



1 **Measurement report: Seasonal, distribution and sources of**
2 **organophosphate esters in PM_{2.5} from an inland urban city in**
3 **southwest China**

4 Hongling Yin, Jiangfeng Liang, Di Wu, Shiping Li, Yi Luo, Xu Deng

5 College of Resources and Environment, Chengdu University of Information Technology, Chengdu,
6 Sichuan, 610025, China

7 *Correspondence: Hongling Yin (yhl@cuit.edu.cn)*

8 **Abstract.** Organophosphate esters (OPEs) are emerging contaminants in recent years and studies
9 concluded that urban centers were a significant source of OPEs. Samples were collected from six
10 ground-based sites located in Chengdu, a typical fast developing metropolitan of southwest China and
11 were analyzed for seven OPEs in atmospheric PM_{2.5}. The concentrations of Σ_7 OPEs in PM_{2.5} ranged
12 from 5.83 to 6.91 ng·m⁻³, with a mean of 6.6 ± 3.3 ng·m⁻³, and the primary pollutants were TBEP,
13 TnBP, TCEP and TCPP which made up more than 80% in the Σ_7 OPEs. The concentrations of Σ_7 OPEs
14 were higher in autumn/winter than that in summer. Nonparametric test showed that there was no
15 significant difference in Σ_7 OPEs concentrations among the six sampling sites, but the occurrence of
16 unexpected high level of individual OPEs at different sites in autumn might indicate that there was a
17 noteworthy emission. Very strong correlation ($R^2 = 0.98$, $p < 0.01$) between the OPEs in soil and in
18 PM_{2.5} suggested the atmospheric PM_{2.5} settlement is an important source of OPEs in soil. The
19 backward trajectory analysis displayed that OPEs in PM_{2.5} were mainly affected by local sources. The
20 principal component analysis (PCA) identified the OPEs in PM_{2.5} were largely sourced from the plastic
21 industry/interior decoration /traffic emission (34.5%) and chemical, mechanical and electrical industry
22 (27.8%), while PMF model found the main sources were the plastics industry/indoor source emissions,
23 the food/cosmetics industry, and industrial emissions. Differed from the coastal cities, the sustained
24 and stable high local emissions in the inland city were identified which were particularly noteworthy.
25 The chlorinated phosphate, especially TCPP and TCEP have a high content, whose usage and source
26 emissions should be controlled.



27 **1. Introduction**

28 With the prohibition of brominated flame retardants, the production and the demand of
29 organophosphate esters (OPEs) have rapidly increased in recent years (Wang et al., 2012). To date,
30 OPEs are widely distributed in the environment and have been detected in air (Bacaloni, A. et al.,
31 2008), water (Wang et al., 2013; Li et al., 2014), soil (Yin et al., 2016), sediment (Cristale J. et al.,
32 2013; Celano R, et al., 2014) and organisms (Araki et al., 2014; Kim et al., 2011). However, many
33 scholars found that OPEs have negative effects on the human body with the characteristics of water
34 resistance, weather resistance, heat resistance and good polymer substrates compatibility (Matthews, et
35 al., 1990; 1993). Some countries have legislated to restrict the usage of OPEs. Nevertheless, the
36 production and usage of OPEs in China is still on the rise.

37 As synthetic substances, the only source of OPEs in the environment is anthropogenic emissions. The
38 detection of OPEs in Arctic and Antarctic snow samples and atmospheric particulate matter samples
39 demonstrated that OPEs can be transported over long distances. Studies on OPEs in oceans were
40 carried out a lot, and the concentrations of particle-bound OPEs ranged from tens to thousands of ng m^{-3}
41 (Covaci et al., 2007; Cristale J & Lacorte S., 2013; Li et al., 2017). Researchers noted that the
42 contribution of air flow originated from the mainland when high concentrations of OPEs (thousands of
43 ng m^{-3}) appeared (Möller et al., 2012; Lai et al., 2015). In addition, studies proved the urban area was
44 the highest pollution area of OPEs. However, until now, only a few papers reported the concentration
45 and distribution of OPEs in urban atmospheric $\text{PM}_{2.5}$. Concentrations of OPEs in most cities were
46 lower than 10 ng m^{-3} , higher concentrations of 19.2 ng m^{-3} were observed at a suburban site in
47 Shanghai, and 49.1 ng m^{-3} were observed in Hongkong (Ohura et al., 2006; Salamova et al., 2014b;
48 Marklund et al., 2005; Shoeib et al., 2014; Yin et al., 2015; Liu et al., 2016; Ren et al., 2016; Guo et al.,
49 2016; Wong et al., 2018). To date, most of studies in China focus on the OPEs in the Yangtze River
50 Delta and Pearl River Delta, especially eastern coastal cities while little attention was paid to the
51 western inland cities.

52 Chengdu is a typical inland city located in the southwest of China. It is the capital and megacity of
53 Sichuan Province, which covers an area of 14335 square kilometers and has a permanent population of
54 16.33 million. As the important national high-tech industrial base, commercial logistics center and
55 comprehensive transportation hub determined by the State Council, Chengdu is the important central



56 city in the western region. Liu et al. (2016) reported an investigation of three chlorinated OPEs in the
57 atmosphere at 10 urban sites in China during 2013–2014 and found that the highest annual mean
58 concentrations were observed in Chengdu ($1,300 \pm 2,800 \text{ ng m}^{-3}$). However, there is still a lack of
59 information regarding the levels, sources, and fate of OPEs in the southwest China which may
60 obviously differed from the coastal cities or over the sea. In this study, we investigated the atmospheric
61 OPEs in $\text{PM}_{2.5}$ through intensive sampling in an economically fast growing city--Chengdu. Sampling
62 was carried out over one year (October 2014 to September 2015) which was a continuous and further
63 project of our previous study from December 2013 to October 2014. The aims of the study were to: a)
64 report the levels and composition profiles of OPEs in urban air in the typical inland city; (b) obtain the
65 seasonal and spatial variation of OPEs in $\text{PM}_{2.5}$; (c) investigate the relationships and correlations
66 among the target compounds or with influence factors; (d) illustrate the potential sources of OPEs in
67 $\text{PM}_{2.5}$.

68 2. Materials and Methods

69 2.1 Chemicals

70 The main reagents, such as ethyl acetate, acetone, hexane and acetonitrile, were High Performance
71 Liquid Chromatography (HPLC) grade (Kelon Chemical, China). The standard solution (Sigma
72 Aldrich) included tri-n-butyl phosphate (TnBP), tris-(2-ethylhexyl)phosphate (TEHP), tris-(2-
73 butoxyethyl) phosphate (TBEP), triphenyl phosphate (TPhP), tris-(2-chloroethyl)-phosphate (TCEP),
74 tris-(2-chloroisopropyl)phosphate (TCPP), and tris-(2,3-dichloropropyl)-phosphate (TDCIPP). Copper,
75 aluminium oxide, silica gel, Na_2SO_4 and other chemicals were purchased from Kelon Chemical.
76 Deionized water was supplied from a Milli-Q equipment.

77 2.2. Sample collection

78 The atmospheric sampling sites were located in the main city area (site B: downtown; site C: south; site
79 D: east; site E: north; site F: west) and suburban area (site A) of Chengdu, as shown in Fig. S1. The
80 atmospheric samples were collected by KC-6120 medium flow atmospheric comprehensive sampler
81 with quartz film. The speed was set at 100 L min^{-1} , and each collection campaign lasted 23 h. The
82 sampling campaign was carried out between October 2014 and September 2015. A total of 149
83 samples were obtained. Most of the weather conditions were cloudy days, with south/north wind at



84 ≤ 5.5 m/s. Temperature ranged from 0 to 35 °C. Weather conditions could represent typical weather
85 conditions of the season.

86 **2.3. Sample preparation and analysis**

87 The shredded PM_{2.5} sample film was placed in a test tube and incubated in 20 mL ethyl acetate/acetone
88 (v:v, 3:2) for 12 hours. After ultrasonic extraction for 30 minutes, the liquid was separated, and the
89 residue was further extracted with 10 mL ethyl acetate/acetone (v:v, 3:2) by ultrasonic extraction for 15
90 minutes. The extracts were combined and concentrated by vacuum-condensing equipment (Buchi
91 Syncore Q-101, Switzerland) to approximately 1 mL, then loaded onto an activated aluminium
92 oxide/silica gel (v: v, 3: 1) column. The column was first eluted with 20 mL hexane to remove
93 impurities, then with 20 mL ethyl acetate/acetone (v: v, 3: 2) and the eluate (ethyl acetate/acetone) was
94 collected. The solvent extracts were concentrated by vacuum-condensing equipment and diluted to 200
95 μ L for gas chromatography-mass spectrometry (GC-MS) (Shimadzu 2010plus, Japan) analysis.

96 The GC is equipped with a capillary column Rti-5MS (30 m \times 0.25 μ m \times 0.25 mm, Kelong), with a
97 280 °C inlet temperature using splitless injection. The MS source was electron impact (EI) and
98 operated in selected ion monitoring (SIM) mode. Helium was used as a carrier gas with a flow rate of
99 1.00 mL min⁻¹. The GC oven temperature was held at 50 °C for 1 minute, increased to 200 °C at 15 °C
100 min⁻¹ and held for 1 minute, increased to 250 °C at 4.00 °C min⁻¹, and then increased to 300 °C at 20 °C
101 min⁻¹ and held for 4 minutes. The interface temperature was 280 °C, and the ion source temperature was
102 200 °C. The respective characteristic ion and reference ions (m/z) of the 7 target compounds were:
103 155/99, 211, 125 (TnBP), 249/63, 143, 251 (TCEP), 125/99, 201, 277, 157 (TCPP), 75/99, 191, 209,
104 381 (TDCPP), 326/325, 77, 215 (TPhP), 85/100, 199, 299 (TBEP), 99/113 and 211 (TEHP).

105 **2.4. QA / QC**

106 The concentrations of the 7 OPEs were determined by an external standard method. The correlation
107 coefficients of the standard curves of the seven OPE monomers were all greater than 0.990. The
108 recoveries of the 7 OPEs ranged from 83.9% to 121.2%. A matrix blank was analysed with each batch
109 of samples. Only TnBP was detected in the blanks, and the level of TnBP found in the blanks was <5%
110 of the concentrations measured in all samples, which means it was negligible. The instrument precision
111 was in the range of 1.9%-8.3%.



112 3. Results and Discussion

113 3.1. Levels of OPEs in PM_{2.5}

114 OPEs were present in PM_{2.5} samples collected across the study area (Fig. S1). Four OPEs (TCPP,
115 TDCPP, TCEP and TnBP) were detected in all samples (n=149), while TBEP was detected in all but
116 one sample. Additionally, TEHP was detected in 96.7% of samples overall and TPhP was detected in
117 98% of samples. The high detection frequencies of most OPEs indicated OPE contamination was
118 ubiquitous in the air of Chengdu city.

119 Concentrations of Σ_7 OPEs in PM_{2.5} across the six sites were in the range of 3.5 - 11.5 ng m⁻³, and the
120 annual median concentration of Σ_7 OPEs was 6.5 ± 3.3 ng m⁻³ (Fig. 1). The average value of OPEs in
121 PM_{2.5} at each site in four seasons was almost at the same level (5.8 ± 1.3 ng m⁻³-6.9 ± 2.5 ng m⁻³).
122 Nonparametric test showed that there was no significant difference in Σ_7 OPEs concentrations among
123 the six sampling sites, indicating that the atmosphere mixed evenly, and there was no particularly
124 heavy or light pollution area in Chengdu city. These data are quite consistent with our previous study
125 that showed the annual median concentration of OPEs in PM_{2.5} from December 2013 to October 2014
126 (Yin et al., 2015). Interestingly, the annual median of total OPEs at the suburban site was not the
127 lowest as might be expected and is instead likewise similar to, or even higher than some urban sites
128 which indicated more local sources of these compounds in the suburban area.

129 The concentrations of OPEs in the particles of Chengdu were comparable to that reported from Beijing
130 (0.257 - 8.36 ng m⁻³) (Wang et al., 2018), 6.6 ng m⁻³ (Σ_6 OPEs) for Shanghai urban site (Ren et al.,
131 2016), 6.5 ng m⁻³ (Σ_6 OPEs) for Bursa, but higher than that in Houston, US (Σ_{12} OPEs, 0.16 - 2.4 ng m⁻³)
132 (Clark et al., 2017), Dalian (Σ_9 OPEs, 0.32-3.46 ng m⁻³, 1.21 ± 0.67 ng m⁻³) (Wang et al., 2019),
133 European Arctic(0.033 - 1.45 ng m⁻³) (Salamova et al., 2014), Northern Pacific and Indian Ocean (0.23
134 - 2.9 ng m⁻³) (Moller et al., 2012), the Yellow Sea and Bohai Sea (0.044 - 0.52 ng m⁻³) (Li et al., 2017),
135 South China Sea (0.047 - 0.161 ng m⁻³) (Lai et al., 2015), North Atlantic and Arctic Oceans(0.035 -
136 0.343 ng m⁻³) (Li et al., 2017). And lower than that in Guangzhou and Taiyuan (Σ_{11} OPEs, 3.10 - 544ng
137 m⁻³) (Chen et al., 2020), in Bursa, Turkey (Σ_6 OPEs, 0.53 - 19.14 ng m⁻³) (Kurtkarakus et al., 2018), 20
138 industrial sites in an urban region (Σ_{12} OPEs, 0.52 - 62.75 ng m⁻³) in Guangzhou, China(Wang, T. et al.,
139 2018).



140 3.2. The composition profiles of OPEs in PM_{2.5}

141 There was clear dominant non-chlorinated OPEs across Chengdu city. The annual median
142 concentrations of total OPEs were fairly uniform at six sites and influenced mainly by the alkylated
143 OPEs. As listed in Table 1, the general trend was found that TBEP was the most abundant OPE (2.3 ng
144 m⁻³, 35.3%), followed by TCEP (1.1 ng m⁻³, 16.3%) ≈ TnBP (1.0 ng m⁻³, 15.6%) ≈ TCPP (1.0 ng m⁻³,
145 15.0%) > TPhP (0.5 ng m⁻³, 8.4%) > TEHP (0.3 ng m⁻³, 5.1%) > TDCPP (0.3 ng m⁻³, 4.3%), with the
146 concentrations of TBEP being approximately 7 - 10 times higher than those of TDCPP and TEHP. The
147 composition profile of OPEs was similar at all sites except for that the east site which has a higher
148 contribution of TnBP. But TBEP, TCEP, TCPP and TnBP were dominant OPEs across the city who
149 contributed more than 80% to Σ₇OPEs. This profile was similar to that in Longyearbyen, Norway, with
150 primary pollutants being TnBP and TBEP (Möller et al., 2012), as well as the OPEs in outdoor urban
151 air being TBEP > TCPP > TCEP > TnBP > TPhP in Stockholm, Sweden (Wong et al., 2018) and
152 TBEP > TCPP > TPhP > TEHP > TCEP in Turkey (Kurtkarakus et al., 2018). However, these results
153 substantially differed from the report of an urban site in Shanghai that showed TCEP (0.1 - 10.1 ng m⁻³,
154 1.8 ng m⁻³) > TCPP (0.1 - 9.7 ng m⁻³, 1.0 ng m⁻³) > TPhP (0.06 - 14.0 ng m⁻³, 0.5 ng m⁻³) > TBP (0.06 -
155 2.1 ng m⁻³, 0.4 ng m⁻³) > TDCPP (Nd.-23.9 ng m⁻³, 0.3 ng m⁻³), whereas TBEP was only detected in 3
156 out of 116 samples (Nd.-0.7 ng m⁻³, Nd.) (Ren et al., 2016), and the reported data over the Bohai and
157 Yellow Seas showed TCPP (43 - 530 ng m⁻³; 100 ng m⁻³, 50 ± 11%) > TCEP (27 - 150 ng m⁻³; 71 ng m⁻³,
158 25 ± 7%) > TiBP (19 - 210 ng m⁻³; 57 ng m⁻³, 14 ± 12%) > TnBP (3.0-37 ng m⁻³; 13 ng m⁻³). Li et al.
159 (2014) determined the primary pollutant of outdoor air in Nanjing was TCEP, and TBEP was not
160 detected. These differences reflected that there were significant differences in OPE production and
161 usage in different regions, even in the same country. It should be noted that concentrations of TCPP
162 and TCEP were in the same level in this study, suggesting the industrial replacement of TCEP by
163 TCPP wasn't identified in the southwest China which differed from that the higher concentration of
164 TCPP in comparison with TCEP was observed due to the industrial replacement of TCEP by TCPP in
165 Europe (Quednow and Püttmann, 2009). This was confirmed by the fact that there are manufacturers
166 and sellers of TCEP and TCPP in Chengdu, indicating that there is production and demand both for
167 TCPP and TCEP in this region.



168 Combined with the data of 2013-2014 year (Yin et al., 2015), TBEP was always the dominant OPEs
169 during the two sampling periods (2013-2014 and 2014-2015). Kruskal Wallis test was used and found
170 that TnBP and TCPP had no significant difference between the two sampling periods, but there were
171 significant differences in other kinds of OPEs between the two sampling periods. This indicated that
172 the production and usage of individual OPEs have certain change suggesting that OPEs should be
173 better investigated and governed for individual compounds.

174 OPEs can be categorized by whether they are halogenated, alkylated or aryl OPEs. Of the OPEs
175 measured in this study, TCEP, TCPP and TDCPP are halogenated, TBEP, TnBP and TEHP are
176 alkylated, and TPhP is aryl OPEs. The OPEs in $PM_{2.5}$ at all sites were dominated by the alkylated
177 compounds ($55.9 \pm 10.1\%$), followed by halogenated OPEs ($35.8 \pm 9.9\%$) and aryl OPEs ($8.3 \pm 4.1\%$).
178 Our results are similar to those observed in Bursa, Turkey (Kurtkarakus et al., 2018), whose alkylated
179 OPEs covered 68% ~ 95% of total OPEs, while halogenated OPEs covered 3.1% ~ 29%, and aryl OPEs
180 covered 1.4% ~ 3.7% of total OPEs. At Longyearbyen, the non-chlorinated OPE concentrations
181 comprised 75% of the Σ_8 OPE concentrations (Salamova et al., 2014a). However, our results are
182 obviously different from many studies with the atmospheric samples collected in urban areas being
183 dominated by chlorinated OPEs (50 ~ 80%) (Salamova et al., 2014b; Liu et al., 2016; Guo et al., 2016).
184 In our study, non-chlorinated OPEs were dominant in urban and suburban area across the city.

185 **3.3. Seasonal and spatial variation of OPEs in $PM_{2.5}$**

186 The mean seasonal concentrations were plotted for six sampling sites in Fig. 2. The data were quite
187 consistent with our previous study from December 2013 to October 2014 (Yin et al., 2015). The
188 concentrations of OPEs in $PM_{2.5}$ have been fairly uniform in the past three years. As shown in Fig. 2,
189 the general order of the decreasing average Σ_7 OPEs concentrations in suburban area was autumn ($8.4 \pm$
190 4.3 ng m^{-3}) \approx winter ($8.4 \pm 4.5 \text{ ng m}^{-3}$) $>$ spring ($7.6 \pm 2.2 \text{ ng m}^{-3}$) $>$ summer ($3.5 \pm 1.1 \text{ ng m}^{-3}$), while in
191 urban area was autumn ($9.30 \pm 3.89 \text{ ng m}^{-3}$) $>$ winter ($6.63 \pm 3.65 \text{ ng m}^{-3}$) $>$ spring ($6.36 \pm 1.72 \text{ ng m}^{-3}$)
192 $>$ summer ($4.60 \pm 1.91 \text{ ng m}^{-3}$). The average concentration of Σ_7 OPEs in autumn/winter was
193 approximately 2 times that in summer. In summer, the turbulent flow accelerated the diffusion of
194 pollutants, leading to the lowest concentration, while the higher concentrations of OPEs appeared in
195 autumn and winter because the inversion layer appeared more frequently in autumn and winter,
196 resulting in the pollutants being more difficult to diffuse and dilute. This seasonal variation was mostly



197 in line with that at the Shanghai urban site of autumn (8.4 ng m^{-3}) > winter (7.6 ng m^{-3}) > spring (5.5 ng
198 m^{-3}) > summer (4.4 ng m^{-3}), of which the maximum value was also approximately twice the minimum
199 (Ren et al., 2016). In addition, this finding was similar to that in Xinxiang that no significant seasonal
200 changes and only exhibited individual high values in winter. On the contrary, Wang et al. (2019) found
201 the $\text{PM}_{2.5}$ -bound fractions of OPEs varied significantly between seasons in Dalian, China, with their
202 concentrations higher in hot seasons, which may due to the temperature-driven emission or gas-particle
203 partitioning. Wong et al. (2018) reported that most of OPEs in outdoor urban air showed seasonality,
204 with increased concentrations during the warm period in Stockholm, Sweden. Sühning et al. (2016)
205 reported temperature dependence of chlorinated OPEs and EHDPP in Arctic air. Liu et al. (2014) did
206 not observe any temperature dependence for the OPEs in urban air in Toronto, Canada. Thus previous
207 reports of temperature dependence of OPEs are not consistent. In this study, the lowest concentrations
208 of $\Sigma_7\text{OPEs}$ and individual compound were observed in summer suggesting the OPEs level was not
209 driven by the temperature-driven emission or gas-particle partitioning, but mainly by the local emission
210 sources.

211 Compared to the coastal cities, the most obvious difference was that concentrations of almost all OPEs
212 monomers in this study were highest in autumn/winter and lowest and concentrated in summer
213 suggesting the sustained and stable high local emissions in the inland city which were particularly
214 noteworthy. No point source was identified in summer and the OPEs level was diluted and diffused in
215 summer due to the higher wind speed than in winter in the inland city. This was different from the
216 coastal cities: Liu et al. (2016) reported that the highest TCPP and TCEP concentrations were observed
217 in the summer in Guangzhou and Javier et al. (2018) found the OPEs in spring generally exhibited the
218 lowest concentrations in Bizerte, Tunisia, probably linked to the influence of local meteorological
219 conditions and air mass trajectories to a lesser extent.

220 Though Kruskal Wallis test showed that there was no significant variation of $\Sigma_7\text{OPEs}$ concentrations
221 across the city, the spatial differences were identified in the study. For example, TnBP and TCPP had
222 significant difference among six sites. In addition, the higher concentrations and more dispersed pattern
223 of most OPEs were observed in autumn and winter than in summer (Fig. 3). The concentrations of
224 TEHP in autumn at the eastern and northern sampling site were more dispersed than others. The same
225 dispersion pattern was observed for TBEP in winter at the western sampling site, TPhP in autumn at
226 the suburban sampling site, TnBP in autumn at the eastern sampling site, suggesting that there existed



227 the extra emission sources in autumn or winter. Considered the layout of Chengdu which develops
228 from the central area with the loop line (the first ring road, the second ring road and the third Ring
229 Road), we could understand the OPEs levels and distribution were quite uniform across the city. But
230 different types of industrial parks in different directions in Chengdu may be the reason for the spatial
231 differences of OPEs. For example, in the east of Chengdu, there are automobile industrial parks and
232 other large industrial parks while logistics and shoemaking industrial parks in the suburbs. The
233 occurrence of unexpected high level of individual OPEs at different sites in autumn might indicate that
234 there was a noteworthy emission. The spatial and seasonal variation of individual OPE suggest that the
235 control and management of OPEs should be taken to the individual OPE.

236 OPEs can be categorized as halogenated, alkylated and aryl OPEs. Of the OPEs measured in this study,
237 TCEP, TCPP and TDCPP are halogenated, TBEP, TnBP and TEHP are alkylated, and TPhP are aryl
238 OPEs. Many studies focused on the halogenated OPEs due to their persistence, bio-accumulation, and
239 potential human health effects, and they dominated the OPEs profile in the air of many cities and other
240 areas (Zhang et al., 2016, Li et al., 2017). Liu et al. (2016) reported that the sum of the concentrations
241 of the three halogenated OPEs at 10 urban sites ranged from 0.05 to 12 ng m⁻³ suggesting the highest
242 production volume and widest applications of OPEs leading to large emissions of OPEs in China in
243 recent years. However, in our study, the mean concentrations of halogenated, alkylated and aryl OPEs
244 were 2.4 ± 1.4 ng m⁻³, 3.7 ± 2.1 ng m⁻³, 0.5 ± 0.4 ng m⁻³, respectively, which showed the alkylated
245 OPEs dominated the profile of OPEs in PM_{2.5} in Chengdu. In different seasons, the most notable
246 seasonal variation was observed for alkyl phosphate, followed by halogenated OPEs and aryl OPEs.
247 These results were significantly different from those in other studies which reported that the
248 halogenated OPEs had the maximum seasonal variability (Guo et al., 2016; Shoeib et al., 2014).

249 **3.4. Correlation analysis of OPEs**

250 **3.4.1 Linkage to environmental factors**

251 Most of OPE monomers concentrations in PM_{2.5} have a strong linear correlation ($R^2 = 0.79$) with their
252 vapor pressure (Fig. 4), suggesting that the vapor pressure is an important factor controlling the levels
253 of OPEs in PM_{2.5} except for TBEP. Generally speaking, the greater the vapor pressure of OPEs, the
254 easier it is to be released into the environment. Therefore, the sources of most OPEs in Chengdu
255 atmospheric PM_{2.5} are mainly both from the production process containing OPEs and the phase



256 transition process before they enter into the atmosphere. The boiling points of OPEs are relatively high,
257 so they tend to be adsorbed in $PM_{2.5}$ after being released to the environment, and their gas-particle
258 distribution determines their concentration in $PM_{2.5}$. Interestingly, the vapor pressure of TBEP is lower
259 than other OPEs, but its concentration in $PM_{2.5}$ was higher which indicated that there were sustained
260 and stable high emission sources to keep its concentration at a high level which may include the traffic
261 emission source (Chen et al., 2020). Sühning et al. (2016) reported non-halogenated OPE
262 concentrations in Canadian Arctic air appeared to have diffuse sources or local sources close to the
263 land-based sampling stations.

264 **3.4.2 Correlation between target analytes**

265 Spearman's ranks correlation coefficients were used to investigate the potential emission sources for
266 OPEs by the relationship between individual OPE in $PM_{2.5}$ (Fig.5, Table 2). Fig. 4 showed no
267 statistically significant positive correlations between OPE monomers ($r < 0.50$, $p < 0.01$). However,
268 Σ_7 OPEs concentrations were closely related to TBEP, TCEP and TnBP ($r = 0.53-0.61$, $p < 0.01$) which
269 further identified the OPEs levels were influenced mainly by the dominated OPEs compounds.
270 Comparatively, weak correlations between most of OPEs were observed in urban regions (Wang et al.,
271 2018) and Turkey (KurtKarakus et al., 2018). However, strong correlations between individual OPEs
272 were found in Guangzhou and Taiyuan (Chen et al., 2020).

273 Further analysis results were shown in Table 2. Only significant correlation between TCPP and TCEP
274 both at downtown ($r = 0.82$, $p < 0.01$) and suburban sites ($r = 0.85$, $p < 0.01$) were observed indicating the
275 high homology between these two compounds. So the inland city in China is still using a large number
276 of products containing chlorinated flame retardants, which was confirmed by our previous study of
277 house dust (Liu et al. 2017; Yin et al., 2019). At downtown site, another significant correlation existed
278 between TEHP and TCEP ($r = 0.50$, $p < 0.01$) while others have weak to moderate correlations ($r < 0.46$,
279 $p < 0.01$). The downtown area mainly focuses on the light industry and software development, and
280 TCPP, TCEP, TnBP, TBEP and TPhP are used in textile, leather, electronic products and other fields.
281 However, the correlation of each OPE monomer at site A (suburb) was stronger than that in the urban
282 area. TnBP and TCEP, TnBP and TDCPP, TCEP and TCPP, TCEP and TDCPP, TCEP and TBEP,
283 TCPP and TDCPP and TBEP were all extremely significant. This result indicated that the pollution in
284 the suburb was commixed and was influenced by many kinds of pollution sources.



285 3.4.3 Correlation analysis of OPEs and PM_{2.5} concentrations

286 The SPSS software scatter diagram was used to analyse the relationship between the concentrations of
287 OPE monomers and PM_{2.5}. As displayed in Fig.S2, only weak to moderate correlation were observed
288 between most of OPEs and PM_{2.5} except significant correlation was found between TDCPP and PM_{2.5}
289 ($r=0.53$, $p<0.01$) which suggest the continuous and relative constant local sources were the main
290 sources. This result was similar with that reported from Taiyuan (Guo et al., 2016), where no
291 correlation was between the concentrations of OPEs and the concentration of particulate matter.
292 However, this result differed from that in Xinxiang (Shen et al., 2016), which showed that the
293 concentrations of OPEs and PM_{2.5} had significant correlation ($r=0.85c$), and a high value of
294 OPEs/PM_{2.5} was related to the contribution of the air mass from the heavily polluted area (Henan and
295 Jiangsu province), while low OPEs/PM_{2.5} was due to the air mass from Shanxi-Gansu and Neimenggu
296 Province. Chen et al. (2020) found there was a significant correlation ($p < 0.05$) between the
297 concentrations of Σ_{11} OPEs and PM_{2.5} in some sampling sites but not a site located in the urban region
298 in Guangzhou with potential additional pollution sources.

299 3.4.4 Correlation analysis of OPEs in PM_{2.5} and soil

300 Due to the low detection frequency of TCPP and TDCPP in the soil (Yin et al., 2016), the relationship
301 of other five OPE monomers in the soil and in atmospheric PM_{2.5} were presented in Fig. 6. A very
302 strong linear relationship was obtained between the OPEs in soil and in PM_{2.5} ($R^2 = 0.98$, $p<0.01$),
303 indicating that the atmospheric PM_{2.5} settlement is an important source of OPEs in the soil, so does the
304 soil be a source for OPEs in the air.

305 3.4.5 Correlation analysis of OPEs indoor and outdoor air

306 The OPEs profile in outdoor air in this study were: TBEP> TCEP > TnBP> TCPP > TPhP> TEHP>
307 TDCPP, which was different with indoor dust reported from our previous study (Liu et al., 2017):
308 TPhP>TCPP>TnBP> TDCPP >TBEP> TCEP > TEHP. TPhP is used as one of important alternatives
309 for technical decabrominated diphenyl ether (deca-BDE) product, which is typically used as a flame
310 retardant in electrical and electronic products. In addition, the use of plastic film and rubber may be an
311 important source of TPhP. Thus OPEs in indoor dust mainly comes from indoor environment and
312 related to human activities, not from outdoor air. Studies in Swedish (Wong, 2018) reported the



313 concentrations of OPEs in indoor air were TCPP > TCEP > TBEP > TnBP > TPhP, and in outdoor
314 urban air were TBEP > TCPP > TCEP > TnBP > TPhP (Wong, 2018) which also indicated the
315 differences of emission sources in indoor and outdoor air due to the different use of OPEs.

316 **3.5 Source apportionment of OPEs**

317 **3.5.1 Analysis of backward trajectory model**

318 The backward trajectory cluster analysis (HYSPLIT4) combines the horizontal and vertical motion of
319 the atmosphere, which can analyse the transport, migration and diffusion of atmospheric pollutants,
320 were used in this study. The height of AGL500m can better represent the characteristics of the process
321 wind field, and HYSPLIT4 was used to obtain the 24 hours backward trajectory of AGL500m during
322 the sampling period of Chengdu. During the sampling period, the air mass was mainly from the
323 northeastern and southern parts of Sichuan Province, including Mianyang, Deyang, Renshou and
324 Chengdu, and a few of the trajectories came from Chongqing and other places in Gansu Province.
325 Therefore, during the sampling period, Chengdu was mainly affected by the air mass of the eastern
326 Sichuan.

327 In different seasons, the air sources always came from the southern or the northern regions of Chengdu.
328 In spring, Chengdu was influenced by air mass from the southern region, which could be divided into
329 three paths: (a) from Ya'an through Renshou to Chengdu; (b) from Leshan and Yibin; and (c) from
330 Chongqing through Ziyang to Chengdu. The concentrations of OPEs at the northern and suburban site
331 were relatively high in spring. During the summer period, Chengdu was mainly influenced by air
332 masses from both the southern areas (Yibin, Zigong and others) and the northern areas (Gansu
333 Province, Guangyuan and Mianyang), but there was no significant difference in OPE concentrations at
334 each sampling site, nor in autumn and winter. Combined with the backward trajectory cluster analysis
335 and the concentrations of OPEs at each sampling site, the concentrations of OPEs had no obvious
336 change. This result suggested that OPEs were not affected by exogenous pollution but were mainly
337 affected by the local sources of Chengdu. These results are consistent with the meteorological and
338 topographic conditions. Chengdu's wind has always been breezy with much smaller strength than
339 coastal cities or other inland cities. The wind direction is relatively constant, mainly from the south and
340 the north. In addition, Chengdu is located in the basin, surrounded by the Qinghai-Tibet Plateau, the
341 Qinling Mountains, etc. These topographic and meteorological conditions block the influence of



342 foreign sources on Chengdu's atmosphere, which further explained that the pollution of OPEs in $PM_{2.5}$
343 was controlled by endogenous pollution, not by exogenous pollution.

344 **3.5.2 Principal Component Analysis**

345 The principal component analysis (PCA) of OPEs was carried out by SPSS. The normalized correlation
346 coefficient matrix of the original data of each sampling site showed that there was a strong correlation
347 between TCPP and TCEP, TCEP and TBEP, and TnBP and TPhP, which satisfied the condition of
348 dimensionality reduction of PCA. Two principal component factors were obtained in this study. The
349 cumulative contribution of the two principal component factors was 62.3%, which can basically reflect
350 the data information. The results were shown in Table S1. For factor 1, there was a large load on
351 TCEP, TCPP, TBEP and a moderate load on TDCPP. Factor 1 can represent the sources of OPEs
352 from the plastic industry, interior decoration and traffic emission, with the contribution ratio of 34.5%.
353 Factor 2 has higher load on TnBP, TEHP and TPhP. The highest load was on TnBP, which is often
354 used as a high-carbon alcohol defoamer, mostly in industries that do not come in contact with food and
355 cosmetics, as well as in antistatic agents and extractants of rare earth elements. TEHP can be used as an
356 antifoaming agent, hydraulic fluid and so on. TPhP is typically used in electrical and electronic
357 products, or plastic film and rubber. Factor 2 can be considered the chemical, mechanical and electrical
358 industry, and its contribution ratio was 27.8%.

359 **3.5.3 PMF model analysis**

360 The basic principle of the PMF method is to decompose the sample matrix into a factor contribution
361 matrix and factor component spectrum. The source type of the factor is judged according to the factor
362 component spectrum, and then the contribution ratio of source is determined. From 149 samples
363 collected in Chengdu, 132 valid samples were selected to participate in the model calculation and three
364 factors were determined. TPhP was the only chemical with residual (4.0) greater than 3. Concentrations
365 of OPEs satisfied the normal distribution. The components of factor 1 were complex. Factor 1
366 contributes 71.0%, 70.7% and 70.9% to TCEP, TCPP and TEHP, respectively, and 58.3% to TPhP.
367 Factor 1 was deduced to be the plastics/electrical industry and indoor source emissions. Factor 2
368 contributed the most to TBEP (78.0%), followed by TDCPP (44.7%), while it did not contribute to



369 TnBP. Therefore, factor 2 was deduced as the food/cosmetics industry and traffic emissions. Factor 3
370 contributes 71.7% of the total TnBP, which can be deduced as chemical industrial source.

371 **4. Conclusions and Implications**

372 Compared to levels of OPEs in other cities, the levels of OPEs measured in this study were comparable
373 or even higher than most of other studies. This suggests that during the shift of labour-intensive
374 manufacturing from the coastal developed areas to inland regions, OPEs were widely used in industrial
375 and manufacturing processes in southwest China which needs concern.

376 This intensive sampling campaign of urban and suburban area found no significant spatial variability of
377 Σ_7 OPEs across Chengdu, China, but the most notable seasonal variation was observed for alkyl
378 phosphate, followed by halogenated OPEs and aryl OPEs. Higher concentrations and more dispersed
379 pattern of OPEs in autumn/winter than that in summer, with TBEP, TCEP, TCPP and TnBP being the
380 dominant compounds. The occurrence of unexpected high level of individual OPEs at different sites in
381 autumn might indicate that there was a noteworthy emission. PCA analysis showed the main sources of
382 OPEs in $PM_{2.5}$ include plastic industry/interior decoration /traffic emission (34.5%) and chemical,
383 mechanical and electrical industry (27.8%). PMF showed the main sources were the plastics/electrical
384 industry and indoor source emissions. OPEs have a wide range of physical and chemical properties,
385 combined with differences in its behavior identified in this study, the management of OPEs as
386 individual compounds instead of a single chemical class should be considered. In addition, due to the
387 special topography and meteorological conditions of the inland city, the distribution and seasonal
388 variation of OPEs in the air in this study were significantly different from that of most coastal cities
389 and over the sea. The sustained and stable high local emissions are particularly noteworthy. The
390 chlorinated phosphate, especially TCPP and TCEP, which are highly toxic and not easy to degrade in
391 the environment, have a high content. Their usage and source emissions should be controlled.

392 **Acknowledgments**

393 We acknowledge financial support from National Natural Science Fund (41773072, 21407014,
394 41831285).



395 **References**

- 396 Araki, A., Saito, I., Kanazawa, A., Morimoto, K., Nakayama, K., Shibata, E., Tanaka, M., Takigawa,
397 T., Yoshimura, T., Chikara, H.: Phosphorus flame retardants in indoor dust and their relation to asthma
398 and allergies of inhabitants, *Indoor Air*, 24, 3-15, <https://doi.org/10.1111/ina.12054>, 2013.
- 399 Bacaloni, A., Cucci, F., Guarino, C., Nazzari, M., Samperi, R., Laganà, A.: Occurrence of
400 organophosphorus flame retardant and plasticizers in three volcanic lakes of central Italy, *Environ. Sci.*
401 *Technol.*, 42, 1898-1903, <https://doi.org/10.1021/es702549g>, 2008.
- 402 Cristale, J., Katsoyiannis, A., Sweetman, A. J., Jones, K. C., Lacorte, S.: 2013. Occurrence and risk
403 assessment of organophosphorus and brominated flame retardants in the River Aire (UK), *Environ.*
404 *Pollut.*, 179, 194-200. <https://doi.org/10.1016/j.envpol.2013.04.001>, 2013.
- 405 Cristale, J., Lacorte, S.: Development and validation of a multiresidue method for the analysis of
406 polybrominated diphenyl ethers, new brominated and organophosphorus flame retardants in sediment,
407 sludge and dust, *J. Chromatogr. A.*, 1305, 267-275, <https://doi.org/10.1016/j.chroma.2013.07.028>, 2013.
- 408 Covaci, A., Voorspoels, S., Ramos, L., Neels, H., Blust, R.: Recent developments in the analysis of
409 brominated flame retardants and brominated natural compounds, *J. Chromatogr. A.*, 1153, 145-171,
410 <https://doi.org/10.1016/j.chroma.2006.11.060>, 2007.
- 411 Chen, Y. Y., Song, Y. Y., Chen, Y. J., Zhang, Y. H., Li, R. J., Wang, Y. J., Qi, Z. H., Chen, F. H., Cai,
412 Z. W.: Contamination profiles and potential health risks of organophosphate flame retardants in PM_{2.5}
413 from Guangzhou and Taiyuan, China, *Environ. Int.*, 134, 105343,
414 <https://doi.org/10.1016/j.envint.2019.105343>, 2020.
- 415 Celano, R., Rodríguez, I., Cela, R., Rastrelli, L., Piccinelli, A. L.: Liquid chromatography quadrupole
416 time-of-flight mass spectrometry quantification and screening of organophosphate compounds in
417 sludge, *Talanta.*, 118, 312-320. <https://doi.org/10.1016/j.talanta.2013.10.024>, 2014.
- 418 Guo, Z. M., Liu, D., Shen, K.J., Li, J. Yu, Z.Q. Zhang, G.: Concentration and seasonal variation of
419 organophosphorus flame retardants in PM_{2.5} of Taiyuan City, China, *Earth and environment (in*
420 *chinese)*, 44, 600-604. <https://doi.org/10.14050/j.cnki.1672-9250.2016.06.002>, 2016.
- 421 Javier, C.J., Richard, S.: Atmospheric particle-bound organophosphate ester flame retardants and
422 plasticizers in a North African Mediterranean coastal city (Bizerte, Tunisia), *Sci. Total Environ.*, 642,
423 383-393. <https://doi.org/10.1016/j.scitotenv.2018.06.010>, 2018.



- 424 Kim, J. W., Isobe, T., Chang, K. H., Amano, A., Maneja, R. H., Zamora, P. B., Siringan, F.P., Tanabe,
425 S.: Levels and distribution of organophosphorus flame retardants and plasticizers in fishes from Manila
426 Bay, the Philippines. *Environ. Pollut.*, 159, 3653-3659. <https://doi.org/10.1016/j.envpol.2011.07.020>,
427 2011.
- 428 Kurtkarakus, P., Alegria, H., Birgul, A., Gungormus, E., Jantunen, L.: Organophosphate ester (opes)
429 flame retardants and plasticizers in air and soil from a highly industrialized city in Turkey, *Sci. Total*
430 *Environ.*, 625, 555-565, <https://doi.org/10.1016/j.scitotenv.2017.12.307>, 2017.
- 431 Li, J.: Occurrence and health risk assessment of organophosphate flame retardants in drinking water
432 and air. Nanjing University, 2014.
- 433 Liu, Q., Yin, H.L., Li, D., Deng, X., Fang, S.H., Sun, J.: Distribution characteristic of OPEs in indoor
434 dust and its health risk, China. *Environ. Sci.*, 37, 2831-2839. <https://doi.org/10.3969/j.issn.1000-6923.2017.08.004>, 2017.
- 436 Liu, R., Lin, Y., Liu, R., Hu, F., Ruan, T., Jiang, G.: Evaluation of two passive samplers for the
437 analysis of organophosphate esters in the ambient air, *Talanta.*, 147, 69-75,
438 <https://doi.org/10.1016/j.talanta.2015.09.034>, 2016.
- 439 Li, J., Xie, Z., Mi, W., Lai, S., Tian, C., Emeis, K.C.: Organophosphate esters in air, snow and seawater
440 in the north atlantic and the arctic, *Environ. Sci. Technol.*, 51, 6887-6896.
441 <https://doi.org/10.1021/acs.est.7b01289>, 2017.
- 442 Lai, S., Xie, Z., Song, T., Tang, J., Zhang, Y., Mi, W.: Occurrence and dry deposition of
443 organophosphate esters in atmospheric particles over the Northern South China Sea, *Chemosphere* 127,
444 195-200. <https://doi.org/10.1016/j.chemosphere.2015.02.015>, 2015.
- 445 Möller, A., Sturm, R., Xie, Z., Cai, M., He, J., Ebinghaus, R.: Organophosphorus flame retardants and
446 plasticizers in airborne particles over the northern pacific and Indian Ocean toward the polar regions:
447 evidence for global occurrence, *Environ. Sci. Technol.*, 46, 3127-3134. 2012
- 448 Marklund, A., Andersson, B., Haglund, P.: Traffic as a source of organophosphorus flame retardants
449 and plasticizers in snow, *Environ. Sci. Technol.*, 39, 3555-3562, <https://doi.org/10.1021/es0482177>,
450 2007.
- 451 Marklund, A., Barbro Andersson, A., Haglund, P.: Organophosphorus flame retardants and plasticizers
452 in Swedish sewage treatment plants, *Environ. Sci. Technol.*, 39, 7423-7429.
453 <https://doi.org/10.1021/es051013l>, 2005.



- 454 Matthews, H. B., Eustis, S. L., Haseman, J.: Toxicity and carcinogenicity of chronic exposure to tris(2-
455 chloroethyl) phosphate, *Fundam. Appl. Toxicol.*, 20, 477-485, <https://doi.org/10.1006/faat.1993.1058>,
456 1993.
- 457 Matthews, H. B., Dixon, D., Herr, D. W.: Sub-chronic toxicity studies indicate that tris (2-chloroethyl)
458 phosphate administration results in lesions in the rat hippocampus, *Toxicol. Ind. Health.*, 6, 1-15,
459 <https://doi.org/10.1177/074823379000600101>, 1990.
- 460 Ma, Y., Cui, K., Zeng, F., Wen, J., Liu, H., Zhu, F.: Microwave-assisted extraction combined with gel
461 permeation chromatography and silica gel cleanup followed by gas chromatography-mass spectrometry
462 for the determination of organophosphorus flame retardants and plasticizers in biological samples,
463 *Analytica. Chimica. Acta.*, 786, 47-53, <https://doi.org/10.1016/j.aca.2013.04.062>, 2013.
- 464 Ohura, T., Amagai, T., Senga, Y., Fusaya, M.: Organic air pollutants inside and outside residences in
465 Shimizu, Japan: levels, sources and risks, *Sci. Total. Environ.*, 366, 485-499,
466 <https://doi.org/10.1016/j.scitotenv.2005.10.005>, 2006.
- 467 Ren, G., Chen, Z., Feng, J., Ji, W., Zhang, J., Zheng, K.: Organophosphate esters in total suspended
468 particulates of an urban city in East China, *Chemosphere.*, 164, 75-83,
469 <https://doi.org/10.1016/j.chemosphere.2016.08.090>, 2016.
- 470 Salamova, A., Ma, Y., Venier, M., Hites, R.A.: High levels of organophosphate flame retardants in the
471 great lakes atmosphere, *Environ. Sci. Technol.*, 46, 8653-8660, <https://doi.org/10.1021/ez400034n>,
472 2014a.
- 473 Salamova, A., Hermanson, M.H., Hites, R.A.: Organophosphate and halogenated flame retardants in
474 atmospheric particles from a European Arctic site, *Environ. Sci. Technol.*, 48, 6133-40,
475 <https://doi.org/10.1021/es500911d>, 2014b.
- 476 Shoeib, M., Ahrens, L., Jantunen, L., Harner, T.: Concentrations in air of organobromine,
477 organochlorine and organophosphate flame retardants in Toronto, Canada, *Atmos. Environ.*, 99, 140-
478 147, <https://doi.org/10.1016/j.atmosenv.2014.09.040>, 2014.
- 479 Sühling, R., Diamond M. L., Scheringer, M., Fiona, Wong., Monika, Pu#ko., Gary. S., Alexis. B.,
480 Hayley. H., Philip. F., Henrik. L., Liisa. M.: Jantunen et al. Organophosphate Esters in Canadian Arctic
481 Air: Occurrence, Levels and Trends, *Enviro. Sci. Technol.*, *acs.est.6b00365*,
482 <https://doi.org/10.1021/acs.est.6b00365>, 2016.



- 483 Shen, K.J., Zhang, X.Y., Liu, D. Geng, X.F., Sun, J.H. Li, J.: Characterization and seasonal variation of
484 carbonaceous aerosol in urban atmosphere of a typical city in North China, *Ecology. Environ. Sci. (in*
485 *chinese)*, 25, 458-463, <https://doi.org/10.16258/j.cnki.1674-5906.2016.03.013>, 2016.
- 486 Tang, R., Keming, M.A., Zhang, Y., Mao, Q.: Health risk assessment of heavy metals of street dust in
487 Beijing, *Acta. Scientiae. Circumstantiae.*, 32, 2006-2015,
488 <https://doi.org/10.13671/j.hjkxxb.2012.08.029>, 2012.
- 489 Wang, X., He, Y., Li, L., Feng, Z., Luan, T.: Application of fully automatic hollow fiber liquid phase
490 microextraction to assess the distribution of organophosphate esters in the Pearl River Estuaries. *Sci.*
491 *Total. Environ.*, 470-471C, 263-269, <https://doi.org/10.1016/j.scitotenv.2013.09.069>, 2013.
- 492 Wang Y., Sun H., Zhu H., Yao Y.M., Chen H., Ren Chao., Wu F.C., Kannan Kurunthachalam.:
493 Occurrence and distribution of organophosphate flame retardants (OPFRs) in soil and outdoor settled
494 dust from a multi-waste recycling area in China, *Sci. Total. Environ.*, 625:1056,
495 <https://doi.org/10.1016/j.scitotenv.2018.01.013>, 2018.
- 496 Wang, Y., Bao. M. J., Tan. F., Qu. Z. P., Zhang. Y. W., Chen. J. W.: Distribution of organophosphate
497 esters between the gas phase and PM 2.5 in urban Dalian, China, *Environ. Pollut.*,
498 <https://doi.org/10.1016/j.envpol.2019.113882>, 2019.
- 499 Wang, T., Ding. N., Wang. T., Chen. S. J., Luo. X. J., Mai. B. X.: Organophosphorus esters (OPEs) in
500 PM_{2.5} in urban and e-waste recycling regions in southern China: concentrations, sources, and emissions,
501 <https://doi.org/10.1016/j.envres.2018.08.015>, 2018.
- 502 Wong, F., Wit, C.A.D, Newton, S.R.: Concentrations and variability of organophosphate esters,
503 halogenated flame retardants, and polybrominated diphenyl ethers in indoor and outdoor air in
504 Stockholm, Sweden, *Environ. Pollut.*, 240, 514-522, <https://doi.org/10.1016/j.envpol.2018.04.086>,
505 2018.
- 506 Yin, H.L., Wu, D., You, J.J., Li, S.P., Deng, X., Luo, Y., Zheng, W.Q.: Occurrence, Distribution, and
507 Exposure Risk of Organophosphate Esters in Street Dust from Chengdu, China, *Arch. Environ. Con.*
508 *Tox.*, 76(04), 617-629, <https://doi.org/10.1007/s00244-019-00602-3>, 2019.
- 509 Yin, H.L., Li, S.P., Ye, Z.X., Liang, J.F., You, J.J.: Pollution characteristics and sources of OPEs in the
510 soil of Chengdu City, *Acta. Scientiae. Circumstantiae.*, 36, 606-613,
511 <https://doi.org/10.13671/j.hjkxxb.2015.0489>, 2016.



- 512 Yin, H.L., Li, S.P., Ye, Z.X., Yang, Y.C., Liang, J.F., You, J.J.: Pollution Level and Sources of
513 Organic Phosphorus Esters in Airborne PM_{2.5} in Chengdu City, Environ. Sci. (in chinese), 36, 3566-
514 3572, <https://doi.org/10.13227/j.hjcx.2015.10.003>, 2015.
- 515 Zhang, Q. H., Yang, W. N., Ngo, H. H., Guo, W. S., Jin, P. K., Dzakpasu, M.: Current status of urban
516 wastewater treatment plants in China, Environ. Int., 92-93, 11-22,
517 <https://doi.org/10.1016/j.envint.2016.03.024>, 2016.



518 Figure Captions:

519 Fig.1. Levels and seasonal variation of Σ_7 OPEs at each sampling site. A:autumn, W:winter, Sp:spring,

520 Su:summer, Sub:suburbs, Dow:downtown, S:south, E:east, N:north, W:west.

521 Fig. 2 Seasonal variation of Σ_7 OPEs at each sampling site.

522 Fig.3 The seasonal variation of monomer OPEs in Chengdu city. A:Autumn, W:Winter, Sp:Spring,

523 Su:Summer, Sub:Suburbs, Dow:Downtown, S:South, E:East,N:North, W:West

524 Fig.4 Relationship of OPE monomer concentration in $PM_{2.5}$ and its vapor pressure

525 Fig. 5 Spearman's ranks correlation coefficients between the concentrations of individual OPEs in

526 $PM_{2.5}$ samples

527 Fig. 6 Relationship between OPEs in atmospheric $PM_{2.5}$ and in soil.

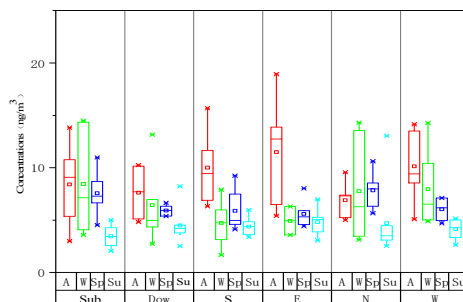
528 Table Captions:

529 Table 1 Table 1 The annual median concentrations of OPEs in $PM_{2.5}$ form Chengdu ($ng\ m^{-3}$).

530 Table 2 The correlation analysis of monomer OPEs in downtown and suburb sampling sites.

531 *. Correlation is significant at the 0.05 level (2-tailed).

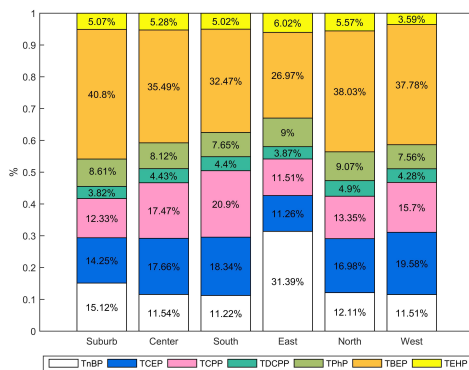
532 **. Correlation is significant at the 0.01 level (2-tailed).



533

534 **Fig.1** Levels and seasonal variation of $\Sigma 7\text{OPEs}$ at each sampling site. A:autumn, W:winter, Sp:spring,

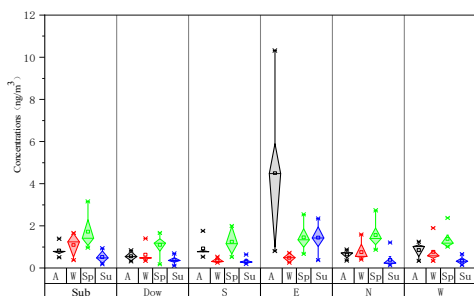
535 Su:summer, Sub:suburbs, Dow:downtown, S:south, E:east, N:north, W:west.



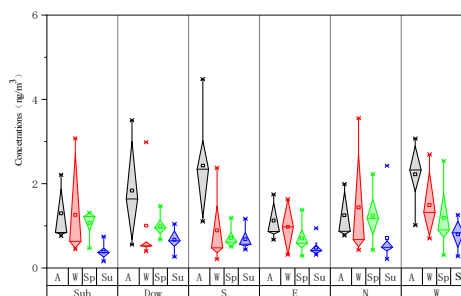
536

537 **Fig.2** Seasonal variation of $\Sigma 7\text{OPEs}$ at each sampling site.

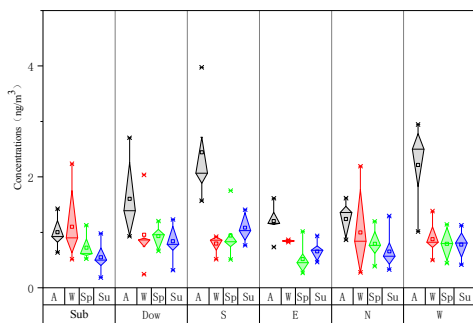
538



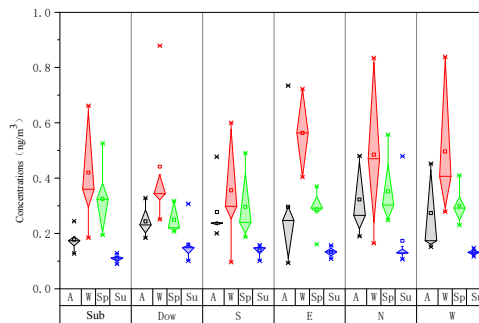
TnBP



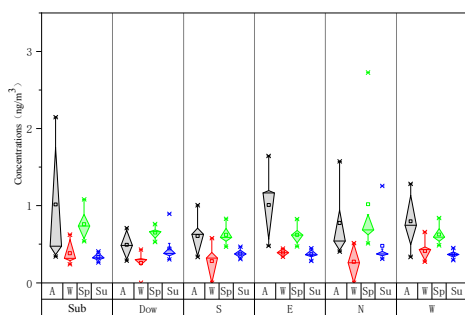
TCEP



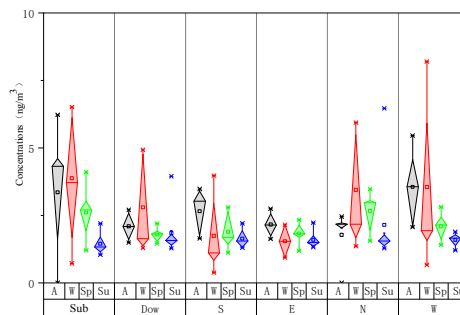
TCPP



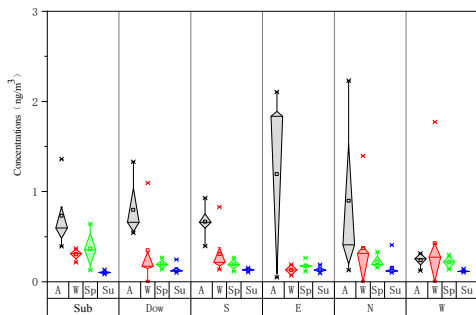
TDCPP



TPhP



TBEP

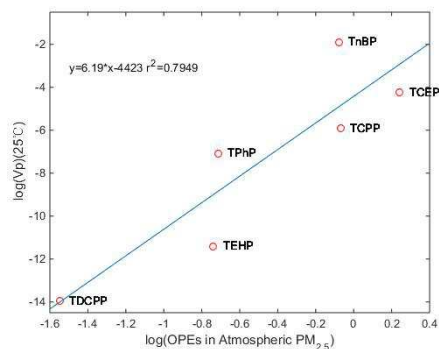


TEHP



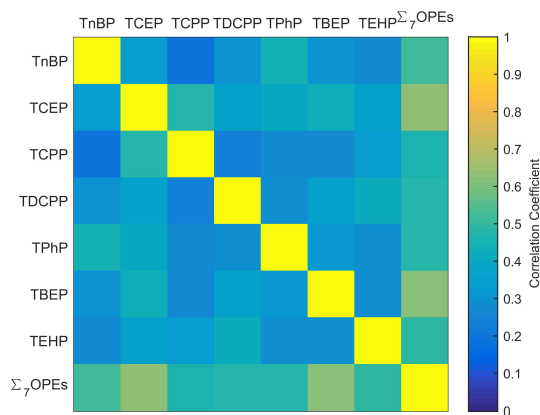
539 **Fig.3** The seasonal variation of monomer OPEs in Chengdu city. A:Autumn, W:Winter, Sp:Spring,

540 **Su:Summer, Sub:Suburbs, Dow:Downtown, S:South, E:East,N:North, W:West.**



541

542 **Fig.4** Relationship of OPE monomer concentration in PM2.5 and its vapor pressure

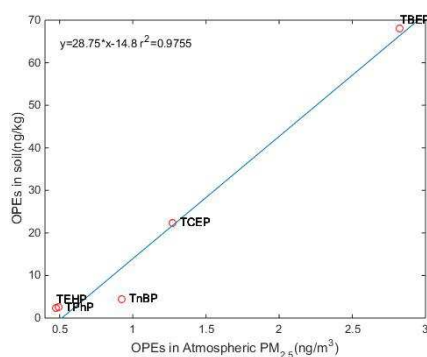


543



544 Fig.5 Spearman's ranks correlation coefficients between the concentrations of individual OPEs in PM2.5

545 samples



546

547 Fig.6 Relationship between OPEs in atmospheric PM2.5 and in soil.

548

549 Table 1 The annual median concentrations of OPEs in PM2.5 from Chengdu (ng m⁻³).

Orientation	TnBP	TCEP	TCPP	TDCPP	TPhP	TBEP	TEHP	Σ ₇ OPEs
suburb	1.0	1.0	0.8	0.3	0.6	2.7	0.3	6.7
downtown	0.7	1.0	1.0	0.3	0.5	2.1	0.3	5.8
south	0.7	1.1	1.2	0.3	0.5	1.9	0.3	5.9
east	2.1	0.8	0.8	0.3	0.6	1.8	0.4	6.6
north	0.8	1.1	0.9	0.3	0.6	2.5	0.4	6.7
west	0.8	1.4	1.1	0.3	0.5	2.6	0.3	6.9
median	1.0	1.1	1.0	0.3	0.5	2.3	0.3	6.4

550

551 Table 2 The correlation analysis of monomer OPEs in downtown and suburb sampling sites.

552

	TnBP	TCEP	TCPP	TDCPP	TPhP	TBEP	TEHP
Downtown	TnBP	1	.408*	0.319	0.15	.455*	0.105
	TCEP	.408*	1	.818**	0.165	.342	.447*
	TCPP	0.319	.818**	1	0.184	.392	.500*
	TDCPP	0.15	0.165	0.184	1	0.053	0.175
	TPhP	.455*	0.342	0.392	0.053	1	-0.081



	TBEP	0.187	.447*	.447*	0.216	0.104	1	0.338
	TEHP	0.105	.449*	.500*	0.175	-0.081	0.338	1
Suburb	TnBP	1	.566**	.476*	.650**	0.269	.417*	0.141
	TCEP	.566**	1	.852**	.683**	0.368	.784**	.423*
	TCPP	.476*	.852**	1	.686**	0.304	.701**	0.297
	TDCPP	.650**	.683**	.686**	1	0.175	.708**	0.158
	TPhP	0.269	0.368	0.304	0.175	1	.512**	.629**
	TBEP	.417*	.784**	.701**	.708**	.512**	1	.434*
	TEHP	0.141	.423*	0.297	0.158	.629**	.434*	1

553 *. Correlation is significant at the 0.05 level (2-tailed).

554 **. Correlation is significant at the 0.01 level (2-tailed).