



1 Molecular fingerprints and health risks of home-use incense burning

2 smoke

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28 Abstract: The burning of incense for home use is a widespread practice that has been shown to have 29 significant negative impacts on human health and air quality. However, there is a lack of 30 understanding regarding its emission profiles and associated health risks. To address this knowledge 31 gap, we utilized a state-of-the-art thermal desorption comprehensive two-dimensional gas 32 chromatography-mass spectrometer (TD-GC×GC-MS) to (semi-)quantify the emission factors (EFs) of 317 volatile compounds and thoroughly investigate the organic profiles of incense burning smoke 33 across a full-volatility range. Results showed that toluene $(70.8 \pm 35.7 \,\mu g \,g^{-1})$ is the most abundant 34 compound in incensing-burning smoke, followed by benzene, furfural, and phenol. Phenol, toluene, 35 36 furfural, 2-furanmethanol, benzene, and benzyl alcohol are the main contributors to ozone and 37 secondary organic aerosol (SOA) estimation. Intermediate volatility organic compounds (IVOCs) 38 accounted for 19.2% of the total EFs, but 40.0% of the estimated SOA. Additionally, a novel 39 pixel-based method, combined with aroma analysis, revealed that furfural can act as a key tracer of incense burning, and is responsible for the distinctive flavor of incense smoke. High bioaccumulation 40 41 potential (BAP) assessment using pixel-based partition coefficient estimation revealed that 42 acenaphthylene, dibenzofuran, and phthalate esters (PAEs) are chemicals of high-risk concern and 43 warrant further control. Our results highlight the critical importance of investigating home-use 44 incense burning and provide new insights into the health impacts of incense burning smoke by novel 45 approaches.

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1 Introduction

49 Incense burning is a prevalent custom in many cultures, especially in East and Southeast Asia 50 (Chen et al., 2021a). In modern times, incense burning for fragrance has become a frequent partice in 51 households (Manoukian et al., 2013), while functional incense burning, such as mosquito coils, is 52 used for specific purposes. Exposure to incense smoke is linked to adverse health effects like eye 53 irritation, carcinogenicity, genotoxicity, and respiratory system damage (Wong et al., 2020; Yang et 54 al., 2007; Chen et al., 2021b; Yang et al., 2017). Incense is composed of fragrant materials, aromatic 55 woods, herbs, and adhesive powders, usually available in the form of sticks and coils (Wong et al., 56 2020; Yadav et al., 2022a). Incense burning releases multiple pollutants into the air, including 57 particulate matter (PM), carbon monoxide (CO), volatile organic compounds (VOCs), and 58 intermediate volatility/semi-volatile organic compounds (I/SVOCs) (Wong et al., 2020; Yang et al., 59 2007; Jetter et al., 2002). 60 Current studies mainly focus on the hazardous VOC and SVOC homologs released from 61 incense-burning smoke. For instance, Lee et al. investigated 8 carbonyls and 11 VOCs emitted from 62 incense burning and found that the emission factors (EFs) of traditional incense burning were higher than aromatic incense (Lee and Wang, 2004). Lu et al. detected 230 kinds of VOCs from 63 64 mosquito-repellent incense burning, elucidating that alkanes, esters, aldehydes, ketones, and 65 aromatics are predominant (Lu et al., 2020). Staub et al. measured 6 methoxy phenolics, 10 monoterpenoids, and other 21 kinds of SVOCs in the burning smoke of incense sticks, and identified 66 67 cedrol as an important odor source (Staub et al., 2011). However, most of the studies have focused on 68 VOC compounds, with less attention given to gaseous organics in the full volatility range 69 (VOC-IVOC-SVOCs). A full-volatility organic characterization may better evaluate the ozone 70 formation potential (OFP) and SOA formation, as I/SVOCs are potentially important precursors of 71 ozone and secondary organic aerosol (SOA) formation (Lu et al., 2018a; Zhao et al., 2007; Tang et 72 al., 2021a). Meanwhile, mapping organics from incense smoke helps to evaluate the potential health 73 risks of toxic compounds. 74 Comprehensive two-dimensional gas chromatography (GC×GC) is a powerful technique dealing 75 with the coelution problem in conventional one-dimensional gas chromatography (1D GC).





76 Pollutants from gasoline exhaust, diesel exhaust, and cooking emissions are well separated and 77 identified (Drozd et al., 2019; Alam et al., 2018; Song et al., 2022a). As much as 50 ~ 98% of the 78 total response in GC×GC chromatograms could be explained (Huo et al., 2021; Song et al., 2022b). 79 Previous work identified 324 compounds from incense smoke by coupling solid-phase 80 microextraction (SPME) with GC×GC, yet chemicals are not quantified (Tran and Marriott, 2007). 81 Thus, a non-targeted and quantitative assessment of incense burning emissions is currently lacking. 82 In this work, two types of incense sticks and three kinds of incense coils were burned in a steel 83 chamber. Gaseous pollutants were trapped by Tenax TA desorption tubes and then analyzed by a thermal desorption comprehensive two-dimensional gas chromatography-mass spectrometer 84 85 (TD-GC×GC-MS). Pixel-based multiway principal component analysis (MPCA) was utilized to 86 identify markers of incense burning. Risk assessment of pollutants from incense burning emissions 87 was then evaluated by pixel-based approaches and high-risk compounds related to incense burning

2 Methodology

were assessed.

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2.1 Sampling and instrumentation

91 Incenses were purchased from the market, including 4 common incense sticks, 2 Thailand 92 incense sticks, 1 mosquito coil, and 2 incense coils (Figure S1). Incenses could also be classified by 93 their material, containing 2 aromatic coils, 4 aromatic sticks, 1 mosquito coil, 1 sandalwood stick, 94 and 1 smokeless sandalwood stick (Figure S1). Incenses were burned in a stainless combustion 95 chamber (1 m³). After ignition, the burning incense changed from flaming to smoldering. Each kind 96 of incense was burned at least twice. Incenses were weighed before and after combustion. 97 Preconditioned Tenax TA desorption tubes (Gerstel 6 mm 97 OD, 4.5 mm ID glass tube) were utilized to trap organics with a sampling flow of 0.2 L min⁻¹. 98 99 A comprehensive two-dimensional gas chromatography-quadrupole mass spectrometer 100 (GC×GC-qMS, GC-MS TQ8050, Shimadzu, Japan) coupled with a thermal desorption system (TDS 101 3 C506, Gerstel, Germany) was used for sample analysis. The desorption temperature was 280 °C.

The cooled injection system (CIS) with a Tenax TA liner was held at 20 °C and ramped to 320 °C

once injecting the gaseous sample into GC columns. The column combination was SH-Rxi-1ms (1st,





 $104~30~m\times0.25~mm\times0.25~\mu m)$ and BPX50 (2 $^{nd},\,2.5~m\times0.1~mm\times0.1~\mu m).$ The modulation period was

6s. See Table S1 and elsewhere (Song et al., 2022a) for more information.

2.2 Chemical identification, quantification, and 2D binning

A series of standard mixtures (2 µg mL⁻¹, 5 µg mL⁻¹, 10 µg mL⁻¹, 20 µg mL⁻¹, 40 µg mL⁻¹ in 107 CH₂Cl₂) was injected into Tenax TA tubes (2 µL). After purging the solvent with nitrogen gas, the 108 109 standards were thermally desorbed. The standard mixture contains 26 n-alkanes (C7 - C32, CNW 110 Technologies, ANPEL Laboratory Technologies (Shanghai) Inc., China), 16 PAHs, 11 phenolic 111 compounds, 9 alcohols, 4 aldehydes, 8 aromatics, 24 esters, 7 ketones, 5 siloxanes, and 39 other 112 compounds. Gaseous organics are quantified by external calibration curves with most of the R² 113 between 0.95 and 0.999 (Table S2). Chemicals with the same retention times and mass spectrums 114 were directly qualified and quantified. The unidentified chemicals were qualified by matching their 115 mass spectrum with library spectrums in the National Institute of Standard Technology library (NIST 17). Reverse factors of more than 700 were acceptable in this work. As homologs on the 116 117 two-dimensional chromatogram (contour plot) were eluted with near-equal one-dimensional intervals, 118 chemicals were then qualified by combing the location of the contour plot and the mass spectrums 119 (Song et al., 2023). Compounds without standards were semi-quantified by *n*-alkanes from the same 120 volatility bin (uncertainty 69%) and surrogates from the same chemical class (uncertainty 27%). 121 Instrument detection limits (IDLs) for organics semi-quantified were unknown, as a result, chemicals 122 with negative values calculated by calibration curves were quantified by the volume-to-mass (ng) 123 ratio of the lowest quantification point of standards (Table S2). A total of 317 chemicals were 124 (semi)-quantified, including 10 acids, 34 alcohols, 19 aldehydes, 25 aromatics, 38 esters, 49 ketones, 125 18 *n*-alkanes, 26 nitrogen-containing compounds, and 10 phenols (Table S3). The compounds identified were sliced into two-dimensional bins (2D bins) (Song et al., 2022a). 126 127 1st retention times are linked to the volatility of species (B8 to B31 with decreasing volatility) while 2nd retention times are associated with polarity (P1 to P8 with increasing polarity). Emission factors 128 129 of compounds in the same 2D bin were aggregated (Table S3).

2.3 Emission factor (EF), ozone formation potential (OFP), and secondary organic aerosol

131 **(SOA) estimation**





The emission factor (EF, $\mu g g^{-1}$) was calculated by the following equation:

$$EF = \frac{m V}{ftM} \tag{1}$$

- m is the absolute mass of pollutants (μ g) captured by Tenax TA tubes. V is the volume of the
- steel chamber (1 m³). The sampling flow and duration of the Tenax TA tube are $f(0.0002 \text{ m}^3 \text{ min}^{-1})$
- and t (min), respectively. M is the combustion mass (g) of the incense. The sampling volume of
- 137 Tenax TA tubes $(0.003 \sim 0.01 \text{ m}^3)$ was significantly smaller than the total volume of the steel
- chamber (1 m³) and the volume change of the chamber could be neglected.
- The ozone formation potential (OFP, $\mu g g^{-1}$) was calculated using equation (2). Where EF_i is the
- emission factor of precursor i (µg g⁻¹) with maximum incremental reactivity (MIR) of MIR_i . The
- 141 OFP was calculated inside the FOQAT packages developed by Tianshu Chen
- 142 (https://github.com/tianshu129/foqat). The MIR used in this work can be found in Table S3.

$$OFP = \sum [EF_i] \times MIR_i \tag{2}$$

Secondary organic aerosol (SOA) was estimated by equation (3).

$$SOA = \sum [EF_i] \times (1 - e^{-k_{OH,i} \times [OH] \times \Delta t}) \times Y_i$$
 (3)

- Where $k_{OH,i}$ and Y_i represent the OH reaction rate and SOA yield of precursor i,
- respectively(Table S3). The SOA yields of precursors were from literature (Loza et al., 2014; Harvey
- 148 and Petrucci, 2015; Tkacik et al., 2012; Shah et al., 2020; McDonald et al., 2018; Chan et al., 2010,
- 149 2009; Wu et al., 2017; Li et al., 2016; Matsunaga et al., 2009; Algrim and Ziemann, 2019, 2016; Liu
- et al., 2018; Charan et al., 2020) or surrogates from *n*-alkanes in the same volatility bins (Zhao et al.,
- 151 2014a, 2017). k_{OH} and Y could be found in Table S3. $[OH] \times \Delta t$ is the OH exposure and was set to
- be 13×10^{10} molecules cm⁻³s (24 hours in OH concentration of 1.5 $\times 10^6$ molecules cm⁻³).

2.4 Pixel-based risk assessments of incense-burning pollutants

- Octanol-air partition coefficient (K_{0-a}) , air-water partition coefficient (K_{a-w}) , and octanol-water
- partition coefficient (K_{o-w}) were estimated by a linear free-energy relationship (LFER) model (Nabi
- 156 et al., 2014; Zushi et al., 2019). Partition coefficients of chemicals are associated with their
- 157 two-dimensional retention times (Song et al., 2022b). Chemicals with high bioaccumulation potential
- 158 (BAP) are defined as contaminants with partition coefficients of $(2 < \log K_{o-w} < 11)$ and $(6 < \log K_{o-w} < 11)$
- 159 < 12). See Zushi et al. (Zushi et al., 2019) for more information. The R source code was obtained





160 from GitHub (https://github.com/Yasuyuki-Zushi).

3 Results and discussions

3.1 Emission profiles of different incense-burning organics

163 Figure S2 is a typical chromatogram of incense burning emissions, which is also set as the 164 reference chromatogram during the pixel-based analysis. As much as 90.2% of the total percent 165 response could be explained. The ratio is similar to a recent study resolving biomass burning 166 emissions (98%) (Huo et al., 2021). The emission factor (EF) of total organics is $791.8 \pm 300.6 \,\mu g \, g^{-1}$, consistent with previous work (100 ~ 19100 µg g⁻¹) (Lee and Wang, 2004), and comparable to rice 167 $(475.9 \pm 61.2 \ \mu g \ g^{-1})$, pine $(558.6 \pm 103.6 \ \mu g \ g^{-1})$ and poplar $(564.6 \pm 124.1 \ \mu g \ g^{-1})$ combustions 168 (Zhu et al., 2022), but much lower than coal combustion (6.3 mg g⁻¹) (Huo et al., 2021). The 169 170 contributions of different chemical categories are displayed in Figure S3. Oxygenated compounds 171 dominate the total EFs, accounting for 48.4%, followed by aromatics (29.8%), b-alkanes (5.3%), nitrogen-containing compounds (4.0%), alkenes (4.0%), and n-alkanes (2.3%). Unresolved complex 172 173 mixtures (UCMs) are further separated into aliphatic, cyclic, and oxygenated UCM due to retention 174 times and mass spectrums. The UCM ratio in this work (2.3% in EFs) is comparable to biomass 175 burning (Huo et al., 2021) and diesel exhaust (He et al., 2022) analyzed by GC×GC-MS, and is much 176 smaller than the UCM ratio (>50%) in biomass burning smoke analyzed by 1D GC-MS (Zhu et al., 177 2022). Ketones are the most abundant oxygenated compounds, accounting for 13.6% of the total EFs, 178 followed by aldehydes (9.7%), esters (8.1%), alcohols (6.9%), phenols (3.6%), and acids (3.1%). The 179 emission profiles are comparable to corncob and wood combustion, which are also dominated by 180 ketones and esters (Huo et al., 2021). However, the abundance of phenol is much lower than in 181 biomass-burning smoke (>15%) (Zhu et al., 2022; Huo et al., 2021), while comparable to coal 182 combustion (5.4%) (Huo et al., 2021). 183 EFs of selected compounds are listed in Table S4, which were comparable with other incense burning studies (Lee and Wang, 2004; Yang et al., 2007; Manoukian et al., 2016), while the EF of 184 benzene (59.6 $\pm 43.1 \,\mu g \,g^{-1}$) is slightly lower than other studies (188 $\sim 1826 \,\mu g \,g^{-1}$) (Lee and Wang, 185 186 2004; Yang et al., 2007; Manoukian et al., 2016). The Tenax TA liner in the CIS system does not 187 capture benzene at an initial temperature of 20 °C, while it is efficient for the trapping of most





189 As a result, the tailing of benzene on the second column (Figure S2) causes an underestimation of 190 blob integration and results in an underestimation of EF. 191 The top 10 compounds are all VOC compounds (Figure S4), accounting for 35.3% of the total EFs. Toluene $(70.8 \pm 35.7 \,\mu g \,g^{-1})$ is the most abundant compound in incensing-burning smoke, 192 193 followed by benzene, furfural, phenol, styrene, 2-oxo-propanoic acid methyl ester, 194 3-methyl-2-butanone, ethylbenzene, 1-hydroxy-2-propanone, and benzyl alcohol. Note that VOC 195 compounds discussed here are part of volatile organics captured by Tenax-TA, not the common 196 VOCs detected by SUMMA-GC-MS. The top 5 IVOCs are B17 b-alkanes, B16 b-alkanes, B18 197 b-alkanes, diethyl phthalate, and 1,6-dioxacyclododecane-7,12-dione. The naphthalene (a typical PAH, 2 rings) EF is $3.0 \pm 1.5 \,\mu g \, g^{-1}$, comparable to rice straw combustion (Zhu et al., 2022). SVOCs 198 199 are all *n*-alkane species and only account for less than 1% of the total EFs. 200 The average VBS distribution of incense burning is displayed in Figure 1, and the 201 volatility-polarity distribution is exhibited in Figure S5. In general, the EF decreases as the volatility 202 decreases, following the trend of VOC-EF (80.8%) > IVOC-EF(19.2%) >> SVOC-EF (<0.1%). The 203 chemical compositions in the VOC-IVOC range are shown in Figure S6. Oxygenated compounds 204 (53.5% of the total VOC EFs) and aromatics (37.6%) are largely detected in the VOC range, while 205 b-alkanes, n-alkanes, and oxygenated compounds are the main components of IVOC compounds. 206 The average VBS distribution is similar to cooking emissions (Song et al., 2022a) and wood 207 combustion (Stewart et al., 2021), but less volatile than gasoline exhausts (Lu et al., 2018b) and more 208 volatile than diesel emissions (Lu et al., 2018b). For example, the proportion of chemicals with saturated vapor concentration (C*) more than 10⁶ µg m⁻³ (Figure 1 a) is 80.8% (incense burning), 209 80.7% (cooking emissions) (Song et al., 2022a), 77.6% (wood combustion) (Stewart et al., 2021), 210 211 94.2% (gasoline exhaust) (Lu et al., 2018b), and 41.0% (diesel exhaust) (Lu et al., 2018b). The 212 polarity of incense burning is dominated by non-polar and intermediate-polarity organics (P1 ~ P5, 213 Figure S5). The volatility-polarity distribution of incense burning is quite similar to cooking 214 emissions (Song et al., 2022a), dominant by VOCs in the volatility range of before B13 and the 215 polarity range of P1 ~ P5.

I/SVOC compounds. Lower CIS temperature may trap benzene while causing water condensation.





216 A similar emission pattern but different EFs of different incense-burning emissions are observed. 217 Similarities among incense burning are more dominant than diversities. First, pixel-based partial 218 least squares-discriminant analysis (PLS-DA) elucidates that there is no systemic difference between 219 different chromatograms of incense burning emission, no matter different incense shapes (Figure S7) 220 or materials (Figure S8). Second, the compositions of different types of incense emissions are indeed 221 quite similar (Figure S9 and Figure S10). Third, the multiway principal component analysis (MPCA) 222 positive loadings are much larger than negative loadings, indicating that the similarities between 223 samples are much more important than the differences (Figure 2). 224 However, the absolute EFs significantly diverge according to different incense forms (p = 0.03, 225 Figure S11) and different materials (p < 0.01, Figure S12). Incense made in stick form (incense stick: $893.2 \pm 335.6 \,\mu g \,g^{-1}$, Thailand incense stick: $877.5 \pm 123.8 \,\mu g \,g^{-1}$) emits more organics than made in 226 227 coil form (incense coil: $835.5 \pm 306.0 \,\mu g \, g^{-1}$). The EF of mosquito coil is the smallest (382.5 ± 175.0 μg g⁻¹). A similar pattern was observed in previous work (Jetter et al., 2002). Concerning the incense 228 229 materials, we spot that the so-called smokeless sandalwood stick emits more abundant organics (1195.8 \pm 83.3 μ g g⁻¹) than common sandalwood sticks (633.7 \pm 6.6 μ g g⁻¹). The emission of 230 smokeless sandalwood sticks is even greater than aromatic sticks (893.2 \pm 335.6 μg g⁻¹) and coils 231 232 $(824.8 \pm 228.5 \,\mu g \,g^{-1})$. Our results demonstrate that although smokeless sandalwood stick is 233 preferred as fewer particulates are generated during the combustion process, the gaseous emissions 234 are enhanced compared to other incenses. 235 3.2 Contributions of home-use incense burning to ozone and secondary organic aerosols 236 (SOA) The total OFP is 1513.4 \pm 551.0 μg g⁻¹ which is 1.91 g O₃/g VOC-IVOCs. The OFP 237 238 enhancement ratio (OFP per mass of precursor) is much smaller than gasoline exhaust (3.53 g O₃/g 239 VOCs) (Wang et al., 2013) and evaporation (2.3 \sim 4.9 g O₃/g VOC) (Yue et al., 2017), showing that 240 incense burning is less efficient on ozone formation than gasoline-related sources. The lack of IVOC 241 measurements in previous work could also cause an overestimation of the OFP enhancement ratio as 242 IVOCs are less efficient in ozone formation. Toluene, furfural, p-xylene, benzyl alcohol, phenol, 243 2-furanmethanol, o-xylene, ethylbenzene, 1-hydroxy-2-propanone, and benzene are top 10 species





244 that contribute most to OFP (Figure S4). Oxygenated compounds take up 48.2% of the total OFP, 245 followed by aromatics (41.0%), and alkenes (6.7%) (Figure S3). VOCs dominate the total OFP, 246 accounting for 92.4% while IVOCs take up 7.6% (Figure 1). Aromandendrene, naphthalene, and 247 α-cedrene are the top 3 IVOC-OFP contributors. The volatility distribution of OFP contribution is 248 comparable to cooking emissions, as VOCs account for 88.8 ~ 99.9% of the total cooking OFP 249 estimation (Song et al., 2022a). Toluene contributes the most OFP in both cooking emissions and 250 incense burning. Short-chain linear aldehydes (pentanal, hexanal, nonanal) originating from the 251 degradation of oils play a more important role in OFP contribution in cooking emissions (Song et al., 252 2022a), while benzenes, furfural, alcohols, and phenols are non-negligible OFP contributors in 253 incense burning. 254 Figure 1 shows the volatility distribution of estimated SOA estimation, with the top 10 255 contributors displayed in Figure S4. IVOCs contribute 19.2% of the EFs while accounting for 40.0% of the total SOA estimation, highlighting the importance of IVOCs in SOA formation. The 256 257 contribution of IVOC species to SOA is higher than EFs due to the relatively higher yields and k_{OH} , 258 which has already been reported in cooking emissions (Song et al., 2022a; Yu et al., 2022), gasoline 259 exhaust (Zhao et al., 2014b; Tang et al., 2021b), diesel exhaust (Zhao et al., 2015), and biomass 260 burning (Stewart et al., 2021). Oxygenated compounds account for 32.9% of the SOA estimation, 261 followed by aromatics (23.7%), and b-alkanes (11.5%) (Figure S3). Phenol, benzyl alcohol, styrene, 262 toluene, B18 cyclic UCM, aromandendrene, 2-furanmethanol, B17 b-alkanes, benzene, and phenyethyne are the top 10 SOA contributors. The incense-burning SOA formation profiles are 263 264 distinct from cooking emissions (Song et al., 2022a) and biomass burning (Huo et al., 2021). Cooking SOA is largely derived from the oxidation of short-chain acids and aromatics (Song et al., 265 266 2022a), while phenols account for more than 65% of the SOA estimation from biomass burning (Huo 267 et al., 2021). Phenols only account for 11.0% of SOA estimation in this work. Alcohols (7.3%) and 268 furans (7.6%) are much more important SOA precursors in incense burning compared to biomass 269 burning and cooking emissions._Compared with other sources, we stress the importance of 270 incense-burning benzenes, furfural, alcohols, and phenols in OFP formation and alcohols and furans 271 in SOA formation. The secondary formation potential of mosquito coils is the lowest, while OFP and





- 272 SOA of burning smokeless sandalwood sticks are the highest. Compared to other incense, the higher
- 273 aromatic contents of smokeless sandalwood sticks burning fumes result in much more ozone and
- 274 SOA formation.

3.3 Identification of molecular markers from incense burning

276 Pixel-based MPCA is utilized to identify tracers of incense burning emissions. In brief, MPCA 277 decomposes a matrix X into a scoring matrix (S) and a loading matrix (L). Similarities and 278 differences in chromatograms are revealed by positive and negative loadings, respectively (Figure 2) 279 (Song et al., 2022b). The similarities of chromatograms could be explained by benzenes (toluene, 280 p-xylene, o-xylene, and ethylbenzene), ketones (3-methyl-2-cyclopenten-1-one, 281 2-hydroxy-2-cyclopenten-1-one, 3-ethyl-2-pentanone), aldehydes (furfural, succindialdehyde, 282 2-methyl-2-butenal), 2-methyl-propanoic acid, 1-methyl-1H-pyrazole, 2(5H)-furanone, and 283 2-furanmethanol. The differences between samples could be largely explained by 2-methyl-2-butenal, 284 2(5H)-furanone, 3,4-dimethylfuran, 2,3-dihydro-1H-inden-1-one, 2-methoxy-naphthalene, and 285 1,2-dihydro-2,2,4-trimethyl-quinoline. The negative loadings (0.006) are significantly smaller than 286 the positive loadings (0.07), confirming the dominance of similarities among chromatograms. The 287 relationship between the EFs of these compounds among different incense types is displayed in 288 Figure S13. Although the total EFs are significantly different (p = 0.03), the EFs of selected 289 compounds (2-hydroxy-2-cyclopenten-1-one, 2-furanmethanol, 3-ethyl-2-pentanone, and furfural are 290 significantly no different (p > 0.08). As a result, we recommend these compounds as incense-burning 291 tracers. It is reported that furfural is formed during the thermal degradation of hemicelluloses (Uhde 292 and Salthammer, 2007), while the oxidation of furfural under harsher conditions forms 293 2(5H)-furanone (Depoorter et al., 2021a). The formation mechanism of furfural from xylose and 294 D-xylopyranose is displayed in Figure S14 (Ahmad et al., 1995; Bonner and Roth1, 1959; Nimlos et 295 al., 2006). The initiation of the degradation of five-carbon sugars is from the acyclic form of pentoses 296 or directly via a 2,3- $(\alpha,\beta$ -)unsaturated aldehyde. The dehydrating of the intermediate compounds 297 finally forms furfural (Figure S14). The addressed tracers, furfural, 2-furanmethanol, and 298 2(5H)-furanone, have already been identified in incense burning smoke in previous work (Depoorter 299 et al., 2021a; Tran and Marriott, 2007).



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300 Furthermore, we compare the chemical profiles with an odor database (Aroma Office 2D, 301 Gerstel). Among the top 20 chemicals contributing to EFs, furfural (bread-like, alcoholic, 302 incense-like), phenol (mushroom, acid, burnt plastics), 1-hydroxy-2-propanone (buttery, caramellic, 303 fruity), benzyl alcohol (burning taste, flower, roasted), limonene (citrus-like, fruity, lemon-like), and 304 2-methyl-propanoic acid (apple-like, cheese-like, sweat) could be the aroma compounds. As for 305 tracers identified above, 2-furanmethanol (burnt sugar, honey, sweet) could also be another aroma 306 compound. Among them, furfural is widely and largely detected which could be the most important 307 molecular marker of incense burning (Silva et al., 2021; Depoorter et al., 2021b; Ho and Yu, 2002). 308 Note that aromandendrene, a cucumber-like, woody, and floral compound, is only detected in one 309 incense coil sample (incense coil 2, Figure S1). Aromandendrene is also detected in plants, such as in 310 dry flowers of Lonicera japonica (Shang et al., 2011). The emission factor of aromandendrene is 311 rather large (4.3 µg g⁻¹, 0.7% of the total EFs), and is a significant SOA precursor (2.3 µg g⁻¹, 3.9% of the total SOA estimation). The importance of aromandendrene on incense flavor and SOA 312 313 formation could not be neglected. Aromandendrene could also be responsible for the distinct flavor 314 of a certain incense coil. As above said, we recommend furfural be used as a molecular indicator of 315 incense burning regardless of the incense type or additives, especially responsible for the flavor of 316 incense burning.

3.4 Risk assessment of incense burning organics

The hazardous compounds from incense burning could cause adverse health effects on human health (Wong et al., 2020; Yang et al., 2007; Chen et al., 2021b; Yang et al., 2017). To evaluate the potential risks of these compounds, we conducted a pixel-based risk assessment (bioaccumulation potential, BAP) for partition coefficient estimation. Chemicals with high BAP concerns are listed in Figure 3. 2-methoxy-naphthalene, acenaphthylene, dibenzofuran, diethyl phthalate, dibutyl phthalate, benzoic acid 2-ethylhexyl ester, C15 – C19 n, and b-alkanes are regarded as high BAP concern (Figure 3). Among them, acenaphthylene is a toxic polycyclic aromatic hydrocarbon (PAH) that is widely detected in incense smoke (Yadav et al., 2022a). Dibenzofuran, an oxygenated compound with detrimental effects on human health (Suzuki et al., 2021), is also detected in the smoke of incense burning (Tran and Marriott, 2007). Diethyl phthalate and dibutyl phthalate are phthalate



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esters (PAEs) widely used as plasticizers which are endocrine disruptors (Wang and Qian, 2021). PAEs are abundant in incense smoke (Tran and Marriott, 2007). We propose that acenaphthylene, dibenzofuran, and PAEs could be chemicals of high-risk concern in incense smoke. We also assess the arctic contamination potential (ACP) as shown in Text S1. Further epidemiologic studies should be carried on to demonstrate the health effect of these hazardous compounds. 4 Implication The non-target approach of GC×GC-MS gives us a full glimpse of incense smoke, spotting a large pool of organics (317 compounds) covering the VOC-IVOC-SVOC range. We have provided a detailed description of both primary emission and secondary estimation of incense burning organics which is ready-to-use in SOA simulation models. IVOCs (130 compounds) are crucial organics accounting for 19.2% of the total EFs and 40.0% of the SOA estimation, highlighting the importance of incorporating IVOCs into SOA models. Further investigation should be carried on to elucidate emission characteristics of short-chain compounds that are lacking in our research, such as alkanes (<C7), alkenes (<C7), and aldehydes (<C5). By combining data obtained from gas-chromatography-flame ionization detector (GC-FID) and proton transfer mass spectrometer (PTR-MS), the emission pattern of incense burning could be well demonstrated. High-time resolution measurement should also be carried on to understand the time-resolved pattern of incense burning. We also suggest furfural as the molecular marker of incense burning as the EFs of furfural among samples are relatively stable. Pixel-based MPCA also indicates that furfural is responsible for the similarities between chromatograms. Furfural may be the key aroma compound of incense smoke. This key component identified in this work could be implemented in source apportionment. Furfural is also a key component contributing to OFP (rank 2). Phenol, toluene, 2-furanmethanol, benzene, and benzyl alcohol are the main contributors to both OFP and SOA. Surprisingly, we find that the EF of burning smokeless sandalwood sticks is the highest, with a remarkable contribution to OFP and SOA, due to the high aromatic contents. We recommend that both gaseous and particulate organics should also be taken into consideration when burning incense.

The single reduction of particles does not mean fewer emissions of gas-phase organics. A

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comprehensive assessment of incense-burning organics in both gas- and particle-phase should be implemented.

Combing pixel-based property estimation and blob identification, the risk assessment analysis of compounds could benefit analysts with less experience with GC×GC. The risk assessment in this work demonstrates that acenaphthylene, dibenzofuran, and PAEs are chemicals of high-risk concern and warrant further control. It was reported that more than half of Chinese residents burn incense every day at home for more than 20 years (Salvi and Apte, 2016). The toxic PAHs detected in indoor air could be 19 times higher than in outdoor air (Salvi and Apte, 2016; Yadav et al., 2022b). Exposure to these hazardous compounds could result in significant health threats. As a result, it is of vital importance to reveal and assess the epidemiological influences of incense burning in future work.

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602	Figures
603	Figure 1. Volatility distributions of EF, OFP, and SOA with chemical class in each volatility bin. The
604	x-axis is the unsaturated vapor concentration in logarithmic form (log C*, μ g m ⁻³). The y-axis is the
605	normalized mass emission factor (100%).
606	Figure 2. Positive (a) and negative loadings (b) of incense burning samples describing similarities
607	and differences between chromatograms. The color bar is the loading.
608	Figure 3. Chemicals with high bioaccumulation potential (BAP) assesses by pixel-based approaches.
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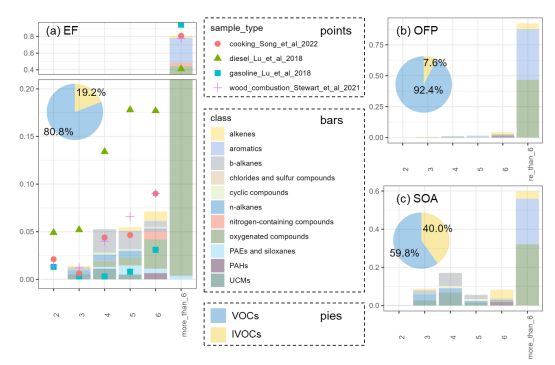


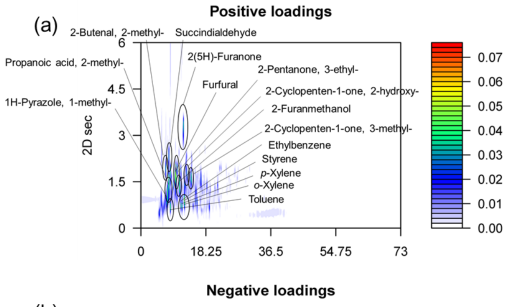
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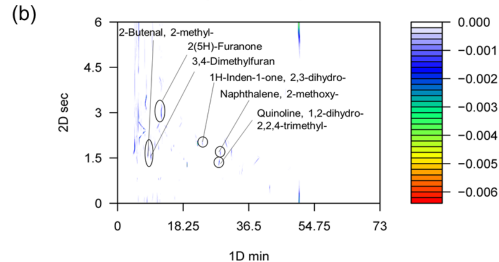


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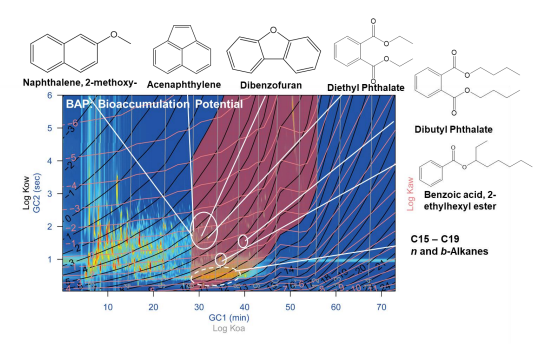


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