

Identification by volatolomics of hydrocarbon, oxygenated, sulfur and aromatic markers of livestock exposure to α -hexabromocyclododecane

jérémy ratel, Frédéric Mercier, Magaly Angénieux, Nathalie Kondjoyan, Saïd Abou El Karam, Patrick Blinet, Angélique Travel, Eric Royer, Elisabeth Baéza, Ronan Cariou, et al.

▶ To cite this version:

jérémy ratel, Frédéric Mercier, Magaly Angénieux, Nathalie Kondjoyan, Saïd Abou El Karam, et al.. Identification by volatolomics of hydrocarbon, oxygenated, sulfur and aromatic markers of livestock exposure to α -hexabromocyclododecane. Food Chemistry, 2022, pp.131504. 10.1016/j.foodchem.2021.131504. hal-03472011

HAL Id: hal-03472011 https://hal.inrae.fr/hal-03472011

Submitted on 8 Jan 2024

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.



Distributed under a Creative Commons Attribution - NonCommercial 4.0 International License

Version of Record: https://www.sciencedirect.com/science/article/pii/S0308814621025103 Manuscript_62c46974d41f0a0ce1c6dd2f16454e35

1	Identification by volatolomics of hydrocarbon, oxygenated, sulfur and
2	aromatic markers of livestock exposure to α -hexabromocyclododecane
3	
4	Jérémy Ratel ¹ *, Frédéric Mercier ¹ , Magaly Angénieux ¹ , Nathalie Kondjoyan ¹ , Saïd
5	Abouelkaram ¹ , Patrick Blinet ¹ , Angélique Travel ² , Eric Royer ^{3#} , Elisabeth Baéza-Campone ⁴ ,
6	Ronan Cariou ⁵ , Catherine Jondreville ^{4#} , Erwan Engel ¹
7	
8	¹ INRAE, UR QuaPA, F-63122 Saint-Genès-Champanelle, France
9	² INRAE, ITAVI, F-37380 Nouzilly, France
10	³ IFIP, F-31500 Toulouse; # IDELE, F-31321 Castanet-Tolosan, France
11	⁴ INRAE, UMR BOA, F-37380, Nouzilly, France; # INRAE UR ASTER, F-88500 Mirecourt,
12	France
13	⁵ INRAE, LABERCA, Oniris, F-44307 Nantes, France
14	* Corresponding author: jeremy.ratel@inrae.fr
15	
16	HIGHLIGHTS
17	• An experimental animal exposure to α -hexabromocyclododecane (HBCDD) was designed
18	• Hens, broilers and pigs were exposed to realistic α -HBCDD doses in their diet
19	• Liver volatolomics evidences exposure of the three farm animals to α -HBCDD
20	• Most candidate markers are hydrocarbon, oxygenated, sulfur and aromatic compounds
21	• Volatolomics, as an option for surveillance of chemical contamination in livestock
22	
23	ABSTRACT
24	Volatile organic Compounds (VOC)-based metabolomics, or volatolomics, was investigated
25	for revealing livestock exposure to chemical contamination. Three farm animals, namely

laying hens, broilers, and pigs, were experimentally exposed to 5 or 50 ng α -HBCDD g⁻¹ 26 feed. Liver and egg yolk for hens were analysed by headspace-SPME-GC-MS to reveal 27 28 candidate markers of the livestock exposure to α -HBCDD. For hens, 2-butanol was found as 29 marker in egg. In liver, twelve VOCs were highlighted as markers, with three aromatic VOCs 30 - styrene, o-xylene, α -methylstyrene - highlighted for the two α -HBCDD doses. For broilers, 31 six markers were revealed, with interestingly, styrene and phenol which were also found as 32 markers in hen liver. For pigs, ten markers were revealed and the seven tentatively identified 33 markers were oxygenated and sulfur VOCs. The candidate markers tentatively identified were 34 discussed in light of previous volatolomics data, in particular from a γ -HBCDD exposure of 35 laying hens.

36

37 KEYWORDS: Volatolomics, α-HBCDD, Headspace-SPME, GC-MS, Livestock, Food
38 safety.

- 39
- 40

41 **1. INTRODUCTION**

42 Hexabromocyclododecane (HBCDD) is a brominated flame retardant that has been a targeted 43 substance in the Stockholm Convention since May 2013. HBCDD is classified as a substance of very high concern (SVHC) on the REACH candidate list due to its persistent 44 45 bioaccumulative and toxic (PBT) properties. For non-occupationally exposed persons, dietary 46 exposure is a major route of total HBCDD intake (Covaci et al., 2006). Food contaminations 47 by HBCDD are especially problematic because of sometimes high levels, so that the ingestion 48 of a single contaminated animal-derived food can significantly increase consumer exposure to 49 it (Ratel et al., 2017). Based on direct HBCDD detection in food, the current analytical 50 methods for the surveillance of HBCDD contamination are efficient and sensitive but are

51 hindered by cumbersome implementation related in particular to the ubiquitous occurrence of 52 HBCDD. These routine monitoring techniques thus do not allow rapid cost-effective large-53 scale methods, which seem essential for strengthening food safety with respect to this 54 contaminant (Meurillon et al., 2018).

55 To overcome these limitations, alternative approaches based on omics have emerged to detect 56 contaminations. They are inspired by research showing the usefulness of the rapid cost-57 effective analysis of expired volatile organic compound (VOC) markers in clinical diagnosis 58 (Hakim et al., 2012). Berge et al. (2011) proved the concept that the signature of the liver 59 metabolome in small compounds, the volatolome, was modified for chickens in response to 60 exposure in their diet to different xenobiotics including brominated flame retardants such as 61 polybrominated diphenyl ethers. Liver is a key target for the determination of markers of 62 exposure to contaminants because of its major function in protecting the organism from 63 potentially toxic chemical insults. Bouhlel et al. (2018) took this concept and identified some 64 VOCs that were impacted in the liver volatolomes of chickens exposed to three types of 65 micropollutants - a pesticide, an environmental contaminant and a coccidiostatic agent. As 66 HBCDD impacts the metabolism of exposed organisms with clear indications of toxicological 67 effects in the liver (Cantón et al., 2008; Germer et al., 2006), liver volatolomics could be useful for identifying markers of HBCDD exposure and back-tracing HBCDD food 68 69 contamination.

Ratel et al. used volatolomics to reveal exposure of laying hens to γ -HBCDD (Ratel et al., 2017) and reported a list of VOCs in livers impacted by γ -HBCDD exposure. The study of Ratel et al. (2017) showed that at least four additional issues needed to be addressed: (i) this first study performed on HBCDD focused on the γ -isomer, whereas the α -HBCDD isomer is probably more relevant because it predominates in the environment, in animal tissues and in animal-derived food (Marvin et al., 2011; Rivière et al., 2014; Koch et al., 2015; DominguezRomero et al., 2016), (ii) this first list of VOC candidate markers needed filtering to screen for
the most robust one, (iii) the proof of concept had to be confirmed on more realistic HBCDD
levels of feed contamination, and (iv) the scope had to be extended to other farm animals.

79 Based on control/test experiments, the present study aimed first to confirm the utility 80 demonstrated by Ratel et al. (2017) of liver volatolomics in demonstrating the exposure of 81 laying hens to HBCDD, by considering the α -HBCDD isomer. In addition, realistic exposure 82 levels of laying hens were implemented and the potential of egg volatolomics was explored. 83 Egg is widely consumed and the literature has reported high HBCDD levels in chicken eggs, sometimes above the mg kg⁻¹ LW level (Hiebl and Vetter, 2017; Dominguez-Romero et al., 84 85 2016). According to its lipid rate, egg yolk may contain many volatile compounds and it is therefore very relevant for considering the implementation of non-invasive volatolomics 86 87 methods for food safety surveillance. Liver volatolomes were also investigated for two other 88 major monogastric farm animals, namely broilers and pigs. The potential markers discovered 89 are discussed especially in the light of the candidates found in the two main studies carried 90 out from liver volatolomics of livestock exposed or not to micropollutants: the study of Ratel 91 et al. (2017), based on the exposure of laying hens to γ -HBCDD, and that of Bouhlel et al. 92 (2018), based on the exposure of broilers to different micropollutants.

93

94 2. MATERIALS AND METHODS

95 **2.1. Animal feeding**

96 Compositions of experimental feeds for the three farm animals were detailed in Table S1. 97 Contaminated diets were obtained by replacing 5 g of clean soy oil in the control diet by 5 g 98 of soy oil spiked at appropriate levels with α -HBCDD. The synthesis of α -HBCDD has been 99 described by Dominguez-Romero et al. (2016). Briefly, technical HBCDD, containing 1, 5, 100 and 93% of α -, β -, and γ -HBCDD, respectively, was enriched in α -isomer by thermal

101 rearrangement (172 \pm 0.4 °C, 6 h), according to a method adapted from that described by 102 Szabo et al. (2011). Purification was performed on neutral silica gel and magnesium silicate 103 manually packed solid phase extraction (SPE) columns and also by preferential precipitation 104 at -20 °C, using dichloromethane and n-hexane. The purity of resulting crystals has been 105 estimated at 99.3% α-HBCDD, η-HBCDD being the only identified impurity. Crystals of α-106 HBCDD were dissolved in acetone used to spike soy oil at the targeted concentration. All 107 spiked feeds were prepared at the same time in the INRAE PEAT feed mixing unit (Nouzilly, France) by using dilutions of a single 50 μ g g⁻¹ contaminated oil prepared by LABERCA 108 109 (Nantes, France). However, for pigs, the volume of contaminated feeds needed for all the 110 experiment was too high to be produced in the INRA PEAT facilities. To overcome this issue, 111 the contaminated feeds for pigs have been concentrated 10 times more than the expected 112 level, then contaminated feed was brought each day of the experiment in the pig trough in a 113 1:10 ratio (w/w) mixed with uncontaminated feed.

114 For the three farm animals, a contaminated diet with the target concentration of 50 ng α -HBCDD g⁻¹ was prepared. This concentration was expected to enable the animal-derived 115 products to reach several hundreds of ng of HBCDD g⁻¹ LW, as previously reported in heavily 116 117 contaminated samples shown by French monitoring plans (Jondreville et al., 2017). For laying hens and pigs, a second lower dose of 5 ng g⁻¹ feed was tested to obtain animal-derived 118 119 products with more realistic α-HBCDD contamination levels. For broilers, the high 120 occupancy of the INRAE facilities did not allow testing two exposure doses with individual 121 animal cages.

122

123 **2.2. Animal testing**

Animal experiments have been designed to study the exposure to α-HBCDD in feed of laying
hens, slow-growing broiler chickens and growing pigs. These experiments were conducted in

compliance with Directive 2010/63/EU in France and approved by the relevant ethics 126 127 committee. Animal experiments were conducted in appropriate facilities with cages allowing 128 feed ingestion of individual animals to be monitored. Laying hens, broilers and pigs were fed 129 with non-contaminated feeds (control groups) or with α -HBCDD contaminated feeds 130 (exposed groups) for 18, 12 and 16 weeks, respectively. The main features of the three 131 experiments conducted in laying hens, broilers and pigs are summarized in Table 1, with the 132 detailed conditions of each animal experiment which are given in Table S2. Information about 133 body weight and feed ingested of animals during animal testing is given in Table S3.

134

135 **2.3 Slaughtering and sampling**

136 At the end of the exposure period, all animals were sacrificed and the weight of each warm 137 carcass was recorded at slaughter (Table S3). Poultry and pigs were killed after a 12-hour and 138 a 18-hour fast, respectively, by electronarcosis followed by exsanguination (electric stunner for poultry, Ducatillon, Cysoing, France; Morphee M4 electric stunner for pigs, Lelong & 139 140 Cie, Savigny, France). "Control samples" and "exposed samples" correspond to the samples 141 collected from control and exposed animal carcasses, respectively. For eggs, the eggs laid 142 before slaughter and on the day of slaughter were collected. The egg laid before slaughter was 143 dedicated to HBCDD analysis. It was weighed and removed from its shell before separating 144 and storing at -20°C the white and yolk. The egg laid on the day of slaughter was dedicated to 145 volatolomics analysis. It was collected, immersed in liquid nitrogen and stored at -80 °C. For 146 fat and muscle samples dedicated to HBCDD quantification, abdominal fat and muscles from 147 one thigh (without skin and bone) for poultry and samples of back fat and semi-membranous muscle for pigs were collected, weighed and stored at -20°C. For liver samples dedicated to 148 149 HBCDD quantification and volatolomics, the whole liver for poultry and a sample prepared 150 from the left lateral and right medial lobes for pigs were collected, weighed, immersed in liquid nitrogen, wrapped in aluminum foil, vacuum-packed, and stored at -80 °C. No visible
sign of pathology was detected during the autopsy carried out during the cutting of the postslaughter animals. Data about weight and lipid content of collected samples are given in Table
S3.

155

156 **2.4 HBCDD quantification**

157 HBCDD isomers (α , β and γ) were analysed in feed and animal samples according to a 158 method covered by the scope of the ISO/IEC 17025:2005 accreditation of the LABERCA 159 laboratory, and described by Dominguez-Romero et al. (2016). Briefly, all glassware and 160 Na2SO4 were baked prior to use at 400 °C for 4 h or at 650 °C for 6 h, respectively. Feed was 161 dried in an oven at 80 °C. Other matrices, animal tissues and egg yolk, were lyophilized. The 162 sample size ranges used for HBCDD quantification was given for each matrix in Table S4. Lipids were extracted by pressurized liquid extraction (PLE) over three successive static 163 164 cycles (100 bar, 120 °C) (SpeedExtractor, Büchi, Switzerland) with a toluene/acetone mixture 165 (70:30, v/v), evaporated to dryness, and weighed. Purification steps were conducted on a SPE 166 column manually packed with Na2SO4, neutral, and acidic (H2SO4) silica gel using hexane 167 and dichloromethane, followed by partitioning between n-hexane and 1 N NaOH. Final 168 extracts were reconstituted into a mixture of methanol/water 80:20 (v/v). HBCDD isomers 169 were analyzed by LC-ESI(-)-MS/MS (6410, Agilent Technologies). Separation was achieved 170 on a Hypersil Gold column (100 mm × 2.1 mm, 1.9 µm) (Thermo Scientific, San Jose, CA, 171 USA) fitted to a 1260 series HPLC pump. The mobile phase was constituted of 20 mM 172 ammonium acetate (A) and a mixture of acetonitrile/methanol 1:1 (v/v) (B) in isocratic 173 conditions (A/B 30:70, v/v). The transitions monitored through the triple-quadrupole mass 174 filter corresponded to $[M - H]^- \rightarrow [Br]^-$. Each analytical series comprised a procedural blank 175 and a quality control sample. Quantification was achieved according to the isotopic dilution principle (13C-labeled isomers as internal standards and 2H18-β-HCBDD as external
standard). Limits of quantification (LOQ) were determined individually for each sample and
isomer, based on a signal-to-noise ratio of 3. A limit of reporting (LoR) higher than each LOQ
was established by matrix, species and isomer (considering procedural contamination as well),
as presented in Table 2.

181

182 **2.5 Volatolome analysis**

183 All of each egg yolk and liver were ground in liquid nitrogen for 1 and 3 min, respectively, 184 into a fine homogeneous powder using a home-made stainless steel ball mill. A 1.2 g aliquot 185 of powder was placed in a 20mL-glass vial (Supelco, Sigma-Aldrich, St. Louis, MO), sealed 186 under a nitrogen flow with magnetic caps with PTFE/silicone septa (Supelco, Sigma-Aldrich), 187 and stored at -80 °C. The volatolome of powdered liver and egg yolk samples was analyzed 188 by headspace-solid-phase microextraction (HS-SPME) coupled to gas chromatography - mass 189 spectrometry (GC-MS) according to Ratel et al. (2017). Briefly, the following steps were 190 carried out with an automated sampler (AOC-5000 Shimadzu, Japan): (i) preheating of the 191 sample for 10 min in the agitator (500 rpm), (ii) SPME trapping (75 µm 192 carboxen/polydimethylsiloxane, 23-gauge needle, Supelco) of the volatile organic compounds 193 (VOCs) for 30 min. For the liver samples, the extraction temperature was set at 40 °C, as 194 recommended by Bouhlel et al. (2017) for better extraction of liver VOCs with a narrower 195 analytical variability and improved sensitivity. For egg yolk samples, this temperature had to 196 be increased to 60 °C to boost extraction rates. After extraction, thermal desorption was 197 performed at 250 °C for 2 min in the GC inlet. Further VOC analysis was performed by 198 GC/MS-full scan (GC2010; QP2010+, Shimadzu). VOCs were injected in splitless mode into 199 a DB-5MS capillary column (60 m \times 0.32 mm \times 1 μ m; Agilent J&W) with Helium as carrier 200 gas at a flow rate of 1 ml.min⁻¹. Oven temperature was held at 40 °C for 5 min, ramped to 201 230 °C at 3 °C.min⁻¹, and held at 230 °C for 10 min. The temperature of the transfer line 202 between GC and MS was set at 230 °C. Temperature was fixed at 180 °C in the MS source 203 and 150 °C in the MS quadrupole. Electron impact energy was set at 70 eV, and data was 204 collected in the range m/z 33–250 at 10 scans per second.

205

206 **2.6. Data treatment**

207 All calculations and statistical analyses applied to animal testing and HBCDD quantification 208 data have been previously detailed (Dominguez-Romero et al., 2016; Jondreville et al., 2017). 209 Peak areas of the VOCs were integrated from the SPME-GC-MS signals using a mass fragment selected for both being specific to the sought-after molecule and free of any co-210 211 elution with a home-made automatic algorithm developed in Boulhel et al. (2017) under 212 Matlab R2017 (The MathWorks, Natick, USA). VOCs were tentatively identified on the basis 213 of both mass spectra, by comparison against the NIST 17 mass spectral library (version 2.3, 214 build May 4 2017), and retention indices (RI), by comparison with published RI values and 215 those of our internal database. For calculation of the experimental RI, an alkane standard 216 solution (Supelco, Sigma-Aldrich) was analyzed by SPME-GC-MS at the end of the 217 analytical campaign. Data were processed using Statistica (version 12, StatSoft) and R 218 (version 2.1.4., http://www.R-project.org) software. Student's t-test and one-way ANOVA 219 were performed on the abundances of the VOCs monitored in volatolomes for the 220 comparisons of control vs. one or two groups of α -HBCDD contaminated animals.

221

222 **3. RESULTS**

223

3.1. HBCDD in feed

225 HBCDD concentrations determined in feed and animal tissues are summarized in Table 2. 226 Control feeds used for the three animals tested were HBCDD-free according to limits of reporting. For contaminated feed intended to contain 50 ng α -HBCDD g⁻¹ spiked feed, 227 compliant concentrations of 40, 38 and 31.8 ng α -HBCDD g⁻¹ feed were measured for laying 228 229 hens, broilers and pigs, respectively. The concentration of α -HBCDD in contaminated feeds 230 was 20–36% lower than expected. For the second lot of contaminated feed intended to contain 5 ng α -HBCDD g⁻¹ spiked feed for laying hens and pigs, concentrations of 3.62 ng and 3.42 231 ng α -HBCDD g⁻¹ feed were measured, respectively. The intended 1:10 ratios between the two 232 233 α -HBCDD levels (5 and 50 ng α -HBCDD g⁻¹) in spiked feed were moderately well kept following the preparation of contaminated feed, with a 1:9.0 ratio for laying hens and 1:9.3 234 235 for pigs. The difference observed in feed between the expected and measured concentrations 236 is probably related to the uncertainty in the weight of the produced crystals of α -HBCDD used 237 for spiking oils. But no degradation product of α -HBCDD has been identified in feed and 238 only α -HBCDD measured values were considered for further calculations. No β -HBCDD was detected in any spiked feed. Some γ -HBCDD was quantified at levels of 0.18 and 0.14 ng g⁻¹ 239 240 fw in the high- and low-level feeds, respectively, representing 0.45 and 3.9% of the total 241 HBCDD.

242

243 **3.2. HBCDD in animal samples**

The results of HBCDD quantification in animal samples obtained at the end of the exposure period are presented in Table 2. The results of tissue distribution and transfer to eggs of ingested α -HBCDD have been detailed in laying hens and in broilers by Dominguez-Romero et al. (2016) and Jondreville et al. (2017), respectively. The exposed animals, at any level, performed as well as the control animals in terms of body weight, growth rate, feed 249 efficiency, and laying rate, and the weight and lipid content of their tissues were not 250 significantly different (Dominguez-Romero et al., 2016, Jondreville et al., 2017). While the 251 HBCDD levels in ground liver samples from control laying hens should have been almost zero, traces of α - and γ -HBCDD isomers have been detected, with 2.2 ± 1.8 ng and 5.0 ± 3.3 252 ng g⁻¹ lw for α - and γ -isomer, respectively (Table 2). The γ -HBCDD content was very similar 253 254 to that from exposed hens (2.6 \pm 1.7 ng and 5.6 \pm 4.1 ng γ -HBCDD g⁻¹ lw for the exposure 255 "dose 5" and "dose 50", respectively; Table 2), which suggested a systematic contamination 256 in all ground liver samples. The profile in HBCDD isomers determined in ground control liver 257 samples was fairly constant. Dominated by the γ -isomer (29%, 5%, 66% of α -, β - and γ -258 HBCDD, respectively), it matched that of a technical mixture. HBCDD is known as a 259 ubiquitous environmental contaminant commonly found in industrial and domestic polymeric 260 materials. The analysis of the polystyrene box used during the liver grinding (Laberca 261 analysis ID 14.1842.3) has revealed a similar profile in HBCDD isomers (26%, 14%, 60% of 262 α -, β - and γ -HBCDD, respectively). HBCDD traces detected in ground control liver samples 263 could thus be ascribed to the polystyrene box. The analyses of HBCDD quantification pointed 264 out that the feeds given to control animals were $\alpha/\beta/\gamma$ -HBCDD-free and that the liver grinding 265 step was the source of the contamination revealed in control ground liver samples. 266 Accordingly, the volatolome of the liver from control hens can then be used to reveal α -267 HBCDD-exposure markers by comparison with the volatolome of liver from exposed animals 268 deliberately fed with α-HBCDD-contaminated feeds. In the experiments conducted on pigs 269 and laying hens, the α -HBCDD concentration measured in tissues was proportional to the 270 level of diet exposure of the pigs and laying hens.

271

272 **3.3.** Change in volatolomes in response to α-HBCDD exposure

To confirm the promising interest of liver volatolomics (Ratel et al., 2017) and to explore the potential of egg volatolomics for revealing livestock exposure to HBCDD, changes in volatolomes were investigated in SPME-GC-MS signals obtained from livers and eggs of control and α -HBCDD exposed animals. Analyses of liver volatolomes found 98, 105, and 134 VOCs for laying hens, broilers and pigs, respectively, and 51 VOCs were detected in egg yolk volatolomes (Table S5).

279 3.3.1. Laying hens

280 The results of liver volatolomics presented in Table 3 show that 8 VOCs were impacted in the 281 livers of laying hens for the two exposure levels. Figure S1 presents the first map of PCA 282 plotted on these candidate markers. The hepatic disturbance was thus visible in the 283 volatolome even for realistic low doses of α -HBCDD. Levels of HBCDD in feed in our study (5 and 50 ng α -HBCDD g⁻¹ feed) were reduced by a factor of 20 and 200 compared to the 284 levels of HBCDD used in Ratel et al. (2017) (0.1 and 10 μ g γ -HBCDD g⁻¹ feed). Table 3 285 286 includes mainly hydrocarbon, oxygenated (alcohols, ketones, acids), aromatic and sulfur 287 compounds. These results are consistent with the list of candidate markers published by Ratel 288 et al. (2017) in response to γ -HBCDD exposure, the authors highlighting in the livers of 289 laying hens mainly hydrocarbon compounds (alkanes and branched alkanes), oxygenated 290 compounds (alcohols, aldehydes, ketones) and aromatic compounds. Hydrocarbons. Heptane 291 (increased in exposed animals) and 2,2,4-trimethylpentane (decreased in exposed animals) 292 were found as significant markers of α -HBCDD exposure. Several hydrocarbons were 293 previously found as markers in the livers of laying hens after a γ -HBCDD exposure, with 294 levels in these compounds also increased or decreased in exposed animals according to the 295 compound considered (Ratel et al., 2017). The changes in the hydrocarbon content of livers in 296 response to α -HBCDD exposure may result from an imbalance between their production, 297 mainly due to unsaturated fatty acid peroxidation by reactive oxygen species (ROS), and their

298 hydroxylation, by detoxifying enzymes resulting in the production of alcohols (Hakim et al., 299 2012). Alcohols. Table 3 includes 2-butanol. In response to the exposure to toxic xenobiotics, 300 the level of alcohols in livers could result from an equilibrium between anabolism (e.g. 301 hydroxylation of hydrocarbons) and catabolism (e.g. activation of CYP-450) reactions. 302 Several alcohols were included in the list of Ratel et al. (2017), and primary and secondary 303 alcohols have been proposed as candidate markers for VOC-based clinical diagnoses (Hakim 304 et al., 2012). 2-Butanol can be thus considered as particularly useful for revealing exposure of 305 laying hens to α -HBCDD. Ketones. Like for alcohols, these compounds are at the crossroads 306 of cell metabolism elicited in response to exposure to toxic xenobiotics. The levels of 2,5-307 octanedione could be modified because of an impact of the toxic exposure on the lipid 308 metabolism with a high oxidation rate of fatty acids or on the protein metabolism with amino 309 acid metabolism-induced ketone formation (Hakim et al., 2012). Sulfur compounds. Two 310 compounds (sulfur dioxide, dimethylsulfone) were identified as markers at the highest 311 HBCDD dose. Among the candidate markers not classified according to their chemical 312 structure ("others" class) by Ratel et al. (2017), we can note two sulfur-containing compounds 313 (thiazole and thiadiazole). Bouhlel et al. (2018) also highlighted dimethylsulfone and carbon 314 disulfide among the most important liver VOC contributors to the separation of control 315 chickens and exposed chickens to micropollutants such as pesticides or polychlorobiphenyls. 316 In the review of Shubert et al. (2004) on the medical diagnostic potential of endogenous 317 VOCs, the authors report a possible relation between impairment of liver function, which 318 could be initiated by exposure to α -HBCCD given its toxicological effects reported in the 319 liver (Cantón et al., 2008; Germer et al., 2006), and level of sulfur-containing compounds. 320 The generation of these compounds may be connected to an incomplete metabolism of 321 methionine in the transamination pathway. Aromatic compounds. Table 3 shows 3 aromatic 322 compounds derived from alkylbenzenes among candidate markers: styrene, o-xylene and α - 323 methylstyrene. Alkylbenzenes were already highlighted by Ratel et al. (2017) when animals 324 were exposed to γ -HBCDD. In their study, the levels of all alkylbenzene candidate markers 325 were higher in exposed laying hens, like in our study. o-Xylene and α -methylstyrene were 326 also among the VOCs identified by Bouhlel et al. (2017) as the major contributors to the 327 separation of control and exposed chickens based on liver volatolomics. Although it is largely 328 agreed that alkylbenzenes have an exogenous origin (Hakim et al., 2012), their level in the 329 liver could be impacted by cellular and enzymatic defense mechanisms elicited to eliminate 330 hazardous compounds such as α -HBCDD. It is of note that we found styrene, o-xylene, α -331 methylstyrene and one unknown VOC (RI 1028) as candidate markers in the livers of laying 332 hens for both α -HBCDD doses. However, the abundance of these VOCs and the α -HBCDD 333 concentration in liver were not significantly correlated according to the R-Pearson test (r 334 Pearson indices between -0.13 and -0.46; p values between 0.26 and 0.75).

335 Regarding the egg volatolome, only 2-butanol was affected by the α -HBCDD exposure 336 (Table 3). This VOC was also identified as a marker in the livers of laying hens exposed to 337 the higher dose of α -HBCDD. 2-Butanol is a secondary alcohol in one of the VOC families 338 reported as likely to be impacted by a chemical risk exposure (Ratel et al., 2017). Given the 339 high utility of eggs in any non-invasive volatolomics-based control, it would be interesting to 340 deepen our knowledge of the potential impact of liver disorders on egg composition. 341 Saraswati et al. (2013) showed that liver functions modified yolk precursor synthesis and 342 depositions in the developing follicles. It would also be of interest to improve egg volatolome analysis by implementing recent advances in sample preparation and volatile fraction 343 344 collection (Majchrzak et al., 2018).

345 **3.3.2.** Broilers

346 The exposure to α -HBCDD generates a detectable metabolic disturbance in the liver 347 volatolome of broilers. The first map of PCA plotted on the candidate markers allows the 348 separation of case/control groups to be visualized (Figure S1). With levels higher in control 349 animals, the candidate markers listed in Table 4 are hydrocarbon (2-methylbutane) or 350 aromatic (phenol, 2-phenoxyethanol, methoxybenzene and styrene) compounds. These 351 chemical families were already highlighted in the liver volatolome of laying hens exposed to 352 α -HBCDD (Table 3) and to γ -HBCDD (Ratel et al., 2017). Concerning aromatic compounds, 353 styrene was already identified as affected in liver of laying hens for the two exposure doses of 354 α -HBCDD (Table 3). Styrene levels were detected significantly higher in control hens than in 355 exposed hens. These results make it promising candidate marker. Phenol was already 356 highlighted in the livers of laying hens as a candidate marker of γ -HBCDD exposure by Ratel 357 et al. (2017), with levels also higher in the livers of controls. To explain changes in liver 358 phenol content in response to y-HBCDD exposure, the authors hypothesize an imbalance 359 between production and degradation of phenol related to (i) the double involvement of CYP-360 450 enzymes in the detoxication process of toxic xenobiotics and in microsomal 361 hydroxylation of phenol in liver, and (ii) the hepatic degradation of tyrosine and tryptophan 362 amino acids responsible for phenol generation, which could be increased in the case of liver 363 function impairment. Our result thus confirms that phenol is a promising candidate marker to 364 reveal differences in liver metabolism after a diet exposure to HBCDD.

365 3.3.3. Pigs

Table 5 shows that 1 and 9 VOCs were significantly impacted when animals were exposed to 5 and 50 ng α -HBCDD g⁻¹ feed, respectively. This result suggests a dose effect and possibly a higher metabolic response threshold in the case of this animal. Figure S1 presents the first map of PCA plotted on the candidate markers revealed for the exposure at 50 ng α -HBCDD g⁻¹ feed. The candidate markers, levels of which were all higher in exposed animals, were 371 mainly oxygenated compounds (alcohols, ketones, lactones) and one sulfur-containing 372 compound (carbon disulfide). Among alcohols, we found the primary alcohol 3-methyl-1-373 butanol. This compound was reported as a hepatic candidate marker for exposure of laying 374 hens to γ -HBCDD (Ratel et al., 2017), with levels also higher in exposed animals. Carbon 375 disulfide was reported in the liver volatolome of chickens by Bouhlel et al. (2017) as a major 376 contributor to the discrimination between control chickens and chickens exposed to pesticide. 377 Further work is needed to identify the unknown VOC (RI 1132), which is a candidate marker 378 for pigs and broilers for the higher α -HBCDD level.

379

380 4. CONCLUSION

381 The present paper confirms that liver volatolome is relevant to highlight metabolic 382 disturbances induced by the exposure of animals to a chemical contamination with α -HBCDD. Based on realistic α -HBCDD exposures, our work supports the study of Ratel et al. 383 384 (2017) by detecting, by liver volatolomics, the α -HBCDD exposure of laying hens. It shows 385 the effectiveness of this approach in the case of two other farm animals, namely broilers and 386 pigs. But given the numbers of animals involved in this proof-of-concept study, further work 387 is needed to assess the robustness of the markers identified before considering to use them for 388 food safety surveillance. The chemical families of most candidate markers are consistent with 389 the two main cellular reactions put forward as affecting the anabolism and catabolism of 390 VOCs studied as marker of pathologies as cancers (Hakim et al., 2012). First, oxidative stress 391 could be induced, with synthesis of reactive oxygen species that leak from the mitochondria 392 or from peroxidated polyunsaturated fatty acids in the cell membranes. Second, detoxifying 393 enzymes like cytochrome P-450 enzymes could be induced with catalysis of the oxidation of 394 organic substances. Additional case/control experiments need to be investigated to go further 395 in assessing the robustness of the candidate markers, understanding of the cellular reaction 396 mechanisms which affect the VOC production and ascertaining the relationship between level 397 of risky exposure and volatolomic response. In this purpose, repeating *in vivo* experiments 398 with higher numbers of animals, especially for volatolomic studies on pig livers and poultry 399 eggs, or implementing *in vitro* experiments with hepatocyte culture cells are two very useful 400 opportunities. The recent advances in sensors for non-invasive and early detection of VOCs 401 (Li et al., 2020) should be followed since they may lead rapidly to the design of routine 402 sensors for easy, rapid detection of volatolomics markers useful for food chemical safety 403 surveillance.

404

405 ACKNOWLEDGMENTS

The authors thank the French Ministry for Food, Agriculture and Fisheries for financial support from the Compte d'Affectation Spéciale Développement Agricole et Rural (CASDAR project 1256). They are also grateful to the technical staff of PEAT and BOA (INRAE Nouzilly) and of GIE VGS (IFIP Villefranche-de-Rouergue) for diet preparation, animal rearing and sample collection.

411

412 **REFERENCES**

Aaslyng, M.D., Elmore, J.S., Mottram, D.S. (1998). Comparison of the aroma characteristics
of acid-hydrolyzed and enzyme-hydrolyzed vegetable proteins produced from soy,
Journal of Agricultural and Food Chemistry, https://doi.org/10.1021/jf9806816.

Amann, A., Mochalski, P., Ruzsanyi, V., Broza, Y. Y., & Haick, H. (2014). Assessment of
the exhalation kinetics of volatile cancer biomarkers based on their physicochemical
properties. Journal of Breath Research, https://doi.org/10.1088/1752-7155/8/1/016003

- Beaulieu, J.C., Grimm, C.C. (2001). Identification of volatile compounds in cantaloupe at
 various developmental stages using solid phase microextraction, Journal of
 Agricultural and Food Chemistry, https://doi.org/10.1021/jf0005768.
- Berge, P., Ratel, J., Fournier, A., Jondreville, C., Feidt, C., Roudaut, B., Le Bizec, B., &
 Engel, E. (2011). Use of Volatile Compound Metabolic Signatures in Poultry Liver to
 Back-Trace Dietary Exposure to Rapidly Metabolized Xenobiotics. *Environmental Science & Technology*, https://doi.org/10.1021/es200747h
- 426 Bouhlel, J., Ratel, J., Abouelkaram, S., Mercier, F., Travel, A., Baéza, E., Jondreville, C., 427 Dervilly-Pinel, G., Marchand, P., Le Bizec, B., Dubreil, E., Mompelat, S., Verdon, E., Inthavong, C., Guérin, T., Rutledge, D. N., & Engel, E. (2017). Solid-phase 428 429 microextraction set-up for the analysis of liver volatolome to detect livestock exposure 430 of micropollutants. Journal Chromatography to Α, 431 https://doi.org/10.1016/j.chroma.2017.03.008
- Cajka, T., Hajslova, J., Cochran, J., Holadova, K., Klimankova, E. (2007). Solid phase
 microextraction comprehensive two dimensional gas chromatography time-of-flight
 mass spectrometry for the analysis of honey volatiles, Journal of Separation Science,
 https://doi.org/10.1002/jssc.200600413.
- Cantón, R. F., Peijnenburg, A. A. C. M., Hoogenboom, R. L. A. P., Piersma, A. H., van der
 Ven, L. T. M., van den Berg, M., & Heneweer, M. (2008). Subacute effects of
 hexabromocyclododecane (HBCD) on hepatic gene expression profiles in rats.
 Toxicology and Applied Pharmacology, https://doi.org/10.1016/j.taap.2008.04.013
- Covaci, A., Gerecke, A. C., Law, R. J., Voorspoels, S., Kohler, M., Heeb, N. V., Leslie, H.,
 Allchin, C. R., & de Boer, J. (2006). Hexabromocyclododecanes (HBCDs) in the
 Environment and Humans: A Review. Environmental Science & Technology,
- 443 https://doi.org/10.1021/es0602492

- 444 de Montellano, P. R. O. (2015). Substrate oxidation by cytochrome P450 enzymes. In
 445 *Cytochrome P450* (pp. 111–176). Springer.
- 446 Dominguez-Romero, E., Cariou, R., Omer, E., Marchand, P., Dervilly-Pinel, G., Le Bizec, B.,

Travel, A., & Jondreville, C. (2016). Tissue Distribution and Transfer to Eggs of

447

- 448 Ingested α -Hexabromocyclododecane (α -HBCDD) in Laying Hens (*Gallus* 449 *domesticus*), https://doi.org/10.1021/acs.jafc.5b05574.
- Engel, E., Ratel, J. (2007). Correction of the data generated by mass spectrometry analyses of
 biological tissues: Application to food authentication. Journal of Chromatography A,
 https://doi.org/10.1016/j.chroma.2007.02.012.
- 453 Fournier, A., Feidt, C., Marchand, P., Vénisseau, A., Bizec, B. L., Sellier, N., Engel, E., Ratel,
- J., Travel, A., & Jondreville, C. (2012). Kinetic study of γ-hexabromocyclododecane
 orally given to laying hens (*Gallus domesticus*). Environmental Science and Pollution
 Research, http://doi.org/10.1007/s11356-011-0573-6
- Germer, S., Piersma, A. H., van der Ven, L., Kamyschnikow, A., Fery, Y., Schmitz, H.-J., &
 Schrenk, D. (2006). Subacute effects of the brominated flame retardants
 hexabromocyclododecane and tetrabromobisphenol A on hepatic cytochrome P450
 levels in rats. Toxicology, http://doi.org/10.1016/j.tox.2005.10.019
- Haick, H., Broza, Y. Y., Mochalski, P., Ruzsanyi, V., & Amann, A. (2014). Assessment,
 origin, and implementation of breath volatile cancer markers. Chemical Society
 Reviews, http://doi.org/10.1039/c3cs60329f
- 464 Hakim, M., Broza, Y. Y., Barash, O., Peled, N., Phillips, M., Amann, A., & Haick, H. (2012).
- 465 Volatile organic compounds of lung cancer and possible biochemical pathways.
 466 Chemical Reviews, https://doi.org/10.1021/cr300174a

- 467 Hiebl, J., Vetter, W. (2007). Detection of hexabromocyclododecane and its metabolite
 468 pentabromocyclododecene in chicken egg and fish from the official food control.
 469 Journal of Agricultural and Food Chemistry, https://doi.org/10.1021/jf063428b.
- 470 Jondreville, C., Cariou, R., Méda, B., Dominguez-Romero, E., Omer, E., Dervilly-Pinel, G.,
- 471 Le Bizec, B., Travel, A., & Baéza, E. (2017). Accumulation of α472 hexabromocyclododecane (α-HBCDD) in tissues of fast- and slow-growing broilers
 473 (*Gallus domesticus*). Chemosphere,
- 474 https://doi.org/10.1016/j.chemosphere.2017.03.064

485

- Koch, C., Schmidt-Kötters, T., Rupp, R., & Sures, B. (2015). Review of
 hexabromocyclododecane (HBCD) with a focus on legislation and recent publications
 concerning toxicokinetics and-dynamics. Environmental Pollution,
 https://doi.org/10.1016/j.envpol.2015.01.011
- Li, H. Y., Zhao, S. N., Zang, S. Q., & Li, J. (2020). Functional metal–organic frameworks as
 effective sensors of gases and volatile compounds. Chemical Society Reviews,
 https://doi.org/10.1039/c9cs00778d
- Majchrzak, T., Wojnowski, W., Piotrowicz, G., Gębicki, J., & Namieśnik, J. (2018). Sample
 preparation and recent trends in volatolomics for diagnosing gastrointestinal diseases.
- 484 TrAC Trends in Analytical Chemistry, https://doi.org/10.1016/j.trac.2018.08.020
- 486 V. (2011). Hexabromocyclododecane: Current Understanding of Chemistry,
 487 Environmental Fate and Toxicology and Implications for Global Management.
 488 Environmental Science & Technology, https://doi.org/10.1021/es201548c

Marvin, C. H., Tomy, G. T., Armitage, J. M., Arnot, J. A., McCarty, L., Covaci, A., & Palace,

489 Meurillon, M., Ratel, J., & Engel, E. (2018). How to secure the meat chain against toxicants?
490 Innovative Food Science & Emerging Technologies,
491 https://doi.org/10.1016/j.ifset.2017.10.004

- 492 Radulovic, N., Blagojevic, P., Palic, R. (2010). Comparative study of the leaf volatiles of
 493 Arctostaphylos uva-ursi (L.) Spreng. and Vaccinium vitis-idaea L.
 494 (Ericaceae), Molecules, https://doi.org/10.3390/molecules15096168.
- Ratel, J., Planche, C., Mercier, F., Blinet, P., Kondjoyan, N., Marchand, P., Fournier, A.,
 Travel, A., Jondreville, C., & Engel, E. (2017). Liver volatolomics to reveal poultry
 exposure to γ-hexabromocyclododecane (HBCD). Chemosphere,
 https://doi.org/10.1016/j.chemosphere.2017.09.074
- Rivière, G., Sirot, V., Tard, A., Jean, J., Marchand, P., Veyrand, B., Le Bizec, B., & Leblanc,
 J. (2014). Food risk assessment for perfluoroalkyl acids and brominated flame
 retardants in the French population: results from the second French total diet study.
 Science of the Total Environment, https://doi.org/10.1016/j.scitotenv.2014.01.104
- Saraswati, T. R., Manalu, W., Ekastuti, D. R., & Kusumorini, N. (2013). Increased egg
 production of Japanese quail (Cortunix japonica) by improving liver function through
 turmeric powder supplementation. International Journal of Poultry Science,
 https://doi.org/10.3923/ijps.2013.601.614
- Schubert, J. K., Miekisch, W., Geiger, K., & Nöldge–Schomburg, G. F. (2004). Breath
 analysis in critically ill patients: potential and limitations. Expert Review of Molecular
 Diagnostics, https://doi.org/10.1586/14737159.4.5.619
- de Simon, B.F., Estruelas, E., Munoz, A.M., Cadahia, E., Sanz, M. (2009). Volatile
 compounds in acacia, chestnut, cherry, ash, and oak woods, with a view to their use in
 cooperage, Journal of Agricultural and Food Chemistry, 2009, 57, 8, 3217-3227,
 https://doi.org/10.1021/jf803463h
- 514 Szabo, D. T., Diliberto, J. J., Hakk, H., Huwe, J. K., & Birnbaum, L. S. (2010).
 515 Toxicokinetics of the Flame Retardant Hexabromocyclododecane Gamma: Effect of

- 516 Dose, Timing, Route, Repeated Exposure, and Metabolism. Toxicological Sciences,
 517 https://doi.org/10.1093/toxsci/kfq183.
- Szabo, D. T., Diliberto, J. J., Hakk, H., Huwe, J. K., Birnbaum, L. S. (2011). Toxicokinetics
 of the flame retardant hexabromocyclododecane alpha: effect of dose, timing, route,
 repeated exposure, and metabolism. Toxicological Sciences,
 https://doi.org/10.1093/toxsci/kfr059.
- Vasta, V., Ratel, J., Engel, E. (2007). Mass spectrometry analysis of volatile compounds in
 raw meat for the authentication of the feeding background of farm animals. Journal of
 Agricultural and Food Chemistry, https://doi.org/10.1021/jf063432n.
- Xie, J., Sun, B., Zheng, F., Wang, S. (2008). Volatile flavor constituents in roasted pork of
 Mini-pig. Food Chemistry, https://doi.org/10.1016/j.foodchem.2007.12.074.

527

Livestock animals	Laying hens			Broi	lers	Pigs			
Strain Novo Brown		ו	JA6	57	(LW x Ld) x Piétrain				
Initial age (day)	210			1			70		
Duration of exposure (week)		18		12	2	16			
Target dose of α -HBCDD (ng g ⁻¹ feed)	50	5	0	50	0	50	5	0	
Number of animals	3 3 2			5	4	3	3	3	

 Table 1. Summary of the experiments conducted in laying hens, broilers and pigs

Samples	Laying hens			Bro	oilers	Pigs		
	dose 50 [°]	dose 5 ª	control	dose 50	control	dose 50	dose 5	control
Feed (ng g ⁻¹ fw)								
LoR ^b		0.04		0	.03		0.04	
α-HBCDD	40	3.62	< LoR	38.0	< LoR	31.8	3.42	< LoR
β-HBCDD	< LoR	< LoR	< LoR	< LoR	< LoR	< LoR	< LoR	< LoR
γ-HBCDD	0.18	0.14	< LoR	< LoR	< LoR	< LoR	< LoR	< LoR
Liver (ng g⁻¹ lw)								
LoR		0.4		(0.2		0.08	
number of samples	3	3	2	5	4	3	3	3
α-HBCDD	142±22*	11.0±2.2*	2.2 ± 1.8	100 ± 16	0.42 ± 0.29	15.6 ± 3.2	1.71 ± 0.17	< LoR
β-HBCDD	< LoR	< LoR	< LoR	< LoR	< LoR	< LoR	< LoR	< LoR
γ-HBCDD	5.6 ± 4.1	2.6 ± 1.7	5.0 ± 3.3	< LoR	< LoR	< LoR	< LoR	< LoR
Egg (ng g ⁻¹ lw)								
LoR		0.1						
number of samples	3	3	2	5	4	3	3	3
α-HBCDD	242±8.5	23.3±1.5	< LoR	N/A	N/A	N/A	N/A	N/A
β-HBCDD	< LoR	< LoR	< LoR	N/A	N/A	N/A	N/A	N/A
γ-HBCDD	< LoR	< LoR	< LoR	N/A	N/A	N/A	N/A	N/A
\mathbf{Fat}^{c} (ng g ⁻¹ lw)								
LoR		0.1		(0.2		0.1	
number of samples	3	3	2	5	4	3	3	3
α-HBCDD	302 ± 6.6	33.1 ± 6.3	0.26 ± 0.10	384 ± 82	0.46 ± 0.18	179 ± 21	14.1 ± 0.3	0.32 ± 0.19
β-HBCDD	< LoR	< LoR	< LoR	< LoR	< LoR	< LoR	< LoR	< LoR
γ-HBCDD	0.89 ± 0.42	0.46 ± 0.29	0.45 ± 0.29	< LoR	< LoR	< LoR	< LoR	< LoR
Muscle ^d (ng g ⁻¹ lw)								
LoR		0.2		(0.3		0.1	
number of samples	3	3	2	5	4	3	3	3
α-HBCDD	378± 41	N/A	N/A	260 ± 63	0.88 ± 0.64	142 ± 15	13.0 ± 2.8	< LoR
β-HBCDD	< LoR	N/A	N/A	< LoR	< LoR	0.7 ± 0.5	< LoR	< LoR
γ-HBCDD	< LoR	N/A	N/A	< LoR	< LoR	< LoR	< LoR	< LoR

Table 2. HBCDD concentrations determined in feeds and animal tissues. Values are means ± standard err

 a Diet contaminated with the target concentration of 5 or 50 ng $\alpha\text{-HBCDD g}^{\text{-1}}$

^b Limit of Reporting

^c Abdominal fat for hens and broilers, back fat for pigs

^d Thigh muscle for hens and broilers, semi-membranous muscle for pigs
 * Each α-HBCDD concentration in liver of exposed hens was corrected in removing the part of α-HBCDD brought by the polystyre

ne box used during liver grinding, according to Dominguez-Romero et al. (2016) and Jondreville et al. (2017).

·or. Levels of α - and γ -HBCDD measured in liver samples of control laying hens were not zero because of a

In accidental contamination during liver sample treatment (see paragraph 3.2. in Results section).

				_		Liver		Egg yolk		
Candidate markers	Tentatively identification ^a	m/z ^b	Exp. RI ^c	Ref. RI ^d	control	dose 50 ^e	dose 5 ^e	control	dose 50 ^e	dose 5 ^e
Hydrocarbons										
2,2,4-trimethylpentane	MS+RI	57	688	687 [1]	5.7 (9.1%)		1.4** (24.7%)			
heptane	MS+RI	100	700	700	13.7 (26.9%)		21.8* (5.3%)			
Alcohols										
2-butanol	MS+RI	59	602	603 [1]	13.4 (32.8%)	23.7* (3.9%)		42.5 (22.0%)		82.8* (1.1%)
Ketones	MS+RI									
2,5-octanedione	MS+RI	99	985	985 [2]	2.0 (47.1%)		6.6* (26.1%)			
Acids										
acetic acid	MS+RI	60	575	602 [1]	512.8 (29.7%)	775.5* (2.6%)				
Aromatic compounds										
styrene	MS+RI	104	898	897 [1]	4.0 (11.2%)	2.7* (15.7%)	2.6* (16.4%)			
o-xylene	MS+RI	91	898	898 [3]	5.8 (1.6%)	3.7* (15.9%)	3.5* (26.6%)			
α -methylstyrene	MS+RI	117	994	994 [4]	1.1 (0.3%)	2.1* (19.6%)	4.1** (13.7%)			
Sulfur compounds										
sulfur dioxide	MS	64	<500		32.7 (7.2%)	20.9* (18.6%)				
dimethylsulfone	MS+RI	79	906	915 [1]	9.7 (6.1%)	38.3* (23.3%)				
Unknown										
unknown		54	1028		0.4 (31.3%)	1.7** (11.7%)	3.5* (26.2%)			
unknown		100	1141		0.3 (39.4%)		1.5** (4.7%)			

Table 3. Laying hens - Volatile compounds in liver volatolome impacted by the exposure to α -HBCDD. Values are the mean of abundances (×10⁴) of each candidate mark

^a MS + RI, mass spectrum and RI agree with literature data; MS, mass spectrum agrees with literature spectrum

^b Mass fragment used for area determination

^{c,d} Retention indices on a DB5 capillary column from experimental run (c) or bibliographic data (d)
[1] Engel and Ratel, 2007; [2] Xie et al., 2008; [3] Vasta et al., 2007; [4] Cajka et al., 2007.

 e $\,$ Diet contaminated with the target concentration of 5 or 50 ng $\alpha\text{-HBCDD g}^{\text{-1}}$

* *p* <0,05

** p<0,01

Level found higher in liver from "control" animals compared to "exposed animals"

ker with its standard deviation (in bracket).

Table 4. Broilers - Volatile compounds in liver volatolome impacted by the exposure to α-HBCDD. Values are the mean of abundances (×10⁴) of each candidate m

Candidata markara	Tentatively	m/z ^b	Exp. RI ^c	Ref. RI ^d	control	dose 50 ^e
Candidate markers	identification ^a					
Hydrocarbons						
2-methylbutane	MS	72	508		63.9 (31.2%)	28.2* (45.1%)
Aromatic compounds						
styrene	MS+RI	104	898	897 [1]	13.7 (19.2%)	8.9* (32.2%)
methoxybenzene	MS+RI	108	922	918 [2]	9.3 (29.5%)	3.2** (21.0%)
phenol	MS+RI	94	976	983 [3]	92.4 (23.2%)	38.3** (54.3%)
2-phenoxyethanol	MS+RI	94	1207	1220 [4]	270.7 (26.2%)	38.9** (100.9%)
Unknown						
unknown		77	1132		10 (16.2%)	3.4** (35.2%)

^a MS + RI, mass spectrum and RI agree with literature data; MS, mass spectrum agrees with literature spectrum

^b Mass fragment used for area determination

^{c,d} Retention indices on a DB5 capillary column from experimental run (c) or bibliographic data (d)
[1] Engel and Ratel, 2007; [2] Leffingwell and Alford, 2011; [3] Vasta et al., 2007; [4] de Simon et al., 2009.

 e Diet contaminated with the target concentration of 50 ng $\alpha\text{-HBCDD}$ g $^{-1}$

* *p* <0,05

** p<0,01

Level found higher in liver from "control" animals compared to "exposed animals"

narker with its standard deviation (in bracket).

Table 5. Pigs - Volatile compounds in liver volatolome impacted by the exposure to α-HBCDD. Values are the mean of abundances (×10⁴) of each candidate mathematical determinants (×10⁴)

Candidate markers	Tentatively identification ^a	m/z ^b	Exp. RI ^c	Ref. Ri ^d	control	dose 50 ^e	dose 5 ^e
Alcohols							
3-methyl-1-butanol	MS+RI	55	733	734 [1]	5.0 (28.5%)	8.7* (20.5%)	
2,6-dimethylcyclohexanol	MS+RI	95	1119	1114 [2]	832.2 (33.0%)	2116.5* (32.7%)	
Ketones							
2,4,4-trimethylcyclopentanone	MS	83	1010		15.1 (19.3%)	62.2* (44.6%)	
2,2,6-trimethylcyclohexan-1-one	MS+RI	82	1042	1047 [3]	104.2 (32.6%)	229.8** (13.9%)	
isophorone	MS+RI	82	1062	1080 [4]	51.5 (36.7%)	151.0** (18.2%)	
Lactones							
dihydro-5-methyl-2(3H)-furanone	MS+RI	56	953	954 [1]	31.1 (6.3%)	50.5* (22.3%)	
Sulfur compounds							
carbone disulfide	MS+RI	76	570	568 [5]	526.8 (46.3%)	1102.0* (9.5%)	
Unknown							
unknown		69	1087		1.7 (41.3%)		3.4* (12.8%)
unknown		77	1132		1.6 (18.6%)	3.1* (28.1%)	
unknown		55	1162		4.7 (18.5%)	15.1** (12.8%)	

^a MS + RI, mass spectrum and RI agree with literature data; MS, mass spectrum agrees with literature spectrum

^b Mass fragment used for area determination

^{c,d} Retention indices on a DB5 capillary column from experimental run (c) or bibliographic data (d)

[1] Engel and Ratel, 2007; [2] Radulovic et al., 2010; [3] Cajka et al., 2007; [4] Aaslyng et al., 1998; [5] Beaulieu and Grimm, 2001.

 e $\,$ Diet contaminated with the target concentration of 5 or 50 ng $\alpha\text{-HBCDD g}^{\text{-1}}$

* *p* <0,05

** *p* <0,01

Level found higher in liver from "control" animals compared to "exposed animals"

arker with its standard deviation (in bracket).

