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Concentrations of Polybrominated Diphenyl Ethers, Hexabromocyclododecanes and Tetrabromobisphenol-A in Breast Milk from United Kingdom Women Do Not Decrease over Twelve Months of Lactation

Harrad, Stuart; Abdallah, Mohamed Abou-Elwafa

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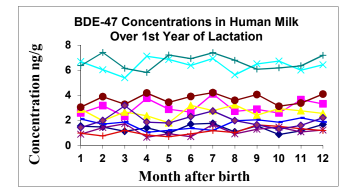
CONCENTRATIONS OF POLYBROMINATED DIPHENYL ETHERS, HEXABROMOCYCLODODECANES AND TETRABROMOBISPHENOL-A IN BREAST MILK FROM UNITED KINGDOM WOMEN DO NOT DECREASE OVER TWELVE MONTHS OF LACTATION

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6	Stuart Harrad ¹ * and Mohamed Abou-Elwafa Abdallah ^{1, 2}
7	
8	*Author for correspondence
9	email: <u>S.J.Harrad@bham.ac.uk</u>
10	Tel: +44 121 414 7298
11	
12	¹ School of Geography, Earth and Environmental Sciences,
13	University of Birmingham,
14	Birmingham, B15 2TT
15	United Kingdom
16	
17	² Department of Analytical Chemistry
18	Faculty of Pharmacy, Assiut University
19	71526 Assiut,
20	Egypt
21	

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25 Abstract

Conflicting evidence exists about whether concentrations of persistent organic chemicals in 26 human milk decrease over the course of lactation. This has implications for the timing of 27 28 sampling human milk for exposure assessment purposes. We examined this issue by measuring concentrations of polybrominated diphenyl ethers (PBDEs), hexabromocyclododecanes 29 (HBCDs), the HBCD degradation products tetrabromocyclododecenes (TBCDs), and 30 tetrabromobisphenol-A (TBBP-A) in human milk collected in 2010-11 from 10 first-time 31 mothers from Birmingham, UK. To evaluate whether concentrations varied significantly over the 32 first 12 months post-partum, 12 samples were taken - one per month - from each mother, 33 amounting to 120 samples overall. While concentrations of most of our target contaminants 34 displayed no significant variation (p>0.1) over the duration of our study, significant increases 35 were detected in concentrations of Σ TBCDs (p=0.029, average increase 1.4%/month) and BDE-36 37 153 (p=0.058, average increase 4.2%/month). When compared to data obtained from a different 38 set of UK mothers from a related but geographically wider catchment area sampled 39 contemporaneously to this study, the ratio of median concentrations of BDE-153 to BDE-99 was markedly lower in the current study (0.46 compared to 1.32). This may reflect unidentified 40 differences in exposure of the participants in the two studies. 41

42

43 Introduction

Polybrominated diphenyl ethers (PBDEs), hexabromocyclododecane (HBCD). 44 and tetrabromobisphenol-A (TBBP-A) are chemicals that have found extensive global use as flame 45 retardants incorporated within a wide range of goods and materials, such as electrical and 46 electronic items, and soft furnishings. The extensive use of such brominated flame retardants 47 (BFRs) has led to demonstrable contamination of both indoor and outdoor environments¹. 48 Contact with such contamination has led to human exposure via pathways such as inhalation of 49 air, and ingestion of both food and indoor dust, and resulted in the ubiquitous presence of BFRs 50 in humans^{2, 3}. As with other persistent organic chemicals, concerns exist about the presence of 51 BFRs in human milk. While studies to date of BFRs in human milk are consistent in 52 demonstrating that breast-fed infants are exposed substantially via ingestion of human milk^{4, 5}; 53 conflicting findings have emerged from the small number of studies that have examined the 54 temporal variation in concentrations of PBDEs in human milk from individual women over 55 extended periods of lactation. Essentially, while some authors have reported no discernible 56 consistent decrease of PBDE concentrations in human milk with increasing duration of lactation⁶, 57 ⁷; others report PBDE concentrations in human milk over the first 6-18 months of lactation to 58 decrease^{8, 9}. Whether such temporal declines in PBDE contamination of human milk occur is of 59 importance, as substantial reduction of concentrations over the course of lactation would mean 60 analysis of milk samples taken soon after birth will both overestimate exposure of the nursing 61 infant over the full period of lactation, as well as the reduction in mothers' body burdens as a 62 consequence of lactation. 63

Given this background, we report here a study of human milk samples taken every month overthe first year of lactation from 10 primiparas from Birmingham, UK. For each of these samples,

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we report concentrations of PBDEs (including BDE-209), α -, β -, and γ -HBCDs, the HBCD degradation products tetrabromocyclododecenes (TBCDs), and TBBP-A. To our knowledge, these are the first data on temporal variation in concentrations in human milk from the same women, of TBCDs, and TBBP-A, as well as the first such data for individual HBCD diastereomers.

71

72 Methodology

73 Sample collection

Breast milk samples (each comprising ~ 50 mL) were obtained from 10 adult volunteers via 74 Birmingham Women's Hospital Milk Bank, following approval of the study protocol by 75 76 Warwickshire Research Ethics Committee and the R&D Department in Birmingham Women's 77 NHS foundation trust. Informed consent was obtained from all participants before sample collection. Recruitment criteria were that mothers were healthy primiparas aged between 18 and 78 79 35, who were prepared to bring samples to the Milk Bank every month for the 12 month duration of the study. Samples collected in 2010-11 were kept in clean screw-capped glass containers and 80 transferred from the Milk Bank to the laboratory in special ice boxes then stored at -20°C until 81 the time of analysis. Due to ethical regulations, the samples were collected in a completely 82 anonymous fashion with all participant information kept strictly confidential. For the purposes of 83 84 this study, a total of 12 milk samples were collected at monthly intervals from each mother 85 commencing in the first month post-partum. All participants completed the study fully, with samples collected according to the same protocol by each participant throughout the course of 86 87 the study.

88

89 *Sample extraction*

Samples were first freeze-dried, following the addition of 25 ng of each of ¹³C-labeled BDE-47, 90 BDE-99, BDE-153, BDE-209, TBBP-A, α -, β - and γ -HBCDs as internal (surrogate) standards. 91 Accurately weighed aliquots of the freeze-dried samples (~ 2 g) were loaded into pre-cleaned 66 92 93 mL Accelerated Solvent Extraction (ASE 300, Dionex Inc., UK) cells containing 1.5 g florisil, 3 g alumina, 5 g anhydrous Na₂SO₄ and hydromatrix (Varian Inc., UK) to fill the void volume of 94 the cells and spiked with 25 ng each of d_{18} - α -HBCD and ¹³C-BDE-154 as QA/QC standards to 95 96 evaluate losses due to extraction and clean-up. The ASE cells were extracted with hexane:dichloromethane (1:9, v/v) at 90 °C and 1500 psi. The heating time was 5 minutes, static 97 98 time 4 min, purge time 90 s, flush volume 50%, with three static cycles. The lipid weight of the 99 studied samples was determined gravimetrically on separate aliquots using a standard procedure 100 (European Standard EN 1528-2, 1996; see supporting information for a summary).

101

102 *Extract purification*

103 The crude extracts were concentrated to 0.5 mL using a Zymark Turbovap® II (Hopkinton, MA, USA) then washed with 3 mL of 98 % sulfuric acid. After phase separation, the hexane layer was 104 transferred onto a florisil column (1.5 g of 5% deactivated florisil, 60-100 mesh, topped with 1g 105 106 of Sigma-Aldrich, UK)anhydrous sodium sulfate (Sigma-Aldrich, UK) and eluted with 25 mL of hexane:dichloromethane (1:1, v/v). The eluate was evaporated to dryness under a gentle stream 107 of N₂ and the dried extract reconstituted in 200 μ L of methanol containing 25 pg μ L⁻¹ of both 108 $^{13}C_{12}$ -BDE-100 and d_{18} -v-HBCD used as recovery determination (or syringe) standards to 109 determine recoveries of internal standards for QA/QC purposes. 110

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112 LC-MS/MS analysis of PBDEs, HBCDs and TBBP-A

Concentrations of target BFRs were determined using an LC-MS/MS system composed of a dual 113 pump Shimadzu LC-20AB Prominence liquid chromatograph equipped with SIL-20A 114 autosampler, a DGU-20A3 vacuum degasser coupled to a Sciex API 2000 triple quadrupole mass 115 spectrometer. The mass spectrometer was operated in atmospheric pressure photoionization 116 mode (APPI) for the determination of PBDEs, and in electrospray ionization mode (ESI) to 117 determine HBCDs, TBCDs, and TBBP-A. Full details of the multi-residue analytical 118 methodology used for separation and quantification of our target compounds can be found 119 elsewhere^{4, 5, 10, 11}. 120

121

122 *Quality assurance/quality control*

Full details of internal standard recoveries, field/method blanks, and method accuracy (measured by comparing our data with certified/indicative values for HBCDs and PBDEs for replicate analyses of NIST SRM2585, and matrix spikes at three concentration levels) have been reported previously^{4,5}. A summary of these data is provided as supplementary information. Limits of quantification (LOQ) are also provided as SI. Where an analyte was <LOQ in a sample, it was substituted for the purposes of statistical analysis by f x LOQ – where f = fraction of samples in which the analyte was present >LOQ.

130

131 **RESULTS AND DISCUSSION**

132 Concentrations and patterns of BFRs in this study compared to other studies

Table 1 presents a statistical summary of concentrations of our target BFRs in this study,together with comparative data from selected other studies of BFRs in human milk. Table SI-1

gives concentrations of our target BFRs in every sample analyzed in this study. Concentrations 135 of HBCDs and TBBP-A in this study fall within the range of those reported previously by our 136 research group for single milk samples collected within the first 3 months post-partum from 35 137 women in the West Midlands conurbation⁴. This confirms that concentrations in UK mothers 138 exceed ~ 5 fold those in Boston, USA¹⁵, and are a little higher than those in Irish mothers²⁰. 139 Moreover, the HBCD diastereomer pattern is consistent with our previous study and most other 140 studies worldwide. Specifically, α -HBCD was the predominant diastereomer observed 141 contributing between 60 and 89% of Σ HBCDs, with an average of 79%. For comparison, α -142 HBCD contributed 62-95% ∑HBCDs in our earlier study⁴. With respect to TBBP-A, 143 concentrations in this study are well below those of PBDEs and HBCDs - likely due to the short 144 human half-life of TBBP- A^{14} – and are consistent with the small number of previous reports of 145 the presence of TBBP-A in human milk^{4, 19, 20}. 146

Our data also confirm previous reports of the presence of BDE-209 in human milk for which 147 previous data are more limited than for other PBDEs, as well as providing only the second report 148 of the presence of TBCDs in humans. As stated previously, the origins of TBCDs in humans are 149 unclear; while they have been detected in indoor dust¹², they have also been shown to be formed 150 as HBCD metabolites in *in vitro* experiments involving cultured human hepatocytes¹³. Of note, is 151 the fact that median concentrations in this study of PBDEs 47, 153, and 209 are all lower than 152 reported in our earlier study of single milk samples⁵. Furthermore, while in our earlier study. 153 BDE-153 was more prevalent than BDE-99; the reverse was true in this study, and indeed 154 median concentrations of BDE-99 are higher in the current study. As all QA/QC criteria in this 155 study were met, and identical sampling, storage, and analytical protocols were followed in both 156 studies, we do not believe these differences to result from measurement artefacts. Moreover, the 157

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relative abundance of BDEs-99 and -153 in samples taken in this study during the first 3 months 158 post-partum (samples in our earlier studies were collected during this period), was not 159 discernibly different to those in later samples. Hence, the different timing of sample procurement 160 in the two studies does not account for the different congener pattern. In addition, while the 161 average age of participating mothers in the current study was 26.3 years, slightly lower than that 162 of the mothers in the earlier study (28.3 years), the difference was not significant (t-test, p>0.1). 163 Moreover, the mothers in both studies were all primiparas. Instead, we believe that the different 164 congener pattern in the current study, reflects unidentified differences in exposure of the 165 participants in the two studies. Pertinently, mothers in the current study did not participate in our 166 earlier survey, and came from a more geographically restricted area close to the Birmingham 167 Women's Hospital compared to participants in the earlier study who were recruited from across 168 the West Midlands conurbation. Interestingly, based on data suggesting that BDEs-47 and -99 169 display shorter human half-lives than BDE-153¹⁴; Thomsen et al¹⁸ identified an exposure 170 scenario consistent with our observations. Specifically, they hypothesized that mothers with low 171 172 background exposure due mainly to diet would be expected to be exposed to a higher proportion of BDE-153. Conversely, mothers receiving major direct exposure via contact with flame 173 retarded products and indoor dust would display higher relative abundance of BDE-47 and BDE-174 99 in breast milk. 175

176 Comparison of our data with recent studies conducted in the US⁷, New Zealand¹⁶, and elsewhere 177 in Europe¹⁷⁻²⁰ (Table 1) reveals mothers in this study to display concentrations of BDEs- 47, -99, 178 and -153 that are in line with those from other regions, with the exception of the USA, in which 179 concentrations of these congeners in humans are much higher than elsewhere in the world⁷. In 180 contrast, as noted previously⁵, the elevated concentrations of BDE-209 in UK indoor dust compared to other countries²¹ are not reflected in similarly elevated concentrations of this congener in human milk from UK mothers compared to women from other locations. While this may indicate poor bioavailability of BDE-209 from indoor dust²², we note that BDE-209 concentrations in West Midlands mothers are lower than those detected in 6 mothers from north east England¹⁷, and further data are required to ascertain whether the low BDE-209 concentrations in this study reflect a specific exposure pattern of our participants that is atypical of UK women in general.

188

189 Temporal variations in concentrations of BFRs in milk from individual mothers

To evaluate whether concentrations of individual BFRs exhibited significant variation over the 190 full year of lactation for our 10 participants, we plotted concentrations of individual BFRs in 191 192 each monthly sample as a percentage of the concentration detected in the first sample from the same mother. The plots obtained for BDE-47, BDE-153, *SHBCDs*, and *STBCDs* are provided as 193 194 Figure 1, with plots for other BFRs provided as supplementary information (Figure SI-1). With the exception of BDE-153 and Σ TBCDs, correlation analysis of these plots (conducted using 195 Excel for Mac 2008) revealed there to be no significant change in concentration with time over 196 the 12 months lactation covered by this study. In contrast, concentrations of both BDE-153 and 197 Σ TBCDs show a significant increase (average increase 4.2% and 1.4%/month respectively) in 198 concentrations during our study (p=0.058 and p=0.029 respectively). 199

A previous study reported concentrations of tri- through-deca-PBDEs in samples of human milk from 10 women in Oslo, Norway at monthly intervals on between 3 and 10 separate occasions per mother⁹. The authors reported that when normalized to concentrations in the first sample of each mother, concentrations of PBDEs 28, 47, 99, 100, 153, and 154 in subsequent samples

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displayed a significant decrease over the period studied. These decreases ranged between 1.7%/month for BDE-153 and 4.7%/month for BDE-154. A similar study of primiparae women from California, reported PBDE concentrations in milk sampled every 4 weeks on 6 occasions from birth for 9 women, and in milk sampled on 2 occasions at varying time intervals between 18 and >85 weeks from birth for a further 9 women⁸. The authors of this study reported concentrations of BDE-47 to decline significantly by 3%/month on average, and 2%/month on average for both BDE-99 and BDE-100.

Clearly, our data contrast with these studies. However, they are consistent with the observations 211 of two other studies^{6, 7}. In the first of these, in which milk was sampled from 9 mothers on 212 between 2 and 4 occasions up to nearly 1 year post-partum, concentrations of BDE-153 showed 213 an increase in 7 mothers (p=0.09), but no clear, consistent decrease or increase was observed for 214 any other targeted PBDEs⁶. In the second study, concentrations of PBDEs were measured in milk 215 samples collected from 83 women at both 3 and 12 months post-partum⁷. As in the first study, 216 while concentrations of BDE-153 were significantly higher in the 12 month samples (p=0.005), 217 no significant change was observed for all other monitored PBDEs. 218

LaKind et al⁶ offered two hypotheses to account for whether concentrations of POPs like PBDEs 219 will change over the duration of lactation. The congener pattern observed in our study, whereby 220 no significant temporal change was observed for most contaminants, but significant increases 221 were seen for BDE-153 and Σ TBCDs; may conceivably be reconciled with the first of these 222 hypotheses, that fluctuations in mothers' intake over the period monitored can influence 223 concentrations in human milk. While little is known about human exposure to TBCDs, as 224 highlighted above, Thomsen et al¹⁸ identified that for PBDEs, a transition from exposure driven 225 principally by indoor pathways such as direct contact with flame-retarded goods and indoor dust. 226

to background exposure driven mainly by diet, could result in an increase in the relative
abundance of BDE-153 compared to BDEs 47 and 99. Given the hypothesized time-lag between
reductions in PBDE exposure via the diet following reductions in indoor contamination²³ it is not
inconceivable that our data is an indication of a response of the exposure of the UK population to
PBDEs as a result of actions taken within the EU in the mid-2000s to restrict manufacture and
use of the Penta- and Octa-BDE products.

The second hypothesis advanced by LaKind et al is that substantial post-partum weight loss can 233 lead to increased (or at least less decreased) concentrations as a result of increased remobilization 234 of contaminants associated with adipose tissue. Although due to the ethical constraints of our 235 study, we do not have any information on the weight of our study participants, this does not at 236 first seem a credible explanation for our data, given that concentrations increased for only two of 237 238 our target contaminants. However, as these contaminants include BDE-153, for which recent human biomonitoring studies consistently indicate is constituting an increasing proportion of the 239 Σ PBDE burden in human tissues^{5, 18, 20}, as a result of its greater persistence relative to other 240 congeners^{14, 24}; it is possible that our data reflect the impact of post-partum weight loss on BFR 241 concentrations in our participants over a year of lactation. This could conceivably result in no 242 overall temporal change for most BFRs, but an increase for BDE-153 given the temporal 243 increase of this congener relative to other PBDEs reported elsewhere. If true, then this implies 244 enhanced persistence in humans of Σ TBCDs relative to the parent HBCDs and the related 245 PBCDs. In summary therefore, while we are unable to provide a definitive explanation for our 246 observations; neither hypothesis outlined by LaKind et al can be ruled out. 247

This study – which has the highest temporal resolution of any conducted hitherto - provides
substantial evidence that in a small group of UK mothers, concentrations in human milk of most

PBDEs, HBCDs, and TBBP-A do not change significantly over the first year of lactation. In 250 contrast, concentrations of the more persistent BDE-153 congener, and the HBCD degradation 251 product TBCDs display a significant increase over the same period. While a larger study 252 253 involving more mothers is required to confirm our findings, our data suggest that for most of the major BFRs included in our study, human milk samples taken at any point in the first year post-254 partum will provide a reasonably representative measure of the exposure of the mother and the 255 nursing infant. The reasons for the observed increase in concentrations of two BFRs are not 256 clear, but may be related to *in vivo* metabolic production of these contaminants. In practical 257 terms, the absence of any significant decline in BFR concentrations over the first year of 258 lactation, suggests that advice to nursing mothers to practice pumping and discarding milk in the 259 early stages of lactation (referred to colloquially as "pump and dump") in order to minimize 260 infant exposure to such contaminants⁸, is unlikely to be successful. 261

262

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268

269 Supplementary Information

270 Concentrations of all target contaminants in every sample analyzed; plots of concentrations of 271 BDE-99, BDE-209, TBBP-A, α -, β -, and γ -HBCDs in each monthly sample as a percentage of 272 the concentration detected in the first sample from the same mother; as well as detailed

- descriptions of analytical methodology and QA/QC data. This information is available free of
 charge via the Internet at http://pubs.acs.org/.
- 275

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Location (n=number of participants)	Year of Sample Collection	Parameter	BDE- 47	BDE- 99	BDE- 153	BDE- 209	α- HBCD	β- HBCD	γ- HBCD	ΣTBCDs	TBBP-A
Birmingham, UK		5 th %ile	0.89	0.38	0.07	0.05	1.64	0.09	0.12	0.01	0.03
(n=10, 12 samples		Median	2.30	1.04	0.48	0.08	4.16	0.40	0.76	0.11	0.04
from each		Average	2.97	1.58	0.51	0.14	5.27	0.48	0.79	0.14	0.06
participant), this study	2010-11	95 th %ile	6.93	4.26	1.09	0.39	15.1	1.49	2.10	0.38	0.17
		DF* (%)	100	100	100	63	100	100	100	92	61
Birmingham, UK $(n=35)^4$	2010	Median	NR	NR	NR	NR	3.17	0.30	0.56	0.14	<0.04
Birmingham, UK (n=34) ⁵	2010	Median	2.80	0.69	0.91	0.25	NR	NR	NR	NR	NR
North East England, $UK (n=6)^{17}$	2011-12	Median	2.05	0.97	0.93	0.70	NR	NR	NR	NR	NR
Boston, MA, USA (n=43) ¹⁵	2005-06	Geometric Mean	NR	NR	NR	NR	0.71	0.08	0.20	0.05	<0.03NR0.55 ^a
Central North Carolina, USA $(n=303)^7$	2004-06	Median	28	5	6	NR**	NR	NR	NR	NR	NR
Norway (n=393) ¹⁸	2001-2009	Median	0.99	0.27	0.45	0.32 ^b	NR	NR	NR	NR	NR
France $(n=23)^{19}$	2005	Median	NR	NR	0.83	1.50	NR	NR	NR	NR	0.17
Ireland $(n=11^{\circ})^{20}$	2010	Median	1.11	0.27	1.00	0.77 ^d	2.59 ^e	0.42^{e}	0.43 ^e	NR	0.05 ^e
New Zealand (n=33) ¹⁶	2010	Median	2.14	0.56	0.75	0.19	NR	NR	NR	NR	NR

Table 1: Summary of Concentrations (ng g⁻¹ lipid weight) of Target BFRs in Human Milk in this Study and Others 359

^a Range reported; geometric mean not reported due to low detection frequency for TBBP-A (35%) ^b BDE-209 measured in a subset of 46 samples 360

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^c 11 pooled samples analyzed comprising milk from 109 primiparas 362

^d BDE-209 analyzed in a subset of 10 pooled samples 363

^e Lower bound average concentrations (i.e. where concentration below detection limit, concentration assumed to be zero) 364

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* DF refers to detection frequency; ** NR indicates the value was not reported.

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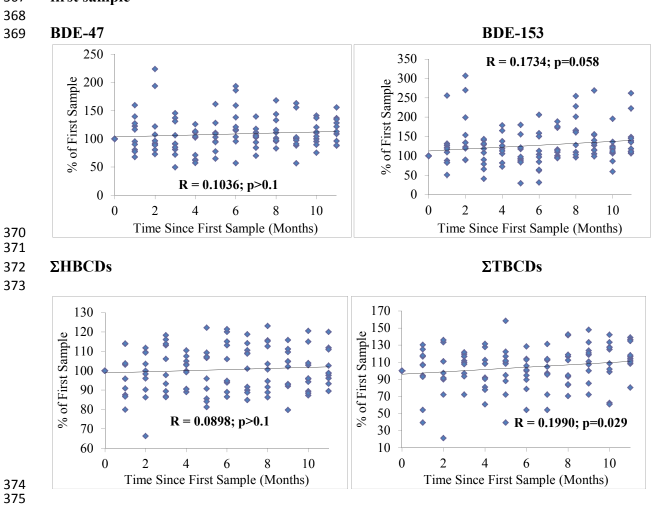


Figure 1: Concentrations of BDE-47, BDE-153, ΣHBCDs, and ΣTBCDs normalized to the first sample