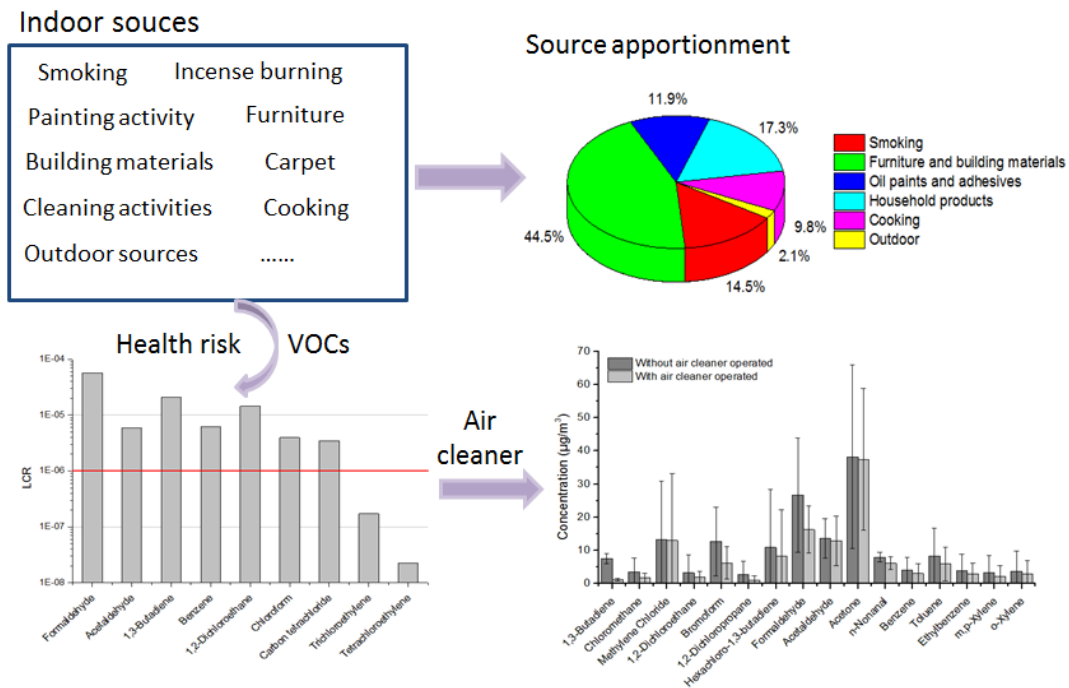


The following publication Huang, Y., Su, T., Wang, L., Wang, N., Xue, Y., Dai, W., ... & Ho, S. S. H. (2019). Evaluation and characterization of volatile air toxics indoors in a heavy polluted city of northwestern China in wintertime. *Science of the Total Environment*, 662, 470-480 is available at <https://doi.org/10.1016/j.scitotenv.2019.01.250>.



Highlights

- Hazardous VOCs and carbonyls were evaluated in typical dwellings in northwestern China
- High levels of the pollutants were associated with characteristic pollution sources
- Paints, adhesives, decoration, and household products are major indoor pollution contributors
- Cancer risks for formaldehyde, 1,3-butadiene and 1,2-dichloroethane exceeded acceptable level
- Household air cleaner can be efficiently reduced pollutant levels in residential airs

1 **Abstract**

2 Hazardous volatile organic compounds (VOCs) and carbonyls were evaluated in typical
3 dwellings in Xi'an in northwestern China in wintertime. High indoor concentrations were
4 observed for formaldehyde, acetone, naphthalene, methylene chloride and acetaldehyde,
5 associated with characteristic pollution sources. In comparison, many of the target VOCs
6 were higher in Chinese dwellings than those in other countries, suggesting the significances
7 of indoor pollutions in China. Source apportionment with receptor model shows that
8 furniture and building materials (44.5%), paints and adhesives (11.9%), household products
9 (17.3%), smoking (14.5%), and cooking (9.8%) are the major contributors to the indoor
10 VOCs and carbonyls. The health risk assessment shows that the cancer risks for
11 formaldehyde (5.73×10^{-5}), 1,3-butadiene (2.07×10^{-5}) and 1,2-dichloroethane (1.44×10^{-5})
12 were much higher than the acceptable level of 1×10^{-6} recommended by International
13 Register for Certified Auditors (IRCA). The hazard quotient (HQ) of target VOCs were far
14 less than the threshold (HQ=1). Moreover, the practical efficiency of household air purifier
15 in removal of the VOCs and carbonyls was examined first time in dwellings in northern
16 China. The results prove that most of the indoor organic pollutants and their cancer risk to
17 humans can be efficiently reduced, particularly for formaldehyde and 1,3-butadiene. The
18 findings of the study offer useful preliminary and updated information on current indoor air
19 toxics levels, dominant pollution sources and their potential health risks to residents in
20 northwest China.

21

22 **Keywords:** *indoor air quality, volatile air toxics (VOCs), Carbonyls, residence, health risk,*

24 **1. Introduction**

25 The importance of indoor air quality (IAQ) alerts public in recent decades
26 world-widely. Surveys show that people spend ~80-90% of their daily time indoors on
27 average (Leech et al., 2002). Volatile organic compounds (VOCs) is one of major class of
28 pollutants due to their ubiquity in indoor environments, significantly impacting on human
29 health. Exposure to hazardous volatile air toxics has potential carcinogenic and other
30 toxicological effects, while long-term exposure can harm the respiratory, neurological and
31 reproductive system, or even lead to death (Delfino, 2002; Windham et al., 2006; Wu et al.,
32 2007).

33 Indoor VOCs and carbonyls can be originated from a variety of sources. Household
34 products were the major contributor (44%), followed by combustion processes and
35 environmental tobacco smoke (ETS) (10.5%), deodorizers (8.4%) and off-gassing of
36 building materials (5.9%) in residences of Edmonton, Alberta (Bari et al., 2015). Seasonal
37 variation on VOCs levels could be seen indoors. Higher indoor levels of alkanes, alkenes,
38 aromatics were reported due to low ventilation rates in heating period (Duan et al., 2014).
39 Pekey et al. (2008) also found that most quantified VOCs had higher concentrations in
40 winter than summer in Turkey (Pekey and Arslanbaş, 2008). The winter values could be
41 even double of those in summer in Edmonton, Canada (Bari et al., 2015). The indoor air
42 can be greatly impacted by coal combustion and biomass burning when household
43 warming is required (Abeleira and Farmer, 2017; Duan et al., 2014). However, in winter,
44 indoor and outdoor air exchange efficiency is much poorer than other seasons, leading to
45 the accumulation of pollutants indoors.

46 A lot of studies have been conducted to screen and measure those priority toxic VOCs as
47 well as to assess their health-related potentials indoors in China. Duan et al., (2014) quantified
48 nearly one hundred VOCs to obtain the seasonal variations, indoor and outdoor
49 relationships, and potential sources at residential units in Beijing, China (Duan et al., 2014).
50 The results showed that formaldehyde, acetone, acetaldehyde, toluene, ethane and propane
51 were the most dominant indoor airborne organic species. Wang et al. (2007) measured
52 carbonyls simultaneously in twelve urban dwellings in Chinese megacities including
53 Beijing, Shanghai, Guangzhou, and Xi'an (Wang et al., 2007). Formaldehyde was the most
54 abundant compound, accounting for ~46.0% of the quantified carbonyls and ranging from
55 the lowest of $19.3 \mu\text{g}/\text{m}^3$ in Xi'an to the highest of $92.8 \mu\text{g}/\text{m}^3$ in Beijing during summer.

56 Higher indoor VOCs and carbonyls levels are always seen in China than other
57 countries. Their concentrations and composition can be varied by interior decorations,
58 activities, ventilations and locations. Guo et al. (2009) conducted a comprehensive study at
59 100 homes in Hong Kong, reporting that the total VOC and formaldehyde concentration
60 was $46.1 \pm 8.8 \mu\text{g}/\text{m}^3$ and $112.3 \pm 9.5 \mu\text{g}/\text{m}^3$, respectively, much higher than other East Asian
61 cities (Guo et al., 2009) . Furthermore, higher levels of 1,2,4-trimethylbenzene, styrene,
62 nonane and heptane were found in gas-use families rather than in electricity-use homes in
63 their study.

64 Due to the importance of VOCs, Du et al. (Du et al., 2014) accessed sixteen highly
65 prevalent Hazardous Air Pollutants (HAPs) in urban cities in China and reported the
66 average total lifetime cancer risks attributable to HAPs are 2.27×10^{-4} and 2.93×10^{-4} for
67 Chinese females and males, respectively. Over 70% of the risk was found due to exposure

68 to indoor air at home and formaldehyde, 1,4-dichlorobenzene, benzene and 1,3-butadiene
69 are the major contributors to health hazard.

70 Xi'an (33°N and 107°E) is a key city in the northwest China and the capital of Shaanxi
71 province. With the supports by the national policies, it has been rapidly developing since
72 1980's. The growth economy elevates the living standard and also alert residents to
73 concern their health regarding air pollutions. To our best knowledge, there is still a lack of
74 comprehensive study to evaluate both VOCs and carbonyls in dwellings in the northwest
75 China. The objectives of this work are to compare indoor and outdoor VOC levels, to
76 explore the potential effects of VOC levels indoor and to quantify exposure risks. This
77 study was designed to cover as many compounds as possible under the premise of
78 experimental condition, because of the lack of VOCs data in Xi'an residence.

79 **2. Methodology**

80 **2.1. Sampling locations**

81 Eleven dwellings in Yanta, Weiyang, Xincheng and Yanliang districts were selected
82 in this study (Fig. 1). The locations represent typical residential areas in urban and
83 suburban Xi'an where the residents are concentrated. The sampling campaign was
84 conducted from mid-November 2016 to mid-February 2017 during the regular regional
85 heating supply period. The average ambient temperature was -1 ± 5 °C. All selected
86 dwellings have not been renovated in the past three years. No particular pollution sources
87 (e.g., industrial sector or power plant) were near the sampling areas.

88 **2.2. Sample collection**

89 Indoor and outdoor samples were collected simultaneously. For indoor, samplers were
90 placed in the center of living room with an inlet height of 1.5 m above the floor. The living
91 room is the center of the room and the place where people undertakes most activities. All
92 doors and windows were closed when the sampling conducted. The sampling time was
93 between 09:00-11:00 when the impact from household cooking was minimized. Additional
94 comparison tests were carried out indoors on the days when an in-house air purifier was
95 operating in each dwelling. The air purifier had worked for 9 h before the first sample was
96 collected. It has operated continuously for four consecutive days (96h) while the samples
97 were collected daily. Other sampling conditions were the same as before. Two sets of four
98 indoor samples were collected in each dwelling when the air purifier was on or off
99 respectively. Outdoor samples were collected spontaneously on the balcony by extending
100 the sampling tubes outside when indoor sampling was conducted.

101 A total of sixty-five VOCs classified as “Air Toxics” by United States Environmental
102 Protection Department (VOC_{Toxic}) (USEPA, 1999b) and seventeen carbonyls (including
103 mono- and di-carbonyls) were quantified in this study. The VOC_{Toxic} was collected into a
104 stainless-steel multi-bed adsorbent tube filled with Tenax-TA, Carbograph I TD and
105 Carboxen 1003 (C3-DXXX-5266, ¼” o.d., Markes International Ltd., Llantrisant, U.K.)
106 using a low-flow module pump (ACTI-VOC, Markes International Ltd.). The sampling
107 flow rate was 50 mL/min and each sample was collected for 120 min. Prior to the sampling,
108 the sorbent tubes were thermally cleaned in a conditioner (TC20, Markes International Ltd.)
109 at 330 °C for 20 min. The pre-conditioned and sampled tubes were sealed with Difok caps

110 (Markes International Ltd.) and stored in pollutant-free desiccators at -4 °C for a maximum
111 of 14 days. The pump was calibrated with a mass flow calibrator (Defender 510, Bios,
112 Torrance, CA, USA) before and after each sampling event. A Teflon filter assembly (47
113 mm, Whatman, Clifton, NJ, USA) and coiled potassium iodide (KI)-coated copper tubing
114 (¼" o.d., 1 m in length) were installed in upstream to remove particle and ozone (O₃)
115 influences, respectively (Ho et al., 2017, 2018).

116 The carbonyls were collected into silica cartridges impregnated with acidified
117 2,4-dinitrophenylhydrazine (DNPH) (Sep-Pak DNPH-silica, 55-105 µm particle size, 125
118 Å pore size; Waters Corporation, Milford, MA) at a flow rate of 0.6 L/min for 120 min
119 (USEPA, 1999a). Detailed sampling procedures were shown in our previous publications
120 (Spaulding et al., 1999) (Ho et al., 2011).

121 An absorbent tube and a cartridge were reserved to serve as field blanks on each
122 sampling trip and were handled in the same way as the samples. The amounts of target
123 compounds were corrected for the field blank. All samples were shipped and stored in a
124 refrigerator at < 4 °C until the chemical analyses.

125 **2.3. Analytical methods**

126 The absorbent tubes for collection of VOC_{Toxic} were analyzed using a thermal
127 desorption (TD) unit (Series 2 UNITY-xr system, Markes International Ltd.) coupled with
128 a gas chromatograph/mass spectrometric detector (GC/MSD, Models 7890A/5977 B,
129 Agilent, Santa Clara, CA, USA). A tube was connected into the TD unit at room
130 temperature (~25 °C) and purged with ultra-high purity (UHP) helium (He) gas at a flow
131 rate of 40 mL/min for 10 s to eliminate air and oxygen intrusion. For the primary

132 desorption stage, the analytes were desorbed at 330 °C for 5 min and refocused onto a
133 cryogenic-trap (U-T1703P-2S, Markes International Ltd.) to capture high volatility target
134 compounds at 15 °C. For the secondary desorption stage, the trap was dry-purged for 10 s
135 and rapidly heated from 15 °C to 320 °C and maintained for 5 min. The analytes were
136 passed via a heated transfer line at 160 °C, and re-refocused onto a cold GC capillary
137 column head (Rtx®-1, 105 m0.25 mm × 1 µm film thickness, Restek Corporation,
138 Bellefonte, PA, USA) at -45 °C with an aid of liquid nitrogen (N₂) in GC oven. Once the
139 second desorption is completed, the oven temperature program started at an initial
140 temperature of -45 °C for 4 min, ramped to 230 °C at a rate of 6 °C /min, and maintained at
141 230 °C for 5 min. The constant flow rate of helium carrier gas was 1.0 mL/min throughout
142 the GC analysis. The MSD was operated in selective ion monitoring (SIM) mode at 230 °C
143 and 70 eV for electron ionization. Identification was achieved by comparing the mass
144 spectra and retention times of the chromatographic peaks with those of authentic standards.
145 Certified Air Toxics standard mixtures (Restek Corporation) were used in calibrations. A
146 multi-point calibration curve was established to quantify each of the target compounds with
147 linearity>0.995. The minimum detection limits (MDL) were in the range of 0.1-0.158 ppbv
148 with a sampling volume of 6 L. The measurement precision for the analysis of eight
149 replicates of standard samples at 2 ppbv were <25%.

150 The carbonyls in DNPH-silica were eluted with acetone-free acetonitrile (ACN) and
151 the extract was injected into a high-pressure liquid chromatography (HPLC) system (1200;
152 Agilent Technology) equipped with a photodiode array detector (DAD). Details on
153 extraction, calibration, and chromatographic conditions were shown elsewhere (Dai et al.,

154 2012). The limit of detections (LOD) of the target carbonyls ranged from 0.002 to 0.010
155 $\mu\text{g/mL}$.

156 **2.4. Questionnaire**

157 Information of selected dwellings characteristics and potential sources for VOCs and
158 carbonyls were obtained from site investigation and self-administered questionnaire (Table
159 S1). It included details of room description (i.e., area, age, type of wall, refurbishment, and
160 pet), ventilation and heating systems, frequency and fuel of cooking, smoking activities,
161 cleaning activities (detergent and frequency). The occupants were further interviewed on
162 their other daily activities to identify any additional potential exposure to the target
163 compounds as show in Table 1.

164 **2.5. Positive matrix factorization (PMF) receptor model**

165 Positive matrix factorization (PMF) (U.S.EPA, PMF3.0) receptor model was applied
166 to distinguish dominant sources in the indoor environments (Mj et al., 2002). The PMF
167 model can be expressed as a chemical mass balance equation in terms of contributions from
168 p independent sources to n chemical species measured in a given sample (Miller et al.,
169 1972):

$$170 \quad x_{ij} = \sum_{k=1}^p g_{ik} f_{kj} + e_{ij} \quad (1)$$

171 Where x_{ij} is the j th chemical species concentration determined in the i th sample,
172 g_{ik} is the species contribution of the k th source to the i th sample, f_{kj} is the loading of j th
173 species on the k th factor, e_{ij} is the residual resulting from bias in the measurement of g_{ik}
174 and f_{kj} , and p represent the total number of independent sources (Paatero, 1997). Every
175 data point can be individually weighed in PMF, so that the retainment of data below

176 detection limit with its associated uncertainty was permissible. The stability of the solution
177 can be evaluated by means of examining the proportion of each source undertaken in terms
178 of the object function Q:

$$179 \quad Q = \sum_{t=1}^n \sum_{i=1}^m \left[\frac{x_{ij} - \sum_{k=1}^p g_{ik} f_{kj}}{\mu_{ij}} \right]^2 \quad (2)$$

180 where μ_{ij} represents the uncertainty of j th species in i th sample. For the PMF input, the
181 uncertainty caused by sampling and analytical errors was calculated using the following
182 equation suggested by Polissar et al. (Polissar et al., 1998):

$$183 \quad U = \sqrt{(EF \times conc)^2 + (MDL)^2} (conc > MDL) \quad (3)$$

184 where EF represent the error fraction, which is the result of the relative standard
185 deviations of the instrument multiply 100, and we set it as 0.10, equal to the average
186 percent uncertainty in our study. For values below detection limit, the uncertainties were
187 replaced by 5/6 times of the detection limit values. Any missing data is replaced with the
188 median concentration of that species and the uncertainty are expressed as four times the
189 median concentration (USEPA, 2008). For the selection of chemical species, typical tracers
190 of different sources and those in high indoor concentrations were taken into account in the
191 receptor modeling. In addition, those species with more than 50% of samples below LODs
192 were screened out.

193 **2.6. Cancer and non-cancer risk calculation model**

194 The risk characterization for indoor VOC inhalation exposure was conducted by
195 combining published toxicity data with the exposure concentrations estimated in this study.
196 To calculate inhalation risks, an adjusted air concentration (EC_i) was calculated using the
197 following equation according to United States Environmental Protection Agency (U.S.EPA)

198 Superfund program (EPA, 2009; Waste, 1991).

$$199 \quad EC_i = \frac{CA_i \times ET \times EF \times ED}{AT} \quad (4)$$

200 Where EC_i is the exposure concentration; CA_i is the measured VOC concentration
201 in the residences ($\mu\text{g}/\text{m}^3$); ET is exposure time (hours/day); EF is the exposure frequency
202 (days/year); ED is exposure duration (years); AT is averaging time (hours). For cancer
203 and chronic hazard assessments, lifetime (70 years) is substituted for AT (lifetime in
204 years \times 365 days/year \times 24 hours/day).

205 We adjusted exposure air concentration by incorporating time-activity data of Chinese
206 residents. Based on Exposure Factors Handbook of Chinese Population (Duan, 2013) and
207 Wang et al.'s (Wang et al., 2012) study, average exposure time was estimated as 15 h/day
208 for residents (Dai et al., 2017). Exposure frequency was estimated as 350 day/year,
209 exposure duration was estimated as 24 years for adults to calculate inhalation cancer risk
210 attributable to indoor VOCs. The indoor inhalation cancer risk at residences was calculated
211 with the methodology proposed by US EPA (USEPA, 2004).

$$212 \quad LCR_i = IUR_i \times EC_i \quad (5)$$

213 Where LCR_i is the cancer risk associated with compound i ; EC_i the daily average
214 inhaled concentration of compound i ; and IUR_i is the estimated inhalation unit risk ($\text{m}^3/\mu\text{g}$)
215 for compound i from U.S.EPA, IRIS (Integrated Risk Information System) or OEHHA
216 (Office of Environment Health Hazard Assessment), which is the excess lifetime cancer
217 risk estimated to result from continuous exposure to an individual VOC via inhalation per
218 $\mu\text{g}/\text{m}^3$.

219 Non-cancer inhalation health impacts were assessed by a direct comparison of the

220 average personal exposure (EC_i) with a substance specific RfC . The hazard quotient (HQ)
221 of each compound was calculated:

$$222 \quad HQ_i = \frac{EC_i}{RfC_i} \quad (6)$$

223 Where HQ_i is the hazard quotient for compound i ; EC_i is the modeled personal
224 exposure to compound i ; and RfC_i is the reference exposure limits for compound i .

225 The results of LCR_i and HQ_i were judged according to U.S.EPA's (2004) approach.
226 Namely, cancer risks no higher than 1×10^{-6} for an “ample margin of safety” and an HQ
227 value of one or less indicates that adverse health effects are not expected to result from
228 exposure to this VOC.

229

230 **3. Results and discussion**

231 **3.1. Characteristics of VOCs and carbonyls**

232 **3.1.1. Indoor and outdoor levels of VOC_{Toxic} and carbonyls**

233 The average concentrations of VOC_{Toxic} and quantified carbonyls indoors and
234 outdoors are listed in Table 2. For the indoors, acetone ($35.55 \pm 24.34 \mu\text{g}/\text{m}^3$) was the most
235 dominant species, followed by formaldehyde ($21.45 \pm 13.72 \mu\text{g}/\text{m}^3$), naphthalene
236 ($16.64 \pm 18.96 \mu\text{g}/\text{m}^3$), methylene chloride ($13.13 \pm 18.64 \mu\text{g}/\text{m}^3$), acetaldehyde (12.92 ± 6.36
237 $\mu\text{g}/\text{m}^3$), hexachloro-1,3-butadiene ($9.41 \pm 15.78 \mu\text{g}/\text{m}^3$), bromoform ($9.21 \pm 8.16 \mu\text{g}/\text{m}^3$),
238 toluene ($7.23 \pm 7.00 \mu\text{g}/\text{m}^3$), n-Nonanal ($6.86 \pm 2.02 \mu\text{g}/\text{m}^3$), methyl butyl ketone (5.45 ± 8.98
239 $\mu\text{g}/\text{m}^3$) and ethyl acetate ($4.59 \pm 3.67 \mu\text{g}/\text{m}^3$). The concentrations of individual target
240 compounds in each dwelling are shown in supporting information (Table S2). The
241 concentrations are associated with specific situations (e.g., size, design, and ventilation rate)

242 and indoor activities. The common indoor sources are known as paints, adhesives, synthetic
243 fragrances and cigarette smoke (Guo et al., 2003; Polzin et al., 2007).

244 Owing to the uniqueness of each dwelling (e.g., size, design, and ventilation rate), it is
245 more appropriate to present the proportion instead of absolute concentration (Fig 2).
246 Acetone had the highest mass proportion of 10-25%. The range is consistent among the
247 dwellings except an extremely high value of 55% at Site 6. Acetone is used as solvent and
248 widely present in many household products (Wang et al., 2007). High proportions of
249 methylene chloride (7-20%) were also found in many dwellings such as Site 2, 4 and 6
250 where occupants frequently conducted cleaning activities with detergents. Methylene
251 chloride is a propellant to form aerosols while spraying (Health et al., 2000). The indoor
252 level of methylene chloride is thus linked with the application of spray products. These
253 could be supported by our results that acetone and methylene chloride were in high
254 proportions at Site 1 and 6 where insecticide had been used (Table 1). The proportion of
255 methylene chloride in Site 1 and 3 were also up to 10% which occupants are not frequently
256 conducted cleaning activities with detergents (1-3 times per week). Methylene chloride is a
257 powerful solvent that often be used as active ingredient in most paint strippers and foaming
258 agent (Riley et al., 2000), so the high proportion of methylene chloride may also from the
259 volatilization of furniture and building materials. Furthermore, high proportion of
260 naphthalene (0-16%) may be related to the use of mothballs but there were difficulties to
261 statistically record their usages in each dwelling in this study (JOWan-Kuen et al., 2008).

262 Formaldehyde widely presents in paints, adhesives, synthetic fragrances and cigarette
263 smoke (Guo et al., 2003; Polzin et al., 2007). The mass proportion of formaldehyde were

264 not greatly varied (5-15%) but high, implying that there were consistent and rich sources in
265 those dwellings in northern China (Salthammer et al., 2010). Wang et al. (2007) revealed
266 that building materials and some combustion activities including tobacco smoke and
267 incense burning are the contributors for indoor carbonyls.

268 Moderate compositions of BTEX (i.e., benzene, toluene, ethylbenzene, m/p-xylene
269 and o-xylene) and styrene (1-20% in total) were shown in most dwellings. Their
270 proportions were up to 20% at Sites 1 and 4, where the occupants were smokers (Table 1).
271 It was reported that tobacco smoking could emit different degrees of benzene, toluene and
272 m,p-xylene (Lee et al., 2002). Besides, BTEX and styrene can be produced in combustion
273 processes, fuel evaporative losses, and uses of solvents (Buczynska et al., 2009; Ilgen et al.,
274 2001). In general, aromatic VOCs were often found higher in China than other countries
275 (Ohura et al., 2009).

276 For the outdoors, the highest concentration was also seen for acetone (26.92 ± 28.09
277 $\mu\text{g}/\text{m}^3$), followed by methylene chloride (18.39 ± 30.48 $\mu\text{g}/\text{m}^3$), methyl butyl ketone
278 (10.80 ± 36.77 $\mu\text{g}/\text{m}^3$). Formaldehyde (8.53 ± 7.94 $\mu\text{g}/\text{m}^3$) and acetaldehyde (7.33 ± 4.54
279 $\mu\text{g}/\text{m}^3$) were also abundant in ambient air as they can be primarily formed in any
280 combustions (i.e., vehicle emission and industrial activities) and secondarily formed
281 through photochemical reactions (Possanzini et al., 2002). Similar explanation was applied
282 in explanation for high abundances of other compounds such as propylene (4.54 ± 6.28
283 $\mu\text{g}/\text{m}^3$), toluene (4.22 ± 4.08 $\mu\text{g}/\text{m}^3$), 1,3-butadiene (3.89 ± 0.64 $\mu\text{g}/\text{m}^3$) and benzene
284 (3.01 ± 2.66 $\mu\text{g}/\text{m}^3$) as well (Li et al., 2017a; Xue et al., 2017). Liu et al. (Liu, 2014)
285 evaluated the carbonyls and BTEX levels in indoor air at 128 residential homes in Beijing,

286 China, and found four major pollution contributors including outdoor incursion, building
287 materials and paints, particle board and plywood, and household cleaning chemicals.
288 Considering that the air can be exchanged physically, the indoor levels can be thus
289 impacted by the outdoor sources particularly during the frequent pollution episodes in
290 northern China.

291 **3.1.2. Comparison of indoor levels with other studies**

292 Table 3 compares the average indoor concentrations of selected compounds between
293 our values and the findings in other literatures. The concentration of formaldehyde was
294 higher in our study than the average value in Japan, but ~50% lower than that report in
295 Beijing. Formaldehyde is mainly released to indoor air from refurbishment materials,
296 wood-based products, flooring materials, smoking, and any indoor combustion
297 (Salthammer et al., 2010). The variations of the indoor levels between the cities could be
298 attributed with the differences of materials used and living styles (i.e., smoking, cleaning,
299 cooking). In addition, formaldehyde is the only carbonyl incorporated in the Chinese
300 national indoor air quality standard. It was found that the concentrations of formaldehyde
301 in all dwellings were well below the standard of $100 \mu\text{g}/\text{m}^3$ on 1-h average (GB/T, 2002).
302 Even though the concentrations in the dwellings were below the standard, the high
303 abundances must be an alarm to the Chinese residents. The level of acetaldehyde in our
304 study was comparative or lower than those in Beijing and Japan, but not for acetone. High
305 abundances of acetone in Xi'an potentially associated with more solvents used such as
306 bleach, laundry detergent, laundry stain remover, floor glue, nail color remover, oil paint,
307 and furniture polish (Kwon et al., 2007). For methylene chloride and chloroform, our

308 values are much lower than the newly renovated apartment in Shanghai and Hong Kong.

309 For aromatics, the concentrations of benzene, toluene, ethylbenzene, m/p-xylene and
310 o-xylene (BTEX) and styrene in the dwellings in Xi'an were relatively low. This could be
311 explained by less or no smoking and cooking activities had been conducted in the
312 residences during the sampling period. In addition, the concentrations of BTEX were much
313 lower than the newly houses renovated in Shanghai, indicating that less evaporation or
314 releases of solvents from the aged walls and furniture.

315 **3.2. Indoor source identification**

316 **3.2.1. Indoor and outdoor ratios**

317 Indoor to outdoor (I/O) ratio is a strong evidence to identify whether indoor or outdoor
318 sources may play decisive roles in IAQ. The I/O ratios of VOC_{Toxic} and carbonyls with air
319 purifier operated and after the use of air purifier in this study were presented in Fig. 3.
320 Higher I/O values (> 1) represent dominant indoor sources. Many VOC_{Toxic} and carbonyls
321 were significantly lower outdoors than those indoors, consistent with the findings in other
322 literatures (Bari et al., 2015). Chlorinated compounds such as hexachloro-1,3-butadiene,
323 1,2,4-trichlorobenzene, 1,1,2-trichloroethane and 1,2-dichlorobenzene had the highest I/O
324 ratios, which were 13-17, 12-17, 6-15 respectively. Bleach is reported to be a contributor
325 for indoor chloroform, and 1,2-dichlorobenzene is used to make mothballs and toilet
326 deodorizer blocks (Shepherd et al., 1996). The I/O ratios of other chlorinated compounds
327 were mostly greater than 3, implying that they were originated from indoor sources such as
328 liquid household products (Kwon et al., 2008). Huang et al. (2014) reported that
329 chlorinated compounds often use as industrial solvent such as pharmaceutical solvents,

330 dyes, pesticides, detergents, rubber, water disinfection, and chemical plants (Huang et al.,
331 2014). For carbonyls, formaldehyde and acetaldehyde had the I/O ratios of ~2. Most
332 aromatic compounds (e.g., BTEX) displayed moderate I/O ratios, revealing the
333 contributions from both indoor and outdoor sources (Edwards et al., 2001; Jia et al., 2008;
334 Wang et al., 2007). It should be noted that the ratios of carbon disulfide, n-hexane,
335 propylene and 1,3-butadiene were below unity, indicating that these VOCs are primarily
336 form outdoor sources. It was reported that propylene and 1,3-butadiene were the major
337 VOCs of vehicle emission (Li et al., 2017a; Xue et al., 2017), explained that the low I/O
338 values were found in our study.

339 **3.2.2. Source apportionment of Indoor VOC**

340 Source apportionment was conducted with U.S.EPA PMF receptor model. The
341 concentrations and uncertainties for the VOCs and carbonyls from those valid samples
342 collected in the eleven dwellings were used. Calibration was run for 3–7 factors and with
343 random seeds. We finally compared those examination results and considered that
344 five-factor PMF solution is the best fit for further analysis, when the relevant Q value
345 equals to 5144 in the robust mode. The residuals of the analytical results are mostly
346 between -3.0 and 3.0 (88%). With the adjustment of the number of factors, the calculated
347 results tend to be stable and the final determination of six factors. Some parameters of PMF
348 model when six major factors are selected were shown in Table S3. The r^2 of majority
349 species were greater than 0.6 and the model fitted well. The selected compounds' average
350 concentrations and mass contributions of each source factor are shown in Fig. 4. In Factor 1,
351 toluene, ethylbenzene, benzene, 1,4-dichlorobenzene and styrene had the highest

352 contribution. Lee et al. (Lee et al., 2002) reported that tobacco smoking could explain
353 indoor levels of benzene, toluene and m,p-xylene. BTEX and styrene are also found in
354 tobacco smoke (Wallace et al., 1987). Source apportionment shows that the dwellings with
355 two smokers (who consumed 6-9 cigarettes per day) had the contribution from this factor
356 being up to 90%. This factor was thus identified as smoking, accounting for 14.5% of the
357 total loading (Fig. 5).

358 Factor 2 should be associated with the off-gases from furniture, floor, building
359 materials and wall coverings. The key characteristic components in this factor were acetone,
360 formaldehyde, and acetaldehyde, also with high contribution of glyoxal, benzaldehyde and
361 methyl isobutyl ketone. Acetone is widely utilized in lacquers for either wooden- or
362 galvanized steel–furniture finishes (WHO, 1998). Hodgson et al. (2002) reported that
363 several cabinetry materials, passage doors, and the plywood subfloor were the predominant
364 sources of formaldehyde and other aldehydes. Wood-based materials used in construction
365 or in furniture production have long been the typical indoor source (Tunga, 2013). These
366 species can be emitted from indoor decorations such as furniture, floor and wall covering
367 materials including carpet, wallpaper, ceiling tiles, sheetrock, concrete and insulation foam
368 (Wallace et al., 1987; Wilke et al., 2010; Yu and Crump, 1998). This common indoor source
369 had an average loading of 44.5%, apportioned to be the most dominant indoor source.

370 Factor 3 was filled with methyl butyl ketone (MBK), acetone and 1,2-dichloroethane,
371 together with high contributions of methyl isobutyl ketone (MIBK) and benzyl chloride.
372 These species are often used as solvent in paints and adhesives (Chin et al., 2014; Yuan et
373 al., 2010). Therefore, this factor has been assigned to oil paints and adhesives. Its

374 contribution was 11.9% of total measured VOCs and carbonyls.

375 Factor 4 was characterized by methylene chloride, acetone, acetaldehyde, with high
376 contribution of chloroethane, 1,1-dichloroethane, cyclohexane. Small amounts of BTEX
377 also contributed to this factor. Previous studies have shown that toluene, xylenes,
378 methylene chloride, acetone, hexane, tetrachloroethylene, 1,1,1-trichloroethane, and
379 trichloroethylene typically exist at high abundances in household products (Sack and Steele,
380 1992). Formaldehyde, acetaldehyde, acetone can be released from cleaning reagents and
381 floor cleaners (Huang et al., 2011a). Wallace et al. (1987) identified 1,1-dichloroethane and
382 methylene dichloride in cleaning agents, pesticides, wallpaper and carpet glues (Wallace et
383 al., 1987). Kwon et al. (2007) also investigated the emission for household products in
384 Korea and found that acetone, m,p-xylenes, toluene, ethylbenzene, and hexane were
385 abundant. Acetone was determined in cleaning products, glues, nail color removers, paints,
386 and polishes. In another study, Kwon et al. (2008) also reported that many liquid household
387 products (e.g., deodorizers, cleaners, color removers, pesticides, and polishes) can release
388 several toxic aromatic and chlorinated organics (Kwon et al., 2008). Hence, factor 4 was
389 interpreted as household products and had a contribution of 17.3%.

390 The major loadings in factor 5 was formaldehyde, other species with high contribution
391 were o-xylene, styrene, benzene, 1,2,4-trimethylbenzene, benzyl chloride, trichloroethene,
392 tetrachloroethene. Formaldehyde had the highest concentration in Hong Kong restaurants
393 (Ho et al., 2006). The highest formaldehyde concentration in smoke from frying was also
394 detected (Xin et al., 2016). Formaldehyde is also produced by combustion processes and
395 heating of foods (Tunga, 2013). The concentration of aromatic hydrocarbons such as

396 benzene, toluene and chlorinated hydrocarbons also increased during the cooking periods
397 (Lin et al., 2014; Wang, 2011). Huang et al. (2011) also reported that the significant
398 increase of aromatic was related to evaporative loss of impurities in cooking fuels (Huang
399 et al., 2011b). As a result, this factor is marked as cooking, accounting for only 9.8% of the
400 total VOCs since the impact from household cooking was minimized.

401 Factor 6 was characterized by toluene, benzene, methylene chloride, propylene,
402 n-hexane, n-heptane, glyoxal, acrolein and freon-11. This series of compounds are highly
403 correlated with vehicle emission, biomass burning, industrial emission and solvent usage
404 (Li et al., 2017b; Xue et al., 2017; Zhang et al., 2012). The I/O value of these species
405 revealed they may also from outdoor. This factor was interpreted as outdoor, accounting for
406 2.1%.

407 **3.3. Health risk assessments**

408 Table 4 lists the 17 health-related chemicals catalogued at different groups by IARC
409 (International Agency for Research on Cancer) and with confirmed IUR or RfC inhalation
410 toxicity according to Integrated Risk Information from U.S. EPA's. The inhalation cancer
411 risk or non-cancer hazard risk were calculated based on these parameters.

412 **3.3.1. Cancer risk assessment**

413 The estimated inhalation cancer risks for nine VOCs are shown in the Fig. 6.
414 Formaldehyde had the highest cancer risk of 5.73×10^{-5} , followed by 1,3-butadiene (2.07
415 $\times 10^{-5}$) and 1,2-dichloroethane (1.44×10^{-5}). They are all higher than the acceptable risk level
416 of 1×10^{-6} but lower than the tolerable risk level of 1×10^{-4} . Formaldehyde and 1,3-butadiene
417 are all classified in group I as a human carcinogen by IARC groups. The major exposure

418 route of formaldehyde is inhalation from indoor air, impacting on nasal and upper airways.
419 Long-term exposure to formaldehyde increases the risk of developing multiple myeloma,
420 myelogenous leukemia and other special cancers. 1,3-butadiene is a characteristic of
421 vehicle exhaust, while 1,2-dichloroethane is often used as a solvent, such as resin, rubber,
422 dry cleaning agent and detergent. Therefore, the best health gains can be realized by
423 reducing both indoor and outdoor emissions of these VOCs.

424 Four chemicals of acetaldehyde (5.84×10^{-6}), chloroform (3.96×10^{-6}), carbon
425 tetrachloride (3.47×10^{-6}) and benzene (1.62×10^{-6}) presented median cancer risks but all
426 were also higher than the acceptable risk of 1×10^{-6} . Trichloroethylene (1.72×10^{-7}) and
427 tetrachloroethylene (2.26×10^{-8}) were well below the acceptable risk level.

428 **3.3.2. Non-cancer hazard risk assessment**

429 1,3-Butadiene presented the highest HQ value at 0.34, followed by acetaldehyde
430 (0.29), but they were below the threshold value (HQ=1). The other target VOCs with HQs
431 values were far less than the 1 (Fig. 7). Adverse health effects are not expected to result
432 from exposure to these VOCs according to the estimation.

433 **3.3.3 Improvement with air purifier**

434 The air purifier combines high efficiency particulate air filter (HEPA) with ambient
435 temperature catalysis technology. The air flow driven by the top fan passes through the
436 HEPA network, catalyst filling layer and inner filter layer and successively be purified
437 hierarchically. At room temperature (15-35 °C), formaldehyde and other VOCs react with
438 catalyst and rapidly decomposes into CO₂ and H₂O, which can effectively remove VOCs

439 (Li et al., 2018). The I/O values of VOCs and carbonyls before air purifier operated and
440 after the use of air purifier were shown in Fig. 3. It was obvious that most compounds' I/O
441 values before air purifier operated were higher than the after the use of air purifier
442 especially 1,1-dichloroethene, bromoform, 1,3-dichlorobenzene, chlorobenzene,
443 formaldehyde and hexanal. The air purifier effectively reduces the indoor concentration of
444 air pollutants. Sixteen selected VOCs and carbonyls at high indoor abundances were
445 selected to compare the impact of operation of air purifiers to purify the indoor air (Fig. 8).
446 Obvious declines in concentration were shown for both target compounds. The greatest
447 improvement was seen for 1,3-butadiene and formaldehyde, which were from of 7.54 ± 1.57
448 and $26.66 \pm 17.22 \mu\text{g}/\text{m}^3$ to 1.26 ± 0.38 and $16.29 \pm 13.41 \mu\text{g}/\text{m}^3$, respectively.
449 Correspondingly, their average estimated cancer risks have been also reduced from
450 4.65×10^{-5} and 7.12×10^{-5} to 7.8×10^{-6} and 4.35×10^{-5} . In addition, good purification
451 efficiencies were also seen for the removal of chloromethane, 1,2-dichloroethane,
452 bromoform, benzene, toluene and m,p-xylene. Even though the health risks for few of them
453 are still higher than the acceptable value, the substantial reduction could benefit the human
454 health.

455

456 **4. Conclusions**

457 Substantially high indoor VOCs and carbonyls concentrations were observed in
458 dwellings in Xi'an during wintertime. Most of the targeted species were more abundant in
459 China than other countries. The results from source apportionment conclude that both
460 smoking, decoration, furniture and household products are dominated sources at the

461 dwellings. The health risk of formaldehyde, 1,3-butadiene and 1,2-dichloroethane were
462 much higher than the acceptable risk level, even though the hazard quotient of few target
463 VOCs were far less than the threshold at non-cancer risk assessment. Preliminary data
464 shows that the use of air purifier can effectively reduce most of the indoor organic
465 pollutants, leading to decline in cancer risk to humans. The findings of this study provide
466 solid data to policy makers for understanding of characteristic pollution sources,
467 importance of IAQ management, and establishment of effective ambient pollution control
468 strategies.

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475

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642

643 **Table 1.** Statistical data of the general information and activities in the sampled dwellings obtained from the questionnaires.

Site#	Floor#	No of Rooms	No of Smokers ^a	Ventilation time per day	Cooking per week	Cleaning per week	Incense burning ^a	Insecticide ^a	Types of household chemicals consumed ^a	Fuel
1	26	4	1	<1 h	3-4	2-3	-	+	Laundry, dishwashing, and toilet detergent	LPG
2	6	4	- ^b	1-3 h	7	1-2	-	-	Laundry, dishwashing, and toilet detergent	natural gas
3	18	5	-	<1 h	3-4	7	-	-	Laundry, dishwashing, and toilet detergent	LPG
4	23	6	1	<1 h	7	7	-	-	Laundry, dishwashing, and toilet detergent	natural gas
5	4	5	-	<1 h	7	7	-	-	Laundry, dishwashing, and toilet detergent	electricity
6	23	4	-	<1 h	7	7	-	+	Dishwashing detergent	Electricity, LPG
7	5	4	-	9-12 h	-	7	-	-	Laundry and dishwashing detergent	-
8	2	2	-	1-3 h	7	4-5	-	-	Laundry and dishwashing detergent	Electricity, natural gas
9	8	3	1	1-3 h	1-2	2-3	-	-	Dishwashing and toilet detergent	LPG
10	2	4	-	3-6 h	7	2-3	-	-	Laundry, dishwashing, and toilet detergent, bleach	Electricity, LPG
11	5	4	2	1-3 h	7	7	+	-	Laundry and dishwashing detergent	Electricity, natural gas

644

645 ^a No record on the quantity consumed daily due to limitation

646 ^b No activity conducted in the dwelling

Table 2. Concentration of different categories of carbonyls and VOCs in indoor and outdoor ($\mu\text{g}/\text{m}^3$)

Compounds ($\mu\text{g}/\text{m}^3$)	Indoor (n=44)		Outdoor (n=37)		Compounds ($\mu\text{g}/\text{m}^3$)	Indoor (n=44)		Outdoor (n=37)	
	Mean	SD	Mean	SD		Mean	SD	Mean	SD
Carbon disulfide	0.63	0.69	1.24	2.41	<i>Halohydrocarbon</i>				
<i>Alkane</i>					Methylene Chloride	13.13	18.64	18.39	30.48
n-Hexane	1.76	1.65	1.83	2.12	Hexachloro-1,3-butadiene	9.41	15.78	0.62	1.74
n-Heptane	0.68	0.86	0.4	0.35	Bromoform	9.21	8.16	2.39	2.65
Cyclohexane	0.51	0.75	0.26	0.23	Carbon Tetrachloride	2.81	5.84	2.12	8.01
<i>Alkene</i>					1,2-Dichloroethane	2.69	4.12	1.78	3.08
1,3-Butadiene	3.36	1.16	3.89	0.64	Chloromethane	2.64	3.31	6.46	14.63
Propylene	2.91	3.38	4.54	6.28	1,2-Dichloropropane	1.93	2.98	1.86	3.35
<i>Carbonyls</i>					cis-1,2-Dichloroethene	1.32	0.8	0.53	0.32
Acetone	35.55	24.34	26.92	28.09	Chloroethane	1.08	1.07	0.89	0.9
Formaldehyde	21.45	13.72	8.53	7.94	1,4-Dichlorobenzene	0.99	1.15	0.16	0.21
Acetaldehyde	12.92	6.36	7.33	4.54	Benzyl Chloride	0.98	1.95	0.23	0.53
n-Nonanal	6.86	2.02	4.9	2.84	Chloroform	0.84	0.51	0.55	0.31
Methyl butyl ketone	5.45	8.98	10.8	36.77	1,1,2,2-Tetrachloroethane	0.74	1.31	0.31	0.66
n-Octanal	4.68	2.1	3.6	1.87	1,1,1-Trichloroethane	0.71	0.56	0.32	0.32
n-Decanal	4.05	1.9	2.77	1.44	1,3-Dichlorobenzene	0.68	0.87	0.14	0.16
Methyl Ethyl Ketone	3.56	1.81	3.01	1.53	1,2-Dichlorobenzene	0.56	1.04	0.06	0.1
Hexanal	3.05	1.66	1.23	1.09	1,1-Dichloroethane	0.47	0.44	0.34	0.32
n-Heptanal	2.49	0.88	1.96	0.72	1,2,4-Trichlorobenzene	0.47	0.38	0.03	0.04
iso-Pentanal	1.78	1.25	1	0.54	Dibromochloromethane	0.45	0.31	0.58	0.17
Methylglyoxal	1.72	0.92	2.16	1.57	Trichloroethene	0.42	0.37	0.48	0.51
Glyoxal	1.56	0.81	2.49	1.44	Tetrachloroethene	0.42	0.32	0.23	0.21
Propanal	1.54	0.66	1.35	0.7	Chlorobenzene	0.39	0.75	0.13	0.13
iso-n-Butanal	1.52	0.76	1.17	0.59	Bromodichloromethane	0.36	0.5	0.46	0.6
Acrolein	1.49	1.14	3.1	3.35	1,1,2-Trichloroethane	0.2	0.1	0.02	-
Benzaldehyde	1.24	0.68	0.9	0.38	1,2-Dichlorobenzene	0.18	0.16	0.26	0.28
n-Pentanal	0.92	0.5	0.53	0.23	1,1-Dichloroethene	0.16	0.11	0.06	0.02
2,5-Dimethylbenzaldehyde	0.9	0.42	0.87	0.46	trans-1,3-Dichloropropene	0.08	0.06	0.04	0.02
Methyl Isobutyl Ketone	0.83	0.76	1.51	1.43	Bromomethane	bd	-	bd	-
o-Tolualdehyde	0.49	0.21	0.43	0.21	trans-1,2-Dichloroethene	bd	-	bd	-
p-Tolualdehyde	0.43	0.19	0.43	0.19	1,4-Dioxane	bd	-	bd	-
m-Tolualdehyde	0.31	0.05	bd	-	cis-1,3-Dichloropropene	bd	-	0.41	0.07
<i>Others</i>					1,2-Dibromoethane	bd	-	bd	-
Ethyl Acetate	4.59	3.67	3.53	2.85	<i>Aromatic</i>				
Isopropyl Alcohol	1.17	1.05	1.71	1.71	Naphthalene	16.64	18.96	5.51	6.49
Vinyl Acetate	0.92	1.86	0.55	1.67	Toluene	7.23	7	4.22	4.08
Tetrahydrofuran	0.91	1.18	0.43	0.59	Benzene	3.58	3.48	3.01	2.66
Methyl-tert-butyl ether	0.61	0.56	0.55	0.52	Ethylbenzene	3.46	4.21	1.5	1.64
Methyl Methacrylate	0.15	0.12	0.22	0.42	o-Xylene	3.29	5.16	1.47	1.61
<i>Freon</i>					m,p-Xylene	2.72	4.41	1.22	1.61
Freon-11	1.67	2.15	0.97	1.8	Styrene	1.74	1.25	0.99	0.69
Freon-12	1.18	1.19	0.65	0.63	1,2,4-Trimethylbenzene	0.87	0.89	0.36	0.39
Freon-113	0.39	0.35	0.21	0.21	4-Ethyltoluene	0.37	0.32	0.15	0.15
Freon-114	0.01	0	0.07	0.03	1,3,5-Trimethylbenzene	0.37	0.35	0.15	0.13

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Table 3. Comparison of selected concentrations ($\mu\text{g}/\text{m}^3$) in dwelling with other relevant studies

Compounds ($\mu\text{g}/\text{m}^3$)	This study	Beijing, China (Duan et al., 2014)	Various cities, Japan (Azuma et al., 2016)	Shanghai, China (Dai et al., 2017)	Hong Kong, China (Lee et al., 2002)	Kocaeli, Turkey (Pekey and Arslanbaş, 2008)
Formaldehyde	21.45±13.72	40.2±26.2	13.00	-	-	-
Acetaldehyde	12.92±636	17.0±10.3	21.10	-	-	-
Acetone	35.55±24.34	23.6±10.7	27.10	-	-	-
Methylene Chloride	13.13±18.64	12.5±78.5	-	47.43±75.66	8.8±0.8	-
Chloroform	0.84±0.51	-	1.10	3.59±6.66	2.6±0.9	-
Benzene	3.58±3.48	7.35±11.6	2.40	2.32±1.19	4.7±0.5	13.06
Toluene	7.23±7.00	23.5±45.6	10.80	200.13±443.89	52.1±8.4	72.44
Ethylbenzene	3.46±4.21	3.68±2.49	5.60	26.33±27.73	0.6±0.8	-
m,p-Xylene	2.72±4.41	6.33±4.41	8.30	39.56±49.81	3.9±1.2	27.46
Styrene	1.74±1.25	1.85±2.13	-	32.59±42.77	-	11.65
o-Xylene	3.29±5.16	2.32±1.57	3.40	-	4.5±0.4	16.24
1,2,4-Trimethylbenzene	0.87±0.89	1.99±2.10	6.40	-	-	4.20

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652 **Table 4.** Health-related VOCs and related toxicity values.

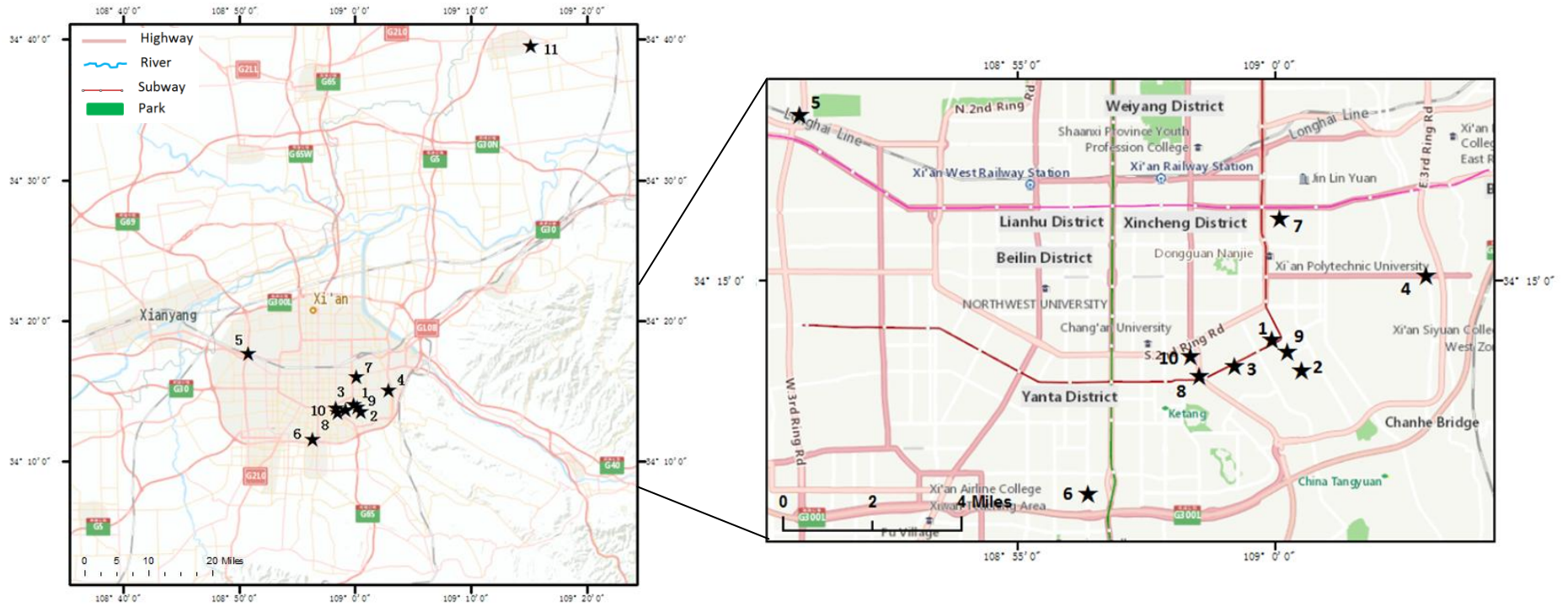
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Compounds	Cas no.	IARC	IUR ($\mu\text{g}/\text{m}^3$) ⁻¹	RfC (mg/m^3)
Formaldehyde	50-00-0	1	1.3×10^{-5}	-
Acetaldehyde	75-07-0	2B	2.2×10^{-6}	0.009
1,3-Butadiene	106-99-0	1	3×10^{-5}	0.002
Benzene	71-43-2	1	2.2×10^{-6}	0.03
Toluene	108-88-3	3	-	5
m/p-Xylene	106-42-3	3	-	0.1
o-Xylene	95-47-6		-	-
Ethylbenzene	100-41-4	2B	-	1
Styrene	100-42-5	2B	-	1
1,4-Dichlorobenzene	106-46-7	2B	-	0.8
Chloromethane	74-87-3	3	-	0.09
Methylene chloride	75-09-2	2A	1×10^{-8}	0.6
1,2-Dichloroethane	107-06-2	2B	2.6×10^{-5}	-
Chloroform	67-66-3	2B	2.3×10^{-5}	-
Carbon tetrachloride	56-23-5	2B	6×10^{-6}	0.1
Trichloroethylene	79-01-6	1	4.1×10^{-6}	0.002
Tetrachloroethylene	127-18-4	2A	2.6×10^{-7}	0.04

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Fig.1 Map shown dwellings in districts in Xi'an.

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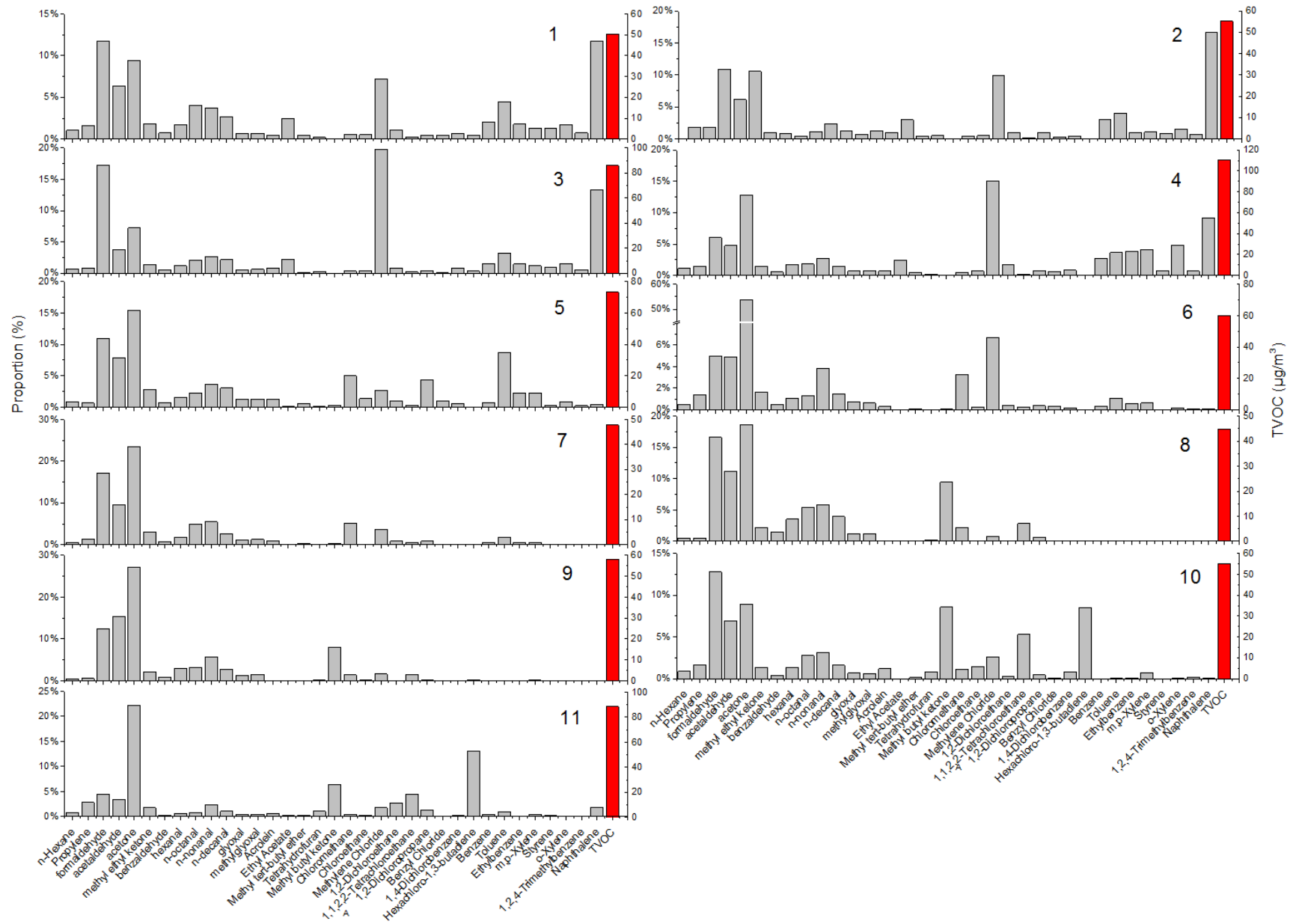


Fig. 2 Indoor mass proportions of typical compounds in each sampling site.

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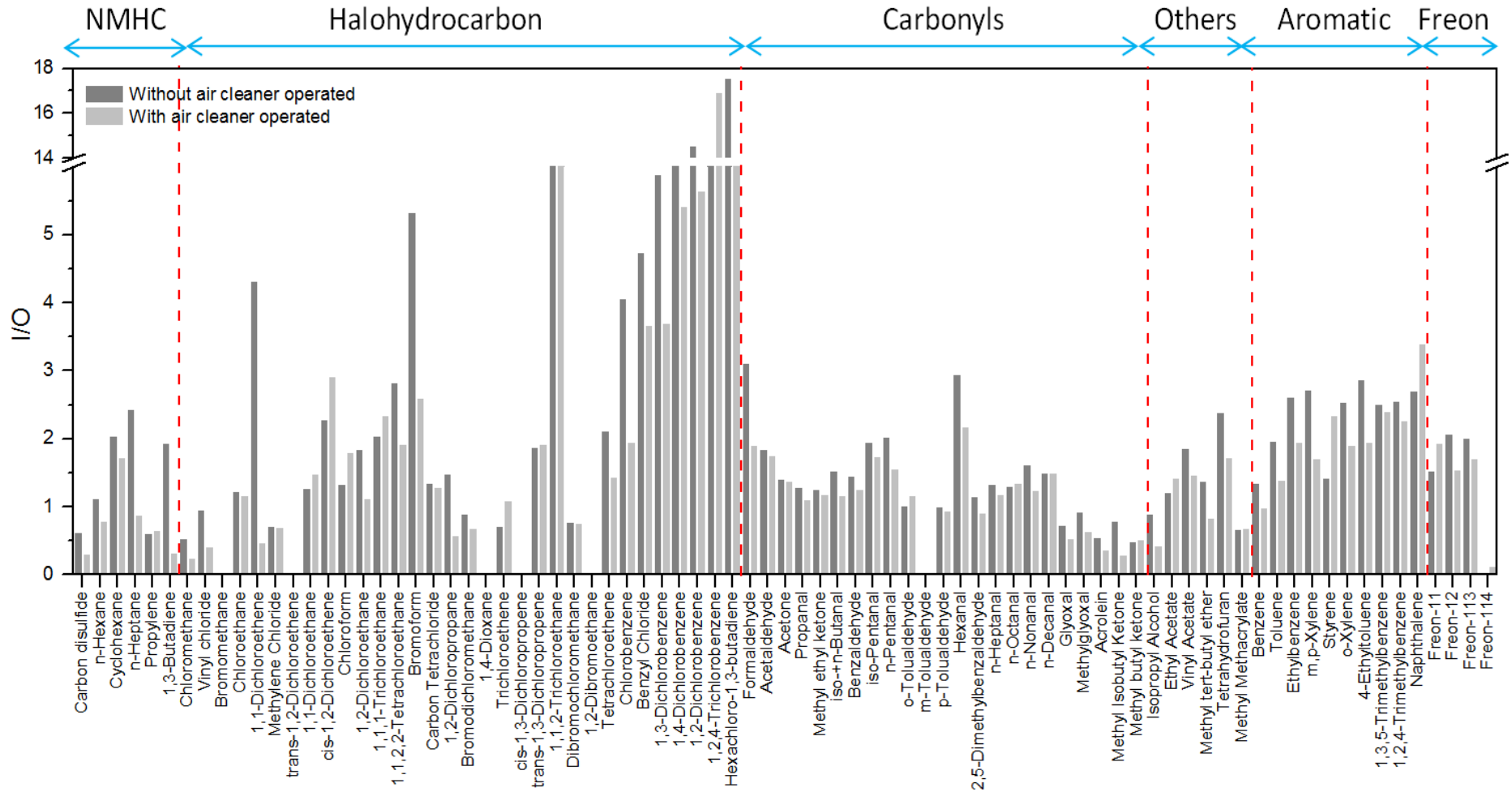
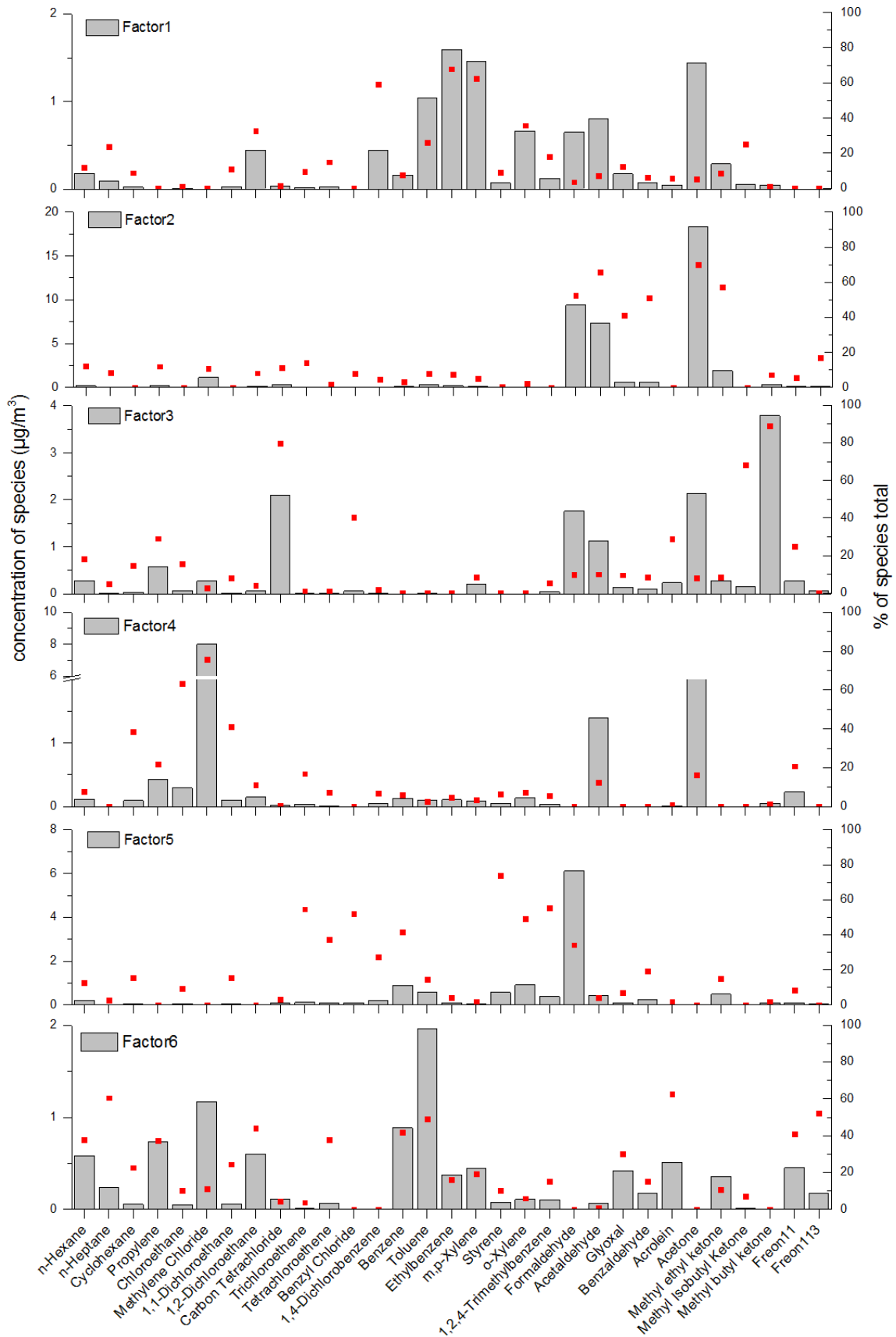


Fig.3 I/O ratios of VOCs and carbonyls

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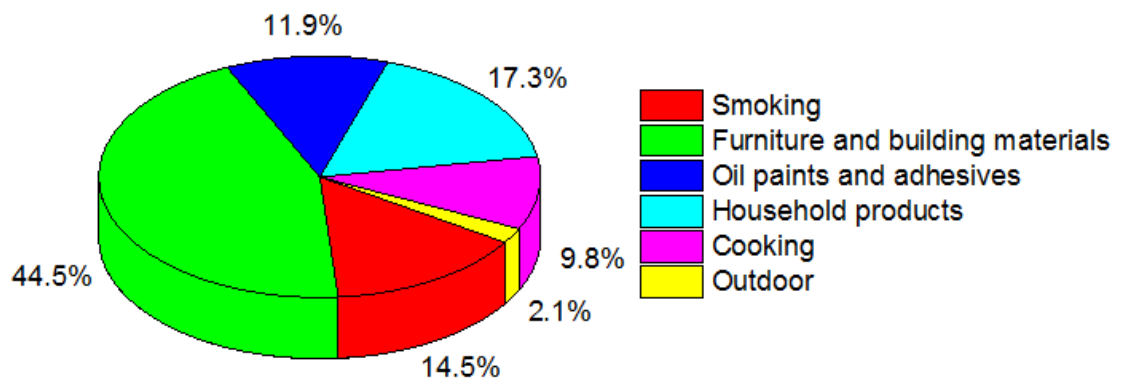


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Fig. 4 PMF-resolved indoor VOC source profiles (concentration of species and % of species apportioned to the factor from base run).

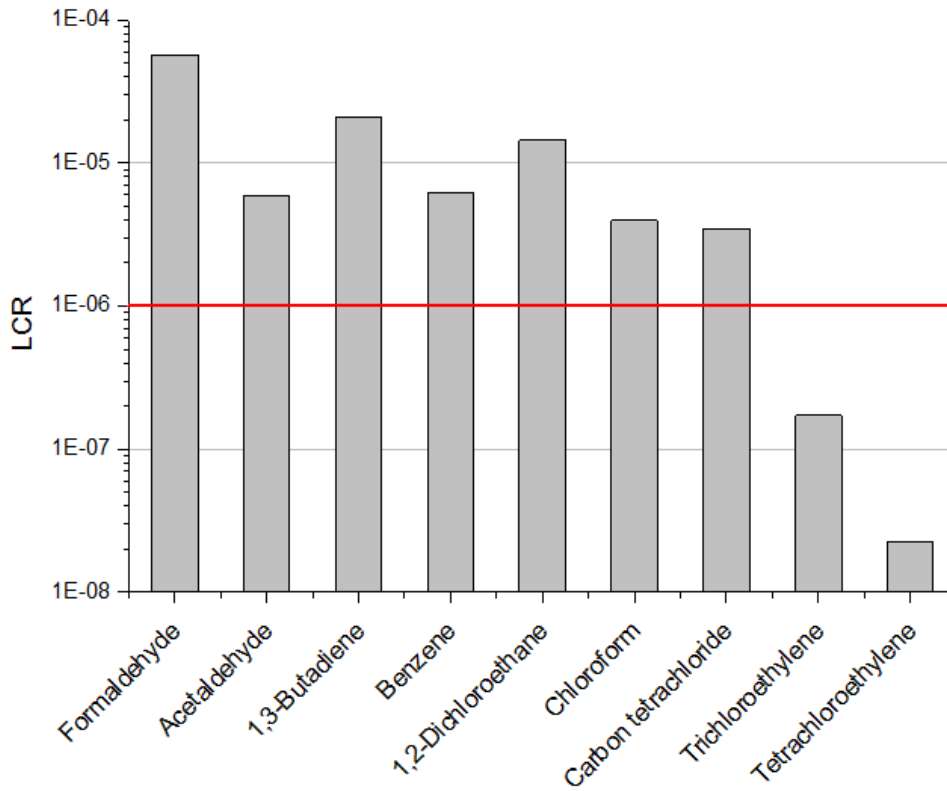


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Fig. 5 Source apportionment of indoor VOCs in Xi'an in winter.

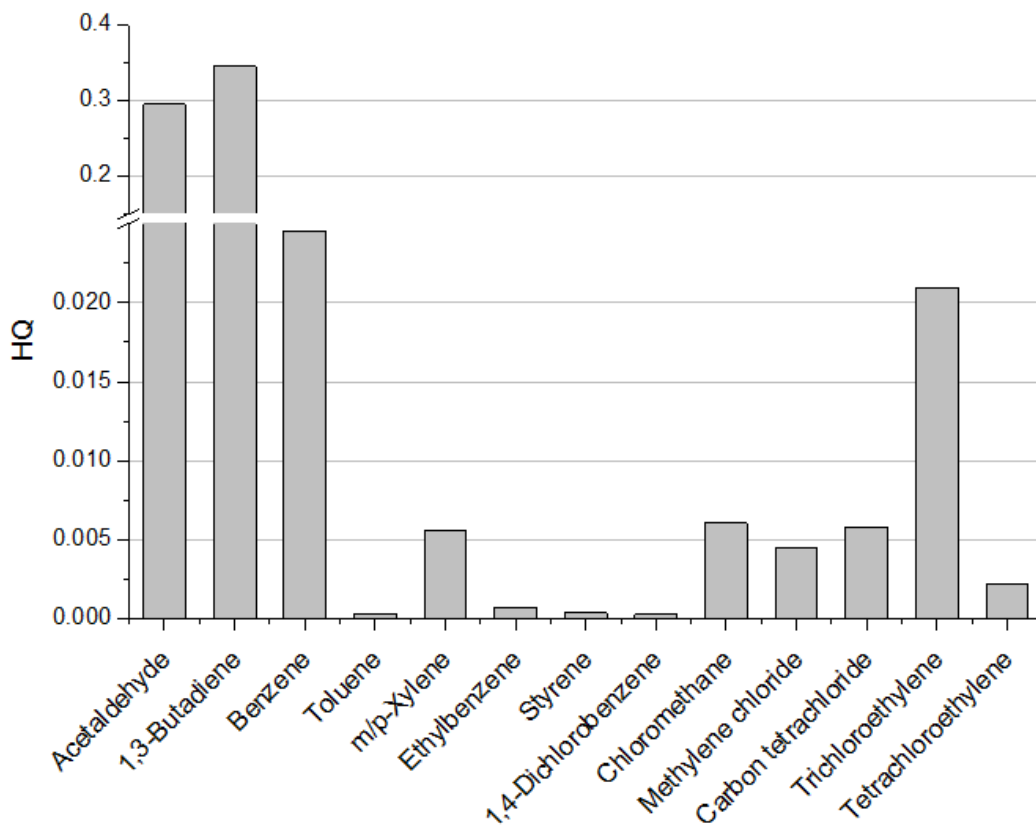
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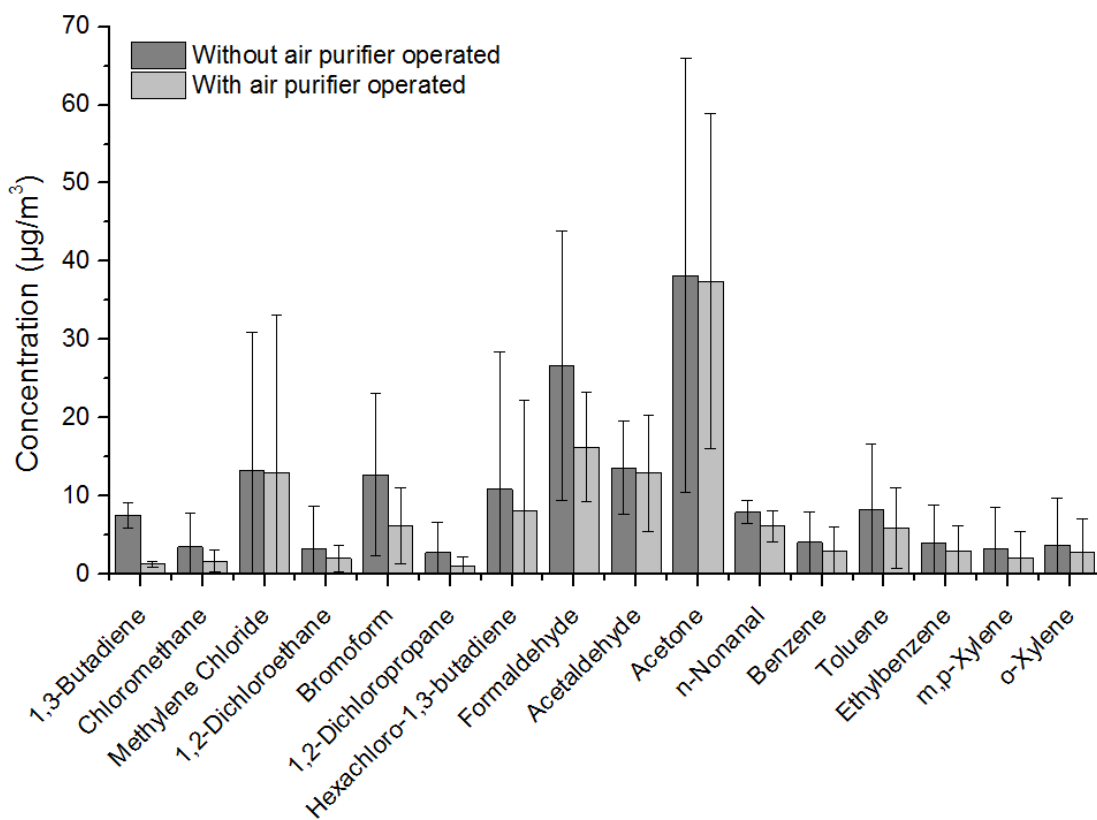
Fig. 6 Inhalation cancer risk evaluation for nine toxic compounds.

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Fig. 7 Non-carcinogenic estimated risk of VOCs using Inhalation Reference Concentration.



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Fig. 8 The comparison of selected VOC concentrations with operation of indoor air purifier.

680 **Table S1.** Indoor air quality questionnaires.

Name of interviewer: _____ Date of interview: _____ Home ID: _____	
I . Household information	
1. Name of respondent: _____	2. Gender: <input type="checkbox"/> Male <input type="checkbox"/> Female
3. Age: _____ 4. Phone: _____	5. Email: _____
6. What is your address? _____	7. Which kind of dwelling are you staying? <input type="checkbox"/> Simple house (Numbers of floor: _____) <input type="checkbox"/> Apartment/Dormitory (Floor: _____) <input type="checkbox"/> Others, please specify: _____
8. What is the number of members in your home/apartment? <input type="checkbox"/> 1 <input type="checkbox"/> 2 <input type="checkbox"/> 3 <input type="checkbox"/> 4 <input type="checkbox"/> 5 <input type="checkbox"/> > 5, please specify _____	9. How many hours do you spend at home/apartment? <input type="checkbox"/> <1-3 hours <input type="checkbox"/> 3-6 hours <input type="checkbox"/> 6-9 hours <input type="checkbox"/> 9-12 hours <input type="checkbox"/> >12 hours, please specify _____
10. How many rooms are there in your home/apartments? <input type="checkbox"/> Living room <input type="checkbox"/> Bedroom <input type="checkbox"/> Kitchen room <input type="checkbox"/> bathroom <input type="checkbox"/> Others, please specify _____	11. During the past 3 months, have any of the following changes been made in your home/apartment? <input type="checkbox"/> No <input type="checkbox"/> New flooring <input type="checkbox"/> New furnishing <input type="checkbox"/> New painted wall <input type="checkbox"/> New partition <input type="checkbox"/> New carpeting <input type="checkbox"/> Others, please spically _____
12. Does your home/apartment has outdoor spaces? <input type="checkbox"/> Garden <input type="checkbox"/> Balcony <input type="checkbox"/> Corridor <input type="checkbox"/> Others, please spically _____	13. Do you have any pets? <input type="checkbox"/> Yes <input type="checkbox"/> No If yes, please specify how many? <input type="checkbox"/> 1 <input type="checkbox"/> 2 <input type="checkbox"/> 3 <input type="checkbox"/> > 3, please specify _____
14. Which of the following activities are nearby your home/department? <input type="checkbox"/> Industrial activities <input type="checkbox"/> Mountain area <input type="checkbox"/> Residential home <input type="checkbox"/> Shopping mall <input type="checkbox"/> Roadside <input type="checkbox"/> Restaurants <input type="checkbox"/> Construction activities <input type="checkbox"/> Beach <input type="checkbox"/> Others, please spically _____	15. 1 How many smokers are there in your home/apartment? <input type="checkbox"/> 0 <input type="checkbox"/> 1 <input type="checkbox"/> 2 <input type="checkbox"/> 3 <input type="checkbox"/> > 3, please specify _____ 2 How many cigarettes do they smoke per day? <input type="checkbox"/> 1 to < 3 <input type="checkbox"/> 3 to < 6 <input type="checkbox"/> 6 to < 9 <input type="checkbox"/> > 9, please specify _____
II . Sources of indoor air pollution	
<input type="checkbox"/> Cooking activities <input type="checkbox"/> Printer/Photocopy <input type="checkbox"/> Cleaning activities <input type="checkbox"/> Carpet/Floor coating <input type="checkbox"/> Chemical usage <input type="checkbox"/> Painting activity <input type="checkbox"/> Incense burning <input type="checkbox"/> Outdoor sources, please spically _____	
III. Air ventilation	
1. What kind of air ventilation/circulation system do you use in your home/apartment? <input type="checkbox"/> Air conditioners <input type="checkbox"/> Fan <input type="checkbox"/> Natural ventilation(opening windows) <input type="checkbox"/> Others, please specify _____	2. How often do you use air ventilation in your home/apartment? <input type="checkbox"/> <1 hour <input type="checkbox"/> 1to < 3 hours <input type="checkbox"/> 3 to < 6 hours <input type="checkbox"/> 6 to < 9 hours <input type="checkbox"/> 9 to < 12 hours <input type="checkbox"/> >12hours, please specify _____
IV. Cooking activity	
1. Do you cook at home/apartment? <input type="checkbox"/> Yes <input type="checkbox"/> No If yes, please answer following question, if No please pass to part V .	2. How often do you cook at home/apartment? <input type="checkbox"/> 1-2 days/week <input type="checkbox"/> 3-4 days/week <input type="checkbox"/> Everyday
3. How many meals do you cook per day? <input type="checkbox"/> 1 <input type="checkbox"/> 2 <input type="checkbox"/> 3 <input type="checkbox"/> > 3	4. What kind of fuel do you use for cooking? <input type="checkbox"/> Electricity <input type="checkbox"/> LPG <input type="checkbox"/> Oil <input type="checkbox"/> Others, please spically _____

V. Cleaning activities	
1. How often do you clean the home/apartment? <input type="checkbox"/> Every day <input type="checkbox"/> 4-5 days per week <input type="checkbox"/> 2-3 days per week <input type="checkbox"/> Others, please specify _____	2. How can you clean the home/apartment? <input type="checkbox"/> Vacuum cleaner <input type="checkbox"/> Brush and broom <input type="checkbox"/> Cleaning towel <input type="checkbox"/> Others, please specify _____
3. What kinds of chemical do you use in the home/apartment? <input type="checkbox"/> laundry detergent <input type="checkbox"/> Insecticide <input type="checkbox"/> Bleach <input type="checkbox"/> Oven cleaners <input type="checkbox"/> Dishwashing detergent <input type="checkbox"/> Antibacterial cleaner <input type="checkbox"/> Toilet detergents <input type="checkbox"/> Others, please spically _____	4. How often do you use these Chemical? <input type="checkbox"/> Every day <input type="checkbox"/> Twice a week <input type="checkbox"/> Once a week <input type="checkbox"/> Once a month <input type="checkbox"/> Others, please spically _____
VI. IAQ-related information about your health	
1. During the past three months, have you had the following symptoms? (Please fill in the number 1: Always; 2: Usually; 3: Sometimes; 4: Occasionally; 5: Rarely; 6: Never)	
1.1 Eye symptoms <input type="checkbox"/> Dryness <input type="checkbox"/> Redness <input type="checkbox"/> Watering 1.2 Nasal symptoms <input type="checkbox"/> Dry nose <input type="checkbox"/> Running nose <input type="checkbox"/> Sneezing <input type="checkbox"/> Stuffy nose or congestion 1.3 Throat symptoms <input type="checkbox"/> Dry cough <input type="checkbox"/> Sore or dry throat	1.4 Skin problems(皮肤症状) <input type="checkbox"/> Dryness <input type="checkbox"/> Itching skin <input type="checkbox"/> Rash 1.5 Other symptoms <input type="checkbox"/> Difficulty in concentrating <input type="checkbox"/> Dizziness <input type="checkbox"/> Fever <input type="checkbox"/> Headache <input type="checkbox"/> Nausea <input type="checkbox"/> Shortness of breath <input type="checkbox"/> Unusual fatigue
VII. Other remarks _____	

Table S2. Concentration of VOC_{Toxic} and carbonyls in each sampling site.

Compounds \ Site#	1	2	3	4	5	6	7	8	9	10	11
Carbon disulfide	0.23	0.50	0.95	0.47	2.96	0.48	1.35	0.39	0.22	0.47	0.49
n-Hexane	1.52	2.91	2.11	3.75	1.66	0.75	0.56	0.59	0.52	1.52	2.18
Cyclohexane	0.27	0.46	0.41	2.46	0.05	0.03	0.03	0.02	0.02	0.21	0.60
n-Heptane	0.50	0.73	0.56	1.38	2.41	0.17	0.15	bd	bd	0.16	0.20
Propylene	2.37	2.92	2.59	4.66	1.32	2.03	1.51	0.59	0.78	2.77	8.84
1,3-Butadiene	bd	bd	bd	bd	bd	3.36	bd	bd	bd	bd	bd
Formaldehyde	17.01	17.63	53.63	20.35	22.67	7.52	18.97	18.38	17.95	21.82	14.01
Acetaldehyde	9.22	9.99	11.77	16.00	16.27	7.38	10.59	12.40	22.13	11.84	10.65
Acetone	13.70	17.01	22.29	43.29	32.09	80.80	25.94	20.64	39.23	15.28	68.93
Propanal	1.31	0.22	1.50	2.37	1.68	1.04	1.84	1.53	1.96	1.43	1.20
Methyl ethyl ketone	2.60	1.54	4.27	4.95	5.78	2.48	3.40	2.43	2.97	2.32	5.36
iso-+n-Butanal	1.23	0.44	2.14	2.03	2.52	0.77	1.30	1.45	1.58	1.38	1.03
Benzaldehyde	1.09	1.36	1.83	1.77	1.52	0.75	0.81	1.61	1.30	0.65	0.80
iso-Pentanal	1.80	0.44	2.33	1.93	3.18	1.07	1.32	0.94	3.65	0.77	1.36
n-Pentanal	0.86	0.18	1.12	1.36	1.52	0.51	0.57	1.05	1.07	0.75	0.60
o-Tolualdehyde	0.63	0.73	0.65	0.71	0.46	0.57	0.30	0.30	0.59	0.27	0.38
m-Tolualdehyde	bd	bd	bd	bd	bd	bd	bd	0.31	bd	bd	bd
p-Tolualdehyde	0.53	0.50	0.66	0.69	0.50	0.42	0.18	0.23	0.36	0.29	0.44
Hexanal	2.52	0.75	3.64	5.87	3.19	1.54	2.02	3.89	4.29	2.27	1.82
2,5-Dimethylbenzaldehyde	0.85	0.66	1.65	1.15	1.37	0.54	0.59	0.71	0.95	0.62	0.61
n-Heptanal	2.45	2.35	3.21	3.39	2.64	2.62	1.27	2.72	2.43	1.81	2.30
n-Octanal	5.89	1.74	6.51	6.09	4.66	1.91	5.53	6.00	4.57	4.70	2.43
n-Nonanal	5.37	3.84	8.21	9.15	7.67	5.72	6.02	6.51	8.13	5.33	7.42
n-Decanal	3.87	2.08	6.72	4.85	6.27	2.16	2.95	4.38	4.00	2.72	3.23
Glyoxal	0.94	1.19	1.60	2.35	2.73	1.07	1.26	1.32	1.84	1.09	1.25

Methylglyoxal	0.98	2.12	2.02	2.65	2.69	0.89	1.43	1.34	2.18	1.03	1.32
Acrolein	0.73	1.63	2.32	2.22	2.53	0.43	1.06	0.11	bd	2.03	1.96
Isopropyl Alcohol	1.20	2.00	1.42	1.05	2.56	0.11	0.68	0.12	bd	0.30	1.26
Ethyl Acetate	3.57	4.78	6.86	8.23	0.10	bd	bd	bd	bd	bd	0.74
Vinyl Acetate	0.16	0.36	0.14	0.41	0.11	0.01	0.30	0.26	0.11	2.16	4.45
Methyl-tert-butyl ether	0.73	0.69	0.50	1.64	1.23	0.14	0.29	0.14	0.03	0.19	0.63
Tetrahydrofuran	0.28	0.79	0.72	0.55	0.18	bd	0.03	0.21	0.19	1.33	3.38
Methyl Methacrylate	0.09	0.13	0.13	0.11	0.18	bd	bd	0.06	0.06	0.31	0.20
Methyl Isobutyl Ketone	0.12	bd	0.21	0.76	0.18	bd	bd	0.20	0.15	2.16	1.56
Methyl butyl ketone	0.04	0.06	0.10	0.19	0.65	0.05	0.26	10.49	11.42	14.60	19.73
Chloromethane	0.82	0.68	1.16	1.35	10.34	4.93	5.75	2.38	1.97	1.99	1.32
Vinyl chloride	0.04	0.12	0.90	0.20	0.11	bd	0.05	bd	bd	0.26	0.17
Bromomethane	bd	bd	bd	bd	bd	bd	bd	bd	bd	bd	bd
Chloroethane	0.76	0.89	1.27	2.30	2.76	0.33	0.09	bd	0.10	2.45	0.71
1,1-Dichloroethene	bd	0.38	bd	0.03	bd	bd	bd	bd	bd	0.02	0.01
Methylene Chloride	10.42	16.03	61.24	50.81	5.49	10.03	4.05	0.81	2.24	4.40	5.43
trans-1,2-Dichloroethene	bd	bd	bd	bd	bd	bd	bd	bd	bd	bd	bd
1,1-Dichloroethane	0.24	0.80	0.46	1.13	0.23	0.03	0.04	0.08	bd	0.65	0.36
cis-1,2-Dichloroethene	bd	bd	bd	1.42	1.46	0.11	0.14	bd	0.09	2.30	2.22
Chloroform	bd	2.75	0.16	0.67	0.79	bd	0.34	bd	bd	bd	bd
1,2-Dichloroethane	1.55	1.64	2.52	5.64	2.12	0.57	0.94	bd	bd	0.39	8.61
1,1,1-Trichloroethane	bd	bd	bd	bd	bd	bd	bd	0.17	0.14	0.73	1.99
1,1,2,2-Tetrachloroethane	0.10	0.06	0.19	0.11	0.06	bd	bd	0.06	0.06	1.70	3.88
Bromoform	bd	bd	bd	bd	bd	bd	bd	1.07	1.40	13.49	19.45
Carbon Tetrachloride	0.34	0.34	0.54	0.64	0.51	0.32	0.53	3.12	2.11	9.02	14.01
1,2-Dichloropropane	0.67	1.58	1.22	2.51	9.13	0.55	0.94	0.70	0.19	0.78	3.90
Bromodichloromethane	0.06	0.06	0.34	0.05	bd	bd	bd	0.09	0.05	0.63	1.32
1,4-Dioxane	bd	bd	bd	bd	bd	bd	bd	bd	bd	bd	bd

Trichloroethene	0.61	0.43	0.69	0.64	0.19	0.09	0.11	bd	bd	0.07	bd
cis-1,3-Dichloropropene	bd	bd	bd	bd	bd	bd	bd	bd	bd	bd	bd
trans-1,3-Dichloropropene	0.04	0.14	0.06	0.07	0.03	bd	bd	bd	bd	bd	bd
1,1,2-Trichloroethane	bd	0.11	0.06	0.05	0.44	bd	bd	bd	bd	bd	bd
Dibromochloromethane	bd	bd	0.07	bd	0.12	bd	bd	bd	bd	0.46	0.94
1,2-Dibromoethane	bd	bd	bd	bd	bd	bd	bd	bd	bd	bd	bd
Tetrachloroethene	0.33	0.46	0.37	0.61	0.21	bd	0.02	bd	bd	bd	1.34
Chlorobenzene	0.07	0.13	0.23	0.23	0.11	bd	bd	bd	bd	0.14	1.65
Benzyl Chloride	0.11	0.18	0.35	0.17	0.04	0.02	bd	0.03	0.05	2.76	4.47
1,3-Dichlorobenzene	0.73	0.49	0.15	1.97	1.97	0.42	0.10	bd	bd	0.08	0.26
1,4-Dichlorobenzene	1.00	0.63	2.62	2.72	1.23	0.27	0.06	0.01	0.01	1.38	0.66
1,2-Dichlorobenzene	0.10	0.10	0.36	0.05	0.01	bd	bd	0.01	0.01	1.55	2.22
1,2,4-Trichlorobenzene	0.89	bd	1.35	0.01	bd	bd	bd	bd	bd	0.21	0.20
Hexachloro-1,3-butadiene	0.71	bd	0.95	0.06	0.00	bd	bd	0.13	0.26	14.47	40.59
Benzene	3.01	4.88	4.82	9.18	1.54	0.45	0.60	bd	bd	bd	1.35
Toluene	6.54	6.51	9.71	12.26	17.97	1.62	1.95	bd	bd	0.06	2.94
Ethylbenzene	2.61	1.63	4.65	12.71	4.73	0.87	0.64	bd	bd	0.05	0.42
m,p-Xylene	1.89	1.76	3.61	13.75	4.54	0.89	0.45	0.10	0.10	1.15	1.27
Styrene	1.94	1.44	2.75	2.64	0.44	bd	0.08	bd	bd	bd	0.77
o-Xylene	2.52	2.40	4.62	16.14	1.69	0.26	0.08	bd	bd	0.03	0.31
4-Ethyltoluene	0.31	0.51	0.52	0.54	0.33	0.02	0.03	0.01	0.02	0.79	0.50
1,3,5-Trimethylbenzene	0.24	0.29	0.43	0.49	0.13	0.01	0.03	bd	0.01	0.95	0.68
1,2,4-Trimethylbenzene	1.04	1.17	1.62	2.18	0.44	0.06	0.07	bd	bd	0.33	0.36
Naphthalene	17.07	26.74	41.31	31.02	0.74	0.10	0.12	bd	bd	0.04	5.57
Freon-11	1.40	2.29	3.34	2.59	0.66	0.28	0.33	0.21	0.61	0.98	5.78
Freon-12	1.95	0.28	bd	2.21	1.29	0.45	0.44	0.23	0.20	1.55	2.69
Freon-113	0.41	0.62	0.51	0.48	0.43	0.21	0.30	0.17	0.05	0.37	0.50
Freon-114	bd	0.01	bd	bd	bd	bd	bd	bd	bd	bd	bd

683 **Table S3.** Parameters of PMF model.

Compounds	Intercept	Slope	r ²
n-Hexane	0.26	0.79	0.90
n-Heptane	0.20	0.36	0.49
Cyclohexane	0.11	0.32	0.65
Propylene	0.79	0.11	0.55
Chloroethane	0.13	0.18	0.60
Methylene Chloride	4.18	0.38	0.48
1,1-Dichloroethane	1.01	0.66	0.79
1,2-Dichloroethane	1.09	0.12	0.09
Carbon Tetrachloride	-0.08	0.99	0.98
Trichloroethene	0.09	0.41	0.45
Tetrachloroethene	0.06	0.49	0.54
Benzyl Chloride	0.09	0.07	0.70
1,4-Dichlorobenzene	0.00	0.86	0.75
Benzene	0.22	0.77	0.96
Toluene	1.40	0.51	0.52
Ethylbenzene	0.16	0.87	0.98
m,p-Xylene	0.38	0.73	0.97
Styrene	0.06	0.80	0.86
o-Xylene	0.67	0.45	0.79
1,2,4-Trimethylbenzene	0.02	0.94	0.97
Formaldehyde	8.16	0.45	0.62
Acetaldehyde	5.47	0.44	0.30
Glyoxal	0.55	0.55	0.33
Benzaldehyde	0.68	0.41	0.28
Acrolein	0.11	0.76	0.88
Acetone	20.76	0.16	0.09
Methyl ethyl ketone	1.64	0.48	0.34
Methyl Isobutyl Ketone	0.05	0.50	0.87
Methyl butyl ketone	-1.00	1.12	0.88
Freon-11	0.44	0.43	0.63
Freon-113	0.07	0.72	0.82

684 **Table S4.** P values of variance analysis at different day in the same room.

P (α :0.05)	Site										
	1	2	3	4	5	6	7	8	9	10	11
Without purifier operated	0.84	0.81	0.15	0.57	0.54	0.93	1.00	0.49	0.34	0.005	0.02
With purifier operated	0.66	0.06	0.66	0.77	0.97	0.95	0.37	0.34	0.29	0.10	0.16

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