# **SCIENTIFIC OPINION**

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# Update of the risk assessment of hexabromocyclododecanes (HBCDDs) in food

# **Panel on Contaminants in the Food Chain (CONTAM Panel)**

Dieter Schrenk, Margherita Bignami, Laurent Bodin, James Kevin Chipman, Jesús del Mazo,
Bettina Grasl-Kraupp, Christer Hogstrand, Laurentius (Ron) Hoogenboom, Jean-Charles
Leblanc, Carlo Stefano Nebbia, Elsa Nielsen, Evangelia Ntzani, Annette Petersen, Salomon
Sand, Tanja Schwerdtle, Heather Wallace, Diane Benford, Peter Fürst, Martin Rose, Sofia
Ioannidou, Marina Nikolic, Luisa Ramos Bordajandi and Christiane Vleminckx

# 9 Abstract

Commission asked EFSA to update its 2011 risk assessment on 10 The European 11 hexabromocyclododecanes (HBCDDs) in food. HBCDDs, predominantly mixtures of the stereoisomers 12  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCDD, were widely used additive flame retardants. Concern has been raised because of 13 the occurrence of HBCDDs in the environment, food, and in humans. Main targets for toxicity are 14 neurodevelopment, the liver, thyroid hormone homeostasis and the reproductive and immune systems. 15 HBCDDs are not genotoxic and the available evidence indicates that they are not carcinogens. The CONTAM Panel concluded that the neurodevelopmental effects on behaviour in mice can be considered 16 17 the critical effects. Based on effects on spontaneous behavior in mice, the Panel identified a LOAEL of 0.9 mg/kg bw as the Reference Point, corresponding to a body burden of 0.75 mg/kg bw. The chronic 18 19 intake that would lead to the same body burden in humans was calculated to be 2.35 µg/kg bw per 20 day. The derivation of a health-based guidance value (HBGV) was not considered appropriate. Instead, 21 the margin of exposure (MOE) approach was applied to assess possible health concerns. Over 6,000 22 analytical results for HBCDDs in food were used to estimate the exposure across dietary surveys and 23 age groups of the European population. The most important contributors to the chronic dietary LB 24 exposure to HBCDDs were fish meat, eggs, livestock meat and poultry. The CONTAM Panel concluded 25 that the resulting MOE values support the conclusion that current dietary exposure to HBCDDs accross 26 European countries does not raise a health concern. An exception are breastfed infants with high milk 27 consumption, for which the lowest MOE values may raise a health concern.

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Keywords: hexabromocyclododecanes, HBCDDs, occurrence, food, toxicology, human exposure, risk
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- 35 Correspondence: contam@efsa.europa.eu
- 36

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Panel members: Margherita Bignami, Laurent Bodin, James Kevin Chipman, Jesús del Mazo, Bettina

57 Panel members: Margherita Bignami, Laurent Bodin, James Kevin Chipman, Jesus del Mazo, Bettina
 58 Grasl-Kraupp, Christer Hogstrand, Laurentius (Ron) Hoogenboom, Jean-Charles Leblanc, Carlo Stefano
 59 Nebbia, Elsa Nielsen, Evangelia Ntzani, Annette Petersen, Salomon Sand, Dieter Schrenk, Tanja
 60 Schwerdtle, Christiane Vleminckx, Heather Wallace.

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# 75 **Summary**

Brominated flame retardants (BFRs) are anthropogenic chemicals, which are used in a wide variety of consumer/commercial products in order to improve their resistance to fire. Concern has been raised because of the occurrence of several chemical compouds from the group of BFRs in the environment,

food, and in humans. This has led to bans on the production and use of certain formulations.

The European Commission asked EFSA to update its 2010–2012 risk assessments on the different families of BFRs, i.e. hexabromocyclododecanes (HBCDDs), polybrominated diphenyl ethers (PBDEs), tetrabromobisphenol A (TBBPA) and its derivatives, brominated phenols and their derivatives, and novel and emerging BFRs.

- The similarities in chemical properties and effects seen in the previous EFSA assessments for the different BFR families warrant the consideration of a mixture approach. The Panel on Contaminants in the Food Chain (CONTAM Panel) will evaluate the appropriateness of applying a mixture approach in an additional Opinion once the risk assessment for each BFR family has been updated. It will be based on the EFSA Guidance on harmonised methodologies for human health, animal health and ecological risk
- assessment of combined exposure to multiple chemicals.

This first Opinion updates the risk assessment of HBCDDs in food previously performed by EFSA and published in 2011. The assessment focuses on 1,2,5,6,9,10-HBCDD, which is predominantly composed of the three stereoisomers  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCDD.

HBCDDs are additive flame retardants used mainly in expanded and extruded polystyrene applied as
 construction and packing material and in textiles. HBCDDs were permitted for general use in the EU
 until August 2015, after which only authorised applications were allowed because of health concerns.

96 The present assessment takes into account the occurrence data in food and biological samples, 97 submitted after the publication of the former Opinion, as well as newly available scientific information 98 on hazard identification and characterisation.

99 The analytical determination of HBCDDs is performed either by gas chromatography (GC)/mass 100 spectrometry (MS) or liquid chromatography (LC)/MS based methods. Both techniques enable the use 101 of isotope-labelled internal standards, which allow for the correction for losses during extraction and 102 clean-up. While GC/MS based methods determine Total HBCDDs as they cannot separate the 103 stereoisomers, LC/MS based methods allow the specific analysis of the individual HBCDD stereoisomers.

# 104 Hazard identification and characterisation

105 In rodents, absorption of HBCDD stereoisomers from the gastrointestinal tract is rapid and almost 106 complete (>80%). HBCDDs and/or their metabolites are distributed to a number of tissues, including 107 adipose tissue, muscle, liver, skin and brain. Following a single oral administration in rats,  $\alpha$ -HBCDD 108 and  $\beta$ -HBCDD accumulate in adipose tissue, while  $\gamma$ -HBCDD accumulates in liver and adipose tissue. In 109 repeated dose experiments in mice,  $\alpha$ -HBCDD and  $\beta$ -HBCDD accumulate mainly in adipose tissue, while 110 y-HBCDD accumulates in the liver. Metabolic debromination and hydroxylation of HBCDDs have been 111 reported. Conversion of  $\gamma$ -HBCDD to  $\alpha$ - and  $\beta$ -HBCDD has been reported, but no stereoisomerisation of 112  $\alpha$ -HBCDD. Elimination of HBCDD stereoisomers in rodents is predominantly via faeces. The elimination 113 half-lives of HBCDD stereoisomers in female mice, based on concentrations in adipose tissue, vary from 114 3–4 days for y-HBCDD, 2.5 days for  $\beta$ -HBCDD, and 17 days for  $\alpha$ -HBCDD.

115 In humans, in the only available study, the half-life was estimated to be 64 days (range 23–219 days)

for the sum of  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCDD stereoisomers. This difference in kinetics affects the extrapolation

117 of animal data to humans.

Transfer of HBCDDs was studied in different animal species (laying hens, broilers, ducks, pigs and fish).
HBCDDs are distributed to a number of tissues and accumulate in tissues with high fat content and
eggs.

121 The acute toxicity of HBCDDs is low. Toxicological studies have been carried out using different 122 experimental designs with single or repeated administration during gestation, postnatally or in adulthood 123 using, in most cases, HBCDDs with no information on the stereoisomer composition specified. Main 124 targets for HBCDD toxicity are neurodevelopment, the liver, thyroid hormone homeostasis and the 125 reproductive and immune systems.

126 HBCDDs induced increased liver weight and hepatocellular hypertrophy in rats and mice. Increased 127 thyroid weight, follicular hypertrophy, hyperplasia and/or colloid depletion and changes in thyroid 128 hormone levels were also reported in these species. HBCDDs decreased the fertility index and increased 129 pup mortality during lactation in a 2-generation toxicity study in rats. In addition, HBCDDs decreased 130 testes weight and delayed vaginal opening in female pups in a 1-generation study in rats. Other 131 endocrine-related effects were changes in levels of sex-steroid hormones (testosterone and oestradiol) 132 in mice, decreased number of primordial ovarian follicles or reduction in the number of growing follicles 133 in rats.

HBCDDs affected the immune system. Changes observed included decreased thymus weight and
increased lesions in the thymus, reduced splenocytes proliferation and increased spleen lesions,
decreased T-cell and increased B-cell populations, increased natural killer (NK) cells, and changes in
immunoglobulin responses.

Exposure of juvenile rats and mice to HBCDDs caused neurodevelopmental effects on behaviour, leading
 to changes in spontaneous behavior, spatial learning and memory as detected in adulthood.

140 HBCDDs are not genotoxic *in vitro* or *in vivo*. The slight induction of DNA strand breaks observed in 141 some *in vitro* tests is most likely due to oxidative stress.

HBCDDs were previously judged not to be carcinogenic in mice and no new carcinogenicity studies havebeen identified. The available evidence indicates that HBCDDs are not carcinogens.

144 A growing number of epidemiological publications were identified assessing the association between 145 exposure to HBCDDs and birth weight/length, neurodevelopment and thyroid dysfunction in children, 146 as well as subfertility, type 2 diabetes, thyroid hormone levels, severe endometriosis (including ovarian 147 endometrioma) and breast cancer metastasis in adults. None of the effects studied in longitudinal 148 studies and using internal exposure measures either reached statistical significance or were replicated 149 in a longer follow-up point in the same study. Considerable limitations exist pertaining to small sample 150 sizes, varying methodological quality, effect inconsistency and considerable heterogeneity in the 151 assessed populations, exposures, and endpoints. Adverse effects of HBCDDs related to 152 neurodevelopment have been assessed in two epidemiological studies; the low volume of prospective 153 data, the differing endpoint measures and the lack of replication render these data insufficient for use 154 in risk characterisation.

155 Regarding the mode of action, effects of HBCDDs on the liver of rodents appear to involve the 156 constitutive and rostane receptor (CAR) and the pregnane-X-receptor (PXR). Mitochondrial energy 157 production is reduced with downstream effects on reactive oxygen species (ROS) production, Na<sup>+</sup>/K<sup>+</sup>-158 ATPase and  $Ca^{2+}$ -ATPase activities and increased cytosolic [ $Ca^{2+}$ ] and [ $Zn^{2+}$ ]. HBCDDs stimulate 159 proliferation and migration of liver cell lines at picomolar to nanomolar concentrations and there is 160 evidence that these effects are related to activation of the estrogen receptor and the PI3K/Akt/mTOR 161 signalling pathway. Oral HBCDD exposure of mice fed a high fat diet aggravates metabolic dysfunction 162 through modifications of lipid and glucose homeostasis. HBCDD-induced lipid accumulation in liver and 163 adipose tissue appears to be associated with a suppression of the  $Wnt/\beta$ -catenin pathway resulting in 164 increased expression of *Pparg* and its adipogenic target genes. Mechanistic studies on the effects of 165 HBCDDs on the nervous system support specific effects on dopaminergic and glutamatergic neurons. 166 This appears to include inhibition of dopamine and glutamate reuptake following synaptic release, and deregulation of Ca<sup>2+</sup> and Zn<sup>2+</sup>. The brain could also be affected by HBCDDs through diminished 167 168 responsiveness to thyroid hormones. HBCDDs may affect the synthesis of sex steroid hormones by 169 interfering with cAMP-dependent cholesterol uptake and by causing dysregulation of enzymes involved 170 in sex steroid metabolism. HBCDDs may also alter the response of tissues to sex steroids. The 171 mechanism(s) involved in HBCDD-induced ROS production may be related to impairment of 172 mitochondrial energy production.

173 The evidence from the available human data was not sufficient to base the risk assessment on. Thus 174 the data from studies on experimental animals were used to identify a Reference Point for the human 175 health risk characterisation. The CONTAM Panel concluded that the neurodevelopmental effects on 176 behaviour can be considered the critical effect for the risk characterisation. The BMD modelling on the 177 neurobehavioural data showed BMD confidence intervals that were relatively wide and that the BMDLs 178 for horizontal locomotion and total activity were far below the lowest dose administered. Therefore, the 179 NOAEL/LOAEL approach was applied. Based on effects on spontaneous behaviour (horizontal 180 locomotion, rearing and total activity), the CONTAM Panel identified a lowest-observed-adverse-effect level (LOAEL) of 0.9 mg/kg bw (single dose) as the Reference Point for the risk assessment of HBCDDs. 181

Because the elimination kinetics for HBCDDs between mice and humans differ, the CONTAM Panel first calculated a body burden of 0.75 mg/kg bw at the LOAEL, considering an oral absorption of 83% in mice. The chronic intake that would lead to the same body burden in humans was calculated assuming an absorption in humans of 100% and the longest half-life identified in humans for HBCDDs of 219 days. This resulted in an estimated chronic human dietary intake of 2.35 µg/kg bw per day.

187 Due to limitations in the database on HBCDDs, the derivation of a health-based guidance value (HBGV) 188 was not considered appropriate. Instead, the margin of exposure (MOE) approach was applied to assess 189 possible health concerns. The CONTAM Panel considered that an MOE higher than 24 would indicate a 190 low health concern. This MOE would be sufficient to cover interspecies and intraspecies differences in 191 dynamics (a factor of 2.5 and 3.2, respectively), as well as extrapolation from a LOAEL to a no-observed-192 adverse-effect level (NOAEL) (a factor of 3). The Panel considered whether an additional factor should 193 be applied to allow for limitations in the database. It was noted that repeated dose reproductive toxicity 194 studies, that include a functional observational battery, showed only sporadic effects. Carcinogenicity 195 was studied only in mice. Based on the mode of action and genotoxicity findings, the CONTAM Panel 196 considered that carcinogenicity is unlikely to be a critical effect. Therefore, it was concluded that an 197 additional uncertainty factor for limitations in the database is not needed.

# 198 Occurrence and dietary exposure assessment for the European population

199 A total of 6,857 analytical results for HBCDDs in food fulfilled the quality criteria applied and from these, 200 6,352 were selected and used for the chronic exposure assessment. Contrary to the data used in the 201 previous Opinion on HBCDDs published in 2011, where most of the data were analysed with GC/MS and 202 reported Total HBCDDs, the current data have mostly been analysed by LC/MS and reported on the 203 specific stereoisomers  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCDD.

The high proportion of left-censored data, partially due to relatively high LOQs, resulted in a large difference between the lower bound (LB) and upper bound (UB) occurrence values for many food groups. The highest mean concentrations of HBCDDs (sum of  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCDD) were recorded for the food category 'Fish meat', with the highest mean concentrations in eel, being 2.33 and 2.35 µg/kg ww at LB and UB, respectively.

- 209 The mean dietary exposure estimates for HBCDDs ranged from 0.07 (minimum LB)/0.17 (minimum UB)
- to 0.79 (maximum LB)/1.52 (maximum UB) ng/kg bw per day across dietary surveys and age groups.
- 211 At the 95th percentile, dietary exposure estimates ranged from 0.23 (minimum LB)/0.45 (minimum UB)

- to 2.30 (maximum LB)/3.61 (maximum UB) ng/kg bw per day. The high proportion of left-censored
- data, partially due to relatively high LOQs, resulted in a 2- to 3-fold difference between the LB and UB
   exposure estimates.
- The most important contributors to the chronic dietary LB exposure to HBCDDs were 'Fish meat', 'Eggs, fresh', 'Livestock meat' and 'Poultry'.

An estimation of the exposure to HBCDDs of breasfed infants via consumption of human milk was also performed. The exposure scenario based on average human milk consumption and the reported UB range for HBCDD (sum of  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCDD) in pooled human milk samples collected in European countries between 2014 and 2016 as part of the World Health Organisation/United Nations Environment Programme (WHO/UNEP) field studies, would result in a median daily exposure of 14.3 ng/kg bw (range: 3.2–73.7 ng/kg bw). For infants with high human milk consumption the median daily exposure would

- 223 result in 21.5 ng/kg bw (range 4.8–110.6 ng/kg bw).
- Non-dietary exposure through intake of dust and dermal contact can substantially contribute, and in some cases even dominate the total human exposure to HBCDDs, especially for toddlers.

# 226 **Risk characterisation**

MOE values were calculated by comparison of the calculated chronic human dietary intake of 2.35 µg/kg bw per day, leading to the body burden at the LOAEL, with the estimated dietary exposure for the different population groups. The MOE values obtained ranged from 34,000 to 650. These MOEs are larger than 24 and the CONTAM Panel concluded that they do not raise a health concern.

The CONTAM Panel also compared the body burden of 0.75 mg/kg bw at the LOAEL with the body burdens estimated in adults based on levels in adipose tissue, blood and milk reported in the literature. The results support the conclusion that current dietary exposure to HBCDDs in European countries does not raise a health concern.

For breastfed infants, the lowest MOE values for high milk consumption are below the value of 24. The CONTAM Panel concluded that these MOEs may raise a health concern for some breastfed infants.

# 237 **Recommendations**

- In order to improve the risk assessment and reduce the uncertainties, the CONTAM Panel made thefollowing recommendations:
- Criteria for the analysis of HBCDD stereoisomers should be set.
- Surveillance of HBCDD stereoisomers in food, in particular in food groups for infants and small children, should continue in order to refine the exposure estimates.
- More data on occurrence of HBCDDs in human milk are needed to enable a more robust exposure assessment for breastfed infants.
- Improved information on toxicokinetics, e.g. half-life values, of HBCDD stereoisomers in humans is needed. More information is needed on the body burden in mothers and relation to transfer of HBCDDs to milk. This information should be used to develop a toxicokinetic model for HBCDDs.
- More information is needed on the transfer into ruminant meat and milk.
- Further toxicological studies should be conducted with individual HBCDD stereoisomers most relevant to human exposure.
- Studies on neurodevelopment (after repeated exposure), including investigations of the mechanisms and mode of action involved, are recommended.
- Studies on reproductive effects are recommended.
- Studies on possible diabetogenic and obesogenic effects are recommended.
- Longitudinal epidemiological studies of sufficient power and appropriate exposure and co-exposure
   assessment are needed.

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#### 1. Introduction 370

371 1.1. Background and Terms of Reference as provided by the European 372 Commission

#### Background 373

374 Brominated flame retardants (BFRs) are anthropogenic chemicals, which are added to a wide variety of 375 consumer/commercial products in order to improve their fire resistance. The major classes of BFRs are 376 brominated bisphenols, diphenyl ethers, cyclododecanes, phenols, byphenyl derivatives and the 377 emerging and novel BFRs.

378 Concern has been raised because of the occurrence of several chemical compouds from the group of 379 BFRs in the environment, including feed and food, and in humans. This has led to bans on the production and use of certain formulations of polybrominated diphenyl ethers (PBDEs). 380

Between September 2010 and September 2012, the Scientific Panel on Contaminants in the Food of 381 382 EFSA adopted six scientific Opinions on different classes of brominated flame retardants<sup>1</sup>. Because in 383 its Opinion EFSA highlighted several data gaps, hampering the consumer risk assessment for these 384 substances, by means of Commission Recommendation 2014/118/EU on the monitoring of traces of brominated flame retardants in food, Member States were recommended to collect in 2014 and 2015 385 386 occurrence data for specific substances in specific foodstuffs.

387 The newly available occurrence data would enable an updated consumer exposure assessment. 388 Furthermore, since the publication of the EFSA scientific Opinions between 2010 and 2012, new scientific 389 information has become available, therefore it would be necessary to verify whether an update of these 390 scientific Opinions would be appropriate, including an update of the consumer risk assessment.

#### **Terms of reference** 391

In accordance with Art. 29 (1) of Regulation (EC) No 178/2002, the European Commission asks the 392 European Food Safety Authority for an updated exposure assessment for the brominated flame 393 retardants, covered by Recommendation 2014/118/EU, taking into account the occurrence data in food, 394 submitted after the publication of the 2010–2012 EFSA scientific Opinions, and an updated consumer 395 risk assessment, taking into account newly available scientific information. 396

1.2. Interpretation of the Terms of Reference 397

398 Following the request from the EC, the CONTAM Panel will update its 2010–2012 risk assessments on 399 the different families of BFRs: hexabromocyclododecanes (HBCDDs), polybrominated diphenyl ethers 400 (PBDEs), tetrabromobisphenol A (TBBPA) and its derivatives, brominated phenols and their derivatives and novel and emerging BFRs (EFSA CONTAM Panel, 2011a,b,c, 2012a,b). 401

402 This first Opinion updates the risk assessment of HBCDDs in food previously performed by EFSA (EFSA 403

CONTAM Panel, 2011a). The assessment covers the transfer from feed to food of animal origin as a 404

Scientific Opinion on Hexabromocyclododecanes (HBCDDs) in Food. EFSA Journal 2011; 9(7):2296. [118 pp.].

relevant source of HBCDDs in food. The assessment focuses on 1,2,5,6,9,10-HBCDD and its major

<sup>&</sup>lt;sup>1</sup> EFSA Panel on Contaminants in the Food Chain (CONTAM); Scientific Opinion on Polybrominated Biphenyls (PBBs) in Food. EFSA Journal 2010; 8(10): 1789. [151 pp.].

Scientific Opinion on Polybrominated Diphenyl Ethers (PBDEs) in Food. EFSA Journal 2011; 9(5):2156. [274 pp.].

Scientific Opinion on Tetrabromobisphenol A (TBBPA) and its derivatives in food. EFSA Journal 2011; 9(12):2477, [61 pp.].

Scientific Opinion on Brominated Flame Retardants (BFRs) in Food: Brominated Phenols and their Derivatives. EFSA Journal 2012; 10(4):2634. [42 pp.].

Scientific Opinion on Emerging and Novel Brominated Flame Retardants (BFRs) in Food. EFSA Journal 2012; 10(10):2908. [125 pp.].

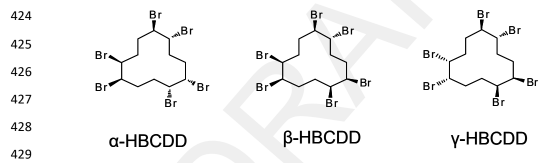
405 constituents, the three stereoisomers  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCDD. The CONTAM Panel noted that in a few 406 publications and dossiers HBCDDs are also mentioned with the substitution pattern 1,3,5,7,9,11. 407 Considering the manufacturing process for HBCDDs, the formation of a structure described by the name 408 1,3,5,7,9,11-hexabromocyclododecane is unlikely. No REACH registration has been requested for a 409 product with this substitution pattern. Therefore, 1,3,5,7,9,11-HBCDD is not further considered in this 410 assessment.

The similarities in chemical properties and effects seen in the previous EFSA assessments for the different BFR families warrant the consideration of a mixture approach. The CONTAM Panel will evaluate the appropriateness of applying a mixture approach in an additional Opinion once the risk assessment for the each BFR family has been updated. It will be based on the EFSA Guidance on harmonised methodologies for human health, animal health and ecological risk assessment of combined exposure to multiple chemicals (EFSA Scientific Committee, 2019).

417 **1.3.** Supporting information for the assessment

# 418 1.3.1. Physicochemical properties

Technical HBCDD is a white odourless solid compound, which is sometimes used in combination with other flame retardants. It is important to note that HBCDDs exist as stereoisomers of the cyclododecane molecule, which is substituted with six bromine atoms as shown in **Figure 1**. The physicochemical properties of HBCDDs were described in section 1.3 (Chemical characteristics) of the previous EFSA Opinion on HBCDDs (EFSA CONTAM Panel, 2011a) and are summarised in **Table 1**.



430 **Figure 1.** General structure of the three major HBCDD stereoisomers; α-HBCDD, β-HBCDD and γ-431 HBCDD (molecular mass = 641.7)

432 **Table 1.** Physico-chemical properties of HBCDDs (summarised from EFSA, 2011a)

Parameter	α-HBCDD	β-HBCDD	γ-HBCDD
Log Kow (ECB, 2008)	5.07	5.12	5.47
Log Kow (Goss et al., 2008) modelled by COSMOtherm	5.59	5.44	5.53
Water solubility <sup>(a)</sup>	48.8 ± 1.9 µg/L	14.7 ± 0.5 µg/L	2.1 ± 0.2 µg/l

433 (a): The full set of physicochemical characteristics is given in the ECB report (ECB, 2008).

434Technical products of HBCDDs consist predominantly of the three stereoisomers shown above and435contain about 9–13% α-HBCDD, <0.5–12% β-HBCDD and 72–90% γ-HBCDD (Peled et al., 1995). Other</td>436stereoisomers, δ- and ε-HBCDD may be present at low concentrations in the technical product (Law et437al., 2005; Arsenault et al., 2007), which may also contain 1–2% tetrabromocyclododecene (Peled et al.,4381995). Each of the stereoisomers of HBCDDs has two enantiomers so for the three main stereoisomers,439the six enantiomers are (+)-α-HBCDD, (+)-β-HBCDD, (-)-α-HBCDD, (-)-β-HBCDD, and (-440)-γ-HBCDD.

441 HBCDDs are relatively stable, but when they degrade the process primarily consists of the elimination 442 of hydrogen bromide leading to the formation of double bonds in the resulting molecule. The y-HBCDD 443 stereoisomer can be rearranged to form  $\alpha$ -HBCDD at temperatures of 190°C (Peled et al., 1995). Trans-444 1,2-dibromo-substituted compounds are known to undergo thermal rearrangements (Peled et al., 1995). 445 β-HBCDD is not affected by the thermal rearrangement process (Fång, 2007). Exposure to UV irradiation 446 may lead to breakdown of the parent compound and/or transformation from y- to  $\alpha$ -HBCDD (Harrad et 447 al., 2009), which has important consequences for sampling and analytical considerations (see Section 448 **1.3.4**). The  $\gamma$ - $\alpha$  transformation contributes to the fact that  $\alpha$ -HBCDD is the predominant of the three 449 main HBCDD stereoisomers in the environment.

Differences in solubility between stereoisomers may also have an impact on the changes that occur in ratio of stereoisomers in the environment and food chain. The  $\alpha$ -HBCDD stereoisomer predominates in food samples, but it is unclear whether this is a result of a biotransformation process, or whether selective absorption as a result of differences in solubility can also contribute to this change (Zegers et al., 2005; Law et al., 2006) (see **Section 1.3.3**).

# 455 **1.3.2. Production and industrial use**

The processes used to produce HBCDDs were described in section 4.1 of the previous EFSA Opinion (EFSA CONTAM Panel, 2011a). Some common trade names under which technical HBCDD was marketed, and some details of historic production volumes are also given in the same section in EFSA CONTAM Panel (2011a).

Prior to concerns about toxicity and environmental safety, HBCDDs were a high production volume chemical. In February 2011, HBCDDs were listed in Annex XIV of REACH regulations, and as such became subject to Authorisation. Their use has been permitted until the 'sunset date' of August 2015, after which only authorised applications were allowed in the EU<sup>2</sup>. In May 2013 the Stockholm Convention on Persistent Organic Pollutants (POPs) decided to include HBCDDs in Annex A of the Convention, which is a list of chemicals whose production and use should be eliminated <sup>3</sup>.

HBCDDs have been used mostly in expanded and extruded polystyrene foams (EPS and XPS) and as insulation material in the construction industry. Because it is so efficient, only low percentages are required to reach desired standards (0.7% in EPS and 2.5% in XPS). HBCDDs have also been used in upholstered furniture, vehicle textiles, car cushions and insulation blocks in commercial haulage vehicles, packaging materials and electrical and electronic equipment (Covaci and Malarvannan, 2016).

HBCDDs are not chemically bound to the polymers where found, and consequently are more likely toleach during production, use, disposal and recycling processes.

# 473 1.3.3. Environmental levels and fate

474 HBCDDs are widespread in the global environment. EFSA CONTAM Panel (2011a) described the main 475 sources of release into the environment and these were primarily from the manufacture and use of 476 insulation boards and manufacture and use of textiles. Due to the properties associated with 477 bioaccumulation and persistence, high levels can be found in top predators. Covaci et al. (2006) reported 478 high concentrations in marine mammals and birds of prey and Zacs et al. (2018a) reported high levels 479 in terrestrial wildlife. Reviews showed that the levels of HBCDDs in the environment were generally 480 increasing in all matrices studied and they seemed to correlate with the use of HBCDDs (UNEP, 2010). 481 Use of HBCDDs was expected to decrease since placement in the Stockholm Convention in 2013 (see 482 Section 1.3.6) but no conclusive evidence of decreasing levels in the environment has been shown in

<sup>&</sup>lt;sup>2</sup> https://echa.europa.eu/substance-information/-/substanceinfo/100.239.157

<sup>&</sup>lt;sup>3</sup> www.pops.int

- the scientific literature (Bao et al., 2015; Cao et al., 2018). There are indications from some studies that concentrations in environmental samples are decreasing, but this is not consistent between studies and is not a statistically robust conclusion. This may be because HBCDDs are still present in many items that are in use, and because of the time lag that is needed before decreasing temporal trends can be verified as a result of the restrictions that have been put in place.
- The paragraphs below, which do not claim to be a comprehensive review of the literature, give an overview of some aspects related to the environmental fate and levels of HBCDDs.

# 490 **1.3.3.1. Biodegradation / transformation**

- Some information on the transformation of HBCDDs in the environment was provided in the previous EFSA Opinion on HBCDDs (EFSA CONTAM Panel, 2011a). As described above in **Section 1.3.1**,  $\alpha$ -HBCDD is the most abundant stereoisomer, and this accounts for the predominance of  $\alpha$ -HBCDD in the environment, biota and in humans. The dominance of  $\alpha$ -HBCDD becomes greater over time and at higher trophic levels.
- 496 β- and γ-HBCDD can be bioisomerized into α-HBCDD by organohalide-respiring bacterium 497 *Dehalococcoides mccartyi* strain 195. β-HBCDD was found to transform to α-HBCDD at faster rate than 498 γ-HBCDD, and the enantiomer fractions of  $(\pm)\alpha$ -,  $(\pm)\beta$ - and  $(\pm)\gamma$ -HBCDD were fairly constant, indicating 499 a lack of enantioselective biotransformation of these diastereoisomers (Zhong et al., 2018).
- Half-lives for the degradation of HBCDDs in water, soil and sediment are shorter than half-lives in fish.
  This may be seen as counterintuitive especially when the magnitude of the differences is considered,
  e.g. 35 days in aerobic sediment and 250 days in a 1 kg fish. However, these data are supported by
  bioaccumulation measurements that demonstrate biomagnification in food webs and low
  biotransformation rates (ECB, 2008; Arnot et al., 2009).
- Thermal degradation of HBCDDs results in the formation of high-molecular-weight aromatic and polyaromatic brominated decomposition compounds (Dirtu et al., 2014). In the same way as polychlorinated dioxins and furans can be produced when organochlorine compounds are combusted, polybrominated dioxins and furans may be generated in low amounts when organobromine-containing products are combusted (D'Silva et al., 2004; Jogsten et al., 2010; Piskorska-Pliszczynska and Maszewski, 2014; IPEN, 2019). Even a very low level of these brominated dioxins could be significant due to the high toxicity.
- 512 It has been reported that pentabromocyclododecanols and isobutoxypentabromocyclododecanes can 513 be formed as environmental transformation products of HBCDDs (possibly formed via hydrolysis in 514 aquatic environments), and they may also be found in technical HBCDD where they are formed as by-515 products. As a result, they may be found in the environment along with parent HBCDDs (Heeb et al., 516 2012).
- 517 Degradation processes have been reported to be important for HBCDDs (Tomy et al., 2011). Whilst they 518 may be considered persistent in abiotic media, it is not necessarily so in biotic media. This difference in 519 degradation rate taking into account the biotransformation of HBCDDs in biological systems may 520 confound interpretation of temporal trend studies (Tomy et al., 2011).

# 521 **1.3.3.2.** Differences in HBCDD profiles and bioaccumulation

522 Previous sections have discussed the transformation of HBCDD stereoisomers from  $\gamma$ -HBCDD, which is 523 predominant in technical HBCDD, to  $\alpha$ -HBCDD, which is more dominant in the environment and biota, 524 including humans. The ratio of stereoisomers relates to the matrix and the time that has elapsed since

525 the contamination occurred.

526 HBCDDs have a strong potential to bioaccumulate and biomagnify. They are persistent in the 527 environment and have a potential for long-range environmental transport. The  $\alpha$ -HBCDD stereoisomer 528 predominates in food samples, but it is unclear whether this is a result of a biotransformation process, 529 or whether selective absorption can also contribute to this change (Zegers et al., 2005; Law et al., 530 2006). Differences in solubility between stereoisomers may also have an impact.

One study that suggested that transformation may take place in the plant, was a report on root uptake 531 532 of HBCDDs (Li et al., 2011). Seeds of cabbage and radish were grown in soil containing 1,000 µg 533 HBCDDs/kg, and the uptake of HBCDDs via the roots was determined. The results showed that after 8 534 weeks, cabbage can contain up to 65 µg/kg in root tissue and 30 µg/kg in shoot tissue. It was also 535 found that the root tissue contained higher levels of  $\gamma$ -HBCDD compared to  $\alpha$ - and  $\beta$ -HBCDD in contrast 536 to what is normally observed in most foods. Up to around 40, 18 and 12 µg/kg, respectively were found 537 in the root issues, whereas in the shoots the more typical pattern was found, i.e.  $\alpha$ -HBCDD was the 538 dominating isomer (18  $\mu$ g/kg) followed by  $\gamma$ - (8  $\mu$ g/kg) and  $\beta$ -HBCDD (3  $\mu$ g/kg).

539 In a study by Wu et al. (2012) concentrations resulting from uptake were greater in shoots than in roots 540 of maize (Wu et al., 2016a), and accumulation was found to be in the order  $\beta$ -HBCDD >  $\alpha$ -HBCDD > y-541 HBCDD in roots, and  $\beta$ -HBCDD >  $\gamma$ -HBCDD >  $\alpha$ -HBCDD in shoots. Diastereomer- and enantiomer-542 specific accumulation and biotransformation of HBCDDs in maize (Zea mays L.) was investigated by 543 Huang et al. (2016a). The  $(-)\alpha$ -,  $(-)\beta$ -, and  $(+)\gamma$ -HBCDD enantiomers accumulated to significantly 544 higher levels in maize compared with their corresponding enantiomers. Bioisomerisation from (+)/(-)-545 β- and y-HBCDDs to  $(-)\alpha$ -HBCDD was frequently observed, and (-)y-HBCDD was most easily 546 converted, with an efficiency of  $90.5 \pm 8.2\%$ .

# 547 **1.3.3.3. Occurrence in the environment**

# 548 Occurrence in soil

549 Most data for HBCDDs in soil originate from urban or industrial areas, and show that soil can contain 550 several hundred  $\mu$ g/kg dry weight (dw) (Covaci et al., 2009). In EFSA CONTAM Panel (2011a) the 551 background concentrations reported were < limit of detection (LOD)-6.6  $\mu$ g/kg dw, median 0.18  $\mu$ g/kg 552 dw.

553 Since the publication of the previous Opinion (EFSA CONTAM Panel, 2011a), Cao et al. (2018) reported 554 levels of HBCDD in soils from China.  $\Sigma$ HBCDDs (sum of  $\alpha$ -,  $\beta$ - and y-HBCDD) levels in the surface soils 555 ranged from 0.88 to 6,901 µg/kg dw with a mean value of 122.57 µg/kg dw. HBCDDs in soil close to a 556 manufacturing facility had the highest level (6,901 µg/kg dw). This value was lower than those reported 557 from HBCDD processing plants in Sweden and Belgium/Germany where concentrations of up to 23,200 558 µg/kg dw were reported (Covaci et al., 2006). It has been shown that there are significant differences 559 among different types of soils (Tang et al., 2014). A study was conducted using six types of soils 560 including samples taken from waste dumping sites, industrial areas, residential areas, traffic areas, 561 vegetable soils, and farmland soils in Ningbo (Zhejiang, China). High *SHBCDD* concentrations were 562 found in the waste dumping sites (60.74  $\mu$ g/kg dw) and the industrial area (37.9  $\mu$ g/kg dw), where 563 HBCDDs could originate from local sources (Tang et al., 2014). The mean concentrations of  $\Sigma$ HBCDDs 564 in traffic areas was 31.8 µg/kg dw, which showed higher concentrations than those in residential areas 565 (14.1  $\mu$ g/kg dw), soils used to grow vegetables (11.0  $\mu$ g/kg dw) and farmland soils (7.75  $\mu$ g/kg dw) 566 (Tang et al., 2014). However, the CONTAM Panel noted that it is difficult to compare these values since 567 the depth of the soil from where these samples were taken from is unknown.

# 568 Occurrence in sediment

569 Sediment is frequently monitored for HBCDDs since it is a recognised sink for hydrophobic chemicals. 570 Due to the relatively high organic content in sediments, and emissions to water, it may be expected to 571 find HBCDDs in sediments close to industry and other sources. It should be noted that aerobic 572 degradation is slower than anaerobic and the status of the sediment is an important factor.

Ganci et al. (2019) studied several BFRs, including HBCDDs, in tidal locations of the river Thames in London (UK). ΣHBCDDs (sum of  $\alpha$ -,  $\beta$ -, and  $\gamma$ -HBCDD) were detected in most samples (91% detection frequency) at an average concentration of 3.7 µg/kg dw (range: <LOD to 38 µg/kg dw). A study on estuarine and marine sediments around the UK reported a comparable range from <LOD to 47.2 µg/kg dw (Barber et al., 2014). Values for lake sediments in the UK ranged from 0.42 to 7.9 µg/kg dw (Ganci et al., 2019; Yang, 2014). The UK has a history of using more BFRs that other European countries, and levels are therefore likely to be higher than from other regions of Europe.

# 580 Occurrence in water

HBCDDs are lipophilic and there are little data in the literature for concentrations in water. Most reported data for water is focussed on suspended particles rather than the water itself. Concentrations of HBCDDs in water can range from a few ng/L to a few hundred  $\mu$ g/L. The major fraction of HBCDDs which reaches wastewater treatment plants is eliminated by adsorption to sludge, and only about 20% is released into the environment in effluents (ECB, 2008). HBCDD concentrations in effluents range from about 10 to 30  $\mu$ g/L (Rüdel et al., 2012; ECB, 2008).

587 Yang et al. (2019) measured HBCDDs in 107 water samples taken from nine English freshwater lakes 588 from a mix of urban, rural, and remote locations between August 2008 and February 2012. 589 Concentrations of HBCDDs declined over the sampling period with half-lives of 5.1 years. Concentrations 590 of HBCDDs ranged from 45 to 890 pg/L, and were generally equally distributed between the freely 591 dissolved and particulate phases in the lakewater Higher concentrations were associated with colder 592 periods at 5 out of 9 English lakes, and γ-HBCDD was consistently the most dominant isomer found. 593 The study found higher levels in the UK comprared with North America, reflecting higher per capita use 594 in the UK.

Ruan et al. (2019) investigated the transport of HBCDD enantiomers by microplastics in a wastewater treatment plant as a potential source of exposure. The release of HBCDDs carried by microplastics potentially reached 15.5 g per day, whereas the dissolved HBCDDs in the effluent may reach 0.067 g per day.

599 Concentrations of HBCDDs in invertebrates and fish can be an indicator of water quality. Data for 600 invertebrates are available for freshwater and marine species and include samples taken from near point 601 sources, from near regional sources, and from remote regions. Concentrations range over four orders 602 of magnitude for marine invertebrates as a result of the importance of proximity to sources (ECB, 2008). 603 Environmental monitoring data for fish includes samples from both urban/industrial and from remote 604 regions. As for invertebrates, the span of data is large covering 5–6 orders of magnitude (ECB, 2008).

# 605 Occurrence in air and dust

Levels of HBCDDs in air and dust were summarised in the previous Opinion (EFSA CONTAM Panel,2011a).

608 In air, HBCDDs are present mainly associated with particulates. Yu et al. (2008) showed concentrations 609 in Guangxhou (China) to range from 0.7 to 3.1  $pg/m^3$  in total with around 90±5% associated with 610 particulates. This study also demonstrated that an urban environment was an important factor with 611 respect to atmospheric concentrations. Roosens et al. (2010) reported the exposure of the Flemish population to BFRs from various sources and included HBCDDs in air. They did not measure 612 613 concentrations but used a figure of 37 pg/m<sup>3</sup> for outdoor air, 180 pg/m<sup>3</sup> for indoor air (homes) and 175 614 pg/m<sup>3</sup> for indoor air (offices) based on findings from 12 European studies reported between 2004 and 615 2008.

There are very little recent data available on HBCDDs in European air. Drage et al. (2016) found that the average  $\Sigma$ HBCDDs air concentration in an urban area in the UK was 100 pg/m<sup>3</sup>, but noted that concentrations were not correlated with distance from the city centre. The authors suggested that the high levels and lack of correlation was likely to be due to the sampling protocol which included locations that were in proximity to buildings possibly containing HBCDDs, which was heavily used as a flame

621 retardant in building insulation at the time of construction.

622 Newton et al. (2015) studied a varity of emerging flame retardants in Swedish indoor and outdoor air, 623 but HBCDDs were only detected in the Central Urban air sample where the concentration for ΣHBCDDs 624 was 0.58 pg/m<sup>3</sup>.

UK public microenvironments had the highest indoor air HBCDD concentrations of 900 pg/m<sup>3</sup> (median)
 compared to 2 pg/m<sup>3</sup> (median) in Swedish houses (Drage et al., 2016).

Levels in dust in Europe that have been reported since the previous Opinion (EFSA CONTAM Panel, 2011a) are summarised in **Appendix A**. For ΣHBCDD, levels reported vary significantly with some as low as <1 ng/g and others as high as around 150,000 µg/kg. Highest concentrations are generally for  $\alpha$ -HBCDD followed by  $\gamma$ -HBCDD and then  $\beta$ -HBCDD, although for some samples, particularly those with the highest concentrations,  $\gamma$ -HBCDD can be the predominant stereoisomer.

632 Concentrations in the dust from homes ranged between <3 (de Wit et al., 2012) to over 100,000  $\mu$ g/kg 633  $\Sigma$ HBCDD (Tay et al., 2017; Tao et al., 2016a), with the majority of data reported being below 1,000 634  $\mu$ g/kg. Levels in offices, day care facilities and educational establishments were in the same broad 635 range, although they were typically higher than in homes (**Appendix A**). Some of these studies may 636 have focussed on places where high exposure may be anticipated, e.g. aircraft and other vehicles, and 637 in some instances values over 1,000,000  $\mu$ g/kg were reported (Allen et al., 2013).

- 638 Malliari and Kalantzi (2017) performed a review on children's exposure to BFRs in indoor environments. 639 In general, large geographical variations were observed for all BFRs, including HBCDDs. The authors 640 estimated median worldwide dust levels in samples collected between 2006 and 2014, ranging from 615  $\mu$ g/kg in Egyptian houses to 340 ng/g in Swedish daycare centers, with most median concentrations 642 in the range of 100–200 ug/kg dust
- 642 in the range of 100–300  $\mu$ g/kg dust.

# 643 Occurrence in wildlife

644 Concentrations of HBCDDs have been reported for terrestrial and marine species of birds, and for marine 645 mammals from both industrialised and remote regions. Concentrations in polar bears from different 646 Arctic regions have also been measured (Muir et al., 2006; ECB, 2008).

647 HBCDD concentrations in wildlife that have been reported since the previous Opinion (EFSA CONTAM 648 Panel, 2011a) support the findings that were previously stated. Most data generated for wildlife concern marine species (e.g. Houde et al., 2017; Jin et al., 2015). There are some data on wild species that are 649 650 sometimes used for food. Zacs et al. (2018a) measured a range of contaminants in the meat and liver 651 of red deer (Cervus elaphus), wild boar (Sus scrofa), and moose (Alces alces) in a study on Latvian 652 wildlife. The wild boar samples contained the highest levels of HBCDDs, with the mean concentration 653 equal to 0.264 µg/kg ww in muscle tissues. In a different study (Zacs et al., 2018b), emerging BFRs 654 and dechlorane-related compounds were measured in European eels (Anguilla anguilla) from Latvian 655 lakes. HBCDDs were found in concentrations up to 6.58  $\mu$ g/kg lipid in the eels.

There are several studies that have been conducted in fish where the primary focus was as an environmental indicator rather than as a food source. Bustnes et al. (2012) investigated the latitudinal distribution of several POPs including HBCDDs, in livers of two species of marine fish, the pelagic saithe (*Pollachius virens*, n = 40) and the demersal cod (*Gadus morhua*, n = 40), along a south-north gradient (59°-70°N) on the Norwegian coastand. It was found that concentrations of especially heavy halogenated compounds (such as HBCDDs) were higher in fish from the southern-most region than inthose taken further north.

663 Bream was sampled annually from several European freshwater sites over a period of 4 years starting 664 from 2007 by Rüdel at al. (2012). Lowest levels of HBCDDs (11 µg/kg lipid weight, sum of three HBCDD 665 diastereomers) were detected in bream sampled in 2009 from Lake Belau, which is situated in a rural area of Northern Germany. Significant decreases of HBCDDs were detected in bream from the rivers 666 667 Rhone (France; -85%, to 205 µg/kg lipid weight) and Western Scheldt (the Netherlands; -60%, to 36 668 µg/kg lipid weight). High concentrations of HBCDDs (9,480–14,500 µg/kg lipid weight) were observed 669 in bream fom the River Tees (UK) with no clear changes in concentration associated with time. A 670 subsequent study (Rüdel et al., 2017) found that for most sites, a decrease in  $\Sigma$ HBCDDs was observed 671 in fish, e.g. in the Rhône and Western Scheldt by about 80 and 60%, respectively, with significantly 672 decreasing trends, p < 0.01. At the sampling site in the River Tees, which was impacted by a former 673 HBCDDs point source, concentrations in fish decreased only after a major flood event in 2013.

Munschy et al. (2013) investigated the temporal trend for HBCDDs in mussel (*Mytilus edulis*) samples harvested at the Seine estuary/France between 1981 and 2011. The concentration for  $\alpha$ -HBCDD ranged between 0.01 and 0.39 µg/kg wet weight (ww <sup>4</sup>). Although concentrations decreased between 2008 and 2010, over the entire study period (1981–2011) the contamination levels revealed a significant exponential upward trend with a doubling time of 8 years. However, since this time the impact on restrictions in use of HBCDDs might have changed this trend.

Fliedner et al. (2016) also studied bream using samples from the German environmental specimen bank. 680 Levels were measured and trends were examined in relation to the EU Water Framework Directive on 681 priority substances in freshwater fish. Samples were mostly compliant with the environmental quality 682 standard (EQS) for HBCDDs (167 µg/kg ww; see Section 1.3.6) but increasing trends were detected 683 for PBDEs and HBCDDs, particularly at one Saar site located downstream of the industries and 684 conurbation of Saarbrücken and Völklingen. The authors suggested that this finding related to new 685 sources of emissions of BFRs. The same authors analysed georeferenced samples of eelpout (Zoarces 686 viviparus) fillet from the German environmental specimen bank for HBCDDs (Fliedner et al., 2018). Two 687 sampling sites were located at German coastel areas of the Central North Sea and one site in the Bodden 688 National Park of Western Pomerania. Catching of the fish was between 2003 and 2017. The levels for 689 HBCDDs ranged between 0.03 and 12.2  $\mu$ g/kg ww with most samples between 0.11 and 1.5  $\mu$ g/kg ww. 690 Between 2003 and 2017 significant decreases were observed at two sampling sites (p = 0.03 and 691 < 0.01). 692

Eels were used as a 'bioindicator' species for the determination of the levels of organic contaminants within different Irish water bodies by McHugh et al. (2010). Levels of HBCDDs were between 1.2 and 15  $\mu$ g/kg (ww), corresponding to 7.4–166  $\mu$ g/kg (lipid weight), which is in line with concentrations reported elsewhere.

Eljarrat and Barceló (2018) reviewed the global scientific literature to compare PBDE and HBCDD levels
in river fish with the European EQSs. Most results reported were compliant, but two European studies
(Rüdel et al., 2012; Malarvannan et al., 2015) and one Chinese study (Tang et al., 2015) reported a
range of concentrations up to 751 µg/kg ww, exceeding the EQS for HBCDDs (167 µg/kg ww; Section
1.3.6) by a factor of about 5.

<sup>&</sup>lt;sup>4</sup> In this Opinion, the terms 'wet weight', 'whole weight' and 'fresh weight' are considered synonymous, unless otherwise stated, are abbreviated as ww.

702 Due to the fact that HBCDDs are persistent, lipohilic and bioaccumulate, they can be found in all animal 703 species including humans, in particular in lipid rich compartments, such as adipose tissue, milk, and liver

# 704 (see **Section 3.1.1.4**)

705 1.3.4. Sampling and methods of analysis

# 706 **1.3.4.1. Sampling**

707 The primary objective is to obtain a representative and homogeneous laboratory sample with no 708 secondary contamination. Therefore, basic rules for sampling of organic contaminants or pesticides 709 should be followed. Respective requirements are laid down, e.g. in Commission Regulation (EU) No 710 2017/644 <sup>5</sup> laying down methods of sampling and analysis for the control of levels of dioxins, dioxin-711 like PCBs and non-dioxin-like PCBs in certain foodstuffs. This Regulation contains, inter alia, a number 712 of provisions concerning methods of sampling depending on the size of the lot, packaging, transport, 713 storage, sealing, labelling, interpretation of analytical results and requirements for assessing the 714 compliance of a lot or sublot with the legislation. Commission Recommendation 2014/118/EU on the 715 monitoring of traces of brominated flame retardants in food lays stipulates that Member States should 716 follow the sampling procedures laid down in Annex II of the predecessor version of the Regulation (EU) 717 No 2017/644 in order to ensure that the samples are representative of the sampled lot.

# 718 **1.3.4.2. Methods of analysis**

719 Detailed information on methods of analysis for HBCDDs in food and parameters that influence the 720 validity of results are described in the previous EFSA Opinion on HBCDDs (EFSA CONTAM Panel, 2011a). 721 In summary, the analytical approaches for the HBCDDs extraction and clean-up generally follow the 722 methodologies for the analysis of persistent lipohilic compounds, such as extraction with organic 723 solvents, pressurised liquid extraction (PLE), and solid phase extraction (SPE). Clean-up is usually 724 performed by different column chromatographic techniques to remove co-extracted interfering matrix 725 constituents and to separate HBCDDs from potentially interfering compounds (Covaci et al., 2007; Xu 726 et al., 2013; Kuc and Grochowalski, 2014). Newer approaches make use of modified QuEChERS (quick, 727 easy cheap, effective rugged and safe) methods (Plassmann et al., 2015; Yuan et al., 2016; Li et al., 728 2017a).

The analytical determination of HBCDDs is performed either by gas chromatography (GC)/mass spectrometry (MS) or liquid chromatography (LC)/MS based methods. Both techniques enable the use of isotope-labelled internal standards, which allow for the correction of losses during extraction and clean-up. GC/MS analyses are, however, hampered by the fact that a separation of the stereoisomers is not possible by this approach. Thus, the result is reported as Total HBCDDs.

734 The separate specific analysis of the HBCDD stereoisomers as well as their enantiomers is performed 735 by LC/MS based methods. The analyses are often integrated into the procedures for the determination 736 of a number of other brominated flame retardants, such as TBBPA and bromophenols (Zhou et al., 737 2010; Butt et al., 2016; Bichon et al., 2018). Baek et al. (2017) reported that  $\delta$ - and  $\epsilon$ -HBCDD can co-738 elute with the predominant stereoisomers  $\alpha$ -,  $\beta$ - and y-HBCDD on columns with C18 stationary phases 739 which are chosen in most studies for HBCDD analysis. In order to avoid false-positive results, they 740 recommend a phenyl-hexyl ultra performance LC (UPLC) column on which ten HBCDD diastereomers 741 can be resolved. The advances in the enantioselective analysis of HBCDDs and other chiral flame 742 retardants with special emphasis on current status, limitations and future perspectives were reviewed 743 by Badea et al. (2016). The CONTAM Panel noted that this potential co-elution is of minor importance

<sup>&</sup>lt;sup>5</sup> Commission Regulation (EU) 2017/644 of 5 April 2017 laying down methods of sampling and analysis for the control of levels of dioxins, dioxin-like PCBs and non-dioxin-like PCBs in certain foodstuffs and repealing Regulation (EU) No 589/2014. OJ L 92, 6.4.2017, p. 9–34.

for the analysis of food, as δ- and  $\epsilon$ -HBCDD may be present at only low concentrations in the technical product.

LC-MS detection is mostly performed on triple-quadrupole instruments (MS/MS) using electrospray
ionisation (ESI) negative mode. Covaci et al. (2007) and Guerra et al. (2011) give extensive overviews
on respective methods used for the determination of HBCDDs in various matrices.

In the past few years new analytical instrumentation for the analysis of HBCDDs and other BFRs was introduced. These include *inter alia* the coupling of supercritical fluid chromatography to positive ion atmospheric pressure ionization mass spectrometry (Riddell et al., 2017), the application of GCatmospheric pressure chemical ionisation (APCI)-MS/MS (Sales et al., 2016), LC-atmospheric pressure photoionisation (APPI)-Orbitrap mass spectrometry (Zacs and Bartkevics, 2015), and ultra high performance LC-time-of-flight (TOF) high resolution MS (Zacs et al., 2014).

# 755 **1.3.4.3. Analytical quality assurance**

The analysis of HBCDDs in food is complex and involves several critical steps. As mentioned earlier, exposure to high temperature and to UV irradiation should be avoided as this may lead to breakdown of the parent compound and/or transformation from  $\gamma$ - to  $\alpha$ -HBCDD. It is therefore recommended to perform all analytical steps in amber glassware or in glassware covered with aluminium foil.

A prerequisite for laboratories to demonstrate that their applied methods of analysis are fit for purpose is the successful participation in proficiency tests or interlaboratory studies. Information on interlaboratory studies perfomed in biota and food samples before 2010 are summarised in the EFSA Opinion on HBCDDs of 2011 (EFSA CONTAM Panel, 2011a). The European Reference Laboratory (EURL) for Halogenated Persistent Organic Pollutants (POPs) in Feed and Food as well as the Norwegian Institute of Public Health (NIPH) regularly offer proficiency tests for analysing various persistent organic pollutants, including HBCDDs in feed and food.

767 The recent proficiency test which included the analysis of HBCDDs were organized by the EURL for 768 National Reference Laboratories (NRLs) and official laboratories, and concerned cod liver and fish liver 769 oil in 2014, bovine liver in 2017, soy bean meal and beef in 2018, and grass in 2019. In the 2014 EURL 770 proficiency test on cod liver and fish oil, assigned values of each 1.1 ng/g ww for  $\alpha$ -HBCDD were 771 estimated. The share of results with a z-score  $^{6} \leq 2$  was 80% for cod liver and 90% for fish liver oil. In 772 the EURL grass PT in 2019 the median concentrations for  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCDD each were around 0.1 773  $\mu q/kq$  (88% dry matter). About 90% of the submitted results were within ±40% of the median value 774 (Schächtele A, personal communication, 2019).

775 The interlaboratory comparison studies offered and organized by the NIPH are open to all laboratories 776 whether official or private. Laboratories were asked in each case to provide results on HBCDDs in 777 addition to data on dioxins and PCBs. Besides a standard solution submitted by the organizers in each 778 round, the matrices involved were reindeer meat, halibut filet and cod liver oil in 2012, poultry meat, 779 crab meat and eggs in 2013, pork, herring and cow's milk in 2014, beef, salmon and cheese in 2015, 780 sheep liver, salmon and fish oil in 2016, sheep meat, cod liver and herring in 2017, and reindeer meat, 781 salmon and fish oil in 2018. Generally, 5–15 laboratories reported one or more of the three isomers in 782 the various food samples. The organizers state that due to the low number of laboratories, no consensus 783 values could be derived and thus the results must be regarded as indicative values.

784In the 2017 NIPH interlaboratory comparison study, the median concentrations (all values) for the sum785of  $\alpha$ -,  $\beta$ - and γ-HBCDD were 10 pg/g ww for the sheep meat sample, 2,950 pg/g ww for cod liver, and786376 pg/g ww for the herring sample. The evaluation of the results indicate that the relative standard

<sup>&</sup>lt;sup>6</sup> A z-score is a numerical measurement used in statistics of a value's relationship to the mean (average) of a group of values, measured in terms of standard deviations from the mean.

deviations substantially increase with decreasing concentrations, being 14%, 33% and 227% for the sum of  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCDD in the cod liver, herring and sheep meat samples.

A similar outcome was reported for the samples analysed within the 2018 NIPH interlaboratory comparison study. The median concentrations (all values) for the sum of  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCDD were 11 pg/g ww for the reindeer meat sample, 378 pg/g ww for the salmon and 1,519 pg/g ww for the fish oil sample. The respective relative standard deviations were 19% for salmon, 45% for fish oil, and 134% for the reindeer sample.

All results of the various interlaboratory comparison studies are published on the home page of the NIPH <sup>7</sup>.

796 **1.3.5. Previous risk assessments** 

797 In 2011, the EFSA CONTAM Panel published its risk assessment on HBCDDs in food (EFSA CONTAM 798 Panel, 2011a). The Panel identified as the critical endpoint neurodevelopmental effects on behaviour 799 and derived a BMDL<sub>10</sub> of 0.93 mg/kg bw, based on the study by Eriksson et al. (2006) in mice, using a 800 single oral administration on postnatal day 10 (PND10). Because the elimination kinetics for HBCDDs 801 between experimental animals and humans differ, the CONTAM Panel converted this BMDL10 into an 802 estimated chronic human dietary intake associated with the body burden at the BMDL<sub>10</sub> as the basis for 803 the risk assessment. For this, the Panel first estimated the body burden at the BMDL<sub>10</sub> (0.79 mg/kg bw, 804 based on 85% oral absorption in mice). Second, the chronic human dietary intake associated with the 805 body burden at the BMDL<sub>10</sub> was calculated, assuming the human absorption of HBCDDs to be 100%, in 806 the absence of robust information on the actual absorption, and using the 'worst-case' longest human 807 half-life identified of 219 days. This resulted in an estimated human dietary intake associated with the 808 body burden at the BMDL<sub>10</sub> of 0.003 mg/kg by per day. Due to the limitations and uncertainties in the 809 database, the CONTAM Panel did not find it appropriate to establish a health-based guidance value 810 (HBGV), and instead used a margin of exposure (MOE) approach for the risk characterisation. The Panel 811 considered that an MOE larger than 8 would imply no health concern, as it would cover interspecies 812 differences on toxicodynamics for the effect observed (uncertainty factor of 2.5) and intraspecies 813 differences for toxicokinetics due to the uncertainty in the elimination half-life in humans (factor of 3.2). The Panel considered that since the MOE approach was based on a body burden comparison between 814 815 animals and humans, the potential kinetic differences between animals and humans had been accounted 816 for. And that by focussing on the body burden associated with a  $BMDL_{10}$  for neurobehavioural effects in 817 mice induced during a relevant period for brain development, and applying this body burden to the 818 entire life span in humans, individual differences in susceptibility had been covered. The chronic human 819 dietary intake of 0.003 mg/kg bw per day was compared with the estimate of the dietary exposure for 820 different age groups based on HBCDD levels in food collected in seven European countries covering the 821 period from 2000 to 2010. The CONTAM Panel concluded that the dietary exposure as estimated at the 822 time of the assessment did not raise a health concern based on the MOEs obtained.

Since then, several bodies have performed risk assessments for HBCDDs. The details of these assessments are reported in **Table 2**.

Some of these assessments used the chronic human dietary intake associated with the body burden at the BMDL<sub>10</sub> for neurodevelopmental effects calculated by the CONTAM Panel in its 2011 assessment, without reviewing the new studies since its publication (Rivière et al., 2014; Petersen et al., 2013). Others did review the new toxicological studies available since the EFSA Opinion but concluded that the new studies did not offer an alternative approach to the Reference Point identified by EFSA and its corresponding chronic human dietary intake (COT, 2015; Anses, 2017).

<sup>&</sup>lt;sup>7</sup> https://www.fhi.no/en/sys/search-result/?term=interlaboratory%20comparison

831 Other bodies identified NOAELs/LOAELs from different studies for their risk assessments. Environment 832 Canada/Health Canada (2011) identified a no-observed-adverse-effect level (NOAEL) of 10 mg/kg bw 833 per day from a two-generation reproductive toxicity study (Ema et al., 2008) to assess the risk for the 834 adult population, while for infants and children identified a lowest-observed-adverse-effect level (LOAEL) 835 of 0.9 mg/kg bw per day based on the study by Eriksson et al. (2006). The National Industrial Chemicals 836 Notification and Assessment Scheme (NICNAS, 2012) identified a NOAEL of 10.2 mg/kg bw per day 837 based on the Ema et al. (2008) study, that was used for both adults and children risk characterisation. None of these bodies established a HBGV but calculated the MOE, i.e. the margin between the BMDL<sub>10</sub> 838 839 or NOAEL/LOAELs identified and the estimated exposure in the corresponding country. All these risk 840 assessments concluded that the calculated MOEs indicate that the respective exposure levels are of no 841 health concern.

The Japanese National Institute of Technology and Evaluation (NITE, 2019) identified a NOAEL of 10 mg/kg bw per day based on the two-generation reproduction toxicity study (Ema et al., 2008), and derived hazard assessment values <sup>8</sup> for general toxicity (0.050 mg/kg bw per day, based on increased absolute and relative liver weight) and for reproductive and developmental toxicity (0.10 mg/kg bw per day, based on a reduction of the number of primordial follicles in the ovary of F1 females).

In 2014, the US-EPA's IRIS (Integrated Risk Information System) released a draft preliminary assessment on HBCDDs prior to the development of the draft assessment (US-EPA, 2014). It included, among others, the draft literature searches and tables summarising the critical available information for HBCDDs, to obtain input from the public prior to developing the draft IRIS assessment. However, in 2019 US-EPA has prioritised its IRIS assessments, and the assessment for HBCDDs among others, have been discontinued <sup>9</sup>.

<sup>&</sup>lt;sup>8</sup> According to the authors (NITE, 2019), the hazard assessment values were derived by dividing the NOAEL by uncertainty factors: the hazard assessment value of 0.050 mg/kg bw per day for general toxicity was derived by dividing the NOAEL by an uncertainty factor of of 200 (10 for animal-to-human extrapolation, 10 for human variability, and 2 in consideration of the use of NOAEL as well as the test period). The hazard assessment value of 0.10 mg/kg bw per day for reproductive and developmental toxicity (reduction of the number of primordial follicles) was derived by dividing the NOAEL by an uncertainty factor of 100 (10 for animalto-human extrapolation and 10 for human variability).

<sup>&</sup>lt;sup>9</sup> IRIS Programme Outlook (April 2019). Available at: <u>https://www.epa.gov/iris/iris-program-outlook</u>

**Table 2.** (Risk) Assessments on HBCDDs since the publication of the EFSA CONTAM Panel (2011a) Opinion, including details about the latter for comparison

Reference	Country	Compounds	NOAEL/LOAEL/BMD	HBGV / MOE	Dietary exposure (ng/kg bw per day)	Conclusion
EFSA CONTAM Panel (2011a)	Europe	Sum of α-, β- and γ-HBCDD	<ul> <li>BMDL<sub>10</sub> = 0.93 mg/kg bw (based on neurodevelopmental effects, single dose at PND10, Eriksson et al., 2006)</li> <li>Associated chronic intake = 0.003 mg/kg bw per day (related to the body burden at the BMDL<sub>10</sub>)</li> </ul>	MOE An MOE larger than 8 was considered to imply no health concern, based on UFs of 2.5 for interspecies differences on toxicodynamics and 3.2 for intraspecies differences for toxicokinetics	Mean exposure <sup>(a)</sup> : Children: 0.15–1.85 Adults: 0.09–0.99 Very Elderly: 0.06–0.54 P95 exposure <sup>(a)</sup> : Children: 0.80–4.46 Adults:0.39–2.07 Very Elderly: 0.27–1.26 Specific population groups: High consumers of fish: 2.76 <sup>(b)</sup> Consumers of fish liver (once a week): 1.94 <sup>(b)</sup>	"The CONTAM Panel concluded that current dietary exposure to HBCDDs in the European Union does not raise a health concern"
Rivière et al. (2014)	France	Sum of $\alpha$ -, $\beta$ - and $\gamma$ -HBCDD	Did not review new literature on toxicity. Used EFSA CONTAM Panel (2011a) chronic dietary intake associated with the body burden at the BMDL <sub>10</sub> (see above)	MOE	Total Diet Study Adults: 0.211 Children: 0.320	"The exposure levels are deemed to be of no public health concern (i.e. MOEs higher than 1000 for children and adults for the 95th percentiles)"
Petersen et al. (2013)	Denmark	Sum of α-, β- and γ-HBCDD	Did not review new literature on toxicity. Used EFSA CONTAM Panel (2011a) chronic dietary intake associated with the body burden at the BMDL <sub>10</sub> (see above)	MOE	Fish Adults mean exposure = 0.19 95th percentile = 0.75 Children (4–14 years old): Mean exposure = 0.23 95th percentile = 1.28	"The estimated MOEs are of no food safety concern"

COT (2015)	UK	Sum of α-, β- and γ-HBCDD	Reviewed new toxicity data since EFSA CONTAM Panel (2011a): "Overall, the new data did not demonstrate a mode of action for HBCDDs or provide an improved basis for extrapolation of experimental animal data to the human situation. Thus, they did not offer an alternative approach to the reference point identified by EFSA from the study by Eriksson et al. (2006), which remained the best available starting point for risk assessment" Reference Point = 3 µg/kg bw per day BMDL <sub>10</sub> body burden = 0.79 mg/kg bw (based on neurodevelopmental effects, Eriksson et al., 2006)	MOE	Breast milk, food for infants Breastfed infants (median): Average consumers (800 mL/day) 0–4 months: 0.018 >4–6 months: 0.014 High consumers: 0–4 months: 0.027 >4–6 months: 0.021 Infants: 4–6 months: 1.39 6–9 months: 1.62 9–12 months: 1.74	"Overall the analysis indicated that estimated exposures via breast milk and food are not a cause for concern, but that high levels found in some samples of domestic dust are"
Anses (2017)	France	Sum of $\alpha$ -, $\beta$ - and $\gamma$ -HBCDD	Reviewed new toxicity data since EFSA CONTAM Panel (2011a). Used EFSA CONTAM Panel (2011a) chronic dietary intake associated with the body burden at the BMDL <sub>10</sub> (see above).	MOE	Mean UB: Infants (1–4 months): 8.27 Infants (1–3 years): 0.505 Children (3–17 years): 0.320 Adults: 0.211 P90 UB: Infants (1–4 months): 43.2 Infants (1–3 years): 0.880 P95 UB: Adults: 0.448 Children (3–17 years): 0.734	"Based on current knowledge and the available data, exposure via food of the children population to the sum of HBCDD is considered tolerable"

Environment Canada/Health Canada (2011)	Canada	HBCDDs	To assess the risk for the adult population: NOAEL = 10 mg/kg bw per day (based on a two- generation reproductive toxicity study, Ema et al., 2008). To assess the risk for infants and children: LOAEL = 0.9 mg/kg bw per day (based on altered behaviour in mice, Eriksson et al., 2006).	MOE	UB exposure = 0.042 μg/kg bw per day (including food, ambient air and dust) Beastfed infants = 0.089 μg/kg bw per day	"Comparison with the NOAEL of 10 mg/kg bw per day resulted in a margin of exposure of 240,000. For breastfed infants, the margin was 10,000. These margins were considered by Environment Canada to be protective for breasfed children and adequate to address uncertainties in the exposure and health effects database."
NICNAS (2012)	Australia	HBCDDs	NOAEL = 10.2 mg/kg bw per day (based on a two-generation reproductive toxicity study, Ema et al., 2008)	MOE	Food consumption Typical exposure: Toddlers = 24 Children = 6 Adults =5.9 Reasonable worst-case: Toddlers = 50 Children = 12 Adults =12	``low risk from exposure through food'
NITE (2019)	Japan	HBCDDs	NOAEL = 10 mg/kg bw per day (based on a two-generation reproduction toxicity study, Ema et al., 2008)	Hazard assessment value <sup>(f)</sup> for general toxicity (increase absolute and relative liver weight) = 0.050 mg/kg bw per day (c) Hazard assessment value for reproductive and developmental toxicity (reduction of the number of	_ (e)	-

primordial follicles in the ovary of F1 females) = 0.10mg/kg bw per day (d)

854 BMDL<sub>10</sub>: benchmark dose lower confidence limit for a benchmark response of 10%. HBGV: health-based guidance value. LOAEL: lowest-observed adverse effect level. MOE: margin of exposure. TDS: 855 total diet study. NOAEL: no-observed adverse effect level. PND: postnatal day

856 (a): Mean exposure across dietary sruveys in European countries.

(b): Maximum UB across European surveys.

(c): Was derived by dividing the NOAEL by an UF of 200 (10 for animal-to-human extrapolation, 10 for human variability, and 2 in consideration of the use of NOAEL as well as the test period (NITE, 2019)

(d): Was derived by dividing the NOAEL by an uncertainty factor of 100 (10 for animal-to-human extrapolation and 10 for human variability) (NITE, 2019).

(e): No dietary exposure was estimated.

(f): According to the authors (NITE, 2019), the hazard assessment values were derived by dividing the NOAEL by an uncertainty factor of 200 (10 for animal-to-human extrapolation, 10 for human variability, and 2 in consideration of the use of NOAEL as well as the test period).

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#### 865 **1.3.6.** Legislation

In this Opinion, where reference is made to European legislation the reference should be understood as relating to the most recent amendment at time of publication of this Opinion, unless otherwise stated.

In order to protect public health, Article 2 of Council Regulation (EEC) No 315/93 <sup>10</sup> of 8 February 1993 laying down Community procedures for contaminants in food stipulates that, where necessary, maximum tolerances for specific contaminants shall be established. A number of maximum levels (MLs) are currently laid down in Commission Regulation (EC) No. 1881/2006 <sup>11</sup>. HBCDDs are not regulated so far under this Regulation or under any other specific European Union (EU) regulation for food.

Council Directive 2002/32/EC <sup>12</sup> regulates undesirable substances in animal feed. HBCDDs are, so far,
 not regulated under this Directive or any other specific EU regulation for feed.

In 2013 and following the recommendation from the Persistent Organic Pollutants Review Committee (POPRC), HBCDDs <sup>13</sup> were listed in Annex A (Elimination) of the Stockholm Convention, with specific exemptions for use or production <sup>14</sup>. For chemicals listed in Annex A, countries must take measures to eliminate the production and use of it.

HBCDDs are listed in Annex I, Part A of Regulation (EU) 2019/1021 of the European Parliament and of
the Council on persistent organic pollutants (POPs) <sup>15</sup>. The objective of this Regulation is to protect
human health and the environment by prohibiting, phasing out as soon as possible, or restricting the
manufacturing, placing on the market and use of POPs.

HBCDDs and all major diastereoisomers identified ( $\alpha$ -,  $\beta$ - and  $\gamma$ -) are included in Annex XIV of Regulation (EC) No 1907/2006 (REACH). According to Art. 57 (d), they are identified as PBT (persistent, bioaccumulative and toxic). The latest application date and sunset date were set as 21 February 2014 and 21 August 2015, respectively, and it is in the candidate list of substances of very high concern (SVHC) for Authorisation<sup>16</sup>.

According to Annex VI of Regulation (EC) No 1272/2008 (Classification, Labelling and Packaging (CLP) Regulation), HBCDDs and 1,2,5,6,9,10-HBCDD are classified (harmonised classification) Repr. 2 H 361 (Suspected of damaging fertility or the unborn child) and Lact. H 362 (May cause harm to breastfed children). For the stereoisomers, there are notified classifications (see **Table 3**) <sup>17</sup>.

Compound	EC/List no	CAS no	Hazard Classification
HBCDD	247-148-4	25637-99-4	The harmonised classifications are Repr. 2 H 361 (Suspected of damaging fertility or the unborn child) and Lact. H 362 (May cause harm to breastfed children).
$\alpha$ -HBCDD	603-801-9	134237-50-6	The notified classifications are Repr. 2 H 361 (Suspected of damaging fertility or the unborn child) and Lact. H 362 (May cause harm to breastfed children).

892 Table 3. Harmonised classification of HBCDDs (source: ECHA)

<sup>10</sup> Council Regulation (EEC) No 315/93 of 8 February 1993 laying down Community procedures for contaminants in food. OJ L 37, 13.2.1993, p. 1

<sup>11</sup> Commission Regulation (EC) No 1881/2006 of 19 December 2006 setting maximum levels for certain contaminants in foodstuffs. OJ L 364/5, 20.12.2006, p. 5-24.

<sup>14</sup> http://chm.pops.int/Implementation/Alternatives/AlternativestoPOPs/ChemicalslistedinAnnexA/tabid/5837/Default.aspx

<sup>15</sup> Regulation (EU) 2019/1021 of 20 June 2019 on persistent organic pollutans. OJ L 169, p.45.

<sup>16</sup> <u>https://echa.europa.eu/documents/10162/471aceac-4e5e-4c53-a4b2-23159a290893</u>

<sup>&</sup>lt;sup>12</sup> Directive 2002/32/EC of the European Parliament and of the Council of 7 May 2002 on undesirable substances in food. OJ L 140, 30.2.2002, p. 10

<sup>&</sup>lt;sup>13</sup> According to the Stockholm Convention, 'Hexabromocyclododecane' means hexabromocyclododecane (CAS No: 25637-99-4), 1,2,5,6,9,10-hexabromocyclododecane (CAS No: 3194- 55-6) and its main diastereoisomers: alpha- hexabromocyclododecane (CAS No: 134237-50-6); beta-hexabromocyclododecane (CAS No: 134237-51- 7); and gamma-hexabromocyclododecane (CAS No: 134237-52-8).

<sup>&</sup>lt;sup>17</sup> https://echa.europa.eu/information-on-chemicals/cl-inventory-database/-/discli/details/104361

β-HBCDD	603-802-4	134237-51-7	The notified classifications are Skin Irrit. 2 H 315 (Causes skin irritation), Eye Irrit. 2 H 319 (Causes serious eye irritation), and STOT SE 3 H 335 (May cause respiratory irritation).
Y-HBCDD	603-804-5	134237-52-8	The notified classifications are Repr. 2 H 361 (Suspected of damaging fertility or the unborn child) and Lact. H 362 (May cause harm to breastfed children).
1,2,5,6,9, 10-HBCDD	221-695-9	3194-55-6	The harmonised classifications are Repr. 2 H 361 (Suspected of damaging fertility or the unborn child) and Lact. H 362 (May cause harm to breastfed children).

HBCDDs are identified as priority hazardous substances under Directive 2013/39/EU of the European 893 Parliament and of the Council amending Directives 2000/60/EC and 2008/105/EC as regards priority 894 substances in the field of water policy <sup>18</sup>. Environmental guality standards (EOS, e.g. annual average 895 and maximum allowable concentrations) are laid down in surface water and biota for HBCDDs <sup>19</sup>. These 896 897 1,3,5,7,9,11-hexabromocyclododecane 20 (CAS 25637-99-4), refer to 1,2,5,6,9,10hexabromocyclododecane (CAS 3194-55-6),  $\alpha$ -hexabromocyclododecane (CAS 134237-50-6),  $\beta$ -898 899 hexabromocyclododecane (CAS 134237-51-7) and y-hexabromocyclododecane (CAS 134237-52-8). In 900 the Annex of this Directive maximum allowable concentrations (MAC-EQS) for inland surface waters 901 (rivers, lakes and related artificial or heavily modified water bodies), and other surface waters of 0.5 902  $\mu$ g/L and 0.05  $\mu$ g/L, respectively are set. For fish, an EQS of 167 ng/g is established.

In Commission Recommendation 2014/118/EU<sup>21</sup>, the EU Commission recommended that Member States 903 904 should perform monitoring on the presence of brominated flame retardants in food following the 905 recommendations of the previous EFSA risk assessment on HBCDDs and other BFRs (EFSA, 2011a,b, 906 2012a,b,c). Besides various other brominated flame retardants, the Recommendation also includes the 907  $\alpha$ -,  $\beta$ -, and  $\gamma$ -stereoisomers of 1,2,5,6,9,10-HBCDD. The aim of the monitoring is to include a wide 908 variety of individual foodstuffs reflecting consumption habits in order to give an accurate estimation of 909 exposure. Regarding HBCDDs, it is recommended to analyse fish and other seafood, meat and meat 910 products, milk and dairy products, eggs and egg products, as well as infant and follow-up formula. The 911 analytical methods should allow the stereoisomeric determination of the above mentioned three HBCDDs 912 with limits of quantification of 0.01 ng/g ww or lower.

# 913 **2. Data and Methodologies**

The current updates of the EFSA risk assessments on BFRs in food, including the current one on HBCDDs, were developed applying a structured methodological approach, which implied developing a priori the protocol or strategy of the full risk assessments and performing each step of the risk assessment in line with the strategy and documenting the process. The protocol in **Annex A** to this Opinion contains the method that was proposed for all the steps of the risk assessment process, including any subsequent refinements/changes made.

The CONTAM Panel used its previous risk assessment on HBCDDs in food (EFSA CONTAM Panel, 2011a)as a starting point for drafting the current Opinion.

<sup>&</sup>lt;sup>18</sup> Directive 2013/39/EU of the European Parliament and of the Council of 12 August 2013 amending Directives 2000/60/EC and 2008/105/EC as regards priority substances in the field of water policy Text with EEA relevance OJ L 226, 24.8.2013, p. 1–17-

<sup>&</sup>lt;sup>19</sup> Referring to 1,3,5,7,9,11-Hexabromocyclododecane (CAS 25637-99-4), 1,2,5,6,9,10- Hexabromocyclododecane (CAS 3194-55-6), α-Hexabromocyclododecane (CAS 134237-50-6), β-Hexabromocyclododecane (CAS 134237-51-7) and γ-Hexabromocyclododecane (CAS 134237-52-8)

<sup>&</sup>lt;sup>20</sup> Considering the manufacturing process for HBCDDs, the formation of a structure described by the name 1,3,5,7,9,11hexabromocyclododecane is unlikely. No REACH registration has been requested for a product with this substitution pattern.
<sup>21</sup> Commission Recommendation of 3 March 2014 on the monitoring of traces of brominated flame retardants in food. OJL 65, 5.3.2014, p. 39



# 922 **2.1.** Supporting information for the assessment

923 Information on physicochemical properties, production and industrial use, environmental fate and levels, 924 analytical methods, previous assessment and legislation was gathered from the previous EFSA Opinion 925 on HBCDDs (EFSA CONTAM Panel, 2011a), assessments by international bodies (by checking their 926 original websites of the relevant organisations), and from current EU legislation. Literature searches 927 were conducted to identify new information in reviews and peer-reviewed publications. Details about 928 the literature searches are given in **Appendix B**. The information was summarised in a narrative way 929 based on expert knowledge and judgement.

# 930 **2.2.** Hazard identification and characterisation

Information relevant for the sections under hazard identification and characterisation was identified by 931 932 a literature search to gather review studies and primary research studies. Details about the literature 933 search are given in Appendix B. The information retrieved was screened and evaluated by relevant 934 domain experts from the EFSA CONTAM Working Group on BFRs in food and used for the present 935 assessment. The selection of the scientific papers for inclusion or exclusion was based on consideration 936 of the extent to which the study was relevant to the assessment or on general study quality 937 considerations (e.g. sufficient details on the methodology, performance and outcome of the study, on 938 dosing, substance studied and route of administration and on statistical description of the results), 939 irrespective of the results. Limitations in the information used are documented in this Scientific Opinion.

Benchmark dose (BMD) analysis was carried out according to the latest EFSA guidance (EFSA Scientific
 Committee, 2017) and using the EFSA BMD modelling application.

# 942 2.3. Occurrence data submitted to EFSA

943 The general steps followed for the acquisition of the food occurrence and consumption data for the 944 exposure assessment of BFRs in food are documented in **Annex A**. Specific details on the occurrence 945 data on HBCDDs in food submitted to EFSA are described below.

# 946 Data collection and validation

Following an EC mandate to EFSA, a call for annual collection of chemical contaminant occurrence data in food and feed, including HBCDDs, was issued by the former EFSA Dietary and Chemical Monitoring Unit (now Evidence Management Unit<sup>22</sup>) in December 2010<sup>23</sup> with a closing date of 1 October of each year. European national authorities and similar bodies, research institutions, academia, food business operators and other stakeholders were invited to submit analytical data on HBCDDs in food.

The data submission to EFSA followed the requirements of the EFSA Guidance on Standard Sample Description (SSD) for Food and Feed (EFSA, 2010a). Occurrence data were managed following the EFSA standard operational procedures (SOPs) on 'Data collection and validation' and on 'Data analysis of food consumption and occurrence data'.

Data on HBCDDs in food available in the EFSA database from 2010 until the end of December 2018 were used for the present assessment.

# 958 Data analysis

Following EFSA's SOP on 'Data analysis of food consumption and occurrence data' to guarantee an appropriate quality of the data used in the exposure assessment, the initial dataset was carefully

<sup>&</sup>lt;sup>22</sup> From 1 January 2014 onwards, Evidence Management Unit (DATA).

<sup>&</sup>lt;sup>23</sup> http://www.efsa.europa.eu/en/consultations/call/190410



961 evaluated by applying several data cleaning and validation steps. Special attention was paid to the
962 identification of duplicates and to the accuracy of different parameters, such as 'Sampling strategy',
963 'Analytical methods', 'Result express' (expression of results, e.g. fat weight), 'Reporting unit', 'Limit of
964 detection/quantification', and the codification of analytical results under FoodEx classification (EFSA,
965 2011a). The outcome of the data analysis is presented in Section 3.2.1.

966 The left-censored data (analytical data below the LOD or limit of quantification (LOQ)) were treated by 967 the substitution method as recommended in the 'Principles and Methods for the Risk Assessment of 968 Chemicals in Food' (WHO/IPCS, 2009). The same method is described in the EFSA scientific report 969 'Management of left-censored data in dietary exposure assessment of chemical substances' (EFSA, 970 2010b) as an option for the treatment of left-censored data. The guidance suggests that the lower 971 bound (LB) and upper bound (UB) approach should be used for chemicals likely to be present in the 972 food. At the LB, results below the LOQ or LOD were replaced by zero; at the UB, the results below the 973 LOD were replaced by the numerical values of the LOD and those below the LOQ were replaced by the 974 value reported as LOQ.

# 975 **2.4. Food consumption data**

The EFSA Comprehensive European Food Consumption Database (Comprehensive Database) provides a compilation of existing national information on food consumption at individual level and was first built in 2010 (EFSA, 2011a; Huybrechts et al., 2011; Merten et al., 2011). Details on how the Comprehensive Database is used are published in the Guidance of EFSA (EFSA, 2011a). The latest version of the Comprehensive Database, updated in 2020, contains results from a total of 69 different dietary surveys carried out in 25 different Member States covering 134,929 individuals.

- 982 Within the dietary studies, subjects are classified in different age classes as follows:
- 983 Infants: < 12 months old

984 Toddlers:  $\geq$  12 months to < 36 months old

- 985 Other children:  $\geq$  36 months to < 10 years old
- 986 Adolescents:  $\geq$  10 years to < 18 years old
- 987 Adults:  $\geq$  18 years to < 65 years old
- 988 Elderly:  $\geq$  65 years to < 75 years old
- 989 Very elderly:  $\geq$  75 years old

990Seven additional surveys provided information on specific population groups: 'Pregnant women'991(Austria:  $\geq$  19 years to  $\leq$  48 years old, Cyprus:  $\geq$  17 years to  $\leq$  43 years old; Latvia:  $\geq$  15 years to  $\leq$ 99245 years old, Spain:  $\geq$  21 years to  $\leq$  46 years old, Portugal: 17 years old to 46 years old) and 'Lactating993women' (Greece:  $\geq$  28 years to  $\leq$  39 years old, Estonia: 18 years old to 45 years old).

994 For chronic exposure assessment, food consumption data were available from different dietary surveys 995 carried out in 23 different European countries. When for one particular country and age class two 996 different dietary surveys were available, only the most recent one was used. This resulted in a total of 997 44 dietary surveys selected to estimate chronic dietary exposure.

In Annex B (Table B.4), these dietary surveys and the number of subjects available for the chronic
 exposure assessment are described.

1000 The food consumption data gathered by EFSA in the Comprehensive Database are the most complete 1001 and detailed data currently available in the EU. Consumption data were collected using single or 1002 repeated 24- or 48-hour dietary recalls or dietary records covering from three to seven days per subject. 1003 Because of the differences in the methods used for data collection, direct country-to-country 1004 comparisons can be misleading.



# 1005 **2.5.** Food classification

1006 Consumption and occurrence data were classified according to the FoodEx classification system (EFSA, 1007 2011b). FoodEx is a food classification system developed by EFSA in 2009 with the objective of 1008 simplifying the linkage between occurrence and food consumption data when assessing the exposure 1009 to hazardous substances. The system consists of a large number of individual food items aggregated 1010 into food groups and broader food categories in a hierarchical parent-child relationship. It contains 20 1011 main food categories (first level), which are further divided into subgroups having 140 items at the 1012 second level, 1,261 items at the third level and reaching about 1,800 endpoints (food names or generic food names) at the fourth level. 1013

# 1014 **2.6. Exposure assessment**

The CONTAM Panel considered that only chronic dietary exposure to HBCDDs had to be assessed. As suggested by the EFSA Working Group on Food Consumption and Exposure (EFSA, 2011c), dietary surveys with only one day per subject were not considered for chronic exposure as they are not adequate to assess repeated exposure. Similarly, subjects who participated only one day in the dietary studies, when the protocol prescribed more reporting days per individual, were also excluded for the chronic exposure assessment. Not all countries provided consumption information for all age groups, and in some cases the same country provided more than one consumption survey.

For calculating the chronic dietary exposure to HBCDDs, food consumption and body weight data at the individual level were accessed in the Comprehensive Database. Occurrence data and consumption data were linked at the relevant FoodEx level.

1025 The mean and the high (95th percentile) chronic dietary exposures were calculated by combining mean 1026 occurrence values for each food collected in different countries (pooled European occurrence data) with 1027 the average daily consumption of each food at individual level in each dietary survey and age class. 1028 Consequently, individual average exposures per day and body weight were obtained for all individuals. 1029 Based on the distributions of individual exposures, the mean and 95th percentile exposures were 1030 calculated per survey and per age class. Dietary exposure was assessed using overall European LB and 1031 UB mean occurrence of HBCDDs. All analyses were run using the SAS Statistical Software (SAS 1032 enterprise guide 7.15).

# 1033 2.7. Risk characterisation

1034 The general principles of the risk characterisation for chemicals in food as described by WHO/IPCS 1035 (2009) will be applied as well as the different EFSA guidance documents relevant to this step of the risk 1036 assessment (see **Annex A**).

# 1037 **3.** Assessment

- **3.1.** Hazard identification and characterisation
- 1039 3.1.1. Toxicokinetics

# 1040 **3.1.1.1. Toxicokinetic studies in experimental animals**

1041 Information on the toxicokinetics of HBCDDs is limited. Since the previous EFSA assessment (EFSA

1042 CONTAM Panel, 2011a), new information has become available regarding the toxicokinetics. The text

below gives a description of previous knowledge complemented with new data. In general, human data

about the toxicokinetics of HBCDDs are less abundant than in experimental animal species.



#### 1045 *3.1.1.1.1. Absorption*

#### 1046 **Mice**

1047 Szabo et al. (2010, 2011a) and Sanders et al. (2013) studied the toxicokinetics of γ-[<sup>14</sup>C]-HBCDD (Szabo 1048 et al., 2010), α-[<sup>14</sup>C]-HBCDD (Szabo et al., 2011a) and β-[<sup>14</sup>C]-HBCDD (Sanders et al., 2013) in female 1049 C57BL/6 mice. Mice (n = 4–8) were given a single dose of HBCDD (3, 10, 30 or 100 mg/kg bw) by oral 1050 gavage, and urine and faeces were collected daily for 4 days. Another group of female C57BL/6 mice 1051 (n = 6–8) received an i.v. single dose (3 mg/kg bw).

By comparing kinetics and disposition between i.v. and oral routes of exposure, the authors calculated the percentage of the three stereoisomers that was absorbed into the systemic circulation four days after treatment. The absorption of the three stereoisomers estimated by the authors after oral gavage was around 90%, >85%, and 83%, for  $\alpha$ -[<sup>14</sup>C]-HBCDD, β-[<sup>14</sup>C]-HBCDD, γ-[<sup>14</sup>C]-HBCDD, respectively.

#### 1056 **Rats**

1057 Hakk (2016) administered  $\alpha$ -,  $\beta$ - and  $\gamma$ -[<sup>14</sup>C]-HBCDD individually at an oral dose of 3 mg/kg bw to 1058 Sprague-Dawley rats (n = 3 rats per isomer). Urine and faeces were collected daily over 96 h and 1059 analysed, including animal tissue for radioactivity quantification. The authors estimated an oral 1060 absorption of 73–83% for the three isomers (79.7%, 83.2% and 72.9% of the administered dose, 1061 respectively for  $\alpha$ -,  $\beta$ - and  $\gamma$ -[<sup>14</sup>C]-HBCDD).

1062 *3.1.1.1.2. Distribution* 

#### 1063 **Mice**

1064 The distribution differs between the isomers (Szabo et al., 2010, 2011a; Sanders et al., 2013). Szabo 1065 et al. (2011a) studied the distribution of  $\alpha$ -[<sup>14</sup>C]-HBCDD in female C57BL/6 mice following a single (3, 1066 10, 30 or 100 mg/kg) and 10-day repeated (3 mg/kg) exposure, and showed that  $\alpha$ -HBCDD was 1067 distributed in the following tissues (for the 10-day repeated exposure at 3 mg/kg): adipose tissue > 1068 liver > skin > blood > muscle > lung > brain > spleen > kidney > thymus (other tissues were not 1069 reported/analysed by the authors).

Szabo et al. (2010) performed the same study design with  $\gamma$ -[<sup>14</sup>C]-HBCDD. After a single oral exposure to  $\gamma$ -[<sup>14</sup>C]-HBCDD (3, 10, 30 or 100 mg/kg) and 10-day repeated exposure (3 mg/kg), the distribution of radioactivity was found in the following tissues (for the 10-day repeated exposure at 3 mg/kg): liver skin > blood > muscle > lung > kidney > brain > adrenals = thymus > adipose tissue > spleen > bladder (other tissues were not reported/analysed by the authors).

1075 Based on the same protocol as Szabo et al. (2010, 2011a), Sanders et al. (2013) showed that after 1076 single oral exposure to  $\beta$ -[<sup>14</sup>C]-HBCDD (3, 10 or 30 mg/kg),  $\beta$ -HBCDD was distributed in the following 1077 tissues (24 h after a single dose of 3 mg/kg  $\beta$ -HBCDD): adipose > muscle > liver > skin > blood > 1078 kidney > lung > brain (other tissues were not reported/analysed by the authors).

# 1079 Neonatal mice

Szabo et al. (2011b) performed a toxicokinetic study on neonatal mice (PND10) with a single oral dose of 3 mg/kg bw of  $\alpha$ - and  $\gamma$ -[<sup>14</sup>C]-HBCDD. They compared their results with a previous experiment on adult mice (PND60) (Szabo et al., 2010, 2011a). They observed a parallel distribution of both isomers (adipose, blood, brain, kidney, muscle and skin) in neonate and adult mice, but  $\alpha$ -[<sup>14</sup>C]-HBCDD and  $\gamma$ -1084 [<sup>14</sup>C]-HBCDD levels were higher in pups than in adults.

#### 1085 **Rats**

1086Hakk (2016) showed that α-, β- and γ-[14C]-HBCDD, when administered individually as a single oral1087dose of 3 mg/kg bw to Sprague-Dawley rats, were distributed preferentially to adipose tissue, adrenals,1088skin and gastronintestinal tract 96 h after administration in the following tissues: For α-[14C]-HBCDD:



1089 adipose > gastrointestinal tract and content > skin > liver > muscle > kidney = testes > adrenals > 1090 lung > thymus = spleen > heart = brain = plasma = heart. For  $\beta$ -[<sup>14</sup>C]-HBCDD: gastrointestinal tract 1091 and content > kidney > liver > adipose > testes > lung > muscle > adrenals > plasma > heart > spleen 1092 > brain = thymus. For  $\gamma$ -[<sup>14</sup>C]-HBCDD: gastrointestinal tract and content > adipose > skin > liver > 1093 muscle > kidney > adrenals > plasma = testes > spleen > thymus = heart > lung > muscle > adrenals 1094 > plasma > brain = thymus > brain.

1095 In a 90-day repeated dose toxicity study with CrI:CD(Sprague-Dawley)IGS BR rats, Chengelis et al. 1096 (2001, as cited by ECB, 2008, EFSA CONTAM Panel, 2011a) administered 0 or 1,000 mg/kg bw per day 1097 of technical HBCDD. The authors found that the concentrations in adipose tissue of the α-stereoisomer 1098 were higher than those of the β- and γ-stereoisomers throughout the study period.

1099 In a 28-day oral dose toxicity study, van der Ven et al. (2006) administered to male and female rats 0, 1100 0.3, 1, 3, 10, 30, 100 or 200 mg/kg bw per day of technical HBCDD (10.3%  $\alpha$ -, 8.7%  $\beta$ - and 81.0% y-1101 HBCDD) by gavage. Only the liver and adipose tissue were analysed at the end of the experiment. The 1102 liver concentrations of HBCDDs were higher in females than in males over the entire dose range (on 1103 average about five times). In adipose tissue, these levels were 2.4 and 1.4 times higher than in liver fat, in females and males, respectively. The authors analysed the HBCDD partitions (ratio  $\gamma$ -/ $\alpha$ -HBCDD) 1104 1105 in the liver, and showed it was dose dependent: the average ratio was 4.2, 4.9, 4.0, 2.2, 3.1, 0.6 and 1106 0.4 in females, and 2.3, 1.7, 3.0, 2.0, 1.5, 1.5 and 0.9 in males according to the increased the dose.

1107 van der Ven et al. (2009) performed a one-generation reproduction study in Wistar rats where technical 1108 HBCDD (10.3%  $\alpha$ -, 8.7%  $\beta$ - and 81.0%  $\gamma$ -HBCDD) was administered via the diet at different 1109 concentrations of 0, 0.1, 0.3, 1, 3, 10, 30 or 100 mg/kg bw per day. Parental male rats were exposed 110 weeks whereas females were exposed for 8 weeks (starting 2 weeks prior to mating). The authors 111 analysed the HBCDD partitions (ratio  $\gamma$ -/ $\alpha$ -HBCDD) in the liver, and showed that the ratio  $\gamma$ / $\alpha$  was 2.2, 1.4, 0.9 and 0.4 in females (for dose between 1 and 100 mg/kg bw per day), and 1.4, 0.8 and 0.2 in 1113 males (for doses between 10 and 100 mg/kg bw per day).

# 1114 *3.1.1.1.3. Metabolism*

#### 1115 **Mice**

Hakk et al. (2012) studied the metabolism of  $\alpha$ - and y-HBCDD in adult C57BL/mice. Six female mice 1116 were exposed to a single oral dose of 3 mg/kg bw ( $\alpha$ - or  $\gamma$ -[<sup>14</sup>C]-HBCDD). Collection of urine and faeces 1117 was done during 4 consecutive days, and an additional group was sacrificed 3 h post-exposure in order 1118 1119 to measure metabolites at tissue levels. The authors found metabolites in liver, blood, fat, brain, bile, 1120 urine and faeces. For  $\alpha$ -HBCDD, the authors found 4 monohydroxylated metabolites in faeces and one of them in liver, brain and adipose tissue. For y-HBCDD, the authors found one monohydroxylated and 1121 three dihydroxylated metabolites in faeces. The monohydroxylated metabolite was also measured in 1122 1123 liver and adipose tissue. Hakk et al. (2012) hypothesised that deiodinase may be involved in the 1124 debromination of HBCCDs in mice. The authors performed enzymatic hydrolysis of urine and did not 1125 detect the presence of glucuronic acid or sulfate conjugates in urine.

1126 In female mice, Szabo et al. (2011a) found different polar metabolites of  $\alpha$ -HBCDD at high levels in the 1127 liver (38% of the dose applied) and faeces (66% of the dose applied). These metabolites were also 1128 observed in the blood and bile.

1129 Szabo et al. (2010) in their study in mice exposed to  $\gamma$ -HBCDD, detected metabolites in the liver, blood,

bile, urine, and faeces between 3 and 24 h after exposure, but did not identify them.

1131 Sanders et al. (2013) demonstrated that  $\beta$ -[<sup>14</sup>C]-HBCDD was metabolised following oral and i.v.

administration of 3 mg/kg. The authors quantified the radioactivity in urine, tissue and faeces extracts. Although metabolites were not identified, the authors found that most of the  $\beta$ -[<sup>14</sup>C]-HBCDD-metabolites



1134 were excreted in urine (hydrophilic metabolites) and that 30% of the  $\beta$ -[<sup>14</sup>C]-HBCDD-metabolites in the 1135 faeces were non-extractable.

#### 1136 **Rats**

1137 Laurence et al. (2010) used incubations of  $\alpha$ -,  $\beta$ -, and  $\gamma$ -HBCDD with liver microsomes from 1138 phenobarbital treated rats. The authors described the formation of hydroxy-HBCDD metabolites, but 1139 without characterisation. In an additional experiment, the authors examined the reductive 1140 debromination of  $\alpha$ - and  $\gamma$ -HBCDD. After incubation with dithiothreitol (DTT), used for debromination, 1141 the authors did not find debrominated metabolites.

1142 Roosens et al. (2011) incubated  $\alpha$ -,  $\beta$ -, and  $\gamma$ -HBCDD with liver microsomes from phenobarbital treated 1143 rats. The authors detected 3 types of metabolites: the major ones were hydroxy-metabolites of HBCDDs, 1144 followed by a hydroxyl-debrominated metabolite (pentabromo) and to a minor extent a debrominated 1145 (pentabromo) metabolite.

1146 Esslinger et al. (2011) studied the metabolism of  $\alpha$ -,  $\beta$ -, and  $\gamma$ -HBCDD in rat liver microsomes. They 1147 reported several hydroxylated metabolites of HBCDDs but no debrominated metabolites.

1148 In a similar experiment, following incubation of  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCDD with rat liver microsomes, Abdallah 1149 et al. (2014) identified four monohydroxylated and two dihydroxylated metabolites. The authors 1150 suggested that metabolism of HBCDDs was mediated by enzymes of the CYP450 system, and that these 1151 are important during the stereo selective phase I oxidative metabolism of HBCDDs (in rat).

In Wistar rats exposed to 30 and 100 mg technical HBCDD/kg bw per day for 28 days, several metabolites were identified including four different types of hydroxylated HBCDD metabolites in adipose tissue, liver, lung and muscle (Brandsma et al., 2009). The authors stated that oxidation and reductive debromination were common metabolic routes for HBCDDs, although it was not possible to determine whether individual stereoisomers underwent the same metabolism.

1157 In male Sprague-Dawley rats dosed orally with 3 mg/kg bw of  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCDD, Hakk (2016) 1158 quantified several metabolites in urine, faeces, adipose tissue, brain, liver and gastrointestinal tract. The 1159 authors showed that  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCDD were metabolised extensively (51%, 65% and 80%, 1160 respectively) via different pathways.  $\alpha$ -HBCDD was metabolised into two monohydroxylated 1161 metabolites, while  $\beta$ - and  $\gamma$ -HBCDD were metabolised via different pathways (stereo isomerisation, 1162 oxidation, dehydrogenation, reductive debromination, and ring opening).

# 1163 **Bio-isomerisation of HBCDDs**

1164 Some authors have observed high concentration of  $\alpha$ -HBCDD after a long period in adipose tissue in 1165 rainbow trout and in rat (Law et al., 2006), and suggested a bio-isomerisation from  $\beta$ - and  $\gamma$ -HBCDD to 1166  $\alpha$ -HBCDD (Szabo et al., 2010). In one study, bio-isomerisation of  $\beta$ - to  $\gamma$ -HBCDD was observed (Sanders 1167 et al., 2013).

- 1168 *3.1.1.1.4. Elimination*
- 1169 **Mice**
- 1170 <u>α-HBCDD</u>

1171 The elimination of  $\alpha$ -[<sup>14</sup>C]-HBCDD-derived radioactivity after oral administration (3, 10, 30, and 100 1172 mg/kg bw) was mainly in the faeces (approximately 40% of the administered dose) and to a lesser 1173 extent in the urine (approximately 30% of the administered dose). As shown by the authors, more than 1174 50% of the cumulative elimination of  $\alpha$ -HBCDD radioactivity following a single oral dose was excreted 1175 via faeces and urine during the first 24 h (Szabo et al., 2011a). The authors also indicated that 66% of 1176 the eliminated radioactivity (in urine and faeces) was  $\alpha$ -HBCDD derived metabolites (after the first day).



- 1177 After administration of 3 mg/kg bw, the authors studied the tissue distribution and elimination over 14
- days. By measuring concentrations in tissues, the authors showed a biphasic profile of elimination with
- an initial steep decline (initial half-life or alpha phase) followed by a second steep (terminal half-life or
- 1180 beta phase) (Szabo et al., 2011a) (see **Table 4**).

Half-lives (days)			
Initial <sup>(a)</sup>	Terminal (a)		
$0.4 \pm 0.1$	$3.0 \pm 0.7$		
$0.1 \pm 0.0$	0.5 ± 0.2		
$0.3 \pm 0.1$	15 ± 12.1		
$0.2 \pm 0.0$	$2.1 \pm 0.4$		
$0.3 \pm 0.1$	8.0 ± 5.0		
$0.1 \pm 0.6$	3.0 ± 2.0		
—	17 ± 6.4		
	Initial (a) $0.4 \pm 0.1$ $0.1 \pm 0.0$ $0.3 \pm 0.1$ $0.2 \pm 0.0$ $0.3 \pm 0.1$		

**Table 4.** Estimated tissue half-lives (in days) of  $\alpha$ -HBCDD in mice (from Szabo et al., 2011a)

1182 (a):  $\pm$  standard deviation considering n = 4–8.

# 1183 Neonatal mice

1184 Szabo et al. (2011b) investigated the distribution and elimination of  $\alpha$ - and  $\gamma$ -HBCDDs in 10-day-old 1185 mice and compared these neonatal levels to adult levels previously reported (Szabo et al., 2010; Szabo 1186 et al., 2011a). According to the authors, 33% of the dose remained in the body in the pups, as compared 1187 to 25% in adult mice, 24 h after oral exposure to  $\alpha$ -[<sup>14</sup>C]-HBCDD.

# 1188 <u>β-HBCDD</u>

Sanders et al. (2013) showed that approximately 90% of the 3 mg/kg bw administered dose of  $\beta$ -HBCDD was excreted in urine and faeces within 24 h of treatment (primarily as  $\beta$ -HBCDD derived metabolites). At higher doses (30 and 100 mg/kg) the elimination of  $\beta$ -[14C]-HBCDD via urine and faeces was less than at the lowest dose. Although the majority of the  $\beta$ -[<sup>14</sup>C]-HBCDD was excreted within 48 h, the authors found measurable amounts of radioactivity in tissues 4 days post administration.

After administration of 3 mg/kg bw, the authors studied the tissue distribution and elimination over 14 days. By measuring concentrations in tissues, the authors showed a biphasic profile of elimination with an initial steep decline (initial half-life or alpha phase) followed by a second steep (terminal half-life or beta phase) (see **Table 5**) (Sanders et al., 2013).

Sanders et al. (2013) stated that the elimination half-lives of  $\beta$ -[<sup>14</sup>C]-HBCDD in mice (see **Table 5**) were

significantly shorter than for  $\alpha$ -HBCDD, but similar to those for  $\gamma$ -HBCDD as calculated by Szabo et al. (2010, 2011a).

**Table 5.** Estimated tissue half-lives (in days) of β-HBCDD in mice (from Sanders et al., 2013)

Tissue	Half-liv	es (days)
lissue	Initial	Terminal
Blood	0.2	2.1
Liver	0.1	1.3
Muscle	0.04	5.3
Adipose tissue	ND	2.5
Brain	0.02	0.2
Lung	0.2	1.5
Skin	0.1	7.0



# 1202 <u>γ-HBCDD</u>

Szabo et al. (2010) showed that the major route of elimination of  $\gamma$ -[<sup>14</sup>C]-HBCDD was via the faeces in mice after oral exposure. From the analyses of faeces and urine 14 days after the administration, nearly 55% of the dose was excreted in faeces and 30% in urine. As shown by the authors, more than 50% of the cumulative elimination of  $\gamma$ -HBCDD radioactivity following a single oral dose was excreted via faeces and urine in the first 24 h. The authors indicated that 95% of the radioactivity that was eliminated, were  $\gamma$ -HBCDD derived metabolites (after the first day).

1209 After administration of 3 mg/kg bw, the authors studied the tissue distribution and elimination over 14 1210 days. By measuring concentrations in tissues, the authors showed a biphasic profile of elimination with

1211 an initial steep decline (initial half-life or alpha phase) followed by a second steep (terminal half-life or

1212 beta phase) (see **Table 6**) (Szabo et al., 2010).

Tissue	Half-live	es (days)
lissue	Initial	Terminal
Liver	0.3 ± 0.0	2.3 ± 0.2
Blood	0.3 ± 0.0	3.5 ± 0.3
Lung	$0.4 \pm 0.1$	2.3 ± 0.2
Kidney	0.2 ± 0.0	2.8 ± 0.2
Muscle	$1.0 \pm 0.1$	3.6 ± 0.3
Skin	$0.4 \pm 0.0$	5.2 ± 0.3
Brain	$0.1 \pm 0.0$	0.8 ± 0.1
Fat	$0.9 \pm 0.1$	3.6 ± 0.2

1213 **Table 6.** Estimated tissue half-lives (in days) of γ-HBCDD in mice (from Szabo et al., 2010)

Szabo et al. (2011b) investigated the distribution and elimination of  $\alpha$ - and γ-HBCDDs in 10-day-old mice and compared these neonatal levels to adult levels previously reported (Szabo et al., 2010, Szabo

1216 et al., 2011a). According the authors, 16% of the dose remained in the body in the pups, as compared

1217 to only 2% in adult mice, 24 h after oral exposure to  $\gamma$ -[<sup>14</sup>C]-HBCDD.

# 1218 **Rats**

1219Hakk (2016) showed after an oral dose of 3 mg/kg bw HBCDDs ( $\alpha$ -,  $\beta$ - and  $\gamma$ -) that faeces were the1220major route of excretion, cumulatively accounting for 42% of the dose for  $\alpha$ -HBCDD, 59% for  $\beta$ -HBCDD1221and 53% for  $\gamma$ -HBCDD after 96 h. In urine, the cumulative excretion accounted for 13% of the dose for

1222  $\alpha$ -HBCDD, 30% for  $\beta$ -HBCDD, and 21% for  $\gamma$ -HBCDD after 96 h.

1223 Elimination was slower in rats than in mice (Szabo et al., 2010, 2011b; Sanders et al., 2013).

1224 The elimination rates in urine and faeces show that  $\alpha$ -HBCDD has a higher propensity to accumulate 1225 than  $\beta$ - and  $\gamma$ -HBCDD.

Hakk (2016) calculated elimination half-lives in rat based on urine/faeces elimination, for  $\alpha$ -HBCDD,  $\beta$ -HBCDD and  $\gamma$ -HBCDD: the corresponding values were 3.3 ± 0.7 days, 1.1 ± 0.2 days and 2.0 ± 0.3

- days, respectively.
- 1229 *3.1.1.1.5. Summary on toxicokinetic studies in rodents*

1230 Results from mice and rat studies showed that the absorption from the gastrointestinal tract was rapid

with an oral absorption for HBCDDs after oral gavage around 85–90% of the total dose (by comparing

1232 kinetics and disposition between i.v. and oral routes of exposure).



1233 HBCDDs are distributed to a number of tissues, including adipose tissue, muscle and the liver. 1234 Elimination of HBCDDs-derived radioactivity is predominantly via faeces but it is also eliminated via 1235 urine. In mice, the studies suggest that more than 66%, 90% and 95% of the radioactivity excreted 1236 from the body correspond to different metabolites of  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCDD, respectively.

1237γ-HBCDD was more rapidly metabolised and eliminated than  $\alpha$ -HBCDD, with a terminal half-life in mice1238of 4 days (Szabo et al., 2010).  $\alpha$ -HBCDD was more biologically persistent due to a lower metabolism1239with a terminal half-life of 17 days (in adipose tissue). Some authors have suggested that debromination1240could be a pathway of HBCDD metabolism: in mice and rat, metabolic debromination and hydroxylation1241of HBCDDs. Conversion of γ-HBCDD to  $\alpha$ - and β-HBCDD have been reported. No stereoisomerisation of1242 $\alpha$ -HBCDD was reported.

1243 After oral administration in mice, elimination of  $\alpha$ -[<sup>14</sup>C]-HBCDD and  $\gamma$ -[<sup>14</sup>C]-HBCDD are mainly in the 1244 faeces (40% and 55% of the administered dose) and in the urine (both 30% of the administered dose). 1245  $\beta$ -[<sup>14</sup>C]-HBCDD was equally excreted in urine and faeces within 24 h of treatment.

1246 Predominance of α-HBCDD in tissue could be explained by either a difference in toxicokinetics, where 1247  $\gamma$ -HBCDD is metabolised and eliminated at a more rapid rate than α-HBCDD, and/or stereo isomerisation 1248 of γ-HBCDD to α-HBCDD.

# 1249 **3.1.1.2. Toxicokinetic studies in humans**

- 1250 *3.1.1.2.1. Absorption*
- 1251 No *in vivo* data were identified.

1252 Abdallah et al. (2012) showed different bioaccessibility values among HBCDD stereoisomers present in 1253 dust. From an *in vitro* test system including human gastrointestinal tract parameters, they reported 1254 92%, 80% and 72% of bioaccessibility for  $\alpha$ -,  $\beta$ - and y-HBCDD, respectively. The bioaccessibility was 1255 calculated from the percentage of the average mass of the three HBCDD stereoisomers in the 1256 supernatant of the test system to the average mass of the HBCDD stereoisomers present in the 1257 extracted dust. It was suggested that this may be attributed to the lower aqueous solubility of the y-1258 isomer (2  $\mu$ g/L) compared to the  $\alpha$ - and  $\beta$ -isomers (45 and 15  $\mu$ g/L, respectively). No significant change in the enantiomeric fractions of HBCDDs was observed in any of the studied samples. 1259

1260 *3.1.1.2.2. Distribution* 

HBCDDs have been detected in human adipose tissue, breast milk, placenta and blood, as well as inhair and faeces (see Section 3.1.1.4).

1263 *3.1.1.2.3. Metabolism* 

1264 Erratico et al. (2016) suggested that CYP3A4 was involved in the *in vitro* metabolism of the HBCDD 1265 stereoisomers. After incubation of human liver microsomes with racemic mixtures of  $\alpha$ -,  $\beta$ - and  $\gamma$ -1266 HBCDD, the authors found formation of several hydroxylated metabolites.

1267 Abdallah et al. (2015) performed an additional metabolic study with human hepatocytes incubated 1268 during 24 h with  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCDD. Chromatographic analysis revealed phase I oxidative metabolism 1269 of HBCDDs (hydroxylated metabolites) and indicated a reductive debromination by the detection of 1270 penta- and tetrabrominated metabolites. They also identified glucuronide metabolites of HBCDDs.



#### 1271 *3.1.1.2.4. Elimination*

1272 In a biomonitoring study in toddlers, Sahlström et al. (2015a) detected the presence of  $\alpha$ -,  $\beta$  and  $\gamma$ -1273 HBCDD in the faeces. However, this study did not distinguish between faecal excretion following biliary 1274 excretion or direct excretion (non-absorption by the gastrointestinal tract).

1275 Geyer et al. (2004, extended abstract) estimated a terminal elimination half-life for HBCDDs of 64 days 1276 (range: 23-219 days) in humans using a linear one-compartment open toxicokinetic model based on 1277 body burden and daily intake in humans of 142 ng per day. The authors estimated this daily intake and 1278 the total body burden in non-occupationally exposed adult humans from a Swedish market basket study. 1279 Assuming a bioavailability of 100%, the corresponding lipid mass would be 13.5 kg for males and 18.7 1280 kg for females. According to the authors, the large range observed in half-lives could be due to the 1281 difference of fat mass in humans that would affect bioaccumulation and elimination. The CONTAM Panel 1282 noted that the method to estimate the half-life is not fully described in the extended abstract and this 1283 leads to uncertainties regarding the half-life calculation (see **Section 3.5.4**).

1284Abdallah and Harrad (2011) compared the concentrations measured in human milk samples with the1285estimated intakes of HBCDDs using a simple one-compartment model. They obtained good agreement1286between the predicted and the observed body burdens of the three HBCDD stereoisomers by using a1287half-life for α-HBCDD in humans of 165 days and a half-life of 55 days for the β- and γ-isomers. For α-1288HBCDD this is representing 75% of the maximum half-life of 219 days estimated previously by Geyer et1289al. (2004, extended abstract), and for the β- and γ-isomers 25% of the maximum half-life of 219 days.

1290 *3.1.1.2.5. Summary on toxicokinetic studies in humans* 

Limited data are available on the toxicokinetics of HBCDDs in humans. As there is no *in vivo* data, an estimate of the oral absortion of HBCDDs cannot be made. Biomonitoring studies report that HBCDDs have been detected in different human fluids (blood, breast milk, faeces) and tissues (adipose, placenta, hair).

*In vitro* studies have suggested that CYP3A4 may be involved in metabolism of the HBCDD stereoisomers
 based on formation of several hydroxylated metabolites. There is also *in vitro* evidence of glucuronide
 formation.

1298 Only one study provided half-life values in humans with a value of 64 days (range of 23–219 days).

# 1299 **3.1.1.3. Transfer of HBCDDs from feed to food producing animals**

Several studies have been identified on the transfer of HBCDDs from feed into several farm animals, i.e.laying hens, broilers, ducks, pigs and fish.

#### 1302 *3.1.1.3.1. Laying hens, broilers and ducks*

1303 Fournier et al. (2012) studied the fate of ingested HBCDDs in laying hens. The authors fed 48 laying 1304 hens with HBCDDs at 1.1 ng/g diet (containing 0.1%, 0.2% and 99.7% of  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCDD, 1305 respectively) for 21 days and analysed the three stereoisomers in egg yolk, abdominal fat and liver after 1306 1, 4, 8, 11, 16, 21, 22, 24, 29 or 39 days. In the control hens,  $\alpha$ -HBCDD was quantified in abdominal fat (32–33 pg/g lipid weight at days 0 and 39, respectively) and in liver (51–69 pg/g lipid weight at days 1307 0 and 39, respectively). In the treated group, concentrations of  $\alpha$ - and y-HBCDD in egg yolk, abdominal 1308 1309 fat, and liver varied between days and between tissues. During the exposure period,  $\alpha$ -HBCDD 1310 concentrations varied in egg yolk (between 23–380 pg/g lipid weight), in abdominal fat (between 25– 1311 108 pg/g lipid weight) and in liver (between 150–235 pg/g lipid weight).  $\gamma$ -HBCDD concentrations varied 1312 in abdominal fat (between 23–490 pg/g lipid weight) and in liver (between 114–273 pg/g lipid weight). 1313 In the treated group, at day 39,  $\alpha$ -HBCDD concentrations of 80, 102 and 3.5 pg/g lipid weight were



quantified in abdominal fat, liver and yolk, respectively. y-HBCDD was not detected in egg yolk nor in 1314 1315 liver of the control group, but concentrations of 17–21 pg/g lipid weight were quantified in abdominal 1316 fat. In comparison, at 39 days, these concentrations were 33, 69 and 3.5 pg/g lipid weight in the control 1317 group. The authors suggested isomerization of y- to  $\alpha$ -HBCDD in laying hens. The authors estimated a 1318 transfer rate of 1.2% from ingested y-HBCDD to egg yolk (at steady state). Over the 21-day exposure 1319 period, the proportions of ingested y-HBCDD stored in abdominal fat and in liver were 0.74% and 1320 0.025%, respectively. Despite the very low level of  $\alpha$ -HBCDD in the feed (0.1% of the 1.1 ng/g),  $\alpha$ -1321 HBCDD was found in egg yolk, abdominal fat, and liver of the hens. The authors suggested isomerization 1322 of y- to  $\alpha$ -HBCDD in laying hens.

1323 Jondreville et al. (2017a) studied the elimination and accumulation of  $\alpha$ -HBCDD in broilers. The authors 1324 exposed 29 fast-growing and 50 slow-growing chickens to  $\alpha$ -HBCDD at 50 ng/g feed over 42 and 84 1325 days. An additional 10 animals for each group were used as control. The authors showed that the 1326 concentrations of  $\alpha$ -HBCDD in leg, breast, liver and abdominal fat were higher in slow-growing than in fast-growing strains. In both strains, the concentration of  $\alpha$ -HBCDD was higher in abdominal fat than 1327 1328 in other tissues. The calculated accumulation ratio (i.e. the ratio of the concentration of  $\alpha$ -HBCDD in 1329 the tissue (lipid based) to its concentration in feed, calculated at market age (6 and 12 weeks in fast-1330 growing and slow-growing broilers, respectively), was 5.2 and 6.7 in leg, 2.2 and 2.1 in breast muscle, 1331 8.4 and 8.9 in abdominal fat, in fast- and slow-growing chicken, respectively. In slow-growing chickens, 1332 the authors estimated half-life values of 53, 24 and 47 days in leg, muscle and abdominal fat, 1333 respectively. In a previous work of this group, Domínguez-Romero et al. (2016) demonstrated that after 1334 exposure to  $\alpha$ -HBCDD (50 and 5 ng/g feed) for 18 or 11 days,  $\alpha$ -HBCDD was transferred to eggs and 1335 significantly accumulated in adipose tissue. The calculated accumulation ratio was 5.2, 3.6 and 9.2 in 1336 eggs, liver and abdominal fat, respectively.

1337 Zheng et al. (2017) measured HBCDDs in diet sources and in chicken tissues in adult (n = 12) and 1338 hatchling chicken (n = 9). In feed samples, the highest HBCDD concentration measured was 0.69 ng/g 1339 dw while the highest concentration in soil was 28.8 ng/g dw. For adult chickens, the order of  $\alpha$ -HBCDD 1340 concentrations within the tissues studied was: gonad > visceral fat > pectoral muscle > intestine > 1341 stomach > kidney > lung > serum > liver. The authors did not detect  $\alpha$ -HBCDD in brain. For hatchling 1342 chicken, the order of  $\alpha$ -HBCDD concentrations was: pectoral muscle > liver (only 2 tissues studied).

Xia et al. (2018) studied the accumulation of HBCDDs in ducks exposed at 0 (n = 8, control group with 1343 1344 blank capsule), 0.8 mg/kg bw (n = 32) and 1.6 mg/kg bw per day (n = 32). At 0, 7, 14 and 21 days 1345 after exposure, eight ducks from each group were sacrified. After a 21-day depuration period, the order 1346 of HBCDD concentrations (in ww) at 0.8 mg/kg bw per day within the tissues studied was: skin > tongue 1347 > intestines > heart > qizzard > muscle > liver > lung > brain > blood. At 1.6 mg/kg per day, the 1348 order of HBCDD concentrations (in ww) within the tissues studied was: skin > tongue > intestines > 1349 gizzard > muscle > liver> heart > lung > brain > blood. The high levels in skin and tongue were 1350 probably attributed to the high fat content. The authors calculated half-lives for the different tissues, 1351 and these varied between 17–70 days for  $\alpha$ -HBCDD, 3.8–9.1 days for  $\beta$ -HBCDD and 8.8–15 days for  $\gamma$ -1352 HBCDD.

1353 *3.1.1.3.2. Pigs* 

1354 Royer et al. (2017) performed a study in order to establish a physiologically based pharmacokinetic model for  $\alpha$ -HBCDD in pigs. The authors exposed 56 pigs during 19, 49, 70, 91 and 112 days to  $\alpha$ -1355 HBCDD at 0, 32 or 297 µg/kg feed (not lipid based). Two control pigs were sacrified at day 1, and three 1356 pigs per treatment were sacrified at 20, 50, 71, 92 and 113 days. The liver, dorsal fat and semi-1357 1358 membranous muscle were collected. The authors found that  $\alpha$ -HBCDD concentrations were higher in 1359 adipose tissue than in muscle and were lower in the liver. The authors estimated accumulation ratios 1360 for  $\alpha$ -HBCDD of 6.6 and 5.3, respectively, in dorsal fat and semi-membranous muscle. The accumulation 1361 ratio for liver was not estimated.



## 1362 *3.1.1.3.3. Fish*

Berntssen et al. (2011) developed a toxicokinetic model in Atlantic salmon. For this purpose, the authors exposed Atlantic salmon to  $\alpha$ -HBCDD at 280 µg/kg feed per day for 2 months, followed by a depuration period of 3 months. Three fish were sacrificed on day 0, 8, 14, 26, 41 and 54 of the uptake period, and on day 0, 1, 3, 5, 7, 14, 28, 42, 55 and 83 of the exposure period. Based on the fat storage in the fillet for  $\alpha$ -HBCDD, the authors showed that their model simulated a concentration of  $\alpha$ -HBCDD in the range of 0.2–1.8 µg/kg with a low background and high levels of  $\alpha$ -HBCDD in the feed at 600 days.

1369 Esslinger et al. (2010) exposed mirror carps (n = 56 per group) to racemic  $\gamma$ -HBCDD in order to study 1370 potential bioisomerisation. The commercial fish feed contained a concentration of 26.0 ± 1.6 ng/g of 1371 (+)- $\gamma$ -HBCDD and 16.8 ± 0.4 ng/g of (-)- $\gamma$ -HBCDD. After treatment for 14 days, an increase of  $\gamma$ -HBCDD 1372 was measured in the fillet (for both racemic) compared to the control group. The authors did not observe 1373 bioisomerisation of  $\gamma$ -HBCDD in this species.

1374 Zhang et al. (2014) studied the accumulation of  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCDD in mirror carps (n = 40 fish per 1375 group) over 30 days followed by a depuration period of 30 days. The authors detected the three HBCDD 1376 stereoisomers in different tissues (gills, viscera, muscle and skin) of the mirror carp with an increased 1377 concentration over the 30 days. The authors calculated kinetic bioconcentration factors (BCFs). For  $\alpha$ -HBCDD the BCFs were  $8.58 \times 10^3$ ,  $1.15 \times 10^4$ ,  $5.57 \times 10^3$  and  $6.40 \times 10^3$  in gills, viscera, muscle and 1378 1379 skin, respectively. For  $\beta$ -HBCDD, the BCFs were 322, 642, 187 and 204, and for  $\gamma$ -HBCDD were 237, 584, 221 and 227, respectively. The accumulation for  $\alpha$ -HBCDD was higher than for the  $\beta$ - and y-1380 1381 stereoisomers. At the end of the depuration period, the authors described transformation of  $\beta$ - and yisomers into  $\alpha$ -HBCDD (53.0–92.9% for  $\beta$ -HBCDD and 96.2–98.6% for  $\gamma$ -HBCDD). 1382

1383 *3.1.1.3.4. Toxicokinetic models for the transfer from feed to food producing animals* 

Meda et al. (2017, 2020) described generic PBPK models for broilers and pigs. These models have been validated using data from Jondreville et al. (2017a) (for broilers) and data from Royer et al. (2017) for pig. These models were calibrated to take into account differences in lipid content according to species, and validated using experimental data where animals were exposed to HBCDDs.

1388 *3.1.1.3.5. Summary on the transfer from feed to food of animal origin* 

1389 In broilers,  $\alpha$ - and  $\gamma$ -HBCDD have been measured in egg yolk, abdominal fat and liver (lipid based). 1390 Isomerisation of  $\gamma$ - to  $\alpha$ -HBCDD in laying hens has been suggested by some authors. In ducks, 1391 accumulation of HBCDDs has been described, with high levels in skin and tongue, probably attributed 1392 to the high fat content.

- 1393 In pigs, HBCDD concentrations seemed to be higher in the adipose tissue than in the muscle and in the 1394 liver.
- 1395 In fish, HBCDDs have been also detected in tissue with high fat content.

1396 In summary, from the few available studies on the transfer of HBCDDs from feed to food of animal 1397 origin, accumulation of HBCDDs mainly happens in adipose tissue among laying hens, broilers, ducks

- 1398 (organ with high fat content), pigs and fish. In the laying hens, transfer to eggs was also described.
- 1399 Some toxicokinetic models have been constructed based on data obtained in broilers and pigs. These 1400 models could be used to assess the risk of meat contamination by HBCDD in animals.

## 1401 **3.1.1.4. Levels in human tissues**

### 1402 *3.1.1.4.1. Human milk*

1403 The previous EFSA Opinion on HBCDDs (EFSA CONTAM Panel, 2011a) summarised the occurrence data 1404 in human milk published in literature until 2010. Except for one study, the levels ranged between 0.13 1405 and 31 ng/g lipid with mean and/or median values generally below 2 ng/g lipid.

1406 **Table 7** summarises the occurrence data on HBCDDs in human milk from European mothers published 1407 in the open domain since the previous EFSA Opinion on HBCDDs. In case of analysis by GC based 1408 methods for which a separation of the diastereomers is not possible, the results are given as Total 1409 HBCDDs. Although the data were published between 2011 and 2020, most of the samples were collected 1410 between 2000 and 2010. In general, the results are in a comparable range with the data reported in 1411 the previous EFSA Opinion. This holds also true for the share of  $\alpha$ -HBCDD which generally contributes 1412 more than 75% to the sum of HBCDDs.

1413 It is noteworthy that Abdallah and Harrad (2011) also detected the degradation products 1414 pentabromocyclododecenes (average = 0.04 ng/g lipid; n = 9) and tetrabromocyclododecadienes 1415 (average = 0.15 ng/g lipid; n = 25) in some samples. Moreover, they reported on the HBCDD 1416 enantiomer profiles in the analysed samples. While the enantiomer fractions (EFs) of  $\beta$ -HBCDD and  $\gamma$ -1417 HBCDD showed no significant deviations from racemic (average EF of 0.49 and 0.51, respectively), 1418 substantial enrichment of the (-)- $\alpha$ -HBCDD enantiomer was evident (EF = 0.29). Similar results have 1419 been reported by Eljarrat et al. (2009).

Harrad and Abdallah (2015) measured concentrations of HBCDDs and the HBCDD degradation products tetrabromocyclododecadienes along with PBDEs, and TTBPA in human milk collected in 2010–2011 from 10 first-time mothers from the UK each over the first 12 months post-partum, amounting to 120 samples overall. While for HBCDDs the correlation analysis revealed no statistically significant change in concentration with time over the 12 months lactation period covered, the concentration of the sum of tetrabromocyclododecadienes showed a significant increase with 1.4% per month (p = 0.02).

1426 Antignac et al. (2016) analysed human milk samples from French, Danish and Finnish women for a 1427 number of POPs, including  $\alpha$ -HBCDD. A statistical evaluation of the results showed no significant 1428 correlations between PBDEs and HBCDD, and between the age of the mother and the respective HBCDD 1429 content.

1430 Thomsen et al. (2010) studied elimination rates of HBCDDs and other persistent organic pollutants in 1431 nine Norwegian single-child primiparous mothers and one mother breast-feeding her second child by 1432 collecting breast-milk samples (n = 70) monthly from about two weeks to up to twelve months after 1433 birth. The number of samples collected by each mother was 3–10. While three mothers sampled breast 1434 milk during 2001 and 2003, six mothers sampled during 2005 and 2006, and one in 2008 and 2009. 1435 Total HBCDDs could be detected in 32 samples with a median of 0.27 ng/g lipid (range: <0.2–3.0 ng/g 1436 lipid). No clear time trend during the laction periods could be observed as the HBCDD levels showed 1437 large variations and the concentrations were below LOQ in several samples.

1438Tao et al. (2017) analysed 10 human milk samples collected in 2014–2015 in the UK and compared the1439results with the data reported by Abdallah and Harrad (2011). No statistically significant change in the1440HBCDD levels in human milk collected between 2014–2015 and 2010 was observed.

1441 Croes et al. (2012) analysed a human milk pool consisting of 84 individual samples collected in 2009– 1442 2010 in a rural area in Belgium and found a 153% higher level compared to the pooled Belgium sample 1443 analysed in 2006 within the World health Organisation (WHO) field study.

1444 Sahlström et al. (2015b) analysed two human milk pools collected from 30 mothers each in 2009 and 1445 2010, respectively within a study where they compared estimated intakes of HBCDDs and other BFRs



1446 via diet and dust to internal concentrations in a Swedish mother–toddler cohort. Levels for  $\alpha$ -,  $\beta$ - and 1447  $\gamma$ -HBCDD as well as for  $\Sigma$ HBCDDs (sum of  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCDD) were given as 8.4, <0.18, <1.3 and 9.4 1448 pg/g ww, respectively. Considering the mean lipid content of 3%, the results are below 0.5 ng/g lipid 1449 and thus are at the lower end of reported occurrence data in human milk. The concentrations in both 1450 pools were similar.

Lignell et al. (2014) determined several POPs including Total HBCDDs in human milk collected 2012 from 30 randomly recruited primiparas from Uppsala (Sweden) and compared the results with respective concentrations in earlier surveys. The temporal trend analysis included results from samples collected in 2002–2004 and 2009–2012. The trend for Total HBCDDs is reported to be uncertain and showed a non-significant decrease of 2.5% per year during the period 2002–2012.

In a follow-up study, Gyllenhammar et al. (2017) analysed 30 human milk samples collected 2015–2016 from Swedish mothers living in Uppsala for Total HBCDDs among other POPs. The results enabled an updated trend analysis. For Total HBCDDs, the authors reported a significant downward trend for the whole study period (1996–2016) with a declining rate of 2.0% per year. A significant change point was observed around year 2002–2003 with an increasing trend before that year and a decreasing trend after.

Wemken et al. (2020) analysed 16 pools from 92 Irish primiparas for α-, β-, and γ-HBCDD, 8 PBDEs, and decabromodiphenyl ethane (DBDPE). Human milk sampling and donor recruitment were comparable with the previous study of Pratt et al. (2013) in order to facilitate elucidation of possible time trends for BFRs in human milk from Irish women. Using a t-test, the authors compared the concentrations of ΣHBCDDs in individual pools of the two studies and reported that the current concentrations are significantly lower (p < 0.05) than in samples analysed by Pratt et al. (2013).

Based on scientific literature published between 1995 and 2011, Fång et al. (2015) performed a global review on spatial and temporal trends of the Stockholm Convention POPs, including HBCDDs in human milk. The levels reported for HBCDDs human milk collected in Africa and Asia are generally in the same range as the occurrence data analysed in samples from Europe at that time. The samples from Japan (collected between 1987 and 2007) and from Sweden (collected between 1987 and 2010) were reported to show increasing trends over the whole period of 5.4% per year (p<0.061) and 7.6% per year (p<0.001), respectively.

In their review, Shi et al. (2018) summarised data on HBCDDs in human milk from China and compared 1475 1476 the results with levels found in other regions of the world. A substantial increase of HBCDD levels in 1477 human milk from China collected in 2007 and in 2011 was observed. While in 24 pooled human milk 1478 samples from 12 provinces collected in 2007 the HBCDD concentrations ranged from <LOQ to 2.78 ng/g 1479 lipid, the respective levels in 29 pools from 16 provinces sampled in 2011 amounted to 1.02 to 81.1 1480 ng/g lipid, with mean and median levels of 10.1 and 6.83 ng/g lipid, respectively. Similar levels were 1481 found in 103 individual human milk samples collected in 2011 in Beijing with mean concentration of 1482 4.29 ng/g lipid (range: <LOQ-78.3 ng/g lipid). A comparison of the HBCDD levels with human milk 1483 analysed in other countries revealed that the contamination of human milk from China is higher than 1484 from most other regions worldwide.

1485Huang et al. (2020) analysed 111 human milk samples collected in 2014 from 27 Beijing mothers for1486HBCDDs and TBBPA, whereby each mother provided one sample per month for 3 months. All mothers1487were primiparous with an average age of 29.7 years. The ΣHBCDDs (sum of  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCDD)1488ranged from <LOD to 36.3 ng/g lipid, with mean and median concentrations of 7.58 and 5.67 ng/g lipid,</td>1489respectively. Compared to the levels analysed in 2011, a statistically significant increase in HBCDD levels1490was observed. A correlation analysis revealed no significant trend in concentrations over the three1491months of lactation.



1492 Since 1987, WHO (partly in cooperation with the United Nations Environment Programme (UNEP)) 1493 conducted several coordinated surveys on the occurrence of various POPs in human milk. These surveys were not primarily intended to compare levels of POPs among countries, but rather to examine levels 1494 1495 within countries over time. Therefore, strict protocols had to be followed with respect inter alia to 1496 selection of donors, location and time of sampling, storage and pooling of samples (generally 1 pool 1497 consists of 50 individual samples), and shipping of the samples to the laboratory (WHO, 2007). To 1498 ensure consistency in the analytical measurements, from the third round all samples were analysed by one laboratory, i.e. the EURL for Halogenated POPs in Feed and Food in Freiburg/Germany. The latest 1499 1500 results of these surveys on HBCDDs in European pooled human milk samples are given in Table 8 1501 (Malisch and Schächtele, 2019, see Documentation provided to EFSA). The table shows the levels of  $\alpha$ -1502 ,  $\beta$ -, and  $\gamma$ -HBCDD as well as the sum of the three isomers ( $\Sigma$ HBCDDs) as upperbound (UB) and lower bound (LB) values, respectively, given as ng/g lipid <sup>24</sup>. The last two columns present the ratio of the 1503 1504 predominant stereoisomer  $\alpha$ -HBCDD in relation to the sum of HBCDDs, calculated as UB and LB, 1505 respectively.

1506 Considering the most recent occurrence data generated as part of the WHO/UNEP coordinated human 1507 milk studies since the last EFSA Opinion on HBCDDs (EFSA CONTAM Panel, 2011a), collected and

analysed between 2014 and 2016, the UB and LB levels for the sum of the three stereoisomers range from 0.70–16.1 and 0.00–16.0 ng/g lipid, respectively. The UB and LB medians for the sum of the three stereoisomers are 3.12 and 3.0 ng/g lipid, respectively. As indicated in **Table 8**,  $\alpha$ -HBCDD is the predominant stereoisomer and generally contributes to more than 75% to the sum of HBCDDs.

1512 In two cases, human milk pools were collected from the same country in 2009 and 2014, and 2009 and

1513 2015, respectively. It is noteworthy that the levels for HBCDDs in the recent samples are substantially 1514 higher than in the earlier samples. However, the low number of samples does not allow deriving a

1515 general time trend from this observation.

<sup>&</sup>lt;sup>24</sup> In literature, both terms 'lipid' and 'fat' are cited, generally without stating which fractions or constituents were actually measured. Therefore, in this Opinion, both terms are considered synonymous, unless otherwise stated.



## 1516 **Table 7.** HBCDDs in human milk. All concentration levels are expressed in ng/g lipid.

Country	Year of	n	α <b>-HBCDD</b>	β-HBCDD	γ-HBCDD	ΣHBCDD	Total HBCDDs	Reference
country	sampling			mean (med	ian), range			
France	2010	106	<lod <sup="">(a)-23.6</lod>	<lod-6.6< td=""><td><lod-6.4< td=""><td>2.7–33.2</td><td>-</td><td>Inthavong et al. (2017)</td></lod-6.4<></td></lod-6.6<>	<lod-6.4< td=""><td>2.7–33.2</td><td>-</td><td>Inthavong et al. (2017)</td></lod-6.4<>	2.7–33.2	-	Inthavong et al. (2017)
France	2011–2014	41	(0.56), 0.22–4.21	-	-		-	Antignac et al. (2016)
Denmark	1997–2002	435	(0.31), 0.02–28.71	-	-	-	-	Antignac et al. (2016)
Finland	1997–2002	22	(0.31), 0.03–2.19	-	-	-	-	Antignac et al. (2016)
UK	2010	34	4.91 (3.17), 0.75–19.71	0.32 (0.30), 0.08–0.75	0.73 (0.56), 0.13–2.29	5.95 (3.83), 1.04–22.37	-	Abdallah and Harrad (2011)
UK	2010–2011	120 <sup>(b)</sup>	5.27	0.48	0.79	-	-	Harrad and Abdallah (2015)
UK	2014–2015	10	-	-	-	3.2 (2.9), 0.69–7.1	-	Tao et al. (2017)
Ireland	2010	11 <sup>(c)</sup>	2.59, (1.50–3.44)	0.46, (0.29–1.56)	0.47, (0.29–0.94)	3.52, (1.67–5.94)	-	Pratt et al. (2013)
Ireland	2016–2018	16 <sup>(d)</sup>	1.5 (1.7), 0.66-3.0	0.22 (0.34), <0.05-0.53	0.27 (0.21), <0.05-1.5	2.0 (1.8), 0.83-3.6	-	Wemken et al. (2020)
Czech Republic	2010	50	<1-76 <sup>(e)</sup>	-	-	-	-	Lankova et al. (2013)
Belgium	2009–2010	84 <sup>(f)</sup>	3.20	0.05	0.55	3.80	-	Croes et al. (2012)
Sweden	2008	18	-	-	-	-	<0.32-1.5	Björklund et al. (2012)
Sweden - Uppsala - Gothenburg - Lund - Lycksele	2000–2004	92 37 36 39	-	-	-	-	0.3 (<0.2–4.4) <0.4 (<0.4–2.4) 0.4 (<0.2–5.9) 0.4 (0.09–10)	Glynn et al. (2011)
Sweden	2012	30					0.40 (0.33), <0.20-1.5	Lignell et al. (2014)
Sweden	2015–2016	30					0.18 (0.15), <0.09-0.72)	Gyllenhammar et al. (2017)

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1517 (a): The LOD and LOQ were reported as 0.5 and 2.5 ng/g lipid for HBCDD diastereomers. α-, β-, and γ-HBCDD diastereoisomers were quantified in 8%, 3% and 3% of samples.

1518 (b): 12 samples of each 10 mothers sampled over a period of 12 months.

1519 (c): 11 pools from 109 first-time mothers.

1520 (d): 16 pools from 92 Irish primiparas.

1521 (e): The LOQ for  $\alpha$ -HBCDD is given as 1 ng/g lipid and the frequency of detection as less than 30%.

1522 (f): Pool of 84 samples from a rural area in Belgium.

1523

1524 **Table 8.** HBCDDs in human milk pools from various European countries as analysed in the frame of the WHO/UNEP coordinated studies

<b>.</b> .	N.	Lipid	α-HBCDD	β-HBCDD	γ-HBCDD	<b>ΣHBCDDs LB</b>	<b>EXAMPLE STATE</b> STATES	ratio α/Σ LB	ratio α/Σ UB		
Country	Year	%		ng/	g lipid			% 0			
Austria	2016	4.1	5.60	<0.10	< 0.10	5.60	5.80	100	97		
Switzerland	2016	4.2	0.50	<0.10	< 0.10	0.50	0.70	100	71		
Belgium	2015	3.6	2.00	< 0.06	< 0.06	2.00	2.12	100	94		
Lithuania	2015	3.4	3.00	0.30	< 0.06	4.00	4.06	75	74		
Malalaura	2015	4.1	8.00	< 0.06	0.20	8.00	8.06	100	99		
Moldova —	2009	4.7	2.84	< 0.10	< 0.10	2.84	3.04	100	93		
Netherlands	2014	3.0	0.60	< 0.06	< 0.06	0.60	0.72	100	83		
Czech Republic	2014	3.8	1.00	< 0.06	< 0.06	1.00	1.12	100	89		
Bulgaria	2014	3.9	2.00	< 0.06	< 0.06	2.00	2.12	100	94		
Croatia	2014	4.0	3.00	< 0.06	< 0.06	3.00	3.12	100	96		
Coordia	2014	4.0	4.00	< 0.06	< 0.06	4.00	4.12	100	97		
Georgia	2009	4.8	1.27	< 0.10	< 0.10	1.27	1.47	100	86		
Romania	2014	3.7	15.00	< 0.06	0.70	16.0	16.1	94	93		
Finland	2007	-	< 0.40	< 0.40	< 0.40	0.00	1.20	-	-		
Luxembourg	2006	3.5	1.10	< 0.40	< 0.40	1.10	1.90	100	58		
Slovak Republic	2006	3.2	1.80	< 0.20	< 0.20	1.80	2.20	100	82		
Hungary	2006	4.1	2.20	< 0.40	< 0.40	2.20	3.00	100	73		

1525 LB: lower bound. UB: upper bound



### 1526 *3.1.1.4.2. Human blood*

Recent analyses of human blood for HBCDDs are scarce, especially from European populations. In the previous EFSA assessment several studies were described reporting levels of HBCDDs in serum and one study in cord blood (EFSACONTAM Panel, 2011a). The results of the few published studies from Europe after 2010 focus on blood from adults and are summarised below.

Ålander et al. (2019) analysed 57 serum pools including 622 individual samples collected between 1996 and 2017 from first-time mothers in Uppsala (Sweden) for Total HBCDDs among other BFRs. The concentrations for Total HBCDDs were in most cases <LOQ which was around 0.5 ng/g lipid. A statistically significant decreasing trend of 4% per year was reported by the authors for the whole study period.

Jansen et al. (2018) analysed serum samples that were collected between 2012–2014 from 63 patients before and one year after bariatric surgery for organochlorine pesticides, PCBs and certain BFRs by GC/low resolution MS (LRMS). The LOD for Total HBCDDs was 0.033 ng/g lipid. Only 3% and 5% of the serum samples were above LOD before and one year after bariatric surgery, respectively. The Total HBCDD levels for the two time points ranged between <LOD–169 ng/g lipid and <LOD–65.2 ng/g lipid, respectively.

Bjermo et al. (2017) analysed 170 serum samples, collected in 2010–2011 from Swedish adults (age 50±17 years old) for HBCDDs and PBDEs by GC/LRMS. The LOD and limit of quantification (LOQ) for Total HBCDDs were 0.017 and 0.50 ng/g lipid, respectively. The share of samples above LOD and LOQ are reported as 63% and 13%, respectively. The levels of Total HBCDDs in serum ranged from <LOD– 77 ng/g lipid. The 95<sup>th</sup> percentile was 1.8 ng/g lipid. By replacing concentrations below LOD by ½ LOD, the median was calculated as 0.10 ng/g lipid.

1548 Darnerud et al. (2015) analysed 36 pooled serum samples from Swedish first-time mothers for HBCDDs 1549 and PBDEs by GC/LRMS. A total of 413 individual serum samples were collected between 1996 and 2010 1550 to prepare the pools. Each pool consisted of 5–25 indidividual samples (approximately 3 pools per year). 1551 The HBCDD levels seemed to be stable up to about 2005 and therafter decreased. Considering the 1552 entire study period from 1996-2010, a significant HBCDD decrease in the pooled serum samples was observed. The change was reported as -6.9% per year (p = 0.002). In addition, serum/breast milk 1553 1554 correlations for HBCDD and PBDE levels in 30 paired samples from individual mothers sampled in 2010 were studied. As most serum samples from 2010 were below LOQ, a calculation of HBCDD correlations 1555 1556 between serum and milk could not be performed.

Sahlström et al. (2014) analysed matched serum samples from 24 Swedish mothers (24–40 years old) and their toddlers (11–15 months of age) for HBCDDs and other BFRs. Samples were collected in 2009– 2010.  $\alpha$ -, β- and γ-HBCDD were not detected at method LOQs, reported as 0.44, 0.20 and 0.34 pg/serum sample <sup>25</sup>, respectively.

Lignell et al. (2013) investigated correlations between PBDEs and Total HBCDDs in paired blood serum and breast milk samples collected in 2010 from 30 mothers living in Uppsala (Sweden) with the aim to evaluate if the concentration of PBDE/HBCDD in breast milk is a good indicator of maternal body burden and possibly also of prenatal exposure of the infant. While the Total HBCDDs concentrations in breast milk ranged between <0.13 and 1 ng/g lipid, most of the levels in serum were below the LOQ which impeded the evaluation of correlations.

Fromme et al. (2016) analysed blood samples collected in 2013 in Germany from 42 randomly selected subjects (20–68 years old) for a number of POPs, including HBCDDs.  $\alpha$ - and β-HBCDD could only be

<sup>&</sup>lt;sup>25</sup> The authors report the method LOQs not per gram serum, but individually for each sample with respect to the different sample intakes of 0.5–5 g.



detected in 3 and 4 samples, respectively, with concentrations between the LOD of 5 ng/g lipid and theLOQ of 16 ng/g lipid. Maximum values were reported as 9 and 15 ng/g lipid, respectively.

Human serum samples (n = 61; age range 20–65 years) were collected in Greece in 2007 and analysed for HBCDDs, PBDEs, PCBs and organochlorine pesticides (Kalantzi et al., 2011). Thirty samples were collected from computer clerks of a large computer company working full-time with computers, and thirty-one from a control population with no computer use. HBCDDs were detected in 70% of the serum samples. The levels ranged between 0.49 and 38.8 ng/g lipid (median: 1.32; mean: 3.39 ng/g lipid). Differences in HBCDD body burden between the two study groups are not reported.

1577 In summary, the available data on HBCDDs in blood from European populations generally point to low 1578 levels with medians around 1 ng/g lipid, however, with some samples containing substantially higher 1579 concentrations. This is similar to global human HBCDD levels in blood as shown, e.g. for Canada (Rawn 1580 et al., 2014a), Korea (Kim and Oh, 2014), China (Shi et al., 2013), and Australia (Drage et al., 2017, 1581 2019).

1582 *3.1.1.4.3. Other human tissues* 

Malarvannan et al. (2013a) analysed HBCDDs in paired visceral fat and subcutaneous abdominal fat samples collected in 2010–2012 from 52 obese individuals in Belgium. Median concentrations of 4.1 ng/g lipid in visceral fat (range: 0.9–53 ng/g lipid) and 3.7 ng/g lipid in subcutaneous fat (range: 0.89– 89 ng/g lipid) for ΣHBCDDs (sum of  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCDD) were reported. The share of  $\alpha$ -HBCDD to the sum of the three HBCDD stereoisomers was 85% and 97%, respectively. The levels in the two fat depots were not significantly different. The HBCDD concentrations in the fat samples did not correlate with age.

Rawn et al. (2014b) determined HBCDDs in Canadian human fetal liver (n = 52) and placental tissue (n1590 = 142). Tissue samples were collected between 1998 and 2010 following elective pregnancy 1591 1592 terminations. While the concentration of  $\Sigma$ HBCDDs in fetal livers ranged from <1–4,500 ng/g lipid, the corresponding concentrations in placenta were determined as <1-5,600 ng/g lipid. No clear temporal 1593 1594 trend was observed in liver samples, nor was a significant relationship found between fetal age and 1595 EXAMPLE 2 STREET S 1596 correlation. The results of the study indicate that HBCDDs are already present at measurable 1597 concentrations in developing fetuses from as early as 6.5 weeks.

1598 Sahlström et al. (2015a) investigated the feasibility of using faeces as a noninvasive matrix to estimate 1599 serum concentrations of BFRs in toddlers for biomonitoring purposes. In their study, faeces samples 1600 from 22 toddlers (11–15-month old) were analysed for HBCDDs and other BFRs, and the results were 1601 compared with previously analysed matched serum samples. The detection frequency for  $\alpha$ -,  $\beta$ - and  $\gamma$ -1602 HBCDD in the faeces was reported to be 100%, 61% and 74%, respectively. The medians for the three 1603 stereoisomers were 0.30 ng/g lipid (range: 0.093-8.1), 0.11 ng/g lipid (range < 0.055-5.8), and 0.18 1604 ng/g lipid (range: <0.083–45), respectively. In the matched toddler serum samples HBCDDs could not 1605 be detected. The authors speculated that the absence of HBCDDs in the serum but high detection 1606 frequency in the faeces of the toddlers could be an indication of low uptake and/or fast excretion 1607 following the uptake from the gut (see **Section 3.1.1.2.4**). It is noteworthy to state that the LC-MS/MS 1608 chromatograms for HBCDDs of the faeces samples showed a further peak, which according to the authors could be  $\delta$ -HBCDD. 1609

1610 Malarvannan et al. (2013b) demonstrated that HBCDDs can be detected in human hair when they 1611 analysed paired human milk and scalp hair samples collected in 2008 from 30 women from the 1612 Philippines. While the median for  $\Sigma$ HBCDDs in human milk was 0.19 ng/g lipid (range: <0.01–0.91), the 1613 corresponding median in scalp hair was 0.93 ng/g hair (range: 0.30–5.4).  $\alpha$ -HBCDD was the major



1614 contributor in both matrices. The authors did not find any association between the levels of HBCDDs1615 measured in human milk and hair samples.

1616 Barghi et al. (2018) analysed human scalp hair for HBCDDs in samples from South Korea and Iran. The 1617 concentrations for  $\Sigma$ HBCDDs in the collected hair samples ranged from <LOD (0.04–0.09 ng/g) to 3.24 1618 ng/g with a mean of 0.46 ng/g hair. As HBCDDs could not be determined in hair samples from Iran, but were present in 13 out of 24 samples in hair samples from South Korea, the results indicate a higher 1619 1620 exposure to these compounds in South Korea. Based on their experiments, the authors concluded that 1621 the HBCDD concentrations in the collected hair samples were mostly coming from endogenous rather 1622 than external exposure, and thus hair can potentially be used as a suitable indicator for monitoring 1623 internal exposure to HBCDDs. Martin et al. (2016) could not detect HBCDDs in human hair from 6 1624 Spanish individuals by LC-MS/MS at an LOD of 11 ng/g hair and an LOQ of 36 ng/g hair.

1625 In summary, the CONTAM Panel concluded that HBCDDs are found in various human tissues and various 1626 concentrations, but reliable correlations and associations between different matrices, which are 1627 important for biomonitoring purposes, are still lacking.

# 1628 **3.1.1.5.** Physiologically Based Kinetic (PBK) modelling

1629 Moreau and Nong (2019) developed a PBK model in rodents and humans. The model includes six 1630 compartments: blood lipoproteins, liver, deep hepatic compartment, adipose tissue, rapidly perfused 1631 tissues and slowly perfused tissues. The model was calibrated with the female C57BL/6 mice data from 1632 Szabo et al. (2011), a toxicokinetic study where mice were orally administered a single dose of 3 mg/kg 1633 of  $\alpha$ -HBCDD (see **Section 3.1.1.1**).

According to the authors, the same study was used for the validation of the model, by using the concentrations found in the liver and the blood. To adapt the mouse PBPK model to humans, the model was scaled to human physiology. The authors performed simulations in human based on the estimations of exposure levels found in the literature. The predicted plasma concentrations were then compared to biomonitoring data from different populations.

According to the authors, the model is '*not ideal for risk assessment for several reasons*'. The same study was used both to calibrate and validate the model. For validation, it is necessary to perform simulations with other independent studies. The validation of the human model was based on estimations of exposure and does not necessarily correspond to the same population where blood concentrations were measured.

- 1644 The CONTAM Panel considers that the model is not valuable for human risk assessment, e.g. the model 1645 was not calibrated in humans (lack of partition coefficients between blood and tissues), and not 1646 validated.
- 1647 3.1.2. **Toxicity in experimental animals**

1648 EFSA CONTAM Panel (2011a) concluded that main targets in sub-chronic and chronic toxicity studies in 1649 rats and mice were the liver, thyroid and the nervous, reproductive and immune systems. This section 1650 provides an overview of the toxicity data described in the previous EFSA Opinion on HBCDDs (EFSA 1651 CONTAM Panel, 2011a) with the new data published since, when available.

Several studies were identified in which the animals were exposed to mixtures of different BFRs including HBCDDs (Ernest et al., 2012; Berger et al., 2014; Lefevre et al., 2016; Tung et al., 2017; Dianati et al., 2017; Allais et al., 2020). The CONTAM Panel did not consider these studies informative for the hazard characterisation of HBCDDs and thus were not considered in the current Opinion.



## 1656 **3.1.2.1. Acute toxicity studies**

Acute toxicity of HBCDDs is very low. The oral lethal dose in rats is >20 g/kg bw and >40 g/kg bw in mice (see EFSA CONTAM Panel, 2011a).

## 1659 **3.1.2.2. Repeated dose toxicity studies**

### 1660 Studies considered in the previous EFSA assessment

1661 In the previous Opinion (EFSA CONTAM Panel, 2011a), several 28 days studies in which HBCDDs were 1662 administered to rats described changes in hepatic weight, drug metabolism, gene expression or function. 1663 A BMDL<sub>20</sub> of 22.9 mg/kg bw per day was reported for increased absolute liver weights and a BMDL<sub>10</sub> of 1664 4.1 mg/kg bw per day was reported for increased glucuronidation of T4 (van der Ven et al., 2006). 1665 Increased liver weights were also indicated in adults and in weanling rats in a 2-generation dietary study 1666 (Ema et al., 2008).

1667 Effects on thyroid were reported in 28-day and 90-day studies, in a 2-generation reproductive toxicity 1668 study and in a developmental toxicity study. The effects observed were increased hyperplasia and 1669 activity of the follicular epithelial cells, increased thyroid weight and follicular cell hypertrophy (Zeller 1670 and Kirsch, 1969, as cited by ECB, 2008; van der Ven et al., 2006; Chengelis, 2001, as cited in ECB, 2008; Ema et al., 2008; Saegusa et al., 2009). The most sensitive effect was increased relative thyroid 1671 1672 weight in female rats in the 28-day study with a BMDL<sub>10</sub> value of 1.6 mg/kg bw per day <sup>26</sup> (van der Ven et al., 2006). However, the CONTAM Panel at that time noted that the dose-response was not clear (at 1673 1674 doses up to 30 mg/kg bw, except at the dose 1 mg/kg bw the increase was less than 6 %) (EFSA 1675 CONTAM Panel, 2011a). The increased relative thyroid weight was accompanied by induction of phase 1676 I and phase II biotransformation enzymes in the same animals (LOEL 3 and 30 mg/kg bw per day in 1677 females and males, respectively) and T4 glucuronidation (BMDL<sub>10</sub> 4.1 mg/kg bw per day  $^{26}$ ; Germer et al., 2006; van der Ven et al., 2006). Endocrine-related effects were reported in several 28-, 90-day and 1678 1679 reproductive toxicity studies, such as changes in thyroid hormone homeostasis (decreased serum T4 1680 level and increased TSH level), inhibition of oogenesis, increased anogenital distance, delayed vaginal 1681 opening, decreased number of ovarian primordial follicles and decreased the fertility index in rats (Zeller and Kirsch, 1969, as cited by ECB, 2008; van der Ven et al., 2006, 2009; Chengelis, 2001, as cited in 1682 1683 ECB, 2008; Ema et al., 2008; Saegusa et al., 2009; see EFSA CONTAM Panel, 2011a). The NOAEL for 1684 reduced fertility index and reduction of the number of ovarian primordial follicles was 10 mg/kg bw per 1685 day (2-generation reproduction toxicity study in rats) and the BMDL<sub>5</sub> for decrease in testes weight was 1686 11.5 mg/kg bw per day (2-generation reproduction toxicity study in rats). The CONTAM Panel concluded 1687 in its previous Opinion that activation of constitutive androstane receptor- (CAR) or pregnane-X-receptor 1688 (PXR)-dependent gene expression was the likely reason for changes in thyroid hormone homeostasis, 1689 and possibly also effects on reproduction, and was considered to be associated with neurodevelopmental 1690 effects on behaviour (EFSA CONTAM Panel, 2011a).

### 1691 Studies published since the previous EFSA assessment

1692 The short-term studies published since the previous EFSA assessment confirm that HBCDDs affect the 1693 liver and the thyroid. In addition, HBCDDs affects sex hormones and lipid and sugar metabolism (see 1694 **Table 9**).

Several studies in rats and mice reported increased liver weight at doses  $\geq$ 20 mg/kg bw per day and lesions at doses of 49.5 µg/kg bw per day (Maranghi et al., 2013; Rasinger et al., 2018; Bernhard et al., 2016; Gannon et al., 2019a).

<sup>&</sup>lt;sup>26</sup> BMD values as calculated and reported by the authors of the study.



In a 28-day dietary study described by Maranghi et al. (2013) and Rasinger et al. (2018), juvenile female 1698 1699 mice were administered HBCDDs (stereoisomers composition not specified) in the diet at doses of 0, 1700 49.5 µg/kg bw per day and 199 mg/kg bw per day. Effects in the liver were increased liver weight at 1701 199 mg/kg bw per day, and lesions (increased vacuolation in hepatocytes, increased pyknotic nuclei, 1702 lymphocytic infiltration and hyperaemic vessels) at both doses. However, no difference was observed 1703 in the incidence of the lesions between the high dose (199 mg/kg bw per day) and the low dose (49.5 1704 µq/kg bw per day) (see Table D.1 in **Appendix D**). These lesions were not considered further due to 1705 the absence of a dose-response relationship. The CONTAM Panel noted that only two doses were tested, 1706 there was a wide range between these two doses (4,000-fold apart), the effects observed at low and 1707 high doses were reported in two different publications, and the effects on certain organs (uterus, 1708 adrenals and brain) were investigated only at the low dose and not at the high dose, precluding the 1709 examination of a dose-response relationship for all the effects. Therefore, the CONTAM Panel did not 1710 use this study for the establishment of a Reference Point.

1711 Benhard et al. (2016) exposed juvenile female mice for 28 days to diets spiked with  $\alpha$ -HBCDD produced 1712 from  $\gamma$ -HBCDD by thermal rearrangement. The doses were 107  $\mu$ g/kg bw per day and 116 mg/kg bw 1713 per day. The control diet contained 46 ng/kg bw per day. In addition, the influence of long-chain omega-1714 3 polyunsaturated fatty acids (LC n3 PUFAs, 4.9 g/kg) on effects induced by  $\alpha$ -HBCDD was examined. 1715 Effects on liver lipid composition and increased body and liver weights were observed at 116 mg/kg bw 1716 per day. There was a marked microvesicular accumulation of lipids at the high dose. These effects were aggravated by LC n3 PUFA intake. Reduction in serum cholesterol occurred at 107 µg/kg bw per day 1717 1718 and reduction of serum triacylglycerol were noted at 116 mg/kg bw per day. At the highest dose, an 1719 increase of serum aspartate aminotransferase (AST) was observed. There was also a decrease in relative 1720 total white adipose tissue weight at high dose. Proteomics of the high dose group indicated effects on 1721 β-oxidation in the liver. The LOAEL was 116 mg/kg bw per day. The CONTAM Panel noted that two 1722 doses were tested, but there was a 1,000-fold range between the doses.

1723 In a 28-day study, female and male F344 rats were fed diets containing HBCDDs at 0, 221, 1,250 and 1724 5,000 mg/kg diet (corresponding to 0, 20, 102 and 430 mg/kg bw per day, and to 0, 19, 94 and 400 mg/kg bw per day, for male and female F344 rats, respectively) (Gannon et al., 2019a). Increased 1725 relative liver weight was reported in females at the two highest doses. No liver lesions were observed. 1726 1727 Increased cholesterol and tryglyceride serum levels were observed in female rats at the two highest 1728 doses (Gannon et al., 2019a). The authors had initially compared several strains of rats, i.e. Sprague-1729 Dawley, Wistar and Fischer F344, to examine strain- and sex related differences in response to exposure 1730 to HBCDDs, and concluded that the F344 strain was the most sensitive to the effects of HBCDDs on 1731 liver and thyroid based on the greatest number of significantly affected endpoints resulting in multiple 1732 health effects. Also that in this strain, sex differences were apparent, especially in tissue concentrations, 1733 number and severity of thyroid and liver lesions, and in immune response parameters, with treatment 1734 eliciting a greater response in males (Gannon et al., 2019a).

1735 In a recent study, male Sprague Dawley rats were exposed by gavage to HBCDDs at 0, 0.06, 0.641, 1736 6.41, 64.1 and 641 mg/kg bw per day for 5 days. Increased liver weight (1.2 times compared to control) 1737 and incidence of centrilobular hepatocyte hypertrophy was observed at 641 mg/kg bw per day. A 1738 significant increase in UDP GT1a1 enzyme was also reported at the two highest doses (Shockley et al., 1739 2020).

1740 Increased thyroid weight, follicular hypertrophy, hyperplasia and/or colloid depletion and vacuolation 1741 were reported in rats after 28 days dietary exposure to HBCDDs (Gannon et al., 2019a). Increased 1742 thyroid weight was observed in males at the lowest dose of about 20 mg/kg bw per day. Desquamation 1743 into follicular lumen, foaming colloid (not statistically significantly increased) and increased ratio of 1744 follicular epithelium areas and number of nuclei were observed in a 28-day study in mice at 49.5 μg/kg



bw per day, but not at the high dose of 199 mg/kg bw per day (Maranghi et al., 2013; Rasinger et al.,
2014, 2018). Therefore, these effects were not considered further.

1747 In the 28-day study in female mice reported by Maranghi et al. (2013) and Rasinger et al. (2014, 2018), 1748 there were statistically significant increased serum testosterone (at 199 mg/kg bw per day) and 1749 decreased serum 17β-oestradiol (at 49.5 µg/kg bw per day) concentrations, and increased ratio 1750 testosterone to 17β-oestradiol (both doses). The relevance of the hormonal changes observed at 49.5 1751 µg/kg bw per day is unclear as no apical endpoints have been investigated at this dose. Reduction in 1752 the number of growing ovarian follicles was also observed in rats at doses ≥20 mg/kg bw per day 1753 (Gannon et al., 2019a). The CONTAM Panel noted that only female mice were tested.

1754 Exposure of male C57BL/6 mice to HBCDDs at a single dose of 0.05 mg/kg bw per week (equivalent to 1755 0.007 mg/kg bw per day) for 31 weeks was associated with hypertrophy and increased relative weight 1756 of epididymal white adipose tissue, with increased lipid accumulation (Xie et al., 2019). The authors 1757 provided compelling evidence that this effect was due to an increased expression of *Pparg* mRNA (see Section 3.1.4). An increased expression of *Pparg* was also observed in liver of male C57BL/6J mice 1758 orally exposed for 14 weeks to an HBCDD dose of 0.035 mg/kg bw per week (equivalent to 5 µg/kg bw 1759 1760 per day) in combination with a high fat diet (containing 330 g lard/kg diet; 506 kcal/kg), which might 1761 explain the observed increased accumulation of liver triglycerides and steatosis (Yanagisawa et al., 1762 2014). The combined treatment of HBCDDs and high fat diet resulted in increased absolute liver weight 1763 and body weight gain compared to controls on normal fat diet (AIN-93M; 360 kcal/kg) and to high fat diet controls (Yanagisawa et al., 2014). The CONTAM Panel noted that both studies administered only 1764 one dose level, the relevance to human health of the effects reported by Xie et al. (2019) is unclear and 1765 1766 for Yanagisawa et al. (2014) the effects were observed in combination with high fat diet.

1767 In summary, HBCDDs affect the liver and lipid and sugar metabolism, and induce endocrine related 1768 effects (thyroid and sex hormone homeostasis).

# 1769 **3.1.2.3. Developmental and reproductive toxicity studies**

## 1770 Studies considered in the previous EFSA assessment

1771 Two dietary reproductive toxicity studies in rats were reported in the previous EFSA assessment (EFSA 1772 CONTAM Panel, 2011a). A NOAEL of 10 mg/kg bw per day was established in the 2-generation toxicity 1773 study based on a reduced fertility index in F0 animals and a reduction of the number of ovarian 1774 primordial follicles in F1 females at 101 mg/kg bw per day (Ema et al., 2008). In the one-generation 1775 study, the most sensitive effect on reproductive organs was a decrease in testes weight (BMDL<sub>5</sub>: 11.5 1776 mg/kg bw per day <sup>26</sup>) (van der Ven et al., 2009). An increased anogenital distance was observed in male pups on PND4 (BMDL<sub>10</sub>: 95.6 mg/kg bw per day Error! Bookmark not defined.) and delayed vaginal 1777 opening (BMDL10: 82.2 mg/kg bw per day Error! Bookmark not defined.) was observed in female pups. The 1778 1779 most sensitive effect observed in this one-generation study was a decrease of the tibia trabecular bone 1780 mineral density in F1 females with a BMDL<sub>10</sub> of 0.056 mg/kg bw per day <sup>26</sup> (van der Ven et al., 2009). 1781 The CONTAM Panel noted at that time the ratio between the BMDL<sub>10</sub> and the BMDU<sub>10</sub> for this effect was 1782 very large (about a factor of 20) indicating a large variation in the dose-response data (EFSA CONTAM 1783 Panel, 2011a). A decrease in epididymal weight was also reported in a 28-day dietary study in rats 1784 (Zeller and Kirsch, 1969, as cited in ECB, 2008).

1785 In developmental toxicity studies in rats, no fetotoxicity, embryotoxicity or teratogenic effects were 1786 reported (Murai et al., 1985; Stump, 1999; see EFSA CONTAM Panel, 2011a). However, an increased 1787 pup mortality during lactation was noted in the F2 generation at 1,008 mg/kg bw per day in the 2-1788 generation reproductive toxicity (Ema et al., 2008).

### 1789 Studies published since the previous EFSA assessment



1790 No developmental and reproductive studies have been identified since.

## 1791 **3.1.2.4. Immunotoxicity studies**

#### 1792 Studies considered in the previous EFSA assessment

1793 In the previous EFSA assessment, effects on the immune system were reported in two studies in rats 1794 (EFSA CONTAM Panel, 2011a). In a 28-day study, reduced splenocyte counts were found (BMDL<sub>20</sub> of 1795 104 mg/kg bw per day <sup>26</sup>) (van der Ven et al., 2006). In a one-generation reproduction study an 1796 increased IgG response was found in male offspring after immunisation with sheep red blood cells 1797 (SRBC) (BMDL<sub>20</sub>: 0.46 mg/kg bw per day <sup>26</sup>) as well as an increase in the fraction of neutrophilic 1798 granulocytes (BMDL<sub>20</sub>: 7.7 mg/kg bw per day <sup>26</sup>) (van der Ven et al., 2009).

### 1799 Studies published since the previous EFSA assessment

In the rodent studies published since the previous EFSA assessment (Maranghi et al., 2013; Rasinger, 2011; Rasinger et al., 2014, 2018; Gannon et al., 2019a) (see **Table 9**), the effects on the immune system were confirmed: decreased thymus weight and increased lesions in the thymus (cortical invasivity or signs of tissue stress), decreased splenocyte proliferation and increased spleen lesions, decreased T-cell and increased B-cell population, increased natural killer (NK) cells, decreased serum IgA or IgG immunoglobulin. These effects were observed at doses around 20 mg/kg bw per day in F344 rats.



## 1807 **Table 9.** Summary of the outcomes of the toxicological studies of HBCDDs in experimental animals on repeated dose

Test compound	Species	Exposure	Outcome	NO(A)EL/LO(A)EL	Reference
			<i>Liver</i> : Increased relative liver weight (high dose). Increased vacuolation in hepatocytes (at both doses), increased pyknotic nuclei (at both doses), lymphocytic infiltration (at both doses) and hyperaemic vessels (at both doses).		
HBCDDs Isomer mixture 'neat' (CAS 3194-55-6) (Promochem) Purity: 99.2% No information on stereoisomer composition	Mice (BALB/c) Juvenile (25 days old at start of the experiment) F	Dietary (fish based), 28 days 10 animals/group Feed ration: 15% (w/w)/kg bw per day 0, 49.5 µg/kg bw/day, 199 mg/kg bw/day Vehicle: DMSO, 0.4 mL/kg feed Control: diet based on casein	Uterus: Increase in incidence of reduction in endometrial glands density and irregular multistratification of the luminal epithelium (low dose). <i>Thymus:</i> Cortical invasivity (high dose) or signs of tissue stress (Hassal's bodies) (low dose). <i>Thyroid</i> : Increased ratio between follicle and colloid areas (at both doses), desquamation into follicular lumen and foaming colloid (low dose), increased ratio of follicular epithelium areas and number of nuclei (low dose). <i>Spleen</i> : Increase lymphocyte hyperplasia (high dose). <i>Hormones:</i> Lower serum concentration of 17β- oestradiol (low dose); increased concentration of testosterone (high dose); and increase ratio testosterone to 17β-oestradiol (at both doses). See further details in <b>Appendix C</b> .	LOEL = 49.5 µg/kg bw/day	Maranghi et al. (2013); Rasinger (2011); Rasinger et al. (2014, 2018). Data from the same animal trial published in three articles and PhD thesis.
HBCDDs Purity: not reported (Sigma-Aldrich) No information on stereoisomer composition	Mice (C57BL/6J) M 5–6 animals/group	0, 1.75, 35, 700 µg/kg bw, once per week by gavage Normal and high-fat diet From 6 to 20 weeks	<i>Metabolic effects</i> : Body and liver weights increases at the two highest doses, along with paralleled increases in blood glucose and insulin levels and microvesicular steatosis and macrophage accumulation in adipose tissue in the high dose group.	Increased body and liver weight in combination with high fat diet: NOEL = 1.75 µg/kg bw/week (or 0.25 µg/kg bw/day)	Yanagisawa et al. (2014)

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				LOEL = 35 µg/kg bw/week (or 5 µg/kg bw/day)	
			Influence of LC n-3 PUFAs on $\alpha$ -HBCDD mediated effects.		
α-HBCDD	Mice (BALB/c)	Dietary, 28 days	The highest dose of $\alpha$ -HBCDD affected liver lipid composition and increased body and liver weight as aggravated by high PUFA intake. The highest dose also reduced thymus weight.	Some effects at the lowest dose (107 µg/kg bw/day) but different	
Produced from γ-	F	46 ng/kg bw (control),		directionality compared	Bernhard et al
HBCDD by thermal rearrangement	3 weeks of age	107 µg/kg bw/day, 116 mg/kg bw/day	The lowest dose slightly reduced body weight gain and liver somatic index. The lowest dose reduced serum cholesterol and the highest dose reduced	with the highest dose (116 mg/kg bw/day).	(2016)
Purity: not reported	8 animals/group	LC n-3 PUFAs: 4.9 g/kg	triacylglycerol. Increased AST serum level was observed at the highest dose and an increase in catalase activity wad noted at the low dose.	LOAEL= 116 mg/kg bw/day	
			Proteomics of the high dose group indicated effects on $\beta$ -oxidation in liver.		
		Dietary, 28 days	Effects on liver, thyroid gland, immune system and endocrine system:		
		0, 250, 1,250, 5,000 mg/kg diet (measured levels in the diet: 0, 221, 1,125, 3,549	Sex differences in metabolism (tissue concentration levels), immune response parameters and in number and severity of thyroid and liver lesions following		
HBCDDs	Rats (F344 M and	mg/kg diet)	exposure to T-HBCDD, with greater responses in M.	LOAEL F344 rats = 19 mg/kg bw/day	
Purity: 95% (Sigma Aldrich)	F, Sprague- Dawley F, Wistar F)	Calculated daily consumption: F F344 rats: 0, 20, 102,	Wistar rats: thyroid (minimal to mild follicular hypertrophy).	LOAEL F Wistar rats = 20 mg/kg bw/day	Gannon et al. (2019a)
1% α-, 1% β-, 98% γ-HBCDD	34–37 days old	430 mg/kg bw/day M F334 rats: 0, 19, 94, 400 mg/kg bw/day	Sprague Dawley rats: liver (increase relative weight), thyroid (mild follicular hypertrophy). Decreased T-cell and increased B-cell population.	LOAEL F Sprague- Dawley rats = 21 mg/kg bw/day	
		F Sprague-Dawley rats: 0, 21, 107, 412 mg/kg bw/day F Wistar rats: 0, 20,	F344 rats: liver (increase relative weight in F); thyroid (increase relative weight in M, mild follicular hypertrophy and colloid depletion); brain (decrease relative weight in F); kidney (mild chronic		
		112, 466 mg/kg bw/day	nephropathy and mild multifocal tubular		



E animala/anam	mineralization in all treated M); spleen (atrophy in
5 animals/group	two high dose M, decrease splenocytes proliferation in
	M and F); ovaries: (significant reduction in the
	number of growing follicles). Decrease in IgG serum
	immunoglobulin in F. Decreased T-cell and increased
	B-cell population, increase in NK-cells in M.
	Changes in multiple haematological parameters in M
	and F at the two highest doses. Changes in clinical
	chemistry parameters (AST and creatine kinase) in all
	strains, more pronounced in F344 rats.
	Decrease serum T4 level at the three dose levels in M
	F344 rats. Increase cholesterol and triglycerides in the
	two high doses in F F344 rats.
	Wistar and Sprague Dawley rats: Concentration of $\gamma$ -
	isomer was more abundant than $\alpha$ -isomer in serum,
	adipose tissues and also in the liver (lower
	concentration); the $\beta$ -isomer was not detected.
	F344 rats: higher $\alpha$ - and $\gamma$ - isomers accumulation in F
	compared to M, except for the $\alpha$ -isomer in serum.

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HBCDDs enriched with $\alpha$ -HBCDD (A- HBCDD) Purity: 95% (Technical HBCDD) (Sigma Aldrich) 81% $\alpha$ -, 7% $\beta$ -, 12% $\gamma$ -HBCDD	Rats (F344) F, M 5 animals/group	Dietary, 28 days 0, 250, 1,250, 5,000 mg/kg diet (measured levels in the diet: 0, 200, 898, 3,938 mg/kg diet) Calculated daily consumption: M: 0, 19, 97, 395 mg/kg bw/day F: 0, 19, 99, 363 mg/kg bw/day	Effects on liver, thyroid gland, immune system and endocrine system: Increased relative liver weight at two high doses in M and F, and thyroid weight at high dose in M and F, decreased thymus and brain weight in two high dose F. Changes in haematological and clinical chemistry parameters in M and F. A-HBCDD residue concentrations: α-isomer was more abundant than the γ-isomer in serum, adipose tissues and also in the liver (lower concentration). Significantly higher A-HBCDD accumulation in F. Histopathological effects: liver (increased incidence and severity zone 3 hepatocellular hypertrophy), thyroid (dose-related increase in incidence and severity of follicular hypertrophy and hyperplasia, colloid depletion), spleen (dose-related increase in lesions: mild to marked PALS atrophy, mild follicular and MZ atrophy), ovaries (increase proportion of growing follicles) Decrease in splenocytes proliferation. Increase in serum immunoglobulin IgA at high dose. Changes in blood lymphocytes populations: decrease in T <sub>H</sub> cells and increase in T <sub>c</sub> cells in high dose F, but no change in T-cell/B-cell ratio; changes in T-cell and B-cell populations at the highest dose in M, decrease in TH cells, T <sub>H</sub> /T <sub>c</sub> ratio and increase in NK and B-cells. Treatment-related effect in thymus lymphocyte subpopulations in M.	NOAEL = 19 mg/kg bw/day	Gannon et al. (2019a)
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HBCDDs Purity: not reported (Sigma-Aldrich) No information on stereoisomer composition	Mice (C57BL/6) M 3–4 weeks of age 6 animals/group	0, 50 μg/kg bw/week For 31 weeks from 3–4 weeks of age	Adipogenic effects. Increased epidymal white adipose tissue weight and hypertrophy.	Increased epidymal white adipose tissue weight and hypertrophy: LOEL = 50 µg/kg bw/week	Xie et al. (2019)
HBCDDs Purity: 96% (Gojira Fine Chemicals) No information on stereoisomer composition	Rat (Harlan Sprague Dawley) M 7-week old 6 animals/group	0, 0.1, 1, 10, 100, 1,000 µM/kg bw per day (corresponding to 0, 0.06, 0.641, 6.41, 64.1, 641 mg/kg bw per day, as reported by the authors) Gavage 5 days	Liver: increased weight (1.2 X) and incidence of centrilobular hepatocyte hypertrophy at HD and significant increase UDP GT1a1 enzyme at two highest doses. Thyroid: no histological changes.	NOAEL = 6.41 mg/kg bw/day	Shockley et al. (2020)

Vehicle: corn oil

AST: aspartate aminotransferase. BMDL: benchmark dose lowest confidence interval. DMSO: Dimethyl sulfoxide. F: female. HBCDDs: hexabromocyclododecanes. IgA: immunoglobulin A. IgG: Immunoglobulin G. M: male. LC n-3 PUFAs: long-chain omega-3 polyunsaturated fatty acids. MZ atrophy: marginal zone atrophy. LOEL: lowest-observed effect. LOAEL: lowest-adverse-effects level. NK-cells: natural killer cells. NOEL: no-observed-effect level. PALS: periarteriolar lymphatic sheaths. T<sub>c</sub> cells: cytotoxic T-cells. T<sub>H</sub> cells: T helper cells. T4: thyroxine. 1808 1809

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## 1813 **3.1.2.5. Neurotoxicity studies**

#### 1814 Studies considered in the previous EFSA assessment

1815 In the previous Opinion, the CONTAM Panel concluded that exposure of rodents to HBCDDs during 1816 development affects the nervous system with subsequent behavioural changes (EFSA CONTAM Panel, 1817 2011a).

Neurodevelopmental effects were described in F1 and F2 offspring of rats receiving 15,000 mg/kg diet HBCDDs (Ema et al., 2008). In a one-generation study in rats, hearing alterations occurred in male rats and changes in haloperidol-induced catalepsy mainly in females (BMDL<sub>5</sub> were 0.2 and 0.9 mg/kg bw per day <sup>26</sup> for increased thresholds in the brainstem auditory evoked potentials (BAEP) and 0.6–4.4 mg/kg bw per day <sup>26</sup> for reduced latencies to movement onset in catalepsy) (Lilienthal et al., 2009).

The critical study selected by the CONTAM Panel at that time involved a single oral administration of 1823 1824 HBCDDs (98% purity, proportions of  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCDD were 3%, 8% and 89%, respectively) via 1825 gavage to 10-day old male and female mice at doses of 0, 0.9 or 13.5 mg/kg bw (Eriksson et al., 2006). 1826 When the mice were three months old, spontaneous behaviour, spatial learning and memory were 1827 assessed and dose-related changes reported. Only male mice were examined. Ten mice per group (from 3-4 litters) were tested for spontaneous behaviour (horizontal locomotion, rearing and total activity) for 1828 1829 three 20-minute intervals of a 60-minute observation period. The HBCDD-exposed mice showed dose-1830 dependent hypoactivity in the first 20 minutes, while they were more active at the end of the observation 1831 period. Groups of 12-17 mice (from 3-4 litters) were assessed for spatial learning and memory in a 1832 Morris swim maze. The mean latencies were longer in mice dosed HBCDDs at 13.5 mg/kg bw but not 1833 at 0.9 mg/kg bw, compared to controls. The NOAEL for spatial learning and memory was 0.9 mg/kg bw. Based on the findings for horizontal locomotion and rearing, the LOAEL in this study was 0.9 mg/kg 1834 1835 bw HBCDDs, the lowest dose tested. The Panel performed dose-response modelling on the results for 1836 horizontal locomotion, rearing and total activity in the first 20-min period. The CONTAM Panel at that 1837 time selected horizontal locomotion and total activity as more reliable parameters than rearing for the analysis. Of these, the lowest BMDL<sub>10</sub> was obtained for horizontal locomotion and this was used by the 1838 1839 then CONTAM Panel as the Reference Point in its previous risk assessment (EFSA CONTAM Panel, 1840 2011a).

Neurobehavioural effects were also reported in reproductive and developmental dietary studies with rats that were reviewed in the previous EFSA assessment (see **Section 3.1.2.3**), but at higher doses than the effects described above. In these studies (Ema et al., 2008; van der Ven et al., 2009), the endpoints measured, and age of observation, differed from those in the study of Eriksson et al. (2006).

### 1845 Studies published since the previous EFSA assessment

Most of the new studies published since the previous EFSA assessment were based on repeated exposures to HBCDDs performed by exposing dams or neonatal rodents at different developmental stages, i.e. GD0-PND21 (Maurice et al., 2015), GD1-PND0 (Miller-Rhodes et al., 2014) and PND10-PND70 (Zhang et al., 2017a), followed by behavioural analyses of the offspring at different ages. One study exposed adult animals for 6 weeks (Pham-Lake et al., 2017) (see **Table 10**).

1851 Only one study provided information on a specific stereoisomer, i.e. alpha-HBCDD (Maurice et al., 2015). 1852 The other studies were conducted with HBCDDs for which the stereoisomer composition was not 1853 reported (Miller-Rhodes et al., 2014; Zhang et al., 2017a).

1854 In Miller-Rhodes et al. (2014), offspring of rats given HBCDDs by gavage from GD1 to PND0 at doses 1855 of 0, 3, 10 or 30 mg/kg bw per day, showed significant neurodevelopmental effects at all doses, but 1856 with no clear dose-response relationships. These effects consisted of increased reactivity to a tailpinch 1857 in neonates, decreased forelimb grip strength in juveniles, impaired sustained attention and loss of



hindleg function, which progressed and increased in severity with age up to 21 months (Miller-Rhodeset al., 2014).

1860 Maurice et al. (2015) found impaired motor activity and reduced anxiety, associated with decreased pup 1861 weight, at 22 ng/kg bw per day in offspring of female rats dosed with hens' egg contaminated with  $\alpha$ -1862 HBCDD by gavage from GD0 to PND21. No effects were reported at a three-fold higher dose of 66 ng/kg 1863 bw per day.

In Zhang et al. (2017a), 10-day old Sprague-Dawley rats (sex not specified, 9 animals/group) were 1864 1865 exposed by gavage to 0, 0.3, 3 and 30 mg/kg bw HBCDDs for 60 consecutive days (from PND10 to 1866 PND70). Using the Morris water maze test, the authors reported a dose-related impaired spatial learning 1867 and memory ability estimated by the increase in the latency time of the animals seeking the platform 1868 (P < 0.05) at 3 and 30 mg/kg bw at all time points (day 3, 4 and 5). This trend was also present for the 1869 0.3 mg/kg bw group, but was significant on day 5, only. In addition, they observed a decrease in the 1870 time remaining in the target quadrant where the platform had been placed and in the number of times 1871 that each rat had crossed the non-exits. Down-regulation of neurotropic factors in the hippocampus in rats dosed with HBCDDs at 3 or 30 mg/kg bw per day HBCDDs was also observed. The CONTAM Panel 1872 1873 noted the lack of information on the protocol, and the poor reporting of the results. For example, the 1874 number of litters, the sex of the animals and the age at observation were not stated. For the last, the 1875 Panel assumed that this was at the end of the 60-days dosing period. The Panel also noted that the 1876 number of animals tested is low, and that the data were very poorly displayed for the Morris water maze (the presentation of the statistics is poor). Therefore, it is difficult to assess whether the increase in 1877 1878 latency is robust or due to chance. Locomotor activity (swim speed) and anxiety were not assessed, 1879 and they can be key confounders of other behaviours. General health condition of the animals was not 1880 reported, e.g. body weight. Learning curves were displayed for only days 3–5. Expression of neurotropic 1881 factors (brain derived neurotropic factor (BDNF), nerve growth factor (NGF) and fibroblast growth factor (FGF)) was investigated. However, the antibodies used for the analysis did not allow for distinction 1882 1883 between the different FGF family members with different function expressed in the brain. Changes in 1884 the expression of these factors were observed at different doses, but no causality can be established. 1885 It is also noted that these data were derived from adult rats (>75 day-old) and thus interpretation 1886 related to early development are difficult to make. Moreover, the litter effect was not investigated. The 1887 Panel concluded that the study shows an effect on neurodevelopment in rats. However, due to the 1888 limitations described above, this study is not considered further for the derivation of a Reference Point.

Pham-Lake et al. (2017) reported that 2-month old male mice dosed with 25 mg/kg bw per day HBCDDs by gavage for 6 weeks, followed by a further 6 weeks without dosing, did not exhibit explicit behavioural abnormalities such as deficits in grooming, isolation, feeding or generalised movement. The absence of these effects does not necessarily contradict those observed by Eriksson et al. (2006) and Zhang et al. (2017a) at an earlier stage of development. They were induced at a different time window and the endpoints evaluated are different.

None of the other new animal neurotoxicity studies with HBCDDs specifically assessed neurobehavioural
endpoints but investigated molecular and cellular effects in the brain (Saegusa et al., 2012; Rasinger et
al., 2014, 2018; Genskow et al., 2015; Pham-Lake et al., 2017; Reffatto et al., 2018) (see Section 3.1.4
and 3.1.5).

The CONTAM Panel re-assessed the study of Eriksson et al. (2006), which had been selected as the critical study in the previous EFSA assessment, and noted several limitations, such as the small number of animals tested (males only) from 3–4 litters, investigation of the litter effect was not reported and the fact that HBCDDs were administered on only one day (PND10). Furthermore, there was a limited range of tests conducted. The mice were tested as adults and key adaptations across adolescence could have altered their behaviour as adults. Anxiety, exploration, motivation and motor ability were not assessed; changes in these parameters could have explained the changes in spontaneous motor



behaviour. However, the CONTAM Panel noted that PND10 does represent the start of a critical period 1906 1907 in the development of the brain in rodents and the observations on spontaneous behaviour (horizontal 1908 locomotion, rearing and total activity) could not be discounted. In the Morris water maze, there were a 1909 limited number of trials, a short trial duration, and only latency to reach the platform was measured. In 1910 addition, swim speed and anxiety were not assessed. Due to these limitations in the assessment of 1911 spatial learning and memory, the CONTAM Panel did not consider this endpoint for the derivation of a 1912 Reference Point. Acknowledging the limitations in the study, the Panel identified a LOAEL of 0.9 mg/kg 1913 bw based on the decrease in spontaneous behaviour (horizontal locomotion, rearing and total activity).

In summary, neurodevelopmental effects were described in offspring of rats given HBCDDs by gavage 1914 from GD1 to PND0 at doses from 3 to 30 mg/kg bw per day (Miller-Rhodes et al. (2014). A dose-1915 dependent decrease in spontaneous behaviour (horizontal locomotion, rearing and total activity) was 1916 1917 noted with a LOAEL of 0.9 mg HBCDDs/kg bw (Eriksson et al., 2006). In addition, impaired spatial 1918 learning and memory were observed in mice exposed by gavage to 13.5 mg/kg bw HBCDDs on PND10 1919 (Eriksson et al., 2006) as well as in rats exposed by gavage to 0.3–30 mg/kg bw per day from PND10 to PND70 (Zhang et al., 2017a). Due to the limitations identified (see above), spatial learning and 1920 1921 memory was not considered for the derivation of a Reference Point.



1922 **Table 10.** Summary of the behavioural studies of HBCDDs in experimental animals

Test compound	Species	Exposure	Outcome	NO(A)EL/LO(A)EL	Reference
HBCDDs (3% $\alpha$ -, 8% β- and 89% γ-HBCDD, as reported in Fång, 2007) Purity: >98% (prepared from a commercial mixture by the research group of Professor Bergman at the Department of Environmental Chemistry, Stockholm University)	– Mice (NMRI) M Neonatal pups	Gavage, at PND10 0, 0.9, 13.5 mg/kg bw (single dose) Vehicle: fat emulsion	Spontaneous behaviour, spatial learning and memory were assessed at age 3 months. At 0.9 and 13.5 mg/kg bw, decreased horizontal locomotion and rearing. Impaired spatial learning and memory (as observed in a Morris water maze) at 13.5 mg/kg bw but not 0.9 mg/kg bw.	LOAEL for spontaneous behaviour = 0.9 mg/kg bw	Eriksson et al. (2006)
HBCDDs Purity: ≥95% (Sigma- Aldrich) No information on stereoisomer composition	Rats (Long- Evans) M, F pregnant 8–11 animals/group	Gavage, from GD1 to PND0 0, 3, 10, 30 mg/kg bw/day Vehicle: corn oil Litters normalised to 8.	<ul> <li>Multiple neurobehavioural tests performed on offspring at different ages up to 21 months.</li> <li>HBCDDs produced age-related effects at 3 mg/kg bw/day (increased reactivity to a tailpinch in neonates, decreased forelimb grip strength in juveniles, impaired sustained attention and progressive loss of hindleg function in aged rats).</li> <li>No effect on locomotor activity.</li> </ul>	No clear dose-response relationships.	Miller-Rhodes et al. (2014)
α-HBCDD Purity: 99.3% (produced by transformation of γ-HBCDD by heat treatment, followed by purification)	Rats (Wistar) F n = 6	Gavage of hen's egg produced by feeding hens with α-HBCDD, from GD0 to PND21 0, 22, 66 ng/kg bw/day Not stated whether litters were normalised	Decreased pup weight at 22 ng/kg bw per day, associated with impaired motor activity and decreased anxiety. No effects at 66 ng/kg bw per day.	No clear dose-response relationships	Maurice et al. (2015)



HBCDDs	Rats (Sprague Dawley) <sup>(b)</sup>	Gavage, from PND10 to PND70	Impaired spatial learning and memory in the Morris water maze test.					
Purity: not reported (Sigma)	Sex not specified	0.3, 3, 30 mg/kg bw/day	Expression of BDNF, NGF and FGF <sup>(c)</sup> in the hippocampus increased at low dose	limitat	identified due t	/hang of a	I.	
No information on stereoisomer composition	10 days of age 9 animals/group	30 mg/kg HBCDD + 300 mg/kg  bw/day taurine	(adaptive) and impaired at higher doses. Taurine appeared to be protective when co- administered.	study	description above)		()	
HBCDDs	Mice (C57BL/6J)	Gavage, 6 weeks	Decreased expression of presynaptic, but not p					
Purity: not reported (Sigma-	М	0, 25 mg/kg bw/day	synaptic, dopaminergic proteins in hypocampus.		LOEL = 25 mg/kg	Pham-Lake	e et al.	
Aldrich) No information on	4 animals/group (control), 6	From 2 months old	No overt behavioural abnormalities (such as defi in grooming, isolation, feeding or generalised movement.		bw/day	(2017)		
stereoisomer composition	animals/group (treatment)	Controls: corn oil						

1923 F: female. M: male. BDNF: brain derived neurotropic factor. NGF: Nerve growth factor. FGF: Fibroblast growth factor. PND: postnatal day. GD: gestational day.

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(a): In the absence of factors for converting concentrations in feed into daily doses for pregnant rats, the default factor of 0.12 for sub-actute studies in rats was used (EFSA, 2012).

(b): The manuscript sometimes refers to mice, which is assumed to be an error.

1926 (c): Expression of specific FGF forms was not investigated.



#### 3.1.2.6. Genotoxicity studies 1927

#### 1928 Studies considered in the previous EFSA assessment

1929 Based on the negative results from the studies reported in EFSA CONTAM Panel (2011a), i.e. a reverse

mutation assay in S. typhimurium, an in vitro chromosomal aberration test in human peripheral blood 1930

1931 lymphocyte, and an *in vivo* micronucleus test in mice after i.p. administration, it was concluded in the 1932

previous assessment that HBCDDs are not genotoxic in vitro and in vivo.

#### 1933 Studies published since the previous EFSA assessment

1934 Since then, the following studies have been published.

Li et al. (2017b) exposed the human breast HBL-100 cell line to HBCDDs at 0, 5, 10 and 50 mg/L for 24 1935 1936 h. The cytotoxicity (as measured by the MTT and LDH assays) was significantly reduced at 50 mg/L 1937 (reduction of 60%). Oxidative stress (measured by reactive oxygen species (ROS) content) was also induced at 50 mg/L. A statistically significant increase in DNA strand breaks (as measured by % tail 1938 DNA) was observed only at the highest and cytotoxic dose (Li et al., 2017b). 1939

1940 No increase of DNA strand breaks was observed in an alkaline Comet assay where human hepatocyte L02 cells (human fetal hepatocyte cell line) were exposed for 24, 48 or 72 h to low concentrations of 1941 1942 HBCDDs ( $10^{-13}$  to  $10^{-7}$  M). However, high concentrations of HBCDDs (20, 40 and  $60 \mu$ M) caused a 1943 significant elevation of DNA strand breaks in a time-dependent manner. High concentrations of HBCDDs 1944 (>20 µM) significantly reduced cell numbers in a concentration and time dependent manner, while lower 1945 concentrations of HBCDDs for 72 h slightly increased cell survival compared with the control. The ROS 1946 level in HBCDD treated cells was elevated in a time-dependent manner. It was also shown that the ROS 1947 level induced by low concentrations of HBCDDs was comparatively lower than that induced by high 1948 concentrations of HBCDDs (An et al., 2013).

High concentration of HBCDDs (50 µM) caused significant increases in DNA strand breaks in L02 cells 1949 1950 (human fetal hepatocyte cell line) as measured by the alkaline Comet assay. Low concentrations of 1951 HBCDDs alone have no obvious effects on the cell viability, while 50 µM HBCDDs significantly suppressed 1952 cell survival. However, a pre-treatment with low concentrations of HBCDDs alleviated the cytotoxicity induced by high concentrations of HBCDDs. When the cells were pre-treated with low concentrations of 1953 1954 HBCDDs for 48 h, the ROS levels induced by subsequent treatment with 50 µM HBCDDs were markedly 1955 lower than those in the cells treated with 50 µM HBCDDs alone. The authors concluded that pre-1956 treatment with low concentrations of HBCDDs induced 'adaptive responses' to high concentrations of 1957 HBCDDs exposure, as evidenced by attenuation of toxicity, of ROS production, and of DNA strand breaks 1958 induction (An et al., 2016).

1959 The CONTAM Panel concluded that this information would not change the previous conclusion that 1960 HBCDDs are not genotoxic in vitro or in vivo. The slight induction of DNA strand breaks observed in 1961 some in vitro tests is most likely due to oxidative stress (see also Section 3.1.4).

#### 3.1.2.7. Carcinogenicity 1962

#### 1963 Studies considered in the previous EFSA assessment

1964 Only one carcinogenicity study was available at the time of the previous EFSA assessment (Kurokawa et al., 1984, as cited in ECB, 2008, EFSA CONTAM Panel, 2011a). In that study mice were exposed to 1965 1966 HBCDDs (information on stereoisomer composition not provided) at 0, 13, 130 or 1,300 mg/kg bw per day in their diet for 18 months. The incidence of liver carcinomas was increased in females, without a 1967 1968 dose-response relationship and the incidences of these tumors were within the background ranges in 1969 this strain of mouse. The CONTAM Panel Panel concluded at that time that "given the lack of genotoxicity



the Panel concluded that carcinogenicity is not a critical effect in the hazard characterisation of HBCDDs"
(EFSA, 2011a).

### 1972 **Studies published since the previous EFSA assessment**

HBCDDs were previously judged not to be carcinogenic in mice and no new carcinogenicity studies havebeen identified. In summary, the available evidence indicates that HBCDDs are not carcinogens.

## 1975 **3.1.3. Observations in humans**

1976 In the previous EFSA Opinion on HBCDDs (EFSA CONTAM Panel, 2011a), two cross-sectional 1977 epidemiological studies on bone mineral density and thyroid stimulating hormone were identified 1978 showing inconclusive results (Weiss et al., 2006; Eggesbø et al., 2011). Since then, 15 additional 1979 epidemiological publications were identified corresponding to 13 study populations assessing the 1980 association between exposure to HBCDDs and any endpoint related to human health (see **Table 11**). 1981 Opting for the totality of the evidence, this section also discusses the two studies already described in 1982 the previous Opinion.

1983 The evidence base includes one cohort study, one birth cohort study (reported in four publications), 1984 and 11 cross sectional studies where the HBCDD exposure was assessed simultaneously or even later 1985 than the endpoint ascertainment. The sample size of the included observational studies ranged from 34 1986 to 71,415 participants. All the evaluated populations came from European countries except for five cross 1987 sectional studies in which populations from the USA (n = 2), China, South Korea and Tanzania were 1988 investigated.

1989 The populations under study were diverse. Four studies recruited younger children or adolescents, while 1990 the remaining studies assessed adult female (n = 6), male (n = 2) or mixed (n = 1) populations. HBCDD 1991 exposure was assessed via serum biomarkers (n = 9), biomarkers in breast milk (n = 1), biomarkers in 1992 adipose tissue (n = 1), HBCDD measurements in dust (n = 1), or through merging dietary patterns and 1993 presence of HBCDDs in food samples (n = 1). Birth weigh/length, neurodevelopment and thyroid 1994 dysfunction were the three endpoint categories used in children. Subfertility, type 2 diabetes, thyroid 1995 hormone levels, severe endometriosis and ovarian endometrioma, and breast cancer metastasis were 1996 the enpoints assessed in the adult populations.

1997 In the following paragraphs, the available evidence base is reported in detail ordered by study design 1998 (cohorts, birth cohorts, cross sectional studies), method of exposure assessment, and year of 1999 publication.

2000 Ongono et al. (2019) reported on the only available observational study conducted on the association 2001 between HBCDDs and type 2 diabetes (T2D); 71,415 middle-aged women were followed for 19 years 2002 in France (3,667 incident T2D cases, 3% attrition). The dietary exposure to HBCDDs was calculated by 2003 merging the usual food consumption over the previous year estimated through a validated 208-item 2004 semi-guantitative dietary questionnaire sent in 1993 and food contamination data available from the 2005 2nd French Total Diet Study (TDS2) published by Anses in 2011. Exposure to PBDEs was also assessed 2006 independently without being included in the analysis for HBCDDs. The dietary exposure to HBCDDs was 2007 calculated for guintiles and these ranged from 0.11–0.35 ng/kg bw per day with a mean of 0.22 ng/kg 2008 bw per day. There was a statistically significant linear association between the dietary exposure per 2009 quintile to HBCDDs and T2D (HR: 1.18; 95% CI: 1.06–1.30 for the 2nd quintile up to HR: 1.47; 95% 2010 CI: 1.29–1.67 for the 5th quintile group). The model was adjusted for several factors including body 2011 mass index (BMI), physical activity, smoking status, level of education. An association with obesity as a 2012 binary endpoint was not investigated. However, due to the fact that BFRs are highly lipophilic and 2013 accumulate in fats, the interaction between HBCDDs (or PBDEs) and BMI was tested; there were no 2014 indications for effect modification of the relation between HBCDDs (ng/kg bw per day) and T2D risk by 2015 BMI ( $\leq 25$  and  $\geq 25$  kg/m<sup>2</sup>) (p<sub>interaction</sub>=0.9).



The Groningen Infant COMPARE <sup>27</sup> (GIC) birth cohort was founded in 2001 in The Netherlands and 2016 2017 consisted of 90 healthy pregnant women, living in the northern provinces of The Netherlands, who delivered a single, full term, healthy infant. HBCDDs were measured through LC/MS-MS in maternal 2018 2019 serum at the 35<sup>th</sup> week of pregnancy and the observed serum levels were relatively low (median, 2020 interquartile range (IQR), range, ng/g lipid; 0.8, 0.47–1.26, 0.3–7.5). Seven additional neutral organohalogen compounds were also assessed (4,4'-DDE, PCB-153, BDE-47, BDE-99, BDE-100, BDE-2021 153 and BDE-154) as well as four phenolic organohalogen compounds (4-hydroxy-2,3,3',4',5-2022 pentachlorobiphenyl (4OH-CB-107), 4-hydroxy-2,2',3,4',5,5'-hexachlorobiphenyl (4OH-CB-146), 4-2023 2024 hydroxy-2,2',3,4',5,5',6-heptachlorobiphenyl (4OH-CB-187) and PCP). Exposure to these compounds 2025 was assessed independently without being included in an adjusted analysis. Four publications reported 2026 on the association between HBCDDs exposure and health-related endpoints in this cohort. Meijer et al. 2027 (2012) addressed the association between prenatal HBCDD exposure and testosterone, free 2028 testosterone, sex hormone-binding globulin (SHBG), LH, FSH, estradiol (E2), free E2 (FE2) and inhibin 2029 B (InhB), at the age of 3 months, and testes volume and penile length at the age of 3 and 18 months. No statistically significant association was found. Ruel et al. (2019) investigated the association between 2030 2031 exposure to organohalogen compounds (including HBCDDs) and child development (mental and motor) at the age of 18 months. No statistically significant association was found. Roze et al. (2009) reported 2032 2033 on an extensive follow-up of the study that assessed neuropsychological functioning including motor 2034 performance (coordination, fine motor skills), cognition (intelligence, visual perception, visuomotor 2035 integration, inhibitory control, verbal memory, and attention), and behaviour at 5–6 years of age. Among 2036 this large number of assessed endpoints, HBCDD exposure (maternal serum) was statistically 2037 significantly positively associated with coordination (rho, 0.29; p value, 0.023), total intelligence (rho, 2038 0.39; p-value, <0.05) and verbal intelligence (rho, 0.48; p value, <0.01). Berghuis et al. (2018) in a 2039 longer follow up reported on endpoints related to cognition and motor performance in adolescents. 2040 Attempting to extend an earlier finding at the 5–6 years follow-up, HBCDD exposure was marginally 2041 statistically significantly and negatively associated with total intelligence (WISC-III-NL <sup>28</sup>; rho -0.355; 2042 95% CI, -5.854, -0.001, p-value 0.05). No statistically significant association was found for performance intelligence, sustained auditory attention, verbal intelligence, auditory-verbal memory, 2043 2044 selective visual attention, or motor endpoint.

2045 In addition to the two cohort studies described in detail above, 11 cross sectional studies (sample size 2046 range: 38–515) were identified, each assessing a different association except for thyroid hormone levels 2047 which were assessed in three studies. The endpoints investigated include breast cancer metastasis (n 2048 = 91, Koual et al., 2019), severe endometriosis (n = 99, Ploteau et al., 2017; Matta et al., 2020), 2049 subfertility (n = 163, Den Hond et al., 2015), congenital hypothyroidism (n = 38, Kim and Oh, 2014), thyroid hormone levels in occupationally exposed adults (n = 80, Li et al., 2014), in pregnant women 2050 (n = 140, Stapleton et al., 2011) and in neonates (n = 193, Eggesbø et al., 2011), sex-related hormones 2051 2052 (n = 62, Johnson et al., 2013), birth weight and birth length (n = 150, Müller et al., 2016), 2053 neurodevelopment in adolescents (n = 515, Kicinski et al., 2012), and bone mineral density (n = 50, Weiss et al., 2006). Of these, statistically significant results were reported for the associations between 2054 2055 lower  $\alpha$ -HBCDD levels in serum of mothers after delivery and the risk of having a child born with congenital hypothyroidism (Kim and Oh, 2014), and between HBCDD levels in house dust from men 2056 2057 recruited through a US infertility clinic and free androgen index and sex hormone binding globulin 2058 (Johnson et al., 2013). Of note, the Müller et al. (2016) study attempted to assess the association 2059 between HBCDDs in breast milk and birth weight and length in Northern Tanzania. HBCDDs were 2060 detected above the LOD in one breast milk samples only and further association analyses were not

<sup>&</sup>lt;sup>27</sup> Comparison of Exposure-Effect Pathways to Improve the Assessment of Human Health Risks of Complex Environmental Mixtures of Organohalogens.

<sup>&</sup>lt;sup>28</sup> The Wechsler Intelligence Scale for Children (WISC) is an intelligence test for children between the ages of 6 and 16. Its output is a Full Scale Intelligence Quotient and five index scores (Verbal Comprehension Index, Visual Spatial Index, Fluid Reasoning Index, Working Memory Index, and Processing Speed Index).



2061 applicable. Stapleton et al. (2011) also attempted to assess the association between HBCDDs ( $\alpha$ -,  $\beta$ -2062 and  $\gamma$ -isomers) and thyroid hormone levels in pregnant women but in none of the samples HBCDDs 2063 were detected above the LOD.

2064 In summary, there is a growing body of epidemiological research in the field of adverse events related 2065 to HBCDDs exposure. The endpoints assessed pertained to type 2 diabetes and neurodevelopment in 2066 the longitudinal studies. In cross-sectional studies the endpoints under consideration (each appraised 2067 in a single study) included breast cancer metastasis, severe endometriosis, subfertility, congenital hypothyroidism, thyroid hormone levels in occupationally exposed adults, in pregnant women, and in 2068 2069 neonates, sex-related hormones, birth weight and birth length, neurodevelopment in adolescents, and 2070 bone mineral density. None of the effects studied in the longitudinal studies assessing internal exposure either reached statistical significance or were replicated in a longer follow-up point in the same study. 2071 2072 The currently available evidence is characterised by relatively small sample sizes in most of the studies, 2073 considerable heterogeneity in the assessed populations, exposures, and endpoints, varying 2074 methodological quality, and effect inconsistency. Moreover, at present no studies were identified with 2075 the same endpoint neither in the pediatric nor in the adult populations. Hence, the ability to confirm the 2076 postulated associations is limited. Moreover, confounding introduced by potential underlying associations with other contaminants was scarcely addressed analytically. 2077



2078 **Table 11.** Summary of epidemiological studies on HBCDDs

Reference	Recruitment year Country	Study design Follow up (years)	N Population group (age) (years) Gender (% males)	Compound Tissue analysed Method (LOD/LOQ)	Levels of Exposure (ng/g lipid) <sup>(a)</sup>	Endpoint health category	Effect 6 (95% 0 value	estimate CI), p-
Ongono et al. (2019)	1990 France	Cohort 19	71,415 Adults (52.87 ± 6.66) 0%	ΣHBCDDs Food samples LC-MS/MS	0.22 ng/kg bw/day (b)	Diabetes type 2	NS; 1.28 NS; 1.39 1.51) NS	
Meijer et al. (2012)	2001–2002 The Netherlands	Birth cohort 0.25, 1.5	34 Infants (0.25, 1.5) 100%	ΣHBCDDs Maternal serum LC-MS/MS (NR)	0.7 (ND-7.4)	Free testosterone	-0.31 (N	IR) NS <sup>(d)</sup>
Ruel et al.	2001–2002	Birth cohort	59 Children (1 5)	ΣHBCDDs	0.02 (0.47, 1.20)	BSID-II-NL	-0.133	(NR) NS <sup>(d)</sup>
(2019)	The Netherlands	1.5	Children (1.5) 63%	Maternal serum LC-MS/MS (NR)	0.82 (0.47–1.26)	MDI	0.004 (NR) NS <sup>(d)</sup>	
						Coordination	0.29 (NI	R) NS <sup>(d)</sup>
		62	ΣHBCDDs		Total intelligence (WISC-III-NL)	0.39 (NI	R) <0.05 <sup>(d)</sup>	
Roze et al. (2009)	2001–2002 The Netherlands	Birth cohort 5–6	Toddlers (5–6) 61%	Maternal serum LC-MS/MS (NR)	0.8 (0.3–7.5)	Verbal intelligence (WISC-III-NL)	0.48 (NI	R) <0.01 <sup>(d)</sup>
						Motor performance and behavior <sup>(e)</sup>	NR (NR)	NS <sup>(d)</sup>
			$\bigcirc$			Performance intelligenc III-NL)	e (WISC-	-0.352 <sup>(d)</sup> (-6.525, 0.029) .052 <sup>(f)</sup>
Berghuis et 2001–2002 al. (2018) The Netherlands	Birth cohort 13–15	31 Adolescents (13-15) 100%	ΣHBCDDs Maternal serum LC-MS/MS (NR)	0.79 (0.46–1.31)	Total intelligence (WISC	C-III-NL)	-0.355 <sup>(d)</sup> (-5.854, -0.001) .050 <sup>(f)</sup>	
						Sustained auditory attention ('Score!', TEA-Ch-NL)		-0.301 <sup>(d)</sup> (-13.739,

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							0.913) .084 (f)
						Neurodevelopment (f)	NR (NR) NS
						Reproductive development (g)	NR (NR) NR
Koual et al. (2019)	2013–2017 France	Cross-sectional NA	91 Adults (64.0, 51.0– 73.0) 0%	α-HBCDD <sup>(h)</sup> Adipose tissue LC-MS/MS (NR)	1.62 (0.97–1.95)	Breast cancer metastasis	NR (NR) 0.18
Ploteau et al. (2017)	2013–2015	Cross-sectional	99 Adults (18-45) 0%	α-HBCDD Adipose tissue LC-MS/MS (NR)	Controls; 0.70, IQR (0.43–1.39)	Severe endometriosis (i)	<b>NR (NR) NS</b> ⑴
Matta et al. (2020)	France		74 Adults (18-45) 0%	α-HBCDD Serum LC-MS/MS (NR)	NA	Severe endometriosis <sup>(i)</sup>	NR (NR) NS <sup>(j)</sup>
den Hond et al. (2015)	NR Belgium	Cross-sectional NA	120 Adults (34.1, 30.0- 38.5) 100%	Total HBCDDs Serum GC-ECNI-MS (NR)	%>LOQ, cases 10 (11.8%), controls 5 (5.3%)	Subfertility	0.65 (0.14– 2.92) 0.57 (j)
				α-HBCDD β-HBCDD γ-HBCDD Serum (Children) LC-MS/MS (0.036)	1.84 ( <mdl-16.6); 0.46 (<mdl-1.3); 14.05 (<mdl- 147.8)</mdl- </mdl-1.3); </mdl-16.6); 	Congenital Hypothyroidism (children)	NR (NR) NS
Kim and Oh (2014)	(im and On NR Cross-sectional II		38 Infants (23–37) NR	β-HBCDD γ-HBCDD Maternal Serum LC-MS/MS (0.036)	0.46 ( <mdl-1.88); 8.86 (<mdl-91.1)< td=""><td>Congenital Hypothyroidism (mothers)</td><td>NR (NR) NS</td></mdl-91.1)<></mdl-1.88); 	Congenital Hypothyroidism (mothers)	NR (NR) NS
			α-HBCDD Maternal Serum LC-MS/MS (0.036)	2.57 ( <mdl-13.8)< td=""><td>Congenital Hypothyroidism (mothers)</td><td>NR (NR) 0.004</td></mdl-13.8)<>	Congenital Hypothyroidism (mothers)	NR (NR) 0.004	
Li et al. (2014)	2009 China	Cross-sectional NA	80 Adults (42, 30–50) 40%	α-HBCDD β-HBCDD γ-HBCDD ΣHBCDD	5.9 (ND-2,702.5)	Thyroid hormones	NR (NR) NS (d)

Serum

0.012) 004



# LC-MS (20, 20, 10)

			62 Adults (18–54) 100%	Total HBCDDs Dust GC-ECNI-MS		Free androgen index	0.46 (NR) 0.004 <sup>(d)</sup>
Jonsson et al. (2013)	2002–2003 USA	Cross-sectional NA			197 (IQR, 107–391)	Sex hormone binding globulin	-0.35 (NR) 0.03 <sup>(d)</sup>
			100 /0	GC-ECNI-M5		Hormones <sup>(k)</sup>	NR (NR) NS
			515 Adolescents (14.9 (0.7)) 53%	Total HBCDDs Serum GC-MS (30ng/L)	<loq, (nd-234)<="" td=""><td>NES-CPT, Reaction Time</td><td>-3.53 (-18.72 - 11.67) NS <sub>(j)</sub></td></loq,>	NES-CPT, Reaction Time	-3.53 (-18.72 - 11.67) NS <sub>(j)</sub>
						NES-CPT, Errors of Omission	27.80% (-17.5% - 97.9%) NS
Kicinski et al. (2012)	2008–2011 Belgium					NES-CPT, Errors of Comission	21.80% (-2.5% to 52.2%) NS
						NES, System, Digit-Symbol test	-0.44 (-6.59 to 5.72) NS <sup>(j)</sup>
						NES, Digit Span test	0.13 (-0.22 to 0.49) NS
Eggesbø et al. (2011)	2003 Norway	Cross-sectional NA	193 Neonates (3 days) 46%	Total HBCDDs Breast milk GC–ECNI-MS (NR)	0.54 (0.10–31)	TSH	0.00 (-0.02, 0.02) NS <sup>(j)</sup>

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Weiss et al. (2006)	2000 Sweden	Cross-sectional NA	50 (25) <sup>(m)</sup> Adults (62, 52–81) 0%	ΣHBCDD α-HBCDD β-HBCDD γ-HBCDD Serum LC-MS/MS (0.12 pg/g; diastereomers 0.03, 0.06, 0.03)	0.46 (<0.24–3.4)	Bone mineral density	NR (NR) NS
Stapleton et al. (2011)	2008–2010 USA	Cross-sectional NA	140 Adults (<20: 23%) 0%	ΣHBCDD α-HBCDD β-HBCDD γ-HBCDD Serum LC-MS/MS (NR)	NA (no detected)	TSH, TT3, fT3, TT4, fT4	NA (NA) NA
Müller et al. (2016)	2012 Tanzania	Cross-sectional NA	150 Neonates (0) 0%	Total HBCDDs Breast milk GC-ECNI-MS (NR)	NA (1 sample above LOQ)	Birth weight Birth length	NA (NA) NA

BSID: Bayley Scales of Infant Development. CI: confidence interval. FU: follow up. GC-ECNI-MS: Gas chromatography-electron capture negative ion-mass spectrometry. HBCDD: hexabromocyclododecane. IQR: interquartile range. LC-MS-MS: liquid chromatography-randem mass spectrometry. LOD: limit of detection. LOQ: limit of quantification. MDI: mental development index. NA: not applicable. ND: not detected. NES-CPT: Neurobehavioral Evaluation System, Continuous Performance test. NS: not statistically significant.

2082 (a): median (range).

2083 (b): Calculated as: Σ[food consumption (g/day) × food contamination (ng/g of food)]/bw (kg); food consumption over the previous year estimated through a validated 208-item semi-quantitative dietary questionnaire sent in 1993; food contamination data extracted from the 2<sup>nd</sup> Anses French Total Diet Study (TDS2) in 2011.

2085 (c): HR for Quintile 2 vs Quintile 1, Qiuntile 3 vs Quintile 1, Quintile 4 vs Quintile 1, Quintile 5 vs Quintile 1, respectively.

2086 (d): rho.

2087 (e): Including motor performance (coordination, fine motor skills), cognition (intelligence, visual perception, visuomotor integration, inhibitory control, verbal memory, and attention), and behaviour.

2089 (f): Including verbal intelligence (WISC-III-NL); auditory-verbal memory (AVLT); selective visual attention ('Sky Search', TEA-Ch-NL); motor outcome (Movement-ABC).

(g): Including testosterone, sex hormone-binding globulin (SHBG); LH, FSH, estradiol (E2), free E2 (FE2) and inhibin B (InhB)], testes volume, penile length were also assessed.

2091 (h):  $\beta$ -HBCDD and  $\gamma$ -HBCDD detection rates were lower than 75% and were excluded by the authors from the statistical analysis.

2092 (i): Ovarian Endometrioma.

2093 (j): OR or beta.

2094 (k): Including follicle stimulating hormone (FSH), serum luteinizing hormone (LH), estradiol, prolactin, free T4, total T3, and thyrotropin (TSH), inhibin B, testosterone; sex hormone binding globulin (SHBG), free androgen index (FAI), free unbound testosterone.

2096 (I): % change.

2097 (m): The PBDE/HBCDD extracts were later pooled (n=25, 125 mL/pool) for stereoisomer determination.



## 2098 **3.1.4.** Mode of action

2099 In the previous EFSA assessment (EFSA CONTAM Panel, 2011a), it was reported that HBCDDs increased 2100 liver weights and induced drug metabolism, mainly CYP2 and CYP3 and phase II conjugating enzymes, 2101 including thyroid hormone conjugating enzymes in the liver. The studies reviewed at the time indicated 2102 that HBCDD toxicity is mediated through activation of PXR and CAR-dependent pathways, with 2103 associated changes in gene expression of biotransformation enzymes with a LOEL of 3 and 30 mg/kg bw per day for female and male rats, respectively (Germer et al., 2006), and increased thyroid hormone 2104 glucuronidation (BMDL<sub>10</sub> 4.1 mg/kg bw per day <sup>26</sup>; van der Ven et al., 2008). These effects on gene 2105 2106 expression and biotransformation in the liver were considered a feasible explanation for changes in 2107 thyroid and sex hormones, with downstream effects on neurodevelopment and reproduction (EFSA, 2108 2011a). It was also indicated that "neurodevelopmental effects might be associated with modulation of 2109 thyroid hormone homeostasis and the involved processes include direct interaction of HBCDDs with 2110 thyroid hormone receptors, and/or perturbations of thyroid hormone transport."

2111 Since the previous EFSA Opinion further studies on the possible mode of action of HBCDDs have been 2112 published (see **Appendix E**) and the most relevant aspects for the risk assessment are described below.

# 2113 3.1.4.1. Hepatotoxicity and metabolic effects

2114 Farmahin et al. (2019) applied RNA sequencing to liver samples from male and female F344 rats exposed 2115 to HBCDDs (11%  $\alpha$ -, 9%  $\beta$ - and 79%  $\gamma$ -HBCDD) at doses of 0, 250, 1,250 and 5,000 mg/kg diet (females: 0, 20, 102 and 430 mg/kg bw per day, males: 0, 19, 94 and 400 mg/kg bw per day) for 28 2116 2117 days. The expression of a total of 428 and 250 gene transcripts was altered by HBCDDs in males and 2118 females, respectively. The number of differentially expressed genes increased with HBCDD dose in both 2119 sexes. The three genes showing the largest change in expression were cytochrome P450 2B1 (Cyp2b1), 2120 nuclear receptor subfamily 1, group D, member 1 (Nr1d1), and metallothionein 1 (Mt1), all showing 2121 dose-dependent upregulation. Gene set enrichment analyses implicated PXR/CAR activation as the 2122 predominantly affected pathway, with genes associated with arylhydrocarbon receptor (AHR) activation 2123 reaching significance in females, only, and no enrichment of PPAR $\alpha$ , DNA damage or cytotoxicity gene 2124 categories. These data suggested that in the rat liver HBCDDs activates the CAR and PXR, and the 2125 upregulation of Mt1 indicates release of cytosolic free  $Zn^{2+}$ , which has also been observed in mouse 2126 brain and neural cells following HBCDDs exposure (Reffatto et al., 2018). The authors suggest that 2127 upregulation of *Mt1* is indicative of oxidative stress. This is a plausible interpretation because oxidative 2128 stress causes release of free Zn<sup>2+</sup> through oxidation or nitrosylation of zinc-thiolate bonds, leading to a 2129 rise in free  $Zn^{2+}$ , activation of metal-responsive factor-1 (MTF1), which induces *Mt1* expression (Chung 2130 et al., 2006; Maret, 2011; Lee, 2018). PXR activation was specifically addressed in human HepG2 cells 2131 and rat H4IIE hepatoma cells, which were stably transfected with a reporter gene driven by PXR 2132 responsive elements (XREMs; Fery et al., 2010). HBCDDs did activate PXR in both human and rat cells, 2133 but only at a relatively high concentration of  $15 \,\mu$ M.

2134 Benchmark Dose Modelling of RNA-sequencing data was carried out by Gannon et al. (2019b) using the 2135 rat liver dataset by Farmahin et al. (2019, see above). The transcriptomic BMD (BMDt) modelling 2136 revealed a bimodal distribution of gene BMDs in both sexes with ranges of 66-104 mg/kg bw per day 2137 in males and 65–94 mg/kg bw per day in females. The most sensitive statistically significant pathway 2138 BMDt was LPS/IL-1 Mediated Inhibition of RXR Function, with BMDt (Benchmark Dose transcriptomics) 2139 of 66 and 71 mg/kg per day for males and females, respectively. To compare with apical effects in the 2140 same experiment, the BMD for increased liver weight in female rats was 99.5 mg/kg bw per day and 2141 the lowest BMD was 15.6 mg/kg bw per day for increased incidence of thyroid colloid depletion in males 2142 (Gannon et al., 2019b).



BMDt was also applied to transcriptomics responses in the liver were also investigated in male Sprague 2143 2144 Dawley rats exposed to 0, 0.064, 0.64, 6.4, 64, and 640 mg/kg bw per day by gavage for five days (n = 6, age not reported; Shockley et al., 2020). There was as dose-dependent increase in gene expression 2145 2146 with doses of 64 and 640 mg/kg bw per day resulting in differential expression (FDR < 0.05) of 23 and 2147 247 genes, respectively. One differentially regulated transcript was found at each of the three lower 2148 doses. In this study, the BMDt was calculated as 38.11 mg/kg bw per day. Pathway analysis of regulated 2149 genes revealed enrichment of liver disease and metabolic pathways, including Cyp2b6, Cyp3a5, Ces2c, 2150 Ugt2a1, and Ugt2b17, which all showed increased expression. There were no changes in serum total 2151 T3 or T4, indicating that in this study an increased expression of biotransformation enzymes in the liver 2152 did not influence thyroid hormone levels.

- 2153 Liver proteomics of HBCDDs exposure was studied in eu- and hypo-thyroid female Wistar rats, generated 2154 through a low iodine diet and sodium perchlorate supplementation, dosed HBCDDs at 0, 3 and 30 mg/kg 2155 bw per day HBCDDs for 7 days (Miller et al., 2016a). Internal concentrations of  $\alpha$ - and  $\beta$ -HBCDD in 2156 adipose tissue were below the LOQ for the 3 mg/kg bw per day groups, whereas accumulation of y-2157 HBCDD (2-3 times) was noted in the highest dose groups. Liver weight was not affected by the seven-2158 day exposure to HBCDDs, but the liver proteome profile was significantly altered with overall changes 2159 in abundance of 13 proteins in euthyroid females and 9 proteins in the hypothyroid animals. The proteins 2160 had mainly mitochondrial cytolocation and were involved in metabolic processes (gluconeogenesis/glycolysis, amino acid metabolism and lipid metabolism) and in oxidative stress 2161 2162 responses in both eu- and hypo-thyroid rats. In contrast, eu- and hypo-thyroid Wistar rats, subjected 2163 to the same treatment regime, showed only very limited alterations of the liver proteome in males only, 2164 without any enriched functions or pathways (Miller et al., 2016b). Also, lower levels of  $\gamma$ -HBCDD 2165 accumulated in white adipose tissue of exposed male rats compared to females.
- 2166 Dietary exposure to  $\alpha$ -HBCDD for 28 days affected whole body lipid metabolism in juvenile female 2167 BALB/c mice, leading to changes in hepatic histology. The effects were aggravated in the presence of a 2168 diet rich in polyunsaturated fatty acids. Hepatic fatty acid profiling and gene expression analysis 2169 indicated that the dietary modulation of the hepatotoxic response to an  $\alpha$ -HBCDD dose of 116 mg/kg bw per day was associated with differential effects on fatty acid β-oxidation (Bernhard et al., 2016). 2170 2171 Oral exposure of male C57BL/6J mice once per week for 14 weeks (starting at 6 weeks of age) doses 2172 of HBCDDs of 0.35 and 0.7 mg/kg bw (equivalent to 0.05 and 0.1 mg/kg bw per day) in combination 2173 with a high fat diet resulted in increased body and liver weight compared to controls on normal diet and 2174 to high fat diet controls (Yanagisawa et al., 2014). These effects were not observed in animals exposed 2175 to HBCDDs in combination with a normal diet. At an HBCDD dose of 0.7 mg/kg bw per week in 2176 combination with high fat diet, there was also an increase in liver triglyceride accumulation and steatosis 2177 along with increased expression of Ppara and Pparg as well as of the PPARy target genes Cd36, Fabp4 2178 and Fsp27, indicating a plausible link to increased lipid droplet formation (Yanagisawa et al., 2014). An 2179 increase in *Pparg* mRNA expression was also observed in mice on a normal diet exposed to HBCDDs, 2180 but only the increase in Cd36 was evident in animals fed high fat diet without HBCDDs. These results 2181 suggest that HBCDDs increase lipid accumulation in tissues of mice fed a high fat diet and that the 2182 effects are linked to increased expression and activity of PPAR $\alpha$  and PPARy. Xie et al. (2019) showed 2183 evidence suggesting that HBCDDs increase expression of *Pparg* and several of its transactivation target 2184 genes in epididymal white adipose tissue (eWAT) through reduced expression of Wnt6. WNT6 inhibits 2185 adipogenesis by suppression of *Pparg* expression via the canonical WNT/ $\beta$ -catenin signalling pathway 2186 (Cawthorn et al., 2012). This effect on adipogenic gene expression was also observed in male C57BL/6 2187 mice (3-4 weeks old at the start of the experiment) exposed to HBCDDs at a weekly oral dose of 0.05 2188 mg/kg bw (equivalent to 0.007 mg/kg bw per day) for 31 weeks with a normal chow diet and was associated with hypertrophy and weight of eWAT (Xie et al., 2019). Using mouse T3-L1 preadipocytes, 2189 2190 it was further shown that HBCDDs reduced nuclearisation of  $\beta$ -catenin, providing mechanistic evidence 2191 for the putative link between reduced *Wnt6* and increased *Pparg* expression (Xie et al., 2019). The



mouse PPARγ was not activated by HBCDDs in a reporter gene assay, suggesting that the PPARγmediated effect of HBCDDs is due to its stimulation of *Pparg* mRNA expression. Experiments on human
preadipocytes from visceral adipose tissue (HPA-V) confirmed that HBCDDs increases lipid accumulation
and expression of *LPL*, but an increased expression of *PPARG* mRNA could not be confirmed in human
cells (Xie et al., 2019).

2197 van den Dungen et al. (2017) found no lipid accumulation in HBCDD-exposed human preadipocytes 2198 (SGBS cells) and bone marrow-derived mesenchymal stem cells (MSC) induced to differentiate into 2199 adipocytes. However, the HBCDDs concentration required to induce lipid accumulation in the study by 2200 Xie et al. (2019) was higher (10  $\mu$ M) than the highest concentration used in this study (1  $\mu$ M) by van 2201 den Dungen et al. (2017). Osteoblastogenesis was induced in MSCs in the presence of 1  $\mu$ M HBCDDs 2202 (van den Dungen et al., 2017). WNT6 is a potent stimulator of MSC fate into osteoblasts (Cawthorn et 2203 al., 2012) and these results, therefore, seem inconsistent with those by Xie et al. (2019) showing 2204 reduced expression of *Wnt6 in vivo* and *in vitro* following HBCDD exposure.

2205 A detailed study on HepG2 cells provided further insight into the mechanisms behind the effects of 2206 HBCDDs on lipid metabolism (Wang et al., 2016a), starting with NMR and following up with hypothesis 2207 driven experiments and analyses. HepG2 cells were exposed to high concentrations of a reagent-grade 2208 HBCDDs formula (purity: >95%) at 50, 1,000 and 10,000 µg/L. Metabolomics analysis using NMR 2209 revealed that HBCDDs exposure resulted in changes in amino acid metabolism, protein biosynthesis, 2210 fatty acid metabolism, and phospholipid metabolism. β-oxidation of long chain fatty acids was 2211 supressed, ATP production reduced, Na<sup>+</sup>/K<sup>+</sup>-ATPase inhibited, and uptake of amino acids and glucose 2212 impaired. Reduced β-oxidation was accompanied by accumulation of free fatty acids and increased 2213 synthesis of phospholipids. The authors suggested that most effects may be a results of reduced ATP production leading to lowered of Na<sup>+</sup>/K<sup>+</sup>-ATPase and Ca<sup>2+</sup>-ATPase activities, resulting in reduced glucose 2214 2215 and amino acid uptake by cells (Wang et al., 2016a). In a related study, Wang et al. (2016b) treated 2216 female CD-1 mice by gavage for 28 days with 10 or 50 mg/kg bw per day of HBCDDs (95% purity; 10% 2217  $\alpha$ -, 10%  $\beta$ -, 80%  $\gamma$ -HBCDD) and used NMR and LC-MS/MS to study changes in the urinary metabolome. 2218 Effects were associated with TCA cycle, lipid metabolism, gut microbial metabolism, and homeostasis 2219 of amino acid. The TCA cycle was concluded to be particularly affected, supporting the results from 2220 studies in HepG2 cells indicating effects on energy metabolism (Wang et al., 2016a,b).

2221 The cellular effects of HBCDDs were investigated on human hepatocyte cell line L02 (An et al., 2013). 2222 Cells were treated with a vast range of HBCDD concentrations  $(10^{-14}-10^{-4} \text{ M})$  for 24, 48 or 72 h. 2223 Treatment with high concentrations (>20 µM) reduced the number of viable cells and increased apoptosis in a dose and time-dependent manner. On the contrary, lower concentrations of HBCDDs (10-2224 2225  $^{13}$ -10<sup>-7</sup> M) for 72 h resulted in increased cell numbers and upregulated the proliferating cell nuclear 2226 antigen (PCNA) protein expression level. High concentrations of HBCDDs also caused significant 2227 decrease of mitochondrial membrane potential in L02 cells. ROS levels, as measured by the DCFH-DA 2228 probe, were also elevated in a concentration and time-dependent manner, with a significant increase observed after 96 h exposure to 10<sup>-13</sup> M of HBCDDs. Intracellular [Ca<sup>2+</sup>] was statistically elevated already 2229 at 10 pM HBCDDs and increased concentration dependently up to the highest concentration of 60 µM. 2230

An increase in liver weight is a robust response observed in many *in vivo* studies with HBCDDs (EFSA CONTAM Panel, 2011a; Maranghi et al., 2013; Yanagisawa et al., 2014; Gannon et al., 2019a; Shockley et al., 2020). Zou et al. (2013) found that nanomolar concentrations of HBCDDs stimulated L02 hepatocyte proliferation in a DNA-dependent protein kinase, catalytic subunit (DNA-PKcs) dependent manner. This was associated with increased NRF2 protein expression and nuclear translocation of NRF2, leading to upregulation of its target gene *HO-1*. They further showed that activation of the NRF2-ARE pathway was dependent on the PI3K/AKT pathway in HBCDD exposed L02 cells.

In an investigation of the ability to induce an adaptive response to HBCDDs, L02 cells were pretreated for 48 h with nanomolar concentrations of HBCDDs followed by exposure for additional 48 h to high



concentration of HBCDDs (50 μM; An et al., 2016). Pretreatment with these low concentrations of HBCDDs protected against the subsequent high concentration of HBCDDs as shown by reduced loss of viable cells and ROS over-production. The compensatory response depended on the activation of the phosphatidylinositide 3-kinase/protein kinase B (PI3K/AKT) pathway, attenuated phosphorylation of adenosine monophosphate-activated kinase (AMPK signaling) and increased phosphorylation of p38 mitogen-activated protein kinases (p38 MAPK pathway) (An et al., 2016).

2246 Cytotoxicity of HBCDDs in two different human hepatoma cell lines, L02 and HepG2, was found to be 2247  $\beta$ -HBCDD >  $\gamma$ -HBCDD >  $\alpha$ -HBCDD (Huang et al., 2016b). HepG2 cells exposed to 0, 10 and 100 nM HBCDDs for 24 or 48 h without effects on viability (Zhong et al., 2015). The lower concentration 2248 2249 stimulated migration and invasion of HepG2 cells, increased abundance of mammalian target of 2250 rapamycin (mTOR) and matrix metalloprotein 9 (MMP9), and suppressed E-cadherin (CDH1) expression. HBCDDs exposure further increased stimulatory phosphorylation of protein kinase B (pPKB/pAKT), 2251 2252 extracellular signal-regulated kinase 32 (pERK), and p-p38 in a dose-dependent manner. Treatment of 2253 PI3K/AKT inhibitors LY294002 and MK-2206 effectively countered the increase of cell migration and 2254 invasion induced by HBCDDs. Together, these results indicate that the enhancement of cell migration 2255 and invasion was through activation of the PI3K/AKT/mTOR signalling pathway (Zhong et al., 2015). 2256 The same pathway also promoted autophagy in L02 cells exposed to 20 µM HBCDDs (Jin et al., 2019).

2257 Thus, studies published since the previous EFSA Opinion on HBCDDs (EFSA CONTAM Panel, 2011a) are 2258 generally consistent with effects on CAR and PXR in the liver of rodents. Several studies report high 2259 sensitivity of lipid metabolism and mitochondrial ATP production to HBCDD with downstream effects on Na<sup>+</sup>/K<sup>+</sup>-ATPase and Ca<sup>2+</sup>-ATPase activities and a rise in intracellular [Ca<sup>2+</sup>]. Effects on mitochondrial 2260 2261 energy production are also consistent with increased ROS production and intracellular Zn<sup>2+</sup> release, 2262 which has been reported in several studies. HBCDD increased adipogenesis in mice by suppressing 2263 WNT/β-catenin signaling, which resulted in increased expression of *Pparg* and its downstream 2264 adipogenic target genes. HBCDDs stimulated proliferation and migration of liver cell lines at picomolar 2265 to nanomolar concentrations and there is evidence that these effects are related to activation of the oestrogen receptor (ER) and the PI3K/AKT/mTOR signalling pathway. 2266

# 2267 **3.1.4.2. Endocrine related effects**

Parallel transcriptomics profiling of mouse brains and two neuronal cell lines (N2A and NSC-19) exposed 2268 2269 to HBCDDs (see Section 3.1.4.2) strongly implicated effects on the hypothalamus-hypophysis-gonadal 2270 axis because the overall most significant predictors of upstream regulators of the observed gene 2271 expression profiles were gonatotropin release hormone, 17β-oestradiol, dihydrotestosterone and 2272 prolactin (Reffatto et al., 2018). The fact that these effects were observed not only in vivo but also in 2273 two different neuronal cell lines is suggesting direct effects on sex steroid target tissue. HBCDDs have 2274 showed both anti-androgenic and anti-oestrogenic activity in CALUX® sex-steroid receptor reporter gene 2275 assays and MCF-7 cell proliferation assays (Hamers et al., 2006; Dorosh et al., 2010; Krivoshiev et al., 2276 2016).

2277 The potential mode of action of reproductive toxicity of HBCDDs (95% pure) was tested in vitro in 2278 Leydig cells of peripubertal (51 days old Wistar) rats during 6 h exposure. Cells were exposed to 0, 1, 2279 2, 5 and 10 µM HBCDDs in the absence or presence of hCG (human chorionic gonadotropin) (0.125, 2280 0.25, 1 and 10 ng/mL). Treatments in the presence of 20  $\mu$ M cholesterol or 20  $\mu$ M 22(R)-2281 hydroxycholesterol were performed to localise the step in the steroidogenic process that is affected by 2282 HBCDDs. Opposite effects of HBCDDs on steroidogenesis were observed in presence or absence of hCG, 2283 in absence of hCG, a concentration dependent increase in androgen and progesterone levels was 2284 observed in culture medium. However, in presence of hCG (1-10 ng/mL), HBCDDs had no effect and in 2285 presence of low hCG (0.125–0.25 ng/mL), HBCDDs significantly decreased androgen production in a 2286 concentration-dependent manner. Inhibition of progesterone was also observed in HBCDD treated cells



2287 in presence of hCG. HBCDDs inhibited human chorionic gonadotropin- and forskolin-supported cAMP 2288 accumulation and steroidogenesis. HBCDDs inhibited also basal cAMP production in presence or absence 2289 of hCG, but elevated basal steroidogenesis. Expression of several cAMP-dependent genes, including 2290 steroidogenic acute regulatory protein, cholesterol side chain cleavage enzyme, and 3-hydroxysteroid 2291 dehydrogenase was also suppressed by HBCDDs, but this was not accompanied by a decrease in 2292 steroidogenic acute regulatory protein expression. HBCDDs caused significant decrease in mitochondrial 2293 membrane potential in untreated and human chorionic gonadotropin-treated cells. The authors 2294 concluded that HBCDDs effect on steroidogenesis in Leydig cells reflect changes in mitochondrial 2295 membrane potential-dependent ATP production, which secondarily reduced cAMP production and basal 2296 and cAMP-regulated cholesterol transport. This in turn facilitates basal but inhibits cAMP-dependent 2297 steroidogenesis (Fa et al., 2013, 2015). Further experiments showed that HBCDD exposure of Leydig 2298 cells caused inhibition in the expression of genes for steroidogenic enzymes, luteinizing hormone 2299 receptor, regulatory and transport proteins, and a decrease in abundance of StAR (Fa et al., 2015). 2300 Enzymatic assays indicated loss of activities of (CYP11A1) and 17β-hydroxysteroid dehydrogenase 2301 (HSD17 $\beta$ ), indicating diminished capacity to synthesise progesterone and testosterone, respectively. 2302 The authors concluded based on these data that conversion of 22-OH cholesterol to pregnenolone and 2303 of progesterone to testosterone may be the sites of HBCDD action on steroidogenesis in Leydig cells 2304 (Fa et al., 2015).

Exposure of human MCF-1 breast cancer-derived cells to HBCDDs at concentrations above 10  $\mu$ M increased proliferation in an oestrogen-dependent manner (Dorosh et al., 2010). Oestrogenic effects of HBCDDs were further assessed by measuring expression of the oestrogen-regulated trefoil factor 1 (*TFF1*) gene with effects starting at 200 nM. Both the cell proliferation and gene expression responses were blocked by the anti-oestrogen, ICI 182,780, suggestive of an oestrogen receptor mediated response.

2311 Changes in thyroid hormone homeostasis have been noted in numerous studies of animals orally 2312 exposed to HBCDDs (see Section 3.1.2.2). Miller et al. (2016a) studied the mechanism of HBCDD-2313 induced thyroid and liver (see above) toxicity. Eu- and hypo-thyroid female Wistar rats were fed 0, 3 2314 and 30 mg/kg bw per day HBCDDs for 7 days. Hypothyroidism was successfully achieved with an 2315 increase in plasma concentration of TSH, a decrease in plasma total T3 and free T3. Exposure over 7 2316 days to HBCDDs did not affect plasma thyroid hormones, TSH, leptin or corticosterone concentrations. However, the increase in TSH concentration was positively correlated with the internal y-HBCDD 2317 2318 concentration in eu-thyroid rats, but not in hypo-thyroid rats. In male rats, subjected to the same 2319 experimental paradigm, corticosterone concentrations increased in unexposed hypothyroid animals and 2320 tended to increase further with HBCDD exposure, however, HBCDD exposure had no effect on serum 2321 leptin concentrations (Miller et al., 2016a).

2322 Thyroid hormones are important for brain development and evidence from two related *in vitro* studies 2323 indicate that HBCDDs might be a direct interruptor of these processes locally in the rat cerebellum. A 2324 very low HBCDDs concentration of 0.1 nM suppressed thyroid hormone-induced dendritic branching of 2325 Purkinje cells in primary cerebellar cultures (Ibhazehiebo et al., 2011a). Thyroid hormone-stimulated 2326 neurite extension of granule cell aggregate cultures was also attenuated by 0.1 nM HBCDDs 2327 (Ibhazehiebo et al., 2011b). Brain derived neurotrophic factor (BDNF) is expressed in response to 2328 thyroid hormone receptor activation and is stimulates granule cell migration and neurite extension. 2329 Addition of BDNF to the culture medium completely rescued HBCDDs inhibition of neurite extension 2330 indicating that the effect might be caused by HBCDDs blocking thyroid hormone-stimulated increase in 2331 BDNF (Ibhazehiebo et al., 2011b). Using a thyroid hormone reporter gene assay it was found that the 2332 same concentration of HBCDDs (0.1 nM) that had effects on dendritic branching and neurite extension 2333 reduced thyroid hormone receptor-mediated transcription (Ibhazehiebo et al., 2011a). These studies 2334 indicate that HBCDDs may block thyroid hormone mediated transcription and associated developmental 2335 processes in the brain.



2336 Extrapolation of effects on thyroid hormone homeostasis observed in rodents to humans is complicated 2337 by differences in levels and binding capacity of transporting proteins and feedback regulation of thyroid 2338 hormone homeostasis. In humans and animals, thyroid hormones circulating in the blood are degraded 2339 by stepwise deiodination, sulfation or glucuronidation. It was shown that sulfation of T3 and T4 in rats 2340 and mice is less than in humans and that glucuronidation predominates in rats (Bartsch et al., 2018). 2341 So, even, if rodents are more sensitive than humans to perturbations of thyroid hormone homeostasis, 2342 rodent data on the effects of HBCDDs on thyroid hormone levels or signalling might be of relevance for 2343 human health risk assessment. Moreover, the human fetus and neonate has a shorter T4 half-life than adults which may make them more vulnerable to a lowering of serum T4 levels (Li et al., 2019a). 2344 2345 Therefore, it is difficult to dismiss a relevance for humans of the thyroid hormones insufficiency observed 2346 in the rat as a result of induction of hepatic metabolism.

- 2347 Effects on insulin and glucose homeostasis have also been observed at low doses of HBCDD exposure. Male C57BL/6J mice were dosed with HBCDDs by gavage for 14 weeks (starting at 6 weeks of age) at 2348 2349 doses of 0.35 and 0.7 mg/kg bw per week (equivalent to 0.05 and 0.1 mg/kg bw per day) in combination 2350 with either a normal or high fat diet (Yanagisawa et al., 2014). The high fat diet by itself did not increase 2351 random blood glucose but did raise random blood insulin levels. Mice exposed to HBCDDs at a dose of 2352 0.7 mg/kg bw per week in combination with the high fat diet had lower insulin sensitivity and higher 2353 random blood glucose levels than the mice on the high fat diet alone, suggesting an effect of HBCDDs. 2354 There were however no clear HBCDD-dependent effects in oral glucose tolerance or insulin sensitivity 2355 tests. Mice on the high fat diet that were exposed to HBCDDs at dose of 0.7 mg/kg bw per week also 2356 showed reduced expression of *Glut4* in epididymal adipose tissue (Yanagisawa et al., 2014).
- 2357 Thus, HBCDDs may interfere with synthesis of sex steroids hormones both by interfering with cAMP-2358 dependent cholesterol uptake and by causing dysregulation of enzymes involved in sex steroid 2359 metabolism. However, there are also evidence that HBCDDs alter the response of tissues to sex steroids. 2360 Cell culture experiments indicate that whilst HBCDD exposure does not remarkably change thyroid 2361 hormone levels, thyroid hormone mediated gene expression and downstream functions in neurons are 2362 exceptionally sensitive to HBCDDs and that this might contribute to the observed effects of HBCDDs on 2363 neural function in animal studies. Relatively low doses of HBCDDs cause glucose dyshomeostasis, with 2364 increases in random blood insulin and glucose, and these effects are exacerbated in combination with 2365 a high fat diet.
- 2366 3.1.4.4. Oxidative stress

Expression of four DNA repair genes was investigated after exposure of human breast cells HBL-100 to HBCDDs at 5, 10 or 50 mg/L (7.8, 16 and 78  $\mu$ M) (Li et al., 2017b). The expression of *ATM*, a key regulator of the DNA repair response, was dose-dependently induced by HBCDD. DNA breaks levels correlated positively with ROS and *ATM*, but a negative correlation was present with *OGG1* and *MTH1*.

Excessive production of ROS has been reported in a variety of studies on HBCDD toxicology. The mechanism(s) involved has/have not been demonstrated but are plausibly related to impairment of mitochondrial energy production as discussed in previous sections.

# 2374 **3.1.4.5. Neurotoxicity**

In its previous Opinion on HBCDDs, the CONTAM Panel concluded that effects of HBCDDs on behaviour may have been caused by changes in thyroid hormone levels, brought about by an increased glucuronidation of thyroid hormone in the liver (EFSA CONTAM Panel, 2011a). An increased T4 UGT activity in the liver was observed in rats with a BMDL<sub>10</sub> of 4.1 mg/kg bw per day <sup>26</sup> (corresponding to a LOAEL of 100 mg/kg bw per day as identified by the CONTAM Panel using ANOVA HSD posthoc test) and this was associated with an increased expression of PXR and CAR-dependent biotransformation



enzymes (Germer et al., 2006; van der Ven et al., 2006). However, effects on thyroid hormone
concentrations and the thyroid gland have only been observed at doses ≥20 mg/kg bw per day, which
is at least an order of magnitude higher than those reported to have effect on neurodevelopment (see **Table 9** and **10**). It is therefore unlikely that neurodevelopmental deficiencies were secondary to
changes in thyroid hormone metabolism and levels.

2386 HBCDDs can pass the blood brain barrier, accumulate in the brain (Covaci et al., 2006; Rasinger et al., 2387 2014), and cause neurotoxicity in juveniles at relatively low doses (Eriksson et al., 2006; Lilienthal et 2388 al., 2009; Zhang et al., 2017a) (see Section 3.1.2 of the current Opinion and EFSA CONTAM Panel, 2389 2011a). Rodents exposed to HBCDDs during early development showed dose-dependent changes in 2390 spontaneous behaviour (hypoactivity followed by hyperactivity in mice, Eriksson et al., 2006), impaired 2391 hippocampus-based spatial memory and learning (in mice and rats, Eriksson et al., 2006; Zhang et al., 2392 2017a). Effects on dopamine-dependent behaviour and auditory evoked potentials were observed in 2393 rats gestationally exposed to HBCDDs (Lilienthal et al., 2009).

2394 Genskow et al. (2015) and Pham-Lake et al. (2017) investigated effects on dopaminergic neurons in the 2395 striatum and hippocampus, respectively, of adult male C57BL/6J mice given HBCDDs at 25 mg/kg bw 2396 per day by gavage for 30 days. In both brain areas they found reduced expression of dopamine 2397 transporters Dat and Wmat2, which are involved in clearing the synapse from dopamine and 2398 reincorporating into presynaptic vesicles. HBCDDs were found to inhibit dopamine uptake with a half maximal inhibitory concentration (IC50) of 4 µM (Mariussen and Fonnum, 2003). In the hippocampus, 2399 2400 there was in addition reduced expression of proteins involved in synthesis (tyrosine hydroxylase) and 2401 degradation (catechol-O-methyltransferase and monoamine oxidase-B) of dopamine (Pham-Lake et al., 2402 2017). These studies are supportive of effects of HBCDDs on pre-synaptic dopaminergic neurons.

2403 Dietary gestational exposure of Sprague-Dawley rats to HBCDDs resulted in an increased number of 2404 mature neurons and apoptotic bodies in the hilus region of the hippocampus in pups (Seagusa et al., 2405 2012). Doses were not reported, but these effects were observed in pups of dams given HBCDDs at 2406 10,000 mg/kg feed and were absent in the two lower HBCDD dietary concentrations of 100 and 1,000 2407 mg/kg feed. Experiments of neuronal cell lines (N2A, NSC19, SH-SY5Y) have confirmed that exposure 2408 to HBCDDs causes cell death by apoptosis at about 1 µM concentration (Reffatto et al., 2018; Shi et al., 2409 2020). Szabo et al. (2017) adopted the experimental paradigm of Eriksson et al. (2006), giving a single 2410 oral dose on PND10 of 3, 10 or 30 mg/kg bw of either  $\alpha$ -HBCDD or  $\beta$ -HBCDD, or 30 mg/kg bw of an 2411 HBCDD mixture. Metabolomic analysis of the serum collected four days after exposure showed dose-2412 dependent reduction in glutamate, which was statistically significant at 10 mg/kg bw.

2413 Rasinger et al. (2014) exposed juvenile Balb/c mice via the diet to HBCDDs at 199 mg/kg bw for 28 days, resulting in an HBCDD accumulation in the brain of 4.7  $\mu$ g/g dw, and reported transcriptomic and 2414 2415 proteomic profiles signifying neurological disease with specific effects on Ca<sup>2+</sup> and Zn<sup>2+</sup> signalling processes. These effects persisted at a lower dose of 49.5  $\mu$ g/kg bw per day from the same experiment, 2416 2417 which were published in a separate report (Rasinger et al., 2018). Similarly, a study carried out in 2418 parallel in brain of female BALB/c mice exposed to 199 mg/kg bw per day of HBCDDs via the diet for 2419 28 days and *in vitro* for 24 h at 1 and 2 µM in two mouse cell lines (N2A and NSC-19) of neuronal origin 2420 implicated Ca<sup>2+</sup> and Zn<sup>2+</sup> dyshomeostasis in glutamatergic neurons as prominent upstream regulators of transcriptomics profiles both in vivo and in vitro (Reffatto et al., 2018). Follow-up experiments in 2421 primary glutamatergic hippocampal neurons showed that HBCDD exposure (1 and 2 µM) inhibited 2422 glutamate-dependent as well as Zn<sup>2+</sup>-evoked intracellular Ca<sup>2+</sup> release (Reffatto et al., 2018). HBCDDs 2423 also triggered intracellular  $Zn^{2+}$  release, an effect which was partially caused by oxidative stress. 2424 2425 Oxidative stress responses were also substantiated by transcriptomics and proteomics profiles (Reffatto 2426 et al., 2018; Rasinger et al., 2018). Both pre- and post-synaptic events are dependent on Ca<sup>2+</sup> signalling 2427 in glutamatergic neurons and many glutamatertic neurons, particularly in the hippocampus, release  $Zn^{2+}$ 2428 along with glutamate to modulate responses by N-methyl-d-aspartate (NMDA) receptors (Sensi et al.,



2429 2011) and AMPA receptors (Kalappa et al., 2015). The NMDA receptor has a GluN2A subunit which has 2430 a high affinity  $Zn^{2+}$ -binding site that mediates allosteric inhibition of post-synaptic Ca<sup>2+</sup> currents (Sensi 2431 et al., 2011). Additionally, in hippocampal neurons,  $Zn^{2+}$  can evoke post-synaptic currents via activation 2432 of GPR39, which is a metabotropic zinc receptor that stimulates Ca<sup>2+</sup> release in cells (Besser et al., 2433 2009).

Intracellular release of Ca<sup>2+</sup> following exposure to HBCDDs has been evidenced in other cell types. 2434 2435 Exposure of PC12 cells to 2 µM HBCDDs inhibited depolarisation-evoked intracellular [Ca<sup>2+</sup>] transients 2436 and associated catecholamine release (Dingemans et al., 2009). Likewise, rat H9C2 cardiomyocyte cells 2437 treated with 2–200 nM HBCDDs showed reduced the cytosolic free Ca<sup>2+</sup> concentrations and increased Ca<sup>2+</sup> accumulation in the sarcoplasmic reticulum in H9C2 rat cardiomyocytes (Wu et al., 2016b). HBCDDs 2438 2439 were also reported to inhibit the sarcoplasmic-endoplasmic reticulum Ca<sup>2+</sup>-ATPase (SERCA) in 2440 neuroblastoma SH-SY5Y cells by preventing its ATP binding with an IC50 of 2.7 µM (Al-Mousa and Michelangeli, 2014). The relative cytotoxicity of HBCDD diastereoisomers in SH-SY5Y cells was  $\beta$ - > y-2441 2442 >  $\alpha$ -HBCDD and this order was mirrored in the potency of the respective diasereoisomer in generating 2443 ROS (Shi et al., 2019).

2444 Ibhazehiebo et al. (2011a,b) found that an exceptionally low HBCDD concentration of 0.1 nM was 2445 sufficient to supress thyroid hormone-induced dendrite branching of Purkinje cells and extension of 2446 granule cell aggregate in primary cerebellar cultures from rat neonates. As these are endocrine-related 2447 effects, they will be discussed further below in Section 3.1.4.3. Possible thryroid hormone related effects on the nervous system were also indicated in a study in male rats maternally exposed from GD10 2448 to PND20 via dams given HBCDDs at 100, 1,000 or 10,000 mg/kg in the diet (corresponding to 2449 2450 approximately 12, 120 and 1,200 mg/kg bw per day; Fujimoto et al., 2013). Pups sampled at PND20 2451 showed an increased abundance of ret proto-oncogene and vimentin positive cells in the white matter 2452 of the cingulum, suggesting an increased number of immature oligodendrocytes. This could potentially be interpreted as an effect of low thyroid hormone as developmental hypothyroidism has been reported 2453 2454 to produce a similar effect (Fujimoto et al., 2012). No remarcable effects on thyroid hormone levels 2455 were observed in pups following maternal HBCDD exposure, which might mean that the effect is brought 2456 about by a reduced responsiveness of the thyroid hormone receptor (Fujimoto et al., 2013) as observed 2457 in cell culture (Ibhazehiebo et al., 2011a,b).

Thus, mechanistic studies on the effects of HBCDDs on the nervous system support specific effects on dopaminergic and glutamatergic neurons. This appears to include inhibition of dopamine and glutamate reuptake following synaptic release, and deregulation of  $Ca^{2+}$  and  $Zn^{2+}$  which are both integral to the function of glutamate transmission. There is also *in vitro* evidence suggesting that thyroid hormone mediated developmental processes in the brain could be affected by HBCDDs through direct effects on developing neurons on their ability to respond to thyroid hormone.

24643.1.5.Consideration of critical effects and dose-response analysis2465for the human risk assessment

# 2466 **3.1.5.1. Consideration of critical effects**

In its previous Opinion on HBCDDs, the CONTAM Panel noted that most of the toxicological studies were 2467 performed with HBCDD preparations for which the purity and stereoisomer composition was not always 2468 2469 indicated. Therefore, a risk assessment of individual stereoisomers was not possible (EFSA CONTAM 2470 Panel, 2011a). The CONTAM Panel concluded that the main targets for toxicity were the liver, thyroid 2471 hormone homeostasis and the reproductive, nervous and immune systems. The most sensitive 2472 endpoints for HBCDDs were considered to be an increased thyroid weight in female rats from a repeated 2473 dose 28-days study with a BMDL<sub>10</sub> of 1.6 mg/kg by per day as calculated by the authors (van der Ven 2474 et al., 2006), and behavioural effects in male mice from a study with a single HBCDD administration at



2475 PND10 (Eriksson et al., 2006) with a BMDL<sub>10</sub> of 0.93 mg/kg bw calculated by the CONTAM Panel at that 2476 time. The effect on the thyroid was concluded to originate from increased T4 glucuronidation in the 2477 liver, leading to a decrease in blood T4 concentration (van der Ven et al., 2006). Changes in thyroid 2478 hormone homeostasis were considered to be associated with the neurodevelopmental deficiencies 2479 caused by HBCDD exposure (EFSA CONTAM Panel, 2011a). In its 2011 Opinion, the Panel expressed 2480 concerns about the reliability of the BMDL<sub>10</sub> of 1.6 mg/kg bw per day, as the dose-response was not 2481 clear (at doses up to 30 mg/kg bw, except at the dose 1 mg/kg bw the increase was less than 6 %), 2482 and decided to use the BMDL10 of 0.93 mg/kg bw for behavioural effects in male mice (Eriksson et al., 2483 2006) as the Reference Point for the hazard characterisation (see Section 1.3.5 on previous risk 2484 assessments).

2485 The new studies published since then have still in most cases been conducted with HBCDDs with no 2486 information on the stereoisomer composition specified. One study was performed with an HBCDD 2487 mixture enriched with  $\alpha$ -HBCDD (81%) (Gannon et al., 2019a), and two studies were performed with 2488  $\alpha$ -HBCDD alone (Maurice et al., 2015; Bernhard et al., 2016). These studies confirmed that in 28-day 2489 studies HBCDD exposure primarily resulted in liver and endocrine related effects.

2490 In the liver, increased weight and hepatocellular hypertrophy were observed at doses  $\geq$  20 mg/kg bw 2491 per day in 28-day studies in rats and mice (Maranghi et al., 2013; Rasinger et al., 2014, 2018; Bernhard 2492 et al., 2016; Gannon et al., 2019a). These effects could be due to increased adipogenesis due to induced 2493 PPARgamma expression (see Section 3.1.4.1). Liver lesions, such as increased vacuolation in 2494 hepatocytes, increased pyknotic nuclei, lymphocytic infiltration and hyperaemic vessels, were seen at 2495 doses 49.5 µg/kg bw per day and 199 mg/kg bw per day in a 28-day mice study (Rasinger et al., 2018). However, these liver lesions were not considered further because there was no difference in the 2496 2497 incidence of the lesions between these doses varying by three orders of magnitude.

2498 Changes in the thyroid observed in the new studies were increased weight, follicular hypertrophy, 2499 hyperplasia and/or colloid depletion. In contrast to the studies available at the time of the previous 2500 CONTAM Opinion on HBCDDs (EFSA CONTAM Panel, 2011a), newer studies reported effects on the 2501 thyroid only at doses  $\geq$  20 mg/kg bw per day. Since changes in thyroid hormone levels have also only 2502 been observed at doses  $\geq$  20 mg/kg bw per day, changes in the thyroid gland and circulating thyroid 2503 hormone concentrations were not considered for the establishment of a Reference Point.

2504 Changes in testosterone (at a dose of 199 mg/kg bw per day) and oestradiol (not dose-related) 2505 concentrations in mice and reduction in the number of growing ovarian follicles in rats (at doses  $\geq$  20 2506 mg/kg bw per day) were also observed.

2507 Two studies in male C57BL/6 mice exposed to HBCDDs by oral gavage once per week provided evidence 2508 of effects on lipid and sugar metabolism at low doses (Yanagisawa et al., 2014; Xie et al., 2019). 2509 Yanigasawa et al. (2014) found dose-dependent increases in body and liver weights at HBCDD doses of 2510 0.035 and 0.70 mg/kg bw per week (5 and 100  $\mu$ g/kg bw per day) when given in combination with a high fat diet. There were also increases in blood glucose and insulin levels and microvesicular steatosis 2511 2512 and macrophage accumulation in adipose tissue in the high dose group, only. However, as these effects 2513 were only observed in combination with a high fat diet, they were not considered relevant for 2514 establishment of a Reference Point. Effects on lipid metabolism were also found in a study exposing the 2515 same mouse strain to a weekly HBCDDs dose of 0.050 mg/kg bw, corresponding to 7.1 µg/kg bw per 2516 day (Xie et al., 2019). Increased epidymal white adipose tissue weight and hypertrophy was observed 2517 and mechanistic evidence suggested that the adipogenic effects in mice was due to increased expression 2518 of Pparg mRNA. However, whilst HBCDD-induced lipid accumulation in human preadipocytes, the 2519 adipogenic effect appeared less potent than that in mice and did not involve stimulation of PPARG-2520 mRNA expression (Xie et al., 2019). The CONTAM Panel noted that both studies (Yanagisawa et al., 2521 2014; Xie et al., 2019) administered only one dose level, that the relevance to human health of the 2522 effects reported by Xie et al. (2019) is unclear, and that for Yanagisawa et al. (2014) the effects were



observed in combination with a high fat diet. Because of these considerations, these studies were notconsidered further for the derivation of a Reference Point.

2525 No new developmental and reproductive study has been identified since the previous EFSA assessment. In the previous assessment (EFSA, 2011a), several reproduction toxicity studies in rats reported reduced 2526 2527 fertility index (at 115 mg/kg bw per day), reduction of the number of ovarian primordial follicles (138 2528 mg/kg bw per day), decrease in testes weight (BMDL<sub>5</sub> of 11.5 mg/kg bw per day  $^{26}$ ), increased 2529 anogenital distance in male pups (BMDL<sub>10</sub> PND4: 95.6 mg/kg bw per day <sup>26</sup>), and delayed vaginal 2530 opening (BMDL<sub>10</sub>: 82.2 mg/kg bw per day <sup>26</sup>) (Ema et al., 2008; van der Ven et al., 2009). No fetotoxicity, 2531 embryotoxicity or teratogenic effects were reported in developmental toxicity studies in rats (up to dietary doses of 750 mg/kg bw per day from GD0-20 or up to 1,000 mg/kg bw per day by gavage from 2532 2533 GD6-9) (Murai et al., 1985; Stump, 1999; EFSA CONTAM Panel, 2011a). Increased pup mortality during 2534 lactation was noted in the F2 generation at 1,008 mg/kg bw per day in males and at 1,363 mg/kg bw per day in females) (Ema et al., 2008). 2535

Regarding the immune system, the rodent studies published since the previous EFSA assessment confirm the effects, such as decreased weight and increased lesions in the thymus, decreased splenocytes proliferation and increased spleen lesions, decreased T-cell and increased B-cell populations, increased NK-cells, decreased serum IgA or IgG immunoglobulin (see **Section 3.1.2.4**). These effects were observed at doses around 20 mg/kg bw per day in F344 rats.

2541 In the previous EFSA assessment (EFSA CONTAM Panel, 2011a), several studies in rodents were 2542 identified demonstrating that exposure to HBCDDs induced neurodevelopmental effects on behaviour 2543 after single (Eriksson et al., 2006) or repeated administration (Ema et al., 2008; Lilienthal et al., 2009; 2544 Saegusa et al., 2009). The study by Eriksson et al. (2006), in which HBCDDs (3%  $\alpha$ -, 8%  $\beta$ - and 89% 2545 y-HBCDD) were administered as a single dose by gavage on PND10 at 0.9 or 13.5 mg/kg bw, provided 2546 the lowest doses leading to neurobehavioural effects. PND10 was selected by Eriksson et al. (2006) as 2547 the optimal day for administering the HBCDDs because it represents a critical period in the development 2548 of the rodent brain, coinciding with a period of rapid brain growth spanning the first 3-4 weeks of 2549 postnatal life and reaching its peak at around day 10 (Eriksson et al., 1992, 2000; Ahlbom et al., 1995; 2550 Eriksson and Fredrikson, 1998; Davison and Dobbing, 1968; Gaitonde and Richter, 1966). They stated 2551 that in humans this period begins in the third trimester of pregnancy and continues throughout the first 2552 2 years of life. A dose-dependent decrease in spontaneous behaviour (horizontal locomotion, rearing 2553 and total activity) was noted (LOAEL of 0.9 mg/kg bw). Impaired spatial learning and memory were 2554 observed in mice exposed to 13.5 mg/kg bw HBCDDs (NOAEL of 0.9 mg/kg bw). In its 2011 Opinion, 2555 the CONTAM Panel had noted limitations and concerns regarding the single administration protocol and 2556 that the litter effect was not taken into account properly, and recommended that the results required 2557 independent verification (EFSA CONTAM Panel, 2011a). However, the CONTAM Panel also noted at that 2558 time that PND10 covered a relevant neurodevelopmental period in experimental animals, and that it 2559 provided the lowest dose leading to neurobehavioural effects. Thus, in its previous assessment the 2560 CONTAM Panel decided to perform dose-response modelling on the results obtained in this study, and used the lowest and most reliable BMDL (for horizontal locomotion) as the Reference Point for the 2561 2562 hazard characterisation of HBCDDs (EFSA CONTAM Panel, 2011a).

2563 Of the new studies published since the previous assessment on effects on the nervous system, most 2564 did not investigate neurobehavioural effects but provided cellular and molecular support for effects on 2565 glutamatergic and dopaminergic neurons in the hippocampus and striatum (Saegusa et al., 2012; 2566 Rasinger et al., 2014, 2018; Genskow et al., 2015; Pham-Lake et al., 2017; Reffato et al., 2018). Only 2567 one study provided information on  $\alpha$ -HBCDD (Maurice et al., 2015), showing significant effects at a very 2568 low dose of 22 ng/kg bw per day, but not at the higher dose of 66 ng/kg bw per day for motor activity 2569 and anxiety in offspring of female rats dosed by gavage from GD0 to PND21. Miller-Rhodes et al. (2014) 2570 administered HBCDDs to pregnant rats and found significant changes at all doses but no clear dose-



response relationship in multiple neurobehavioural tests performed on the offspring at ages up to 21 months. The study of Zhang et al. (2017a) investigated the effects of HBCDDs (0.3, 3 and 30 mg/kg bw per day) on spatial learning and memory after exposure of rats from PND10 to PND70. Effects were observed at all doses. This study is supportive of the Eriksson et al. (2006) study in that HBCDDs induced neurobehavioral effects, but it was not considered further for the derivation of a Reference Point due to limitations in the study.

2577 The current assessment highlighted additional limitations in the study of Eriksson et al. (2006), including 2578 possible alternative explanations/drivers for the observed changes in spontaneous behaviour that were 2579 not assessed, and that the mice were only tested when adults so key adaptations across adolescence, 2580 for example, could have altered their behaviour as adults. However, the CONTAM Panel concluded that 2581 these results could not be discounted, and the LOAEL was determined to be 0.9 mg/kg bw, based on 2582 changes in spontaneous behaviour (horizontal locomotion, rearing and total activity). Due to limitations in the assessment of spatial learning and memory, this endpoint is not considered suiltable for the 2583 2584 establishment of a Reference Point in the current assessment.

The new *in vitro* genotoxicity studies available do not change the previous conclusion that HBCDDs are not genotoxic *in vitro* or *in vivo*. The slight induction of DNA strand breaks observed in some *in vitro* tests is most likely due to oxidative stress.

HBCDDs were previously judged not to be carcinogenic in mice and no new carcinogenicity studies have been identified. In summary, the available evidence indicates that HBCDDs are not carcinogens.

2590 Concerning effects in humans, since the previous EFSA assessment fifteen new epidemiological 2591 publications were identified assessing the association between exposure to HBCDDs and several human 2592 health endpoints. Neurodevelopment and thyroid dysfunction were assessed in children, while in adults, 2593 subfertility, type 2 diabetes, severe endometriosis and ovarian endometrioma and breast cancer 2594 metastasis were assessed. In the single available prospective study using measures of internal exposure, 2595 statistically significant - yet opposite-direction - results were reported for endpoints related to 2596 neurodevelopment (coordination, and WISC-III total and verbal intelligence). No concordant findings 2597 were identified in the single available cross-sectional study on the same research question (Kicinski et 2598 al., 2012). In another large longitudinal study in a European population assessing the association 2599 between HBCDDs and type 2 diabetes, statistically significant results were reported but there was no 2600 assessment of internal exposure (Ongono et al., 2019). Among the available cross-sectional studies, 2601 statistical significance was recorded for the associations between lower  $\alpha$ -HBCDD concentrations and 2602 the risk of having a child born with congenital hypothyroidism (Kim and Oh, 2014), and between HBCDD 2603 levels and free androgen index and sex hormone binding globulin (Johnson et al., 2013). Although the 2604 number of epidemiological studies has grown in the last years, limitations related to exposure 2605 assessment, study design, sample size, effects direction and lack of validity hinder the possibility to 2606 consider any of the assessed endpoints as the basis for the risk assessment.

Neurodevelopmental effects on behavior are supported by mechanistic studies with HBCDDs, indicating specific effects on dopaminergic and glutamatergic neurons, which could be the mode of action of the neurobehavioural effects. These include inhibition of dopamine and glutamate reuptake following synaptic release, and deregulation of Ca<sup>2+</sup> and Zn<sup>2+</sup> which are both integral to the function of glutamate transmission. There is also *in vitro* evidence suggesting that thyroid hormone mediated developmental processes in the brain could be affected by HBCDDs through direct effects on developing neurons in their response to thyroid hormones (see **Section 3.1.4.2**).

2614 Overall, the CONTAM Panel concluded that the neurodevelopmental effects on behaviour still can be 2615 considered the critical effect for the risk characterisation.



#### 2616 **3.1.5.2. Dose-response analysis**

2617 Since the evidence from the available human data was not sufficient to base the risk assessment on, 2618 the CONTAM Panel considered the data from studies on experimental animals to identify Reference 2619 Points for the human hazard characterisation.

The Panel performed BMD modelling according to the EFSA Guidance on the use of the BMD approach in risk assessment (EFSA SC, 2017) for the data on horizontal locomotion, rearing and total activity in mice from Eriksson et al. (2006). Since this study was performed with a mixture of HBCDD stereoisomers, a dose-response analysis for individual stereoisomers was not possible. The results of the modelling are summarised in **Table 12** and details of the BMD analysis are reported in **Annex C**.

Table 12. Benchmark dose (BMD) modelling for the effects of HBCDDs in Eriksson et al. (2006) with a
 benchmark dose response (BMR) of 10%. Detailed results of the BMD modelling can be found in Annex
 C.

	Madal	BMDL10	BMDU <sub>10</sub>	Reference
Critical effect	Model	(mg/kg b		
l levinentel le cometion	Model averaging	0.08	2.09	
Horizontal locomotion	Lowest	0.05	2.15	_
Desular	Model averaging	0.09	0.76	Eriksson et al.
Rearing	Lowest	0.06	0.75	(2006)
Tatal activity	Model averaging	0.58	8.41	_
Total activity	Lowest	0.51	12.7	-

2628

2629 In its previous Opinion on HBCDDs (EFSA CONTAM Panel, 2011a), the CONTAM Panel performed dose-2630 response analysis and the calculation of the BMD and the BMDL of the Eriksson et al. (2006) study were based on the previous EFSA guidance on the use of the BMD approach in risk assessment (EFSA SC, 2631 2009). Re-analysis of the Eriksson et al. (2006) data with the new EFSA guidance for BMD modelling 2632 2633 (EFSA SC, 2017), led to wider intervals around the BMD. This is mainly due to differences in methods 2634 recommended by the two guidance documents. In the current guidance, for continuous data only four models are used, and only two models per nested family (Hill and exponential). Moreover, the new BMD 2635 guidance (EFSA SC, 2017) does not recommend constraining the steepness/shape parameter in the 2636 models. Therefore, if the shape of the dose-response curve is not sufficiently constrained by the data 2637 itself in the region of the BMR (e.g. due to the low number of dose groups, and/or the dose spacing, 2638 2639 and/or limited sample size) a large BMD confidence interval can result as a consequence. The CONTAM 2640 Panel noted that the BMDLs for horizontal locomotion and total activity are far below the lowest dose administered. Overall, the Panel decided to identify the Reference Point based on the NOAEL/LOAEL 2641 2642 approach instead of BMD modelling. From the Eriksson et al. (2006) study, a LOAEL of 0.9 mg/kg bw 2643 was identified for spontaneous behaviour.

2644 Daily exposures to HBCDDs result in increasing levels in the body. For this reason, the accumulated 2645 concentrations in the body or body burden, rather than the daily exposure, should be considered as the 2646 proper starting point for the risk assessment. In mice (Eriksson et al., 2006), the body burden was 2647 calculated assuming an oral absorption of 83%. This value corresponds to the calculated absorption of 2648 the  $\gamma$ -stereoisomer in mice (Szabo et al., 2010, 2011a) (See **Section 3.1.1.1**). The steady state body 2649 burden is usually estimated in mice and human following the equation:

2650 body burden =  $(F_{abs} \times dose)/K_{el}$ 

2651 Where,

2652dose = daily dose applied (mg/kg bw per day), NOAEL or chronic human dietary2653 $K_{el}$  = elimination rate [In(2)/( $T_{1/2}$  in days)] (1/days)



2654  $F_{abs}$  = fraction of the chemical absorbed into the body

For mice, the body burden was not calculated at steady state, but at PND10, i.e. the first day of HBCDDs administration and corresponding to the start of a critical period in the development of the rodent brain (see **Section 3.1.5.1**). Consequently, the CONTAM Panel decided not to apply the elimination rate in the calculation of the mouse body burden.

Taking the LOAEL of 0.9 mg/kg bw per day for spontaneous behaviour and adjusting by the mouse oral bioavailability (83%) of  $\gamma$ -HBCDD (Szabo et al., 2010, 2011a), the body burden in mice at the LOAEL is estimated to be 0.747 mg/kg bw.

- The chronic human dietary intake corresponding to the calculated body burden at the LOAEL (0.747 mg/kg bw) in the mice is calculated from the same equation:
- 2664 Chronic human dietary intake = (body burden  $\times$  K<sub>el</sub>) / F<sub>abs</sub>

In the absence of robust information, the Panel assumed the human absorption of HBCDDs ( $F_{abs}$ ) to be 100%. Due to the uncertainties in the method to estimate the half-life in Geyer et al. (2004, extended abstract, see **Section 3.1.1.2** and **3.5.4**), the worst-case longest half-life for HBCDDs of 219 days was used ( $K_{el} = 0.00315 \text{ days}^{-1}$ ). Considering these values, the chronic human dietary intake would be:

2669 Chronic human dietary intake =  $0.747 \text{ mg/kg bw } \times 0.00315 \text{ days}^{-1} = 2.35 \mu\text{g/kg bw per day}$  (or 2670 0.00235 mg/kg bw per day)

# 3.1.5.3. Derivation of a health-based guidance value or margin of exposure approach

The CONTAM Panel concluded that the derivation of a health-based guidance value is not appropriate due to limitations in the current database on HBCDDs, including in most studies the lack of information on the stereoisomer composition of the HBCDD mixture tested. Repeated dose reproductive toxicity studies, that include a functional observational battery, showed only sporadic effects.

Instead the margin of exposure (MOE) approach was used for the risk characterisation by comparing the calculated chronic human dietary intake leading to a body burden at the LOAEL of 0.9 mg/kg bw in the mouse study of Eriksson et al. (2006) (see **Section 3.1.5.2**) with the estimated dietary exposure reported in **Section 3.3.1**.

Usually a MOE of 100, covering uncertainties and variability with respect to kinetic and dynamic differences between animal species and humans (factor  $4 \times 2.5 = 10$ ) and within the human population (factor  $3.2 \times 3.2 = 10$ ), is considered sufficient to conclude that there is no health concern.

Since the MOE approach for HBCDDs is based on a body burden comparison between animals and humans, the potential toxicokinetic differences can be considered as sufficiently covered. Therefore, the calculated MOE should be sufficient to cover interspecies differences in dynamics for the effects observed (a **factor 2.5**).

Considering the assumption of 100% absorption of HBCDDs in humans, and use of the worst-case maximum half-life estimated in humans of 219 days, the CONTAM Panel concluded that it was not necessary to apply an uncertainty factor to cover individual differences in kinetics. The Panel recognised that there could be differences in the individual susceptibility in the sub-population of infants and children. Therefore, the Panel concluded that the MOE should also cover individual differences in dynamics (a **factor 3.2**).

The Panel applied an additional uncertainty **factor of 3** to extrapolate from a LOAEL to a NOAEL. This default factor is considered sufficiently conservative (ECHA, 2012; EFSA SC, 2012).



The Panel considered whether an additional factor should be applied to allow for limitations in the database. It was noted that repeated dose reproductive toxicity studies, that include a functional observational battery, showed only sporadic effects. Carcinogenicity was studied only in mice. Based on the mode of action and genotoxicity findings (see **Section 3.1.2.6** and **Section 3.1.4**), the CONTAM Panel considered that carcinogenicity is unlikely to be a critical effect. Therefore, it was concluded that an additional uncertainty factor for limitations in the database is not needed.

Thus, the Panel concluded that an MOE higher than **24** ( $2.5 \times 3.2 \times 3$ ) would indicate a low health concern.

- 2704 **3.2.** Occurrence data
- 2705 3.2.1. Occurrence data submitted to EFSA

An initial number of 11,770 analytical results (5,122 samples) on HBCDDs in food were available in the EFSA database. Data were reported by 11 European countries. The analytical results were obtained between 2000 and 2018. Because the previous Opinion included data collected up to 2010, and in order to have a data set representing current occurrence levels while retaining a sufficiently high number of samples, the present assessment used data submitted to EFSA between 2010 and 2018.

The occurrence data were carefully evaluated, and a number of validation steps were applied before being used to estimate the dietary exposure (see **Annex B**, Table B.1 for further details). The resulting dataset included a total of 7,593 analytical results (2,530 samples) on HBCDDs ( $\alpha$ -HBCDD,  $\beta$ -HBCDD,  $\gamma$ -HBCDD and Total HBCDDs) in food.

- 2715 Data providers were contacted to clarify inconsistencies identified during the data check. The following 2716 modifications were made to the initial data set based on the feedback received:
- The product description of several records allowed a more accurate FoodEx classification. In these cases, the samples were reclassified to a more specific, lower level.
- 2719 Special attention was given to the analytical method reported. There were samples on Total • 2720 HBCDDs which were reported being analysed by LC/MS. After clarifying with the data providers, 2721 the analytical method was changed to GC/MS. In addition, there were samples on the individual 2722 stereoisomers with reported analytical method 'unspecified' but containing some information in the analytical method text. These were clarified further with the data providers and the 2723 2724 analytical method was adjusted accordingly. In the final dataset, all samples reported on Total 2725 HBCDDs were clarified to be analysed by GC/MS and the three major HBCDD stereoisomers by 2726 LC/MS.
- 303 occurrence values were reported as 'HBCDDs unspecified'. Since data on the same samples
   for the three major stereoisomers were also reported, it was decided to exclude them.
- 187 samples data were reported on both Total HBCDDs and individual stereoisomers. Checking
   these data in detail, it was clarified that the Total HBCDDs reported was not analysed separately
   but it was the calculated sum of the values provided for the individual stereoisomers. These
   187 occurrence values on Total HBCDDs were excluded from the dataset.
- Given that  $\alpha$ -HBCDD is the predominant stereoisomer in food, three samples that reported only values on  $\beta$ - and  $\gamma$ -stereoisomers, i.e. six occurrence values in total, were excluded from the dataset. On that basis, two samples that reported values only on the  $\alpha$ - isomer were kept and used for the estimation of exposure to HBCDDs.
- Some occurrence values were expressed on fat weight, but the fat content of the food item was not provided. However, the fat content was required for the calculation of the values on whole



weight. For these occurrence values, the median fat content reported for the same foodcategory by other countries was used.

- A high variability was noticed in fat content reported within certain FoodEx categories. For these entries, as well, the same approach of using the median values was followed. One of the affected food categories was the group of 'Salmon and trout (Salmo spp.)'. Since this is a single category at the most detailed level of the FoodEx with substantially different fat content, it was decided to split it whenever possible to identify the species from the product description. This resulted in having two new food categories: 'Salmon' and Trout'.
- For 141 samples, occurrence data were reported only for Total HBCDDs. The majority of these
   samples were left-censored (analytical data below LOD or LOQ) with high LOQs. Therefore,
   these values were excluded from the dataset.
- 99 occurrence values were reported as 'suspect sampling' and were excluded.
- In the final dataset (6,857 occurrence values from 2,287 samples), 67.7% of the data were reported as 'selective sampling', 32.1% as 'objective sampling', while the remaining 0.2% were reported as 'convenient sampling'<sup>29</sup>. It was decided to retain all samples regardless of the sampling strategy.

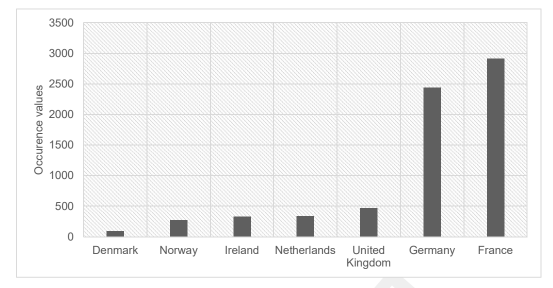
For all samples in the final dataset, values for the three major stereoisomers were available. Therefore, values for the exposure to HBCDDs were calculated as sum of LB/UB of the individual stereoisomers ( $\alpha$ - $\beta$ - and  $\gamma$ -HBCDD).

- As shown in **Figure 2**, occurrence values from the final dataset were reported by seven European countries, most of them by France (42.5%) and Germany (35.6%). The majority of the data (79%) were reported between 2012 and 2016 (**Figure 3**).
- The left-censored data accounted for 71% of the occurrence values. Both LOQ and LOD were provided for 46% of all left-censored data, while only LOD or only LOQ were provided for 26% and 28% of the reported left-censored occurrence values, respectively. Among those, six food categories were 100% left-censored as shown in **Figure 4**. On the other side, the food group with the highest percentage of quantified data was 'Fish and other seafood (including amphibians, reptiles, snails and insects)'.

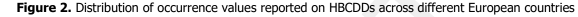
Based on the FoodEx classification, 12 food categories at FoodEx Level 1 were represented (**Figure 4**). After the most represented group, 'Fish and other seafood' with 3,554 occurrence values (1,186 samples) reported, the food group with the highest number of samples was 'Meat and meat products (including edible offal)' with 1,656 occurrence values (552 samples). The food groups 'Eggs and egg products' and 'Milk and dairy products' followed with 666 occurrence values (222 samples) and 645 occurrence values (215 samples) reported, respectively.

<sup>&</sup>lt;sup>29</sup> Definitions on different sampling strategies can be found in the EFSA Guidance on Standard Sample Description (SSD) for Food and Feed (EFSA, 2010a, 2013).

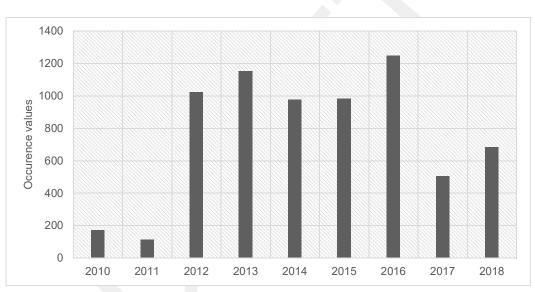




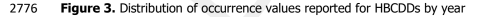




2774



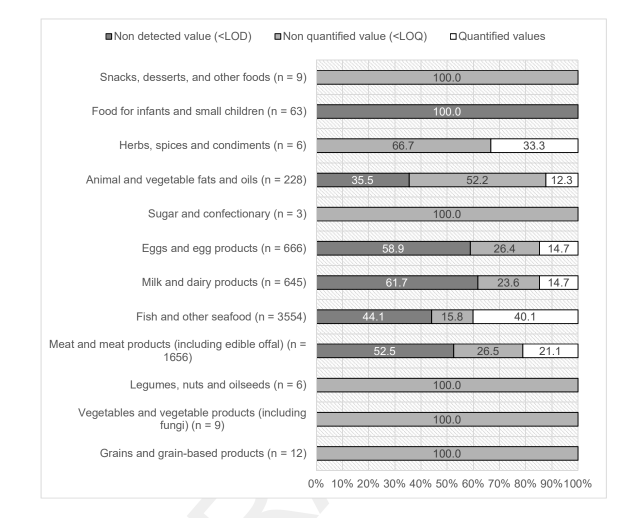
2775



2777 In view of the exposure assessment, food data were grouped at different FoodEx levels, and other 2778 merged categories used as supplements to FoodEx, taking into consideration several factors including

the similarities between food categories, the number of samples, and the concentrations observed.





2780 2781

Figure 4. Percentage of analytical results below LOD, below LOQ and quantified values in the final dataset across the different food categories (FoodEx Level 1)

At the most detailed level (FoodEx Level 3), the food category was retained if more than six samples were available for that category. If less than six samples were available, the levels were compared with similar foods belonging to other categories:

- If the levels were similar, the samples were either grouped together with the similar food, resulting in a new category, or the food was taken into account at a higher (parent) level (FoodEx Level 2) if this broader category was well-represented by the available categories at FoodEx Level 3. For example, 'Goose meat (Anser, Branta, Chen)' found at FoodEx Level 3, was taken into account at the parent FoodEx Level 2 'Poultry', as data reported were 100% left-censored but the broader category ('Poultry') was well-represented by other samples.
- Samples in the categories of the least detailed FoodEx classification (FoodEx Level 1) were excluded if no further information was available for a more specific reclassification. The exception to this were 27 samples reported as 'Meat and meat products (including edible offal)' which were considered at the second level of the FoodEx, i.e. as 'Livestock meat'.
- Categories where the analytical results were 100% left-censored and HBCDDs contamination
   was not expected, were not taken into account in the assessment. If quantified values were
   found in similar food categories at the same FoodEx level, the samples were grouped together
   with the similar food according to the same principle as in the first bullet point. Based on these



criteria, 'Snacks, deserts and other foods', 'Herbs, spices and condiments', 'Sugar and confectionary', 'Legumes, nuts and oilseeds', 'Vegetables and vegetable products (including fungi)' and 'Grains and grain-based products' (15 samples in total) were not included in the estimation of exposure to HBCDDs because they were 100% left-censored and/or represented with less than six samples. In addition, 135 samples were excluded from other categories because they were either 100% left-censored or represented with less than six samples at the level 3 of the FoodEx.

Mean occurrence values for the food categories considered at the third level of FoodEx are calculated as average of all samples reported within the same food category. However, for the samples reported/grouped at the second level of the FoodEx, mean occurrence values are calculated as average of all samples reported at second level and all associated samples reported at the more detailed level. For example, mean occurrence values for the samples reported/grouped as 'Fish meat', were calculated using all samples for which the second level was 'Fish meat' including those reported at more detailed levels.

For the detailed list of the categories considered see **Annex B** (Table B.2). Overall, 169 food samples (505 occurrence values) were excluded from the assessment from the 2,287 samples (6,857 occurrence values) as they did not fulfil the above selection criteria. In total, 51 food categories were considered for the linking between food occurrence and food consumption data (see **Annex B**, Table B.3).

# 2819 Occurrence data by food category

Mean values of the LB and UB by food category and FoodEx level, used for the estimation of exposure from 'Fish and other seafood' are presented in **Table 13**. The highest mean concentrations among Fish meat samples were reported for 'Eel' (2.33/2.35  $\mu$ g/kg ww, LB/UB) and 'Grey mullet (Mugil)' (2.06/2.07  $\mu$ g/kg ww, LB/UB). Mean concentrations reported for the rest of samples belonging to the food category of 'Fish meat' ranged from 0.00/0.01  $\mu$ g/kg (LB/UB) to 0.80/0.83 01  $\mu$ g/kg (LB/UB). Details on the concentrations reported for the food categories of 'Crustaceans', 'Fish offal' and 'Water molluscs' are presented in **Table 13**.

Mean values of the LB and UB for food categories other than 'Fish and other seafood' used for the estimation of exposure are presented in **Table 14**. Mean concentrations for the LB ranged from 0.00 µg/kg ww for 'Butter', 'Pork kidney', 'Game mammals' and 'Cow milk' to 0.53 µg/kg ww for 'Turkey meat (*Meleagris gallopavo*)'. The lowest mean occurrence value for the UB has been reported for 'Game mammals' (0.01 µg/kg ww) while the highest value was reported for 'Turkey meat (*Meleagris gallopavo*)' (0.58 µg/kg ww) (**Table 14**).

More details on grouping of samples within the food category are provided in **Annex B** (Table B.3). In addition, frequency distribution of LB and UB for relevant food categories are presented as violin plots with integrated box plots on the log10 scale (**Annex B**, Figure B.5 and B.6, respectively). The plot related to the LB shows only quantified values.

Table 13. Mean LB and UB values, as used for the exposure assessment, for the food category 'Fish and other seafood'

FoodEx Level 3	FoodEx Level <sup>(a)</sup>	Mean LB <sup>(b)</sup> (µg/kg ww)	Mean UB <sup>(b)</sup> (µg/kg ww)
Crustaceans	2	0.01	0.05
Shrimps (Crangon crangon)	3	0.00	0.05
Crab ( <i>Cancer</i> spp.)	3	0.02	0.07
Fish meat	2	0.19	0.25
Plaice ( <i>Pleuronectes</i> )	3	0.00	0.11

Sole ( <i>Limanda</i> ; <i>Solea</i> )	3	0.01	0.09
Sea catfish and wolf-fish (Anarhichas)	3	0.01	0.04
Lophiiformes ( <i>Pediculati</i> )	3	0.01	0.02
Cod and whiting ( <i>Gadus</i> spp.)	3	0.02	0.06
Tuna ( <i>Thunnus</i> )	3	0.02	0.09
Salmon and trout (Salmo spp.)	3	0.12	0.17
Sardine and pilchard (Sardina)	3	0.13	0.14
Trout	3	0.14	0.20
Hake ( <i>Merluccius</i> )	3	0.16	0.17
Salmon	3	0.17	0.27
Mackerel (Scomber)	3	0.49	0.51
Herring ( <i>Clupea</i> )	3	0.51	0.54
Bass ( <i>Marone</i> )	3	0.80	0.83
Grey mullet ( <i>Mugil</i> )	3	2.06	2.07
Eels	3	2.33	2.35
Fish offal	2	0.93	1.18
Water molluscs	2	0.07	0.07
Scallop ( <i>Pecten</i> spp.)	3	0.01	0.01
Clam ( <i>Mya arenaria</i> )	3	0.01	0.01
Queen scallop (Chlamys opercularis)	3	0.01	0.01
Oyster ( <i>Ostrea edulis</i> )	3	0.06	0.07
Mussel (Mytilus edulis)	3	0.12	0.12

2839 2840 2841 (a): FoodEx level eaqual to 2 means that samples were reported/grouped at the second level of the FoodEx; FodEx level equal to 3 means that samples were reported at the third level of the FoodEx.

(b): The values are rounded to two decimals.

#### 2842 Table 14. Mean LB and UB values, as used for the exposure assessment, for food categories other than 'Fish and other seafood' 2843

FoodEx Level 3	FoodEx Level <sup>(a)</sup>	Mean LB <sup>(b)</sup> (µg/kg ww)	Mean UB <sup>(b)</sup> (µg/kg ww)
Animal fat	2	0.06	0.46
Tallow	3	0.10	0.57
Pork lard (Schmaltz)	3	0.01	0.46
Chicken fat	3	0.01	0.37
Butter	3	0.00	0.49
Edible offal, farmed animals	2	0.01	0.03
Mutton / lamb liver	3	0.02	0.03
Pork kidney	3	0.00	0.03
Eggs, fresh	2	0.17	0.20
Whole egg, chicken	3	0.16	0.19
Game mammals	2	0.00	0.01
Liquid milk	2	0.01	0.03
Cow milk	3	0.00	0.03
Livestock meat	2	0.03	0.05
Beef meat ( <i>Bos</i> spp.)	3	0.05	0.08
Pork / piglet meat ( <i>Sus scrofa</i> )	3	0.04	0.05
Mutton / lamb meat ( <i>Ovis aries</i> )	3	0.02	0.04



Rabbit meat (Lepus cuniculus)	3	0.01	0.03
Poultry	2	0.18	0.21
Turkey meat (Meleagris gallopavo)	3	0.53	0.58
Chicken meat (Gallus domesticus)	3	0.01	0.03

(a): FodEx level eaqual to 2 means that samples were reported/grouped at the second level of the FoodEx; FodEx level equal to

2845 3 means that samples were reported at the third level of the FoodEx.

(b): The values are rounded to two decimals.

#### 2847 3.2.2. Previously reported occurrence data

The previous EFSA Opinion on HBCDDs (EFSA CONTAM Panel, 2011a) summarised the occurrence data 2848 on HBCDDs in food submitted by European countries as well as those published in peer-reviewed 2849 journals. Fish and other seafood were the food groups with the highest number of samples. Depending 2850 on the fishing ground, the HBCDD levels covered a guite broad contamination range. The occurrence 2851 data in other food groups were characterised by a high proportion of non-detects. Generally,  $\alpha$ -HBCDD 2852 was the predominant stereoisomer found in food of animal origin. Exceptions were, e.g. fish samples 2853 caught in rivers downstream of highly industrialised areas which showed higher concentrations for y-2854 HBCDD, the major stereoisomer in technical HBCDD. 2855

The following paragraphs, which does not claim for completeness, give a short summary on HBCDDs 2856 occurrence in food collected in Europe and published since the previous EFSA Opinion on HBCDDs. The 2857 primary focus of the recently published occurrence data was on fish and marine sea food. As already 2858 indicated in the previous Opinion,  $\alpha$ -HBCDD prevailed in most studies and the median levels, especially 2859 in fish caught for consumption was generally below 1  $\mu$ g/kg ww (Munschy et al., 2013; Kuc et al., 2014; 2860 Munschy et al., 2015; Aznar-Alemany et al., 2017; Malysheva et al., 2018, Nøstbakken et al., 2018; 2861 Vénisseau et al., 2018; Fliedner et al., 2018). However, a number of samples were detected where y-2862 HBCDD was the dominating stereoisomer. The reason for this, whether due to point sources or 2863 eventually influenced by the species, is not known. In addition, a number of substantially higher 2864 contamination levels were found, especially in mackerel and mussels. In general, the contamination 2865 levels in samples from the Mediterranean were higher than from the North Sea, the English Chanel and 2866 the North Atlantic. 2867

The HBCDD contamination of 25 fish oils for feed and food of different origin was assessed by Ortiz et al. (2011). Total HBCDDs ranged from 0.09 to 26.8  $\mu$ g/kg, with higher concentrations in fish oil for feed (average of 9.69  $\mu$ g/kg) than those for food (1.14  $\mu$ g/kg). In general, fish oils from Northern Atlantic presented higher levels (4.78–15.49  $\mu$ g/kg) than those from the Southern Pacific (3.34–5.49  $\mu$ g/kg).

Vénisseau et al. (2018) studied the occurence of legacy and novel brominated flame retardants, 2872 including HBCDDs in more than 600 food and feed samples from France collected for the period 2014 2873 to 2016 in the context of French monitoring plans. The broadest contamination range for the prevailing 2874  $\alpha$ -HBCDD stereoisomer was found in fish, fish derived products, such as fish meal and fish oil for feed, 2875 and shellfish. The range for  $\alpha$ -HBCDD in fish meal (n = 15), fish oil for feed (n = 15), fish (n = 114) 2876 and crustaceous/molluscs (n = 154) was determined as <LOD-1.08 (median: 0.024), 0.061-2.1 2877 (median: 0.697), <LOD-3.29 (median: 0.0235) and <LOD-0.368 (median: 0.021) µg/kg ww, 2878 respectively. While the frequency of detection in these food and feed commodities was between 67 and 2879 100%, the respective ranges for  $\beta$ -HBCDD and  $\gamma$ -HBCDD were between 24–40 and 33–87%, 2880 respectively. Compared to fish, fish derived products and marine sea food, the range of HBCDD levels 2881 in food of animal origin was substantially smaller. The concentration ranges for  $\alpha$ -HBCDD in milk (n = 2882 72), eggs (n = 57), sheep liver (n = 28), and meat (n = 152) were reported as <LOD-0.00987 (median: 2883 <LOD), <LOD-0.4152 (median: 0.00204), <LOD-0.1337 (median: <LOD), and <LOD-0.2325 (median:</p> 2884



 $\begin{array}{ll} 2885 \\ 2886 \\ 2886 \\ 2887 \end{array} 0.00118) \ \mu g/kg \ ww, \ respectively. \ The frequency of detection ranged between 21 and 54% \ and thus is lower compared to fish, fish derived products and shellfish. For <math display="inline">\beta$ -HBCDD and  $\gamma$ -HBCDD the frequency of detection in the food samples of animal origin ranged between 1 and 32%.

Fernandes et al. (2016) measured HBCDDs in most commonly consumed foods (n = 156) and animal feeds (n = 51) sampled in the UK. The food samples comprised fish and shellfish, milk, dairy products, eggs, meat, processed meat, offal, processed and "other" foods. Fish, shellfish, and processed meats showed the most frequent detections and also the highest HBCDD concentrations, where  $\alpha$ -HBCDD was predominant. The ranges for HBCDDs in the various food groups were generally between <0.01 and 0.1 µg/kg ww with higher levers found in fish and shellfish where maximum concentrations for  $\alpha$ -HBCDD of 10.13 and 3.41 µg/kg ww were determined.

- García López et al. (2018) assessed the HBCDD contamination of 53 composite food samples from Ireland collected between July and December 2015. The composite samples contained several hundred subsamples of eggs, milk, fish, fat (ovine, bovine, porcine and avian fat), and liver. Fish showed the broadest contamination range for the sum of HBCDDs (0.07–0.41 µg/kg ww) with a mean value of 0.28 µg/kg ww. A somewhat lower mean value of 0.23 µg/kg ww was found for fat (range: 0.06–0.20 µg/kg ww). Substantially lower HBCDD levels were found in egg yolk and liver with mean values of 0.01 and 0.05 µg/kg ww, respectively. In milk samples, HBCDDs were not detected above 0.01 µg/kg ww.
- Poma et al. (2018) reported on the occurrence of HBCDDs and other flame retardants in a total of 183 composite food samples collected in Belgium. Mean levels for the sum of the three HBCDD stereoisomers in fish, meat and meat products, milk and milk products, and eggs and egg products were 0.131, 0.043, 0.0001, and 0.971  $\mu$ g/kg ww, respectively. The high mean value in eggs and egg products was mainly attributed to the presence of  $\alpha$ -HBCDD (3.885  $\mu$ g/kg ww) in one organic egg sample. In food for infants and small children, animal and vegetable fat, grains and grain products, and potatoes and derived products, HBCDDs could not be detected.
- HBCDD levels in chicken eggs are generally low. Nevertheless, in a few cases concentrations of more than 3,000 µg/kg lipid were reported. As a plausible reason, Jondreville et al. (2017b) have demonstrated that hens can ingest HBCDD-containing extruded polystyrene (XPS) which is occasionally used as insulating material in rearing buldings. Due to the addition of HBCDDs to XPS which can be in the percent range, ingestion of this material by hens can lead to extraordinary HBCDD levels in chicken eggs.
- Dervilly-Pinel et al. (2017) assessed the chemical contamination levels of both conventional and organic 2915 meats. Besides  $\alpha$ -,  $\beta$ -, and y-HBCDD the study included a number of other inorganic and organic 2916 environmental contaminants together with chemical residues arising from production inputs. The type 2917 of farming investigated was conventional standard, Label Rouge, and organic farming. Label Rouge 2918 which is a French national quality assurance scheme for meat production (e.g. prescribing outdoor 2919 access, age and weight at slaughter) is closer to organic than to standard meat production. Median 2920 HBCDD contamination levels (sum of three stereoisomers) in bovine from conventional (n = 42) and 2921 organic farming (n = 43) were 0.03 and 0.04  $\mu$ g/kg lipid, respectively. Discrimination between the two 2922 farming practices was not possible. Regarding porcine meat, the median value for the sum of the 3 2923 stereoisomers in samples from conventional standard farming (n = 41) being 0.11 µg/kg lipid was lower 2924 compared to products from Label Rouge farming (n = 12) and organic farming (n = 43) with medians 2925 of 0.16 µg/kg lipid and 0.30 µg/kg lipid, respectively. The highest concentration of 194.8 µg/kg lipid 2926 was determined in a sample from conventional standard farming. Analyses of broiler meat for the 3 2927 stereoisomers revealed a median of 0.17 µg/kg lipid for samples from conventional standard farming 2928 (n = 31) which was significantly lower (p<0.001) than the medians for meat samples from Label Rouge 2929



farming (n = 13) and organic farming (n = 41) with medians of 0.34 and 0.33  $\mu$ g/kg lipid, respectively.

2931 In a monitoring study conducted in France from 2013 to 2015, Huneau-Salaun et al. (2020) measured 2932 the levels of HBCDDs in 60 hen egg farms (34 without an open-air range and 26 free-range), 57 broiler 2933 farms (27 without an open-air range and 30 free-range) and 42 pig farms in relation to their rearing 2934 environments. The authors measured the levels of HBCDDs in eggs, broiler muscle and pig muscle and 2935 found that  $\alpha$ -HBCDD was the most frequently detected stereoisomer found in eqgs, broiler muscle and 2936 pig muscle, while  $\beta$ -HBCDD was reported in broilers muscles, and  $\gamma$ -HBCDD in broiler and pig muscle. 2937 The authors concluded that the contamination of free-range eggs and broilers was more frequent than 2938 that of conventional ones, suggesting that access to an open-air range could be an additional source of 2939 exposure to HBCDDs. The authors could not establish any direct relationship between the occurrence 2940 of HBCDDs in eggs and meat and the characteristics of farm buildings (such as age, building materials).

- The above data from European countries show a broad range of HBCDD concentrations depending on the food commodity and its origin of sampling. This is also true for samples from non-European countries. A number of studies were conducted especially in China (Xia et al., 2011; Qiu et al., 2012; Shi et al., 2017; Meng et al., 2012; Yin et al., 2015; Zhu et al., 2013), the US (Schecter et al., 2010, 2012), Canada (Su et al., 2018), and Japan (Ueno et al., 2010; Kakimoto et al., 2012), which confirm the ubiquitous distribution of HBCDDs and their broad contamination of food.
- A compilation of global occurrence levels on HBCDDs and other flame retardants in food is presented in the recent publication by Aznar-Alemany and Eljarrat (2020).
- The impact of e-waste recycling plants on the HBCDD concentrations in eggs from free ranging chicken were demonstrated in several papers (Huang et al., 2018; Zeng et al., 2016; Zheng et al., 2012; Tao et al., 2016b). Levels of several thousand  $\mu$ g/kg fat were reported resulting in considerable dietary exposure when consumed by the inhabitants living in these areas.
- The Panel noted that the data that were submitted by European countries on HBCDD concentrations in food from official food control are in a comparable range as the results published during the last decade in the peer-reviewed literature. In general, the HBCDD concentrations in food for human consumption indicate to levels below 1 µg/kg ww. However, literature data also showed some substantially higher HBCDD concentrations in several specimens depending on the origin of sampling, such as in samples collected in contaminated areas, or in eggs and egg products from hens that were housed in rearing buildings where HBCDD containing extruded polystyrene (XPS) was used as insulating material.
- 2960 **3.2.3**. Food processing
- Data on effects of food processing on the HBCDD concentrations in food are scarce. Regarding food of animal origin, and considering that HBCDDs are lipophilic compounds, it can be assumed that all processing steps which alter the lipid content of a food commodity, such as trimming of visible fatty tissue will have an impact on the HBCDD levels in the processed sample. As HBCDDs are relatively stable compounds degradation at typical temperatures during food and feed processing seem unlikely. However, the  $\gamma$ -HBCDD stereoisomer can be rearranged to form  $\alpha$ -HBCDD at temperatures of 190°C (Peled et al., 1995) (see **Section 1.3.1**).
- The effect of steaming at 105°C for 15 minutes for fish, and at 105°C for 5 minutes for bivalves, each wrapped in aluminium foil, on the levels of various contaminants was investigated by Alves et al. (2017). In the case of  $\alpha$ -HBCDD, concentrations were similar in raw and steamed plaice and mussel, respectively (p>0.05).
- Aznar-Alemany et al. (2017) analysed 14 fish samples both raw and steamed with no further details on



steaming conditions given. Total HBCDD levels seemed to behave differently depending on the species
with increased levels for mackerel, mixed results for mussel and loss of contamination for plaice, tuna,
salmon and seabream.

2976Rasmussen et al. (2017) investigated the effect of industrial cold smoking on various contaminant levels,2977including α-, β- and γ-HBCDD, in Greenland halibut and farmed Atlantic salmon. The mean α-HBCDD2978concentration in the raw and processed Greenland halibut fillets (0.06 vs. 0.05 µg/kg ww) and the2979farmed Atlantic salmon (0.18 vs. 0.20 µg/kg ww) were not significantly different (p>0.05). β- and γ-2980HBCDD could not be detected at an LOD of 0.05 µg/kg ww.

- Zhang et al. (2017b) systematically studied the changes of HBCDDs during the refining process, 2981 including neutralization, bleaching, and deodorization in peanut, corn, and pressed and extracted 2982 soybean oils. The concentrations of HBCDDs in the crude oils of peanut, corn, pressed soybean, and 2983 extracted soybean oils were 0.04, 0.06, 0.52, and 0.10  $\mu$ g/kg, respectively. In all oils  $\alpha$ -HBCDD was by 2984 far the predominant stereoisomer. During the refining process, the mean concentrations of  $\alpha$ -HBCDD 2985 detected in peanut, corn, and extracted soybean oils were reduced by 25%, 17% and 100%, 2986 respectively. In contrast to the above oils, all three diastereoisomers were detected in all the pressed 2987 soybean oil samples. While  $\alpha$ - and  $\beta$ -HBCDD were reduced during the refining process by 90% and 2988 82%, respectively, there was no significant reduction of the y-HBCDD levels. 2989
- Several authors studied the presence of HBCDDs in kitchen hoods and utensils for food preparation. Bendig et al. (2013) analysed fat residue from 15 kitchen hoods for several polyhalogenated compounds. Total-HBCDD was detected in 3 samples at 8.8–11 µg/kg fat. Additionally, one sample contained the HBCDD metabolite pentabromocyclododecene at 350 µg/kg fat, which was about 40 times higher than HBCDDs. The results indicate that kitchen hood fat may serve as a valuable alternative matrix for the assessment of indoor contaminations with polyhalogenated compounds.
- 2996 Samsonek and Puype (2013) investigated brominated flame retardants in black thermo cups and 2997 selected kitchen utensils purchased on the European market. They did not find HBCDDs in any of the 2998 products tested using X-ray fluorescence spectrometry (XRF) to screen products for total bromine 2999 followed by thermal desorption GC-MS. Li et al. (2019b) also applied portable XRF combined with 3000 chemical analysis to investigate the occurrence, profile and distribution of the most important BFRs 3001 (PBDEs, HBCDDs and TBBPA) in 120 consumer products on the Chinese market, with the aim of 3002 identifying the possible reservoirs of persistent BFRs in products in China. Of these, 30 were daily-use 3003 household items, including storage boxes, fruit bowls, and cups and bottles. Bromine was only detected 3004 in four daily-use household products within a low concentration range.
- In summary, the data on the fate of HBCDDs during food processing are scarce. Although one report showed degradation of HBCDDs in kitchen hoods, due to their thermal stability it seems unlikely that they are formed or degraded during temperatures typical for food processing. Thus, potential reduction of their concentration in processed foods may be mainly caused by loss of fat, rather than degradation. Nevertheless, the  $\gamma$ -HBCDD stereoisomer can be rearranged to form  $\alpha$ -HBCDD at 190°C resulting in a different stereoisomeric pattern during food processing. It cannot be excluded that contact with BFR containing kitchen consumer goods may lead to contamination of the respective food due to migration.
- 3012 **3.3.** Dietary exposure assessment

### 3013 3.3.1. Current dietary exposure assessment

3014 The CONTAM Panel assessed the dietary exposure to HBCCDs (sum of  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCDD) following 3015 the methodology described in **Section 2.5**. A summary of the HBCDD occurrence data including the



3016 number of results and mean concentrations across the FoodEx level food categories as used for 3017 exposure assessment in presented in **Section 3.2.1**.

#### 3018 Mean and high dietary exposure

The mean and 95<sup>th</sup> percentile chronic dietary exposure to HBCDDs (ng/kg bw per day) was estimated separately for each consumption survey using data recorded at the individual level from the Comprehensive Database (see **Section 2.3**). Due the methodological differences among the surveys, chronic dietary exposure was estimated separately for each of them.

It was not possible to assess the dietary exposure to HBCDDs for 'Infants' using the data submitted to EFSA through the call for data since the occurrence data available in the food group 'Food for infants and small children' <sup>30</sup> was limited (only 19 samples) and was all left-censored. Exposure of breastfed infants to HBCDDs was assessed using data from a global WHO/UNEP study (see **Section 3.1.1.4**).

**Table 15** shows the summary statistics for the assessment of chronic dietary exposure to HBCDDs. Detailed mean and 95<sup>th</sup> percentile dietary exposure estimates calculated for all population groups for each of the 44 dietary surveys are presented in **Annex D** (Table D.1 and D.2). The total dietary intake was estimated using the LB and UB HBCDD concentrations from the calculated sum of  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCCD for all selected food groups.

# 3032 **Table 15.** Summary statistics for the chronic dietary exposure to HBCDDs (sum of α-, β- and γ-HBCDD) 3033 (ng/kg bw per day) across European countries

_	Mini	mum	Med	dian	Maxi	mum
Age group -	LB	UB	LB	UB	LB	UB
	Mean	dietary expos	sure in total p	opulation (ng	g/kg bw per o	lay) <sup>(b)</sup>
Toddlers	0.14	0.63	0.33	0.98	0.79	1.52
Other children	0.11	0.37	0.28	0.77	0.63	1.21
Adolescents	0.07	0.19	0.17	0.36	0.34	0.64
Adults	0.08	0.17	0.16	0.29	0.28	0.44
Elderly	0.10	0.19	0.16	0.30	0.23	0.40
Very elderly	0.07	0.19	0.15	0.30	0.30	0.49
	P95 d	lietary expos	ure in total po	opulation (ng	/kg bw per d	ay) <sup>(b)</sup>
Toddlers <sup>(a)</sup>	0.58	1.54	0.90	2.16	2.30	3.61
Other children	0.37	1.01	0.78	1.66	2.05	2.95
Adolescents (a)	0.23	0.51	0.44	0.86	1.20	1.54
Adults	0.29	0.48	0.46	0.73	1.18	1.37
Elderly	0.30	0.50	0.49	0.74	0.74	0.90
Very elderly <sup>(a)</sup>	0.30	0.45	0.44	0.74	0.78	0.97

bw: body weight; LB: lower bound; UB: upper bound.

(a): The 95<sup>th</sup> percentile estimates obtained on dietary surveys/age groups with fewer than 60 observations may not be

3036 statistically robust (EFSA, 2011a) and are therefore not included in this table.

3037 (b): The values are rounded to two decimals.

The highest estimated chronic dietary exposure to HBCDDs (except for breastfed infants) was in the younger age groups, in particular in 'Toddlers'. For this age group, the mean dietary estimates range

3040 from 0.14 to 1.52 ng/kg bw per day (minimum LB and maximum UB, respectively). The 95<sup>th</sup> percentile

3041 exposure ranges from 0.58 to 3.61 ng/kg bw per day (minimum LB and maximum UB, respectively).

3042 Dietary exposure in specific groups of the population, namely 'Pregnant women' and 'Lactating women',

<sup>&</sup>lt;sup>30</sup> Description as in FoodEx.



were within the range of exposure estimates for the adult population (see **Annex D**, Table D.1 andD.2).

3045 In the previous Opinion (EFSA CONTAM Panel, 2011a), due to lack of consumption data, an exposure 3046 estimation for toddlers was not possible which precludes a comparison with the current assessment. 3047 The age group 'Other children' was identified in the former Opinion to have the highest dietary exposure 3048 to HBCDDs. The mean and P95 dietary exposure then ranged between 0.15–1.85 and 0.80–4.46 ng/kg 3049 bw per day, respectively. In the current assessment, the respective estimated dietary exposure is 3050 somewhat lower, being 0.11–1.21 and 0.37–2.95 ng/kg bw per day, for mean and P95 dietary exposure, 3051 respectively. The same holds true for dietary HBCDD exposure for adults. While the mean and P95 3052 dietary exposure across dietary surveys in European countries in the former HBCDD Opinion was 3053 estimated as 0.09-0.99 and 0.39-2.07 ng/kg bw per day, respectively, the current estimates range 3054 between 0.08–0.44 and 0.29–1.37 ng/kg bw per day, for mean and P95 dietary exposure, respectively.

In the previous Opinion (EFSA CONTAM Panel, 2011a), a scenario for high and frequent fish consumers was estimated as high consumption of fish meat was considered as a special diet with specific concern for dietary exposure to HBCDDs. In the current Opinion the intake estimations of HBCDDs for high and frequent fish meat and fish offals consumers are represented by the 95th percentile reported in **Table 15**.

It can be assumed that the dietary HBCDD exposure for vegetarians is lower than that for people consuming a mixed diet. This is because HBCDDs are persistent and lipophilic compounds with low water solubility that bioaccumulate in the food chain. Thus, consumption of food of animal origin represents the main route of human dietary exposure to HBCDDs. Since uptake of HBCDDs by plants from soil is low, the contamination of food of plant origin is generally of minor importance. This is substantiated by the occurrence data on HBCDDs in food samples of plant origin submitted by several European countries to EFSA which were almost completely below LOD/LOQ.

#### 3067 **Contribution of different food categories to the chronic dietary exposure**

The contribution of individual food categories to the LB and UB mean chronic dietary exposure to HBCDDs varied between the dietary surveys and is shown in **Annex D** (Table D.4 and D.5). This is explained by the specific food consumption patterns in the individual European countries and even in different regions within a country. The detailed contribution to the mean LB chronic dietary exposure to HBCDDs of the different food categories at FoodEx Level 2 and grouped by age class, country and survey as used for the exposure assessment and grouped by age classes is shown in **Annex D** (Table 0.3).

In **Table 16** the relative contribution (%) of each food category to the overall mean LB and UB exposure of HBCDDs as median and range (minimum and maximum) for each age class across all European dietary surveys is presented. Dietary exposure reflects the pattern of consumption figures of each age class and the respective country as well as occurrence values.

When a high proportion of left-censored data produces a large difference between UB and LB occurrence values, this results in a commensurate uncertainty in dietary exposure estimates. Appraising the contribution of the respective food groups to the total LB exposure is based on measured values not influenced by the percentage and magnitude of the left-censored data. Appraising the contribution of food groups to the total UB dietary exposure should be done with care as the high contribution of certain food groups can be the result of high LOQs or high consumption rather than an actual high contribution.

Four main food categories ('Fish meat', 'Eggs fresh', 'Livestock meat' and 'Poultry') are the main contributors for the LB estimates. Six food categories ('Fish meat', 'Eggs fresh', 'Livestock meat', 'Poultry', 'Liquid milk' and 'Animal fat') are the main contributors for the UB estimates (highlighted in grey in the table).



Considering the LB estimates, which are less influenced by the values of the LOD/LOQ, the contribution of 'Fish meat' to the median intake of HBCDDs across European dietary surveys varies from 22% in 'Adolescents' to 47% in 'Elderly'. A considerably lower contribution to the dietary intake of HBCDDs is observed when the UB estimates are considered, where the contribution of 'Fish meat' to the median intake of HBCDDs varies from 11% in 'Toddlers' to 30% in 'Elderly'.

The food category of 'Eggs, fresh' is also significantly contributing to the dietary intake of HBCDDs with a median LB contribution across European dietary surveys ranging from 18% in 'Very elderly' to 27% in 'Other children'. The median UB contribution is lower, ranging from 8% in 'Toddlers' to 13% 'Other children'.

For the food group 'Livestock meat', the median LB contribution ranges from 14% in 'Elderly' to 21% in 'Adolescents', whereas considering the median UB, the contribution ranges from 7% in 'Toddlers' to 13% in 'Adults'.

The food category 'Poultry' is also contributing to the dietary intake of HBCDDs with a median LB contribution across European dietary surveys ranging from 16% in 'Adolescents' to 19% in 'Very elderly', whereas considering the median UB, the contribution ranges from 9% in 'Toddlers' to 14% in 'Adults'.

For the food group 'Liquid milk', the median LB contribution ranges from 1% in 'Elderly' to 9% in 'Toddlers'. The median UB contribution is much higher ranging from 14% in 'Elderly' to 44% in 'Toddlers'.

The category of 'Animal fat' is only contributing in the UB estimates with values ranging from 8% in 'Toddlers' to 17% in 'Very elderly'.

The distinct influence of left-censored data on the calculation of the distribution of different food categories to the chronic dietary exposure can best be demonstrated by the contribution of 'Liquid milk'. While the LB contribution ranges from 1.3–8.7% across the different age classes, the corresponding UB values amount to 14.3 to 42.8%. As a result, the contribution of other food groups decreased.

In the previous HBCDD Opinion (EFSA CONTAM Panel, 2011a), the contribution of 'Fish meat and products' to the median LB intake of HBCDDs across European dietary surveys varied from 83 to 88.2% for the different age groups. In the current assessment, the contribution of fish meat to the LB chronic dietary exposure only ranges between 22 and 47.1%. The CONTAM Panel notes that a comparison of the current data with results from the previous Opinion is hampered by a number of facts, such as improvements in instrumental analysis, different percentage of left-censored data, consideration of further food commodities, and use of more detailed consumption data.



**Table 16.** Relative contribution (%) to the overall mean dietary intake of HBCDDs of the food categories across different age classes of the general population. 3120 The median and the range (minimum and maximum) lower (LB) and upper bound (UB) contributions of each food category to the overall diet across the 3121 European dietary surveys are given. Percentages refer to the intake of HBCDDs estimated from LB and UB mean values occurrence from the calculated sum of 3122  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCDD for all selected food categories. The values are rounded to two decimals

3123

						Age c	lass					
	Т	oddlers	Othe	er children	Ado	lescents	ŀ	dults	E	lderly	Ver	y elderly
	Median	(Min-Max)	Median	(Min-Max)	Median	(Min-Max)	Median	(Min-Max)	Median	(Min-Max)	Median	(Min-Max)
Food category												
						LB (%	<b>b)</b> <sup>(a)</sup>					
Animal fat	0.04	(0.00-0.15)	0.04	(0.00-0.35)	0.03	(0.00-0.57)	0.04	(0.00-1.58)	0.05	(0.00-1.80)	0.05	(0.00-3.61)
Crustaceans	0.04	(0.00-0.18)	0.04	(0.00-0.19)	0.14	(0.01-0.31)	0.08	(0.00-0.51)	0.09	(0.01-0.43)	0.13	(0.02-0.22)
Edible offal, farmed animals	0.06	(0.01-0.63)	0.10	(0.00-0.64)	0.08	(0.01-0.68)	0.13	(0.03-1.11)	0.20	(0.09-0.86)	0.19	(0.04-2.23)
Eggs, fresh	24.09	(9.06-56.38)	27.15	(10.60-43.55)	20.52	(5.52-45.70)	18.77	(5.78-37.34)	17.72	(4.49-37.19)	16.63	(8.95-37.64)
Fish meat	22.90	(12.37-50.01)	25.70	(13.48-45.70)	22.36	(12.19-51.75)	33.54	(9.06-68.71)	47.11	(6.46-65.45)	44.41	(5.03-66.20)
Fish offal	0.24	(0.03-25.81)	0.30	(0.02-9.70)	0.36	(0.04-5.50)	0.50	(0.08-13.67)	1.01	(0.04-7.48)	0.71	(0.02-9.90)
Game mammals	0.01	(0.00-0.08)	0.01	(0.00-0.07)	0.01	(0.00-0.02)	0.01	(0.00-0.08)	0.01	(0.00-0.08)	0.01	(0.00-0.04)
Liquid milk	8.73	(1.77-28.09)	4.91	(1.43-12.80)	3.36	(1.26-12.26)	1.44	(0.60-5.23)	1.30	(0.53-2.90)	1.36	(0.74-3.95)
Livestock meat	15.45	(6.25-31.77)	18.34	(7.30-36.47)	21.18	(10.64-48.17)	18.47	(9.95-34.03)	13.86	(7.38-31.57)	18.28	(7.07-32.36)
Poultry	18.54	(1.85-39.81)	17.32	(4.50-41.62)	16.10	(2.57-55.21)	18.31	(4.06-40.44)	15.84	(3.26-35.84)	19.22	(1.78-29.37)
Water molluscs	0.26	(0.01-1.10)	0.38	(0.00-4.99)	0.51	(0.01-8.15)	0.61	(0.00-4.89)	1.31	(0.04-3.81)	0.37	(0.02-2.59)
						UB (%	6) <sup>(a)</sup>					
Animal fat	7.85	(0.63-31.48)	9.09	(0.59-36.13)	9.65	(1.37-41.14)	10.43	(1.94-38.93)	15.81	(0.60-32.71)	16.80	(5.44-36.44)
Crustaceans	0.07	(0.01-0.36)	0.12	(0.00-0.58)	0.31	(0.02-1.09)	0.29	(0.00-1.30)	0.30	(0.07-0.88)	0.33	(0.07-0.56)
Edible offal, farmed animals	0.07	(0.01-0.41)	0.09	(0.00-0.55)	0.10	(0.00-0.59)	0.15	(0.05-1.36)	0.25	(0.12-1.02)	0.27	(0.07-2.34)
Eggs, fresh	8.23	(4.10-33.79)	13.28	(4.67-21.10)	9.16	(2.96-21.66)	11.40	(4.87-24.14)	10.17	(3.31-22.18)	9.23	(5.87-22.57)
Fish meat	11.04	(5.09-34.53)	12.56	(6.17-30.19)	13.33	(6.88-29.87)	24.90	(6.56-53.70)	29.50	(4.58-53.18)	27.77	(3.31-53.32)
Fish offal	0.09	(0.02-8.62)	0.12	(0.01-4.39)	0.21	(0.03-2.91)	0.31	(0.06-10.00)	0.80	(0.04-5.88)	0.46	(0.01-7.41)
Game mammals	0.01	(0.00-0.12)	0.02	(0.00-0.10)	0.01	(0.00-0.04)	0.02	(0.00-0.18)	0.03	(0.00-0.21)	0.03	(0.00-0.12)
Liquid milk	44.26	(15.02-70.64)	37.06	(13.38-68.16)	30.31	(13.30-67.59)	15.18	(6.22-30.69)	14.30	(6.12-25.48)	17.80	(7.67-27.48)

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#### Hexabromocyclododecanes (HBCDDs) in food

Livestock meat	7.80	(3.11-12.72)	8.24	(4.31-14.58)	12.43	(6.48-24.52)	12.66	(7.93-21.92)	11.32	(6.37-22.55)	12.44	(5.77-21.68)
Poultry	9.43	(2.58-22.21)	10.81	(3.68-26.49)	10.67	(4.36-34.78)	14.31	(5.66-33.59)	12.54	(4.53-25.84)	13.74	(2.64-22.32)
Water molluscs	0.11	(0.00-0.63)	0.12	(0.00-2.64)	0.24	(0.00-4.86)	0.33	(0.00-3.07)	0.94	(0.02-2.36)	0.20	(0.01-1.46)

3124

(a): Due to the high proportion on non-detects across the individual stereoisomers, the calculation of the UB sum might be overestimated.

96



#### 3125 Breastfed infants

For the exposure assessment of breastfed infants, an age of three months was selected, equivalent to a weight of about 6.1 kg, with an estimated average daily consumption of about 800 mL and a high consumption of 1,200 mL of human milk, each with a mean fat content of 3.5% (EFSA CONTAM Panel, 2011a).

The exposure scenario based on average human milk consumption and the reported UB range for  $\Sigma$ HBCDD (predominantly  $\alpha$ -HBCDD) in pooled human milk samples collected in European countries between 2014 and 2016 as part of the WHO/UNEP field studies (see **Table 8**), would result in a median daily exposure of 14.3 ng/kg bw (range: 3.2–73.7 ng/kg bw). For infants with high human milk consumption the respective median daily exposure would result in 21.5 ng/kg bw (range 4.8–110.6 ng/kg bw).

Using the LB data, the median daily exposure would result in 13.8 ng/kg bw (range: 2.3–73.4 ng/kg bw). For infants with high human milk consumption the respective LB median daily exposure would result in 20.7 ng/kg bw (range 3.4–110.2 ng/kg bw).

3139 The Panel noted that since these were pooled samples, it was not possible to estimate specific values 3140 for individuals.

Compared to the previous Opinion on HBCDDs (EFSA CONTAM Panel, 2011a) where daily exposures to HBCDDs from average and high milk consumption of 0.60–142 and 0.90–213 ng/kg bw, respectively, were calculated, the respective estimations in the current assessment, in particular at the upper end, are substantially lower.

3145 **3.3.2. Previously reported dietary exposure assessment** 

In 2011, the CONTAM Panel summarised the occurrence data submitted to EFSA by European countries 3146 and assessed chronic dietary exposure based on the concentration of Total HBCDDs in the food group 3147 of 'Fish and other seafood' and the sum of the individual HBCDD stereoisomers ( $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCCD) 3148 for 'Eggs and egg products', 'Milk and dairy products' and 'Meat and meat products' (EFSA CONTAM 3149 Panel, 2011a). The highest mean estimated dietary exposure to HBCDDs across the European dietary 3150 surveys was for children from three to ten years old ('Other children') and was between 0.15 to 1.85 3151 ng/kg bw per day. The range reflects the minimum LB and the maximum UB exposure, respectively. 3152 Total dietary exposure for adults was around half the exposure for 'Other children', with minimum LB 3153 and maximum UB of respectively 0.09 and 0.99 ng/kg bw per day. For high consumers (95<sup>th</sup> percentiles), 3154 the dietary intake of HBCDDs across European countries for 'other children' were between 0.80 and 3155 4.46 ng/kg bw per day (minimum LB and maximum UB, respectively). The corresponding data for 3156 'adults' were between 0.39 and 2.07 ng/kg bw per day, respectively. 3157

As contamination of food samples of plant origin is generally lower than that of food samples of animal origin, the CONTAM Panel assumed that the dietary exposure to HBCDDs for vegetarians is lower than that for people consuming a mixed diet.

The exposure scenario based on average consumption of human milk (800 mL per day) and the reported range for Total HBCDDs resulted in daily exposures of about 0.6–142 ng/kg bw. For infants with high human milk consumption (1,200 mL per day) the respective daily exposures ranged from 0.90 to 213 ng/kg bw (EFSA CONTAM Panel, 2011a).

The following, which does not claim for completeness, gives a short summary on assessments on dietary exposure to HBCDDs in Europe published in open literature since the previous EFSA Opinion on HBCDDs.



Tao et al. (2017) collected samples of 14 different food groups (3 samples each) from two supermarkets 3167 representing national chains and one local market in Birmingham, UK during May and June 2015. 3168 Besides a number of emerging flame retardants, the study also included HBCDDs. Dietary intakes of the 3169 flame retardants were calculated for UK toddlers and adults based on the analytical results of the 3170 samples and the food consumption data from the latest national diet and nutrition survey report. Body 3171 weight values were assumed to be 70 kg for adults and 10 kg for toddlers, respectively. The estimated 3172 average dietary intakes of the sum of  $\alpha$ -,  $\beta$ - and y-HBCDD for UK adults and toddlers are reported to 3173 be 8.8 and 31 ng/day, respectively. 3174

Based on the analysis of 42 commercial sea food samples (10 species) and the respective seafood consumption pattern, Aznar-Alemany et al. (2017) estimated a mean daily intake via fish of 0.49 ng/kg bw for Spanish adults. The P99 (high seafood consumers) exposure for the UB scenario resulted in a mean daily intake of 1.1 ng/kg bw.

Coelho et al. (2016a) measured a number of persistent organic pollutants including HBCDD in 7-days duplicate diet samples from 21 Portuguese volunteers. Estimated daily intakes (EDIs) of the target compounds were calculated considering that a person ingests daily 1,867.2 g of food. For HBCDDs, the levels of α-, β- and γ-HBCDD were mostly below LOD, with α-HBCDD being the most frequent isomer detected (23.8%). The estimated daily mean LB and UB intake for the sum of the three HBCDD isomers was reported as 2.0 ng/kg bw (range: 0–37, median: 0 ng/kg bw), and 2.5 ng/kg bw (range: 0.37–37, median: 0.70 ng/kg bw).

Sahlström et al. (2015b) determined the concentrations of HBCDDs among other brominated flame retardants in diet and house dust for a mother-toddler cohort and estimated exposure via these two media. Market basket samples comprised the food categories fish, meat, vegetable oils, dairy products, and eggs (4 homogenates for each category) collected in 2010. The median daily intake of the sum of  $\alpha$ -,  $\beta$ -, and  $\gamma$ -HBCDD from diet for Swedish mothers and toddlers (n = 20) was estimated as 11 ng per day (range: 2.7–21 ng/day) for mothers and 5 ng per day (range: 1.3–9.7 ng per day) for toddlers, respectively.

Rivière et al. (2014) reported on the results of the total diet study (TDS) performed in France 2007-2009. The TDS covered the most important foods in terms of consumption, selected nutrients and contribution to contamination. The mean daily dietary exposure to the sum of  $\alpha$ -,  $\beta$ -, and  $\gamma$ -HBCDD was 0.211 ng/kg bw and 0.32 ng/kg bw for adults and children, respectively. The P95 was estimated at 0.448 ng/kg bw per day and 0.734 ng/kg bw per day for adults and children, respectively.

Based on consumption data statistics, food items from six food groups, i.e. fish and seafood, meat, animal fat, dairy products, eggs, and vegetable oils, Eljarrat et al. (2014) assessed the dietary daily exposure of the Spanish adult population to HBCDDs. The daily intake to the sum of  $\alpha$ -,  $\beta$ -, and  $\gamma$ -HBCDD was estimated at 2.58 ng/kg bw. The HBCDD exposure mainly came from fish and seafood (56%), dairy products (14%) and meat (12%).

In summary, the dietary exposure assessments performed in the past decade in European countries generally point to mean intake values for the sum of  $\alpha$ -,  $\beta$ -, and  $\gamma$ -HBCDD exposure between <1 and 2.5 ng/kg bw per day, which is in general accordance with the exposure estimates performed by EFSA based on the occurrence data submitted by the European countries. The major categories contributing to total dietary exposure are food of animal origin, such as fish and fish derived products, meat, dairy products and eggs.



#### 3209 3.3.3. Non-dietary sources of exposure

3210 Non-dietary human exposure to HBCDDs can occur via inhalation of gas-phase HBCDDs and HBCDDs 3211 on particles, as well as oral intake of dust, and via dermal contact with consumer products that may 3212 contain HBCDDs. HBCDD levels in dust are summarised in **Appendix A**. In the former EFSA Opinion on 3213 HBCDDs, the CONTAM Panel concluded that non-dietary exposure, mainly through dust in homes, 3214 offices, schools, cars and public environment can substantially contribute, and in some cases even 3215 dominate the total human exposure to HBCDDs, especially for toddlers and other children (EFSA 3216 CONTAM Panel, 2011a). The following paragraphs give some examples on non-dietary exposure 3217 assessments for HBCDDs published since the previous EFSA Opinion, and where available, compares 3218 the non-dietary with dietary exposure. The focus is on studies where the samples are collected after 3219 2010.

### 3220 Non-dietary oral exposure

3221 Kalachova et al. (2012) determined several BFRs in dust collected in 25 Czech households and 27 car 3222 interiors (see levels **Appendix A**). The exposure was estimated assuming that the mean dust ingestion 3223 rates for adults and toddlers (6-24 months) were 20 and 50 mg per day, and high dust ingestion rates 3224 were 50 and 200 mg per day, respectively. For adults, and considering a mean household dust ingestion, 3225 the exposure (mean (median, P95)) was 2,600 (1,300, 8,600) pg per day, while for high dust ingestion 3226 it was 6,400 (3,300, 21,500) pg per day. The exposure from car dust was considerably lower: for a 3227 mean car dust ingestion the exposure was estimated as 50 (30, 130) pg per day, and for a high car 3228 dust ingestion it was 110 (60, 320) pg per day. For toddlers, the exposure from both dust sources was 3229 higher. Considering a mean holsehold dust ingestion, the exposure was estimated as 8,500 (4,400, 3230 28,600) pg per day, and with a high household dust ingestion it was 34,000 (17,700, 114,500) pg per 3231 day. As for adults, car dust exposure was lower: for a mean ingestion, the exposure was 110 (60, 320) 3232 pg per day, and for a high car dust ingestion it was 460 (250, 1,300) pg per day.

3233 Fromme et al. (2014) collected 20 residences' dust samples from vacuum cleaner bags and analysed 3234 them for several brominated flame retardants, including HBCDDs in order to assess human exposure 3235 through dust (see levels in **Appendix A**). Two scenarios were examined, an average intake on the basis 3236 of median concentrations in house dust, and a high intake on the basis of 95<sup>th</sup> percentiles. In addition, the exposure was determined for the group of adults and that of toddlers. Body weights of 70 kg and 3237 3238 12 kg for adults and toddlers, respectively, were assumed. The average daily intake of house dust was 3239 assumed to be 30 mg for adults and 60 mg for toddlers. The average and high intakes for adults are 3240 given as 0.15 pg/kg bw and 0.76 pg/kg bw, respectively. For toddlers, the respective daily intakes via 3241 dust amount to 1.7 pg/kg bw and 8.9 pg/kg bw.

3242 Strid et al. (2014) investigated personnel exposure to BFRs both during flights and maintenance of 3243 aircrafts. High HBCDD concentrations in dust and air were detected in relation to the insulation material 3244 in the aircraft (see **Appendix A**), something that was not reflected in the serum samples of the involved 3245 personnel since the presence of HBCDDs was only indicated in a few samples. Allen et al. (2013) also 3246 investigated exposure to flame retardant chemicals from commercial aircraft and found levels in dust of 3247 up to 1,100,000 ng/g (see **Appendix A**).

Sahlström et al. (2015b) determined the concentrations of HBCDDs among other BFRs in diet and house dust for a mother-toddler cohort and estimated exposure via these two media. Dust samples (n = 27) were collected on surfaces at least 1 m above the floor in the living room, kitchen, bedroom and/or hallway (see levels **Appendix A**). Based on dust ingestion rates of 30 mg per day for adults and 60 mg per day for toddlers, median (range) intakes for the sum of  $\alpha$ -,  $\beta$ - and  $\gamma$ -HBCDD via dust of 3.4 (0.86– 180) ng per day, and 6.9 (1.7–360) ng per day for mothers and toddlers, respectively, were estimated. While for the mothers, the median dietary intake for the sum of the three HBCDD isomers was around



three times higher compared to dust (11 vs. 3.4 ng per day), for toddlers, the median intake via dust exceeded the respective dietary intake (6.9 vs. 5.0 ng per day).

Coelho et al. (2016b) analysed 28 house dust samples (vacuum cleaner bags) from two Portuguese cities, for HBCDDs and other organic contaminants (see levels in **Appendix A**). Human exposure through dust ingestion was evaluated by calculating the estimated daily intakes based on the high dust ingestion rates of 100 mg per day for adults and 200 mg per day for children, and considering 70 and 12 kg as the average body weights of adults and children, respectively. While for adults the estimated daily intake ranged from 22 to 2,900 pg/kg bw (mean: 540, median: 220 pg/kg bw), the corresponding intakes for children ranged from 260 to 33,000 pg/kg bw (mean: 6,300, median: 2,600 pg/kg bw).

Tay et al. (2017) estimated human exposure to HBCDDs and other flame retardants via inhalation and dust ingestion. For this assessment, 60 indoor stationary air samples, 13 personal air samples, and 60 settled dust samples were collected from a Norwegian cohort during winter 2013 (see levels in **Appendix A**). The median (P95) daily exposure via dust ingestion was estimated as 68 (500) pg/kg bw. The estimated median (P95) daily inhalation from stationary air sampling and personal air sampling were 0.083 (4.2) pg/kg bw and 8.1 (14) pg/kg bw, respectively.

Besis et al. (2017) investigated the occurrence of brominated flame retardants in the dust from the interior of private cars in Thessaloniki/Greece, aged from 1 to 19 years with variable origin and characteristics (see levels in **Appendix A**). Median and 95<sup>th</sup> daily exposure to  $\Sigma$ HBCDDs through dust ingestion for adults were 0.00731 and 0.0253 ng/kg, respectively. For toddlers, the respective daily exposures were 0.0512 and 0.177 ng/kg bw. The median and 95<sup>th</sup> daily exposures through dermal contact for adults were 0.00310 and 0.0107 ng/kg bw, respectively. For toddlers, the respective daily exposures were 0.0402 and 0.139 ng/kg bw.

3277 Kurt-Karakus et al. (2017) analysed several BFRs in indoor and outdoor air and indoor dust in urban, 3278 semi-urban and rural locations in Istanbul/Turkey, including HBCDDs (reported as γ-HBCDD) (see levels 3279 in **Appendix A**). The median low and high intake of HBCDDs via dust for adults and children was 3280 estimated to be 16–379 pg/kg bw per day and 952–3,460 pg/kg bw per day, respectively. The median 3281 intake via inhalation and dermal contact for adults and children were estimated as 25 and 6 pg/kg bw 3282 per day, and as 5 and 12 pg/kg bw per day, respectively.

3283 Larsson et al. (2018) analysed several brominated flame retardants in dust collected in 2015 from a 3284 total of 100 preschools in Sweden. In addition, also 100 hand wipe samples were collected and analysed 3285 for BFRs (see levels in **Appendix A**). For the exposure calculation of the 4-year old children, a dust 3286 intake of 30 mg and a body weight of 17.6 kg was assumed. The daily intake of ΣHBCDD in preschool 3287 dust via dermal contact was calculated using various assumptions, such as exposed body area (hands, 3288 arms, legs) of children of 3,380 cm<sup>2</sup>, amount of dust adhered to the skin, absorbtion factors, and fraction 3289 of the time spent in the preschool. The estimated geometric mean (P95) daily intake of **SHBCDD** in 3290 preschool dust via oral and dermal exposure in 4-year old children were 200 (1,900) pg/kg bw and 7 3291 (60) pg/kg bw, respectively.

Wemken et al. (2019) measured HBCDDs and other flame retardands in indoor air and dust collected from Irish homes, cars, offices, and primary schools during 2016–2017.  $\alpha$ -HBCDD was the dominant isomer in homes, offices, and schools, with  $\gamma$ -HBCDD dominant in cars (see levels in **Appendix A**). Using median  $\Sigma$ HBCDD occurrence concentrations, the daily total exposure via air and dust for adults, toddlers, and school children was estimated as 140, 2,500 and 1,400 pg/kg bw, respectively. Assuming ingestion/inhalation of the 95<sup>th</sup> percentile HBCDD concentrations, the respective daily exposure scenarios via air and dust was 7,800, 170,000, and 86,000 pg/kg bw.

The broad range of dust contamination with HBCDDs and its impact on non-dietary exposure is supported in the following investigations conducted in non-European countries.



3301 Ali et al. (2012) analysed 50 indoor dust samples collected from different homes in selected rural and 3302 urban areas of New Zealand. HBCDD concentrations ranged between 20-4,100 ng/g (median: 190, 3303 mean 460 ng/g). For the exposure assessment, average adult and toddler dust ingestion figures of 20 3304 and 50 mg per day, and high dust ingestion intakes for adults and toddlers of 50 and 200 mg per day, 3305 respectively ,were assumed. Body weights used were 70 kg for adults and 12 kg for toddlers. Considering mean and high dust intake, the median (P95) daily exposure for adults was 50 (500) and 3306 3307 130 (1,260) pg/kg bw, respectively. For toddlers, the corresponding daily exposures were calculated as 3308 780 (7,320) and 3,130 (29,300) pg/kg bw, respectively.

Tue et al. (2013) investigated the occurrence of several brominated flame retardants in indoor dust and air from two Vietnamese e-waste recycling sites (EWRSs) and an urban site in order to assess the relevance of these media for human exposure. The EWRSs were rural communes of approximately 250 households, with about 40–50% regularly involved in recycling of metals and plastics from e-waste. The levels of HBCDDs in settled house dust from the EWRSs of 5.4–400 ng/g were significantly higher than in dust from houses in urban and surban areas which ranged from 0.99–61 ng/g, indicating the importance of informal recycling of e-waste.

3316 Barghi et al. (2017) investigated BFR concentrations in 124 vacuum dust samples of six categories of 3317 indoor environments (homes, offices, kindergartens, cars, schools, and public indoor environments) and 3318 32 surface dust samples from five cities in South Korea, and based on these results, estimated exposures 3319 for adults and children. The median  $\Sigma$ HBCDD concentrations ranged from 106.30 ng/g in home dust to 3320 496.13 ng/g in office dust. Due to the broad electrical equipment in offices, the measured concentrations 3321 were significantly higher than those determined in schools and homes (p<0.05). The exposure via dust 3322 was estimated using the time activity in the different indoor places, and high and average dust intake of >3 years olds and 1–3 year olds of 50 and 20 mg dust per day, and 200 and 50 mg dust per day, 3323 3324 respectively. The estimated dust intake for adults and toddlers was 10.66 and 4.27 ng/day, and 35.75 3325 and 8.94 ng/day, respectively. A comparison with dietary intake revealed that food is the major 3326 contributor for HBCDD intake in all subgroups except 1–3-year olds in the high dust intake scenario.

Wang et al. (2018) analysed dust samples from 30 homes and 27 offices from Beijing (China) for a number of BFRs, including HBCDDs. While the concentrations for  $\Sigma$ HBCDD in house dust ranged from 73.6–995 ng/g (median: 156 ng/g), the respective concentrations in office dust were determined as 110–394 ng/g (median: 258 ng/g). The median exposure for  $\Sigma$ HBCDD via dust intake for adults was estimated as 48 pg/kg bw per day, which is substantially lower than the estimated daily dietary intake of 1,050 pg/kg bw per day based on a Chinese total diet study. For toddlers, the median intake via dust was reported as 649 pg/kg bw per day.

3334 The CONTAM Panel made a rough scenario to estimate the potential ingestion from dust in toddlers and 3335 adults. The US-EPA assumed that while an adult ingests an average of 20 mg per day, a child of 1-2years old ingests an average of 50 mg per day (US-EPA, 2017). Values found in dust from European 3336 3337 countries ranged from very low to around 150,000 µg/kg in homes and up to 1,000,000 µg/kg in some 3338 vehicles (see Section 1.3.3). The majority of data reported for dust in homes had concentrations below 3339 1,000  $\mu$ g/kg, and so this was taken as a high but realistic value for  $\Sigma$ HBCDD to make an estimate of 3340 exposure from dust. Considering a body weight of 12 kg for a 1–3-years old child and 70 kg for adults 3341 (EFSA SC, 2012a), the resulting daily exposures via dust ingestion would be around 4 ng/kg bw per day 3342 for toddlers and around 0.3 ng/kg bw per day for adults. Whilst there is a wide variation in exposure 3343 from dust reported in the literature (see above), this estimate is within the range of those reported. 3344 The CONTAM Panel noted that this is only a very crude estimate of the exposure via dust and has a 3345 large associated uncertainty, and that there is a wide range in estimates of exposure from dust reported 3346 in the scientific literature. It is nevertheless evident that for all population groups, exposure from dust 3347 could make a substantial contribution to the overall HBCDDs exposure.



A further so far not intensively studied potential oral exposure can arise from unintentional ingestion of parts of plastic toys by small children. Fatunsin et al. (2020) analysed 23 plastic samples from 20 new and second-hand children's toys that had been previously shown to be bromine positive by XRF. HBCDDs wer detected in 14 cases with concentrations between 0.25 and 840 mg/kg. Besides exposure from mouthing, exposure arising from accidental ingestion of plastic from toys can be significant for young children.

#### 3354 *Dermal exposure*

Pawar et al. (2017) investigated the dermal bioaccessibility of flame retardants from indoor dust and 3355 the influence of topically applied cosmetics. The authors performed an *in vitro* physiologically based 3356 extraction test of various BFRs from indoor dust to a synthetic sweat/sebum mixture (SSSM). The SSSM 3357 3358 was prepared using over 25 different chemical components. In general, the bioaccessibility of HBCDDs 3359 increased with increasing sebum content of the SSSM. At 100% sweat, the bioaccesibility of y-HBCDD 3360  $(1.4 \pm 0.1\%)$  was less than that of  $\beta$ -HBCDD  $(1.6 \pm 0.6\%)$  and  $\alpha$ -HBCDD  $(2.3 \pm 0.2\%)$ . However, the reverse trend was observed at 100% sebum, where the bioaccessibility was highest for y-HBCDD (67.2 3361 3362 ± 3.37%), followed by  $\beta$ -HBCDD (60.4 ± 10.1%) and  $\alpha$ -HBCDD (50.5 ± 7.0%). This behaviour is 3363 consistent with the lower water solubility of the y-stereoisomer compared with that of  $\beta$ -HBCDD and  $\alpha$ -3364 HBCDD. When tested, 6 mg of cosmetics (moisturising cream, sunscreen lotion, shower gel and body 3365 spray) were each examined separately. The presence of cosmetics decreased the bioaccessibility of HBCDDs from indoor dust, whereas shower gel and sunscreen lotion enhanced the bioaccessibility of 3366 HBCDD. 3367

3368 Miyake et al. (2010) performed a comparative study of human health risks posed by a flame retarded 3369 curtain with HBCDDs. The concentrations of  $\alpha$ -HBCDD,  $\beta$ -HBCDD,  $\gamma$ -HBCDD, and  $\Sigma$ HBCDDs in the curtain were 340,000, 150,000, 1,000,000, and 1,500,000 ng/g, respectively. The amounts of  $\Sigma$ HBCDDs, 3370 3371 determined after drawing the curtain in triplicate, on the hand were 0.9±0.6 ng. The lifetime average 3372 daily dose (LADD) by the dermal exposure via direct contact with the curtain was calculated as 3373 0.0000025 mg/kg bw. The LADD by the ingestion exposure via hand-to-mouth contact assigning default 3374 values, such as contact frequency with the surface, hand-to-mouth events, dermal absorption efficiency, and body weight, was calculated as  $2.3 \times 10^{-9}$  mg/kg bw. This value was a 100-fold lower than the 3375 3376 LADD by the inhalation exposure of dust.

3377 Tay et al. (2018) assessed dermal exposure to BFRs and compared direct measurements from hand 3378 wipes with an indirect estimation from settled dust concentrations. Sixty-one hand wipe samples were 3379 collected from a Norwegian adult cohort using gauze pads immersed in isopropanol. The range (median) 3380 of  $\Sigma$ HBCDDs in the hand wipe samples were 49–8,900 (180) ng per participant, corresponding to 27– 3381 11,000 (640) pg/cm<sup>2</sup> hand surface. A positive statistically (p<0.05) correlation was found for y-HBCDD 3382 between settled dust (ng/g) and hand wipe (ng/participant). Concentrations of  $\Sigma$ HBCDDs in hand wipes 3383 were positively correlated (p < 0.05) to the number of electronic consumer products at home (i.e. TV, 3384 DVD players, tablets, laptops, cell phones and PC screens). Positive correlations were also found between the number of people living in the house and the concentrations of  $\alpha$ - and  $\beta$ -HBCDDs in hand 3385 wipes. The median (P95) daily dermal exposure to  $\Sigma$ HBCDDs was estimated as 840 (12,000) pg/kg bw 3386 3387 per day.

An association of BFRs between children's hand wipes and house dust was also demonstrated by Stapleton et al. (2014). The  $\Sigma$ HBCDD levels in 43 hand wipes of children ranged from <0.05–10.8 ng with a geometric mean of 0.97 ng. In the 30 house dust samples, the  $\Sigma$ HBCDD concentrations were determined as 77.6–2,658 ng/g with a geometric mean of 338 ng/g. In general, higher levels of BFRs in house dust were associated with higher handwipe levels.



Another potential source of dermal intake is the contact with used dishcloths. Gallistl et al. (2017) analysed 19 dishcloths after use for 14 days in kitchens for a number of polyhalogenated compounds. The median surface concentration for total-HBCDDs was determined as 550 (P95: 5,300) ng/m<sup>2</sup>. Based on a skin absorption factor of 11–13%, a median intake of 5.4 (P95: 53) ng per day via skin was estimated.

3398 In summary, the data on intake of HBCDDs via dust and dermal contact indicate that these routes of 3399 exposures may be substantial, especially for small children and toddlers and must be considered when 3400 total exposure to HBCDDs are assessed as they can even exceed the dietary exposure.

# 3401 **3.4.** Risk characterisation

The chronic human dietary intake that would lead to a body burden at the LOAEL of 0.9 mg/kg bw was estimated (see **Section 3.1.5.2**). The MOE was calculated by dividing this intake by the estimated dietary exposure (see **Section 3.3.1**). Although the LOAEL was identified from a study involving a single administration on PND10, it is not viewed as an acute effect because HBCDDs are persistent in the body. PND10 marks the start of a critical period in the development of the rodent brain, which corresponds in humans to the period beginning in the third trimester of pregnancy and continues throughout the first 2 years of life.

As described in **Section 3.1.5.3**, an MOE higher than 24 would indicate a low health concern. The MOEs were calculated for all age groups across all food consumption surveys in the EFSA Consumption Database (see **Table 17**). The lowest MOE was 650, which was calculated from the UB high level dietary exposure in the toddler age group. The CONTAM Panel concluded that these MOE values do not raise a health concern.

			MEA	N			
	M	EN	Med	ian	MAX		
	LB	UB	LB	UB	LB	UB	
Toddlers	17,000	3,700	7,100	2,400	3,000	1,600	
Other children	21,000	6,400	8,400	3,100	3,700	1,900	
Adolescents	34,000	12,000	14,000	6,600	7,000	3,700	
Adults	29,000	14,000	15,000	8,100	8,400	5,300	
Elderly	24,000	12,000	15,000	7,800	10,000	5,900	
Very elderly	34,000	12,000	16,000	7,800	7,800	4,800	
		7	P95	;			
Toddlers	4,100	1,500	2,600	1,100	1,000	650	
Other children	6,400	2,300	3,000	1,400	1,200	800	
Adolescents	10,000	4,600	5,300	2,700	2,000	1,500	
Adults	8,100	4,900	5,100	3,200	2,000	1,700	
Elderly	7,800	4,700	4,800	3,200	3,200	2,600	
Very elderly	7,800	5,200	5,300	3,200	3,000	2,400	

**Table 17.** Margin of exposure (MOE) values across age groups <sup>(a)</sup> (rounded to two significant figures)

(a): It was not possible to assess the dietary exposure to HBCDDs for 'Infants' (not breastfed) using the data submitted to EFSA
 (see Section 3.3.1).

3417 The CONTAM Panel noted that exposure to HBCDDs via dust and dermal contact can be an additional

source of exposure to HBCDDs especially for children and can exceed dietary exposure (see **Section 3419 3.3.3**).



For breastfed infants with average human milk consumption (800 mL per day) the LB and UB range of 3420 3421 occurrence in pooled human milk samples would result in daily exposures (median, range) of 13.8 (2.3-3422 73.4) ng/kg bw and 14.3 (3.2–73.7) ng/kg bw, respectively. The chronic human dietary intake that 3423 would lead to a body burden at the LOAEL of 0.9 mg/kg bw was estimated (see Section 3.1.5.2). The 3424 MOE was calculated by dividing this intake by the exposure resulting from breastfeeding (see Section 3425 3.3.1). This resulted in MOE values of 170 (1,022–32) and 164 (734–32), respectively. For infants with 3426 high milk consumption (1,200 mL per day), the LB and UB intakes are 20.7 (3.4–110.2) ng/kg bw and 3427 21.5 (4.8-110.6) ng/kg bw, respectively. This resulted in MOE values of 114 (691-21) and 109 (490-3428 21), respectively. The lowest MOE values for breastfed infants with high milk consumption and highest 3429 breast milk levels are below the value of 24 mentioned above. It was noted that since these were pooled 3430 samples, it was not possible to estimate specific values for individuals and MOE values will be lower for 3431 some of them. The Panel concluded that these MOE values may raise a health concern for some 3432 breastfed infants.

#### 3433 Comparison of body burdens in adults

3434 In the previous Opinion on HBCDDs (EFSA CONTAM Panel, 2011a), the CONTAM Panel compared the body burden at the Reference Point calculated from animal studies with the estimated body burdens in 3435 3436 humans as an additional approach to assess the health risk of exposure to HBCDDs in addition to the 3437 MOE calculated for exposure via the diet. For this comparison, the Panel considered at that time the 3438 available data on levels in human tissues, and in particular the levels in adipose tissue reported in the 3439 literature, as they were considered to best reflect long-term exposure to HBCDDs. The reported 3440 concentrations in adipose tissue in humans were converted to an overall body burden assuming an 3441 average fat content in adult women of 25% (van der Molen, 1998).

Following a similar approach, for the current assessment the CONTAM Panel used the HBCDD results 3442 3443 from the study of Malarvannan et al. (2013a) in visceral and abdominal fat of obese patients. In addition, 3444 serum data on HBCDDs from the study of Kalantzi et al. (2011) were converted to overall body burden. 3445 As the HBCDD levels in human breast milk are likely to be related to the body burden (on a lipid basis) 3446 of women of child-bearing age, the reported UB range for HBCDDs in human milk collected in European 3447 countries between 2014 and 2016 as part of the WHO/UNEP field studies were also converted to overall 3448 body burden. The CONTAM Panel is aware that the body fat content is dependent inter alia on gender, age and physical condition. Deurenberg et al. (2001) reported mean body fat concentrations of  $31.2 \pm$ 3449 3450 7.8% and 20.1  $\pm$  7.6% for females (n = 234) and males (n = 182), respectively. In order to compare 3451 the results of the present assessment with the former HBCDDs Opinion (EFSA CONTAM Panel, 2011a), 3452 an average fat content of 25% was used. The results are depicted in Table 18. The MOE values were 3453 calculated by dividing the body burden at the LOAEL of 0.747 mg/kg bw by the estimated body burden based on the different human matrices. 3454

Table 18. Overall body burden and margin of exposure (MOE) values for adults calculated on the basis
 of various human matrices (MOE values rounded to two significant figures)

Basis	Concentration range (medians) (ng/g lipid)	Body burden (µg/kg bw)	MOE
Adipose tissue	0.89–89 (4.1/3.7) <sup>(a)</sup>	0.223–22.3 (1.03/0.93)	3,400–34 (730, 810)
Serum	0.49–38.8 (1.32) <sup>(b)</sup>	0.123–9.7 (0.33)	6,100–77 (2,300)
Human milk	0.7–16.1 (3.12) <sup>(c)</sup>	0.175–4.03 (0.78)	4,300–190 (960)

3457 (a): Data from Malarvannan et al. (2013a). Medians are for visceral and abdominal fat, respectively.

3458 (b): Data from Kalantzi et al. (2011).

(c): UB range for HBCDDs in pooled human milk collected in European countries between 2014 and 2016 as part of the WHO/UNEP

field studies (see **Table 8**).



The CONTAM Panel concluded that these results support the conclusion that current dietary exposure to HBCDDs in European countries does not raise a health concern.

# 3463 **3.5.** Uncertainty analysis

The evaluation of the inherent uncertainties in the assessment of exposure to HBCDDs in food has been performed following the guidance of the Opinion of the Scientific Committee related to Uncertainties in the Dietary Exposure Assessment (EFSA, 2007). In addition, the report on 'Characterizing and Communicating Uncertainty in Exposure Assessment' has been considered (WHO/IPCS, 2008). The new guidance on uncertainties of the Scientific Committee (EFSA SC, 2018) was not implemented in this Opinion as the draft was developed before the implementation of the guidance by the CONTAM Panel in its risk assessments.

3471 3.5.1. Assessment objectives

3472 The objectives of the assessment were clarified in **Section 1.2.** on Interpretation of the Terms of 3473 Reference.

3474 3.5.2. Exposure scenario/exposure model

### 3475 Occurrence data

The exposure assessment was based on HBCDDs occurrence data reported by seven European countries. However, most of the data (78%) were reported by France and Germany. There is uncertainty around possible regional differences in HBCDDs contamination and the data set is likely not to be fully representative for the EU market.

3480 Exclusion of the occurrence data reported as Total HBCDDs (analysed by GC-MS) due to the high LOQ 3481 values and high proportion of left-censored data resulted in excluding three food categories: 'Vegetable 3482 oils', 'Ready-to-eat meal for infants and young children' and 'Dietary supplements', which may result in 3483 underestimation of exposure. In case of occurrence data expressed on a fat weight basis, the 3484 contamination level was combined with the fat content as reported for the related sample. Where the fat content was missing, the median value of the samples reported for the same category was used. 3485 3486 This could lead to both over- and underestimation of the exposure. In addition, exclusion of the food 3487 categories 'Infant formulae', 'Follow-on formulae' and 'Cereal-based food for infants and young children' 3488 due to the data reported for the individual stereoisomers being 100% left-censored, may lead to an underestimation of dietary exposure of 'Toddlers' to HBCDDs. This did not allow to estimate the 3489 exposure of 'Infants' from food. 3490

3491 A high proportion of left-censored data (71%) was reported across all food categories including the 3492 main contributors to the dietary exposure and consequently the difference between LB and UB estimates 3493 is 2- to 3-fold. The use of the LB in this Opinion tends to underestimate, while the UB tends to 3494 overestimate the dietary exposure. The limited number of available analytical results for some food 3495 categories adds uncertainty to the representativeness of the mean concentration values used to 3496 estimate the exposure. For some highly consumed food groups ('animal fat', 'liquid milk') the sensitivity 3497 of the methods seems too low, resulting in a large uncertainty in the exposure from these food groups, thereby increasing the difference between LB and UB exposure. 3498

Exposure to HBCDDs from food supplements containing special fatty acids (e.g. fish oil) could not be considered due to lack of occurrence data. Moreover, the contribution from several food categories, especially of plant origin, could not be considered due to lack of data. This will result in an underestimation of exposure.



#### 3503 Consumption data

Uncertainties and limitations related to the use of the EFSA Comprehensive Food Consumption Database have already been described by EFSA (EFSA, 2011a) and are not further detailed in this Opinion. Generally, these limitations and uncertainties relate to the use of different dietary survey methodologies, standard portion sizes, representativeness of samples included in surveys, or to the inclusion of consumption surveys covering only few days to estimate high percentiles of chronic exposure.

#### 3509 Dietary Exposure Assessment

3510 Important sources of uncertainty are related to the different assumptions done, mainly for the linkage 3511 between the occurrence and consumption data. In particular:

- 3512 Sampled foods were assumed to represent consumed foods, which could lead to both over- and
   3513 underestimation of the exposure.
- A number of foodstuffs were grouped mainly at the 2nd level of the FoodEx system, assuming homogeneity of the contamination levels. This was the case for food categories represented with less than six samples or 100% left-censored at the third level, but enough data were available at the upper level. This could lead to both over- and underestimation of the exposure.
- For the samples reported as 'Salmon and trout (Salmo spp.)' without additional information on the species, and therefore the reclassification not possible, homogeneity of the contamination was assumed. This could lead to both over- and underestimation of the exposure.

3521 Chronic dietary exposure is based on the sum of the three major HBCDD stereoisomers ( $\alpha$ -,  $\beta$ - and  $\gamma$ -3522 HBCDD) for all food categories. Due to the exclusion of  $\delta$ - and  $\epsilon$ -HBCDD, exposure might be slightly 3523 underestimated. The CONTAM Panel noted that these stereoisomers  $\delta$ - and  $\epsilon$ -HBCDD are only present 3524 at low concentrations in the technical product, and in food.

The exposure of breastfed infants was estimated based on pooled samples from several European 3525 3526 countries, showing a wide range of concentrations. Since these were pooled samples, it was not possible 3527 to estimate the distribution within the population. Some breastfed infants showed an exposure that 3528 resulted in an MOE smaller than 24, based on high milk intake and HBCDD levels in pooled milk. This 3529 may imply a health concern, also considering that HBCDD levels in milk of individual mothers are likely 3530 to be higher than that in the pooled sample. This higher exposure is caused by the body burden of the 3531 mothers due to chronic exposure. Smaller MOEs in these infants may be in contrast to the large MOEs 3532 for adults, even at the P95 (1,700-8,100). The transfer to infants should be taken into account when 3533 deriving the safe intake level in humans but there is currently too little information to base the risk 3534 assessment on the body burdens of mothers and transfer to the infant.

3535 Effect of cooking/processing was not taken into account. The human exposure estimations in this 3536 Opinion are based on consumption data and occurrence levels in raw food commodities. It is known 3537 that typical household cooking practices neither lead to degradation nor to generation of appreciable 3538 amounts of HBCDDs. However, changes in the fat content of food commodities during 3539 cooking/processing practices may lead to changes of the lipophilic contaminants in the processed food compared to the raw food commodity. Due to lack of representative data on these possible changes the 3540 3541 effect of cooking and processing could not be taken into account. This may have added to the 3542 uncertainty in the exposure estimation to some extent.

3543 **3.5.3.** Hazard identification and characterisation

No data were identified regarding absorption in human, and sparse data were found on distribution and metabolism.



Only one study reported on the terminal elimination half-life of HBCDDs in humans. Geyer et al. (2004, extended abstract) estimated a half-life of 64 days (range 23–219 days) by using several estimations (daily intake, fat mass). However, the method is not fully described. This leads to uncertainties regarding the human half-life value as follows:

- The authors indicated that 'the whole body (total body burden) half-lives in humans (t½H in days) of BFRs were estimated from the daily intake (DI in ng per day -1) and the total body burden under steady state conditions in non-occupationally exposed adult humans". It is not clear for the CONTAM Panel if the estimate of daily intake of HBCDDs corresponded to the same population in which HBCDD concentrations were measured.
- The demographic characteristics (number of subjects, weight, age, gender, etc.) are not provided. According to the equation used by Geyer et al. (2004, extended abstract), the human half-life was estimated by considering a fat mass value (m<sub>f</sub>) of 13.5 kg for adult man and 18.7 kg for an adult woman:

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$$t_{\rm 15H} = \frac{\ln 2 \times m_{\rm ss}}{DI \times f} = \frac{0.693 \times c_{\rm ss} \times m_f}{DI \times f}$$

3563Where f is the fraction of dose absorbed from food,  $m_{ss}$  is the total amount (in ng) of the3564chemical in the whole human body at steady-state,  $c_{ss}$  is the concentration (ng/kg lipid) in adult3565humans, and  $m_f$  is the fat mass (13.5 kg for an average adult man and 18.7 kg for an average3566adult woman).

- The authors assumed that the HBCDD concentrations were converted into total body burdens assuming a balance between all lipid compartments in the body. Nevertheless, the fat content of the human body varies with age, sex, weight and height, as previously described by Deurenberg et al. (1991). The CONTAM Panel noted that information about the demographic characteristics data on the subject used for the estimation was not available.
- In their equation, Geyer et al. (2004, extended abstract) used as internal dose metric the concentration (ng/kg lipid) in adult humans measured in breast milk. Further information on how the calculation of the half-life was performed is not available.

As noted in the previous Opinion (EFSA CONTAM Panel, 2011a), most of the toxicological studies with HBCDDs were performed with HBCDDs preparations for which the purity and stereoisomer composition was not always indicated, rather than with purified individual stereoisomers. As the toxicity of the individual stereoisomers is not known, and the toxicological Reference Point is based on information from exposure to HBCDDs as mixtures of isomers with unclear relevance to the distribution of isomers in food, this adds a considerable uncertainty to the risk assessment.

The studies on neurobehavioural effects of HBCDDs all have limitations, which introduce uncertainty into the Reference Point. The studies of Eriksson et al. (2006) and Zhang et al. (2017a) did not assess anxiety and swim speed, although these parameters might also explain the increase in latency to reach the platform in the Morris Water Maze. Thus, there is uncertainty as to whether HBCDDs have effects on spatial learning and memory. Changes in the anxiety level and horizontal locomotion could also explain observed effects on spontaneous behaviour as recorded in Eriksson et al. (2006).

In the Eriksson et al. (2006) study a single dose of HBCDDs was administered to mice on PND10, which marks the start of a critical period in the development of the rodent brain. It is unclear whether PND10 is the most critical day and if exposure at another time point would produce a response at a lower dose. There is also uncertainty as to whether an effect would have been observed at a lower dose following



3591 repeated dosing. However, the current assessment was done based on body burden, and repeated 3592 dosing even at lower levels might have resulted in a higher body burden.

3593 Although occurring at a dose (49.5 µg/kg bw per day) below that used to derive a Reference Point, the 3594 CONTAM Panel chose not to further consider effects on sex hormones due to a lack of clear dose 3595 response and information on related apical effects at the LOEL (Maranghi et al., 2013; Rasinger et al., 3596 2014, 2018). Likewise, effects on metabolism, and in particular lipid metabolism, were observed at a 3597 dose of 7.1 µg/kg bw per day (Xie et al., 2019), but because of uncertainty regarding relevance to 3598 human health, this endpoint was also not selected for risk characterisation. There is remaining 3599 uncertainty regarding the relevance of these findings and therefore of the selected endpoints for establishment of a Reference Point. 3600

Exposure of adult rats caused an increase in hepatic glucuronidation of T4 hormone (van der Ven et al., 2006). Whilst increased T4-UGT was only significant at a relatively high dose of 100 mg/kg per day, the authors reported a BMDL<sub>10</sub> of 4.1 mg/kg per day (van der Ven et al., 2006). There is an uncertainty as to whether this effect on T4 conjugation might be different in early infancy and, thus, whether effects on neurodevelopment might be secondary to changes in thyroid hormone homeostasis. Furthermore, there is uncertainty about the relevance of glucoronidation for thyroid hormone homeostasis in humans.

The CONTAM Panel conducted BMD modelling on the neurobehavioural data of Eriksson et al. (2006) according to the latest EFSA guidance (EFSA SC, 2017), and found that the BMD confidence intervals were relatively wide and that the the BMDLs for horizontal locomotion and total activity were far below the lowest dose administered. Therefore, the NOAEL/LOAEL approach was applied, since the BMD modelling generally indicated that determination of the Reference Point based on the key data is uncertain.

Regarding the available epidemiological evidence, the assessed evidence base is characterised by relatively small studies, heterogenous populations and lack of replicated endpoints (no studies identified on the same endpoint). Therefore, there is uncertainty as to whether the observed epidemiological associations are causal.

3617 **3.5.4.** Summary of uncertainties

3618 In **Table 19**, a summary of the uncertainty evaluation is presented, highlighting the main sources of 3619 uncertainty and indicating an estimate of whether the respective source of uncertainty might have led 3620 to an over- or underestimation of the exposure or the resulting risk.

3621 **Table 19.** Summary of qualitative evaluation of the impact of uncertainties on the risk assessment of3622 HBCDDs in food

Sources of uncertainty	Direction				
Hazard identification and characterisation					
Limited information on the stereoisomer composition and purity of the HBCDDs used in toxicological studies	+/-				
Isomeric profiles of HBCDDs used in toxicological studies are not the same as the profiles found in food	+/-				
Limited data on the toxicokinetics of HBCDDs in humans, including uncertainty in the data on the half-life of HBCDDs in humans	+/-				
Lack of information on the toxicity of the individual stereoisomers	+/-				
Whether exposure on a day other than PND10 in mice would produce a response at a lower body burden	-				
Whether repeated exposure of HBCDDs will produce neurobehavioural effects in rodents at a lower body burden	-				

Lack of large-scale prospective epidemiological data to support the toxicological effects	+/-
Determination of the Reference Point based on the key data is uncertain	+/-
ccurrence and exposure assessment	
Uncertainty of analytical results due to different LODs/LOQs	+/-
Some food categories could not be considered due to lack of occurrence data	-
Extrapolation of occurrence data from a few countries to whole Europe	+/-
Impact of upper-bounds for non-detects on dietary exposure estimate	+
Distribution of HBCDDs levels in individual human milk samples is not known because only pooled samples analysed across Europe	+/-
Lack of information on the impact of food processing	+/-
Consumption data: different methodologies / representativeness / underreporting / misreporting / no portion size standard	+/-

(a) +=uncertainty with potential to cause over-estimation of exposure/risk; - =uncertainty with potential to cause underestimation of exposure/risk.

3625 The CONTAM Panel considered that the impact of the uncertainties on the risk assessment of HBCDDs in food is substantial. Without a quantitative uncertainty analysis, the Panel was unable to determine 3626 3627 the direction of the overall uncertainty.

#### 4. Conclusions 3628

3629 HBCDDs, composed of mixtures of stereoisomers of 1,2,5,6,9,10-hexabromocyclododecane, were 3630 widely used as additive flame retardants in a number of applications. Due to the widespread use, 3631 HBCDDs have found global distribution. HBCDDs were permitted to be used in the EU until August 2015, after which only authorised applications were allowed because of health concerns. 3632

3633 The present assessment is an update of the former EFSA CONTAM Panel Opinion on HBCDDs published 3634 by EFSA in 2011. It takes into account the occurrence data in food and biological samples submitted 3635 after the publication of the former Opinion, as well as the newly available scientific information on 3636 hazard identification and characterisation.

3637 The analytical determination of HBCDDs is performed either by GC/MS or LC/MS based methods. Both 3638 techniques enable the use of isotope-labelled internal standards, which allow for the correction for losses 3639 during extraction and clean-up. While GC/MS based methods determine Total HBCDDs as they cannot 3640 separate the stereoisomers, LC/MS based methods allow the specific analysis of the individual HBCDD 3641 stereoisomers.

#### 4.1. Hazard identification and characterisation 3642

#### 4.1.1. **Toxicokinetics** 3643

- 3644 In rodents, absorption of HBCDD stereoisomers from the gastrointestinal tract is rapid and 3645 almost complete (>80%).
- 3646 HBCDDs and/or their metabolites are distributed to a number of tissues, including adipose • 3647 tissue, muscle, liver, skin and brain. Following a single oral administration in rats,  $\alpha$ -HBCDD and 3648 β-HBCDD accumulate in adipose tissue, while γ-HBCDD does in liver and adipose tissue. In 3649 repeated dose experiments in mice,  $\alpha$ -HBCDD and  $\beta$ -HBCDD accumulate mainly in adipose tissue, while y-HBCDD accumulates in the liver. 3650
- 3651 Metabolic debromination and hydroxylation of HBCDDs have been reported.



3652 3653	•	Conversion of $\gamma\text{-HBCDD}$ to $\alpha\text{-}$ and $\beta\text{-HBCDD}$ has been reported, but no stereoisomerisation of $\alpha\text{-HBCDD}.$
3654 3655 3656	•	Elimination of HBCDD stereoisomers in rodents is predominantly via faeces. The elimination half-lives of HBCDD stereoisomers in female mice, based on concentrations in adipose tissue, vary from 3–4 days for $\gamma$ -HBCDD, 2.5 days for $\beta$ -HBCDD, and 17 days for $\alpha$ -HBCDD.
3657 3658 3659	•	In humans, in the only available study, the half-life was estimated to be 64 days (range 23–219 days) for the sum of $\alpha$ -, $\beta$ - and $\gamma$ -HBCDD stereoisomers. This difference in kinetics affect the extrapolation of animal data to humans.
3660 3661 3662	•	Transfer of HBCDDs was studied in different animal species (laying hens, broilers, ducks, pigs and fish). HBCDDs are distributed to a number of tissues and accumulate in tissues with high fat content and eggs.
3663	4.1.2.	Toxicity in experimental animals
3664	•	The acute toxicity of HBCDDs is low.
3665 3666 3667 3668 3669	•	Toxicological studies have been carried out using different experimental designs with single or repeated administration during gestation, postnatally or in adulthood using, in most cases, HBCDDs with no information on the stereoisomer composition specified. Main targets for toxicity are neurodevelopment, the liver, thyroid hormone homeostasis and the reproductive and immune systems.
3670 3671 3672	•	HBCDDs induce increased liver weight and hepatocellular hypertrophy in rats and mice. Increased thyroid weight, follicular hypertrophy, hyperplasia and/or colloid depletion and changes in thyroid hormone levels were also reported in these species.
3673 3674 3675 3676 3677	•	HBCDDs decreased the fertility index and increased pup mortality during lactation in a 2- generation toxicity study in rats. In addition, HBCDDs decreased testes weight and delayed vaginal opening in female pups in a 1-generation study in rats. Other endocrine-related effects are changes in levels of sex-steroid hormones (testosterone and oestradiol) in mice, decreased number of primordial ovarian follicles or reduction in the number of growing follicles in rats.
3678 3679 3680 3681	•	HBCDDs affected the immune system. Changes observed included decreased thymus weight and increased lesions in the thymus, reduced splenocyte proliferation and increased spleen lesions, decreased T-cell and increased B-cell populations, increased NK-cells, and changes in immunoglobulin responses.
3682 3683 3684	•	Exposure of juvenile rats and mice to HBCDDs caused neurodevelopmental effects on behaviour, leading to changes in spontaneous behavior, spatial learning and memory as detected in adulthood.
3685 3686	•	HBCDDs are not genotoxic <i>in vitro</i> or <i>in vivo</i> . The slight induction of DNA strand breaks observed in some <i>in vitro</i> tests is most likely due to oxidative stress.
3687 3688	•	HBCDDs were previously judged not to be carcinogenic in mice and no new carcinogenicity studies have been identified. The available evidence indicates that HBCDDs are not carcinogens.
3689	4.1.3.	Observations in humans
3690 3691 3692 3693	•	A growing number of epidemiological publications were identified assessing the association between exposure to HBCDDs and birth weight/length, neurodevelopment and thyroid dysfunction in children, as well as subfertility, type 2 diabetes, thyroid hormone levels, severe endometriosis (including ovarian endometrioma) and breast cancer metastasis in adults.



- None of the effects studied in the longitudinal studies and assessing internal exposure either
   reached statistical significance or were replicated in a longer follow-up point in the same study.
   Considerable limitations exist pertaining to small sample sizes, varying methodological quality,
   effect inconsistency and considerable heterogeneity in the assessed populations, exposures,
   and endpoints.
- Adverse effects of HBCDDs related to neurodevelopment have been assessed in two
   epidemiological studies; the low volume of prospective data, the differing endpoint measures
   and the lack of replication render these data insufficient for use in risk characterisation.
- 3702 **4.1.4.** Mode of action
- Effects of HBCDDs on the liver of rodents appear to involve CAR and PXR. Mitochondrial energy production is reduced with downstream effects on reactive oxygen species (ROS) production, Na<sup>+</sup>/K<sup>+</sup>-ATPase and Ca<sup>2+</sup>-ATPase activities and increased cytosolic [Ca<sup>2+</sup>] and [Zn<sup>2+</sup>].
- HBCDDs stimulate proliferation and migration of liver cell lines at picomolar to nanomolar
   concentrations and there is evidence that these effects are related to activation of the estrogen
   receptor and the PI3K/Akt/mTOR signalling pathway.
- Oral HBCDD exposure of mice fed a high fat diet aggravates metabolic dysfunction through
   modifications of lipid and glucose homeostasis.
- HBCDD-induced lipid accumulation in liver and adipose tissue appears to be associated with
   suppression of the Wnt/β-catenin pathway resulting in increased expression of *Pparg* and its
   adipogenic target genes.
- Mechanistic studies on the effects of HBCDDs on the nervous system support specific effects on dopaminergic and glutamatergic neurons. This appears to include inhibition of dopamine and glutamate reuptake following synaptic release, and deregulation of Ca<sup>2+</sup> and Zn<sup>2+</sup>. The brain could also be affected by HBCDDs through diminished responsiveness to thyroid hormones.
- HBCDDs may affect the synthesis of sex steroids hormones by interfering with cAMP-dependent
   cholesterol uptake and by causing dysregulation of enzymes involved in sex steroid metabolism.
   HBCDDs may also alter the response of tissues to sex steroids.
- The mechanism(s) involved in HBCDD-induced ROS production may be related to impairment
   of mitochondrial energy production.
- 3723 4.1.5. Critical effects and dose-response analysis
- The evidence from the available human data was not sufficient to base the risk assessment on.
   Thus the data from studies on experimental animals were used to identify a Reference Point for
   the human health risk characterisation.
- The CONTAM Panel concluded that the neurodevelopmental effects on behaviour can be considered the critical effect for the risk characterisation.
- The BMD modelling on the neurobehavioural data showed BMD confidence intervals that were
   relatively wide and that the BMDLs for horizontal locomotion and total activity were far below
   the lowest dose administered. Therefore, the NOAEL/LOAEL approach was applied.
- Based on effects on spontaneous behaviour (horizontal locomotion, rearing and total activity), the CONTAM Panel identified a LOAEL of 0.9 mg/kg bw (single dose) as the Reference Point for the risk assessment of HBCDDs.



3735 3736 3737	•	Because the elimination kinetics for HBCDDs between mice and humans differ, the CONTAM Panel first calculated a body burden of 0.75 mg/kg bw at the LOAEL, considering an oral absorption of $83\%$ in mice.				
3738 3739 3740	•	The chronic intake that would lead to the same body burden in humans was calculated assuming an absorption in humans of 100% and the longest half-life identified in humans for HBCDDs of 219 days. This resulted in an estimated chronic human dietary intake of 2.35 $\mu$ g/kg bw per day				
3741 3742 3743	•	Due to limitations in the database on HBCDDs, the derivation of a health-based guidance value (HBGV) was not considered appropriate. Instead, the margin of exposure (MOE) approach was applied to assess possible health concerns.				
3744	•	The CONTAM Panel considered that an MOE higher than 24 would indicate a low health concern.				
3745	4.2.	Occurrence and exposure for the European population				
3746	4.2.1.	Occurrence in food				
3747 3748 3749	•	A total of 6,857 analytical results for HBCDDs in food fulfilled the quality criteria applied and from these 6,352 were selected and used in the assessment after grouping the categories and matching with the food consumption data.				
3750 3751 3752	•	Contrary to the data used in the previous Opinion on HBCDDs published in 2011, where most of the data were analysed with GC/MS and reported Total HBCDDs, the current data have mostly been analysed by LC/MS and reported on the specific stereoisomers $\alpha$ -, $\beta$ - and $\gamma$ -HBCDD.				
3753 3754 3755	•	The high proportion of left-censored data, partially due to relatively high LOQs, resulted in a large difference between the lower bound (LB) and upper bound (UB) occurrence values for many food groups.				
3756 3757 3758	•	The highest mean concentrations of HBCDDs (sum of $\alpha$ -, $\beta$ - and $\gamma$ -HBCDD) were recorded for the food category 'Fish meat', with the highest mean concentrations in eel, being 2.33 and 2.35 µg/kg ww at LB and UB, respectively.				
3759	4.2.2.	Exposure assessment				
3760 3761 3762 3763 3764	•	The mean dietary exposure estimates for HBCDDs ranged from 0.07 (minimum LB)/0.17 (minimum UB) to 0.79 (maximum LB)/1.52 (maximum UB) ng/kg bw per day across dietary surveys and age groups. At the 95th percentile, dietary exposure estimates ranged from 0.23 (minimum LB)/0.45 (minimum UB) to 2.30 (maximum LB)/3.61 (maximum UB) ng/kg bw per day.				
3765 3766	•	The high proportion of left-censored data, partially due to relatively high LOQs, resulted in a 2- to 3- fold difference between the LB and UB exposure estimates.				
3767 3768	•	The most important contributors to the chronic dietary LB exposure to HBCDDs were 'Fish meat', 'Eggs, fresh', 'Livestock meat' and 'Poultry'.				
3769 3770 3771 3772 3773 3774	•	The exposure scenario based on average human milk consumption and the reported UB range for HBCDDs (sum of $\alpha$ -, $\beta$ - and $\gamma$ -HBCDD) in pooled human milk samples collected in European countries between 2014 and 2016 as part of the WHO/UNEP field studies, would result in a median daily exposure of 14.3 ng/kg bw (range: 3.2–73.7 ng/kg bw). For infants with high human milk consumption the median daily exposure would result in 21.5 ng/kg bw (range 4.8–110.6 ng/kg bw).				



- Non-dietary exposure through intake of dust and dermal contact in homes, offices, schools, cars
   and public environment can substantially contribute, and in some cases even dominate the total
   human exposure to HBCDDs, especially for toddlers and other children.
- 3778 **4.3.** Risk characterisation
- MOE values were calculated by comparison of the calculated chronic human dietary intake of 2.35 µg/kg bw per day, leading to the body burden at the LOAEL, with the estimated dietary exposure for the different population groups. The MOE values obtained ranged from 34,000 to 650. These MOEs are larger than 24 and the CONTAM Panel concluded that they do not raise a health concern.
- The CONTAM Panel also compared the body burden of 0.75 mg/kg bw at the LOAEL with the body burdens in adults based on levels in adipose tissue, blood and milk reported in the literature. The results support the conclusion that current dietary exposure to HBCDDs in European countries does not raise a health concern.
- For breastfed infants, the lowest MOE values for high milk consumption are below the value of
   24. The CONTAM Panel concluded that these MOEs may raise a health concern for some
   breastfed infants.

### 3791 **5.** Recommendations

3792 In order to improve the risk assessment and reduce the uncertainties, the CONTAM Panel made the 3793 following recommendations:

- 3794 Criteria for the analysis of HBCDD stereoisomers should be set. • 3795 Surveillance of HBCDD stereoisomers in food, in particular in food groups for infants and small ٠ 3796 children, should continue in order to refine the exposure estimates. More data on occurrence of HBCDDs in human milk are needed to enable a more robust 3797 3798 exposure assessment for breastfed infants. Improved information on toxicokinetics, e.g. half-life values, of HBCDD stereoisomers in humans 3799 3800 is needed. More information is needed on the body burden in mothers and relation to transfer 3801 of HBCDDs to milk. This information should be used to develop a toxicokinetic model for HBCDDs. 3802 More information is needed on the transfer into ruminant meat and milk. 3803 3804 Further toxicological studies should be conducted with individual HBCDD stereoisomers most • 3805 relevant to human exposure. Studies on neurodevelopment (after repeated exposure), including investigations of the 3806 • mechanisms and mode of action involved, are recommended. 3807
- Studies on reproductive effects are recommended.
- Studies on possible diabetogenic and obesogenic effects are recommended.
- Longitudinal epidemiological studies of sufficient power and appropriate exposure and co exposure assessment are needed.



### 3812 **Documentation provided to EFSA**

Malisch R and Schächtele A, 2019. Data provided to EFSA by Rainer Malisch and Alexander
 Schächtele on the HBCDDs-related results of the WHO/UNEP coordinated exposure studies
 2000–2015 performed in European countries, and used in Section 3.1.1.4.

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4790 4791 4792 4793 4794 4795 4796 4797 4798 4799 4800 4801 4802 4803 4804 4805 4806 4807 4808 4807 4808 4809 4810 4811 4812 4813 4814	<ul> <li>chicken whole blood by a modified quick, easy, cheap, effective, rugged, and safe method with liquid chromatography and tandem mass spectrometry. Journal of Separation Science, 39, 2846-2852.</li> <li>Zacs D and Bartkevics V, 2015. Analytical capabilities of high performance liquid chromatography - Atmospheric pressure photoionization - Orbitrap mass spectrometry (HPLC-APPI-Orbitrap-MS) for the trace determination of novel and emerging flame retardants in fish. Analytica Chimica Acta, 898, 60-72.</li> <li>Zacs D, Rjabova J, Pugajeva I, Nakurte I, Viksna A and Bartkevics V, 2014. Ultra high performance liquid chromatography-time-of-flight high resolution mass spectrometry in the analysis of hexabromocyclododecane diastereomers: Method development and comparative evaluation versus ultra high performance liquid chromatography coupled to Orbitrap high resolution mass spectrometry and triple quadrupole tandem mass spectrometry. Journal of Chromatography A, 1366, 73–78.</li> <li>Zacs D, Rjabova J, Ikkere LE, Bavrins K and Bartkevics V, 2018a. Brominated flame retardants and toxic elements in the meat and liver of red deer (<i>Cervus elaphus</i>), wild boar (<i>Sus scrofa</i>), and moose (<i>Alces alces</i>) from Latvian wildlife. Science of the Total Environment, 621, 308-316.</li> <li>Zacs D, Ikkere LE and Bartkevics V, 2018b. Emerging brominated flame retardants and dechlorane-related compounds in European eels (<i>Anguilla anguilla</i>) from Latvian lakes. Chemosphere, 197, 680-690.</li> <li>Zegers BN, Mets A, Van Bommel R, Minkenberg C, Hamers T, Kamstra JH, Pierce GJ and Boon JP, 2005. Levels of hexabromocyclododecane in harbor porpoises and common dolphins from western European seas, with evidence for stereoisomer-specific biotransformation by cytochrome p450. Environmental Science and Technology, 39, 2095-2100.</li> <li>Zeller H and Kirsch P, 1969. Hexabromocyclododecane: 28-day Feeding Trials with Rats. BASF Institute for Industrial Hygiene and Pharmacology, Federal Republic of Germany, EPA/OTS Doc. #86–900000376. As ci</li></ul>
4790 4791 4792 4793 4794 4795 4796 4797 4798 4799 4800 4801 4802 4803 4804 4805 4806 4807 4808 4807 4808 4809 4810 4811 4812 4813 4814 4815	<ul> <li>chicken whole blood by a modified quick, easy, cheap, effective, rugged, and safe method with liquid chromatography and tandem mass spectrometry. Journal of Separation Science, 39, 2846-2852.</li> <li>Zacs D and Bartkevics V, 2015. Analytical capabilities of high performance liquid chromatography - Atmospheric pressure photoionization - Orbitrap mass spectrometry (HPLC-APPI-Orbitrap-MS) for the trace determination of novel and emerging flame retardants in fish. Analytica Chimica Acta, 898, 60-72.</li> <li>Zacs D, Rjabova J, Pugajeva I, Nakurte I, Viksna A and Bartkevics V, 2014. Ultra high performance liquid chromatography-time-of-flight high resolution mass spectrometry in the analysis of hexabromocyclododecane diastereomers: Method development and comparative evaluation versus ultra high performance liquid chromatography coupled to Orbitrap high resolution mass spectrometry and triple quadrupole tandem mass spectrometry. Journal of Chromatography A, 1366, 73–78.</li> <li>Zacs D, Rjabova J, Ikkere LE, Bavrins K and Bartkevics V, 2018a. Brominated flame retardants and toxic elements in the meat and liver of red deer (<i>Cervus elaphus</i>), wild boar (<i>Sus scrofa</i>), and moose (<i>Alces alces</i>) from Latvian wildlife. Science of the Total Environment, 621, 308-316.</li> <li>Zacs D, Ikkere LE and Bartkevics V, 2018b. Emerging brominated flame retardants and dechlorane-related compounds in European eels (<i>Anguilla anguilla</i>) from Latvian lakes. Chemosphere, 197, 680-690.</li> <li>Zegers BN, Mets A, Van Bommel R, Minkenberg C, Hamers T, Kamstra JH, Pierce GJ and Boon JP, 2005. Levels of hexabromocyclododecane in harbor porpoises and common dolphins from western European seas, with evidence for stereoisomer-specific biotransformation by cytochrome p450. Environmental Science and Technology, 39, 2095-2100.</li> <li>Zeller H and Kirsch P, 1969. Hexabromocyclododecane: 28-day Feeding Trials with Rats. BASF Institute for Industrial Hygiene and Pharmacology, Federal Republic of Germany, EPA/OTS Doc. #86–900000376. As ci</li></ul>
4790 4791 4792 4793 4794 4795 4796 4797 4798 4799 4800 4801 4802 4803 4804 4805 4806 4807 4808 4807 4808 4809 4810 4811 4812 4813 4814 4815 4816	<ul> <li>chicken whole blood by a modified quick, easy, cheap, effective, rugged, and safe method with liquid chromatography and tandem mass spectrometry. Journal of Separation Science, 39, 2846-2852.</li> <li>Zacs D and Bartkevics V, 2015. Analytical capabilities of high performance liquid chromatography - Atmospheric pressure photoionization - Orbitrap mass spectrometry (HPLC-APPI-Orbitrap-MS) for the trace determination of novel and emerging flame retardants in fish. Analytica Chimica Acta, 898, 60-72.</li> <li>Zacs D, Rjabova J, Pugajeva I, Nakurte I, Viksna A and Bartkevics V, 2014. Ultra high performance liquid chromatography-time-of-flight high resolution mass spectrometry in the analysis of hexabromocyclododecane diastereomers: Method development and comparative evaluation versus ultra high performance liquid chromatography cupled to Orbitrap high resolution mass spectrometry and triple quadrupole tandem mass spectrometry. Journal of Chromatography A, 1366, 73–78.</li> <li>Zacs D, Rjabova J, Ikkere LE, Bavrins K and Bartkevics V, 2018a. Brominated flame retardants and toxic elements in the meat and liver of red deer (<i>Cervus elaphus</i>), wild boar (<i>Sus scrofa</i>), and moose (<i>Alces alces</i>) from Latvian wildlife. Science of the Total Environment, 621, 308-316.</li> <li>Zacs D, Ikkere LE and Bartkevics V, 2018b. Emerging brominated flame retardants and dechlorane-related compounds in European eels (<i>Anguilla anguilla</i>) from Latvian lakes. Chemosphere, 197, 680-690.</li> <li>Zegers BN, Mets A, Van Bommel R, Minkenberg C, Hamers T, Kamstra JH, Pierce GJ and Boon JP, 2005. Levels of hexabromocyclododecane in harbor porpoises and common dolphins from western European seas, with evidence for stereoisomer-specific biotransformation by cytochrome p450. Environmental Science and Technology, 39, 2095-2100.</li> <li>Zeller H and Kirsch P, 1969. Hexabromocyclododecane: 28-day Feeding Trials with Rats. BASF Institute for Industrial Hygiene and Pharmacology, Federal Republic of Germany, EPA/OTS Doc. #86–900000376. As cit</li></ul>
4790 4791 4792 4793 4794 4795 4796 4797 4798 4799 4800 4801 4802 4803 4804 4805 4806 4807 4808 4807 4808 4809 4810 4811 4812 4813 4814 4815	<ul> <li>chicken whole blood by a modified quick, easy, cheap, effective, rugged, and safe method with liquid chromatography and tandem mass spectrometry. Journal of Separation Science, 39, 2846-2852.</li> <li>Zacs D and Bartkevics V, 2015. Analytical capabilities of high performance liquid chromatography - Atmospheric pressure photoionization - Orbitrap mass spectrometry (HPLC-APPI-Orbitrap-MS) for the trace determination of novel and emerging flame retardants in fish. Analytica Chimica Acta, 898, 60-72.</li> <li>Zacs D, Rjabova J, Pugajeva I, Nakurte I, Viksna A and Bartkevics V, 2014. Ultra high performance liquid chromatography-time-of-flight high resolution mass spectrometry in the analysis of hexabromocyclododecane diastereomers: Method development and comparative evaluation versus ultra high performance liquid chromatography coupled to Orbitrap high resolution mass spectrometry and triple quadrupole tandem mass spectrometry. Journal of Chromatography A, 1366, 73–78.</li> <li>Zacs D, Rjabova J, Ikkere LE, Bavrins K and Bartkevics V, 2018a. Brominated flame retardants and toxic elements in the meat and liver of red deer (<i>Cervus elaphus</i>), wild boar (<i>Sus scrofa</i>), and moose (<i>Alces alces</i>) from Latvian wildlife. Science of the Total Environment, 621, 308-316.</li> <li>Zacs D, Ikkere LE and Bartkevics V, 2018b. Emerging brominated flame retardants and dechlorane-related compounds in European eels (<i>Anguilla anguilla</i>) from Latvian lakes. Chemosphere, 197, 680-690.</li> <li>Zegers BN, Mets A, Van Bommel R, Minkenberg C, Hamers T, Kamstra JH, Pierce GJ and Boon JP, 2005. Levels of hexabromocyclododecane in harbor porpoises and common dolphins from western European seas, with evidence for stereoisomer-specific biotransformation by cytochrome p450. Environmental Science and Technology, 39, 2095-2100.</li> <li>Zeller H and Kirsch P, 1969. Hexabromocyclododecane: 28-day Feeding Trials with Rats. BASF Institute for Industrial Hygiene and Pharmacology, Federal Republic of Germany, EPA/OTS Doc. #86–900000376. As ci</li></ul>



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#### 4853 **Abbreviations**

- 4854 AMPK Adenosine monophosphate-activated kinase
- 4855 APCI Atmospheric pressure chemical ionisation
- 4856 APPI Atmospheric pressure photoionisation
- 4857 BDNF Brain derived neurotropic factor
- 4858 BFRs Brominated Flame Retardants
- 4859 BMD Benchmark dose
- 4860 BMDL Benchmark dose lower confidence limit
- 4861 BMDL<sub>10</sub> Benchmark dose lower confidence limit for a benchmark response of 10%
- 4862 BMI Body mass index
- 4863 BMR Benchmark response
- 4864 CAR Constitutive and rostane receptor
- 4865 CDH1 Suppressed E-cadherin
- 4866 CYP Cytochrome P450
- 4867 DNA-PKcs DNA-dependent protein kinase
- 4868 DTT Dithiothreitol
- 4869 dw dry weight
- 4870 E2 Estradiol
- 4871 EC European Commission
- 4872 EF Enantiomeric fraction
- 4873 EFSA European Food Safety Authority
- 4874 EPS Expanded polystyrene foams
- 4875 EQS Environmental quality standard
- 4876 ESI Electrospray ionisation
- 4877 EU European Union

4879	EED	Free estradiol
	FE2	
4880	FGF	Fibroblast growth factor
4881	FSH	Follicle stimulating hormone
4882	GC	Gas chromatography
4883	GIC	Groningen Infant COMPARE birth cohort
4884	HBCDDs	Hexabromocyclododecanes
4885	hCG	human chorionic gonadotropin
4886	HSD17β	17β-hydroxysteroid dehydrogenase
4887	IC <sub>50</sub>	half maximal inhibitory concentration
4888	i.p.	intraperitoneal
4889	InhB	inhibin B
4890	IQR	median interquartile range
4891	IRIS	Integrated Risk Information System
4892	LADD	Lifetime average daily dose
4893	LB	lower bound
4894	LC	Liquid chromatography
4895	LH	Luteinizing hormone
4896	LOAEL	Loest-observed-adverse-effect level
4897	LOD	Limit of detection
4898	LOEL	Lowest-observed-effect level
4899	LRMS	low resolution MS
4900	MAPK	Mitogen-activated protein kinases
4901	MDL	Method detection limit
4902	MOE	Margin of exposure
4903	MMP9	Matrix metalloprotein 9
4904	MS	Mass spectrometry
4905	MSC	Mesenchymal stem cells
4906	mTOR	Mammalian target of rapamycin
4907	NGF	Nerve growth factor
4908	NICNAS	National Industrial Chemicals Notification and Assessment Scheme
4909	NIPH	Norwegian Institute of Public Health
4910	NITE	Japanese National Institute of Technology and Evaluation
4911	NMDA	N-methyl-d-aspartate
4912	NOAEL	No-observed-adverse-effect level
4913	NOEL	No-observed-effect level
4914	NRLs	National Reference Laboratories
4915	PAHs	
4916		Polycyclic aromatic hydrocarbons
	-	Polycyclic aromatic hydrocarbons Polybrominated dinbenyl ethers
	PBDEs	Polybrominated diphenyl ethers
4917	PBDEs PBDD	Polybrominated diphenyl ethers Polybrominated dioxins
4917 4918	PBDEs PBDD PBPK	Polybrominated diphenyl ethers Polybrominated dioxins Physiologically based pharmacokinetic
4917 4918 4919	PBDEs PBDD PBPK PBT	Polybrominated diphenyl ethers Polybrominated dioxins Physiologically based pharmacokinetic Persistent, bioaccumulative and toxic
4917 4918 4919 4920	PBDEs PBDD PBPK PBT PCBs	Polybrominated diphenyl ethers Polybrominated dioxins Physiologically based pharmacokinetic Persistent, bioaccumulative and toxic Polychlorinated biphenyls
4917 4918 4919 4920 4921	PBDEs PBDD PBPK PBT PCBs PCNA	Polybrominated diphenyl ethers Polybrominated dioxins Physiologically based pharmacokinetic Persistent, bioaccumulative and toxic Polychlorinated biphenyls proliferating cell nuclear antigen
4917 4918 4919 4920 4921 4922	PBDEs PBDD PBPK PBT PCBs PCNA PI3K	Polybrominated diphenyl ethers Polybrominated dioxins Physiologically based pharmacokinetic Persistent, bioaccumulative and toxic Polychlorinated biphenyls proliferating cell nuclear antigen Phosphatidylinositide 3-kinase
4917 4918 4919 4920 4921 4922 4923	PBDEs PBDD PBPK PBT PCBs PCNA PI3K PLE	Polybrominated diphenyl ethers Polybrominated dioxins Physiologically based pharmacokinetic Persistent, bioaccumulative and toxic Polychlorinated biphenyls proliferating cell nuclear antigen Phosphatidylinositide 3-kinase Pressurised liquid extraction
4917 4918 4919 4920 4921 4922 4923 4923	PBDEs PBDD PBPK PBT PCBs PCNA PI3K PLE PND	Polybrominated diphenyl ethers Polybrominated dioxins Physiologically based pharmacokinetic Persistent, bioaccumulative and toxic Polychlorinated biphenyls proliferating cell nuclear antigen Phosphatidylinositide 3-kinase Pressurised liquid extraction Postnatal day
4917 4918 4919 4920 4921 4922 4923 4924 4925	PBDEs PBDD PBPK PBT PCBs PCNA PI3K PLE PND POPRC	Polybrominated diphenyl ethers Polybrominated dioxins Physiologically based pharmacokinetic Persistent, bioaccumulative and toxic Polychlorinated biphenyls proliferating cell nuclear antigen Phosphatidylinositide 3-kinase Pressurised liquid extraction Postnatal day Persistent Organic Pollutants Review Committee
4917 4918 4919 4920 4921 4922 4923 4924 4925 4926	PBDEs PBDD PBPK PBT PCBs PCNA PI3K PLE PND POPRC POPs	Polybrominated diphenyl ethers Polybrominated dioxins Physiologically based pharmacokinetic Persistent, bioaccumulative and toxic Polychlorinated biphenyls proliferating cell nuclear antigen Phosphatidylinositide 3-kinase Pressurised liquid extraction Postnatal day Persistent Organic Pollutants Review Committee Persistent Organic Pollutants
4917 4918 4919 4920 4921 4922 4923 4924 4925 4926 4927	PBDEs PBDD PBPK PBT PCBs PCNA PI3K PLE PND POPRC POPS PPAR	Polybrominated diphenyl ethers Polybrominated dioxins Physiologically based pharmacokinetic Persistent, bioaccumulative and toxic Polychlorinated biphenyls proliferating cell nuclear antigen Phosphatidylinositide 3-kinase Pressurised liquid extraction Postnatal day Persistent Organic Pollutants Review Committee Persistent Organic Pollutants Peroxisome proliferator-activated receptors
4917 4918 4919 4920 4921 4922 4923 4924 4925 4926 4927 4928	PBDEs PBDD PBPK PBT PCBs PCNA PI3K PLE PND POPRC POPs PPAR PXR	Polybrominated diphenyl ethers Polybrominated dioxins Physiologically based pharmacokinetic Persistent, bioaccumulative and toxic Polychlorinated biphenyls proliferating cell nuclear antigen Phosphatidylinositide 3-kinase Pressurised liquid extraction Postnatal day Persistent Organic Pollutants Review Committee Persistent Organic Pollutants Peroxisome proliferator-activated receptors Pregnane-X-receptor
4917 4918 4919 4920 4921 4922 4923 4924 4925 4926 4927 4928 4929	PBDEs PBDD PBPK PBT PCBs PCNA PI3K PLE PND POPRC POPS PPAR PXR ROS	Polybrominated diphenyl ethers Polybrominated dioxins Physiologically based pharmacokinetic Persistent, bioaccumulative and toxic Polychlorinated biphenyls proliferating cell nuclear antigen Phosphatidylinositide 3-kinase Pressurised liquid extraction Postnatal day Persistent Organic Pollutants Review Committee Persistent Organic Pollutants Peroxisome proliferator-activated receptors Pregnane-X-receptor Reactive oxygen species
4917 4918 4919 4920 4921 4922 4923 4924 4925 4926 4927 4928 4929 4930	PBDEs PBDD PBPK PBT PCBs PCNA PI3K PLE PND POPRC POPs PDAR PXR ROS SERCA	Polybrominated diphenyl ethers Polybrominated dioxins Physiologically based pharmacokinetic Persistent, bioaccumulative and toxic Polychlorinated biphenyls proliferating cell nuclear antigen Phosphatidylinositide 3-kinase Pressurised liquid extraction Postnatal day Persistent Organic Pollutants Review Committee Persistent Organic Pollutants Peroxisome proliferator-activated receptors Pregnane-X-receptor Reactive oxygen species Sarcoplasmic-endoplasmic reticulum Ca <sup>2+</sup> -ATPase
4917 4918 4919 4920 4921 4922 4923 4924 4925 4926 4927 4928 4929 4930 4931	PBDEs PBDD PBPK PBT PCBs PCNA PI3K PLE PND POPRC POPs PDAR PXR ROS SERCA SHBG	Polybrominated diphenyl ethers Polybrominated dioxins Physiologically based pharmacokinetic Persistent, bioaccumulative and toxic Polychlorinated biphenyls proliferating cell nuclear antigen Phosphatidylinositide 3-kinase Pressurised liquid extraction Postnatal day Persistent Organic Pollutants Review Committee Persistent Organic Pollutants Peroxisome proliferator-activated receptors Pregnane-X-receptor Reactive oxygen species Sarcoplasmic-endoplasmic reticulum Ca <sup>2+</sup> -ATPase Sex hormone-binding globulin
4917 4918 4919 4920 4921 4922 4923 4924 4925 4926 4927 4928 4929 4930 4931 4932	PBDEs PBDD PBPK PBT PCBs PCNA PI3K PLE PND POPRC POPs PDAR PXR ROS SERCA SHBG SPE	Polybrominated diphenyl ethers Polybrominated dioxins Physiologically based pharmacokinetic Persistent, bioaccumulative and toxic Polychlorinated biphenyls proliferating cell nuclear antigen Phosphatidylinositide 3-kinase Pressurised liquid extraction Postnatal day Persistent Organic Pollutants Review Committee Persistent Organic Pollutants Peroxisome proliferator-activated receptors Pregnane-X-receptor Reactive oxygen species Sarcoplasmic-endoplasmic reticulum Ca <sup>2+</sup> -ATPase Sex hormone-binding globulin Solid phase extraction
4917 4918 4919 4920 4921 4922 4923 4924 4925 4926 4927 4928 4929 4930 4931 4932 4933	PBDEs PBDD PBPK PBT PCBs PCNA PI3K PLE PND POPRC POPS PPAR PXR ROS SERCA SHBG SPE SSD	Polybrominated diphenyl ethers Polybrominated dioxins Physiologically based pharmacokinetic Persistent, bioaccumulative and toxic Polychlorinated biphenyls proliferating cell nuclear antigen Phosphatidylinositide 3-kinase Pressurised liquid extraction Postnatal day Persistent Organic Pollutants Review Committee Persistent Organic Pollutants Peroxisome proliferator-activated receptors Pregnane-X-receptor Reactive oxygen species Sarcoplasmic-endoplasmic reticulum Ca <sup>2+</sup> -ATPase Sex hormone-binding globulin Solid phase extraction Standard Sample Description
4917 4918 4919 4920 4921 4922 4923 4924 4925 4926 4927 4928 4929 4930 4931 4932	PBDEs PBDD PBPK PBT PCBs PCNA PI3K PLE PND POPRC POPs PDAR PXR ROS SERCA SHBG SPE	Polybrominated diphenyl ethers Polybrominated dioxins Physiologically based pharmacokinetic Persistent, bioaccumulative and toxic Polychlorinated biphenyls proliferating cell nuclear antigen Phosphatidylinositide 3-kinase Pressurised liquid extraction Postnatal day Persistent Organic Pollutants Review Committee Persistent Organic Pollutants Peroxisome proliferator-activated receptors Pregnane-X-receptor Reactive oxygen species Sarcoplasmic-endoplasmic reticulum Ca <sup>2+</sup> -ATPase Sex hormone-binding globulin Solid phase extraction



4940ODOpperboland4941UKUnitek Kingdom4942UNEPUnited Nations Environment Programme4943UPLCUltra performance LC4944UVUltraviolet4945WHOWorld health Organization4946wwwet weight4947XPSExtruded polystyrene foams49484949495049514952	4942 4943 4944 4945 4946 4947 4948 4949 4950 4951	UNEP UPLC UV WHO WW	United Nations Environment Programme Ultra performance LC Ultraviolet World health Organization wet weight
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# Appendix A – Levels in dust

#### 4953 **Table A.1**. Occurrence of HBCDDs in dust in Europe

Country	Sampling year		ΣHBCDD	α-HBCDD	β-HBCDD	γ-HBCDD	
		Sampling place	Median (range) (ng/g)				Reference
		Houses	100 (15–990)				
		Apartments	45 (<3–2,400)				
Sweden	2006	Offices	300 (190–1,600)	-	-	-	de Wit et al. (2012)
		Day care centres	340 (190–1,600)				()
		Cars	54 (6.8–170)				
Sweden	2000	Vacuum cleaner bags	89 (6.4–95,000)				Björklund et al. (2012)
<b>Sweden</b> 2008	2008	Researcher-collected dust	8.9 (5.9–62)		-	-	
Sweden	2015	Preschool	100 (7.6–16,000)	52 (4.9–3,200)	15 (1.2–1,200)	26 (1.3–12,000)	Larsson et al. (2018)
Sweden	2012	Offices, appartments, stores and schools	150 (17–2,900)	-	-	-	Newton et al. (2015)
Sweden	2010–2011	House dust	110 (20–6,000)	56 (14–1,400)	18 (3.4–730)	37 (2.5–4,000)	Sahlström et al. (2015b)
		Houses	100 (15–990)				
		Apartments	45 (<3–2,400)				
Sweden	2006	Offices	300 (190–5,700)	-	-	-	Thuresson et al (2012)
		Day care centres	340 (190–1,600)				<u> </u>
		Cars	54 (6.8–170)				



Country	Sampling year		ΣHBCDD	α-HBCDD	β-HBCDD	γ-HBCDD		
		Sampling place		Reference				
Norway	2013–2014	Homes	190 (0.43–150,000)	94 (<0.14–74,000)	23 (<0.017–60,000)	43 (<0.28–13,000)	Tay et al. (2017)	
Germany	2010	House dust	345 (53–4,041)	180 (32–1,063)	35 (8–496)	114 (14–3,563)	Fromme et al. (2014)	
		Houses	1,125 (636–1,865)	559	144	422		
France	2008	Offices	4,493 (1,096-	2,722	442	1,329	Abdallah et al. (2016)	
		Cars	4,539 (1,458–7,900)	2,221	629	1,689	(2010)	
Dolaium	2008	House dust	130 (5–42,692)				D'Hollander et	
Belgium		Office dust	367 (256–1,153)	-	-	-	al. (2010)	
		Kitchen	-	110 (5.2–3,800)	29 (2.3–1,100)	35 (1.7–13,000)		
UK		Living room/bedroom	-	280 (75–4,900)	67 (6.4–1,600)	110 (14–21,000)	Kuang et al. (2016)	
		Matched kitchen-living room/bedroom dust	-	0.37 (0.05–2.88)	0.41 (0.08–1.86)	0.37 (0.003–34.85)		
UK		Homes	610 (50–110,000)	320 (21–28,000)	85 (6.1–12,000)	93 (23–71,000)	Tao et al.	
UK		Offices	1,700 (150–6,400)	980 (100–2,800)	330 (22–590)	350 (31–3,700)	(2016a)	
		Office dust	380					
Ireland	2016–2017	School dust	800	-	-	-	Wemken et al. (2019)	
		Car dust	490				(2015)	
Portugal	2010–2011	House dust	150 (16–2,000)	91 (11–1,800)	16 (1.9–230)	35 (2.8–900)	Coelho et al. (2016b)	
	2000	Car dust	(<0.3–241)	(<0.3–45)	(<0.3–44)	(<0.3–152)	Kalachova et	
Czech Republic	Czech Republic 20	2008	Household dust	(<0.3–950)	(<0.3–275)	(<0.3–57)	(<0.3–740)	al. (2012)



Country	Sampling	<sup>g</sup> Sampling place	ΣΗΒCDD	α-HBCDD	β-HBCDD	γ-HBCDD		
	year			- Reference				
Czech Republic	2013	House dust	122 (34.1–714)	-	-	-	Lankova et al. (2015)	
Romania	2010	Indoor dust	325 (4–2,190)	200 (4–1,325)	30 ( <loq-165)< td=""><td>55 (<loq-1,910)< td=""><td>Dirtu et al. (2012)</td></loq-1,910)<></td></loq-165)<>	55 ( <loq-1,910)< td=""><td>Dirtu et al. (2012)</td></loq-1,910)<>	Dirtu et al. (2012)	
Greece	2016	Car dust	-	90.3 (<5–1,288)	15.8 (<5–294)	46.4 (<5–260)	Besis et al. (2017)	
		<i>Urban location</i> Office dust Home dust	(140–94,000) (50–8,800)					
Istanbul/Turkey	2012	<i>Semi-urban location</i> Office dust Home dust	( <lod-20,000) (110-1,200)</lod-20,000) 		-	-	Kurt-Karakus et al. (2017)	
		<i>Rural location</i> Office dust Home dust	not analysed (230–29,000)					
Airplanes	2010	Floor	7,600 (180– 1,100,000)	2,300 (4.7–290,000)	310 (1.2–75,000)	4,500 (130–700,000)	Allen et al.	
Anplanes	2010	Vent	10,000 (370–97,000)	1600 (17–32,000)	230 (0.8–11,000)	7,600 (99–59,000)	(2013)	
Airplanes	2212		difcfall	(301.6–231,010)	_	_	_	Strid et al.
	2012	During flight (ventilation outlets in aircraft toilets)	(243.85–173,260)	-	-	-	(2014)	
Several countries	2002–2017	Dust (children's indoor exposure)	World-wide dust levels: ranging from 6 (Egypt) to 340 ng/g (Sweden)	-	-	-	Malliari and Kalantzi (2017	



### Appendix B – Literature search to identify studies on HBCDDs

The details of the literature searches performed to identify studies on HBCDDs are shown in **Table B.1** The search strings were run in the databases indicated, and the CAS numbers in SciFinder.

The literature searches were performed in February 2019. The outcome of the searches in the different databases was saved in separate EndNote files and an automatic duplicate detection run. The references were then transferred to DistillerSR (a web-based systematic review software) where another automatic duplicate detection was made (**Table B.2**). The selection for relevance based on tittle and abstract and full text was done in duplicate by EFSA staff.

An update of the literature available in the public domain was done in April 2020, and since that date, the literature was monitored to identify studies relevant for the risk assessment until the time of endorsement.

4965 **Table B.1.** Details of the literature search for HBCDDs

Databases interrogated:	WoS - Web of Science PubMED SciFINDER (including CAS numbers)	
Year:	2010-date of the search	
Languages:	English	
Type of documents:	Peer reviewed articles, reviews, books	
Search string:	TS=(hexabromocyclododecane <u>OR</u> hexabromocyclododecane* <u>OR</u> HBCDD <u>OR</u> HBCDDs <u>OR</u> *hexabromocyclododecane OR alpha-hexabromocyclododecane <u>OR</u> beta-hexabromocyclododecane <u>OR</u> gamma-hexabromocyclododecane <u>OR</u> delta- hexabromocyclododecane <u>OR</u> epsilon-hexabromocyclododecane)	
CAS numbers:	25637-99-4, 3194-55-6, 134237-50-6, 134237-51-7, 134237-52-8, 878049-08-2, 878049-05-9, 878049-04-8, 678970-17-7, 678970-16-6, 678970-15-5, 169102- 57-2, 138257-19-9, 138257-18-8, 138257-17-7	
Grey literature:	<ul> <li>Assessments done by national and international bodies (e.g. in google or from papers, ECHA, NTP).</li> <li>Dedicated search in <u>Organohalogen Compounds database</u> (extended abstracts from DIOXIN conferences). To provide list in a table.</li> <li>Dedicated search in the BFR conference abstracts available from its website.</li> </ul>	

#### 4966 **Table B.2.** Outcome of the literature searches

	Date	Search string	N hits
WOS (all databases)	05.02.2019	TS=(hexabromocyclododecane OR hexabromocyclododecane* OR HBCDD OR HBCDDs OR *hexabromocyclododecane OR alpha- hexabromocyclododecane OR beta-hexabromocyclododecane OR gamma-hexabromocyclododecane OR delta-hexabromocyclododecane OR epsilon-hexabromocyclododecane) AND P=(2010-2019) Refined by: [excluding] DOCUMENT TYPES: (EDITORIAL OR DATA STUDY OR LETTER OR DATA SET OR UNSPECIFIED OR DATA PAPER ) Search language=English	1,002 (after automatic duplicate removal in EndNote)



PUBMED	05.02.2019	(((hexabromocyclododecane OR hexabromocyclododecane* OR HBCDD OR HBCDDs OR *hexabromocyclododecane OR alpha- hexabromocyclododecane OR beta-hexabromocyclododecane OR gamma-hexabromocyclododecane OR delta-hexabromocyclododecane OR epsilon-hexabromocyclododecane))) AND (("english"[Language]) AND ("2010"[Date - Publication] : "3000"[Date - Publication]))	613 (after automatic duplicate removal in EndNote)
SciFinder	05.02.2019	hexabromocyclododecane OR hexabromocyclododecanes (1166) 25637-99-4 (820) 3194-55-6 (70) 134237-50-6 (439) 134237-51-7 (363) 134237-52-8 (384) 878049-08-2 (1) 878049-08-2 (1) 878049-04-8 (1) 678970-17-7 (25) 678970-16-6 (18) 678970-15-5 (26) 169102-57-2 (23) 138257-18-8 (20) 138257-18-8 (20) 138257-17-7 (1) Selected on: Journal, review and book 2010-2019 English	1,211 (after automatic duplicate removal in EndNote)
ALL databases combined			1,412 (after automatic duplicate removal in DistillerSR)



# Appendix C – Histological evaluation in Maranghi et al. (2013), Rasinger et al. (2014, 2018)

**Table C.1.** Histological evaluation of potential target tissues of HBCDDs toxicity in mice after 28 days of repeated dietary exposure to HBCDDs at daily doses of 49.5  $\mu$ g/kg bw per day or 199 mg/kg bw per day (Maranghi et al., 2013; Rasinger et al., 2014, 2018). Significant effects at the p<0.05 and p<0.01

4972 levels are indicated by (+) and (++), respectively.

Tissue	Observations	Control	HBCDDs low dose (49.5 µg/kg bw/day)	HBCDDs high dose (199 mg/kg bw/day)
Thyroid	Desquamation into follicular lumen	0/7 (0%)	3/7 (43%)	1/7
	Foaming colloid	1/7 (14%)	2/7 (29%)	not analysed
	Irregular multistratification of the luminal epithelium	1/8 (12%)	2/7 (29%)	not analysed
Uterus	Reduction in endometrial glands density	0/8 (0%)	4/7 (+) (57%)	not analysed
	Marked vacuolization in hepatocytes	0/10 (0%)	5/8 (++) (62%)	5/8 (++)
Liver	Lymphocytic infiltration	0/10 (0%)	6/8 (++) (75%)	6/8 (++)
	Hyperaemic vessels	0/10 (0%)	6/8 (++) (75%)	6/8 (++)
	Picnotic nucleus	0/10 (0%)	2/8 (25%)	2/8
Adrenals	Thickness external capsule	2/10 (20%)	3/7 (43%)	not analysed
	Hassal's bodies	2/10 (20%)	5/10 (50%)	2/9
-	Cortical invasivity	2/10 (20%)	2/10 (20%)	8/9 (++)
Thymus	Minerals	2/10 (20%)	5/10 (50%)	not analysed
	Stress	0/10 (0%)	5/10 (+) (50%)	not analysed
	Lymphocyte hyperplasia	4/10 (40%)	2/10 (20%)	5/10
Spleen	Infiltration in red pulp and periarteriolar zones (PALS)	0/10 (0%)	2/10 (20%)	0/10
	Eosinophilic cells in the cortex	2/10 (20%)	4/9 (44%)	not analysed
Brain	Eosinophilic cells in the thalamus	3/10 (30%)	4/9 (44%)	not analysed
Brain	Condensed cells in the hypothalamus	0/10 (0%)	0/9 (0%)	not analysed

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Reference	Comments
Li et al. (2017b). Hexabromocyclododecane- induced Genotoxicity in Cultured Human Breast Cells through DNA Damage.	Largely observational study. HBL-100 cells were exposed to high concentrations (0, 5, 10 and 50 mg/L) for 24 h. LDH leakage observed at 5 mg/L and significant ROS and comet tail migration at 50 mg/L. Expression of transcripts for ATM, and DNA repair genes OGG1 and MTH was increased and that for tumour suppressor gene BRCA1 downregulated.
Krivoshiev et al. (2015). Elucidating toxicological mechanisms of current flame retardants using a bacterial gene profiling assay.	Unclear what concentration was used and genes <2-fold regulated by HBCDDs.
Wielogorska et al. (2015). Endocrine disruptor activity of multiple environmental food chain contaminants.	Mammalian reporter gene assay for ER activation by $\beta\text{-HBCDD}.$ No activation found.
Kovarich et al. (2011). Quantitative structure- activity relationship classification models for prediction of endocrine disrupting activity of brominated flame retardants.	γ-HBCDD predicted to be anti-oestrogenic by QSAR.
Papa et al. (2013). QSAR prediction of the competitive interaction of emerging halogenated pollutants with human transthyretin SAR QSAR.	$\gamma$ -HBCDD used in training set as a BFR without T4-TTR competing potency.
Anisuzzaman and Whalen (2016). Tetrabromobisphenol A and hexabromocyclododecane alter secretion of IL- 1β from human immune cells.	Human natural killer (NK) cells, monocyte-depleted (MD) peripheral blood mononuclear cells (MD-PBMC) and PBMC exposured to 0.05–5.0 mM of HBCDD for 24 h, 48 h and 6 days. Interfere with the ability of immune cells to secrete IL-1B. Exposure to HBCDD from 0.5–5.0 mM caused increases in IL-1B secretion. Inhibitors of ERK1/2 and Casp1 ameliorated HBCDD induced IL-1B release.
Almughamsi and Whalen (2016). Hexabromocyclododecane and tetrabromobisphenol A alter secretion of interferon gamma (IFN-gamma) from human immune cells.	Human natural killer (NK) cells, monocyte-depleted (MD) peripheral blood mononuclear cells (MD-PBMC) and PBMC exposured to 0.05–5.0 mM of HBCDD for 24 h, 48 h and 6 days. interfere with the ability of immune cells to secrete IFN-gamma. HBCDD stimulated secretion if IFN gamma but response very different in cells from different donors. Also baseline values were different.
Cato et al. (2014). Brominated flame retardants, tetrabromobisphenol A and hexabromocyclododecane, activate mitogen- activated protein kinases (MAPKs) in human natural killer cells.	HBCDDs interfere with NK-cell(s) lytic function. HBCDDs at 2.5 uM and above increased phosphorylation of p44/42; 1 uM and above increase phospho-MEK1/2. p44/42 is required for NK lysis of tumour target cells
Hinkson and Whalen (2010). Hexabromocyclododecane decreases tumor- cell-binding capacity and cell-surface protein expression of human natural killer cells.	Exposure of NK-cells to 10 $\mu$ M HBCDDs for 24 h caused > 90% loss of lytic function of NK-cells and 71% decrease in NK-cell binding function, and in expression of cell surface markers CD16 (58%) and CD56 (25%). NK-cells exposed to 10 $\mu$ M HBCDDs for 1 h followed by 24 h in HBCDD-free media showed 89% loss of lytic function and decreased binding function (79.2%), and CD 16 expression (48.1%).
Koike et al. (2016). Brominated flame retardants, hexabromocyclododecane and tetrabromobisphenol A, affect proinflammatory	HBCDDs exposure (10 µg/mL) increased the expression of ICAM-1 and the production of IL-6 and -8 in human bronchial epithelial cells (BEAS-2B). HBCDDs caused activation of nuclear factor-kappa B (p50, p65) and

# Appendix D – Mode of action studies



protein expression in human bronchial epithelial cells via disruption of intracellular signalling.	activator protein 1 (c-Jun). HBCDDs showed no binding to nuclear oestrogen or thyroid hormone receptors.
Steves et al. (2018). Ubiquitous Flame- Retardant Toxicants Impair Spermatogenesis in a Human Stem Cell Model.	Human stem cell-based model of spermatogenesis, indicating that HBCDDs affect spermatogonia and primary spermatocytes through mitochondrial membrane potential perturbation and ROS generation, resulting in apoptosis. HBCDD concentrations: 1 uM, 10 uM, 25 uM, 50 uM, 100 uM, and 200 $\mu$ M dissolved in dimethyl sulfoxide (DMSO). Reduced viability at 25 $\mu$ M but still 80% viability at 200 $\mu$ M. Loss of marker (PLZF) for stem and progenitor spermatogonia observed at 1 $\mu$ M. Primary spermatocyte marker piwi like RNA-mediated gene silencing 2 (HILI) increased in expression from 1 to 25 $\mu$ M and then declined.
Yasmin and Whalen (2018). Flame retardants, hexabromocyclododecane (HCBD) and tetrabromobisphenol a (TBBPA), alter secretion of tumor necrosis factor alpha (TNFa) from human immune cells.	Effects of HBCDDs on secretion of tumour necrosis factor alpha (TNF $\alpha$ ) from human immune cells. In presence of T-cells, HBCDDs increased TNF $\alpha$ secrection, but without T-cells HBCDD caused reduction. HBCDD-induced increases in TNF $\alpha$ secretion utilized the p38 MARK pathway.
Zhang et al. (2015). Transcriptomic and metabolomic approaches to investigate the molecular responses of human cell lines exposed to the flame retardant hexabromocyclododecane (HBCD).	Transcriptomic and metabolomic effects of HBCDDs on A549 and HepG2/C3A cells. MTT assay for A549 and HepG2 cells indicated EC50 values of 27.4 $\mu$ M and 63.0 $\mu$ M, respectively. Little effect on gene expression or metabolome in either cell type up to the highest dose of 4 $\mu$ M.
Zou et al. (2013). PI3K/Akt pathway mediates Nrf2/ARE activation in human L02 hepatocytes exposed to low-concentration HBCDs.	Nanomolar concentrations of HBCDDs stimulate L02 cell proliferation in a DNA-PKcs-dependent manner, increase protein levels and nuclear translocation of Nrf2, leading to upregulation of its target gene HO-1. The PI3K/Akt pathway is essential for HBCDD activation of the Nrf2-ARE pathwa L02 cells.
Dorosh et al. (2010). Assessing oestrogenic effects of brominated flame retardants hexabromocyclododecane and tetrabromobisphenol A on MCF-7 cells.	E-screen assay in MCF-7 cells. HBCDDs increased cell proliferation and dose-dependently increased gene expression of TFF1 with effects starting at 20–200 nM. Blocked by the anti-oestrogen, ICI 182,780.
Arini et al. (2017). A cell-free testing platform to screen chemicals of potential neurotoxic concern across twenty vertebrate species.	High throughput cell-free neurochemical assay based on receptors and enzymes. Not much specific information on HBCDDs.
An et al. (2014a). Hexabromocyclododecane and polychlorinated biphenyls increase resistance of hepatocellular carcinoma cells to cisplatin through the phosphatidylinositol 3- kinase/protein kinase B pathway.	HBCDDs reduces sensitivity of HCC cells (HepG2, MHCC97H, and MHCC97L) to cisplatin through modulation of the NF-kB pathway activation and p53 and associated with the PI3K/Akt pathway activity.
Al-Mousa and Michelangeli (2014). The sarcoplasmic-endoplasmic reticulum Ca(2+)- ATPase (SERCA) is the likely molecular target for the acute toxicity of the brominated flame retardant hexabromocyclododecane (HBCD).	Six BFRs assessed for cytotoxicity and potency to inhibit SERCA. Strong Direct correlation (r=0.94) between the potencies of inducing cell death and inhibiting SERCA. Ki of HBCDDs for SERCA was 2.7 uM. Mechanistic studies indicate that HBCDDs precent ATP binding to SERCA.
Al-Mousa and Michelangeli (2012). Some commonly used brominated flame retardants cause Ca2+-ATPase inhibition, beta-amyloid peptide release and apoptosis in SH-SY5Y neuronal cells.	HBCDDs causeed dose-dependent increase in [Ca2+] <sub>i</sub> and inhibits the SERCA with an apparent Ki of 3.5 $\mu$ M in SH-SY5Y human neuroblastoma cells. Increase in ROS formation, cytochrome C release and apoptosis.
Bastos Sales et al (2013). Effects of endocrine disrupting chemicals on in vitro global DNA methylation and adipocyte differentiation.	No effect of 10 $\mu\text{M}$ HBCDDs on DNA methylation and adipocyte differentiation.



Christen et al. (2010). Some flame retardants and the antimicrobials triclosan and triclocarban enhance the androgenic activity in vitro.	Studied androgenic and antiandrogenic activity of BFRs and antimicrobials <i>in vitro</i> in MDA-kb2 cells. HBCDDs very low activity but enhances androgenic activity.
Fa et al. (2013). Acute effects of hexabromocyclododecane on Leydig cell cyclic nucleotide signaling and steroidogenesis in vitro.	HBCDDs inhibited basal and hCG-stimulated cAMP production, but elevated basal steroidogenesis. HBCDDs decrease in mitochondrial membrane potential in untreated and hCG-treated cells.
Fa et al. (2015). HBCDD-induced sustained reduction in mitochondrial membrane potential, ATP and steroidogenesis in peripubertal rat Leydig cells.	HBCDDs caused a sustained reduction in ATP level. Accumulation of cAMP and androgen were also reduced. There was inhibition in the expression of genes for steroidogenic enzymes, luteinizing hormone receptor, regulatory and transport proteins, and a decrease in abundance of StAR. Enzymatic experiments indicated loss of activities of (CYP11A1) and 17 $\beta$ -hydroxysteroid dehydrogenase (HSD17 $\beta$ ).
Fa et al. (2014). Hexabromocyclododecane facilitates FSH activation of ERK1/2 and AKT through epidermal growth factor receptor in rat granulosa cells.	HBCDDs potentiate FSH-stimulated phosphorylation of EGFR, ERK1/2 and AKT, indicating a direct effect on EGFR.
Huang et al. (2016b). In vitro study on the biotransformation and cytotoxicity of three hexabromocyclododecane diastereoisomers in liver cells.	Cytotoxicity of HBCDDs in L02 and HepG2 cells was $\beta$ -HBCDD > $\gamma$ -HBCDD > $\alpha$ -HBCDD.
Ibhazehiebo et al. (2011a). 1,2,5,6,9,10- aHexabromocyclododecane (HBCD) impairs thyroid hormone-induced dendrite arborization of Purkinje cells and suppresses thyroid hormone receptor-mediated transcription.	HBCDDs at 0.1 nM suppressed TR-mediated transcription in a reporter gene assay, and suppressed TH-induced dendrite arborization of Purkinje cells in primary cerebellar culture derived from rat neonates.
Ibhazehiebo et al. (2011b). Brain-derived neurotrophic factor (BDNF) ameliorates the suppression of thyroid hormone-induced granule cell neurite extension by hexabromocyclododecane (HBCD).	HBCDDs at 0.1 nM suppressed TH-induced neurite extension of granule cell aggregate in primary rat cerebellar granule cell aggregate cultures. BDNF ameliorated this effect in presence of T3. Results indicate that T3- stimulated increase in BDNF may be involved in HBCDD-induced impairment of TH-mediated neuritogenesis of granule cells.
Kim et al 2016. Influence of hexabromocyclododecane and 4-nonylphenol on the regulation of cell growth, apoptosis and migration in prostatic cancer cells <i>Toxicol In</i> <i>Vitro</i> , 32: 240-7.	HBCDDs may enhance progression of prostate cancer by modulating growth and migration of LNCaP prostate cells by acting on cell cycle and apoptosis.
Koike et al. (2013). Brominated flame retardants stimulate mouse immune cells in vitro. <i>J Appl Toxicol</i> , 33(12): 1451-9.	HBCDDs increased expressiom of T-cell receptor, MHC class II and CD86 as well as IL-4 production in splenocytes. Authors suggested that HBCDDs may induce or enhance immune/allergic responses by increasing antigen presentation-related molecule expression and IL-4 production.
Krivoshiev et al. (2016). Assessing in-vitro estrogenic effects of currently-used flame retardants.	HBCDDs stimulated proliferation of MCF-7 cells with an EC20 of 5.5 $\mu$ M. HBCDDs also inhibited 17B-oestradiol stimulated proliferation with an IC20 of 17.6 $\mu$ M.
Park et al. (2012). Cell growth of BG-1 ovarian cancer cells is promoted by di-n-butyl phthalate and hexabromocyclododecane via upregulation of the cyclin D and cyclin-dependent kinase-4 genes.	0.2 $\mu$ M HBCDDs stimulated proliferation of BG-2 ovarian cancer cells. HBCDDs upregulated mRNA for cyclin D and cyclin-dependent kinase-4 (cdk-4), which are downstream target genes of ER.
Reffatto et al. (2018). Parallel in vivo and in vitro transcriptomics analysis reveals calcium	Neurotoxic potential of HBCDDs was studied in mouse brain and, to separate direct effects from system responses (e.g. via hormones) two cell lines of mouse neuronal origin, NSC-19 and N2A, as well as in primary



and zinc signalling in the brain as sensitive targets of HBCD neurotoxicity.	hippocampal neuronal cultures. Transcriptome profiling of mouse brains and the two cell lines indicated that sex steroid regulation, and $Ca^{2+}$ and $Zn^{2+}$ homeostasis in glutamatergic neurons were preferentially affected. Follow up experiments on the two cell lines and isolated primary hippocampal neurons confirmed effects of HBCDDs on free [Zn <sup>2+</sup> ], glutamate-induced post-synaptic [Ca <sup>2+</sup> ], and zinc-stimulated post-synaptic [Ca <sup>2+</sup> ] release, which was almost completely blocked by 1 uM HBCDDs. The increase in cytosolic free [Zn <sup>2+</sup> ] could be partially blocked by an antioxidant, indicating ROS formation as a contributing factor.
Saegusa et al. (2012). Transient aberration of neuronal development in the hippocampal dentate gyrus after developmental exposure to brominated flame retardants in rats.	Female pregnant SD rats were fed a diet containing 0 (control), 100, 1,000 or 10,000 mg/L of HBCDDs from GD 10 to PND20. Effects on neuronal development was observed at 1,000 and 10,000 mg/L. Reelin <sup>+</sup> and NeuN <sup>+</sup> cells in interneurons of the dentate hilus increased temporarily,suggestive of abberant neuronal migration. Effect on reelin <sup>+</sup> cells was only seen at 1,000 mg/L (not at 10,000 mg/L). An increase in apoptotic bodies was observed at PND20 in the subgranular zone. At PND20 there was a small reduction in serum T3.
Suzuki et al. (2013). Similarities in the endocrine-disrupting potencies of indoor dust and flame retardants by using human osteosarcoma (U2OS) cell-based reporter gene assays.	CALUX reporter gene assays, based on U2OS cells were used to evaluate reporter gene assays used to evaluate activities of flame retardants on the human androgen receptor (AR), estrogen receptor a (ER $\alpha$ ), progesterone receptor (PR), glucocorticoid receptor (GR), and peroxisome proliferator-activated receptor $\gamma$ 2 (PPAR $\gamma$ 2). HBCDD showed relatively mild antagonistic effect on the AR and PR receptors.
Wilson et al. (2016). Do persistent organic pollutants interact with the stress response? Individual compounds, and their mixtures, interaction with the glucocorticoid receptor.	HBCDDs were not tested alone but in chemical mixture.
Wu et al. (2016b). Hexabromocyclododecane exposure induces cardiac hypertrophy and arrhythmia by inhibiting miR-1 expression via up-regulation of the homeobox gene Nkx2.5.	Zebrafish and H9C2 rat cardiomyocyte cells. Zebrafish embryos were exposed to 0, 2, 20 and 200 nM HBCDD in the water. Exposure to 20 and 200 nM resulted in cardiac hypertrophy and increased deposition of collagen. miR-1 might mediate HBCDD induced cardiac hypertrophyand arrhythmia via its target genes Mef2a and Irx5. H9C2 rat cardiomyocyte cells exposed to HBCDDs showed reduced Ca <sup>2+</sup> -ATPase activity and elevated Ca <sup>2+</sup> in the cytosol.
Zhang et al. (2016). Gene expression and metabolic responses of HepG2/C3A cells exposed to flame retardants and dust extracts at concentrations relevant to indoor environmental exposures.	Multiomics approach used to interrogate effects of dust extract and BFR mixture (including HBCDDs) in HepG2 cells. Exposure of cells to dust extacts induced expression of several CYP biotransformation enzymes. Such effects were not observed in the mixture of BFRs and it was concluded that other components of the dust extract were likely causing the observed effects.
Zhong et al. (2015). HBCD and PCBs enhance the cell migration and invasion of HepG2 via the PI3 K/Akt pathway.	HepG2 cells exposed to HBCDDs were assessed for cell viability, apoptosis, cell migration and invasion using cell counting kit-8 (CCK-8) assay, flow cytometry, cell scratch assay, respectively. 10 nM HBCDDs stimulated migration and invasion of HepG2 cells, increased protein expression of MMP9 and suppressed E-cadherin (CDH1) expression. HBCDD exposure increased stimulatory phosphorulation of protein kinase B and extracellular signal-regulated kinase 32 (ERK), and expression of mammalian target of rapamycin (mTOR).
Ziemiliska et al. (2012). Acute cytotoxicity evoked by tetrabromobisphenol A in primary cultures of rat cerebellar granule cells	Mild effect of 25 $\mu\text{M}$ HBCDDs on calcium uptake in CGCs.



outweighs the effects of polychlorinated biphenyls.	
Zimmer et al. (2011). In vitro steroidogenic effects of mixtures of persistent organic pollutants (POPs) extracted from burbot (Lota lota) caught in two Norwegian lakes.	HBCDDs were not tested alone but in chemical mixture.
An et al. (2013). The cytological effects of HBCDs on human hepatocyte L02 and the potential molecular mechanism.	High concentration of HBCDDs (>20 $\mu$ M) decreased survival of L02 hepatocytes, lower dose of HBCDDs ( $10^{-13}$ – $10^{-7}$ M) stimulated cell proliferation and up-regulation of PCNA protein expression level.
An et al. (2016). The "adaptive responses" of low concentrations of HBCD in L02 cells and the underlying molecular mechanisms.	L02 cells were able to acquire tolerance to a high concentration of $\alpha$ -HBCDD through pre-exposure to a lower dose. They provided evidence that the mechanism of this is through activation of the PI3K/Akt pathway, reduced phosphorylation of AMPK and increased phosphorylation of p38 MAPK.
An et al. (2014b). Oligomeric proanthocyanidins alleviate hexabromocyclododecane-induced cytotoxicity in HepG2 cells through regulation on ROS formation and mitochondrial pathway.	This study examined effects of oligomeric proanthocyanidins (OPCs) on cytotoxicity induced by HBCDDs. Whilst interesting it is not considered relevant to the RA of HBCDDs.
Canbaz et al. (2016). Indoor pollutant hexabromocyclododecane enhances house dust mite-induced activation of human monocyte- derived dendritic cells.	Simultaneous exposure of monocyte dendritic cells to house mite dust allergens and HBCDDs, and HBCDDs increased the expression of HLA-DR, co-stimulatory molecule CD86 and pro-inflammatory cytokine IL-8 depending on the dose of HBCDDs (1–20 $\mu$ M). An increase in IL-8 was observed at 1 $\mu$ M but other effects required higher concentrations (10–20 $\mu$ M).
Wang et al. (2016a). New Insights into the Cytotoxic Mechanism of Hexabromocyclododecane from a Metabolomic Approach.	Detailed study on HepG2 cells, starting with NMR and following up with hypothesis driven experiments and analyses. HepG2 cells exposed to reagent-grade HBCDD formula (purity: > 95%, Aladdin Industrial Corp.) at 0.05, 1 and 10 mg/L. HBCDD exposure resulted in amino acid metabolism, protein biosynthesis, fatty acid metabolism, and phospholipid metabolism. Beta oxidation of long-chain fatty acids was supressed, ATP production reduced, Na/K-ATPase inhibited, and uptake of amino acids and glucose impaired. Reduced beta oxidation was accompanied by increase in free fatty acids and increased synthesis of phospholipids. The authors suggest that most effects may be a results of reduced ATP production and consequential loss of Na/K-ATPase and Ca-ATPase activities, leading to reduced glucose and amino acid uptake by cells.
Georgantzopoulou et al. (2014). P-gp efflux pump inhibition potential of common environmental contaminants determined in vitro.	The calcein-acetoxymethyl ester (calcein-AM) assay in P-glycoprotein– overexpressing Madin–Darby canine kidney cells (MDCKII–MDR1) was exploited to study the inhibition of P-gp efflux pumps by HBCDDs (technical mixture) and other aquatic contaminants. Data suggested that HBCDDs may stimulate the P-gp efflux pump.
Kamata et al. (2018). Agonistic effects of diverse xenobiotics on the constitutive androstane receptor as detected in a recombinant yeast-cell assay	HBCDDs have a relatively low potential to activate CAR in a recombinant yeast assay.
Lille-Langoey et al. (2015). Environmental contaminants activate human and polar bear ( <i>Ursus maritimus</i> ) pregnane X receptors (PXR, NR1I2) differently.	COS-7 cells transfected with luciferase reporter gene construct was used to assess activation of polar bear and human PXR. A technical HBCDD mixture containing 81% $\gamma$ -HBCDD was tested. The HBCDD mixture had a moderate potency to activate PXR compared with other chemicals tested



	and maximum activation was 41% of that obtained by rifampicin exposure.
Montano et al. (2011). Effects of mixtures of persistent organic pollutants (POPs) derived from cod liver oil on H295R steroidogenesis.	HBCDDs were not tested alone but in chemical mixture.
van den Dungen et al. (2017). Persistent organic pollutants alter DNA methylation during human adipocyte differentiation.	DNA methylation and gene expression were studied in response to exposure to different POPs during human adipocyte differentiation. HBCDDs did not affect lipid accumulation in adipocytes. 1 µM HBCDD induced ALP activity in adipocytes.
van den Dungen et al. (2015). Steroid hormone related effects of marine persistent organic pollutants in human H295R adrenocortical carcinoma cells.	Effects of various POPs and mixtures thereof were studied in H295R adrenocortical carcinoma cells in relation to endocrine effects. HBCDDs did not affect any steroid hormone levels. 100 $\mu$ M HBCDDs induced expression of CYP19A1.
Kang et al. (2012). Induced growth of BG-1 ovarian cancer cells by 17beta-estradiol or various endocrine disrupting chemicals was reversed by resveratrol via downregulation of cell cycle progression.	Treatment of BG-1 cells with HBCDDs resulted in an increase of cell growth.
Li et al. (2017c). Neuroprotection by Taurine on HBCD-Induced Apoptosis in PC12.	Exposure of PC12 cells to 10 $\mu$ M HBCDDs reduced protein expression of Bcl-2, increased expression in Bax protein and activity of caspase-3. Taurine protected against these effects.
Liu et al. (2017). Taurine Alleviate Hexabromocyclododecane-Induced Cytotoxicity in PC12 Cells via Inhibiting Oxidative Stress.	Taurine protected against cytotoxicity induced by HBCDDs in PC12 cells through inhibition of oxidative stress.



# Annex A – Protocol for the risk assessments for human health related to the presence of brominated flame retardants (BFRs) in food

4980 The word file containing the protocol or strategy on the problem formulation and approach selected by

- 4981 the CONTAM Panel to update the previous risk assessments of brominated flame retardants (BFRs) in
- food is available on the EFSA Knowledge Junction community on Zenodo at: xxx 4982

# Annex B – Occurrence data a of HBCDDs in food submitted to EFSA

The excel file containing the occurrence data submitted to EFSA and dietary surveys per country and 4983 age group is available on the EFSA Knowledge Junction community on Zenodo at: xxx 4984

# Annex C – Benchmark dose (BMD) analysis

4985 The word file containing the BMD analysis is available on the EFSA Knowledge Junction community on Zenodo at: xxx 4986

### Annex D – Dietary surveys per country and age group available in the EFSA Comprehensive Database, considered in the exposure assessment

- The excel file containing the dietary surveys per country and age groups, and the exposure assessment 4987
- 4988 estimations is available on the EFSA Knowledge Junction community on Zenodo at: xxx