US EPA, 1990).
50 Atmospheric O₃ shows profound impacts on the atmospheric oxidizing capacity, agriculture production and ecosystem (Liu, 1987; Carter, 1994; Mousavinezhad et al., 2021; Sadeghi et al., 2021). Additionally, O₃ and PM_{2.5}, together with their VOC precursors can damage human health and lead to many diseases such as cancers, respiratory and nervous systems (Rumchev et al., 2007; Amor-Carro et al., 2020). Therefore, the VOC-related studies can provide scientific-based information for the decision-makers to draw up strategies to control PM_{2.5} and O₃.

55 The long-term VOC emission inventory highlighted that there were significant spatial discrepancies of VOC emissions (Li et al., 2019a). The observation campaign also showed that VOC concentrations varied with the sampling sites. These phenomena were attributed to the fact that VOC concentrations were closely correlated with the land-use types. For instance, the VOC concentrations varied approximately from 20 to 40 ppb at the urban area (Geng et al., 2008; Cai et al., 2010b; Huang et al., 2015; Dai et al., 2017; Liu et al., 2019; Xu et al., 2019; Ren et al., 2020; Wang et al., 2020; Liu et al., 2021),
60 and ~ 90 ppb at the industrial area in Shanghai, China (Zhang et al., 2018; Wang et al., 2021). Kumar et al. (2018) found that the VOC concentration at the urban area was approximately twice higher than that at the rural area in Delhi, India, as the finding of Tang et al. (2008). However, the reported VOC concentrations were widely discussed by single-site measurements, the limited knowledge is available on the multi-site research at a city level. Besides, the land-use types not only influence the VOC concentrations but also the sources especially the anthropogenic sources (Yoo et al., 2015; Chen et al., 2017; Wang et al., 2017; Jookjantra et al., 2022). For example, in Shanghai, China, vehicle-related source was regarded as the predominate
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VOC source accounting for 34 % at the urban area (Cai et al., 2010a), while the industrial emission accounted for over 50 % of the measured VOCs near an industrial site (Zhang et al., 2018). In Zhengzhou, China, the solvent utilization was the largest contributor (34 %) at the urban area (Li et al., 2020a), whereas coal and biomass burning was the major source for ambient VOCs (30.6 %) at the Gongshui site surrounded by several manufacturing plants (Li et al., 2019b). Such great discrepancies among the different land-use types results in the fact that it is necessary to perform concurrent multiple-site measurements of VOC characteristics at a city level.

The diversities of VOC concentrations among the different land-use types could affect the ozone and SOA formation potentials (OFP and SOAFP), resulting in the variations of O₃ and SOA concentrations. For instance, Song et al. (2021) found that the OFP values at the commercial and residential areas were approximately twice higher than that at the area dominated by natural vegetation in Xi'an, China, which was in agreement with the finding of Kumar et al. (2018). Such scenario could influence the atmospheric O₃ concentration in that VOCs is an important precursor of O₃ formation. In detail, atmospheric VOCs undergoes degradation to produce oxidants (HO₂ and RO₂), which further oxidizes atmospheric NO, followed by producing NO₂ and the formation of O₃ finally *via* the photochemical pathways (Wang et al., 2017). In terms of SOAFP, there was higher value at the urban area relative to the suburban area, increased by 54.84 % in Beijing, China (Zhan et al., 2021; Liu et al., 2022). This discrepancy further had profound impacts on the concentration, physical property and chemical composition of SOA. As the key precursor of SOA, VOCs can be oxidized to produce the low VOCs, followed by the formation of SOA *via* homogeneous nucleation (Merikanto et al., 2009). Moreover, the partitioning of semi-volatile products from VOCs and oxidants gas-phase photochemical reactions to form SOA (Pankow, 1994; Lim et al., 2010). Additionally, low VOCs are produced *via* the aqueous-phase reactions in atmospheric waters e.g., clouds, fogs, and aerosol water which are largely retained in the particle-phase to generate SOA (Lim et al., 2010).

As the junction of transportation and a center of industry, finance, economy and technology of East China, Shanghai covers an area of 6340 km² and the population over 24 million. The expanding urbanization and industrialization jointly aggravate the VOC pollution. Moreover, the O₃ concentration at the urban area in Shanghai increased by ~ 67 % from 2006 to 2015 with the growth rate of 1.1 ppbv per year (Gao et al., 2017). The maximum 1-hour concentration of O₃ could exceed 380 µg m⁻³ during polluted days (Shi et al., 2015; Gu et al., 2020). Such scenario suggested that O₃ played an important role in atmospheric pollution and Shanghai was suffering from heavy O₃ pollution. Additionally, the large changes of land-use occurred in Shanghai due to the rapid development e.g., many cultivated areas became urban and/or industrial zones, resulting in the diverse land-use types (Tian et al., 2017). Therefore, Shanghai is regarded as an ideal area to perform atmospheric measurements with the different land-use types. However, many studies were mainly focused on the single-site measurements, particularly conducted at the urban site in Shanghai, resulting that the impact of land-use type on VOC characteristics is still unclear to date. Besides, they mainly concentrated on the O₃ characteristics, while the O₃ and SOA formations from VOCs and the relationship between VOCs together with PM_{2.5} and O₃ were rarely analyzed. Given the factors mentioned above, in this study, the concurrent multiple-site and high time-resolution measurement of VOCs with three typical land-use types in Shanghai for their concentrations, sources and ozone and SOA formation potentials were

100 performed. The objectives of this study are to (1) figure out the VOC characteristics varying with the land-use types, (2) identify the predominate VOC emission sources, and (3) quantitatively evaluate the roles of VOCs in the formation of PM_{2.5} and O₃. The findings could advance our knowledge in the atmospheric characteristics of VOCs and provide more control strategies for policymakers to alleviate atmospheric pollution.

2. Measurements and methods

105 2.1 Sampling site description

The three sampling sites are located at the different land-use types of Shanghai, China: industrial district (the Jinshan site: JS, 121.18 °N, 30.44 °E), residential and commercial mixed districts (the Pudong site: PD, 121.54 °N, 31.23 °E) and background district (the Qingpu site: QP, 120.98 °N, 31.09 °E) (Fig. S1). The online instruments were 30 m above the ground level to avoid airflow obstruction.

110 The JS site is located in the Second Jinshan Industrial Area of Shanghai as an industrial site. The site is situated in ~ 20 km southwest of the Shanghai Chemical Industrial Park and Garbage Disposal Incinerator-2. Moreover, a large number of chemical factories, such as rubber factories, paint solvent factories and oil refineries are located within 2 km of the JS site. In particular, Shanghai Petrochemical Industrial Limited Company (one of the largest refining-chemical integrated petrochemical companies in Shanghai) is 1.5 km southeast of the JS site. Additionally, two arterial roads are located 100 and
115 300 m away from the JS site, respectively. Thus, the land-use type of JS site could be regarded as industrial district (Tab. S1). The PD site is located in the Pudong New Area as an urban site, ~ 59 km northeast of the JS site. The Pudong New Area is the most populous district of Shanghai with a population up to 5.6 million (the Seventh Census of China). The Baosteel Group Corporation, International Cruise Port wharf, Waigaoqiao Shipbuilding LTD and Domestic Waste Incinerator totally are located within 30 km of the PD site. The Yangshan Harbor (the biggest harbor of Shanghai) and Garbage Disposal
120 Incinerator-1 are located in 51 and 37 km southeast of the PD site, respectively. Furthermore, the PD site is surrounded by three main airports of Shanghai, including Hongqiao International Airport, Longhua Airport and Pudong International Airport. In addition, the site is a junction of roads, commercial and financial areas with intense human activities, which is ~ 1.5 km away from Jinxiu Road, Jincai High School, Jinnan High School, and Jincai Experimental Primary School. Therefore, the land-use type of PD site is residential and commercial mixed districts (Tab. S1).

125 The QP site is located near the southeast of Dianshan Lake where is a tourist attraction in the Qingpu district as a city background site, ~ 50 km northwest of the JS site and 54 km southwest of the PD site. Surrounded by vegetation, the QP site is situated in ~ 7.6 and 4.6 km northeast of the Grand View Garden and Chenguang Garden, respectively. Moreover, this site is 3.6 km northeast of the Qingxi Country Park and 7.5 km southwest of the Zhuxi Garden. Besides, the QP site is close to Hu-Yu Highway. Hence, the land-use type of QP site is considered as background district (Tab. S1). Since the distinct
130 land-use types among the sampling sites were observed, it is quite necessary to perform atmospheric measurements herein.

2.2 Sampling and analysis

The concentrations of VOCs, CO, NO-NO₂-NO_x, PM_{2.5}, and meteorological factors including temperature, wind speed and relative humidity (RH) were measured in one-hour resolution. The sampling campaign was performed simultaneously at the three sites from January 1 to March 31 2019.

135 At the JS site, the VOCs was collected and analyzed by an online gas chromatography (GC866, Chromato, France) equipped with flame ionization detector (FID). In brief, after the removal of water, the samples were separated for low-carbon (C₂-C₅) and high-carbon (C₆-C₁₂) compounds at the temperature of -5°C and 25°C, respectively. Then the gas was analyzed by FID after high temperature desorption (380°C) and column chromatographic separation. At the PD site, VOCs was measured by gas chromatography (GC580-FID, PE, USA) and TD300 (a transformer driver). The samples were separated at -30°C after
140 the removal of water. Then the gas was determined by FID, after high temperature desorption (325°C) and column chromatographic separation. At the QP site, VOCs were determined by gas chromatography (GC5000 BTX/VOC, AMA, German) and a flame ionization detector (FID). The samples were condensed for low-carbon (C₂-C₅) compounds at 15°C and high-carbon (C₆-C₁₂) compounds at 30°C. Then the gas was analyzed by FID after high temperature desorption (230°C) and column chromatographic separation. The curve correlation coefficient (R²) of all of the VOCs were ≥ 0.995. The accuracy of
145 more than 95 % of compounds was ≤ ± 20 %. The precision of all target compounds was ≤ 10 %. Totally 43 species of VOCs were observed, including 16 alkanes, 11 alkenes, 16 aromatics and 1 alkyne. The minimum detection limit (MDL) of most VOC components was ≤ 0.15 ppb (Tab. S2).

The O₃, NO-NO₂-NO_x were characterized by trace gas instruments (49i ozone analyzer and 42i nitrogen oxide analyzer, produced by Thermo Environmental Instruments Inc., USA) with the detection limits of 0.50 and 0.40 ppb, respectively.

150 PM_{2.5} was monitored by a TEOM 1405-F. The meteorological variables including temperature, RH and wind speed were simultaneously acquired from a weather station about 10 km northwest of the Shanghai Academy of Environmental Sciences.

2.3 Spatial analysis

The spatial heterogeneity of VOC concentration between two different sites was determined by the coefficient of divergence (COD) (Wongphatarakul et al., 1998; Sawvel et al., 2015). The COD was calculated by Eq. (1):
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$$COD_{jk} = \sqrt{\frac{1}{p} \sum_{i=1}^p \left(\frac{x_{ij} - x_{ik}}{x_{ij} + x_{ik}} \right)^2} \quad (1)$$

where x_{ij} presents the mass concentration in i time, j and k are two datasets, p presents the number of observations. The values of COD represented the degree of similarity of the two datasets, i.e., the value of COD approached 1, illustrating that the big difference exist in the two datasets (Song et al., 2017).

160 2.4 Positive matrix factorization (PMF) model

PMF was a receptor model designed for the source identification of atmospheric pollutants (Li et al., 2015b; Hopke et al., 2016; Pallavi et al., 2019). In brief, it measured the contributions and sources of pollutant by the least squares, and was based on the mass balance instead of the spectrum of source component (Hui et al., 2018). In the PMF 5.0, the mass balance equation was calculated by Eq. (2):

$$165 \quad x_{ij} = \sum_{k=1}^p g_{ik} f_{kj} + e_{ij} \quad (2)$$

where x_{ij} is the j th species concentration in the sample i , g_{ik} represents the species contribution of the k th source to the i th sample, f_{kj} is the j th species fraction from the k th source, e_{ij} is the residual result for j th species in i th sample, p is the number of independent sources (Paatero, 1997). The function Q was an important factor of PMF (Brown et al., 2015), and calculated by iterative minimization algorithm (Hui et al., 2018). The objective function Q was shown in the Eq. (3):

$$170 \quad Q = \sum_{i=1}^n \sum_{j=1}^m \left[\frac{x_{ij} - \sum_{k=1}^p g_{ik} f_{kj}}{u_{ij}} \right] \quad (3)$$

where u_{ij} is the uncertainty of j th species in the sample i .

The uncertainty of sampling was calculated by the Eq. (4).

$$Unc = \sqrt{(EF \times conc)^2 + (0.5 \times MDL)^2} \quad (conc > MDL) \quad (4)$$

175 where MDL is the minimum detection limit, EF is the error fraction and can be set to 0.05-0.2 (Song et al., 2007). It was 0.1 in this study. In this study, four to eleven factors were utilized to determine the option solution. Q_{true}/Q_{robust} and $Q_{true}/Q_{expected}$ are important parameters for characterizing the rationality of the PMF results (Brown et al., 2015). Seven factors were regarded as the optimal solution, comparing the ratios of Q_{true}/Q_{robust} , $Q_{true}/Q_{expected}$ and PMF results. The Q_{true}/Q_{robust} values were set to 1.0 at the three sampling sites. The $Q_{true}/Q_{expected}$ values were 1.3, 1.1, and 1.0 at the JS, PD and QP sites, respectively.

180 **2.5 Potential source contribution function (PSCF) and Cluster**

PSCF and Cluster were widely used to determine the back trajectories, source and direction of pollutants (Draxier and Hess, 1998; Hong et al., 2019; Liu et al., 2019), and designed to measure the potential VOC source and primary transport pathway of trace elements (Ashbaugh et al., 1985; Xie et al., 2007; Zheng et al., 2018; Liu et al., 2020).

185 This study was determined by the 24-h back trajectories (one hour interval) at the height of 500 m via the MeteoInfoMap software. Relevant parameters were acquired from the National Oceanic and Atmospheric Administration. The study area covered by back trajectories was divided into an array of $0.25^\circ \times 0.25^\circ$ grid cell. The higher PSCF indicated that the area was a great contributor to the VOC pollution. The PSCF could be defined as the Eq. (5).

$$PSCF_{ij} = \frac{m_{ij}}{n_{ij}} \quad (5)$$

190 where the m_{ij} is the number of polluted trajectories through the grid, n_{ij} is all of the trajectories through the grid. In order to distinguish the value of PSCF and increase the accuracy, the weight function W_{ij} was applied to reveal the uncertainty of small values of n_{ij} (Polissar et al., 1999). The W_{ij} could be calculated using the Eq. (6) as follows:

$$W_{ij} = \begin{cases} 1.00 & 80 < n_{ij} \\ 0.70 & 20 < n_{ij} \leq 80 \\ 0.42 & 10 < n_{ij} \leq 20 \\ 0.05 & n_{ij} \leq 10 \end{cases} \quad (6)$$

Therefore, the $PSCF_{ij}$ can be calculated using the Eq. (7) as follows:

$$PSCF_{ij} = \frac{m_{ij}}{n_{ij}} \times W_{ij} = \frac{m_{ij}}{n_{ij}} \times \begin{cases} 1.00 & 80 < n_{ij} \\ 0.70 & 20 < n_{ij} \leq 80 \\ 0.42 & 10 < n_{ij} \leq 20 \\ 0.05 & n_{ij} \leq 10 \end{cases} \quad (7)$$

195 2.6 Ozone formation potential (OFP)

The photochemical activity of VOCs was normally assessed by the O₃ formation potential (OFP) (Niu et al., 2016). The value of OFP was affected by many factors like meteorological conditions, VOC concentrations and VOC sources. The OFP was calculated by Eq. (8):

$$OFP_i = [conc]_i \times MIR_i \quad (8)$$

200 where OFP_i is the ozone formation potential of the i species, $[conc]_i$ is the concentration of VOC species i , MIR_i is the maximum incremental reactions of the i VOC species, as reported by Carter (1994).

2.7 Secondary organic aerosol formation potential (SOAFP)

205 As discussed by Grosjean and Seinfeld (1989), SOAFP could be used to evaluate quantitatively the VOC influences on the secondary aerosol formation based on a variety of assumed interactions between VOCs and OH under the sunlight irradiation (8:00-17:00).

$$SOAFP_i = [VOC_i] \times FAC_i \quad (9)$$

where VOC_i is the concentration of the i VOC species, FAC_i is fraction aerosol coefficient of the i compound. The FAC_i was obtained from the previous studies (Grosjean and Seinfeld, 1989; Zhu et al., 2017; Mozaffar et al., 2020).

2.8 Sensitivity analysis

210 In order to further understand the relationship between VOCs and PM_{2.5}, a gradient model was applied to investigate the sensitivity of PM_{2.5} pollution to the VOC concentration. The Eq. (10) was below:

$$\eta = \frac{\Delta_{VOCs}}{\Delta_{PM_{2.5}}} \quad (10)$$

215 where Δ_{VOCs} and Δ_{O_3} is the concentrations of VOCs and PM_{2.5} in the specific PM_{2.5} gradients, respectively. The characteristic structure and reactivity could influence the contribution of VOCs to PM_{2.5} formation (Cater, 1994). To determine the quantitative relationship between the VOCs and PM_{2.5}, the VOCs-sensitivity coefficient (VOCs-S_{PM_{2.5}}) was used as the Eq. (11):

$$VOCs-S_{PM_{2.5}} = \frac{\Delta_{VOCs}/B_{VOCs}}{\Delta_{PM_{2.5}}/B_{PM_{2.5}}} \quad (11)$$

220 where B_{VOCs} and $B_{PM_{2.5}}$ are the background concentrations of VOCs and PM_{2.5}. The PM_{2.5} concentrations below 20 $\mu\text{g m}^{-3}$ were averaged to be the background level (Han et al., 2017). The background concentration of VOCs (B_{VOCs}) was the corresponding VOC concentrations. The high value of VOCs-S_{PM_{2.5}} implied that the concentration of VOCs was greatly affected by the variations of PM_{2.5} concentration. In order to quantify the sensitivity of VOCs to PM_{2.5}, VOCs and PM_{2.5} were classified into different groups with a PM_{2.5} concentration interval of 5 $\mu\text{g m}^{-3}$. It could be calculated by the Eqs. (12-15):

$$y = a \cdot x^b \quad (12)$$

225 $\ln y = \ln a + b \cdot \ln x \quad (13)$

$$\ln VOCs - S_{PM_{2.5}} = b \cdot \ln a + b \cdot \ln \frac{\Delta_{PM_{2.5}}}{B_{PM_{2.5}}} \quad (14)$$

$$\ln \frac{\Delta_{VOCs}}{B_{VOCs}} = k \cdot \ln \frac{\Delta_{PM_{2.5}}}{B_{PM_{2.5}}} + c \quad (15)$$

where k represents the linear coefficient between $\ln(\Delta_{VOCs}/B_{VOCs})$ and $\ln(\Delta_{PM_{2.5}}/B_{PM_{2.5}})$, c is the intercept. This method was appropriate for understanding the sensitivity of VOC pollution level to PM_{2.5} at polluted environments.

230 3. Results

3.1 Characteristics of air pollutants

3.1.1 Data overview

Figure 1 shows the variations of meteorological factors, VOC categories, PM_{2.5} and O₃ at the sampling sites. The average temperatures were 8.69 ± 3.24 , 9.02 ± 3.24 and 7.73 ± 2.92 °C, and the mean RHs were 83.77 ± 11.38 , 75.37 ± 13.29 and 71.80 ± 9.28 % at the JS, PD and QP sites, respectively. The wind speed at the QP site (4.37 ± 1.47 m s⁻¹) was 2.29 and 1.36 times higher than those at the JS (1.91 ± 0.49 m s⁻¹) and PD (1.30 ± 0.62 m s⁻¹) sites, respectively, indicating the decreased dilution and diffusion conditions at the latter two sites. Besides, the land-use types of the JS (industrial district) and PD (residential and commercial mixed districts) sites could lead to high anthropogenic emission. These phenomena result in the fact that the VOC concentrations at the JS (21.88 ± 12.58 ppb) and PD (21.36 ± 8.58 ppb) sites were higher than that at the QP site (11.93 ± 6.33 ppb). Compared with the relevant measurements performed previously in Shanghai at the same sampling sites, this study generally presented lower VOC concentrations (Cai et al., 2010b; Zhang et al., 2018; Zhang et al., 2020a). In detail, at the JS site, the VOC concentration was approximately 4 times lower than the measurement of Zhang et al., (2018) (94.14 ppb). At the PD and QP sites, the results in this study were slightly lower than those reported by Cai et al. (2010b) (24.3 ppb) and Zhang et al. (2020a) (15.41 ppb). A variety of control strategies, such as prohibiting of fireworks in the open air, improving VOC detection standards and strengthening control technology were implemented, thus resulting in the low VOC concentrations herein. Particularly, the policy of “one factory, one strategy”, targeted at mitigating VOC emissions, was published by Shanghai government in 2018.

During the observation period, the average PM_{2.5} values were 45.57 ± 27.59 , 48.51 ± 27.22 and 40.27 ± 27.78 µg m⁻³, and the mean O₃ concentrations were averaged to be 73.59 ± 23.59 , 57.48 ± 20.49 and 99.30 ± 24.00 µg m⁻³ at the JS, PD and QP sites, respectively. VOCs was found to be positively correlated with PM_{2.5}, and the pearson correlation coefficients ($R_{Pearson}$) were 0.58, 0.71 and 0.25 at the JS, PD and QP sites, respectively. It was well documented that the elevated VOC concentrations led to the increasing rate of PM_{2.5} production *via* photochemical oxidation, gas-particle partition and/or heterogeneous absorption (Seinfeld et al., 2001; Yang et al., 2015; Han et al., 2017). Moreover, PM_{2.5} and VOCs present similar emission sources. For example, traffic exhaust was proven as the predominant contributor for both of them (Li et al., 2009; Cai et al., 2010a; Cai et al., 2010b; Wang et al., 2013; Kuo et al., 2014; Liu et al., 2019). However, the VOC concentrations were negatively correlated with O₃ ($R_{Pearson} = -0.24$ at the JS site, $R_{Pearson} = -0.48$ at the PD site and $R_{Pearson} = -0.25$ at the QP site, respectively). The termination and titration ($\text{NO} + \text{O}_3 \rightarrow \text{NO}_2 + \text{O}_2$) were more efficient and lots of factors including sunshine duration, temperature and relative humidity rather than the emission of precursors, impacted on the surface O₃. Li et al. (2019b) emphasized that the absolute concentration of precursor was not the only factor during the O₃ formation in Zhengzhou, China.

3.1.2 Chemical compositions

The VOC compositions at the JS, PD and QP sites are presented in Fig. 2. During the observation campaign, 43 VOC species including 16 alkanes, 11 alkenes, 16 aromatics and 1 alkyne were measured and the contributions of total VOCs (TVOCs) > 1 % were marked. The most abundant specie was alkanes, followed by aromatics at the JS, PD and QP sites. The contributions of alkenes and alkyne were comparable at the above sites. Specially, ethane, propane, iso-butane, n-butane,

ethylene, benzene, toluene and ethyne were the abundant individual species and they totally contributed to greater than 75 % of the TVOCs.

Among four major organic classes, the fractions of alkanes were 49.32, 71.48 and 60.88 % at the JS, PD and QP sites, respectively, in line with the previous result (59.36 %) (Zhang et al., 2020a). Such high fraction of alkanes can be attributed to their widespread emission sources and long atmospheric lifetimes. In terms of VOC species, the ethane, propane and C₄-C₆ branches alkanes contributed greatly to the alkanes at the sampling sites, in accordance with the findings of Zhang et al. (2020a) and Song et al. (2021).

The second-largest group was aromatics accounting for 21.00, 10.70 and 13.75 % at the JS, PD and QP sites, respectively. As reported by Cai et al. (2010b), the aromatics at the Xujiahui site in Shanghai, China accounted for 24.9 % of the TVOCs which was slightly higher than the results obtained in this study. It is interesting to note that the dominating compounds of aromatics were toluene, m/p-xylene, benzene, ethylbenzene and o-xylene (BTEX), in line with the results at some Chinese cities such as Shanghai (Zhang et al., 2018) and Xi'an (Song et al., 2021).

The alkenes accounted for 15.30, 10.13 and 10.57 % at the JS, PD and QP sites, respectively, and the discrepancies of alkenes contributions to TVOCs among three sites were narrow. The ethene, propylene and butene were predominant compounds in the composition of alkenes, and their contributions were comparable to the results of Shao et al. (2016). The alkyne contributed 14.38, 7.69 and 14.80 % on the TVOCs at the above sites. Relative to the result reported by Liu et al. (2017) in Hangzhou, China, the contribution of alkyne was comparable to the present study, accounting for 16.6 % of the TVOCs.

3.1.3 “Weekend effects”

The “weekend effects” of VOCs at the JS, PD and QP sites are shown in Fig. 3. On the weekdays, the average VOC concentrations were 21.03 ± 2.66 , 22.00 ± 4.88 and 11.09 ± 1.58 ppb at the JS, PD and QP sites, respectively. The morning peak values were observed during the rush-hour traffic at 8:00 to 10:00 LT, reaching 26.61, 35.11 and 13.54 ppb, respectively due to the morning traffic. The peak was more significant at the PD site than those at the JS and QP sites, illustrating that the significant impact of land-use type, leading to the high vehicle emission at the PD site. During the rush-hour traffic at 18:00 to 21:00 LT, the VOC concentrations also tended to increase, and the evening peak values were 18.46, 20.82 and 10.22 ppb at the JS, PD and QP sites, respectively. Such scenario could be attributed to the strong influence of the vehicle exhaust. The VOC concentrations dropped to the minimum at the 16:00 LT with the values of 16.96, 15.62 and 8.77 ppb at the JS, PD and QP sites, respectively. The strong radiation, high temperature and boundary layer resulted in the most intense air convection condition and promoted the photochemical loss and dilution of VOCs, leading to the decreased VOC concentration (Zhang et al., 2018). On the weekends, the JS site exhibited the highest average VOC concentration of 20.36 ± 2.23 ppb, followed by the PD and QP sites with the concentrations of 19.96 ± 2.37 and 10.96 ± 0.67 ppb, respectively. The VOC concentrations on the weekends were 3.31, 10.19 and 1.19 % lower than those on the weekdays at the JS, PD and QP sites, respectively. Similarly, the elevated VOC concentrations on the weekdays were found in the

other studies including Shanghai, China (Geng et al., 2008), Sacramento, California (Murphy et al., 2007) and Los Angeles, California (Nussbaumer and Cohen, 2020), due to the low anthropogenic activities occurring on the weekends, reflecting the low VOC emission (Murphy et al., 2007). The minimum VOC concentrations on the weekends appeared on the 16:00 LT at the JS and PD sites and 15:00 LT at the QP site, respectively, with the values of 17.14, 17.56 and 10.33 ppb which occurred around the similar time as those on the weekdays.

3.1.4 Discrepancy of clean and haze day

Referring to the previous documents (Li et al., 2017; Hui et al., 2019), haze pollution was defined as the condition with visibility < 10 km and RH > 80 %. During the observation period, there were three sequential haze pollution events at the three sites on the basis of this criteria, i.e., 18 to 25 January, 23 February to 3 March and 21 to 26 March. The VOC concentrations between the clean and haze days are shown in Fig. 4. The VOC concentrations on the clean days were averaged to be 20.53 ± 12.10 , 19.29 ± 7.60 and 11.04 ± 6.67 ppb at the JS, PD and QP sites, respectively. For the haze days, the VOC concentrations increased by 27.15, 32.85 and 22.42 % at the above sites. Wu et al. (2016) also emphasized the increased VOC concentrations during the haze days relative to clean days in Beijing, China. Obviously, the ‘haze-clean’ differences of VOC concentrations were more significant at the JS and PD sites compared with the QP site. Such scenario could be attributed to the locations of JS and PD sites which implicated stagnant weather conditions and high anthropogenic emissions, therefore inducing the severe haze pollution. In detail, the ‘clean-haze’ discrepancy was dominated by the aromatics (*m*-ethyltoluene, *p*-ethyltoluene, *m*-ethylbenzene, 1, 2, 4-trimethylbenzene) at the JS and PD sites, as reflected by the 44.54 and 36.05 % higher concentrations on haze days relative to clean days. The elevated concentrations reflected the concentrated emission sources of industrial production, painting/coating and vehicle exhaust. Zhang et al. (2021) studied the VOC characteristics during the haze pollution in Zhengzhou, China, and showed that aromatics present an upward tendency (~ 34.04 % uplift) on the haze days, which was slightly lower than the results shown herein. At the QP site, alkanes (2, 2, 4-trimethylpentane, *n*-hexane, *n*-heptane) presented significant ‘clean-haze’ discrepancy (~ 36.58 % uplift), implying the great impacts of vehicle exhaust and fuel evaporation. Hui et al. (2018) also observed the elevated concentration of alkanes during the haze days in Wuhan, China, and found that there were 37.28 % increment of alkanes, which was comparable to the result in the present study. Thus, it could be deduced that, the haze occurrence is mainly associated with vehicle exhaust, industrial production and painting/coating at the JS and PD sites, whereas the vehicle exhaust and fuel evaporation at the QP site. Such deduction on the source of haze events agreed well with the studies about the haze pollution in Shanghai, China (Li et al., 2019b; Wei et al., 2019).

3.2 Source apportionment

3.2.1 Special VOC ratio analysis

Different VOC species have different sources, and the ratio of different species can be used to preliminary distinguish the

330 emission sources (An et al., 2014). Herein, the characteristics of toluene/benzene (T/B), *iso*-pentane/*n*-pentane (P/P) and *m*,
p-xylene/ethylbenzene (X/E) varying at the sampling sites are utilized to identify the VOC sources. The results are shown in
Fig. 5 and Fig. S2-4.

It is well documented that the varied VOC sources could be identified by the different T/B ratios (Song et al., 2021). The
mean T/B ratios were 4.59 ± 4.3 ($R_{Pearson} = 0.41$) and 1.61 ± 0.79 ($R_{Pearson} = 0.65$), respectively at the JS and PD sites. The
335 most of the ratios (68.89 and 84.15 %) distributed the range of 0.9-2.2 and 1.4-5.8, suggesting that vehicle emissions and
industrial emissions exerted the significant impact on VOC concentrations (Fig. 5a-b), which was well agreement with the
previous studies (Dai et al., 2013; Zhang et al., 2013; Mo et al., 2015; Shi et al., 2015; Yao et al., 2015; Deng et al., 2018).
The average ratio at the QP site was 1.01 ± 0.66 ($R_{Pearson} = 0.70$) with 43.02 % of T/B ratios distributing the range of 0.9-2.2
and 0.2-0.4, suggesting that vehicle emissions and burning emissions contributed significantly to VOC pollution (Fig. 5c) (Li
340 et al., 2011; Dai et al., 2013; Zhang et al., 2013; Yao et al., 2015; Mo et al., 2016; Deng et al., 2018). The temporal variations
of T/B ratio and TVOC concentrations are shown in Fig. S2. The T/B ratios were mainly located in the scopes of vehicle
emission and industrial source at the JS and PD sites when the VOC concentrations were high, respectively, while the
specific values at the QP site were only concentrated in the scope of vehicle emission. Such scenario showed that vehicle
exhaust was a significant contributor to the VOC pollution at the sites we studied, and VOC concentrations were less
345 influenced by the industrial emission at the QP site relative to the JS and PD sites, due to the land-use type of QP site where
is normally taken as a city background site and far from the industrial area.

The fossil fuel-derived sources (vehicle exhaust, fuel evaporation and coal combustion) could be further distinguished by the
P/P ratio. *Iso*-pentane and *n*-pentane have similar atmospheric lifetimes, therefore varied with a similar trend and were highly
correlated ($R_{Pearson} = 0.91, 0.91$ and 0.81 at the JS, PD and QP sites, respectively), indicating the similar emission sources of
350 two compounds (Jobson et al., 1998; Yan et al., 2017). Lower P/P ratios (0.56-0.80) are often identified for coal combustion
(Li et al., 2019c), and the P/P ratios in the range of 2.2-3.8 are characterized by vehicle emission (Liu et al., 2008; Wang et
al., 2013). The average P/P ratios between the *iso*-pentane and *n*-pentane were $0.96 \pm 0.30, 1.36 \pm 0.22$ and 2.46 ± 1.49 at the
JS, PD and QP sites, respectively (Fig. 5d). The result at the QP site was comparable with that measured at a Pearl River
tunnel (2.93), suggesting that the vehicle emission was an important VOC source (Liu et al., 2008). The lower ratios at the JS
355 and PD sites indicated that VOC concentrations were affected by coal combustion sources. The temporal variations of P/P
ratio and TVOC concentrations are shown in Fig. S3. The P/P ratios were distributed within the range of 2.2-3.8 at the
sampling sites during the VOC pollution, indicating the great impact of vehicle emission on VOC pollution, which was in
agreement with the report of Song et al. (2021).

Besides the local emissions influencing the VOC pollution characteristics, regional transport has been probed as a potential
360 VOC source. Herein, the ratio of X/E was used to evaluate the transport impacts. *M*, *p*-xylene and ethylbenzene are found to
be similar in emission sources, while the former exhibits ~ 3 times greater reactivity toward OH radical than the latter
(Nelson and Quigley, 1983; Chang et al., 2006; Vardoulakis et al., 2011). Hence, lower X/E ratios normally suggested more
significant air mass aging, that is, more influences from external transport. The X/E ratios were averaged to be 2.33 ± 0.37

($R_{Pearson} = 0.98$), 2.18 ± 0.42 ($R_{Pearson} = 0.97$) and 2.03 ± 1.52 ($R_{Pearson} = 0.74$) at the JS, PD and QP sites, respectively (Fig. 365 5e). The results showed that the X/E ratios were slightly lower at the QP site relative to the JS and PD sites, indicating that there were less primary emissions from anthropogenic sources at the QP site. The temporal variations of the X/E ratio and TVOC concentrations are shown in Fig. S4. During the VOC pollution, the X/E ratios were approximately 2.3, 2.5 and 1.8 at the JS, PD and QP sites, respectively. Obviously, the X/E ratio was lower at the QP site than those at the JS and PD sites. In particular, the high VOC concentration (27.17 ppb) was observed in 7 February at the QP site, corresponding to the 370 minimum X/E ratio (0.27), while the X/E ratios were 3.07 and 1.95 at the JS and PD sites, corresponding to the VOC concentrations of 12.45 and 22.45 ppb. Such results illustrated that the impact of external transport during the VOC pollution was greater at the QP site compared with the JS and PD sites.

3.2.2 The PMF analysis

The PMF analysis can quantitatively determine the VOC source contributions (Hui et al., 2019). In this study, 39 VOC 375 species were put into the PMF model, followed by the output of seven resolved factors including vehicle exhaust, industrial source, LPG usage, paint solvent, fuel evaporation, coal combustion and biogenic source. The source profiles of the resolved factors by the PMF model are shown in Fig. 6.

Vehicle exhaust was characterized by the high proportions of alkanes, some alkenes and certain percentage of aromatics (benzene, xylene and trimethylbenzene) (Liu et al., 2008; Cai et al., 2010a; Ling et al., 2011; An et al., 2017; Hui et al., 380 2018). In this study, the high contributions of the ethane (69.59 and 48.01 %), *iso*-butane (35.14 and 26.12 %) and some alkenes especially the propylene (34.72 and 19.74 %) and *trans*-2-butene (50.49 and 17.40 %) at the JS and PD sites and the ethane (42.41 %), ethylene (49.14 %) and ethyne (45.94 %) at the QP site were observed. These compounds are widely regarded as vehicular emission tracers (Cai et al., 2010a; Ling et al., 2011; An et al., 2017; Hui et al., 2018). The contributions of vehicle exhaust were calculated to be 23.18, 33.37 and 32.12 % at the JS, PD and QP sites, respectively. 385 The relevant high results indicate that vehicle exhaust was the predominant source of VOCs, in accordance with the results of T/B and P/P ratios mentioned above. Shanghai owns the motor vehicles more than 4.4 million in 2019. Vehicle exhausts had been proved to be one of the main causes of the local air pollution in Shanghai (Huang et al., 2015; Dai et al., 2017; Liu et al., 2019; Cai et al., 2010b).

Industrial emission was featured by the high percentage of aromatics (benzene, toluene and trimethylbenzene) and certain 390 percentages of alkanes and alkenes (Guo et al., 2011; Dumanoglu et al., 2014; Sun et al., 2016). At the JS and PD sites, the high contributions of *n*-nonane (17.84 and 33.99 %), benzene (13.89 and 39.16 %), toluene (21.11 and 19.54 %), trimethylbenzene (94.18 and 84.05 %), hexane (14.81 and 38.96 %) and ethylene (14.99 and 36.91 %) were observed. It was widely acknowledged that asphalt application could release *n*-nonane (Brown et al., 2007; Liu et al., 2008). Moreover, benzene and toluene are primarily applied for industrial solvent production, and trimethylbenzene is frequently used in 395 manufacturing (Morrow, 1990; Ling et al., 2011). Therefore, the recognized factor can be identified as industrial source.

Industrial events were calculated to contribute 22.39 % and 18.81 % to VOCs at the JS and PD sites, respectively, indicating the considerable contributions of industrialization next to the aforementioned traffic factors. The QP site was far away from the industrial regions, and therefore was less influenced by industrial factors compared with JS and PD sites.

Coal combustion factor was characterized by propane, ethylene, ethyne, and benzene (Liu et al., 2008; Ling et al., 2011; Song et al., 2018). Moreover, the proportions of propene, *n*-hexane, *n*-heptane and toluene were low, but these compounds could not be neglected yet (Hui et al., 2018). The PMF analysis illustrated that coal combustion was the third-largest source of VOCs at the three sites. Coal combustion was responsible for 14.95 % and 13.85 % of the VOC emission at the PD and QP sites, respectively. However, the contribution reached up to 21.48 % at the JS site, reflecting more frequent fossil usage in the industrial area of Shanghai.

LPG usage was determined by some alkanes (propane, *n*-butane and iso-butane) and alkenes (ethylene, propylene and butene). There were high proportions of propane (21.24 and 14.41 %), *n*-butane (15.53 and 13.79 %), *iso*-butane (12.61 and 11.40 %), ethylene (31.18 and 10.08 %) and propylene (34.94 and 31.38 %) observed at the JS and PD sites, coupled with *iso*-butane (43.88 %), propylene (25.08 %) and *cis/trans*-2-butene (53.10 and 57.65 %) at the QP site, highlighting the LPG-related sources (Yang et al., 2013; Lyu et al., 2016). Herein, LPG usage accounted for 13.17, 14.76 and 12.14 % of the VOCs monitored at the JS, PD and QP sites, respectively. At the end of 2015, the amount of LPG users about household and catering achieved 3.3-6.5 million in Shanghai (Hui et al., 2018; Zhang et al., 2018a). Overall, LPG usage was one of the important VOC sources at the sampling sites.

Fuel evaporation could be identified by C₃-C₇ alkanes, especially *n*-pentane and *iso*-pentane (Zheng et al., 2020), and the C₃-C₅ alkenes such as *trans/cis*-2-butene (Geng et al., 2009; Hui et al., 2018; Zhang et al., 2018a; Zheng et al., 2020). There were high contributions of *n*-pentane (37.94, 25.20 and 71.95 %), *iso*-pentane (41.45, 28.18 and 23.16 %) and butene (26.73, 70.83 and 22.52 %) at the JS, PD and QP sites, respectively. It was well documented that alkanes like *n*-pentane and *iso*-pentane were gasoline tracers, and some low-carbon alkanes could evaporate from the unburned fuels (Guo et al., 2004; Wang et al., 2013). The VOC contributions from fuel evaporation were calculated to be 4.62, 10.35 and 20.15 % at the JS, PD and QP sites, respectively.

Paint solvent usage was normally characterized by C₆-C₈ alkanes and some aromatics like toluene, ethylbenzene and *x/p*-xylene (Wang et al., 2013; An et al., 2017; Hui et al., 2018; Song et al., 2019a). There were high contributions of toluene at the JS site (65.90 %), toluene and ethylbenzene at the PD site (47.95 and 33.85 %) and *x/p*-xylene at the QP site (24.69 %), respectively. Herein, emission factors that meet these characteristics were identified as paint solvent usages, which accounted for 15.15, 7.77 and 10.36 %, respectively at the JS, PD and QP sites.

Biogenic source was distinguished by the great contribution of isoprene (Liu et al., 2008; Wu et al., 2016). Factors that coincide to the specific characteristic were regarded as biogenic source in this study. Our analysis showed that the contribution of biogenic source to VOC concentrations was 11.39 % at the QP site, reflecting the vegetation surrounding this site. However, this emission factor could not be reproduced at the JS and PD sites, implying the limited impacts of biogenic sources in the population- and industrialization-concentrated areas that were primarily controlled by the anthropogenic

3.2.3 Back trajectories and PSCF results

Besides the direct impact of local source, regional transport could also contribute to VOC pollution (Hui et al., 2018). The 24-h backward trajectories, the PSCF results and the VOC proportion of each trajectory throughout the observation at the JS, PD and QP sites are shown in Fig. 7. There were 6 clusters (northern trajectories: Cluster 1 + 2; eastern trajectories: Cluster 3; southern trajectories: Cluster 4; southwestern trajectories: Cluster 5; northwestern trajectories: Cluster 6) at the JS site, as shown in Fig. 7a. The contributions of northern trajectories to the total trajectories (Fig. 7a left) and that of the northern pollution trajectories to the total pollution trajectories (Fig. 7a right) were significantly higher than those of the other four cluster trajectories, comprising 45.74 and 46.46 %, respectively (Tab. 1). Identified as the north direction cluster, the north long-distance trajectory (Cluster 1) and short-distance trajectory (Cluster 2) accounted for 16.67 and 29.07 % of the VOC transportation. The results indicated that the VOC concentrations at the JS site was mainly impacted by the trajectories from the junction of the Bohai Sea, Shandong Province, the East China Sea and Jiangsu Province in addition to the local emission. The contributions of VOCs by each air mass trajectories indicated that alkanes tended to dominate the transported VOC community at the JS site due to its weak reactivity in the long-range transport.

The air mass trajectories could be cluster into northern trajectories (Cluster 1 + 2), northeastern trajectories (Cluster 3), southeastern trajectories (Cluster 4), southwestern trajectories (Cluster 5) and northwestern trajectories (Cluster 6) at the PD site, as shown in Fig. 7b. It is evident that the contributions of northern (Cluster 1) and northeastern (Cluster 3) trajectories to the total trajectories (Fig. 7b left) and that of the northern and northeastern pollution trajectories to the total pollution trajectories (Fig. 7b right) were greatly higher than other four cluster trajectories, accounting for 52.23 and 54.72 % of VOC concentrations, respectively. The results suggested that VOCs at the PD site was significantly influenced by the pollution transportation from East China Sea, Zhejiang Province and Jiangsu Province, coupled with the local sources. The proportions of VOCs by each air mass trajectories indicated that alkanes and alkenes both contributed to the transported VOC community at the PD site.

At the QP site (Fig. 7c), the trajectories were clustered into the northern trajectories (Cluster 1 + 2), northeastern trajectories (Cluster 3), southeastern trajectories (Cluster 4), southwestern trajectories (Cluster 5) and northwestern trajectories (Cluster 6). The proportion of northeastern trajectories to the total trajectories (Fig. 7c left) and that of northeastern pollution trajectories to the total pollution trajectories (Fig. 7c right) were higher than those of the other trajectories, accounting for 31.87 and 27.81 % of the VOC concentrations, respectively. This result indicated that the concentration of VOCs was mainly impacted by the northeastern trajectory from the junction of East China Sea, Zhejiang Province and Jiangsu Province in addition to local sources. Although the contribution of Cluster 2 was relatively small, the VOC concentration was the highest, reaching 23.63 ± 10.75 ppb (Tab. 1). Therefore, attention should be paid to the long-distance transmission of highly polluting from Beijing, Tianjin and Liaoning Province. The alkanes contributed most of VOCs in air mass trajectories at the QP site, which was in line with the results of the JS site.

The region with high PSCF levels indicates highly potential regional transport sources (Hui et al., 2018). Based on the PSCF results, at the JS and QP sites, high values were observed in the north of Shanghai. At the PD site, high values were observed in the northeast of Shanghai. All of the three sampling sites presented that the highest PSCF levels appeared in areas near the JS, PD and QP sites, suggesting that local source was a significant contributor to the VOC pollution.

3.3 Ozone and SOA formation potentials

3.3.1 Ozone formation potential

The atmospheric VOCs was considered as the important precursor of O₃ due to their photochemical reactivity (Alghamdi et al., 2014; Kumar et al., 2018). Fig. 8 displays the OFP contribution proportions of the primary VOC categories and species varying with sampling sites. Hereby, the OFPs were calculated to be 50.89 ± 2.63 , 33.94 ± 1.52 and 24.26 ± 1.43 ppb for the VOCs measured at the JS, PD and QP sites, respectively. The photo-induced transformation of VOCs could account for 69.15, 59.05 and 24.43 % of the O₃ concentrations in the above sampling sites. The VOC-related O₃ contributions were lower than those in Delhi, India (Kumar et al., 2018), and Chinese cities like Shanghai (Zhang et al., 2018a), Wuhan (Hui et al., 2018), Taiwan (Vo et al., 2018), Nanjing (Mozaffar et al., 2020) and Xi'an (Song et al., 2021). The relatively low OFP values were related with the low VOC concentrations in this study (Tab. S3). The predominate OFP contributor was alkenes especially ethylene and propylene, which exhibited the considerable photochemical reactivity, accounted for 50.21, 49.39 and 60.00 % of the OFPs at the JS, PD and QP sites, respectively (Li et al., 2019b). Such levels were generally comparable with the result obtained in Shanghai, China by Zhang and co-workers (2018), comprising 57 %. Moreover, the high photochemical reactivity of aromatics (Li et al., 2019b), especially the toluene, *o*-xylene and *m/p*-xylene, makes them become the second largest contributor of OFP, in accordance with the finding of Hui et al. (2018). The fraction of aromatics therein accounted for 33.59, 23.21 and 20.02 % of the OFPs at the JS, PD and QP sites, respectively. In comparison, the contributions of alkanes and alkyne were much lower, accounting for 13.11 and 3.09 % at the JS site, 22.90 and 4.50 % at the PD site and 13.07 and 6.91 % at the QP site, respectively. These results indicated that the VOC photochemical reactivity was closely related to the contribution of O₃ formation which was in agreement with other studies (An et al., 2014; Guo et al., 2017; Hui et al., 2018). Hence, alkenes and aromatics played crucial roles in the formation of O₃. According to the source identification results discussed above, the relevant emission sources, which are thought to be the industrial production and vehicle exhaust at the sampling sites, should be controlled in priority.

3.3.2 SOA formation potential

The close linkages between VOCs and new particle formation induced the quantitative discussion on SOAFP (Chen et al., 2007; Guo et al., 2017; Tan et al., 2018), and the Grosjean's methodology has been widely recommended as the standard reference (Fig. 9). Following the reported methods, the SOAFP was calculated only for daytime (8:00-17:00 LT) to avoid the emission from the human activities during nighttime (Grosjean and Seinfeld, 1989; Zhang et al., 2020b). During the

495 observation campaign, the SOAFP was averaged to be 1.00 ± 2.03 , 0.46 ± 0.88 and $0.41 \pm 0.58 \mu\text{g m}^{-3}$ for the VOCs
measured at the JS, PD and QP sites, respectively. Such levels were comparable with those obtained in Jinan, China (Zhang
et al., 2017) and Taiwan (Vo et al., 2018), China, while higher than that of Nanjing, China (Mozaffar et al., 2020) (Tab. S3).
The SOAFP values accounted for 2.19, 0.95 and 1.02 % of the $\text{PM}_{2.5}$ concentrations at the above sampling sites, all of which
were lower than the results in Nanjing, China (3.46 %) (Mozaffar et al., 2020) and Wangdu, China (8.4 and 17.84 % under
high- NO_x and low- NO_x conditions, respectively) (Zhang et al., 2020b). It is worthwhile to mention that, there were some
500 uncertainties leading to the conservative estimation of the SOAFP results, as discussed below. First, only 21 VOC
species were included in the current estimation, and the VOC concentrations during the daytime were only 20.80, 9.43 and
7.30 ppb at the JS, PD and QP sites, respectively. Second, the published FAC values were obtained by merely considering the
OH-related interactions (de Gouw et al., 2011; Hennigan et al., 2011; Spracklen et al., 2011; Zhang et al., 2020b). Third, the
discrepancy is additionally dependent on the different atmospheric conditions and study area. In the current study, aromatics
505 was determined to be the main SOA contributor, accounting for 86.07, 96.21 and 86.38 % at the JS, PD and QP sites,
respectively, in accordance with the findings in Jinan, China (Zhang et al., 2017). The top five abundant VOC species were
toluene, *x/p*-xylene, ethylbenzene, benzene and *o*-xylene at the above sites. Toluene was determined to be the most abundant
VOC specie for SOA formation, with the SOAFP values of 9.31, 3.38 and $2.06 \mu\text{g m}^{-3}$ at the JS, PD and QP sites,
respectively, in agreement with the results of Mozaffar et al. (2020). Obviously, the value at the JS site was significantly
510 higher than those at the PD and QP sites, due to a large number of chemical factories surrounding the JS site, resulting in the
high concentration of VOCs especially the aromatics (Hui et al., 2018). Therefore, to achieve the better SOA reduction
effects, the concentration of aromatics should be controlled in priority in Shanghai, and more efficient strategies should be
developed to limit the emissions of industrial emissions.

4. Discussion

515 In this section, the effects of land-use type on VOC concentrations, sources and ozone and SOA formation potentials are
discussed. Previous studies pointed out that the VOC characteristics (e.g., concentrations and diurnal/weekly variations) and
sources varied with the land-use types (Yoo et al., 2015; Jookjantra et al., 2022). Further, the discrepancies of VOC
characteristics among the land-use type could affect the ozone and SOA formations (Cai et al., 2010a; Cai et al., 2010b;
Jookjantra et al., 2022). Herein, the JS site is located in the Second Jinshan Industrial Area of Shanghai as the industrial
520 district, and surrounded by many chemical factories. The PD site is located in the Pudong New Area as the residential and
commercial mixed districts, and surrounded by residences and administrative areas. The QP site is located near the southeast
of Dianshan Lake as the background district, and surrounded by many farmlands and forests (Tab. S1). Since the distinct
land-use types among the sampling sites were observed, it is highly desirable to discuss the effects of the land-use type on
concentrations, sources and O_3 and SOA formation potentials of VOCs in Shanghai.

525 4.1 The effect of land-use type on VOC concentrations

The mean VOC concentrations at the JS (21.88 ± 12.58 ppb) and PD (21.36 ± 8.58 ppb) sites were 1.83 and 1.79 times higher than that at the QP site (11.93 ± 6.33 ppb), implicating the impact of land-use type. In more detail, the JS and PD sites are located at the industrial and urban regions with numerous anthropogenic activities. The enhanced emission strengths at the JS and PD sites could induce severe atmospheric pollution e.g., haze and O₃ pollution (Cai et al., 2010a, b; Zhang et al., 2018). Further, the meteorological conditions including ambient temperature and sunlight irradiance changed, the dispersive dilution and convection phenomenon decreased, which contributed to high VOC concentrations (Kumar et al., 2018). By contrast, the emission strength dropped at the QP site as a consequence of the land-use type of this site (background district). Besides, the land-use types could not only affect the VOC concentrations but also compositions. Specifically, the JS site exhibited higher fractions of aromatics and alkenes, particularly toluene and propylene, than those at the PD and QP sites. This result did necessarily correlate with the fact that the JS site is close to the industrial area with heavy industrial emissions (the dominating source of toluene and propylene), as reflected by the increased toluene and propylene proportions (Zhang et al., 2018; Hui et al., 2019; Li et al., 2019b). However, the contribution of ethane at the JS site was ~ 50 % lower than those at the PD and QP sites. As is well known that ethane is typically tracer of incomplete combustion especially vehicle emission (Guo et al., 2011). It was thus supposed that the vehicle emission exerted a relatively weak influence on VOC emission at the JS site (23.18 %), as compared to the PD (33.37 %) and QP sites (32.12 %). The fraction of alkanes especially the propane was remarkable in the land-use type with residential and commercial activities (the PD site). This site is the junction of roads, metros, commercial and financial areas that had a significant impact on the enrichment in vehicle source and hence could have participate in the high contribution of alkanes (Cai et al., 2010a; Ling et al., 2011; An et al., 2017; Hui et al., 2018). The fraction of alkyne at the PD site was 1.88 and 1.75 times lower than those at the JS and QP sites, which might be caused by the weak combustion sources including chemical- and bio-fuels burning (Zhu et al., 2016; Li et al., 2019b). Note that the distinct spatial heterogeneity of VOCs was also observed with the highest value of the coefficient of divergence (COD = 0.36) between the JS and QP sites, followed by the PD and QP sites (COD = 0.33), with that between the JS and PD sites (COD = 0.20) being the lowest. Hence, the spatial heterogeneity of VOCs between the JS and PD sites was narrow, while the QP site was largely different from other two sites. This result was ascribed to the fact that there were similar pollutant concentrations, meteorological factors, emission intensities and atmospheric conditions at the JS and PD sites, while these indexes in the QP site were rather different from those of the other sites.

The VOC diurnal variations were analyzed with respect to each of the land-use types. The result showed that the VOC concentration in the early morning (5:00 LT) at the JS site was 17.06 and 52.91 % higher than those at the other two sites, which was attributed mostly to the land-use types. There were a large number of rubber factories, paint solvent factories and oil factories surrounding the JS site, whereby some factories usually worked continuously for 24 hours, thus inducing the increased VOC concentrations. Compared with the other studies (Velasco et al., 2007; Zhang et al., 2018; Wang et al., 2021), no evident bimodal features can be observed at the sampling sites. The industrial processes and biogenic emission

surrounding the sampling sites also had great impact on the VOC variations except for the traffic exhaust, which was the potential reason for such scenario (Yoo et al., 2015). In addition to the diurnal variations, the “weekend effects” of VOCs also appeared to be variable among the different land-use types. Specifically, at the JS site, there were narrow discrepancies of VOC concentrations between the weekdays and weekends except for the rush hour in the morning, which was related to the influence of the full-day operation of industrial processes. At the PD site, the highest value during the morning peak on the weekends (26.03 ppb) was lower than that on the weekdays (35.11 ppb), decreased by ~ 25.86 %, suggesting that there were obvious anthropogenic sources especially the vehicle exhaust. At the QP site, the elevated VOC concentrations (~ 18.69 % uplift) were presented during the 18:00-20:00 LT on the weekends relative to the weekdays. The phenomenon might be associated with the joint effect of VOC intensive emissions by anthropogenic activities at the QP site on the weekends in that this site is located around the Dianshan Lake where is a tourist attraction.

4.2 The effect of land-use type on VOC sources

VOC sources in this study were sensitive to the local emission with the different land-use types. Specifically, the vehicle exhaust was responsible for nearly one third of the VOC emission and was a primary source of VOCs. It is interesting to note that the vehicle contribution was expected to be lower at the JS site than those at the PD and QP sites due to the density of automobiles per unit area. This finding reflected the fact that the JS site is located in the industrial regions favouring the contribution accumulations of industrial production and coal combustion (Yoo et al., 2015). As reported, vehicle emission was a dominating VOC source in other cities, and the contributions were higher than 20 % except in Calgary, Canada (17.2 %) (Fig. 10). Therefore, the effective and continuous control strategies on vehicle exhaust are still of priority to alleviate the VOC pollution. Apart from vehicle exhaust, the industrial emission was also an important VOC source at the JS and PD sites, the contributions of which herein were higher than those in other studies, except for Xi’an, China (29.7 %) and Paris, France (35 %). It is supported by locations of JS and PD sites i.e., the JS site is located at Second Jinshan Industrial Area of Shanghai as an industrial site, and Waigaoqiao Ship building LTD and BaoSteel Group Corporation are located within 30 km of the PD site, illustrating the great influence of industrial processes on VOC pollution at the both sites. However, limited influence of industrial source at the QP site was observed, which because the QP site is surrounded by natural environment and far away from the industrial regions. The discrepancies of combustion contribution also could be illustrated by the land-use types. In this case, the contribution of combustion was more pronounced at the JS site than the PD and QP sites. As mentioned above, the JS site is surrounded by the industrial cluster and hence induced high emission of industrial combustion (Zhang et al., 2018). Moreover, the contribution of combustion varied greatly among different studies, from 4 % (Song et al., 2019b) to 46 % (Bari and Kindzierski, 2018) (Fig. 10), which was partially ascribed to the different definition of combustion. For instance, Song et al. (2019b) only regarded the wood combustion as the combustion source in the study of Seoul, South Korea, while the sum of fuel combustion and oil/natural gas extraction/combustion was summarized as combustion source in the study of Calgary, Canada (Bari and Kindzierski, 2018).

590 The contribution of fuel evaporation showed the obvious discrepancies among the different site locations. In detail, the contribution at the JS site (4.62 %) was lower than those in the previous studies (6.28-20 %), while it was relatively average at the PD site (10.35 %) (Li et al., 2018; Hui et al., 2018; Liu et al., 2019; Song et al., 2019a, b; Song et al., 2021). At the QP site, the result (20.15 %) was comparable with those from the other studies (11.65-20 %) (Hui et al., 2018; Liu et al., 2019; Song et al., 2019b; Gaimoz et al., 2011) (Fig. 10). The proportions of LPG usage were approximately twice higher at the
595 sampling sites than those reported by the other studies (Hui et al., 2018; Song et al., 2019a). The fractions of paint solvent usage here were in the range of estimates by other cities (8.2-41 %) (Gaimoz et al., 2011; Bari and Kindzierski, 2018; Li et al., 2018; Liu et al., 2019; Hui et al., 2018; Song et al., 2019a, b; Liu et al., 2021; Song et al., 2021). The contribution of biogenic source at the QP site was relatively average compared to the estimates from other studies (4.9-15 %) (Cai et al., 2010b; Gaimoz et al., 2011; Bari and Kindzierski, 2018; Hui et al., 2018; Li et al., 2018; Song et al., 2019 a, b) (Fig. 10). It is
600 worth noting that VOC concentrations at the JS and PD sites were less influenced by the biogenic source relative to the QP site. Since the VOC concentrations were related to the emission strength and the mixing ratio of high intensity was determined by anthropogenic operation, it could be concluded that the human influences at the JS and PD sites were higher than that at the QP site which was consistent with the regional characteristics of anthropogenic activities controlled by land-use types. Overall, the results of source contribution estimates varied with different studies (Lyu et al., 2016; Li et al.,
605 2020b). The VOC sources also showed difference at a regional scale within the city (Tang et al., 2008; Li et al., 2019b; Cai et al., 2010; Song et al., 2021). These findings were particularly link to the land-use types, resulting in varying VOC species, sources, and emission strength (Li et al., 2019b; Cai et al., 2010; Li et al., 2020b; Song et al., 2021). Thus, the source apportionment for the different land-use types at a regional scale among various studies is necessary to identify the high-quality localized source.

610 **4.3 The effect of land-use type on ozone and SOA formation potentials**

The OFP values were closely related with the land-use types. In detail, the higher OFPs at the JS and PD sites (50.85 ± 2.63 and 33.94 ± 1.52 ppb) relative to that at the QP site (24.26 ± 1.43 ppb) were observed. Under the high OFPs, the concentrations of O_3 at the JS and PD sites (73.59 ± 23.59 and $57.48 \pm 20.49 \mu\text{g m}^{-3}$) were unexpectedly lower than that at the QP site ($99.30 \pm 24.00 \mu\text{g m}^{-3}$), indicating the poor O_3 formation conditions, since the estimated OFP was proceeded
615 under the optimum conditions. These results were due to the fact that the locations of JS and PD sites resulted in the high emission strength, which could release to high pollutant concentrations and lead to severe atmospheric pollution (Cai et al., 2010a, b; Zhang et al., 2018). This phenomenon could change the strength of solar radiation and further decreased the intensity of photochemical reactions (Kumar et al., 2018). By contrast, the favorable meteorological conditions especially the intensive solar radiation at the QP site surrounded by vegetation were observed which contributed to the high O_3
620 concentration (Yoo et al., 2015). Such results were in good agreement with the findings in Zhengzhou, China (Li et al., 2019b).

The discrepancies of SOAFP values among the sampling sites were observed since they could be significantly influenced by the land-use types. At the JS site, the SOAFP value ($1.00 \pm 2.03 \mu\text{g m}^{-3}$) was 2.17 and 2.44 times higher than those at the PD ($0.46 \pm 0.88 \mu\text{g m}^{-3}$) and QP ($0.41 \pm 0.58 \mu\text{g m}^{-3}$) sites, which was consistent with variations of VOC concentrations especially aromatics and connected with the land-use types (Zhang et al., 2017; Jookjantra et al., 2022). The JS site is surrounded by heavy industrial activities in paint and printing factory, manufacturing factory and rubber factory, resulting in the concentrated emission sources of industrial production (Zheng et al., 2010; An et al., 2014; Debevec et al., 2021). Because the close associations between VOCs and SOA could induce the sensitive response of VOC concentrations to the different pollution degree of $\text{PM}_{2.5}$, the VOCs- $\text{PM}_{2.5}$ sensitivity ($\text{VOCs-S}_{\text{PM}_{2.5}}$) was applied to understand the plausible influence on the atmospheric $\text{PM}_{2.5}$ abundance. Atmospheric pollution here can be divided into five levels by the mass concentration of $\text{PM}_{2.5}$: clean level ($\text{PM}_{2.5} < 35 \mu\text{g m}^{-3}$), slight pollution level ($35 < \text{PM}_{2.5} < 75 \mu\text{g m}^{-3}$), medium pollution level ($75 < \text{PM}_{2.5} < 120 \mu\text{g m}^{-3}$), heavy pollution level ($120 < \text{PM}_{2.5} < 180 \mu\text{g m}^{-3}$) and extreme pollution level ($\text{PM}_{2.5} > 180 \mu\text{g m}^{-3}$) (Han et al., 2017). The variations of $\text{VOCs-S}_{\text{PM}_{2.5}}$ as a function of $\text{PM}_{2.5}$ concentration at the JS, PD and QP sites are shown in Fig. 11. The low values of $\text{VOCs-S}_{\text{PM}_{2.5}}$ basically remain constant, while the high values displayed a descending trend. In the clean level, $\text{VOCs-S}_{\text{PM}_{2.5}}$ varied greatly from 0.19 to 2.50, 0.26 to 2.17 and 0.11 to 3.26 at the JS, PD and QP sites, respectively. In the pollution level, the scope of $\text{VOCs-S}_{\text{PM}_{2.5}}$ was narrow, especially in the extreme pollution level with the values fluctuating from 0.10 to 0.16, 0.13 to 0.17 and 0.17 to 0.28 at the JS, PD and QP sites, respectively. It is worth noting that the $\text{VOCs-S}_{\text{PM}_{2.5}}$ values displayed a steady decreasing trend, and the ratios of $\text{PM}_{2.5}$ to $B_{\text{PM}_{2.5}}$ increased in the clean and slight polluted levels. Such result demonstrated that the VOC concentrations were sensitive to that of $\text{PM}_{2.5}$ in the above two episodes. In order to quantify the sensitivity of VOCs to $\text{PM}_{2.5}$, we classified VOCs and $\text{PM}_{2.5}$ into different groups with a $\text{PM}_{2.5}$ concentration interval of $5 \mu\text{g m}^{-3}$, on the basis of the study reported by Han et al. (2017). The mean $\text{VOCs-S}_{\text{PM}_{2.5}}$ values followed an exponential function, and the values of k were determined to be 0.39, 0.46 and 0.56 at the JS, PD and QP sites, respectively, according to the Eqs. (12-15). The VOCs at the QP site could be more sensitive to $\text{PM}_{2.5}$ relative to other two sites because greater k normally indicated the rapider increment of VOC concentrations. The four groups of VOCs displayed similar linkages with VOCs, and the higher values of k were attributed to the aromatics at the JS and PD sites, while the alkanes at the QP site (Fig. S5). Thus, the optimal choices of controlling VOC species varied with the land-use types. Accordingly, in order to efficiently control the VOC-induced haze pollution, the concentrations of aromatics at the JS and PD sites, as well as alkanes at the QP site should be controlled in priority, as the results of the discrepancy of clean and haze day.

650 5. Conclusions

Herein, a concurrent atmospheric observation campaign was performed at the three supersites with the different land-use types of Shanghai from January to March 2019. Based on the observation data, this study carefully discussed the concentration variations, primary sources, ozone and SOA formation potentials of the atmospheric VOCs influenced by

land-use types.

655 The VOC concentrations in the order of the JS (21.88 ± 12.58 ppb) > PD (21.36 ± 8.58 ppb) > QP (11.93 ± 6.33 ppb) sites were locally sensitive to the land-use types. Alkanes dominated the VOC community and accounted for 50.33, 71.48 and 60.88 % of the TVOCs at the JS, PD and QP sites, respectively because of their widespread emission sources and longer atmospheric lifetime. The relatively high VOC concentrations during the rush traffic hours in the morning and evening were observed at the sampling sites due to the vehicle emission. The “weekend effects” of VOCs showed that the VOC
660 concentrations on the weekends were 3.31, 10.19 and 1.19 % lower than those on the weekdays at the JS, PD and QP sites, respectively, due to the low anthropogenic activities occurring on the weekends. Overall, the variations of VOC characteristics were closely related to the land-use types. The average PM_{2.5} values were 45.57 ± 27.59 , 48.51 ± 27.22 and 40.27 ± 27.78 $\mu\text{g m}^{-3}$ at the JS, PD and QP sites, respectively. The highest O₃ concentration was observed at the QP site (99.30 ± 24.00 ppb), followed by the JS (73.59 ± 23.59 ppb) and PD (57.48 ± 20.49 ppb) sites. The occurrence of haze
665 pollution was characterized by the elevated concentrations of aromatics at the JS and PD sites, while the alkanes at the QP site.

The special ratios and PMF were used to investigate the local sources. Local anthropogenic emissions contributed largely to the VOC pollution and varied with land-use types. The vehicle exhaust was determined as the predominate source at the three sites. The second largest VOC contributor was identified as industrial production at the JS and PD sites, whereas it
670 proved to be the fuel evaporation at the QP site. The limited influence of industrial source at the QP site was observed which was related to the land-use type of this site i.e., the QP site is surrounded by natural environment and far away from the industrial regions. Relative to the QP site, JS and PD sites were less affected by the biogenic source which was consistent with the regional characteristics of anthropogenic activities dominated by land-use types. Apart from the local sources, the influence of regional transport was analyzed *via* 24-h back trajectories and PSCF. The results showed that VOCs were
675 impacted by the north or/and northeast trajectories at the sampling sites. The highest PSCF values were observed in the area near the above sites, indicating that local source was a significant contributor to the VOC pollution.

The formation potentials of O₃ and SOA induced by the studied VOCs were also discussed. During the observation period, relative to the SOA formation, the formation of O₃ was more dependent on the VOC concentrations, and the sensitivity analysis of VOC compositions varied with the land-use type. Specifically, the OFP results illustrated that alkenes were the
680 major OFP contributor, and the aromatic compounds were also positively associated with the high OFP. The higher OFPs at the JS and PD sites (50.85 ± 2.63 and 33.94 ± 1.52 ppb) relative to that at the QP site (24.26 ± 1.43 ppb) were observed, while the O₃ concentrations were low. The result was related to the land-use types of both sites which led to high pollutant concentrations and decreased the solar radiation and the intensity of photochemical reactions. In terms of SOAFP, BTEX was the greatest contributor to SOAFP, comprising 86.07, 96.21 and 86.38 % at the JS, PD and QP sites, respectively. The
685 SOAFP values of each land-use types turned out to be in order of JS (1.00 ± 2.03 $\mu\text{g m}^{-3}$) > PD (0.46 ± 0.88 $\mu\text{g m}^{-3}$) > QP (0.41 ± 0.58 $\mu\text{g m}^{-3}$) sites, in connection with the land-use types. The VOCs-PM_{2.5} sensitivity analysis showed that VOCs at the QP site was more sensitive to PM_{2.5} compared with other two sampling sites. Of the four VOCs categories, aromatics at

690 the JS and PD sites and alkanes at the QP site were more sensitive to PM_{2.5}. This finding indicated that alkenes and aromatics were key concerns in reducing the atmospheric secondary pollution in Shanghai. The relevant results herein reveal the influences of land-use type on atmospheric pollution, and set an example for the future VOC observation research at a city level.

Data availability

Measurement data in this study are available in the data repository maintained by Mendeley Data <https://doi.org/10.17632/mf4gf36r9n.1> (Han, 2022).

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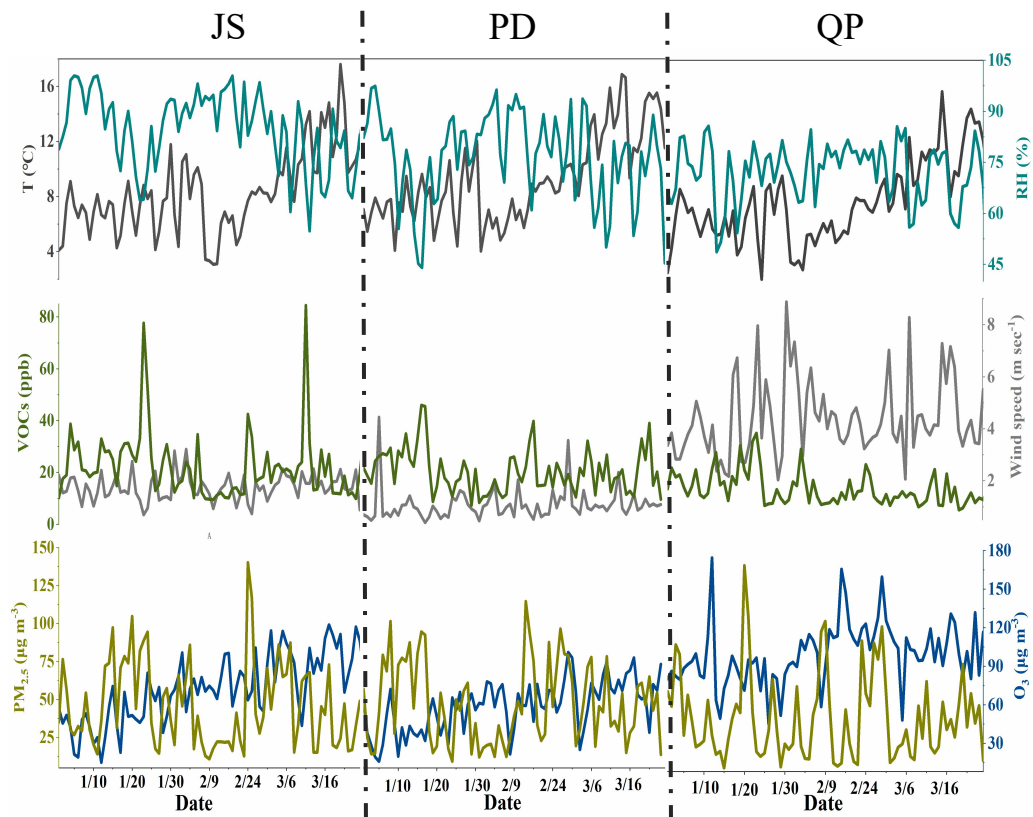


Figure 1: Time series of temperature (T), relative humidity (RH), wind speed, VOC categories, PM_{2.5} and O₃ at the (a) JS, (b) PD and (c) QP sites.

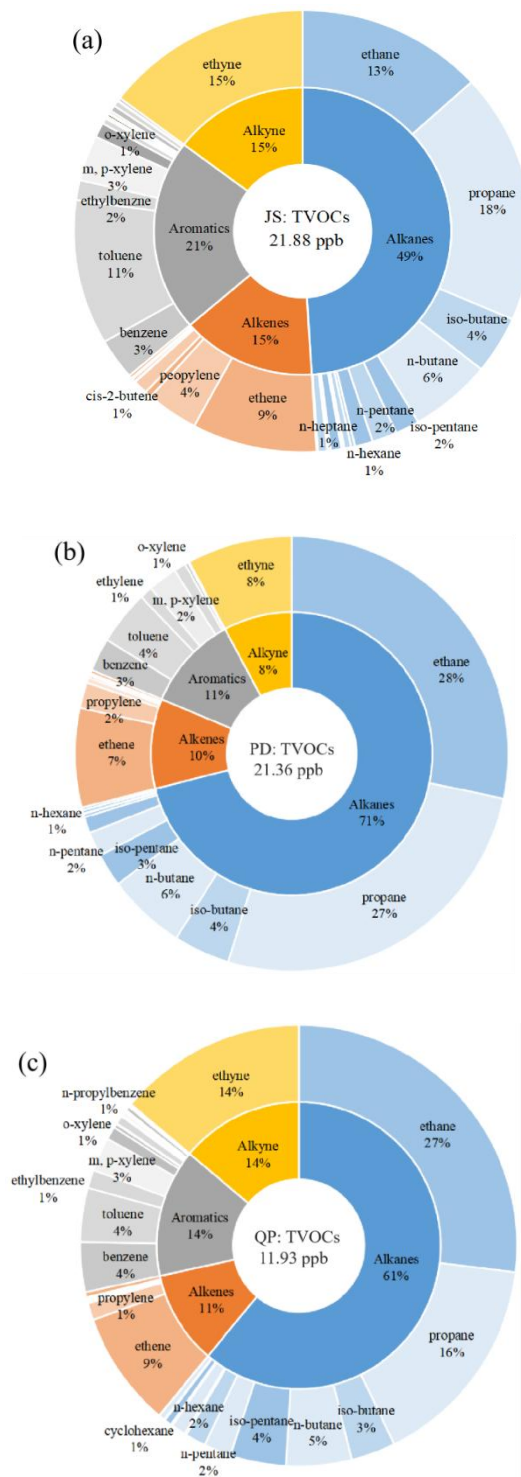


Figure 2: Contributions of the VOC categories (43 species) at the (a) JS, (b) PD and (c) QP sites (the contributions > 1 % were marked).

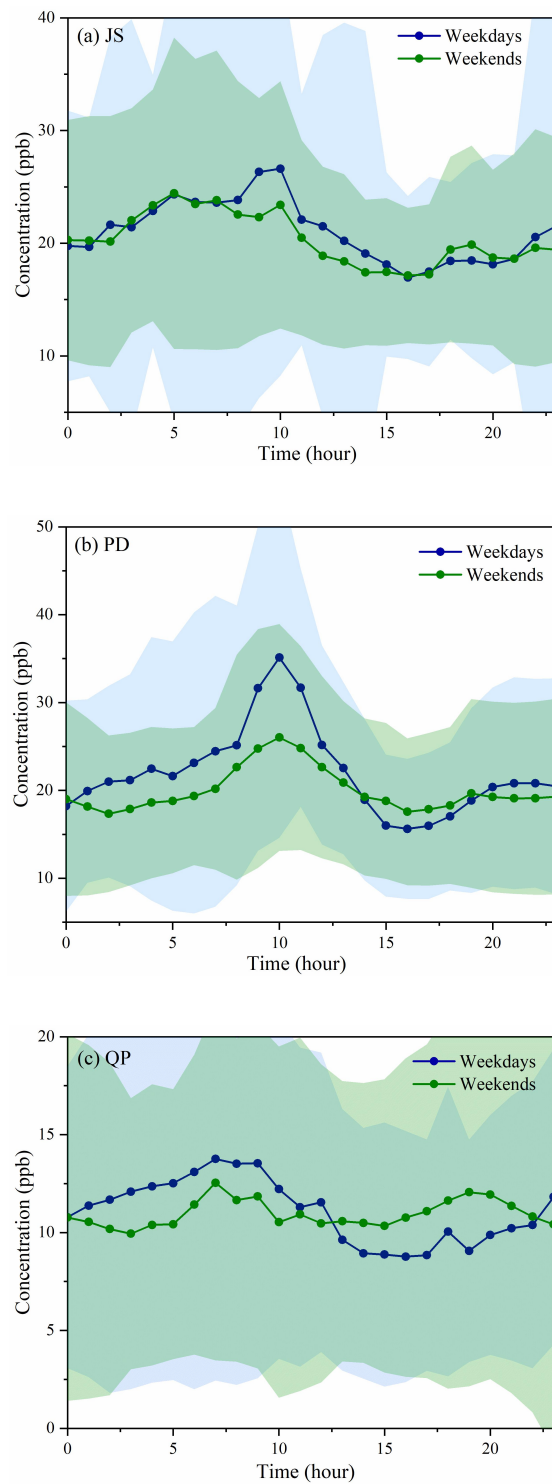


Figure 3: “Weekend effects” of the TVOC concentrations at the sampling sites.

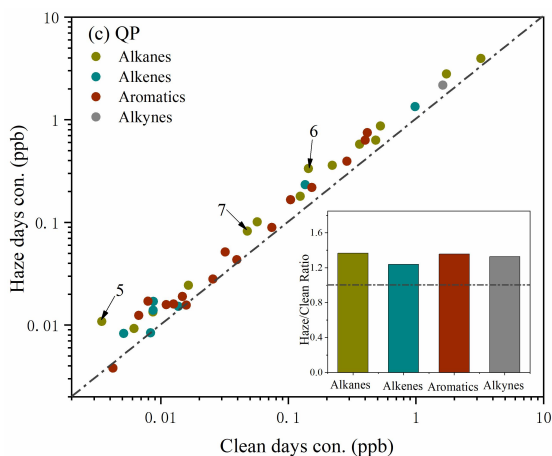
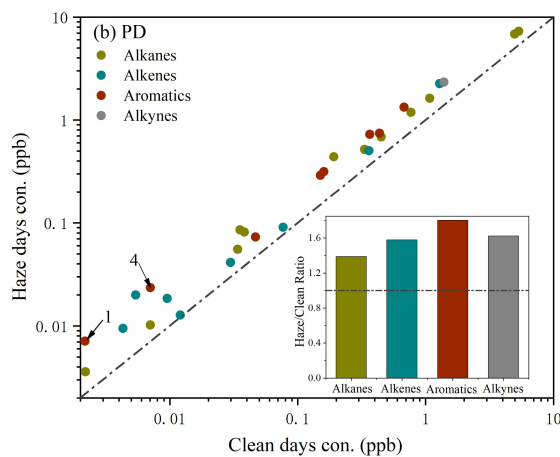
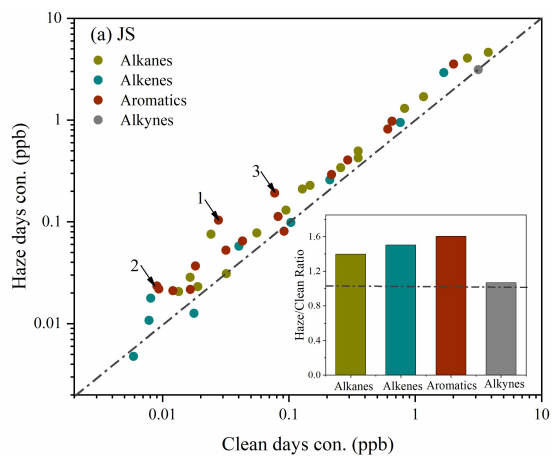


Figure 4: Differences in VOC concentrations between the clean and haze days at the (a) JS, (b) PD and (c) QP sites. 1: 1, 2, 4-trimethylbenzene; 2: *p*-ethylbenzene; 3: *m*-dimethylbenzene; 4: *m*-ethylbenzene; 5: 2, 2, 4-trimethylpentane; 6: *n*-hexane; 7: *h*-heptane.

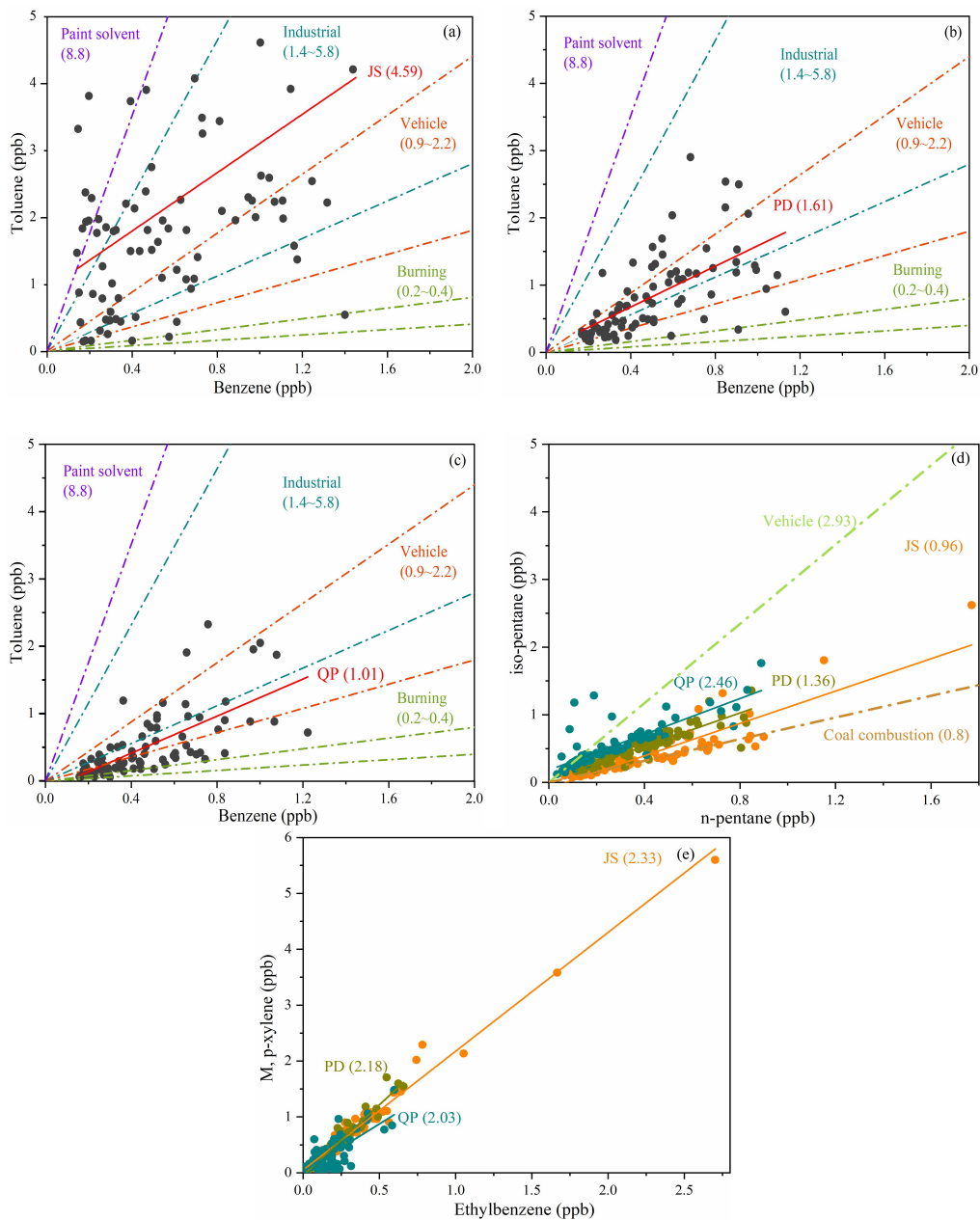
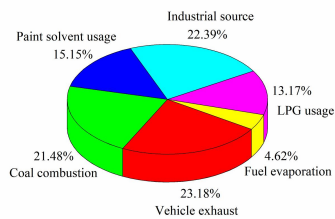
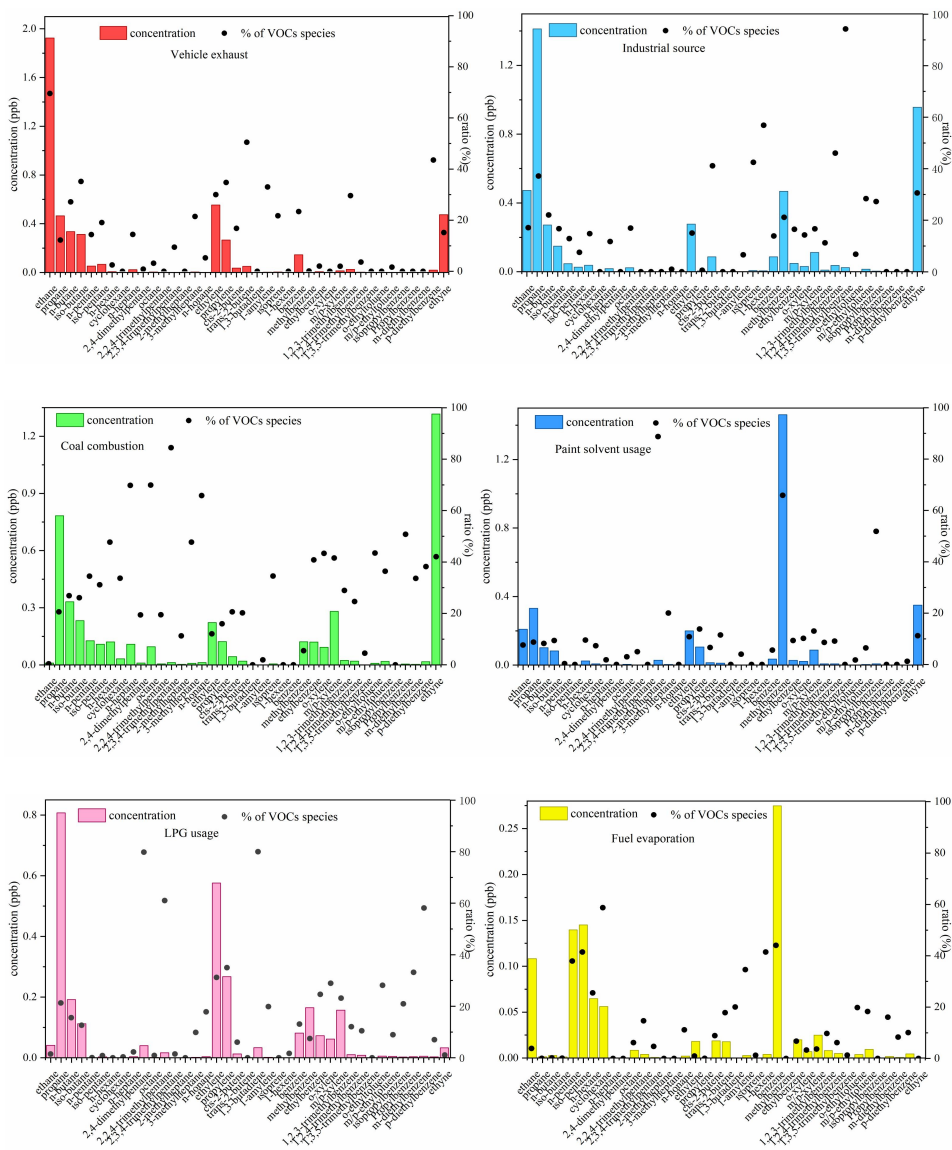
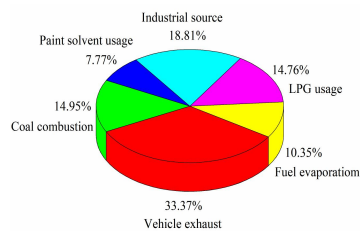
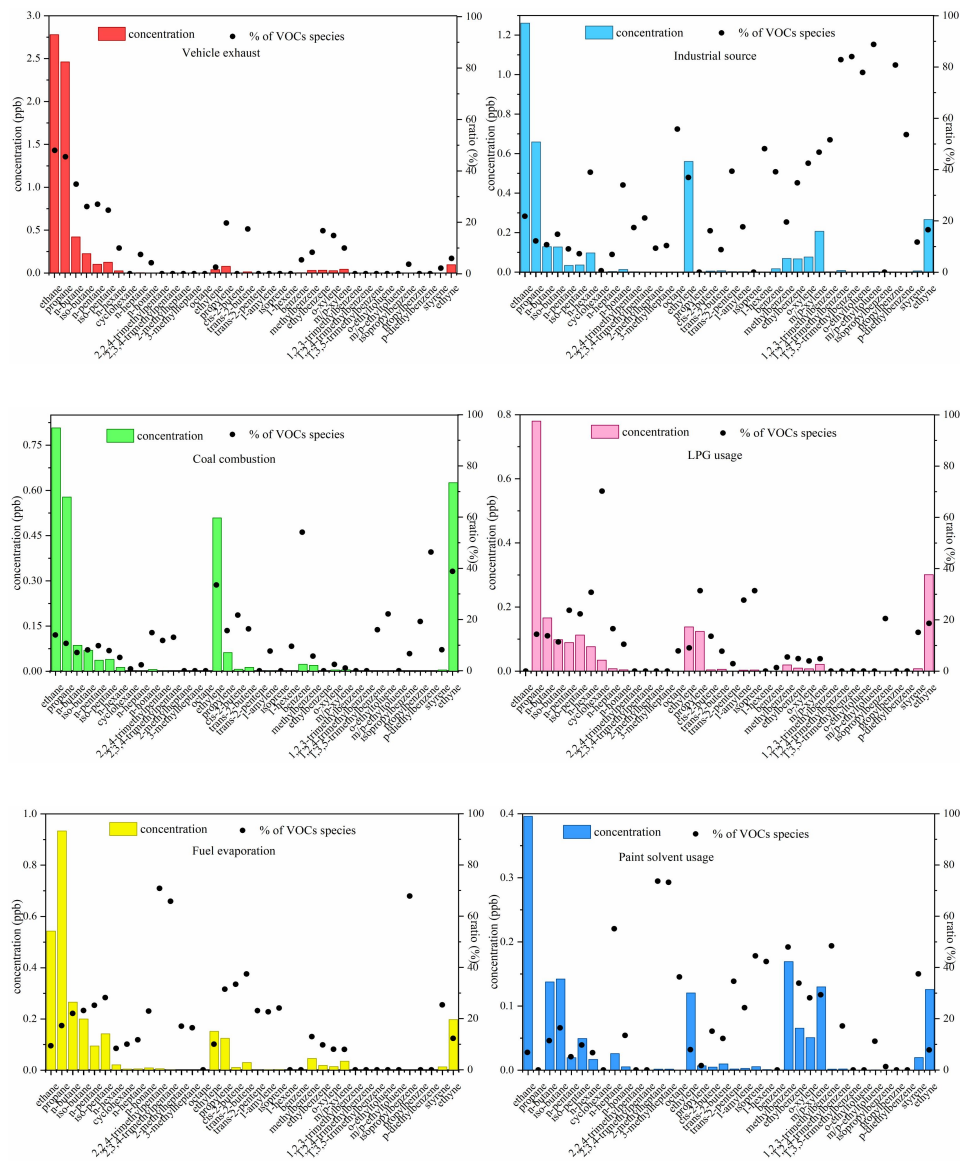


Figure 5: Specific ratios of the target VOC species for the source identification. The linear correlations between toluene and benzene at the (a) JS, (b) PD and (c) QP sites, and (d) *m, p*-xylene and ethylbenzene and (e) *iso*-pentane and *n*-pentane at JS (orange), PD (yellow) and QP (blue).

a. JS



b. PD



c. QP

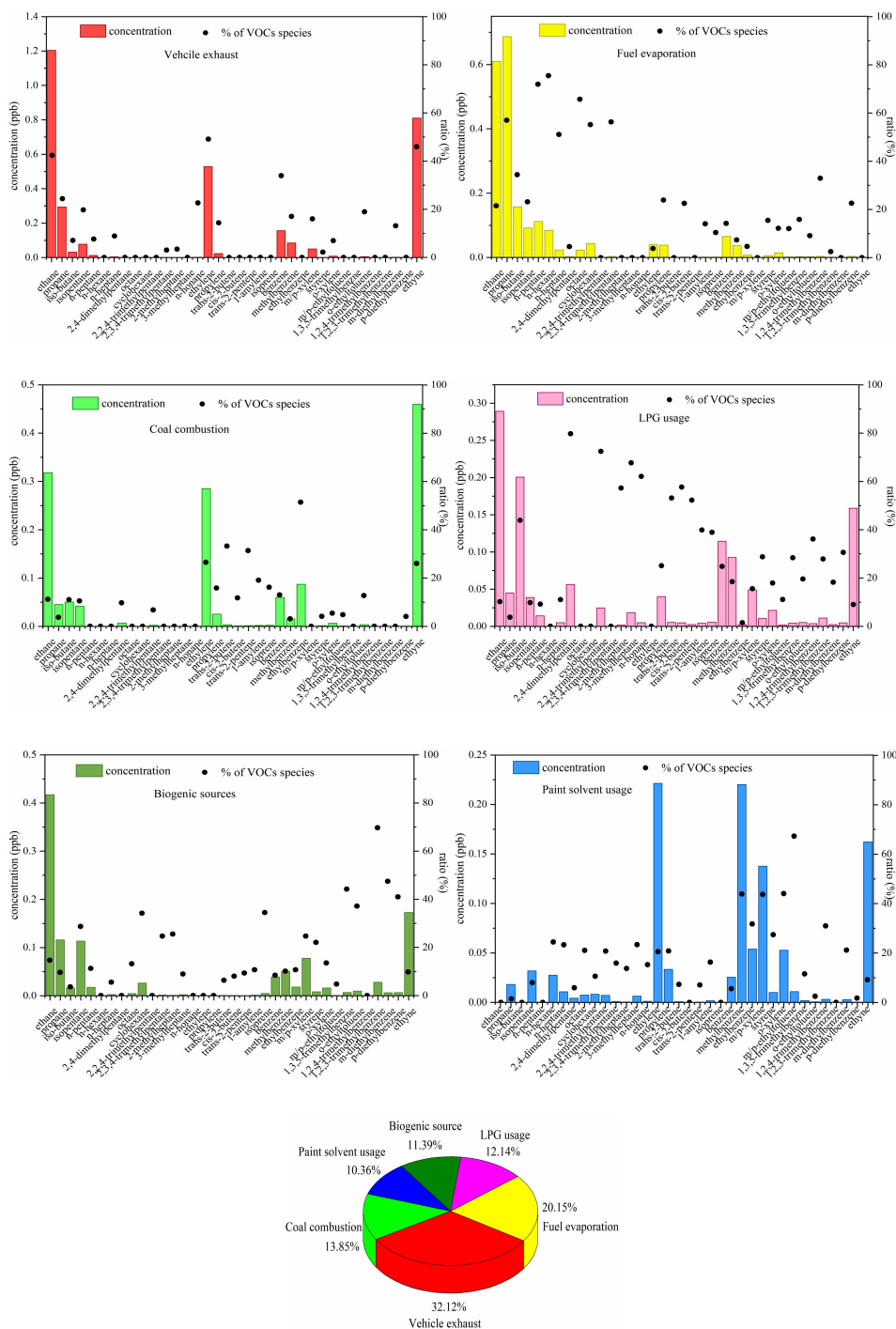
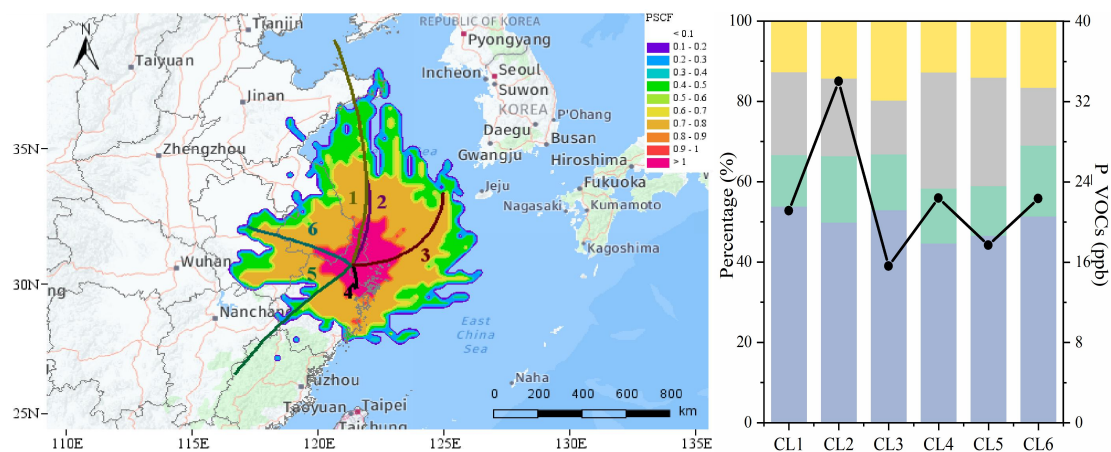
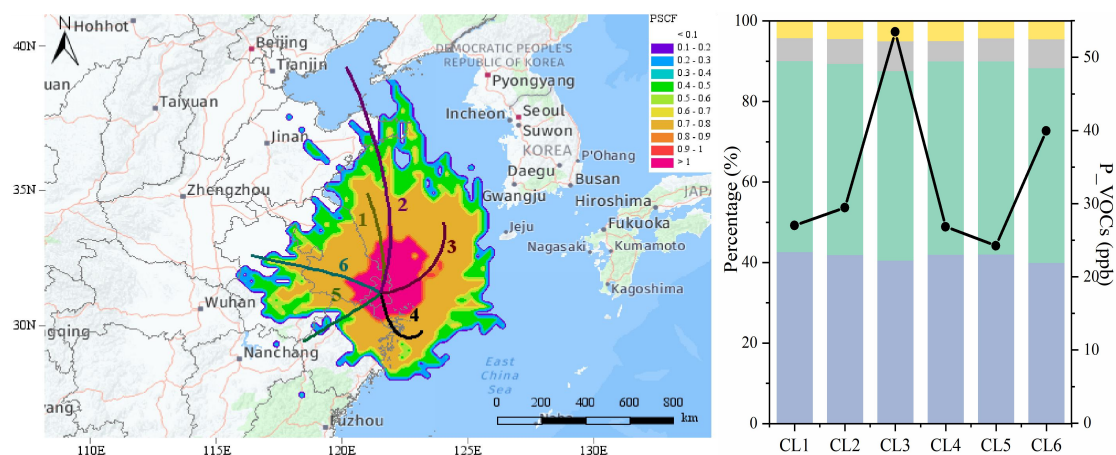


Figure 6: Sources profiles and contributions of VOCs at the (a) JS, (b) PD and (c) QP sites.

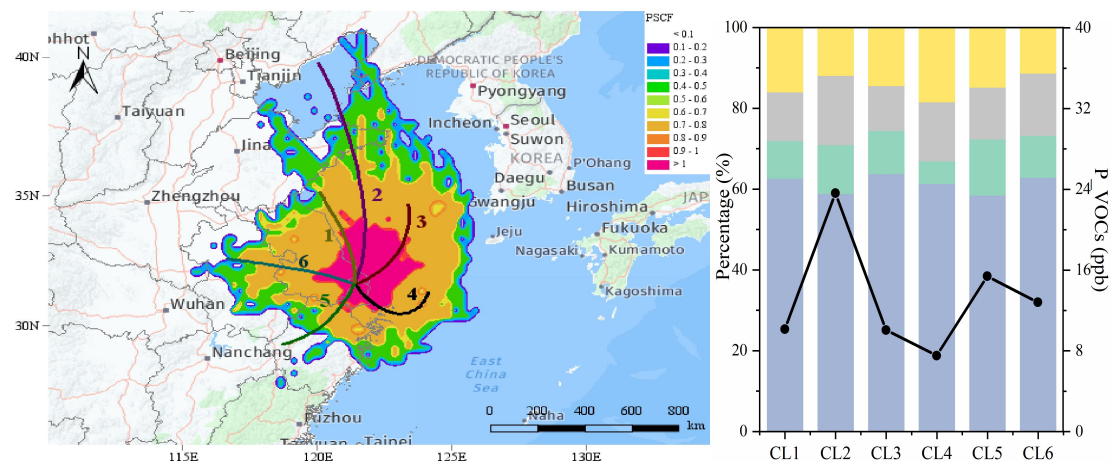
a. JS



b. PD



c. QP



Alkanes Alkenes Aromatics Alkyne P_TVOCs

Figure 7: Backward trajectory cluster analysis (24 h) and PSCF analysis at the JS, PD and QP sites.

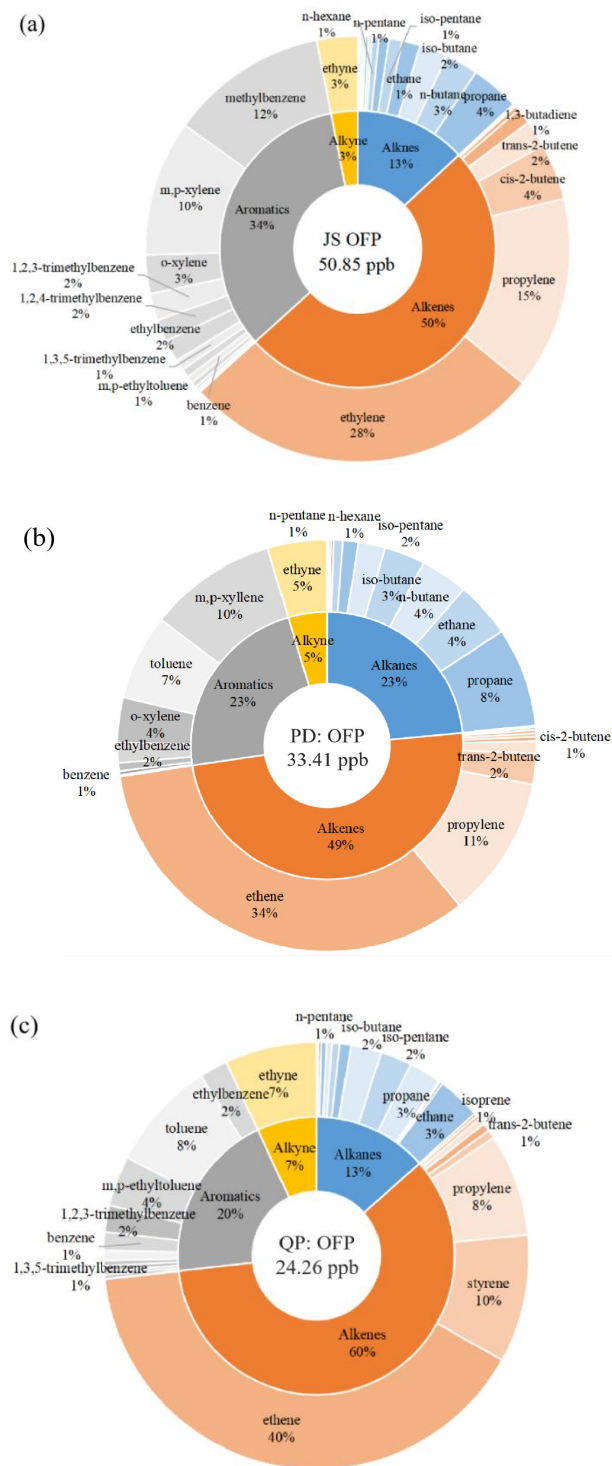


Figure 8: Contributions of VOCs to OFP at the (a) JS, (b) PD and (c) QP sites (the contributions > 1% were marked).

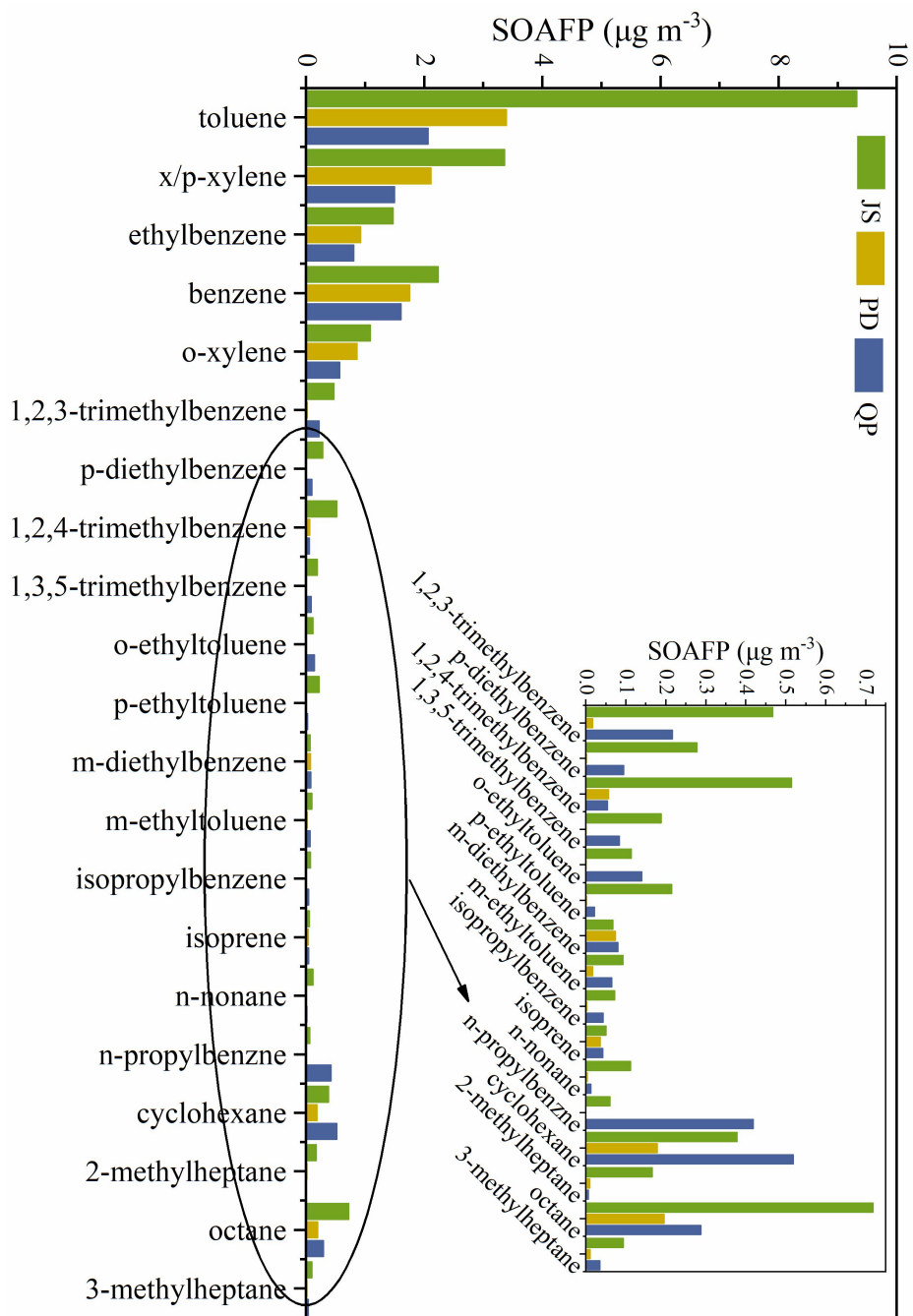


Figure 9: The SOAFP analysis of different VOC species at the JS, PD and QP sites.

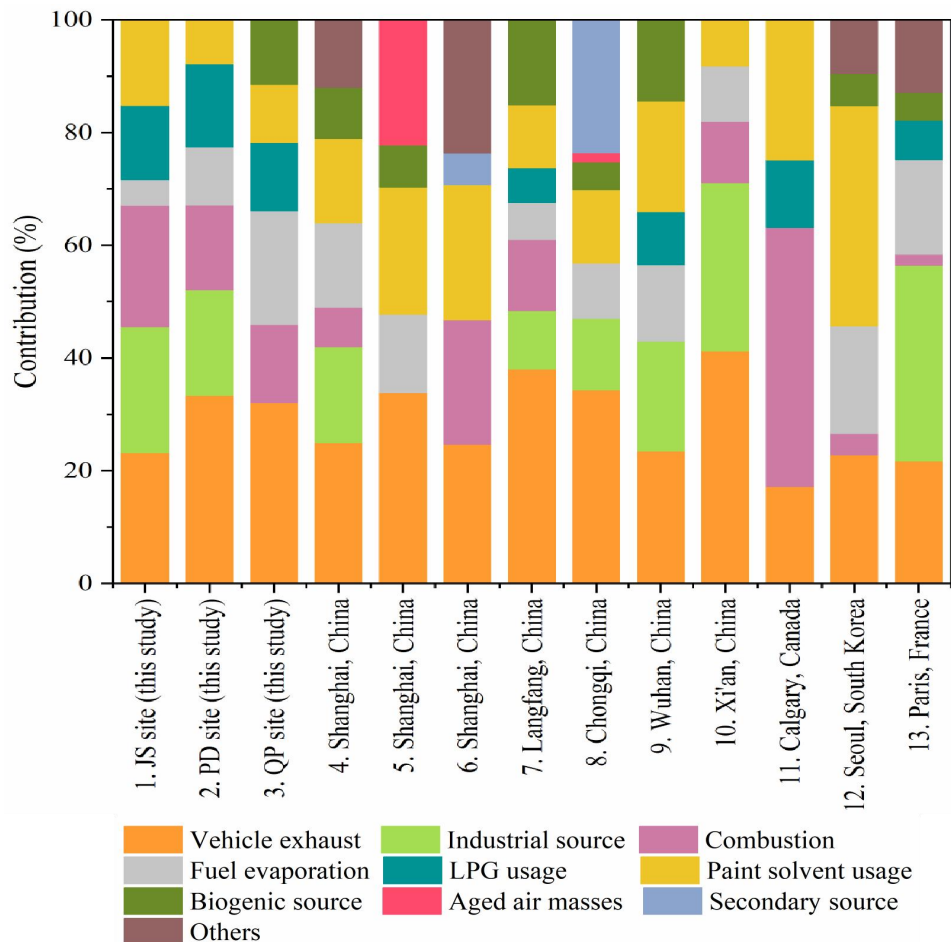


Figure 10: Primary emissions of the observed atmospheric VOCs of this study, and those measured at different locations or during different periods: 1, 2, 3. Shanghai, China (this study); 4. Shanghai, China (Cai et al., 2010b); 5. Shanghai, China (Liu et al., 2019); 6. Shanghai, China (Liu et al., 2021); 7. Langfang, China (Song et al., 2019a); 8. Chongqi, China (Li et al., 2018); 9. Wuhan, China (Hui et al., 2018); 10. Xi'an, China (Song et al., 2021); 11. Calgary, Canada (Bari and Kindzierski, 2018); 12. Seoul, South Lorea (Song et al., 2019b); 13. Paris, France (Gaimoz et al., 2011)

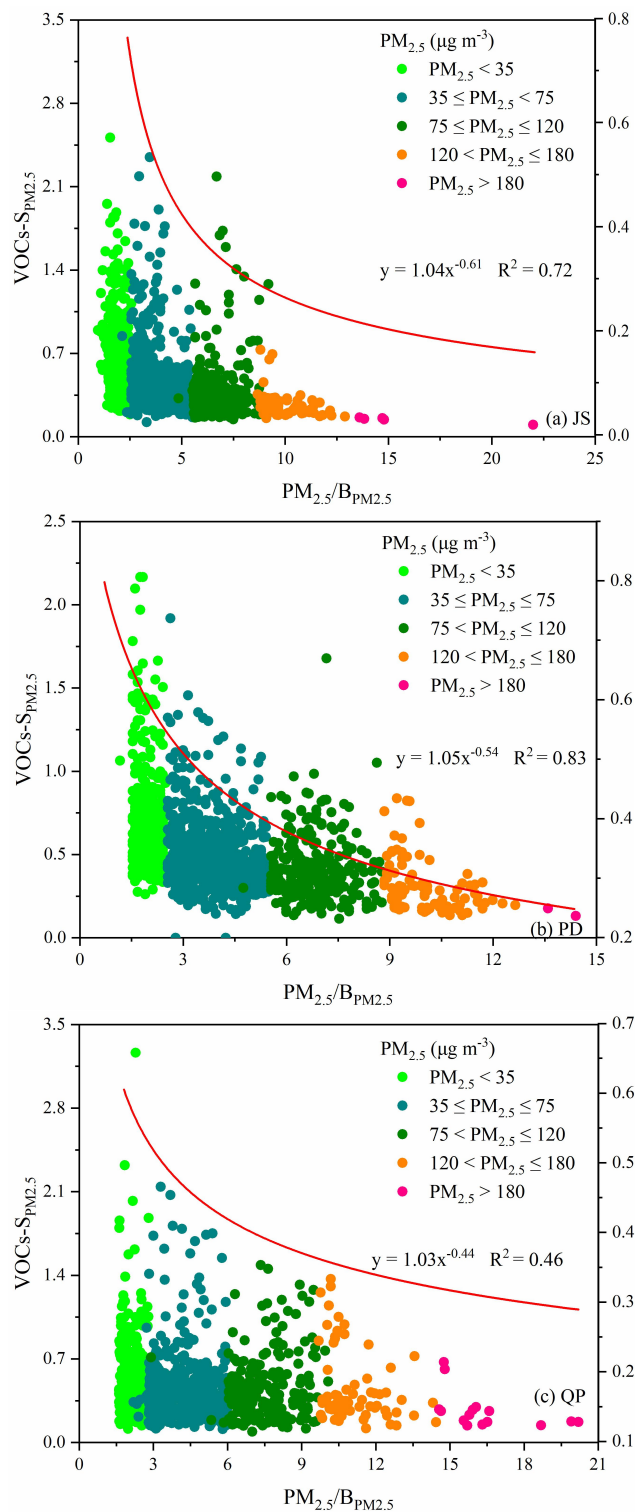


Figure 11: The variations of $VOCs-S_{PM_{2.5}}$ at the different level of $PM_{2.5}$ at the JS, PD and QP sites.

Table 1: Air mass cluster trajectories at the JS, PD and QP sites.

Site	Cluster	Ratio (%)	P_Ratio (%)	P_TVOCs (ppb)
JS	1	16.67	28.04	21.13
	2	29.07	18.03	34.03
	3	19.46	16.62	15.62
	4	12.26	12.75	22.40
	5	5.92	16.62	17.70
	6	16.62	5.93	22.34
PD	1	18.04	33.13	27.03
	2	17.80	10.42	29.45
	3	34.19	21.59	53.48
	4	8.57	15.28	26.85
	5	10.84	8.53	24.25
	6	10.56	11.04	39.95
QP	1	14.20	18.47	10.15
	2	17.57	13.50	23.63
	3	31.87	27.81	10.07
	4	15.20	16.79	7.53
	5	8.71	14.52	15.39
	6	12.45	8.91	12.81