

Fate of polychlorobiphenyls in Tenebrio molitor larvae: consequences for further use as food and feed

J. Ratel, F. Mercier, J. Rivas, H. Wang, M. Angénieux, B. Calmont, S.

Crépieux, Christelle Planche, E. Engel

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1. Introduction

 The United Nations estimate that the global human population is expected to reach 9.7 billion in 2050 increasing the consumption of animal products and the growing demand for feed ingredients (United Nations, 2019). Available resources are currently limited and may raise 49 environmental, societal, and economic questions (Tran *et al.*, 2015). The quest for alternative sustainable animal protein sources is therefore expected to become a considerable issue in the feed market. Edible insects could be an interesting solution with a high nutritional value (Van Huis, 2021). Processed animal protein derived from insects are currently authorised for aquafeeds (Commission Regulation (EU) 2017/893) (EU, 2017), poultry and swine feeds in the EU (Commission Regulation (EU) 2021/1372) (EU, 2021). Among the broad variety of insect 55 species, the yellow mealworm (*Tenebrio molitor*) is one of the most widely bred and traded 56 insect species in Europe (Bordiean et al., 2020).

 To develop this new sector in future years, it is essential to assess and control the risks associated with the use of insects. Some studies are already available about the microbiological hazards (Vandeweyer et al., 2021). However, there is a lack of knowledge concerning the chemical safety while some chemical contaminants may be found in insect farming environment or in their feeding substrates, can be produced during processing methods or can 62 migrate from packaging material to insect products (Meyer *et al.*, 2021). While some authors have reported that insects may degrade some mycotoxins (Niermans et al., 2019), some antimicrobials (Cai et al., 2018) or even some pesticides (De Almeida et al., 2017), other chemical contaminants like heavy metals can have a negative impact on insect development inducing, for example, an increase in larval mortality (Bulak et al., 2018). Moreover, previous 67 studies have shown that insects like T. molitor can bioaccumulate toxic trace elements during rearing, with very variable bioaccumulation factors up to 1.7 for cadmium, 6.2 for mercury and 69 even 34 for lead (Truzzi *et al.*, 2019).

 Among chemical contaminants, persistent organic pollutants (POPs) can be transferred from environment or feed to animal products and therefore are of serious concern in animal-72 derived food products (Engel et al., 2015). In the black soldier fly (Hermetia illucens), Van der 73 Fels-Klerx et al. (2020) have determined that the bioaccumulation factors of POPs from their substrates (different whole meals and different types of snack products, both with paperboard carton and with plastic packaging materials) ranged between 0.3 and 1.2 for polycyclic aromatic hydrocarbon (PAHs) and between 1.0 and 2.0 for the sum of dioxins and dioxin-like polychlorinated biphenyls (PCBs). However, to our knowledge, there are no studies regarding 78 the bioaccumulation of POPs in T. molitor.

79 Using T. molitor larvae as a model and setting up a pilot experimental farming under controlled and safe conditions allowing the implementation of an exposure study with chemical contaminants, the present work explores the ability of insects to bioaccumulate PCBs from their feeding substrate during the larvae growth. For this purpose, the first part of this paper is 83 focused on the set up of an analytical method to monitor PCBs in T . *molitor* larvae and the assessment of the performance of this method in terms of linearity (coefficients of 85 determination, R^2) and sensitivity (limits of detection, LOD). In a second part, the impact of a feed contamination by a mix of 6 non-dioxin like PCBs (nDL-PCBs) on the growth of larvae was studied. The bioaccumulation factors of PCBs in larvae were determined at the end of the rearing and will be discussed in the context of a further use of larvae as food and feed.

2. Materials and Methods

Chemicals

 Hexane, dichloromethane, acetone and toluene were organic trace analysis grade solvents (Sigma-Aldrich, Saint-Quentin Fallavier, France). For PCB extraction, Florisil® and activated aluminium oxide (acidic, Brockmann I) were from Sigma-Aldrich. Diatomaceous earth was obtained from Thermo Fisher Scientific (Waltham, MA, USA). The certified reference material AE-00059-H-2X of AccuStandard Europe (Niederbipp, Switzerland) was used for the development of PCB analysis. This mix is composed of the 6 nDL-PCBs (IUPAC numbers 28, 52, 101, 138, 153, and 180) with individual concentrations of 20 ppm. Fluorinated internal standards 3-F-PCB-52 and 5'-F-PCB-126 (Chiron, Trondheim, Norway) were used for the accurate quantification of target compounds.

Larvae 12 100

Feeding 14 101

 Larval feed was dry wheat bran produced by local flour mills in Puy de Dome (France). Larvae were fed with non-contaminated feeds (control group) or with PCB contaminated feeds (exposed groups). For exposed groups, the bran was spiked according to a protocol adapted from Planche *et al.* (2015), based on a contaminant addition to the feed via a volatile solvent. For each PCB exposure condition, the spiking was carried out on the total quantity of wheat bran necessary for all the individual rearing (45g), allowing to ensure the quality of replicates. Briefly, 45 g of dry wheat bran were set in a glass jar (9.5cm high \times 7cm wide) then immersed in acetone (125 mL) containing the nDL-PCBs. The mixture was evaporated down under a hood and roughly homogenised. Three spiking concentrations, 0.67, 4.0 and 24.4 ppb (ng 111 PCB/g wheat bran), were tested for each PCB congener. This corresponds to levels of $\Sigma(6 \text{ nDL-}$ PCBs) that are 0.4, 2.4 and 14.6 times the maximum level (ML) set at 10 ppb by the Regulation ₁₃ (EU) No 277/2012 in feed materials of plant origin. The spiking concentrations are consistent both with the regulation issues and with the scarce literature data dealing with PCB contamination in food of plant origin, like wheat bran, with for example some concentrations of PCBs in bran that can reach 1.84 ppb (Roszko et al., 2014). For the control groups, the bran was either unspiked (bran without solvent or PCB) or spiked with neat solvent (bran with solvent but without PCB). 19 104 20 105 24 108 25 109 26 110 $29 \overline{)}$ 112 30 113 31 114 35 117 36 118

Rearing

120 Larvae of T. molitor were produced by the startup INVERS (Saint-Ignat, France). Using a balance (Precisa 410 AM-FR, Precisa Instruments Ltd., Switzerland; d: 0.0001), 10g of larvae at L3 growth stage (approximatively 200 individuals) were set on a glass jar containing 45 g of control or contaminated wheat bran. Three glass jars for each modality were prepared and put 124 in an individual incubator (18 L HerathermTM Compact Microbiological Incubator, Thermo Scientific) customised for sufficient ventilation (300 L/h) (Figure 1). In total, 15 rearing glass jars were distributed in the 5 incubators. These replicates were not randomly distributed over 127 the incubators to avoid a cross contamination between the different exposure conditions. The humidity was maintained at 60% and the temperature regulated at 26° C with a monitoring humidity was maintained at 60% and the temperature regulated at $26\degree$ C with a monitoring throughout the rearing (probe: VAISALA HMP110; data acquisition system: AOIP SA32) (Supplementary Figure S1). A water spray was added to each jar containing larvae two times per day. The rearing of larvae was carried out for 20 days in incubators. From the 6th day of the test, the evaluation of larval weight was performed for each trial twice per week. Larvae were separated by using a 1.5 mm mesh sieve. After sieving, the weight of 20 larvae was determined. After each weighting, the sieved content and the larvae were put back in the jar. 42 121 43 122 44 123 48 126 49 127 53 130 54 131 55 132 58° 134

Figure 1: Pilot experimental farming of Tenebrio molitor larvae. 60 135

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Analysis of PCBs

Sample preparation

139 At the end of the rearing period, all insects were sacrificed by freezing at -20° C. Larvae were then microwave-dried according to the usual conditions carried out by INVERS company: the power of the microwave was fixed (600 W) and the duration of the drying adapted so that the final relative humidity of the larvae was inferior to 10% (between 5 and 7 min). Prior to extraction, all samples were powdered with a liquid nitrogen grinder. Extraction process was carried out according to Planche *et al.* (2017) with slight modifications. Briefly, 1 g of larvae 13 144 powder was extracted by accelerated solvent extraction (ASE) using a Dionex ASE 350 14 145 15 146 extractor (Sunnyvale, CA). Stainless-steel extraction cells (22 mL volume) were used, with 5 g of acidic alumina and 5 g Florisil® placed at the bottom of the cells. Paper filters were placed at the bottom and top of the alumina layer. The cells were then filled with 1 g of ground larvae dispersed in diatomaceous earth and hexane was used as extraction solvent at a temperature of 19 149 20 150 100 °C and pressure of 1500 psi. ASE extraction included heating (5 min), static time (5 min) and purging (90 s) with two extraction cycles per sample. After filtration through a 1.2 µm glass fiber prefilter and a 0.45 um nylon filter (Phenomenex, Torrance, CA), the extract was 24 153 evaporated (Rocket; Genevac Ltd, Ipswich, UK) using toluene as a keeper to minimise analyte losses during the evaporation step, then 4 mL of dichloromethane were added. To clean up 25 154 26 155 extracts, gel permeation chromatography (GPC) (Gilson, Middleton, WI) was carried out on an S-X3 Bio-Beads column (Bio-Rad, Philadelphia, PA) using dichloromethane as eluting solvent at a flow rate of 5 mL/min. The fraction collected between 15 and 37 min was evaporated to 30 158 dryness (Rocket, Genevac Ltd), and a mix of 100 µL of hexane and internal standard was added $(Figure 2)$. 31 159

Figure 2: Simplified diagram of the workflow used to analyse PCBs in *Tenebrio molitor*
larvae analyse PCBs in Tenebrio molitor
and the same analyse PCBs in Tenebrio molitor larvae

PCB detection by GC-ToF/MS

 For PCB identification in insect matrix, larvae extracts (obtained according to the conditions detailed in the previous section) were spiked with nDL-PCBs then analyzed with a time-offlight mass spectrometer (Pegasus 4D, Leco) coupled to a gas chromatograph (6890, Agilent Technologies). 1µL of extract was injected at 280°C in splitless mode into a DB-5MS capillary 168 column (60 m \times 0.32 mm \times 1 µm; Agilent J&W) with Helium (purity of 99.99995%) as carrier gas at 1 mL/min. Oven temperature was held at 120°C for 1 min, then ramped up to 240°C at a gradient of 20°C/min and to 300°C at a gradient of 2°C/min, and held at 300°C for 15 min. The MS-temperatures were set at 230 °C, 150 °C and 180 °C in the transfer line, the source and the quadrupole, respectively. Electron impact energy was set at 70 eV, and data was collected in full scan in the range of 45 to 600 m/z at a scan range of 10 scans per second. 44 166 45 167 50 171 51 172

174 PCB quantification by GC - μECD

The PCB levels in the larvae experimentally reared were determined by μ ECD (Agilent) coupled to a gas chromatograph (6890, Agilent Technologies). The GC parameters were the 177 same as for the GC-MS coupling. The micro-ECD system was operated at 300 °C using data 57 175 58 176

 acquisition rate of 50 Hz, with nitrogen as make-up gas at a flowrate of 40 ml/min. Calibration curves were preliminary built from dried larvae extracts spiked with the 6 nDL-PCBs (see Table 1) at the following concentrations: 0.2, 0.5, 1.3, 3.0, 8.0, 20.0, 50.0 ng/g. Larvae extracts without PCB added were run in triplicate to check the absence of targeted analytes. Each concentration level was analysed 4 times by GC-µECD. Peak areas were determined, normalized by internal standard, then used for calculations of the standard deviations and the calibration curve equations. The linearity of the calibration curves was assessed for each PCB 185 by calculating the coefficients of determination (R^2) . The limit of detection (LOD), using the definition 3s/m (s is the standard deviation of the intercept, and m is the slope of the linear calibration curve), was determined from the calibration curves for each individual PCB studied. 11 187

Data treatment

 Data were processed with the TIBCO's Statistica software (version 13.0, TIBCO Software Inc.) and the R software (version 3.5.1, The R Foundation for Statistical Computing). The " lm " (linear model) function of R was used on the calibration curve data for the determination of R^2 as well as s and m requested for LOD calculation. Student's t test or one-way analyses of 193 variance (ANOVA) followed by *post hoc* Newman–Keuls test were performed to compare 194 means data. The differences were considered significant when $p < 0.05$. 17 190 18 191 19 192

3. Results

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PCB analysis

The recovery rates of the 6 nDL-PCBs measured after spiking in dried powder of T . molitor (at 200 ng/g dry matter), extraction, purification and concentration are presented in 202 Table 1. All the recovery rates lay in the classically accepted range of 70–130% according to the EPA Method 8000C (2003), with RSD between 10% and 18% .

Table 1: Recovery rates (%) obtained after spiking, extraction, purification, 206 concentration and analysis of PCBs in dried powder of *Tenebrio molitor* larvae $(n=3)$.

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209 The LOD measured with GC-µECD for the 6 nDL-PCB congeners in dried powder of T. molitor larvae varied from 1.67 ng/g for PCB 153 to 2.98 ng/g for PCB 101, with limits of ²¹¹ quantification (LOQ) from 5.56 to 9.93 ng/g respectively (Supplementary Table S1). ²¹² Coefficients of determination ranged between 0.94 for PCB 101 and 0.98 for PCBs 138, 153 213 and 180. 1

²¹⁶ Impact of feed contamination with PCBs on larval growth

Table 2 and Figure S2 present the weight of T. molitor larvae at days 1, 7, 10, 14, 17 and ²¹⁹ 20 of rearing depending on their feeding substrate: wheat bran unspiked, spiked with neat solvent without PCB or spiked with nDL-PCBs at 0.67, 4.0 or 24.4 ng/g corresponding to 0.4, 221 2.4 and 14.6 times the maximum level (ML) for the nDL-PCBs in feed materials of plant origin, respectively (Commission regulation (EU) No 277/2012). One-way ANOVA (p <0.05) revealed no significant difference between the different conditions with a mean weight of 54.1 mg per larvae at the beginning of the study and a continuous growth throughout the rearing up to 95.6 mg per larvae on average after 20 days of rearing. Visual observations throughout the rearing 226 period did not reveal differences in terms of larval mortality between the different conditions.

Table 2: Weight of *Tenebrio molitor* larvae (mg) throughout their rearing on wheat 229 bran either unspiked (Control), spiked with neat solvent (Control + acetone) or spiked at $0.67, 4.0$ or 24.4 ng PCB/g wheat bran. Data represent mean larvae weight (mg) \pm SD which 0.67, 4.0 or 24.4 ng PCB/g wheat bran. Data represent mean larvae weight (mg) \pm SD which is determined from the 20 larvae measurement for each rearing glass jar ($n=3$ glass jars for each exposure condition).

NS: not significant $(p>0.05)$

 1_p values were calculated for one-way analyses of variance (ANOVA) followed by post hoc Newman-Keuls test.

Bioaccumulation factors

²⁴⁰ Bioaccumulation factors (BAF) were calculated for each congener of PCB in fresh (Table 3) and dried (Table S4) T. molitor larvae according to Truzzi *et al.* (2019) and Van der Fels-

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242 Klerx *et al.* (2020), with BAF = concentration of PCB in the organism (fresh or dried larvae) / concentration of PCB in the feed provided (dried wheat bran).

 For a spiking concentration of 0.67 ng PCB/g wheat bran, concentrations in dried larvae extracts of the 6 nDL-PCBs were below the detection limits detailed in Supplementary Table S1. For a spiking concentration of 4 ng PCB/g wheat bran, the concentrations of PCB 28, 52 247 and 101 were between LOD and LOQ (Supplementary Table S2). For all other conditions, PCB congener concentrations are above the LOQ. Regarding these last values, Supplementary Table S4 shows that BAF obtained in dried larvae ranged from 1.0 to 2.8 (raw data enabling BAF calculation given in Table S3). For PCB 138, there is no significant difference in BAF obtained between the two highest spiking levels whereas for PCBs 153 and 180, BAF significantly increased with the spiking level. Moreover, BAF obtained for PCB 52 are significantly lower than for other congeners.

 Table 3 shows the BAF of the 6 nDL-PCBs congeners in fresh larvae. BAF in fresh larvae are 2.8 times lower than in dried larvae.

257 Table 3: Bioaccumulation factors (BAF) of PCB congeners in Tenebrio molitor larvae on a fresh weight basis after 20 days of rearing on wheat bran spiked at 0.67, 4.0 or 24.4 ng PCB/g. These values were estimated from data obtained on dried larvae, taking into account the impact of drying on larval weight. Data represent mean $BAF \pm SD$ which is determined 261 from 3 analytical replicates of each rearing glass jar $(n=3)$ glass jars for each exposure condition). condition).

NA: non-available (PCBs non-detected in larvae extracts)

a-b: different superscript letters within the same row indicate significant differences among values $(p<0.05)$. BAF are determined from quantification values in larvae extracts either between LOD and LOQ $(*)$ or greater than LOQ.

4. Discussion

273 In order to assess the fate of the 6 nDL-PCBs in T. molitor larvae during rearing with a contaminated substrate, the first step of this study was to determine their recovery rates after the spiking, extraction, purification, concentration and analysis protocol. Based on a protocol 276 adapted from Planche *et al.* (2015) with a delipidation strengthened by the addition of fat retainers, the PCB recoveries in larvae samples (from 98% to 129%) are of the same order of retainers, the PCB recoveries in larvae samples (from 98% to 129%) are of the same order of magnitude as those in raw beef samples (from 87% to 133%) (Planche et al., 2015). This demonstrates that the high fat level of T. molitor larvae $(56-61\%$ in whole dried T. molitor $(EFSA, 2021)$ vs 11% in raw ground beef) does not decrease the recovery rates obtained even

281 if PCBs are hydrophobic compounds (Log Kow $= 4.09 - 8.18$ according to Hawker and Connell ²⁸² (1988)), thereby confirming the relevance of the protocol used.

283 Based on this protocol, the impact of a contamination by PCBs on T. molitor was assessed ²⁸⁴ using a pilot experimental farming where larvae were fed for 20 days with wheat bran either ²⁸⁵ unspiked, spiked with neat solvent without PCB or spiked with nDL-PCBs at 0.67, 4.0 or 24.4 286 ng g^{-1} . No significant differences were observed in terms of larval growth between the group 287 fed with unspiked wheat bran and the group fed with wheat bran spiked with neat solvent ²⁸⁸ suggesting that the spiking step had no impact on the further dietary intake of larvae. This 289 confirms that the use of acetone as an inert PCB "vehicle" volatile solvent is relevant contrary, for example, to a mix of chloroform and methanol which has led to a lower T . *molitor* body weight when it was used as a vehicle to spike their feed with Aflatoxin B1 (Bosch et al., 2017). 11 290 12 291

Moreover, a high PCB concentration in feed has no more impact than a low concentration 293 on the larvae body weight of T. molitor contrary to previous results obtained with antimicrobials 294 by Gao et al. (2019). These authors showed that the weight of H . illucens larvae declined gradually with the increase in sulfonamide concentration from 0 to 10 mg/kg feed (Gao et al., 2019). The present study indicates that a contamination of T . *molitor* by PCBs in an insect farm 297 could not be revealed by monitoring the larvae weight, suggesting that a systematic control of ²⁹⁸ the quality of insect feed substrates used is essential. It would be interesting to determine if these results would be similar with dioxin-like PCBs that have a different mechanism of action than nDL-PCBs on cellular pathways (Elnar *et al.*, 2012). 13 292 17 295 18 296 22 299 23 300

301 In order to determine the capacity of the 6 nDL-PCBs to be transferred from feed to T. 302 molitor larvae, bioaccumulation factors (BAF) were determined for each PCB congener. If we 303 take into consideration BAF determined from quantification values in larvae extracts greater than LOQ, only BAF results for PCBs 138, 153 and 180 can be discussed for a spiking ³⁰⁵ concentration of 4 ng PCB/g wheat bran (Tables 3 and S4). For PCBs 153 and 180, BAF related 306 to the highest spiking concentration (24.4 ng PCB/g wheat bran) are significantly higher ³⁰⁷ compared to lower spiking concentration (4.0 ng PCB/g wheat bran), suggesting that, for these 308 congeners, BAF increases with the contamination level. 28 304 29 305

Knowing that a BAF greater than 1 suggests a bioaccumulation of PCBs from wheat bran into T. molitor larvae, the BAF obtained with fresh larvae (from 0.4 to 1.0; Table 3) indicate ³¹¹ that there is no bioaccumulation of PCBs in larvae. However, PCBs are transferred from the ³¹² diet to the larvae during rearing and are then concentrated in larvae during the drying process. This concentration effect explains why the BAF calculated on the basis of dried larvae are 2.8 times higher (from 1.0 to 2.8 ; Supplementary Table S4). 33 34 309 35 310 36 37 38 39 313 40 314

³¹⁵ Regarding previous data on nDL-PCBs, Van der Fels-Klerx et al. (2020) reported BAF in ³¹⁶ H. illucens dried larvae from 0.8 to 1.2 (based on upper bound values) after rearing on meat or ³¹⁷ vegetarian substrate containing 3-6% of plastic or paperboard carton packaging material. Their lower values of BAF compared to those obtained in the present study may be explained by the differences in terms of insect species and feeding substrate used. Moreover, the starvation 320 period of 24h set up by Van der Fels-Klerx *et al.* (2020) before harvesting may allow to renew ³²¹ the digestive tract content and thus may decrease the concentration of nDL-PCBs in larvae and therefore the BAF. 41 42 43 44 317 45 318 46 319 47 48 49 50 322

PCBs are hydrophobic compounds (Log Kow $= 4.09 - 8.18$ according to Hawker and Connell (1988)) which implies that they may be transferred in the lipid fraction of larvae during 325 rearing. Since the drying process does not impact this lipid fraction, it leads to a concentration ³²⁶ of PCBs in dried larvae and therefore an increase of BAF. When a PCB contamination is detected in larvae feed substrate, it could therefore be envisaged not to use the whole dried larvae for food and feed but to exploit separately the protein and lipid fractions. In order to limit ³²⁹ the risks linked to the presence of chemical contaminants, the lipid fraction (where PCBs are 51 323 52 324 53 54 55 326 56 327 57 328 58

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 accumulated) could then be used for non-food applications (biofuel production for example) while the protein fraction could possibly be used for food and feed. 1 331

5. Conclusion

 The pilot experimental farming set up enabled to assess the bioaccumulation of PCBs in T. molitor larvae during rearing. A feed substrate contamination with PCBs doesn't seem to have any significant impact on larval development without negative effects on growth. It could be supposed that a much higher PCB level than those assessed in this study could have a significant impact on the development of T . *molitor*, but these concentrations would be unrealistic in insect feed substrate. In addition, it would be interesting to assess the impact on larval development of a longer rearing period on a contaminated substrate. The present study revealed that there is no bioaccumulation of PCBs in T. molitor larvae during rearing. However, PCBs can be transferred from the diet to the larvae and are then concentrated in larvae during the drying process. This highlights the significant impact of the drying process which induces a concentration by a factor of almost 3 of PCBs in larvae. Knowing that the nDL-PCB concentration maximum limit is set at 10 µg/kg either for feed materials of plant origin and for feed materials of land animal origin (Commission regulation, EU No 277/2012), the use as larvae feeding substrate of wheat bran containing a residual amount of nDL-PCBs could result, after the rearing period, in dried larvae with an almost 3-fold higher nDL-PCB concentration that could potentially exceed the maximum limit for its use as feed. This highlights the importance of monitoring the quality of the feed substrates used for farming insects for food and feed in order to guarantee a safe content in chemical contaminants including persistent organic pollutants like PCBs. When a PCB contamination is detected in larvae feed substrate, it could therefore be envisaged to use the larvae lipid fraction (where PCBs are accumulated) for non-food applications. Finally, in order to ensure food safety throughout the food chain, it would be interesting to assess the PCB biomagnification factors 357 if T. molitor larvae are used as feed, for example for aquaculture since it is expected that most insect production will be targeted towards aquafeed by 2030 (Van Huis, 2022). 11 339 12 340 13 341 17 344 18 345 23 349 24 350 28 353 29 354 34 358

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- 64 65

- Microwave: moisture content <10%
(800W; 6min)
- Cryogenic grinding: 45s

- 22ml cells
- 1g of larvae
- 5g Alumine + 5g Florisil
- Internal standard
- Hexane 100%

- Low pressure
steam heat transfer
- 40°C / 1h

- S-X3 bioBeads column
- DCM
- 5mL.min⁻¹

Figure 2

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Fate of polychlorobiphenyls in the insect-Tenebrio molitor larvae: consequences for further use as food and feed

J. Ratel¹, F. Mercier¹, J. Rivas¹, H. Wang¹, M. Angénieux¹, B. Calmont², S. Crépieux³, C. Planche^{1*}, E. Engel¹

Figure S1: Temperature (C) and relative humidity (C) readings of each incubator during the larvae rearing.

Figure S2: Weight of *Tenebrio molitor* larvae (mg) throughout their rearing on wheat bran either unspiked (Control), spiked with neat solvent (Control + acetone) or spiked at 0.67, 4.0 or 24.4 ng PCB/g wheat bran. Data represent mean larvae weight (mg) \pm SD which is determined from the 20 larvae measurement for each rearing glass jar ($n=3$ glass jars for each exposure condition). exposure condition).

Table S1: Performance of GC-µECD for quantification of the 6 nDL-PCB congeners in dried powder of Tenebrio molitor larvae (linearity range: 0.2-50.0 ng/g).

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Table S2: Concentrations (ng/g) of PCB congeners in Tenebrio molitor larvae on a dried weight basis after 20 days of rearing on wheat bran spiked at 0.67, 4.0 or 24.4 ng PCB/g. For each exposure condition, three analytical replicates of each rearing glass jar were carried out $(n=3$ glass jars for each exposure condition).

NA: non-available (non-detected in larvae extracts)

LOD: Limit of detection

* Quantification values between LOD and LOQ (see Table S1)

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Table S3: Raw data enabling the calculation of bioaccumulation factors (BAF) of PCB congeners in Tenebrio molitor larvae on a dry weight basis after 20 days of rearing on wheat bran spiked at 0.67, 4.0 or 24.4 ng PCB/g.

NA: non-available (PCBs non-detected in larvae extracts)

^a For each exposure condition, three analyses (R1 to R3) of approximatively 1 g of dried larvae sampled in each of the 3 glass jars (1 to 3) were carried out.
^b Internal standard (5'F-PCB-126) used for the accurate qu

Interna standard on the state of the formulation of the formulation $\frac{1}{2}$. (Area PCB_x/Area IS) – b)/a]/(Sample weight × CF)

SPCB concentration in larvae was calculated according to the formula: Concentration = [((A

⁴ BAF was calculated according to the formula: BAF = $(Lawae PCB_x concentration) / (When the PCB_x concentration)$

Table S4: Bioaccumulation factors (BAF) of PCB congeners in Tenebrio molitor larvae on a dry weight basis after 20 days of rearing on wheat bran spiked at 0.67, 4.0 or 24.4 ng PCB/g. Data represent mean BAF \pm SD which is determined from 3 analytical replicates of each rearing glass jar $(n=3)$ glass jars for each exposure condition).

NA: non-available (PCBs non-detected in larvae extracts)

a-b: different superscript letters within the same row indicate significant differences among values (p <0.05). BAF are determined from quantification values in larvae extracts between LOD and LOQ (*) or greater than LOQ.

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Supplementary material

Fate of polychlorobiphenyls in Tenebrio molitor larvae: consequences for further use as food and feed

J. Ratel¹, F. Mercier¹, J. Rivas¹, H. Wang¹, M. Angénieux¹, B. Calmont², S. Crépieux³, C. Planche^{1*}, E. Engel¹

Figure S1: Temperature $(°C)$ and relative humidity $(°_o)$ readings of each incubator during the larvae rearing.

Figure S2: Weight of *Tenebrio molitor* larvae (mg) throughout their rearing on wheat bran either unspiked (Control), spiked with neat solvent (Control + acetone) or spiked at 0.67, 4.0 or 24.4 ng PCB/g wheat bran. Data represent mean larvae weight (mg) \pm SD which is determined from the 20 larvae measurement for each rearing glass jar ($n=3$ glass jars for each exposure condition). exposure condition).

Table S1: Performance of GC-µECD for quantification of the 6 nDL-PCB congeners in dried powder of Tenebrio molitor larvae (linearity range: 0.2-50.0 ng/g).

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Table S2: Concentrations (ng/g) of PCB congeners in Tenebrio molitor larvae on a dried weight basis after 20 days of rearing on wheat bran spiked at 0.67, 4.0 or 24.4 ng PCB/g. For each exposure condition, three analytical replicates of each rearing glass jar were carried out $(n=3$ glass jars for each exposure condition).

NA: non-available (non-detected in larvae extracts)

LOD: Limit of detection

* Quantification values between LOD and LOQ (see Table S1)

CONJFIDENTIVAL

Table S3: Raw data enabling the calculation of bioaccumulation factors (BAF) of PCB congeners in Tenebrio molitor larvae on a dry weight basis after 20 days of rearing on wheat bran spiked at 0.67, 4.0 or 24.4 ng PCB/g.

NA: non-available (PCBs non-detected in larvae extracts)

^a For each exposure condition, three analyses (R1 to R3) of approximatively 1 g of dried larvae sampled in each of the 3 glass jars (1 to 3) were carried out. ^b Internal standard (5'-F-PCB-126) used for the accurate quantification of the nDL-PCBs

PCB concentration in larvae was calculated according to the formula: Concentration = $[((\text{Area PCR})/\text{Area IS}) - b)/a]/(\text{Sample weight} \times \text{CP})$
where CF is the concentration factor due to the extraction procedure (CF=10), and a and b are the slo

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⁴ BAF was calculated according to the formula: BAF = $(Lawae PCB_x concentration) / (When the PCB_x concentration)$

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NA: non-available (PCBs non-detected in larvae extracts)

a-b: different superscript letters within the same row indicate significant differences among values (p <0.05). BAF are determined from quantification values in larvae extracts between LOD and LOQ (*) or greater than LOQ.

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