

# Atmospheric concentrations of polychlorinated biphenyls, brominated flame retardants, and novel flame retardants in Lagos, Nigeria indicate substantial local sources

Akinrinade, Olumide Emmanuel; Stubbings, William A; Abdallah, Mohamed Abou-Elwafa; Ayejuyo, Olusegun; Alani, Rose; Harrad, Stuart

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1 **Atmospheric Concentrations of Polychlorinated Biphenyls,**  
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3 **Brominated Flame Retardants, and Novel Flame Retardants in**  
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6 **Lagos, Nigeria Indicate Substantial Local Sources**  
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12 5 Olumide Emmanuel Akinrinade<sup>1,2</sup>, William A. Stubbings<sup>1</sup>, Mohamed Abou-Elwafa

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14 6 Abdallah<sup>1</sup>, Olusegun Ayejuyo<sup>2</sup>, Rose Alani<sup>2</sup>, Stuart Harrad<sup>1\*</sup>  
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19 8 <sup>1</sup>School of Geography, Earth, and Environmental Sciences,  
20

21  
22 9 University of Birmingham,  
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24 10 Birmingham B15 2TT,  
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27 11 UK  
28

29  
30  
31 13 <sup>2</sup>Department of Chemistry,  
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33  
34 14 University of Lagos,  
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43 18 \*Corresponding Author

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45 19 E-mail address: S.J.Harrad@bham.ac.uk  
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26 **Abstract**

27 Polychlorinated biphenyls (PCBs), brominated flame retardants (BFRs) like polybrominated  
28 diphenyl ethers (PBDEs), hexabromocyclododecane (HBCDD), and novel flame retardants  
29 (NFRs) like decabromodiphenyl ethane (DBDPE) are ubiquitous environmental pollutants.  
30 Despite this, little is known about their concentrations in outdoor air in the African continent.  
31 To address this knowledge gap, concentrations of BFRs, NFRs, and PCBs were measured in  
32 outdoor air at 8 sites located within the metropolitan area of Lagos, Nigeria. Concentrations  
33 of  $\sum_8$ BDEs,  $\sum$ HBCDD,  $\sum_7$ NFRs and  $\sum_8$ PCBs were: 21 – 750 (median = 100)  $\text{pg}/\text{m}^3$ , < 12 –  
34 180 (median = < 12)  $\text{pg}/\text{m}^3$ , 34 – 900 (median = 300)  $\text{pg}/\text{m}^3$  and 85 – 460 (median = 300)  
35  $\text{pg}/\text{m}^3$ , respectively. Decabromodiphenyl ether (BDE-209, range: <16 - 620  $\text{pg}/\text{m}^3$ , median =  
36 71  $\text{pg}/\text{m}^3$ ) and DBDPE (range: <37 - 890  $\text{pg}/\text{m}^3$ , median = 280  $\text{pg}/\text{m}^3$ ) were the dominant  
37 BFRs detected, while the non-Arochlor PCB 11 (range: 49 – 220  $\text{pg}/\text{m}^3$ , median = 100  $\text{pg}/\text{m}^3$ )  
38 was the dominant PCB. To the authors' knowledge, these are the first data on the non-  
39 Arochlor PCB 11 in outdoor air in Africa. In general, concentrations of all target  
40 contaminants in this study were within the range reported elsewhere in Africa and worldwide.  
41 Likely due to the tropical climate of Lagos, no seasonal variation in concentrations was  
42 discernible for any of the target contaminants. While concentrations of PBDEs and some  
43 NFRs were correlated with population density, concentrations of PCBs appear more impacted  
44 by leaks from electrical transformers and for PCB 11 to proximity to activities like textile  
45 factories that produce and use dyes.

46  
47 **Key words:** PCBs, BFRs, PBDEs, HBCDD, DBDPE, urban air, Africa

## 52 1. Introduction

53 Between 1929 and the mid-1980s, polychlorinated biphenyls (PCBs) were  
54 manufactured as various commercial mixtures around the world. Applications included as  
55 dielectric fluids in electric power transformers and capacitors, as well in lubricating fluids,  
56 adhesives, building sealants, plasticisers, fire proofing agents, paints, and ink products  
57 (Breivik et al., 2016, Wong and Warner, 2010; Zhang et al., 2011).

58 Polybrominated diphenyl ethers (PBDEs) and hexabromocyclododecane (HBCDD)  
59 have been two of the most widely used brominated flame retardants (BFRs) applied to  
60 consumer products such as soft furnishings and electrical and electronic equipment (EEE), to  
61 help meet fire safety regulations (BSEF, 2020; Fatunsin et al., 2020; Muenhor and Harrad,  
62 2018). Commercial PBDE formulations include: pentaBDE (comprising primarily tetra-,  
63 penta-, and hexa-BDEs), octaBDE (comprising mainly hexa-, hepta- and octa-BDEs), and  
64 decaBDE (principally decabromodiphenyl ether (BDE-209)), with the latter the most widely  
65 used (Abb et al., 2011; Wemken et al., 2019). HBCDD was applied mainly as an additive to  
66 expanded and extruded polystyrene (EPS/XPS) building insulation foam. To a lesser degree it  
67 was also applied to high impact polystyrene (HIPS) used in enclosures for electronic  
68 equipment as well as a back-coating for fabrics like sofa covers and curtains (Desborough et  
69 al., 2016; Stubbings and Harrad, 2019). Technical HBCDD comprises predominantly  $\gamma$ -  
70 HBCDD (70%), along with lower proportions of  $\alpha$ - and  $\beta$ -HBCDD (de Wit et al., 2010).

71 In light of evidence of their toxicity, persistence, and potential to undergo  
72 bioaccumulation and long-range atmospheric transport; PCBs, PBDEs, and HBCDD are  
73 listed as persistent organic pollutants (POPs) under the Stockholm Convention of the United  
74 Nations Environment Programme (UNEP) (UNEP, 2020). Listing of these chemicals as POPs  
75 led to ceasing of their manufacture and deliberate use at various points since 1979. With  
76 respect to PBDEs and HBCDD (referred to collectively as POP-BFRs), restrictions on their

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77 manufacture and use without concomitant changes in fire safety regulations has led to  
78 increased use of so-called ‘alternative’, ‘emerging’, or ‘novel’ flame retardants (NFRs) in a  
79 similar range of applications (Covaci et al., 2011; Zuiderveen et al., 2020). Examples of  
80 NFRs include: decabromodiphenyl ethane (DBDPE), dechlorane plus (DP),  
81 hexabromobenzene (HBBz), pentabromobenzene (PBBz), pentabromoethylbenzene (PBEB),  
82 and pentabromotoluene (PBT). Both DBDPE and DP have found use as replacements for  
83 decaBDE in HIPS, back-coated furniture fabrics, electrical hard plastic connectors in  
84 televisions and computer monitors, and wire coatings (de Wit et al., 2010; Sverko et al.,  
85 2011). Meanwhile, HBBz, PBEB, and PBT are all additive flame retardants applied to paper,  
86 thermoset resins in circuit boards, textiles, adhesives, wire and cable coating, polyurethane  
87 foams, thermoplastic resins, polyester resins, and paint emulsions (de Wit et al., 2010).

88 Information regarding the toxicity of such NFRs remains limited. However, the  
89 USEPA’s 2014 assessment of alternatives to decaBDE, rates DBDPE a similarly high hazard  
90 to decaBDE with respect to developmental toxicity (USEPA, 2014). This is concerning as the  
91 endpoint that drives the EPA’s reference dose for decaBDE is neurodevelopmental toxicity  
92 (USEPA, 2008). Moreover, *in silico* assessment suggests HBBz, PBBz, PBEB, and PBT are  
93 potentially persistent, bioaccumulative, and toxic (PBT) compounds (Gramatica et al., 2016),  
94 with DP thought capable of long-range atmospheric transport (Wang et al., 2016).

95 While data on concentrations of PCBs, PBDEs, and HBCDD in outdoor air from East  
96 Asia, Europe, and North America are relatively abundant; such information is far scarcer  
97 elsewhere, particularly in Africa (Akinrinade et al., 2020). This dearth of information for  
98 Africa is an important oversight, given concerns about issues such as: poor management of  
99 waste electrical and electronic equipment (WEEE) containing FRs, and of obsolete electric  
100 transformers containing PCBs. Moreover, there exist increasing reports and concerns about  
101 the environmental presence of non-Aroclor PCBs like PCB 11 (3,3’-dichlorobiphenyl)

102 thought to be formed from diarylide dyes and/or degradation of more highly chlorinated PCB  
103 congeners (Anh et al., 2021; Basu et al., 2009). To our knowledge, no data yet exist on  
104 atmospheric concentrations of PCB 11 in Africa.

105 With respect to recent data for outdoor air in Africa, the GAPS (Global Atmospheric  
106 Passive Sampling) monitoring program has recently reported concentrations of selected  
107 PBDEs, HBCDD, and NFRs in Egypt and Nigeria (Saini et al., 2020), while the MONET  
108 (MONnitoring NETwork) programme has reported PCB concentrations in a semi-urban  
109 location in Sheda, Abuja, Nigeria, in addition to several locations in Africa (Klánová et al.,  
110 2009; White et al., 2021). While studies elsewhere have demonstrated both seasonal and  
111 within-conurbation spatial variability in concentrations of PCBs and BFRs in outdoor air  
112 (Currado and Harrad, 2000; Hoh and Hites, 2005; Carlson and Hites, 2005; Harner et al.,  
113 2006; Motelay-Massei et al., 2005; Harrad and Hunter, 2006), such data are limited for Africa  
114 (e.g. Katima et al, 2018), and it is plausible that the temperature-driven summer peaks in  
115 concentrations observed in more temperate locations, will not be replicated in tropical and  
116 sub-tropical African cities where seasonal temperature differences are not pronounced.

117 Against this backdrop, the objectives of this study are to provide baseline data on the  
118 concentrations of BFRs, NFRs, and PCBs between May 2019 and May 2020 at 8 locations in  
119 the metropolitan area of the largest city in Nigeria (Lagos). Measurement of concentrations in  
120 these samples allowed us to place the pollution levels in Lagos within an international  
121 context. Moreover, by measuring concentrations in outdoor air during up to four 3-month  
122 periods at each location, we were able to examine seasonal and spatial variation in  
123 atmospheric concentrations of BFRs, NFRs, and PCBs within a major African urban centre.

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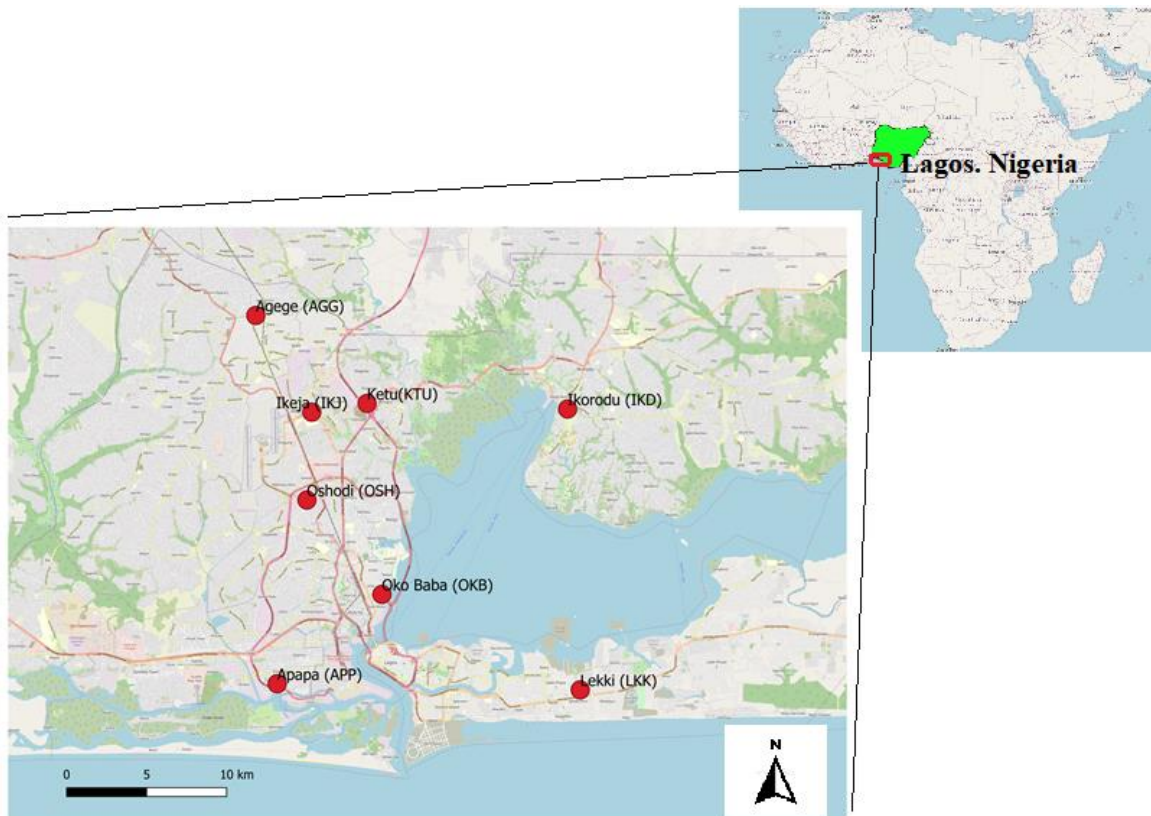
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127 **3. Materials and methods**

128 **3.1 Sample locations**

129 Lagos, Nigeria, is an urban centre with an approximate population of 24 million, housing  
130 numerous industrial activities. For the purposes of this study, we selected 8 locations across  
131 Lagos that reflected different levels of industrialisation, population density, and other  
132 anthropogenic activities. The sampling locations are: Agege (AGG), Apapa (APP), Ikeja  
133 (IKJ), Ikorodu (IKD), Lekki (LKK), Ketu (KTU), Oshodi (OSH), and Oko baba (OKB). The  
134 geographical location of these sites is shown in Figure 1, with relevant information related to  
135 population density and putative sources of each centre provided in Table 1. In brief, AGG,  
136 APP, IKJ, and OSH are mixed industrial – residential locations, while IKD, KTU, OKB, and  
137 LKK are more residential in character, with KTU and OKB additionally potentially impacted  
138 by an inactive waste dump/landfill and large-scale wood burning activities respectively.



140  
141 **Figure 1:** Sampling locations in Lagos, Nigeria

142 **Table 1: Sample locations and attributes**

	<b>AGEGE (AGG)</b>	<b>APAPA (APP)</b>	<b>IKEJA (IKJ)</b>	<b>IKORODU (IKD)</b>	<b>KETU (KTU)</b>	<b>LEKKI (LKK)</b>	<b>OSHODI (OSH)</b>	<b>OKO BABA (OKB)</b>	<b>LAGOS (IN GENERAL)</b>
24									
25									
26	Mix residential and industrial areas	Mix residential and industrial areas	Mix residential and industrial area	Majorly a residential area	Majorly a residential area	Majorly a residential area	Mix residential and industrial area	Majorly a residential area	Large urban population
27									
28									
29									
30									
31	Prominent abattoir	Proximate to port location	Home to Government offices	Sample location proximate to electric transformers and textile factory	High vehicular spots	Well-developed urban site	High vehicular spots	Sample location proximate to the University of Lagos	High vehicular spots
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33									
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42									
43	Large local market	Large local market	State capital	-	Proximate to inactive dumpsite/landfill	-	Market Centre	Mass burning of wood wastes from local wood factories	High industrial complex
44	Number of samples collected		Major industrial complex	-			Major industrial complexes	-	Wastes mostly managed by open land burning
45	Population <sup>a</sup>	1,415,547	715,792	888,903	944,158	1,280,646	1,554,604	862,524	24,051,762
46	Population density <sup>a</sup>	60,768.47	13,568.42	12,995.19	1,997.23	11,073.63	152.43	32,083.03	4,906.78
47									
48	143								
49	144								

<sup>a</sup> Lagos Bureau of Statistics (2017)



### 145 3.2 Sampling procedures

146 Passive sampling of outdoor air utilised polyurethane foam (PUF) discs enclosed in stainless  
147 steel double dome housing (Klanova et al., 2009; Melymuk et al., 2011; Drage et al., 2016).  
148 Prior to deployment, PUF discs were pre-cleaned by soxhlet extraction for ~ 24 hours with  
149 HPLC grade hexane. Samplers were deployed for 90 days in each of the following periods:  
150 May-July 2019, August-October 2019, and November 2019-January 2020, and additionally  
151 for 100 days between February 2020-May 2020. At the end of each sampling event, samplers  
152 were harvested by careful removal of the PUF disc, which was wrapped in hexane-rinsed  
153 aluminium foil before sealing in a polythene zip lock bag and storing in a refrigerator at ~ 4  
154 °C before transport to the University of Birmingham for chemical analysis.

### 157 3.2 Extraction and clean-up

158 High purity (HPLC grade) solvents and reagents were used for all analytical procedures and  
159 were purchased from Fisher Scientific (Loughborough, UK). Individual native BFR and NFR  
160 standards (BDE-28, BDE-47, BDE-99, BDE-100, BDE-153, BDE-154, BDE-183, BDE-209,  
161 DBDPE, *anti*-DP, *syn*-DP, PBT, PBBz, HBBz,  $\alpha$ -HBCDD,  $\beta$ -HBCDD, and  $\gamma$ -HBCDD),  
162 internal (surrogate) standards (BDE-77, BDE-128,  $^{13}\text{C}_{12}$ -BDE-209,  $^{13}\text{C}_{12}$ - $\alpha$ -HBCDD,  $^{13}\text{C}_{12}$ - $\beta$ -  
163 HBCDD, and  $^{13}\text{C}_{12}$ - $\gamma$ -HBCDD) and recovery determination (syringe) standards ( $^{13}\text{C}_{12}$ -BDE-  
164 100 and  $\text{d}_{18}$ - $\gamma$ -HBCDD) were purchased from Wellington Laboratories (Guelph, ON,  
165 Canada). Similarly, individual native PCBs (PCB-11, PCB-28, PCB-52, PCB-101, PCB-118,  
166 PCB-138, PCB-153, PCB-180), internal (surrogate) standards (PCB-14, PCB-34, PCB-62,  
167 PCB-119, PCB-131, PCB-147, and PCB-173), and recovery determination (syringe)  
168 standards (PCB-29 and 129) were purchased from Greyhound Chromatography (Birkenhead,  
169 Merseyside, UK). Silica gel and florisil of pesticide grade (60–100 mesh) were bought from  
170 Acros Organics (Geel, Belgium).

171 Prior to extraction, each PUF disk was spiked with an internal (surrogate) standard  
172 mixture comprising 20 ng of each of PCB-14, 34, 62, 119, 147, and 173 and 50 ng of each of  
173 BDE-77, BDE-128,  $^{13}\text{C}_{12}$ -BDE-209;  $^{13}\text{C}_{12}$ - $\alpha$ -HBCDD,  $^{13}\text{C}_{12}$ - $\beta$ -HBCDD, and  $^{13}\text{C}_{12}$ - $\gamma$ -  
174 HBCDD. PUF samples were subjected to pressurised liquid extraction with HPLC grade  
175 hexane using an ASE-350 instrument (Dionex, CA, USA) under the following conditions: 90  
176 °C for 5 mins, static time = 4 mins, minimum cycles = 3, rinse volume = 60%, and purge time  
177 = 90 s. (Abdallah et al., 2013). The extract was evaporated to ~ 2 mL on a TurboVap®,  
178 before purification on a 9 mm i.d. glass column packed with 1 g 44% H<sub>2</sub>SO<sub>4</sub>-impregnated  
179 silica topped with a 1 cm layer of anhydrous sodium sulfate. The column was pre-conditioned  
180 with 5 mL hexane before introduction of the crude extract and elution with 20 mL hexane.  
181 The consequent eluate was concentrated under a stream of nitrogen using a TurboVap®  
182 solvent evaporator to ~ 0.5 mL. The crude concentrate was then extracted with three 2 mL  
183 aliquots of dimethyl sulfoxide (DMSO). The combined DMSO extracts were diluted with  
184 milli-Q water (4 mL) containing 1 g of sodium chloride, and the target compounds back-  
185 extracted into hexane (using three 4 mL rinses) (Ayris et al., 1997; Folarin et al., 2018). The  
186 combined hexane back-extracts were concentrated to incipient dryness and reconstituted into  
187 100  $\mu\text{L}$  toluene containing 20 ng (PCBs-29 and -129) and 50 ng ( $^{13}\text{C}_{12}$ -BDE-100 and  $d_{18}$ - $\gamma$ -  
188 HBCDD) as recovery determination (or syringe) standards (RDSs) (Ayris et al., 1997; Drage  
189 et al., 2016).

190

### 191 3.3 Instrumental Analysis

192 Target FRs were analysed on a TRACE™ 1310 Gas Chromatograph interfaced with  
193 an ISQ™ single quadrupole mass spectrometer (Thermo Fisher Scientific, Austin, TX, USA)  
194 operated in electron ionisation and selected ion monitoring (SIM) mode. One  $\mu\text{L}$  extract was  
195 introduced via splitless injection onto a Restek Rxi-5Sil MS column (15 m x 0.25 mm x 0.25

196  $\mu\text{m}$  film thickness). Analysis of the target PCBs was carried out on a GC Agilent  
197 HP6850/5975 MS with 1  $\mu\text{L}$  of extract introduced via splitless injection onto a Restek Rxi-  
198 5Sil MS column (30 m  $\times$  0.25 mm  $\times$  0.25  $\mu\text{m}$  film thickness. Analyses of both PCBs and FRs  
199 were operated under 1.5 mL/mins flow of pure Helium as carrier gas. Details of the GC oven  
200 programmes, and the ions monitored for FR and PCB analyses are provided in Table S1 and  
201 Table S2 respectively.

202 Once PCB and FR analysis was complete, sample extracts were reconstituted in  
203 methanol and analysed for HBCDDs. This was conducted using a dual pump Shimadzu LC-  
204 20AB Prominence liquid chromatograph (Shimadzu, Kyoto, Japan)/ Sciex API 2000 triple  
205 quadrupole mass spectrometer (Applied Biosystems, Foster City, CA) equipped with an ESI  
206 ion source operated in negative ionisation mode. 10  $\mu\text{L}$  extract was separated on a Varian  
207 Pursuit XRS3 C18 reversed phase analytical column (150 mm  $\times$  4.6 mm i.d., 3  $\mu\text{m}$  particle  
208 size). Full details of the instrumental analysis conditions have been reported previously in  
209 Abdallah et al. (2016) and Drage et al. (2020).

210

### 211 3.4 Quality Assurance / Quality Control

212 All chemicals used were of high purity-grade, with all glassware baked at 425  $^{\circ}\text{C}$   
213 overnight before use. To validate the methods used, ten aliquots of standard reference  
214 material SRM-2585 (NIST organics in house dust) were analysed. Concentrations recorded  
215 were within acceptable values (Table S3). Field blanks consisting of a PUF disc transported  
216 to and from a sampling site but not exposed to air and treated as a sample were analysed. A  
217 field blank was analysed for every 5 samples making a total of 5 blanks. Where the  
218 concentration of a target compound in a field blank was between 5 and 20% of that in a  
219 sample analysed in the same batch, the corresponding sample concentration was corrected by  
220 subtraction of the concentration detected in the blank. Where concentrations in the blank

221 exceeded 20% of those detected in samples from the corresponding batch, data for the  
222 compound(s) affected are discarded for that sample. However, in this study none of the  
223 blanks contained concentrations of any target compounds that exceeded 20%. Blank  
224 correction was made for BDEs 47 (4 samples), 99 (7 samples), 100 (7 samples), 153 (6  
225 samples), 154 (4 samples); 209 (10 samples); *anti*-DP (13 samples), PBT (5 samples); PCBs  
226 11 (20 samples), 28 (11 samples), 52 (11 samples), 101 (9 samples), 118 (7 samples), 153 (4  
227 samples), 138 (3 samples); 180 (3 samples). For most target compounds, method detection  
228 limits (MDLs) were calculated from the instrument detection limit (IDL); while for those  
229 target compounds detected in blanks, MDLs were calculated based on the average + 3 x  
230 standard deviation of the concentration detected in the blank. (Tables S6a – S9a). Average  
231 internal standard (surrogate) recoveries were between 43% and 103% (Table S4 (a-c)).

232

### 233 3.5 Passive Sampling Rates

234 A passive sampling rate (PSR) of ~ 3.5 - 4 m<sup>3</sup>/day has been applied for calculating  
235 concentrations in outdoor air of BFRs and PCBs in recent studies using fully-sheltered PUF  
236 disc PAS (Herkert et al., 2018, White et al., 2021). More specifically, calibration of the PAS  
237 design deployed in this study yielded PSR values of 3.92 m<sup>3</sup>/day for BFRs except for BDE-  
238 209 and DBDPE for which the PSR was 2.23 m<sup>3</sup>/day (Drage et al., 2016). These values were  
239 thus used here for BFRs. For PCBs, we used a value of 3.5 m<sup>3</sup>/day, consistent with studies  
240 conducted in Accra, Ghana (Bohlin-Nizzetto et al., 2020) and Birmingham, UK (Jamshidi et  
241 al., 2007) (Table S5).

242

### 243 3.6 Statistical Analysis

244 Box plots were obtained using IBM SPSS Statistics v26. Pearson correlations and  
245 scatter plots were obtained using Microsoft Excel 2019. For statistical purposes, non-detect

246 values were replaced by a value equivalent to the fractional detection frequency x MDL. A p  
247 value of <0.05 was considered statistically significant.

248

## 249 **4. Results and Discussion**

### 250 **4.1 BFRs and PCBs in Lagos Air**

#### 251 **4.1.1 Concentrations of PBDEs and HBCDD in Lagos Air**

252 Table 2 and Figure 2 summarise the concentrations of  $\sum_8$ BDEs and HBCDD measured at the  
253 eight locations in this study. Concentrations of  $\sum_8$ BDEs across all samples in this study fell in  
254 the range 21 – 750 pg/m<sup>3</sup> (median = 100 pg/m<sup>3</sup>). The highest concentrations were reported at  
255 mixed residential – industrial locations and those with higher population densities (APP,  
256 OSH, AGG and IKJ), while the lowest concentrations were reported at sites with relatively  
257 low population densities (IKD and LKK) (Table S6b).

258 Possible sources of PBDEs to the atmosphere in Lagos include ventilation of indoor  
259 air from buildings containing BFR-treated consumer products, as well as volatilisation from  
260 polluted surfaces and e-waste as a result of activities like open waste treatment including  
261 burning. We explored the relationship between PBDE contamination and population density  
262 by plotting the former against the concentrations of individual PBDE congeners against  
263 population density. As we were only able to collect samples at all of our 8 locations during  
264 the 1<sup>st</sup> sampling period, we only plotted concentrations for this period. This analysis revealed  
265 significant positive correlations for BDEs- 28, 47, 99, and 183 ( $p < 0.05$ ) (Figure S1a).  
266 Interestingly, significant positive correlations ( $p < 0.05$ ) were also observed for BDEs- 100,  
267 153, 154, and 209 when the samples from the two sites with the highest population densities  
268 (AGG and OKB) were excluded as outliers (Figure S1b). Overall, our data suggest that  
269 sources of PBDEs in the Lagos atmosphere are generally diffuse and related to population  
270 density.

271 The arithmetic mean concentrations of individual PBDEs observed in this study  
272 (Table 2) are consistent with those reported recently for a single sample at one location in  
273 Lagos (Nigeria) (Saini et al., 2020) but - except for BDE-209 - exceed those reported in  
274 single samples in the same (GAPS) study for London, UK and Beijing, China (Saini et al.,  
275 2020), as well as those reported in an earlier study for Entebbe, Uganda (Arinaitwe et al.,  
276 2014). Compared to concentrations reported in 2016 for the West Midlands of the UK (Drage  
277 et al, 2016), those reported here for Lagos are in a similar range for BDEs 28, 47, 99, and  
278 100, but at the lower end of the range of those detected in the West Midlands for BDEs 153,  
279 154, 183, and 209.

280 HBCDD was only detected in a single sample collected at OKB (11 pg/m<sup>3</sup>) as well as  
281 in all 3 samples at APP at concentrations ranging between 13 and 180 pg/m<sup>3</sup> (Table S7b). In  
282 all other samples, it was not detected. Such low concentrations of HBCDD are likely  
283 attributable to the fact that its principal application is in expanded and extruded polystyrene  
284 building insulation foam (Stubbings and Harrad, 2019), for which there is little need in  
285 tropical climates like Lagos. Interestingly, the APP sampling site is in a location close to port  
286 activities. Similar observations of elevated HBCDD concentrations at a location near a port in  
287 Chile have been reported (Rauert et al., 2018). A possible explanation may be a greater  
288 abundance of polystyrene foam containers at ports for transporting foodstuffs at cold  
289 temperatures, as such containers have been shown previously to contain HBCDD (Abdallah  
290 et al., 2018; Rani et al., 2014).

291 Although HBCDD was infrequently detected in our study, Table 2 shows that the  
292 range of concentrations observed where it was detected, exceed those reported for the Great  
293 Lakes in North America (Olunkunle et al., 2018), are consistent with those for Concepción,  
294 Chile (Rauert et al., 2018) and Gauteng, South Africa (Katima et al., 2018), but are at the low  
295 end of the reported range of concentrations for Birmingham (Drage et al., 2016) and China

296 (Hu et al., 2011).

#### 297 **4.1.2 Congener/diastereomer profiles of PBDEs and HBCDD in Lagos air**

298 Similar to previous observations elsewhere (Table 2), BDE-209 is the dominant PBDE  
299 congener observed in this study with a median contribution to  $\Sigma_8$ PBDEs of 70%. By  
300 comparison, the median contributions of BDE-47 and BDE-99 are 14% and 7.9%  
301 respectively (Table S6d). The dominance of BDE-209 suggests that Deca-BDE has been the  
302 major PBDE formulation used in Nigeria. Meanwhile, as observed elsewhere (Harrad and  
303 Hunter, 2006), while BDE-99 is present at higher concentrations than BDE 47 in commercial  
304 Penta-BDE formulations, the higher vapour pressure of BDE-47 results in more facile  
305 volatilisation of this congener, leading to the congener pattern observed here. The %  
306 contribution of BDE-47 to  $\Sigma_{\text{penta}}$ BDEs (BDE-47, -99, -100) in this study (range 43% – 79%,  
307 median = 58%) (Table S6c) is consistent with the range (44% – 78%) reported at a variety of  
308 locations worldwide in the GAPs program (Saini et al., 2020). Other congeners targeted in  
309 this study – namely BDEs -28, -100, -153, -154, and -183 - are – while frequently detected  
310 (78-100%) (Table S6a) - minor contributors (1.1-1.7%) to  $\Sigma_8$ PBDEs (Table 6c).

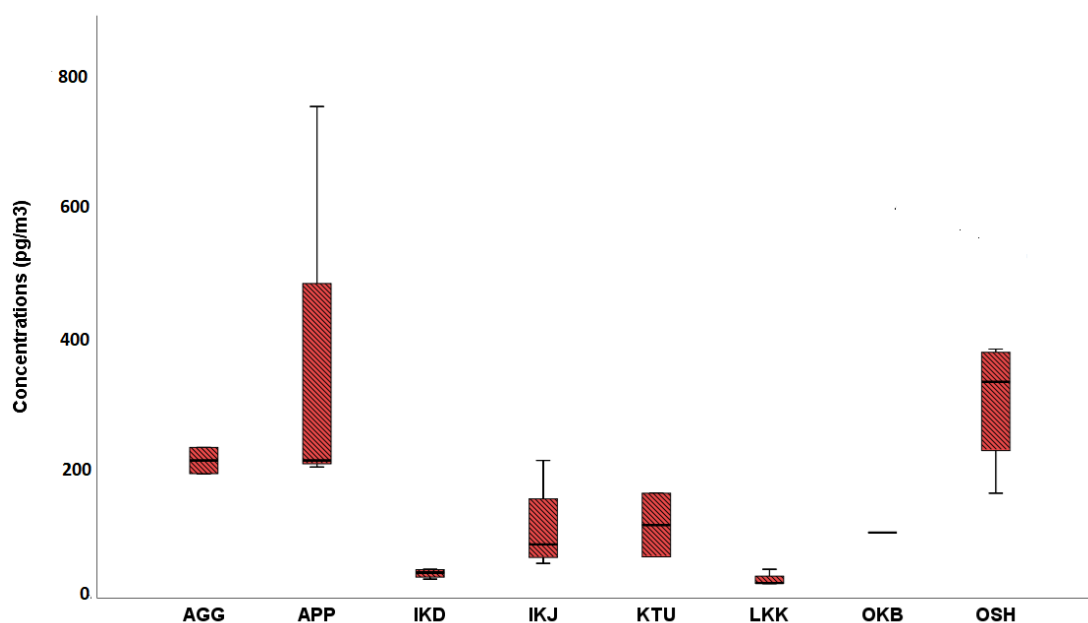
311 While HBCDD was detected in only four samples from two locations, we observed  
312 far higher median contributions for  $\alpha$ -HBCDD (81%  $\Sigma$ HBCDD) compared to  $\gamma$ -HBCDD (19  
313 %  $\Sigma$ HBCDD) with  $\beta$ -HBCDD not detected in any sample (Table S7c). This is a reversal of  
314 the diastereomer pattern present in the technical mixture of HBCDD in which  $\gamma$ -HBCDD  
315 dominates, comprising > 70%  $\Sigma$ HBCDD (de Wit et al., 2010). A similar dominance of  $\alpha$ -  
316 HBCDD has been reported in outdoor air from the UK West Midlands (Drage et al, 2016),  
317 which may be attributable to the higher volatility and lower octanol-air partition coefficient  
318 ( $K_{OA}$ ) of  $\alpha$ -HBCDD (log  $K_{OA}$  = 9.96) relative to  $\beta$ - (log  $K_{OA}$  = 10.47) and  $\gamma$ -HBCDD (log  
319  $K_{OA}$  = 10.40) (Marvin et al., 2011), favouring partitioning to air of  $\alpha$ -HBCDD, especially in  
320 hot climates.

**Table 2:** Comparison of concentrations of PBDEs and HBCDD in outdoor air in Lagos, Nigeria with those reported in other similar studies

Location	Location classification	Sampling period (Sample number)	Concentrations (pg/m <sup>3</sup> ) - Range (mean, median)										ΣHBCDDs	References		
			BDE-28	BDE-47	BDE-99	BDE-100	BDE-153	BDE-154	BDE-183	BDE-209						
29																
30	Lagos, Nigeria	Urban 2018 / 2019 (n=23)	<0.3 - 4.7 (1.9, 1.3) *96%	2.2 - 49 (20, 16) *100%	<1.7 - 53 (13, 11) *96%	<0.5 - 10 (2.7, 1.5) *78%	0.25 - 7.6 (1.9, 1.2) *100%	0.28 - 4.8 (1.7, 1.4) *100%	0.41 - 5.8 (1.9, 1.6) *100%	<16 - 620 (120, 71) *96%	11 - 180 (11, 0.7) *17%	This Study				
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32																
33	African region	Rural to urban 2001 - 2019	-	-	-	-	-	-	-	-	-	0.6 - 11	0.1 - 1.8	White et al., 2021		
34																
35																
36	New York, USA	Urban 2018 (n=1**)	3.3	36	12	3.2	0.41	0.3	-	53	<dl					
37																
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39	London, UK	Urban 2018 (n=1**)	0.73	1.1	0.46	0.09	0.02	<dl	-	85	3.7	Saini et al., 2020				
40																
41																
42	Beijing, China	Urban 2018 (n=1**)	2.3	1.7	0.71	0.1	0.35	0.15	-	48	53					
43																
44	Lagos, Nigeria	Urban 2018 (n=1**)	5.1	32	18	3.8	1.7	1.4	-	64	4.7					
45																
46	Gauteng, South Africa	Urban 2016 / 2017 (n= 8)	<dl - 120 (31, 0.01)	<dl - 212 (145, 183)	12 - 410 (206, 201)	<dl - 94 (49, 50)	<dl - 97 (35, 22)	<dl - 103 (36, 20)	5.6 - 35 (20, 20)	83 - 154 (108, 99)	31 - 70 (50, 50)	Katima et al., 2018				
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50	Entebbe, Uganda	Urban 2008 - 2010 (n=56)	<0.4 - 61 (2.9, 1.4)	<2.1 - 50 (9.8, 7.0)	<1.7 - 77 (4.4, 0.9)	<0.3 - 4 (0.6, 0.3)	<0.1 - 2.3 (0.4, 0.1)	<0.1 - 1.7 (0.3, 0.1)	<0.1 - 4.5 (0.8, 0.3)	<1.6 - 170 (8.3, 1.9)	<0.3 - 6.2 (0.6, 0.2)	Arinaitwe et al., 2014				
51																
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53	Great Lakes	Urban 2014 (n=92)	-	-	-	-	-	-	-	-	# 1.7 - 5.2	Olukunle et al., 2018				
54																
55																
56	Concepción, Chile	Urban 2014 (n=4)	-	-	-	-	-	-	-	-	110 - 180	Rauert et al., 2018				
57																
58	Birmingham, UK	Urban 2012 - 2013 (n= 8)	<0.2 - 26	< 0.3 - 31	< 0.5 - 43	< 0.5 - 30	< 1.1 - 70	<0.9 - 57	< 1.5 - 57	< 2.2 - 1500	< 0.1 - 408	Drage et al., 2016				
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**Figure 2:** Concentrations of  $\Sigma_8$ BDEs in outdoor air samples at various locations in Lagos, Nigeria

\* The boxes indicate the 25<sup>th</sup> to 75<sup>th</sup> percentiles, the whiskers indicate the max. and min. concentrations, while the black bar indicates the median concentrations.

#### 4.1.3 Concentrations of NFRs in Lagos Air

Concentrations of NFRs observed in this study are summarised in Table 3 and Figure 3. All target NFRs were detected at each location in at least one sample, at average concentrations for all 23 samples of: DBDPE (330 pg/m<sup>3</sup>), *anti*-DP (2.7 pg/m<sup>3</sup>), *syn*-DP (2.1 pg/m<sup>3</sup>), HBBz (1.4 pg/m<sup>3</sup>), PBBz (0.88 pg/m<sup>3</sup>), PBEB (0.56 pg/m<sup>3</sup>), and PBT (2.3 pg/m<sup>3</sup>). Interestingly, while  $\Sigma_7$ NFRs were observed at a higher mean concentration (330 pg/m<sup>3</sup>) than  $\Sigma_8$ BDEs (120 pg/m<sup>3</sup>), a similar trend in concentrations between locations was observed – i.e. APP > OSH > AGG > IKJ > KTU > OKB > IKD > LKK (Table S8b).

Plots of population density against NFR concentrations for each location in the first sampling period revealed significant positive correlations for PBBz and PBEB, but not for any other targeted NFRs (Fig S2). This suggests that emissions of most of our targeted NFRs

350 to the Lagos atmosphere are less diffuse and related to population density than are PBDEs.

351 This may reflect lower use of NFRs than PBDEs currently in Lagos.

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#### 353 **4.1.5 Relative abundance of NFRs in Lagos air**

354 Table S8d expresses the relative abundance of our target NFRs as their % contribution  
355 to  $\sum_7$ NFRs. While only PBT was detected in all samples, DBDPE was overwhelmingly  
356 dominant with a median contribution of 97%. Median % contributions followed the order:  
357 DBDPE > *anti*-DP > PBT > *syn*-DP > HBBz > PBBz > PBEB. This dominance of DBDPE  
358 likely reflects its widespread use as a replacement for deca-BDE (de Wit et al., 2010). With  
359 respect to the relative abundance of the two DP isomers, the *anti*-isomer of the DP is known  
360 to be more abundant ( $f_{anti-DP} = 0.59 - 0.80$ ) in the technical mixture (Sverko et al., 2011). In  
361 this study,  $f_{anti-DP}$  fell in the range 0.26 – 0.97 (mean = 0.68, median = 0.58) (Table S8c)  
362 consistent with values reported elsewhere, i.e.: 0.72 (Li et al., 2016), 0.69 (Vorkamp et al.,  
363 2015) and 0.67 (Schuster et al., 2021).

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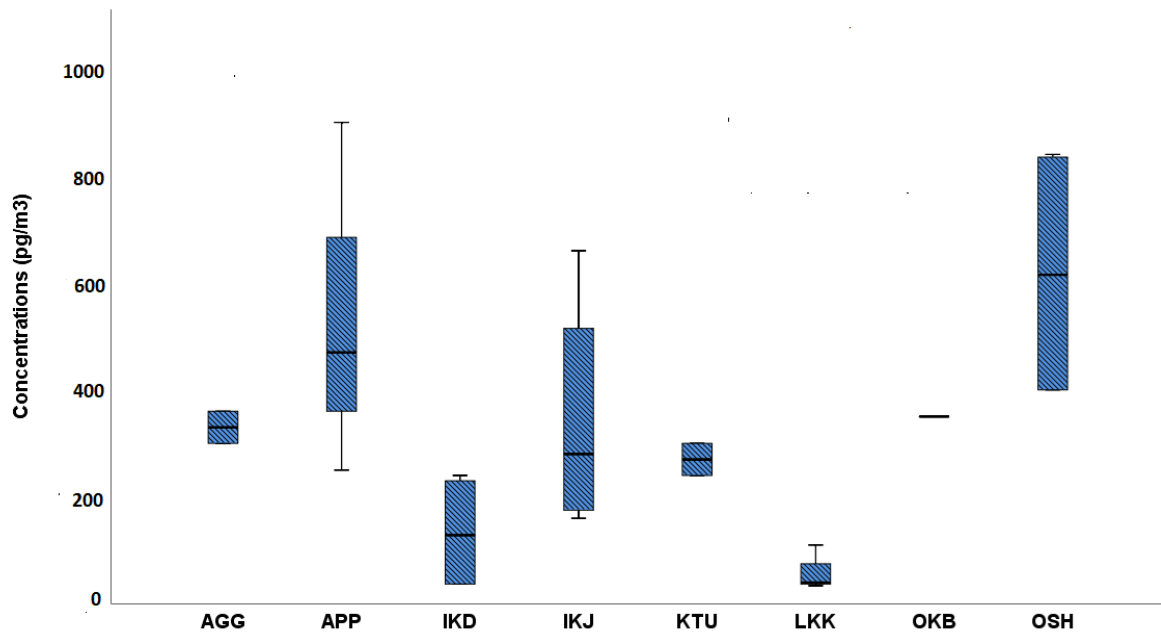
**Table 3:** Comparison of concentrations of target NFRs in outdoor air in Lagos, Nigeria, with those reported in other similar studies

Location	Location classification	Sampling period (Sample number)	Concentrations (pg/m <sup>3</sup> ) – Range (mean, median)							Reference
			PBBz	HBBz	PBT	PBEB	syn - DP	anti - DP	DBDPE	
Lagos, Nigeria	Urban	2019 / 2020 (n=23)	<0.28 – 2.4 (0.90, 0.56) *83%	<0.30 – 6.5 (1.4, 0.60) *87%	0.31 – 10 (2.3, 1.5) *100%	<0.28 – 3.2 (0.56, 0.15) *57%	<0.18 – 8.5 (2.1, 0.83) *57%	<1.8 – 8.1 (2.7, 2.2) *57%	<37 – 890 (330, 280) *87%	This Study
New York, USA	Urban	2018 (n=1**)	6.5	4.5	8.6	6.8	3.6	10.4	-	-
London, UK	Urban	2018 (n=1**)	2.2	1.0	3.4	3.6	0.81	0.44	-	Saini et al., 2020
Beijing, China	Urban	2018 (n=1**)	7.9	0.95	8.7	0.32	0.50	0.88	-	-
Lagos, Nigeria	Urban	2018 (n=1**)	4.6	9.0	7.5	0.86	16.7	11	-	-
Entebbe, Uganda	Urban	2008 – 2010 (n=56)	<0.05 – 2.3 (0.1, 0.11)	<0.09 – 1.5 (0.06, 0.05)	<0.06 – 22 (0.08, 0.03)	<0.06 – 1.8 (0.05, 0.03)	<0.05 – 4.4 (0.04, 0.03)	<0.05 – 1.4 (0.05, 0.03)	<5.9 – 220 (3.4, 3)	Arinaitwe et. al., 2014
Guangzhou, China	Urban	2014 (n= not provided)	-	19±8.8	-	6.2±10.3	4.9±4.5	5.0±4.1	316±285	Li et al., 2017
Harbin, China	Urban	2008 – 2013 (n=227)	-	-	-	-	<dl–52 (1.9, 0.46)	<dl–190 (5.8, 0.99)	-	Li et al., 2016
Chicago, USA	Urban	2005 – 2013 (n= not provided)	-	** 0.54 ± 0.08	-	0.44 ± 0.04	0.96 ± 0.15	1.8 ± 0.4	1.5 ± 0.5	Liu et al., 2016
Station Nord, Greenland	Arctic	2012 (n=13)	-	-	-	-	<1 - 9.0 (2.3)	<1 – 33 (5.2)	-	Vorkamp et al., 2015
Harbin, China	Urban	2007/2008 (n = 57)	<dl – 1.9 (0.46, 0.36)	<dl – 170 (4.8, 1.3)	0.15 - 37 (6.1, 4.0)	0.02 – 1.4 (0.34, 0.23)	-	-	<dl – 370 (1.1, 0.69)	Qi et al., 2014

\* Detection frequency    \*\* Geometric mean ± standard deviation    dl – detection limit    \*\* denotes number of locations

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**Figure 3:** Concentrations of  $\Sigma_7$ NFRs in outdoor air samples at various locations in Lagos, Nigeria

\* The boxes indicate the 25<sup>th</sup> to 75<sup>th</sup> percentiles, the whiskers indicate the max. and min. concentrations, while the black bar indicates the median concentrations.

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#### 4.1.6 Comparison of concentrations of NFRs in Lagos air with reports from other locations worldwide

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The dominance of DBDPE amongst our target NFRs in this study is consistent with most other studies reporting concentrations of NFRs in outdoor air around the world (Table 3). The mean concentration of DBDPE observed in this study exceeds those reported for Chicago, USA (Liu et al., 2016), Entebbe, Uganda (Arinaitwe et al., 2014), and Harbin, China but is lower than that reported for Guangzhou (China) (Li et al., 2017). In general, concentrations of target NFRs in this study exceed those reported for Entebbe (Uganda) (Arinaitwe et al., 2014), are lower than those detected in Guangzhou (China) (Li et al., 2017), while are broadly of similar magnitude to those reported in a single sample in Lagos (Saini et al., 2020) and in other cities in Asia, Europe, and North America.

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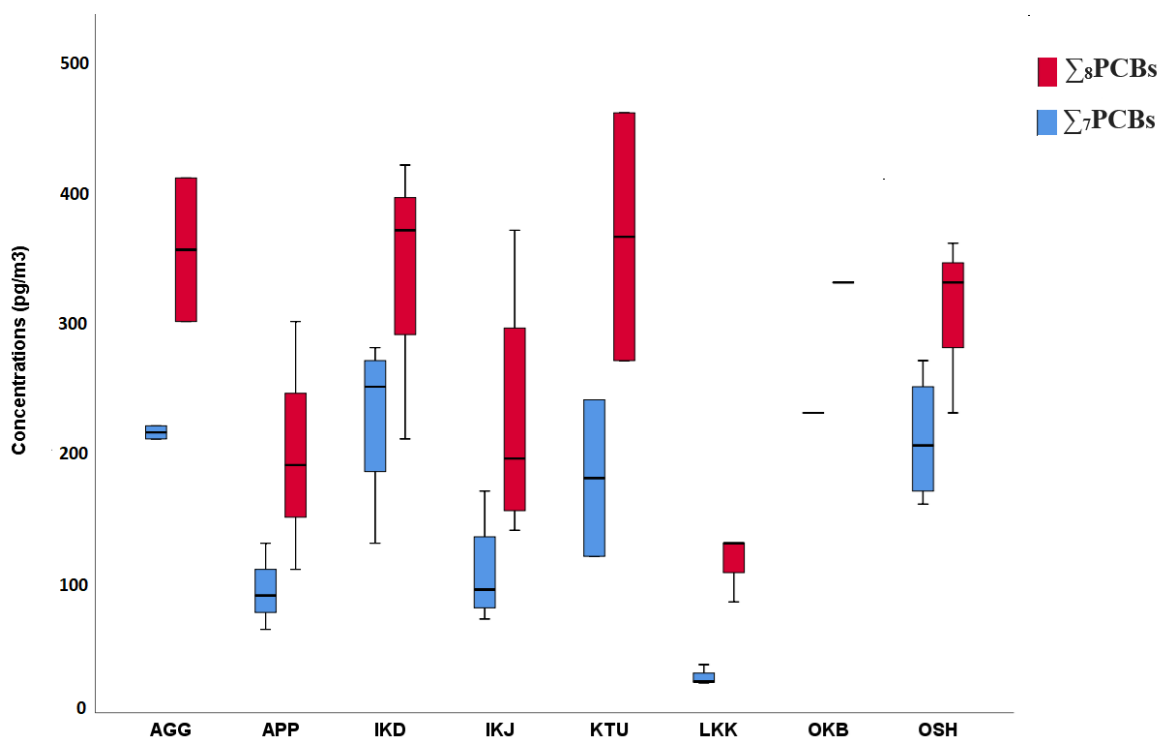
#### 395 4.1.7 Concentrations of PCBs in Lagos air

396 Tables 4a and 4b and Figure 4 summarise the concentrations of PCBs detected in this  
397 study. Those of  $\Sigma_7$ PCBs (comprising only congeners from Aroclor sources) range between 23  
398 – 280  $\text{pg/m}^3$  (median = 160  $\text{pg/m}^3$ ) while those of PCB 11 (considered to arise from non-  
399 Arochlor sources - Guo et al., 2014) range between 49 – 220  $\text{pg/m}^3$  (median = 100  $\text{pg/m}^3$ ).  
400 Similar to the situation for BFRs, higher concentrations of both PCB 11 and  $\Sigma_7$ PCBs were  
401 observed in more densely populated and urbanised settings, with the important exception of  
402 IKD where the sample location was proximate to transformer locations and a textile factory.  
403 As the production of dyes has been highlighted previously as a source of PCB 11 (Vorkamp  
404 et al., 2016); the likely use of dyes at the local textile factory is a plausible source of the  
405 elevated concentrations of this non-Aroclor congener at IKD. The lowest concentrations of  
406 both PCB 11 and  $\Sigma_7$ PCBs were observed at the least densely populated location (LKK)  
407 (Table S9b).

408 Plotting the population density of our sampling locations against the concentrations of  
409 individual PCB congeners detected at each location for the first sampling period indicated no  
410 significant correlation for any of our target PCBs (Figure S3). This suggests that unlike  
411 PBDEs and some NFRs, concentrations of PCBs in Lagos are less influenced by diffuse  
412 sources related to population density and more impacted by point source activities such as  
413 transformer leaks from electricity sub-stations and for PCB 11, activities such as the  
414 production of dyes and activities using such dyes like textile factories. Similar weak  
415 correlations between atmospheric concentrations of PCBs and population density have been  
416 previously reported (Du et al., 2009).

#### 417 4.1.8 Congener profiles of PCBs in Lagos air

419 The dominant congener detected out of our 8 target PCBs was PCB 11, making a  
 420 median contribution to  $\Sigma_8$ PCBs of 47% (Table S9c), followed by PCB 28 (13%) and 138  
 421 (13%), with the least abundant PCB 118 (2.8%). That PCB 28 is the most abundant Aroclor  
 422 congener is consistent with other reports on Africa (Bodgal et al., 2013; White et al., 2021)  
 423 and is likely due to its higher vapour pressure which facilitates volatilisation compared to  
 424 higher molecular weight congeners, especially in hotter climates.



426 **Figure 4:** Concentrations of PCBs in outdoor air samples at various locations in  
 427 Lagos, Nigeria –  $\Sigma_8$ PCBs =  $\Sigma_7$ PCBs + PCB-11  
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429 \* The boxes indicate the 25<sup>th</sup> to 75<sup>th</sup> percentiles, the whiskers indicate the max. and min.  
 430 concentrations, while the black bar indicates the median concentrations.  
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**Table 4a:** Comparison of concentrations of Arochlor-derived PCBs in outdoor air in Lagos, Nigeria with those reported in other similar studies

Location	Location classification	Sampling period (Sample number)	Concentrations (pg/m <sup>3</sup> ) – Range (mean, median)								Σ-PCBs	References
			PCB-28	PCB-52	PCB-101	PCB-118	PCB-138	PCB-153	PCB-180	PCB-180		
Lagos, Nigeria	Urban	2018 / 2019 (n=23)	7.6 – 95 (42, 34) *100%	2.9 – 52 (21, 17) *100%	2.4 – 44 (17, 15) *100%	0.5 – 26 (10, 9) *97%	3.8 – 80 (38, 35) *100%	1.8 – 43 (21, 19) *100%	0.60 – 12 (6.7, 6.9) *100%	23 – 280 (160, 160)	This study	
Entebbe, Uganda	Urban	2008 – 2010 (n=53)	-	-	-	-	-	-	-	4.4 – 48	Arinaitwe et al., 2018	
Gambia	Not specified	2008 (n=4)	-	0.8 – 18	-	-	0.9 – 6.2	-	0.4 – 2.3	30 – 130		
Ghana	Not specified	2008 (n=5)	-	0.8 – 2.1	-	-	<dl- 2.3	-	<dl- 1.1	3.1 – 12	Gioia et al., 2011	
Ivory Coast	Not specified	2008 (n=5)	-	7.0 – 20	-	-	4.9 – 20	-	1.7 – 6.5	6 – 110		
Bamako/Kati, Mali	Urban	2001 / 2008 (n=49)	-	-	0.1 – 18	-	0.01 – 13	0.01 – 13	-	-	Garrison et al., 2014	
Sheda, Nigeria	Semi-urban	2008 (n=6)	-	-	-	-	-	-	-	**14 – 130 (15, 38)	Klánová et al., 2009	
Dakar, Senegal	Urban	2008 (n=6)	-	-	-	-	-	-	-	550 – 1300 (950, 1100)		
Africa region	Not specified	-	-	-	-	-	-	-	-	8 – 2100 (180, 84)	Bogdal et al., 2013	
Canadian Great Lakes Basin	Remote to suburban	2012 (n ≅ 1300)	1.4 – 9.3 *** (4.1)	1.0 – 5.8 (2.8)	0.43 – 3.6 (1.5)	0.11 – 0.88 (0.40)	0.07 – 1.3 (0.53)	0.09 – 1.1 (0.42)	0.03 – 0.29 (0.12)	18 – 110 (55)	Shunthirasingham et al., 2016	
Birmingham, UK	Urban	2003/2004 (n=11)	-	0.4 – 33 *** (7.0)	<dl- 44 (5.5)	-	0.2 – 28 (2.9)	0.2 – 33 (3.4)	0.1 – 15 (2.0)	-	Jamshidi et al., 2007	
India	Rural to urban	2006 (n=18)	31 – 175	0.1 – 137 (27)	-	-	0.3 – 9.0 (2)	0.5 – 3.0 (2)	-	-	Zhang et al., 2008	



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**\* Detection frequency**

**\*\* using 98 m<sup>3</sup> as air volume**

**\*\*\* Arithmetic mean**

**dl – detection limit**

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**Table 4b:** Comparison of Atmospheric Concentrations of PCB 11 in Lagos, Nigeria with those reported elsewhere

Location	Location classification	Sampling period (Sample number)	Mean	Median	Range	References
Lagos, Nigeria	Urban	2018 / 2019 (n= 23)	120	100	49 – 220	This study
China	Remote to urban	2016 – 2017 (n=62)	35	-	<dl – 249	Zhao et al., 2020
Northern Vietnam	Urban	2013 – 2015 (n=not specified)	-	140	68 - 300	Anh et al., 2020
Antarctica	Remote	2011 - 2014 (n = not provided)	21	-	4.8–62	Wang et al., 2017
Chicago, USA	Urban	2006/2007 (n = 184)	24	-	<dl - 140	Hu et al., 2008
Philadelphia/Camden, USA	Urban	2005 (n = 32)	-	-	* 2 – 22	Du et al., 2009
Cleveland, USA	Urban	2004 – 2007 (n = 106)	-	-	<dl - 307	Basu et al., 2009

\* using 315 m<sup>3</sup> as air volume dl – detection limit

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#### 4.1.9 Comparison of concentrations of PCBs in Lagos with those reported elsewhere

Concentrations of  $\sum_7$ PCBs observed in this study are in the middle of the range reported in various previous studies conducted in Africa and other global studies (Table 4a). Similarly, concentrations of PCB 11 reported in this study are – while to our knowledge, the first reported for Africa - within the range reported from various studies around the world (Table 4b).

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#### 4.1.10 Seasonal variation in concentrations of BFRs and PCBs in Lagos air

The Lagos climate is a typical tropical climate, divided mainly into the wet (rainy) season between April and September and dry season between October and May. There is little difference in temperature year-round, with the Nigerian Meteorological Agency (NIMET) reporting temperatures to average 27.3 °C in the wet period and 28.6 °C in the dry

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456 period. It is therefore not surprising our study does not detect a seasonal peak in  
 457 concentrations of PCBs in outdoor air similar to that reported in summer for North America  
 458 (Carlson and Hites, 2005) and Northern Europe (Currado and Harrad, 2000) that have been  
 459 attributed to higher summer temperatures facilitating volatilisation from surfaces.

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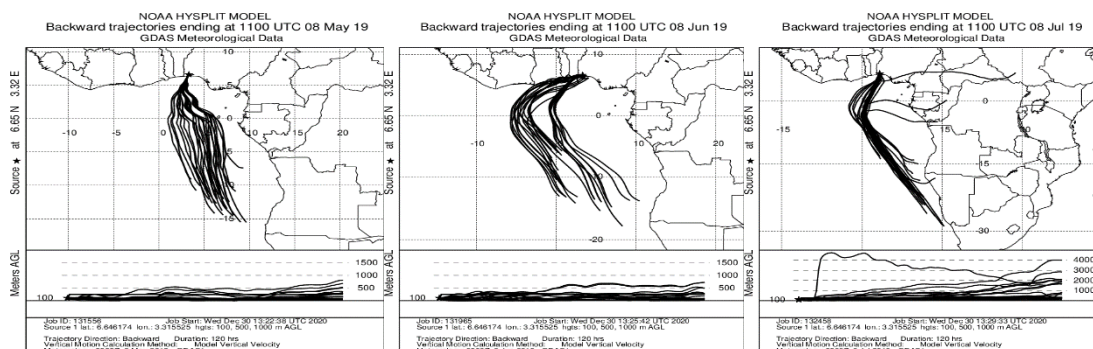
## 461 4.2 Potential Sources of BFRs and PCBs to Lagos Air

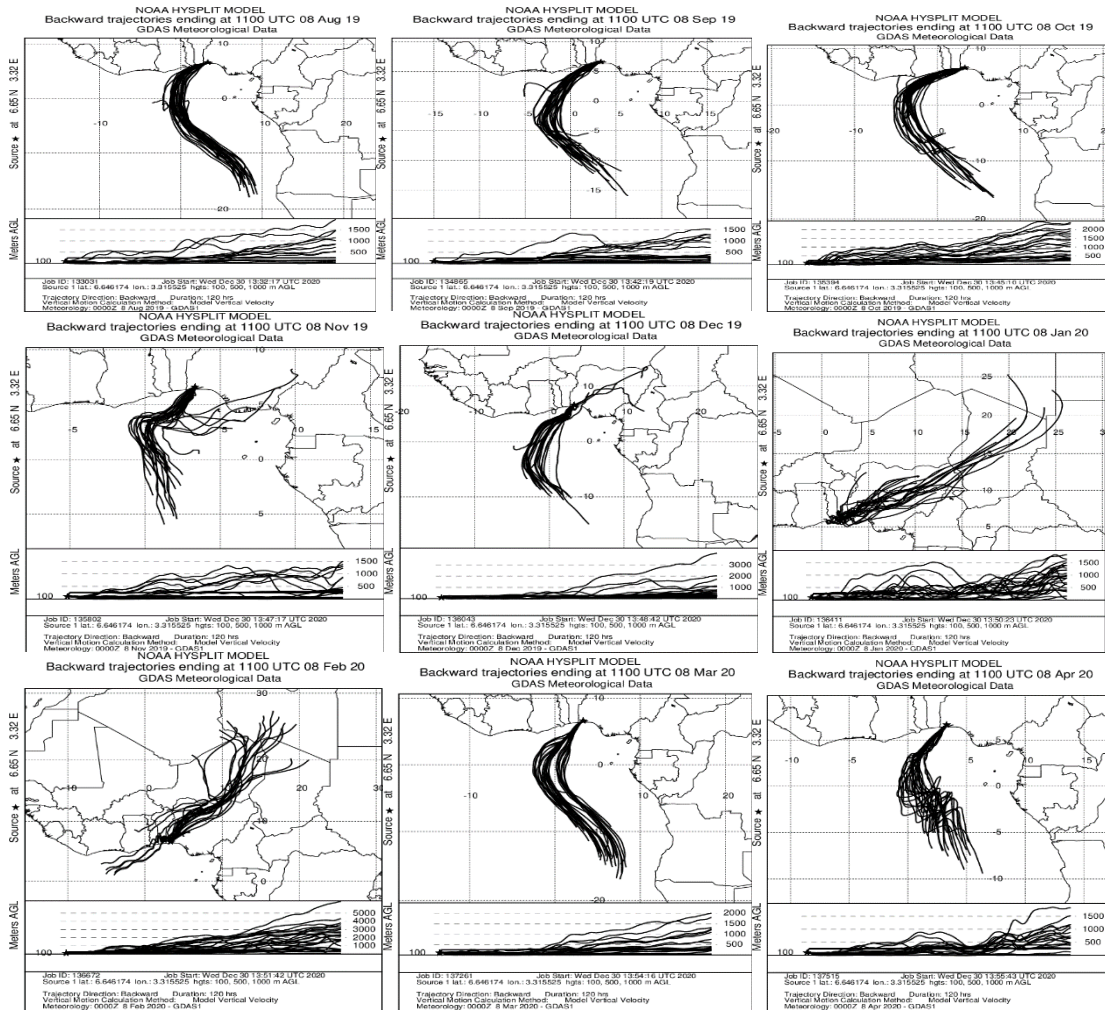
462 Use of National Oceanic and Atmospheric Administration (NOAA) Hypslit back  
 463 trajectory models (Figure 5) indicated that the air masses arriving at our sampling locations  
 464 are predominantly sourced from the Atlantic Ocean within the sampling period. Coupled with  
 465 the consistently low concentrations reported at the least urbanised and densely populated  
 466 LKK location, this indicates that the elevated concentrations observed at the urban locations  
 467 in our study, do not arise from long-range atmospheric transport but instead originate within  
 468 the Lagos environment.

469 The presence of PBDEs and some NFRs in the Lagos atmosphere are therefore likely  
 470 related to diffuse sources linked with urbanisation such as ventilation of contaminated indoor  
 471 air, whereas the sources of PCBs appear more related to point activities such as historical or  
 472 current leakages from electric power transformers, and industrial activities related to pigment  
 473 production and use such as textile manufacture.

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**Figure 5:** HYPPLIT back dispersion models (<https://www.ready.noaa.gov/hypub-bin/trajsrc.pl>) run at 120 hours at 100m, 500m and 1000m above ground arrival height.

## 5.0 Conclusions

This study examines concentrations, profiles, and potential sources of BFRs and PCBs in the atmosphere of Lagos, Nigeria. Our data reveal concentrations of our target pollutants to be comparable or higher than those reported elsewhere in the world and provide a valuable baseline against which the efficacy of actions to reduce environmental contamination with these chemicals can be evaluated. The concentrations of PCBs reported here exceed substantially those reported previously for semi-urban locations in Sheda and Abuja, Nigeria. Importantly, the non-Arochlor PCB congener 11 was reported in outdoor air for the first time in Africa.

493 Back trajectory analysis reveals that the observed target contaminants likely originate  
1  
2 494 within Lagos itself, rather than via long-range atmospheric transport. Concentrations of  
3  
4 495 PBDEs and some NFRs correlated with population density while for PCBs point sources  
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6  
7 496 appear to play a more important role. Combined, this suggests urbanisation and  
8  
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10 497 industrialisation exert important influences on BFR and PCB contamination in Lagos. The  
11  
12 498 absence of any substantial seasonal variation in temperature in Lagos likely explains why no  
13  
14  
15 499 seasonal variation is observed in concentrations of any of our target contaminants.  
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17 500

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26 504

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## **Credit author statement**

**Olumide Emmanuel Akinrinade:** Investigation; Writing – Original manuscript preparation; Funding acquisition

**William A. Stubbings:** Investigation; Writing - Review & Editing

**Mohamed Abou-Elwafa Abdallah:** Writing - Review & Editing

**Olusegun Ayejuyo:** Writing - Review & Editing

**Rose Alani:** Supervision; Writing - Review & Editing

**Stuart Harrad:** Supervision; Writing - Review & Editing; Funding acquisition