

Serum measures of hexabromocyclododecane (HBCDD) and polybrominated diphenyl ethers (PBDEs) in reproductive-aged women in the United Kingdom

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1 **SERUM MEASURES OF HEXABROMOCYCLODODECANE (HBCDD) AND**
2 **POLYBROMINATED DIPHENYL ETHERS (PBDES) IN REPRODUCTIVE-AGED**
3 **WOMEN IN THE UNITED KINGDOM**

4
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22

23 **Abstract**

24

25 We investigated the serum concentrations of two brominated flame retardants (BFRs) –
26 polybrominated diphenyl ethers (PBDEs) and hexabromocyclododecane (HBCDD) –in 59
27 women aged between 23 and 42 from the United Kingdom. We also collected demographic
28 data, including age, bodyweight and height in order to test for associations with BFR levels.
29 Temporal and global differences were also assessed using previously published data.

30 HBCDD was detected in 68% of samples with a mean concentration of 2.2 ng/g lipid (range
31 = <0.3 – 13 ng/g lipid). The dominant stereoisomer was α -HBCDD with an average
32 contribution of 82% (0-100%) towards Σ HBCDD, was followed by γ -HBCDD (average
33 contribution = 17%). PBDEs were detected in 95% of samples with a mean Σ PBDE (sum of
34 BDEs -28, -47, -99, -100, -153, -154 and -183) concentration of 2.4 ng/g lipid (range = <0.4 –
35 15 ng/g lipid). BDEs -153 and -47 were the dominant congeners, contributing an average of
36 40% and 37% respectively, to the average Σ PBDE congener profile.

37 Data from this study suggests that HBCDD levels decrease with age, it also suggests a
38 positive association between bodyweight and HBCDD levels, which likewise requires a
39 large-scale study to confirm this. The data also show that 10 years after their European ban,
40 PBDE body burden has begun to decrease in the UK. Whilst it is too early to draw any firm
41 conclusions for HBCDDs, they appear to be following a similar pattern to PBDEs, with levels
42 decreasing by a factor of >2.5 since 2010. Whilst the human body burden appear to be
43 decreasing, both PBDEs and HBCDD are still consistently detected in human serum, despite
44 legislative action limiting their production and use. This highlights the need to continuously
45 assess human exposure and the effectiveness of policy aimed at reducing exposure.

46 **1.0 Introduction**

47 Hexabromocyclododecane (HBCDD) and polybrominated diphenyl ethers (PBDEs) have
48 been used extensively worldwide as brominated flame retardants (BFRs) in a wide variety of
49 commercial, domestic and industrial applications. There are three commercial PBDE
50 formulations – Penta-, Octa- and DecaBDE. The main PBDE applications include electrical
51 and electronic equipment (EEE - such as TVs, PCs and small domestic appliances) ([European
52 Commission, 2011](#)), soft furnishings (e.g. sofas, mattresses, pillows and curtains) ([United
53 Nations Environment Programme \(UNEP\), 2010](#)) and in polyurethane foam (PUF) seat
54 fillings used in automobiles ([European Chemicals Bureau, 2000](#)). The primary use of
55 HBCDD is to flame retard expanded and extruded polystyrene (EPS/XPS) used in building
56 insulation foam ([European Chemicals Agency, 2009](#)). As of 2001 (the last reliable figures
57 publicly available), Europe accounted for 2 %, 16 %, 14 % and 57 % of the annual global
58 demand for Penta-, Octa-, DecaBDE and HBCDD respectively ([Bromine Science and
59 Environmental Forum \(BSEF\), 2003](#)).

60

61 Both PBDEs and HBCDD are lipophilic and resistant to metabolism allowing them to
62 bioaccumulate in the liver and other fatty tissues. They have long half-lives in humans of
63 approximately 664 – 2380 days and 64 days for PBDEs and HBCDD, respectively ([Geyer et
64 al., 2004](#)), and have been associated with adverse health effects in humans.. For example,
65 PBDEs are thought to disrupt levels of sex hormones, including luteinising hormone and
66 follicle stimulating hormone in men ([Meeker et al., 2009](#)), in addition to other toxic effects
67 including disruption to the liver, kidneys and thyroid gland; neurodevelopmental deficits
68 including inhibited foetal and infant development; and various cancers ([Costa, 2008](#)).
69 Furthermore, *in vitro* studies have demonstrated that doses as low as 5µM can induce
70 oxidative stress and disrupt steroidogenesis, with high level PBDE exposure resulting in

71 pregnancy failure ([Lefevre et al., 2016](#)). Exposure to the Penta-BDE formulation can activate
72 the aryl hydrocarbon (Ah) –receptor ([Gu et al., 2012](#)), cause a reduction in hepatic vitamin A
73 levels, impair neurodevelopment, and induce carcinogenesis ([D'Silva et al., 2004](#), [Hornung et
74 al., 1996](#)). Similarly, the OctaBDE formulation causes developmental toxicity, whilst the
75 DecaBDE formulation is believed to be the least toxic as it contains higher molecular weight
76 congeners that have relatively decreased cell membrane permeability, and are more readily
77 metabolised ([D'Silva et al., 2004](#), [Chevrier et al., 2013](#)). However, it is also believed that
78 higher brominated congeners (such as BDE-209, which makes up >95% of the Deca-BDE
79 formulation ([La Guardia et al., 2006](#))) can be broken down by physical and biological
80 processes to form lower brominated PBDE congeners that are found readily in Penta- and
81 Octa-BDE formulations ([D'Silva et al., 2004](#)). Data on human health effects of HBCDD
82 exposure is limited - [Eggesbø et al., 2011 reported that](#) it does not appear to have an effect on
83 the human thyroid ([Eggesbø et al., 2011](#)). However, [Dorosh et al. \(2011\)](#) suggested its
84 potential endocrine disrupting ability by altering oestrogenic activity.. Further, [Genskow et
85 al. \(2015\)](#) has suggested that HBCDD exposure damages dopaminergic neurons, with
86 consequences for neurological and endocrine system function, and there is evidence for
87 reduced birthweight and significant adverse neurodevelopment, including impaired motor
88 skills and increased anxiety levels in rodent models ([Maurice et al., 2015](#)).

89

90 Concerns over the toxicity of these BFRs led to bans on Penta- and Octa-BDE technical
91 products within Europe in 2003, and globally in 2009 under the UNEP Stockholm
92 Convention (SC) ([Stockholm Convention, 2009](#)). Significant restrictions were placed on the
93 DecaBDE technical product in 2008 (Deffree, 2008), and it was included in the SC in 2017
94 ([Chemical Watch, 2017](#)), alongside HBCDD in 2013 ([Health and Environment Alliance,
95 2013](#)). Whilst these bans will eventually lead to reduced exposure, they only prevent the new

96 manufacture and new use of these chemicals, meaning that BFRs will still be incorporated
97 into products already on the market, and currently in circulation. Both PBDEs and HBCDDs
98 are still regularly found in various indoor microenvironments across the world ([Sahlstrom et](#)
99 [al., 2015](#), [Johnson et al., 2013](#), [Ni and Zeng, 2013](#), [Harrad and Abdallah, 2015](#)), meaning that
100 humans will continue to be exposed to them for the foreseeable future. Given that exposure to
101 these chemicals can lead to a plethora of toxic health effects, it is vital that they are
102 continually monitored in general populations across the globe.

103

104 The aims of this study are to provide the first data on HBCDD exposure in the UK population
105 using human sera, and to provide updated assessment of human exposure to PBDEs and
106 HBCDDs in reproductive-aged women in the UK. The relationship between these BFRs and
107 various demographics (weight, body mass index (BMI), and age) will also be assessed to gain
108 insight into any potential health effects caused by target compounds. We include a temporal
109 assessment of HBCDD and PBDE body burdens in the UK, and a comparison of UK body
110 burdens with available data from other cross-sectional populations, globally.

111

112 **2.0 Materials & Methods**

113 *2.1 Sample Collection and Preparation*

114 This prospective cohort study was performed within the Hull IVF Unit, UK in 2014,
115 following approval by The Yorkshire and The Humber NRES ethical committee, UK
116 (approval number 02/03/043). A total of 59 women were recruited into the study, whose
117 baseline characteristics are shown in Table 1. Inclusion criteria were age 20-45 years, BMI
118 ≤ 35 and undergoing *in vitro* fertilisation. Patients with known immunological disease,
119 diabetes, renal or liver insufficiency, acute or chronic infections, or inflammatory diseases
120 were excluded from the study.

121

122 A fasting blood sample was collected on day 21 of the luteal phase of the cycle, and prior to
123 commencing IVF treatment. Samples were centrifuged, aliquoted, and stored at -80 °C.
124 Samples were shipped on dry ice to The Queensland Alliance for Environmental Health
125 Sciences at The University of Queensland, Australia for further analysis.

126

127 *2.2 Lipid Analyses of Samples*

128 Serum (300µL) was analysed for cholesterol (TC) and triglycerides (TG) by Sullivan
129 Nicolaides Pathology (SNP), Australia. Total lipid (TL) concentration (mg/dL) was
130 calculated using the following equation ([Phillips et al., 1989](#)).

$$TL = 2.27.TC + TG + 62.3$$

131

132 *2.3 Sample Extraction & Clean-up*

133 Five mL of serum was aliquoted into a 50 mL polypropylene centrifuge tube. Samples were
134 spiked with 5 ng each of internal standards (¹³C₁₂-labelled BDEs -28, -47, -99, -100, -153, -
135 154, -183, ¹³C₁₂-labelled α-, β- and γ-HBCDD). Samples were vortexed for approximately 1
136 minute and left to stand for 30 minutes. 6 mL acetonitrile, 3 mL milliQ, 5 g anhydrous
137 MgSO₄ and 1 g NaCl were added along with a ceramic homogenizer. Samples were manually
138 shaken for 1 minute prior to centrifuging at 4500 RPM for 8 minutes at 10 °C. The
139 supernatant layer was collected and transferred to a glass tube. The extract was evaporated to
140 near-dryness on a hot plate using a gentle stream of nitrogen and reconstituted in
141 approximately 1 mL hexane. 1 mL >98% concentrated sulfuric acid was added and the
142 sample was vortexed for at least 30 seconds. The aqueous and organic layers were left to
143 separate overnight at <4 °C. The supernatant layer was transferred directly onto a silica solid
144 phase extraction cartridge (Supelco LC-Si 3mL/500 mg), preconditioned with 6 mL

145 dichloromethane, followed by 6 mL hexane. The sample was allowed to load onto the
146 cartridge gravimetrically. Target compounds were eluted into a glass tube using 6 mL
147 hexane, followed by 8 mL dichloromethane at approximately 2 mL/min. The sample was
148 evaporated to near-dryness and reconstituted in 100 μ L iso-octane containing 2.5 ng $^{13}\text{C}_{12}$ -
149 PCB-141 and $^{13}\text{C}_{12}$ -TBBPA as recovery standards. After analysis for PBDEs by high
150 resolution gas chromatography coupled with high resolution mass spectrometry
151 (HRGC/HRMS) extracts were solvent exchanged into 100 μ L methanol and analysed for
152 HBCDD via liquid chromatography tandem mass spectrometry (LC-MS/MS).

153

154 *2.4 Instrumental Analysis*

155 For PBDE analysis by HRGC/HRMS, a Thermofisher TRACE 1300 gas chromatograph was
156 coupled to a Thermofisher DFS mass spectrometer. The injector was operated in splitless
157 mode with separation achieved on an Agilent DB-5ms column (30 m length x 0.25 mm in
158 diameter x 0.25 μ m film thickness). Experiments were conducted in MID mode at 10,000
159 resolution (10% valley definition). The inlet, transfer line and source were held at 250 $^{\circ}\text{C}$,
160 280 $^{\circ}\text{C}$ and 280 $^{\circ}\text{C}$ respectively. The flow rate was maintained at 1.0 mL/min. Details of
161 acquisition ions for PBDEs are outlined in the supporting information (SI, (Tables S1 and S2
162 respectively).

163

164 HBCDDs (α -, β - and γ -) were measured in serum samples using an AB/Sciex API 5500Q
165 mass spectrometer (AB/Sciex, Concord, Ontario, Canada) coupled to a Shimadzu Nexera
166 HPLC system (Shimadzu Corp., Kyoto, Japan). The mass spectrometer (MS) was operated in
167 multiple reaction monitoring mode using negative electrospray ionisation. A volume of 5 μ L
168 was injected. Separation was achieved using a Kinetex XB C18, 50 x 2.0 mm 1.7 μ m column
169 (Phenomenex, Torrance CA) using a mobile phase gradient of 85% methanol, ramping up to

170 100% methanol over 6 min and then holding for 4 min at a flow rate of 0.3 mL/min. Full MS
171 parameters have been provided previously ([Drage et al., 2017](#)).

172

173 *2.5 Quality Control*

174 A blank sample was extracted as every 6th sample (n=10), alternating between 5 mL of
175 MilliQ water (n=5) and 5 mL bovine calf serum (n=5). If a target compound was detected in
176 a blank at less than 5% of measured sample concentration, then no correction occurred; if
177 blank concentration was 5–25% of measured sample concentration, the blank concentration
178 was subtracted from that of the sample.

179

180 In the absence of a certified QC sample, method precision and accuracy were determined
181 using bovine serum (5mL, n=5) fortified with target compounds. 30 µL of a solution
182 containing 2 ng/mL of all target compounds in methanol was added to each aliquot, which
183 was then vortexed for 1 minute and left at <4 °C overnight. Good accuracy and precision was
184 found for all target analytes with average recoveries between 80-120% and a relative standard
185 deviation <15% (Table S2).

186

187 Internal standard recoveries of ¹³C-labelled HBCDDs were estimated by expressing their
188 ratio with ¹³C₁₂-TBBPA in the samples as a percentage of the same ratio in a non-extracted
189 side-spike (NESS). The recoveries of the remaining internal standards was calculated using
190 their ratio with ¹³C₁₂-PCB-141. Average recoveries ranged from 59% (¹³C₁₂-BDE-28) to 84
191 % (¹³C₁₂-BDE-154). Details of recoveries of all internal standards are provided in the SI
192 (Table S3).

193

194 *2.6 Statistical Analysis*

195 For the purposes of calculations of averages and all statistical testing where a compound was
196 below the limit of quantification (LOQ), values were set to half the limit of detection (LOD).
197 All statistical tests were computed using Microsoft Excel 2010 and SPSS for Windows
198 version 22.0.

199

200 **3.0 Results & Discussion**

201 This study reports the first data for HBCDD in human serum from the UK. Sum of α -, β -, and
202 γ -HBCDD (Σ HBCDD) was detected in 40 out of 59 samples at a concentration range of <0.3
203 – 13 ng/g lipid. The average concentration measured was 2.2 ng/g lipid, the geometric mean
204 was 0.75 ng/g lipid and the median was 1.8 ng/g lipid (Table 2).

205

206 The dominant stereoisomer was α -HBCDD with an average contribution of 82% (0-100%)
207 towards Σ HBCDD, was followed by γ -HBCDD (average contribution = 17%). β -HBCDD
208 was only detected in one sample where it contributed 25% to a Σ HBCDD concentration of 11
209 ng/g lipid. This stereoisomer pattern in human sera is consistent with previous studies from
210 Australia ([Drage et al., 2017](#)), India ([Devanathan et al., 2012](#)), Sweden ([Weiss et al., 2006](#)),
211 Canada ([Ryan et al., 2006](#)) and Japan ([Kakimoto et al., 2008](#)). The dominance of α -HBCDD
212 in human and other biotic samples is likely due to more effective transformation of β - and γ -
213 HBCDD to α -HBCDD through increased metabolic rate, combined with preferential
214 accumulation of the α -stereoisomer ([Fonnum and Mariussen, 2009](#)).

215

216 PBDEs were detected in measurable concentrations in 56 out of 59 samples with a Σ PBDE
217 (sum of BDEs -28, 47, -99, -100, -153, -154 and -183) concentration range of <0.4 – 15 ng/g
218 lipid. The average concentration was 2.4 ng/g lipid, the geometric mean was 1.4 ng/g lipid
219 and the median was 1.9 ng/g lipid (Table 3). BDEs -153 and -47 were the dominant

220 congeners, contributing an average of 40% and 37% respectively, to the average Σ PBDE
221 congener profile. The remaining PBDE content came from BDEs -100, -99 and -28 with
222 average contributions of 12%, 8.5% and 2.6% respectively. BDEs -154 and -183 were not
223 detected in any of the samples. The dominance of BDEs -47 and -153 in human serum is
224 consistent with much of the previous literature including previous measurements of serum
225 from the UK, USA ([Sjödín et al., 2004](#), [Sjödín et al., 2008](#)), Japan ([Akutsu et al., 2008](#)),
226 Greece ([Kalantzi et al., 2011](#)), Romania ([Dirtu et al., 2006](#)) and France ([Brasseur et al.,](#)
227 [2014](#)).

228 *3.1 Demographic trends: Age, Weight and BMI*

229 Despite the narrow age range of participants (23-42 years), Figure 1 suggests that there is a
230 decrease in HBCDD levels with age ($R^2 = 0.105$). However, a linear regression analysis
231 shows this to be insignificant ($p = 0.08$). There were no observed associations between PBDE
232 levels of participants and their age. This may be due to the limited sample size and age range
233 of participants in the study. Previous studies have demonstrated higher levels of PBDEs in
234 children and infants (Toms et al. 2009), however this study only investigated mothers of
235 child-bearing age.

236 A linear regression suggested a weak positive association between HBCDD levels and
237 bodyweight of the participant ($R^2 = 0.075$, $p = 0.036$; Figure S1a). However, when corrected
238 for height by using BMI instead of weight (Figure S1b), this association was no longer
239 significant ($R^2 = 0.057$, $p = 0.068$). There were no observed associations between bodyweight
240 or BMI and PBDE levels in participants from this study.

241 *3.2 Temporal Trends: Exposure in the United Kingdom*

242 Data on human exposure to HBCDDs in the UK is scarce, with only two previous studies
243 measuring breast milk concentrations from samples collected between 2008 and 2011
244 ([Harrad and Abdallah, 2015](#), [Abdallah and Harrad, 2011](#)), and prior to legislative ban.

245 Median Σ HBCDD concentrations from this study (1.8 ng/g lipid, 2014) were significantly
246 lower (ANOVA, $p < 0.0001$) than samples from 2008-2010 and 2010-2011 (3.8 and 5.2 ng/g
247 lipid, respectively) ([Abdallah and Harrad, 2011](#), [Harrad and Abdallah, 2015](#)). A recent study
248 of breastmilk from 10 women in UK collected in 2014-2015 by [Tao et al. \(2017\)](#) reported
249 similar HBCDD levels as the serum measures in our study (median: 2.9 ng/g lipid, range: 0.7-
250 7.1 ng/g lipid) (**Figure 2**). This is suggestive of a temporal trend to decreasing HBCDD
251 exposure in UK women. While there is some precedent for comparing serum and breast milk
252 biomarker concentrations as indicative of overall body burden, the samples were collected
253 over a relatively short period of time (2008 to 2015, across the 4 different studies), for a
254 comprehensive temporal assessment of exposure. Furthermore, HBCDDs were only subject
255 to legislative bans in 2013 – one year before samples were collected for this study ([Health
256 and Environment Alliance, 2013](#)), meaning that it is too early to assess the impact of
257 legislative action on HBCDD exposures in the UK population.

258

259 The range of Σ PBDE concentrations in this study are similar to those found in Newcastle-
260 Upon-Tyne, UK in the same year (1.0-16 ng/g lipid ([Bramwell et al., 2014](#))) and from
261 Birmingham in 2010, 2010-11 and 2014-15 ([Abdallah and Harrad, 2014](#), [Harrad and
262 Abdallah, 2015](#), [Tao et al., 2017](#)). Median Σ PBDE concentrations are approximately 3 times
263 lower than those found in serum (5.6 ng/g lipid ([Thomas et al., 2006](#))) and breast milk (6.3
264 ng/g lipid ([Kalantzi et al., 2004](#))) collected from Lancaster and London from 2001 to 2003
265 (**Figure 3**). This would suggest PBDE levels have fallen since the 2004 bans of Penta- and
266 Octa- BDE in the EU ([Birnbaum and Staskal, 2004](#)). However, breastmilk samples collected
267 in 2014-15 by [Tao et al. \(2017\)](#) contradict this finding with median concentrations of 5.8 ng/g
268 lipid. This is likely due to small sample size ($n=10$), and high variability both between-
269 individuals, and between geographical regions of the UK. However, it is pertinent to note

270 that in our study, there was a 95% detection rate of PBDEs in UK human serum 8 years after
271 these bans, and [Tao et al. \(2017\)](#) had a 100% detection rate in human milk more than a
272 decade later. This demonstrates that UK populations are still continuously exposed to PBDEs
273 despite legislative bans, and further action may be required to reduce body burden at the
274 population level. Similar temporal declines over a period of 10 years have also been
275 suggested for HBCDDs in Australia ([Drage et al., 2017](#)), ([Toms et al., 2012](#)), and Canada
276 ([Ryan and Rawn, 2014](#)), however both compounds are still regularly detected in humans
277 highlighting the need for constant monitoring of their concentrations in humans and the
278 environment.

279

280 *3.3 Comparison with global biomonitoring data*

281 Literature of serum measures of HBCDD is scarce, however there are a number of studies
282 reporting HBCDDs in milk from various countries (Table 1). The average concentration of
283 HBCDDs from this study (2.2 ng/g lipid) is at the lower end of the range of concentrations
284 found across the world (not detected – 43 ng/g lipid) and half the average concentration
285 worldwide (4.6 ng/g lipid). Concentrations were similar to breast milk collected in Canada in
286 1992-2005 (Ryan and Rawn, 2014) and serum from Belgium in 2007 (Roosens et al. 2009),
287 whilst they were 3-10 times higher than milk collected from the Philippines in 2008
288 (Malarvannan et al. 2013b), and India in 2009 (Devanathan et al. 2012). Furthermore,
289 Sahlström et al. (2014) did not detect HBCDD in any serum collected from 48 individuals in
290 Sweden between 2009 and 2010. Average HBCDD concentrations in serum collected in
291 South Korea from 2009-2010 (Kim and Oh, 2014) was approximately 4 times higher than
292 serum from this study, whilst milk collected in Spain from 2006-2007 was almost 20 times
293 higher (Eljarrat et al 2009).

294

295 Human biomonitoring studies for PBDEs are more prevalent in the literature than for
296 HBCDDs. The mean Σ PBDE (2.4 ng/g lipid) concentration from this study was at the lower
297 end of the range of Σ PBDE levels measured between 2009 and 2015 internationally (Table
298 2), but similar to (lipid normalised) Σ PBDE concentrations of breastmilk and serum from
299 other regions of the UK (([Bramwell et al., 2014](#), [Tao et al., 2017](#), [Harrad and Abdallah,](#)
300 [2015](#)), Norway ([Cequier et al., 2015](#)), Denmark ([Vorkamp et al., 2014](#)), and some regions of
301 China ([Wu et al., 2017](#), [Wang et al., 2016](#)). Serum levels of Σ PBDEs in this study were
302 approximately 2.5 times higher than breastmilk from Sweden ([Darnerud et al., 2015](#)), but
303 between 3 and 20 times lower than serum collected across USA ([Watkins et al., 2011](#), [Butt et](#)
304 [al., 2016](#), [Makey et al., 2014](#), [Zota et al., 2013](#), [Hurley et al., 2017](#)). Furthermore, serum from
305 6 individuals in Laizhou Bay, China, with no known occupational exposure were up to 300
306 times higher than from this study ([Wang et al., 2014](#)).

307

308 Major strengths of this study include relatively large sample size (59) as well as the the
309 pairing of BFR body burdens with demographic data such as age, weight and height. A
310 potential weakness of the study is the fact that all participants were undergoing *in vitro*
311 fertilisation. However, this was overcome by the fact that they were an otherwise normal
312 population, and patients with any known conditions were excluded from the study, making it
313 an otherwise normal population.

314

315 **4.0 Conclusions**

316 Here we present data confirming that reproductive aged women from the UK continue to be
317 exposed to both HBCDDs and PBDEs. Data from this study suggests that HBCDD levels
318 decrease with age, however further sampling of a wider age range would be required to
319 further investigate this. It also suggests a positive association between bodyweight and

320 HBCDD levels, which likewise requires a large-scale study to confirm this. The data suggests
321 that 10 years after their European ban, PBDE body burden has begun to decrease in the UK.
322 Whilst it is too early draw any firm conclusions for HBCDDs, they appear to be following a
323 similar pattern to PBDEs, with levels decreasing by a factor of >2.5 since 2010, a trend that
324 has also been observed in Australia. Whilst human body burdens appear to be decreasing,
325 both PBDEs and HBCDD are still consistently detected in human serum, despite legislative
326 action limiting their production and use, and highlighting the need to continuously assess
327 human exposure and the effectiveness of policy aimed at reducing exposure.

328

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333

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626 **Figures and Tables**

627 Table 1: Summary population characteristics

Number of participants	59
Age (years)	32 23-42
Height (cm)	165 148-191
Weight (kg)	70 (50-108)
BMI	
Normal (18.5-24.9)	22
Overweight (25-29.9)	32
Obese (30-34.9)	5
Pregnancy status	
Nulliparous	42
Primiparas	6
Miscarried/terminated	11
Smoking status	
Regular smoker	6
Non-smoker	53

628

Table 2 Σ HBCDD concentrations (ng/g lipid) in humans from this study and other studies internationally from 2002-2015

Country	Matrix	n	Mean	Range	Ref
<i>Europe</i>					
UK	Serum	59 individuals	2.2	<0.3 - 12.6	This Study
UK	Milk	10 individuals	3.2	0.7 - 7.1	Tao et al. (2017)
UK	Milk	25 individuals	5.95	1 - 22	Abdallah and Harrad (2011)
UK	Milk	10 individuals	6.5	0.3 - 21	Harrad and Abdallah (2015)
Belgium	Serum	16 individuals	2.9	<0.5 - 11	Roosens et al. (2009)
Belgium	Milk	1 pooled sample	1.5	n/a	Colles et al. (2008)
Czech Republic	Adipose	98 individuals	1.2	<0.5-7.5	Pulkrabova et al. (2009)
France	Milk	26	n/a	<1-5	Antignac et al. (2006)
France	Adipose	26	n/a	1-3	Antignac et al. (2006)
Greece	Serum	61 individuals	3.39	0.49-39	Kalantzi et al. (2011)
Ireland	Milk	11 pools	3.5	1.7-5.9	Pratt et al. (2013)
Netherlands	Cord Serum	12	0.2	0.2-4.3	Meijer et al. (2008)
Netherlands	Serum	91	0.2	0.1-0.36	Peters (2004)
Norway	Milk	10 individuals	n/a	nd-0.13	Polder et al. (2008a,b)
Norway	Milk	393 individuals	1.7	<0.2-31	Thomsen et al. (2009a)
Norway	Milk	12 individuals	n/a	0.25-2	Thomsen et al. (2003)
Norway	Milk	85 individuals	n/a	0.4-20	Thomsen et al. (2005)
Norway	Milk	67 Individuals	n/a	nd-3	Thomsen et al. (2009b)
Norway	Milk	193 individuals	1.1	0.1-31	Eggesbø et al. (2011)
Russia	Milk	23 individuals	0.71	nd-1.67	Polder et al. (2008a)
Russia	Milk	14 individuals	0.47	nd-1.15	Polder et al. (2008a)

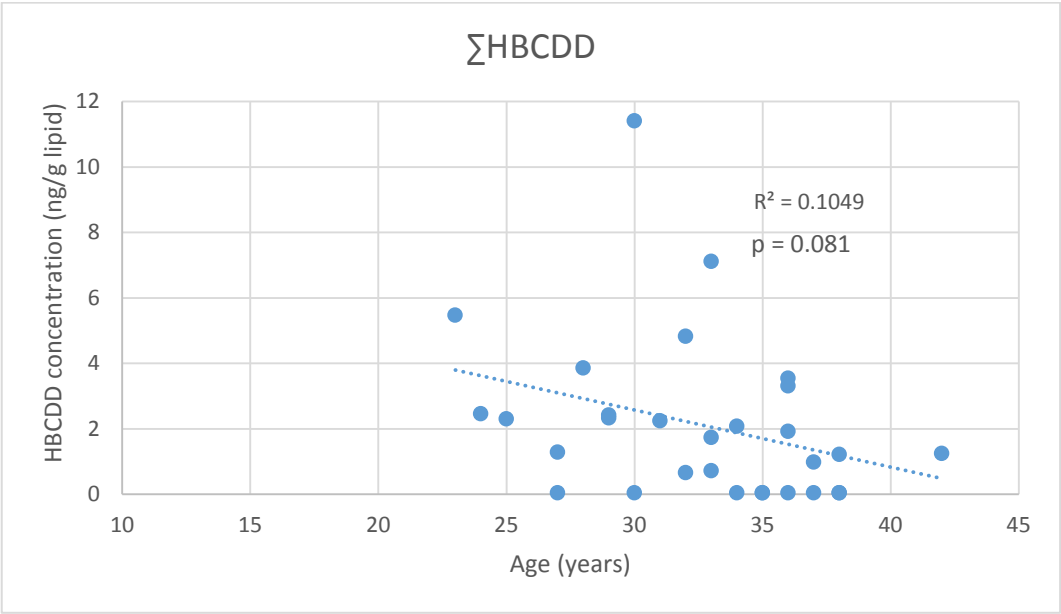
Spain	Milk	33 individuals	43	<LOQ-190	Eljarrat et al. (2009)
Sweden	Milk	14 pools	n/a	0.1-0.6	Fangstrom et al. (2008)
Sweden	Milk	204 individuals	n/a	0.09-10	Glynn et al. (2011)
Sweden	Serum	50 individuals	0.46	<0.24-3.4	Weiss et al. (2006)
Sweden	Serum	48 individuals	0	not detected	Sahlström et al. (2014)
<i>Asia</i>					
India	Milk	55 individuals	0.53	<0.05 - 13	Devanathan et al. (2012)
China	Milk	103 individuals	4.29	<LOQ-78	Shi et al. (2013a)
China	Serum	42 pools	0.86	<LOQ - 7.2	Shi et al. (2013b)
China	Milk	12 individuals	2.2	<LOQ - 5.5	Shi et al. (2013b)
Philippines	Milk	33 individuals	0.86	0.13 - 3.2	Malarvannan et al. (2009)
Philippines	Milk	30 individuals	0.21	<0.01-0.91	Malarvannan et al. (2013b)
South Korea	Serum	76 individuals	8.6	<dl-166	Kim and Oh (2014)
Vietnam	Milk	9 individuals	n/a	0.07 - 1.4	Tue et al. (2010)
Vietnam	Milk	4 individuals	n/a	0.11 - 0.97	Tue et al. (2010)
<i>Africa</i>					
South Africa	Milk	28 individuals	0.55	<0.23 - 1.4	Darnerud et al. (2011)
<i>North America</i>					
Canada	Milk	8	3.8	0.4-19	Ryan et al. (2006)
Canada	Serum	59 pools	1	0.33 - 8.9	Rawn et al. (2014)
Canada	Milk	34 individuals	1.8	0.1-28	Ryan and Rawn (2014)
USA	Milk	9	0.5	0.2-0.9	Ryan et al. (2006)
<i>Oceania</i>					
Australia	Serum	63 pools	3.1	<0.5-36	Drage et al. 2017
Australia	Milk	12 pools	6.6	<LOQ - 19	Toms et al. (2012a)
Australia	Serum	40 pools	0.45	<0.1-1.9	Drage et al. 2019

Table 3 Σ PBDE concentrations (ng/g lipid) in humans from this study and other studies internationally from 2009-2015

Country	Year	Matrix	n	Mean	Median	Range	Ref
<i>Europe</i>							
UK	2014	Serum	59 individuals	2.4	1.9	<0.2 - 15	This Study
UK	2012	Serum	20 individuals	N/A	2.4	1 - 16	Bramwell et al. 2014
UK	2012	Milk	8 individuals	N/A	4.8	1 - 28	Bramwell et al. 2014
UK	2010	Milk	25 individuals	5.9	5	0.2 - 26	Abdallah & Harrad 2014
UK	2010-11	Milk	10 individuals	5.1	3.7	1.3 - 13	Harrad & Abdallah 2015
UK	2014-2015	Milk	10 individuals	6.5	5.8	1.7 - 14	Tao et al. 2017
Denmark	2011	Serum	100 individuals	7.7	7.7	<LOQ - 18	Vorkamp et al. 2014
Norway	2012	Serum	46 individuals	3.6	2.3	0.1 - 23	Cequier et al. 2015
Sweden	2010	Milk	3 pools	0.73	0.77	0.58 - 0.84	Darnerud et al. 2015
<i>Asia</i>							
China	2011	Serum	12 pools	190	N/A	80-780	Wang et al. 2014
China	2012	Serum	6 individuals	N/A	13	4.3 - 42	Chen et al. (2014)
China	2013	Serum	10 pools	25	26	13 - 41	Li et al. 2017
China	2014	Serum	32 individuals	7.8	5.6	1.1 - 39	Wang et al. 2016
China	2014	Serum	9 individuals	5.6	N/A	0.42 - 27	Wu et al. 2017
<i>North America</i>							
USA	2009	Serum	31 individuals	28	N/A	3.5 - 350	Watkins et al. 2011
USA	2008-2010	Serum	43 individuals	28	N/A	0.71 - 250	Butt et al. 2016

USA	2010-2011	Serum	52 individuals	6.2	N/A	0.25 - 97	Makey et al. 2014
USA	2011-2012	Serum	36 individuals	52	N/A	N/A	Zota et al. 2013
USA	2011-2015	Serum	1253 individuals	23	N/A	N/A	Hurley et al. 2017

Figure 1 Individual Concentrations (ng/g lipid) of (a) Σ HBCDD and (b) Σ PBDEs vs their age (years)



ΣPBDEs

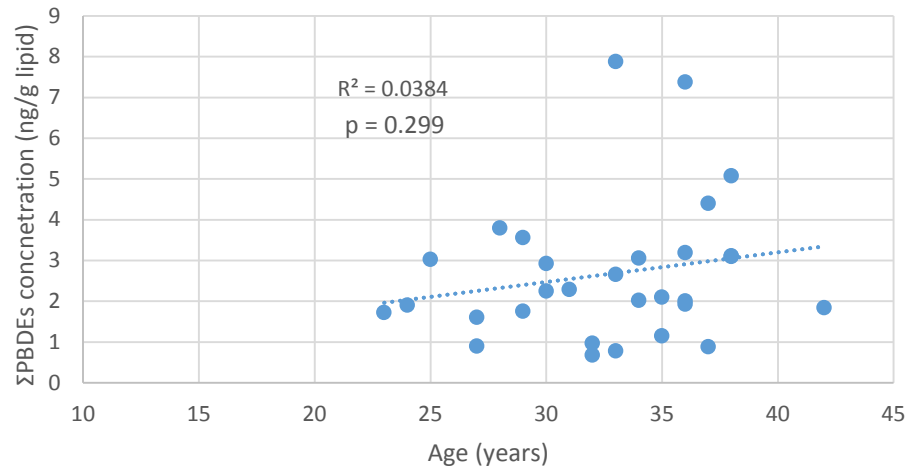


Figure 2 Temporal variation of mean HBCDD concentrations of serum and breast milk from UK women. Error bar denotes maximum concentration.

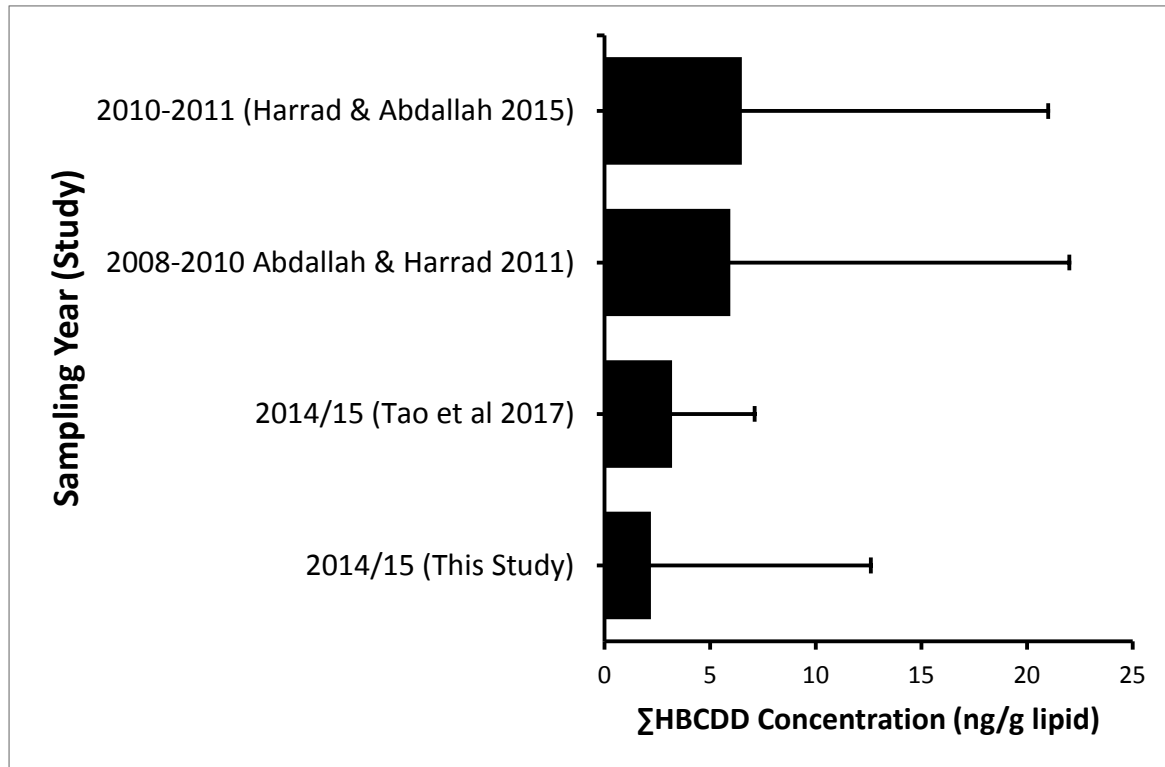


Figure 3 Temporal variation of mean PBDE concentrations of serum from UK adults from this study and previous studies. Error bar denotes maximum concentration.

