

Can polychlorinated biphenyl (PCB) signatures and enantiomer fractions be used for source identification and to age date occupational exposure?

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Abstract

Detailed polychlorinated biphenyl (PCB) signatures and chiral Enantiomer Fractions (EFs) of CB-95, CB-136 and CB-149 were measured for 30 workers at a transformer dismantling plant. This was undertaken to identify sources of exposure and investigate changes to the PCB signature and EFs over different exposure periods. Approximately 1.5 g of serum was extracted and PCB signatures were created through analysis by comprehensive two-dimensional gas chromatography with time-of-flight mass spectrometry (GCxGC-TOFMS) and EFs calculated by analysis by gas chromatography with high resolution mass spectrometry (GC-HRMS). A total of 84 PCBs were identified in the serum samples with concentrations of the 7 indicator PCB ranging from 11 - 350 ng g⁻¹ of serum (1.2 - 39 µg g⁻¹ lipid). The PCB signatures were interpreted using principal component analysis (PCA) which was able to distinguish workers with background or recent minimal exposure, from those with prolonged occupational exposure. Occupationally exposed individuals had a similar PCB profile to Aroclor A1260. However, individuals with prolonged exposure had depleted proportions of several PCB congeners that are susceptible to metabolism (CB-95, CB-101 and CB-151), and elevated proportions of PCBs that are resistant to metabolism (CB-74, CB-153, CB-138 and CB-180). The results also identified a third group of workers with elevated proportions of CB-28, CB-60, CB-66, CB-74, CB-105 and CB-118 who appeared to have been exposed to an additional source of PCBs. The results show near complete removal of the CB-95 E2 enantiomer in some participants, indicating that bioselective metabolism or preferential excretion of one enantiomer occurs in humans. By considering PCB concentrations along with detailed congener specific signatures it was possible to identify different sources of contamination and gain an insight into both the magnitude and duration of exposure.

Keywords

Polychlorinated biphenyls (PCBs); Human exposure, GCxGC-TOFMS, Chemical fingerprinting, PCB atropisomers, Chiral, Enantiomer Fractions

Highlights

- Eighty four different PCBs detected in human serum samples
- PCB signatures used to distinguish recent and prolonged exposure
- PCB signatures used to identify different sources of exposure
- Near complete removal of the CB-95 E2 enantiomer recorded in some humans

1 **1 Introduction**

2 Polychlorinated biphenyls (PCBs) are a group of 209 chlorinated organic compounds that
3 were widely used throughout the 20th century. While PCBs have been largely phased out of
4 commercial/industrial use, they remain an important legacy contaminant (O'Sullivan and
5 Sandau, 2013). They are highly persistent and can still be found in closed systems in some
6 countries, e.g. as dielectric fluids in electrical equipment and transformers. Many of these
7 transformers containing PCBs are in the process of being replaced, and this process therefore
8 presents the potential of PCB exposure for humans working in dismantling plants. In these
9 instances it is important not only to determine the extent of the exposure, along with the
10 potential risks to human health, but also to establish the source of the contamination and age
11 date the exposure.

12 As PCBs were produced as commercial mixtures, such as Aroclors, each blend has a specific
13 congener profile (signature) based on the relative proportions of each PCB in the total
14 mixture. This signature can be used to easily distinguish commercial mixtures, however
15 environmental investigations involving humans are more complex as there are often multiple
16 potential sources of PCB exposure. The signature can be altered by changes such as
17 volatilization, dissolution and biodegradation (Jaspers et al., 2013; Johnson et al., 2006). The
18 signature in humans can also vary depending on different exposure pathways e.g. oral,
19 inhalation or dermal, and can be altered through post uptake processes such as
20 biotransformation and elimination (Jaspers et al., 2013; Megson et al., 2013a). If a dominant
21 source of exposure can be identified then alterations to this signature from post uptake
22 processes such as biotransformation and elimination may provide useful information to
23 distinguish between recent and prolonged exposure. The sera of an individual who has been
24 historically exposed may contain higher proportions of the PCBs that are more resistant to

25 biotransformation and elimination. Due to all of the subtle changes that can occur to the PCB
26 profile, when attempting to identify the source of exposure it is imperative that signatures are
27 created using detailed congener specific datasets. Analysis using comprehensive two
28 dimensional chromatography has proven to be an excellent technique for this purpose as it is
29 able to separate over 190 individual PCB congeners (Focant et al., 2004; Korytar et al., 2006;
30 Harju et al., 2003; Zapadlo et al., 2011; Megson et al., 2013b).

31 Of the 209 PCBs there are 19 which are predicted to exist as stable atropisomers (Oki, 1983).
32 They have a high degree of *ortho* chlorine substitution which inhibits rotation, and an
33 unsymmetrical *meta* and *para* substitution on each biphenyl, resulting in two optical isomers.
34 In commercial mixtures both enantiomers are produced in equal proportions; however in
35 animals, metabolic processes such as enzyme mediated oxidation have been proven to
36 preferentially target one stereoisomer, resulting in atropisomeric enrichment (Harrad et al.
37 2006; Wong et al., 2002; Wu et al., 2014). Therefore, the sera of an individual who has
38 recently been exposed to a commercial PCB mixture may be expected to contain near equal
39 proportions of each stereoisomer whereas the sera of a historically exposed individual may
40 show a greater degree of fractionation. However, this signal is likely to be complicated by
41 interferences such as other background sources of PCBs.

42 The goal of this study was to determine if PCB signatures and enantiomer fractions could be
43 used to identify the source of contamination and distinguish between recent and prolonged
44 exposure periods for 30 workers at a transformer dismantling plant.

45 **2 Experimental**

46 **2.1 Sample collection**

47 Samples of whole blood were collected from 30 people working at a transformer dismantling
48 plant in Europe. Samples were obtained from workers performing a range of different roles at
49 the plant, including workers on the dismantling floor who were likely to have had direct
50 contact with PCBs and those who were not expected to have had any direct contact with
51 PCBs such as administrative staff and a security guard. Samples were obtained from
52 employees who had been working at the plant from 3 - 21 years. However, information on
53 occupation and length of time at the plant was not available for three participants (id no. S028,
54 S029 and S030). A 10 mL sample of whole blood was obtained from each worker, the blood
55 was collected in vacutainers, then centrifuged and the serum collected and stored at -20 °C.

56 Total lipid concentrations were determined by enzymatic analysis which was performed by a
57 sub-contractor clinical laboratory on a dedicated 2 mL serum sub-sample. Four types of lipids
58 were targeted and measured; triglycerides, total cholesterol, non-esterified (free) cholesterol,
59 and phospholipids. Sample sizes were as follows: triglycerides (2 µL), total cholesterol (2
60 µL), non-esterified (free) cholesterol (50 µL), and phospholipids (20 µL). Total lipid
61 concentrations were estimated using the summation method of Akins et al. (1989). The total
62 lipid content was expressed in g L⁻¹. For the inter-conversion of volumetric and gravimetric
63 data, a value of 1.026 g mL⁻¹ for serum specific gravity was used.

64 **2.2 Sample preparation**

65 All reagents required for extraction and clean-up were sourced specifically for dioxin, furan
66 and PCB analysis or of the closest grade available of similar quality. Approximately 1.5 g of
67 serum was accurately weighed (to 4 decimal places) and transferred to a vial and 5 µL of
68 ¹³C₁₂ labelled CB-60, CB-127 and CB-159 at a concentration of 100 pg µL⁻¹ (CIL-EC-5370

69 EN-1948-4 PCB sampling standard) was added to determine recovery. A volume of formic
70 acid equal to the mass of the sample was added to the serum followed by the same volume of
71 high purity water. During each addition the solution was vortexed and allowed to degas for
72 several minutes. The whole extract was then applied to a C₁₈ SPE cartridge (1 g / 6 mL) and
73 the PCBs were eluted with hexane. The eluent was treated using EPA method 3665A
74 sulphuric acid / permanganate clean-up followed by EPA method 3620 Florisil clean-up.
75 Extracts were reduced to approximately 50 µL by nitrogen evaporation and 100 µL of ¹³C₁₂
76 labelled PCBs (CIL-EC-5367 CDC PCB Spiking Standard), each at a concentration of 7.5 pg
77 µL⁻¹, were added. Extracts were left overnight to evaporate to incipient dryness (the spiking
78 standard contained a dodecane keeper). Samples were reconstituted with 10 µL of hexane
79 prior to analysis.

80 **2.3 Congener specific analysis by GCxGC-TOFMS**

81 Analysis was conducted based on the method described by Megson et al. (2013b) which is
82 summarised below. Samples were analysed using a time-of-flight mass spectrometer (LECO,
83 St. Joseph, MI Pegasus 4D) coupled to a two dimensional gas chromatograph (Agilent
84 Technologies 7890A) equipped with a thermal modulator (LECO, St. Joseph, MI). The gas
85 chromatograph was fitted with a Rtx-PCB (60 m x 0.18 mm x 0.18 µm) ¹D column and a
86 Rxi-17 Sil MS (1.5 m x 0.18 mm x 0.18 µm) ²D column. One µL of sample was injected in
87 splitless mode.

88 Procedural blanks were prepared for each batch of 8 samples. Contamination with CB-11 was
89 identified in the blanks and so this congener was excluded from the results. All samples were
90 spiked with two sets of ¹³C₁₂ labelled internal standards which were used to quantify PCB
91 concentrations and calculate recovery. Quantification was undertaken through isotope
92 dilution; calibration data was produced for 41 of the most commonly encountered congeners

93 (CIL-EC-4133 DSJ PCB Mixture). Quantification of other congeners present in the samples
94 was undertaken using the calibration data from the closest eluting calibrated congener with
95 the same level of chlorination. Recovery for all samples was within the accepted range
96 specified by EPA method 1668C (10 % to 145 %), the mean recovery was 55 % (± 16 %; 1σ).
97 Concentrations were recovery corrected and lipid normalised and reported as ng g^{-1} lipid
98 weight. The instrument limit of detection (LOD) was estimated empirically using the
99 calibration standard mixtures, LODs for CB-18 and CB-206 were calculated at a
100 concentration of at 1 and $50 \text{ pg } \mu\text{L}^{-1}$ respectively. Accuracy and precision were measured for
101 the sum of the European Union 7 indicator congeners (EC7; CB-28, CB-52, CB-101, CB-118,
102 CB-138, CB-153, CB-180) by analysing a $10 \text{ ng } \mu\text{L}^{-1}$ Aroclor 1248 standard three times. The
103 accuracy of the sum of the EC7 congeners for the three samples was 105% ($\pm 0.9\%$; 1σ).

104 **2.4 Chiral analysis by GC-HRMS**

105 The Enantiomeric Fractions (EFs) of CBs 95, 136 and 149 were analysed according to the
106 method of Robson and Harrad (2004), using an Agilent 7890 Gas Chromatograph coupled to
107 a Micromass Auto Spec Premier High Resolution MS tuned to greater than 10000 mass
108 resolution. The two most abundant isotopes of each enantiomer were recorded in Single Ion
109 Recording Mode (SIR). This was 325.88040 and 327.87750 for CB 95 and 359.84150 and
110 361.83850 for CBs 136 and 149. These PCBs were chosen because they are; (a) able to be
111 baseline separated on the Chirasil Dex column, (b) free from any co-eluting congeners that
112 may bias the results and (c) normally present in the environment in high enough
113 concentrations to be accurately measured.

114 The chromatographic performance of the method was assessed prior to each run of 8 samples
115 by analysing a 1:1:1 mixture of Aroclors 1248, 1252 and 1260.

116 Enantiomeric Fractions were calculated as per Harner et al. (2000). Whereby

117 $EF=E1/(E1+E2)$

118 Where E1 equals the first eluting or the (+) enantiomer and E2 the second eluting enantiomer.

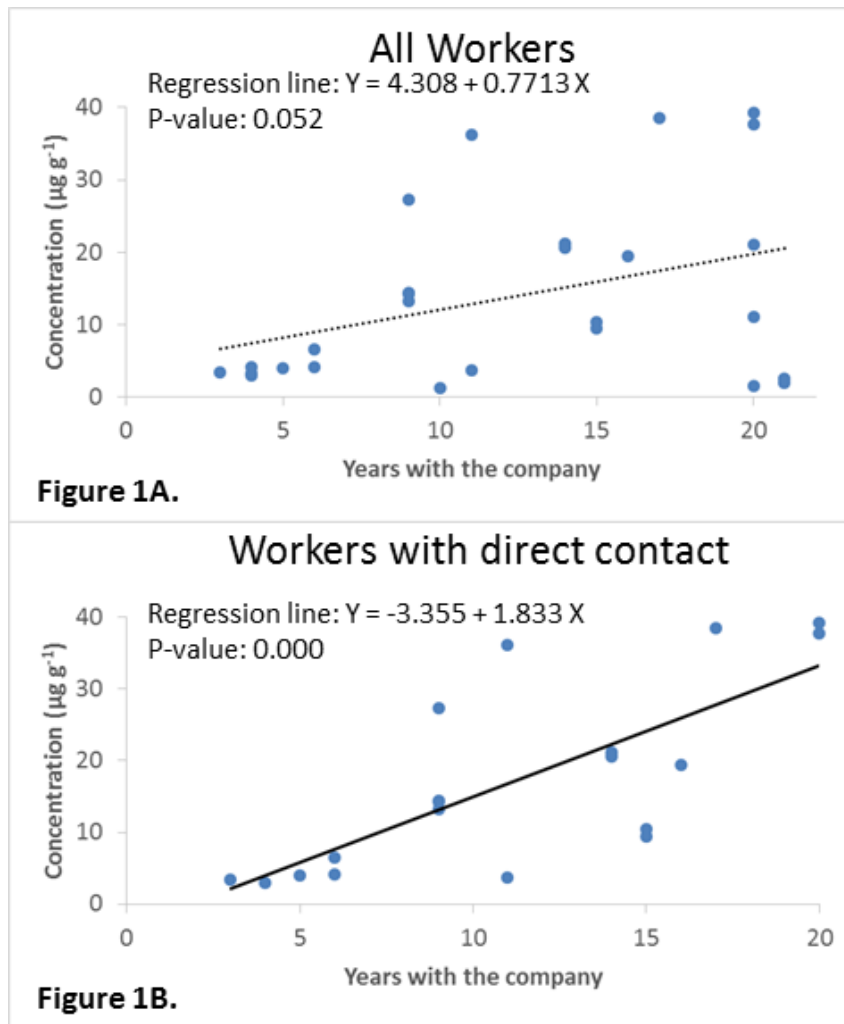
119 Samples were only accepted for quantitation if the enantiomeric fractions of the three
120 atropisomers studied were 0.50 (± 0.01) in the Aroclor mixture, the least abundant enantiomer
121 of the pair had a signal to noise (S:N) ratio greater than 10:1, the isotope ratios were within
122 20% of their theoretical values and the analytical recovery of the sample was greater than
123 30%. The instrument LOD was estimated by analysing a standard mixture of CB-95 and CB-
124 149, LODs were calculated at a concentration of 2.5 pg μL^{-1} per enantiomer. Procedural
125 blanks were prepared for each batch of 8 samples; no chiral PCBs were detected in the blanks
126 above the limits of detection.

127 **3 Results and discussion**

128 **3.1 Concentrations of PCBs in workers at a transformer dismantling plant**

129 A total of 84 different PCB congeners were identified in the serum of the 30 workers at the
130 transformer dismantling plant. Concentrations of the 7 indicator PCBs ranged from 11 - 350
131 ng g^{-1} of serum (1.2 - 39 $\mu\text{g g}^{-1}$ lipid). Background concentrations of these congeners in
132 humans are in the range of 0.1 - 10 ng g^{-1} serum (0.01 - 1 $\mu\text{g g}^{-1}$ lipid) (Longnecker, 2001).
133 This shows that some workers had close to background exposure whereas others had elevated
134 PCB concentrations indicating that they may have been exposed to PCBs through their
135 occupation. The PCB concentrations in the different workers are summarised in Table 1. The
136 results show that the workers with job roles involving direct contact with PCBs generally had
137 higher PCB concentrations than those with no direct contact. The mean concentration of the
138 EC7 PCBs in workers with direct contact was 17.2 $\mu\text{g g}^{-1}$ lipid (± 13.0 ; 1σ) which was
139 significantly greater (P-value 0.004) than workers with no direct contact (5.83 $\mu\text{g g}^{-1}$ lipid

140 (± 6.90 ; 1σ). However, this was not true for all workers, as a concentration of $21.0 \mu\text{g g}^{-1}$
141 lipid was recorded in a chief of operations (chief) who was believed to have had no direct
142 contact with PCBs. PCB concentrations generally increased with the number of years the
143 employee had worked with the company (Figure 1A), although this increase was not
144 statistically significant (P-value of 0.052). However, when PCB concentrations of the 18
145 workers who were working at the dismantling plant with direct contact dismantling
146 transformers were considered, this increase was statistically significant (P-value of 0.000)
147 (Figure 1B). Although the PCB concentration is well correlated with the number of years an
148 individual has worked at the company it should not be used in isolation as proof that
149 occupational exposure has occurred. Higher PCB concentrations have been reported in older
150 individuals as a result of accumulation of the more persistent congeners and exposure to
151 higher historical background concentrations (Megson et al., 2013a; Quinn and Wania 2012).
152 There are also several physiological characteristics, such as body fat, serum albumin and age,
153 that can influence the uptake and retention of PCBs along with social preferences such as diet
154 and smoking (Axelrad et al., 2009; Brown and Lawton, 2001; Jain and Wang, 2011;
155 Weintraub and Birnbaum, 2008). This highlights the importance of looking at the specific
156 PCB signature of each individual to determine their exposure source rather than relying
157 solely on a total PCB concentration.



158

159 **Figure 1. Relationship between PCB concentration and number of years a worker has**
 160 **been at the company. Results are presented for all workers (Figure 1A) and only those**
 161 **with direct contact dismantling transformers (Figure 1B).**

162 **Table 1. PCBs concentrations and enantiomer fractions recorded in workers. DPW =**
 163 **dismantling plant worker and DPC = dismantling plant chief, Concentrations are**
 164 **presented to 3 significant figures and EFs to 2 decimal places <LOD = below level of**
 165 **detection (i.e. S:N ratio <10)**

	Sample ID	Job role	Years with the company	Concentration in serum (ng g ⁻¹)	Lipid corrected concentration in serum (µg g ⁻¹ lipid)		Enantiomer fractions					
					Per sample	Mean [±1 σ]	CB-95	Mean [±1 σ]	CB-136	Mean [±1 σ]	CB-149	Mean [±1 σ]
No direct contact	S001	DPC	21	18.4	2.49	5.83 [± 6.90]	0.51	0.63 [± 0.14]	<LOD	0.44 [± 0.00]	0.47	0.37 [± 0.08]
	S002	Administrative	20	13.7	1.49		0.74		<LOD		<LOD	
	S003	Workshop DPC	10	11.2	1.23		0.41		<LOD		0.41	
	S004	Cleaner	4	29.4	4.08		0.55		<LOD		0.26	
	S005	DPC	20	166	21		0.59		<LOD		0.31	
	S006	DPC	20	86.3	11.1		0.66		0.45		0.39	
	S007	Guard	21	26.6	1.93		0.78		0.44		0.41	
	S008	Maintenance	4	21	3.32		0.82		<LOD		<LOD	
Direct contact	S009	Pumping oil	11	28.4	3.76	17.2 [± 13.0]	0.59	0.74 [± 0.11]	0.45	0.47 [± 0.05]	0.47	0.36 [± 0.07]
	S010	DPW	15	62.8	9.41		0.51		<LOD		0.48	
	S011	DPW	6	73.1	6.53		0.88		0.51		0.37	
	S012	DPW	9	70.1	13.2		0.88		<LOD		0.29	
	S013	DPW	3	27.7	3.37		0.87		0.57		0.44	
	S014	DPW	11	246	36.2		0.87		0.43		0.21	
	S015	DPW	14	163	20.6		0.82		<LOD		<LOD	
	S016	DPW	9	138	14.4		0.79		0.47		0.32	
	S017	DPW	16	121	19.4		0.72		0.43		0.30	
	S018	DPW	17	353	38.5		0.66		0.46		0.35	
	S019	DPW	4	16.8	3		0.60		<LOD		0.33	
	S020	DPW	20	300	37.7		0.87		<LOD		<LOD	
	S021	DPW	9	95.2	14.2		0.74		<LOD		0.37	
	S022	DPW	20	231	39.3		0.67		0.47		0.35	
	S023	DPW	15	62.5	10.4		0.67		0.43		0.35	
	S024	DPW	6	37.5	4.16		0.82		0.51		0.32	
	S025	DPW	5	23.4	4.05		0.66		<LOD		0.44	
	S026	DPW	9	134	27.3		0.66		<LOD		0.44	
	S027	DPW	14	149	21.2		0.77		<LOD		<LOD	
Unknown	S028	Unknown	unknown	26.1	3.55	13.7 [± 14.6]	0.82	0.78 [± 0.15]	0.52	0.55 [± 0.048]	0.36	0.35 [± 0.08]
	S029	Unknown	unknown	187	30.4		0.91		0.27			
	S030	Unknown	unknown	50.7	7.14		0.61		<LOD		0.43	

166 **3.2 Statistical evaluation of PCB signatures**

167 Fifty four PCBs were consistently detected in the > 60% of samples. These were quantified,
168 percent normalised and presented as bar charts to show the PCB signature in each participant
169 (Supplementary Information 1). Only three different Aroclor blends were understood to have
170 been used in transformers; these were A1242, A1254 and A1260 (Johnson et al., 2006). As
171 the dismantling plant was in Europe, transformers were likely to have contained a variety
172 manufacturers products from the European market, including Aroclors, Phenoclor, Pyralenes
173 and Clophens, however there is a very high degree of similarity in the signature from blends
174 with an equivalent chlorine content (Johnson et al., 2006). The signature for the majority of
175 workers was visually similar to the signature of A1260 (Supplementary Information 1) which
176 provides further evidence to suggest occupational exposure had occurred. However, further
177 assessment was undertaken to confirm if this was the source of contamination or if other
178 potential sources were important.

179 The PCB signatures of the workers were assessed using principal component analysis (PCA).
180 Where a PCB was not detected it was included in the dataset as a '0'. As part of the data
181 quality check, other values such as LOD/2 were substituted for '0', but these had no
182 observable effect on the data output and so the '0's were retained. To reduce any bias from a
183 high proportion of non-detects for a specific congener, PCBs that were not detected in over
184 60% of samples (i.e. PCBs present in less than 18 out of the 30 samples) were removed from
185 the analysis following the guidance of Helsel (2006). This resulted in a dataset of 54 PCBs in
186 30 participants. Before performing PCA the data were normalised by transformation to a
187 percent metric to remove concentration/dilution effects. The data were then mean centred and
188 scaled using a Z-transform (autoscale transform) to prevent high concentration variables from
189 dominating the analysis (Johnson et al., 2007). Principal component 1 accounted for 33.1% of

190 the variation and principal component 2 accounted for a further 18.2%. The scores plot is
191 presented as Figure 2 and the loadings plot as Figure 3.

192 The scores plot (Figure 2) displayed a three end member system, showing that there were
193 three groups of workers, each with a different PCB signature. Two of the three groups look as
194 if they are linked to the duration that the participant had been working at the plant and if their
195 job involved direct contact with PCBs. One group may therefore represent participants with
196 prolonged occupational exposure to A1260 and the other group exposure to background
197 levels or a recent minimal exposure to A1260. However, the third group was comprised of
198 participants who all had different ages, jobs and years at the company and may be linked to
199 an additional source(s) of contamination. Bar charts representing the PCB signature were
200 produced to identify differences in the PCB signature between the three groups and aid with
201 the source identification process. Figure 4 displays the signature of A1260 along with the
202 signatures of three participants (selected using the PCA scores plot) to represent the three
203 groups. These were; participant number S020 who had worked at the plant for 21 years
204 (representing a prolonged exposure to A1260), participant number S013 who had been at the
205 company for 3 years (representing a recent minimal exposure to A1260), and participant
206 number S021 (representing a suspected additional exposure source).

207 **3.3 Source identification and age dating exposure**

208 **3.3.1 Occupational exposure to A1260**

209 Participants with a negative score on PC1 and PC2 were comprised of workers who had been
210 working at the dismantling plant for more than 10 years, with direct contact dismantling
211 transformers. The signatures from these samples is similar to the profile of A1260 but with
212 depleted proportions of several of the less chlorinated biphenyls, CB-88 & 95, CB-90 & 101
213 and CB-151 (Figure 4). These are congeners predominantly containing a phenyl group with

214 un-chlorinated *meta* and *para* positions (i.e. 2,5- chlorine substitution), which are particularly
215 susceptible to metabolic attack by P450 cytochromes (Letcher et al., 1999). The signature
216 also contained elevated proportions of CB-74, CB-153, CB-138 and CB-180 (Figure 4).
217 These are congeners containing a phenyl group with 2,4,5- substitution which are particularly
218 resistant to biotransformation and elimination (Megson et al., 2013a). Interestingly this group
219 also included a chief of operations who had low PCB concentrations ($2.49 \mu\text{g g}^{-1}$ lipid) but a
220 signature similar to A1260. The signature shows that this individual appears to have been
221 exposed to PCBs through occupational exposure, although the total PCB concentrations
222 indicate this was only a minimal exposure.

223 **3.3.2 Background or recent minimal occupational exposure**

224 Participants with a positive score on PC2 comprised of workers who had been working at the
225 plant for a relatively short period of time (< 6 years), along with those with jobs that did not
226 involve direct exposure, such as a maintenance worker, cleaner and guard. It also included a
227 chief of operations who had been working at the plant for 10 years but had the lowest PCB
228 concentration of all participants ($1.26 \mu\text{g g}^{-1}$ lipid). All of these individuals contained
229 significantly lower (P-value 0.000) EC7 PCB concentrations (mean value of $3.2 \mu\text{g g}^{-1}$ lipid
230 (± 1.0 ; 1σ)) than the rest of the samples (mean value of $19.1 \mu\text{g g}^{-1}$ lipid (± 12.2 ; 1σ)). The
231 signature from this group contained higher proportions of several episodic congeners such as
232 CB-8, CB-18, CB-31, CB-52 and CB-151 (Figure 3), indicating a recent exposure. The
233 signature also displayed slightly elevated proportions of many of the lower chlorinated PCBs
234 that are not present in high concentrations in A1260 such as CB-28, CB-74, CB-99 and CB-
235 118 (Figure 4), indicating a background exposure (Figure 2).

236

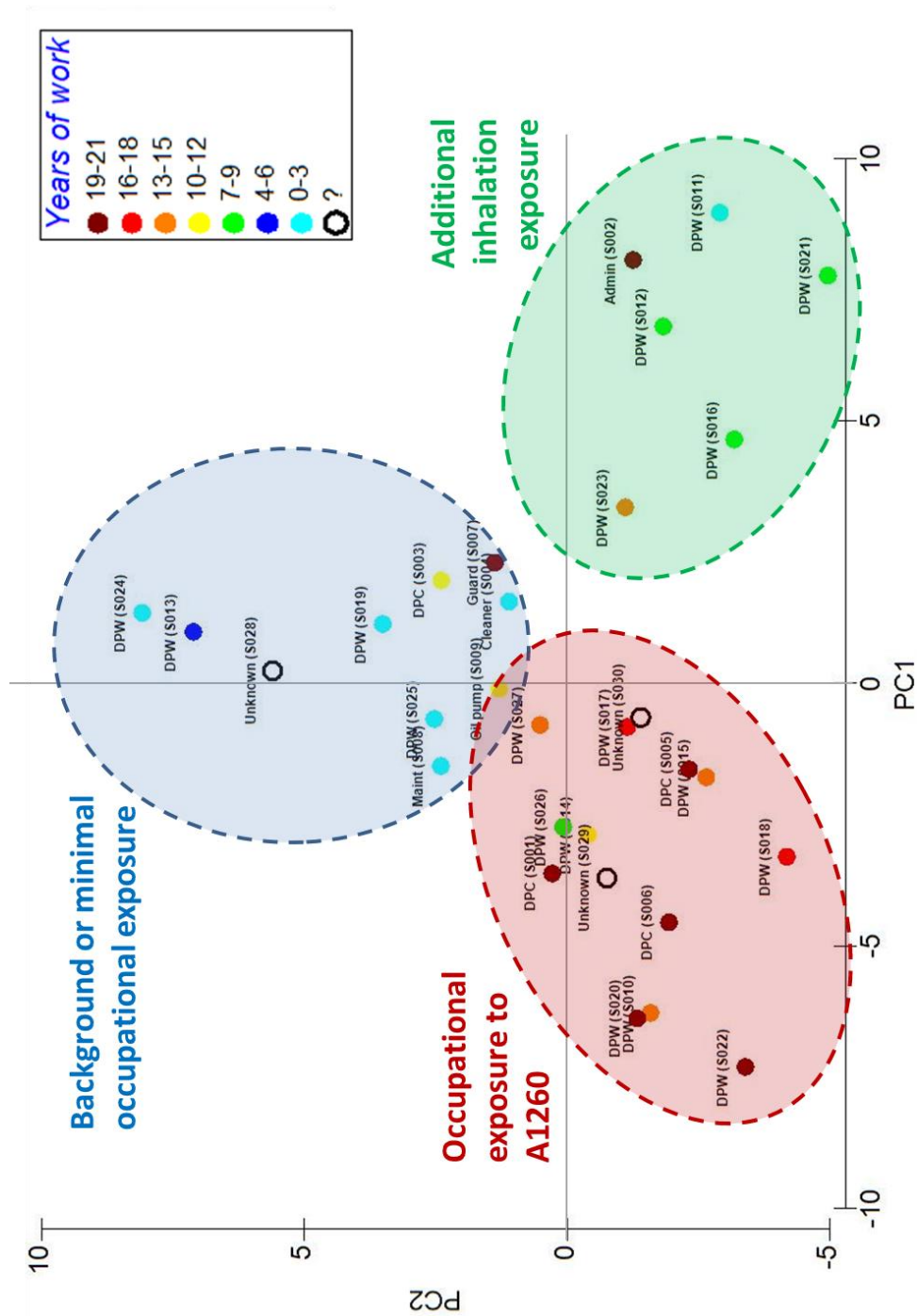


Figure 2. PCA scores plot produced from the PCB signature of workers at the dismantling plant. Data are displayed to show the number of years a worker had been at the plant. Sample identification numbers and job roles are also displayed in short hand; DPW = dismantling plant worker, DPC = dismantling plant chief. For a full list of job roles refer to Table 1

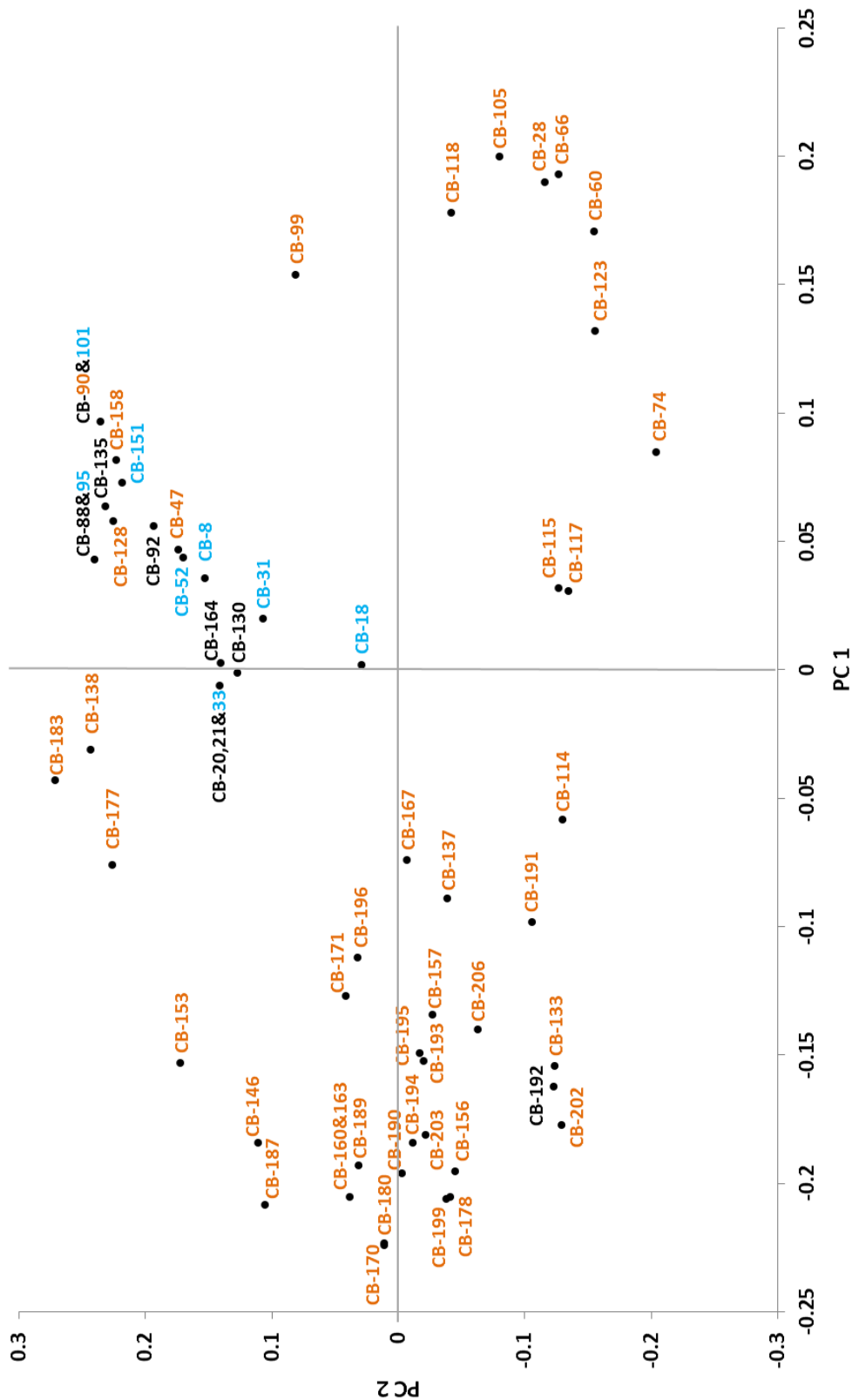
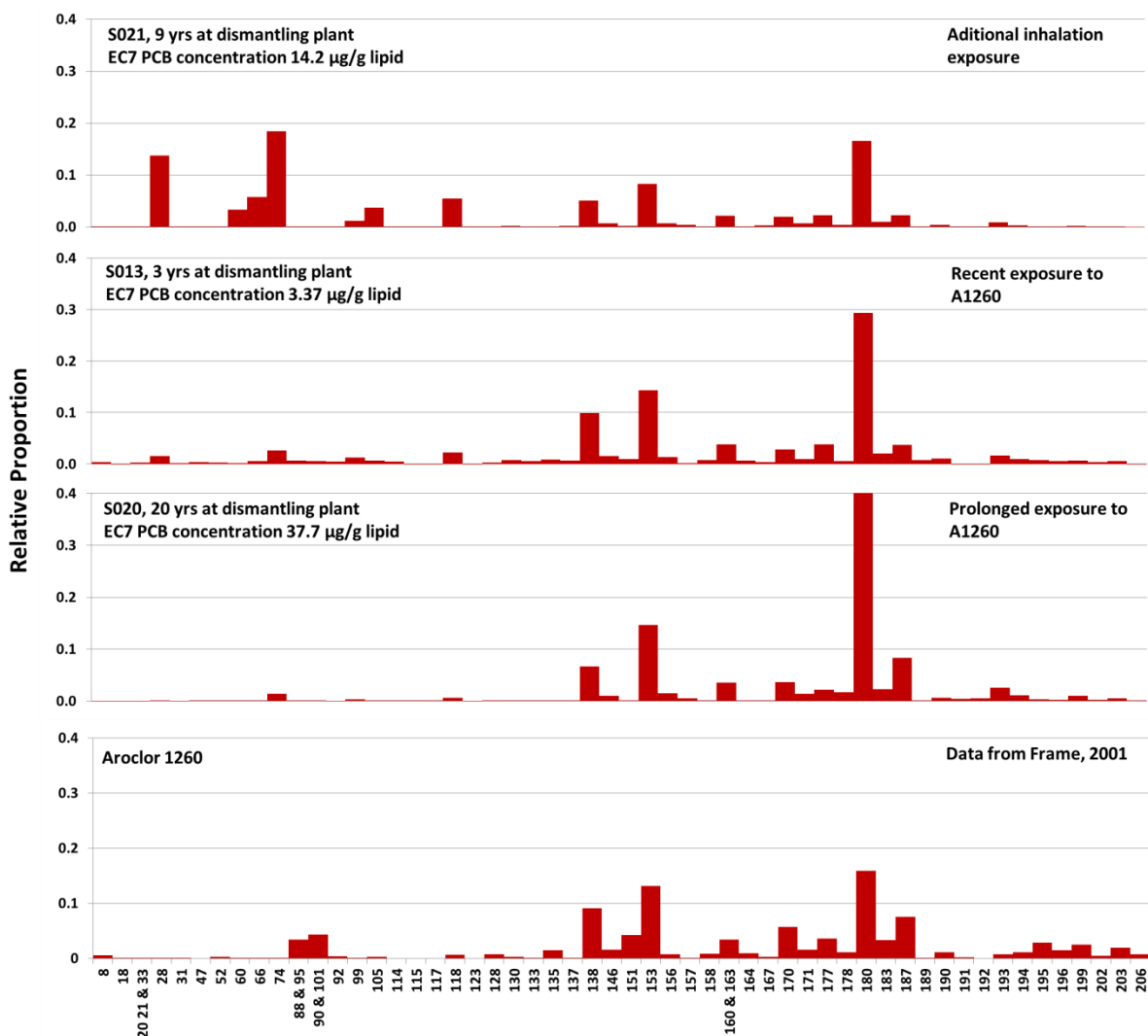


Figure 3. PCA loadings plot, PCBs identified as episodic (Megson et al 2013a) are presented in blue and PCBs identified as steady state (Megson et al 2013a) are presented in orange.



241

242 **Figure 4. PCB signature for the serum samples obtained from participants S020, S013**
 243 **and S021 compared with the signature from A1260.**

244 3.3.3 Additional inhalation exposure

245 In some instances inhalation has proven to be an important PCB exposure pathway
 246 (DeCaprio et al., 2005; Herrick et al., 2011). In a school in Boston (U.S.) inhalation of PCBs
 247 leaching from caulking materials and sealants was determined as the main route of exposure
 248 for teachers. This exposure resulted in the teachers having a distinctive PCB signature, with
 249 proportions of the less chlorinated PCBs such as CB-8, CB-33, CB-37, CB-41, CB-47 and
 250 CB-136 up to five times higher than the control group (Herrick et al., 2011). In this current

251 study, participants with a positive score on PC1 and negative score on PC2 had a signature
252 similar to A1260, but with the addition of high proportions of several less chlorinated PCBs
253 such as CB-28, CB-60, CB-66 and CB-74, along with CB-105 and CB-118. This group was
254 comprised of participants who all had different ages, jobs and years at the company. The
255 signatures all contained higher proportions of more volatile PCBs which have been
256 previously linked to indoor air sources such as leaching from sealants and caulking materials
257 (Harrad et al., 2005; Herrick et al., 2011; Kohler et al., 2005). Several of the congeners
258 present in higher proportions (CB-28, CB-60 and CB-66 and CB-74) have also been linked to
259 exposure from capacitors in electrical equipment (Luotamo et al., 1993). Concentrations of
260 PCBs in individuals from this group were similar to those from the group with prolonged
261 exposure to A1260. Therefore, the results indicate that this group was exposed to an
262 additional source of PCBs through the inhalation of PCBs, possibly through leaching of
263 materials at the home rather than from their workplace (Figure 2).

264 **3.3.4 Enantiomeric fractions**

265 To further elucidate the potential sources of exposure to these individuals we also examined
266 the enantiomer signatures of three chiral PCBs (CB-95, CB-136 and CB-149) in the samples
267 to ascertain if there were any trends in enantio-specific processing of these congeners that
268 could be related to variations in the exposure of the workers, as well as potentially providing
269 further insight into the pharmacokinetics of these pollutants in humans. To our knowledge
270 this is the first time that analysis of this type has been done for human serum samples despite
271 the widespread use of this matrix in human bio-monitoring programs (e.g. Canadian Health
272 Measures Survey, United States Human Bio-monitoring Program).

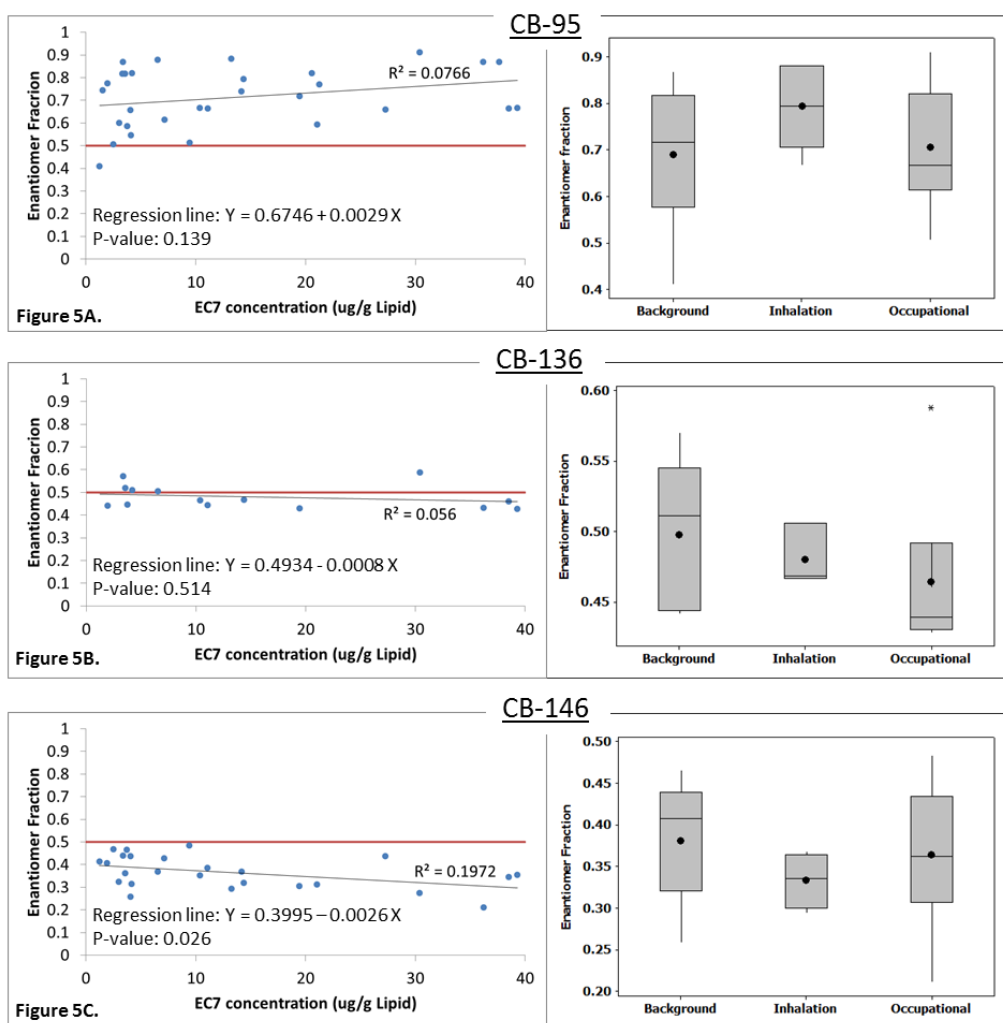
273 The enantiomeric fractions of CB-95 and CB-149 recorded in the workers varied
274 considerably (0.41 to 0.91 and 0.21 to 0.48, respectively), whereas fractions of CB-136
275 remained close to racemic (0.43 to 0.59) (Table 1). For CB-95 most participants contained

276 sera with higher proportions of the *E1* or (+) enantiomer than the *E2* or (-) enantiomer
277 indicating that significant enantioselective metabolism or excretion of this enantiomer has
278 occurred. In one worker (S029) the proportion of *E1* enantiomer was over 10 times greater
279 than the *E2* enantiomer. This mirrors the work of Chu et al (2003) who found similar results
280 for CB-95 in human liver samples. However in this study the extent of change was much
281 great than in Chu et al (2003) with the results here showing evidence of near complete
282 removal of the *E2* enantiomer of in some participants. For CB-149 all participants contained
283 sera with higher proportions of *E2*. In one worker the proportion of *E2* was over 3.5 times
284 greater than *E1* (S014).

285 There were no clear trends in enantiomeric signatures with the exposure type and duration
286 identified in this study (Figure 5). However there was a weak correlation between EFs of CB-
287 95 and CB-149 with EC7 PCB concentrations (Figure 5). Participants with higher PCB
288 concentrations tended display a greater degree of enrichment of the *E1* enantiomer of CB-95
289 (P-value 0.139) and *E2* enantiomer of CB-149 (P-value 0.026). This suggests that there may
290 be a concentration dependent element to the metabolism of these congeners in humans. The
291 high variability in the data for these two congeners also indicates that there is significant
292 intraindividual variation in the enantiospecific processing of these contaminants.

293 Importantly this data also indicates that PCB profiles measured in sera may not fully match
294 those of other bodily tissues, as the data recorded here is in contrast to that recorded by Chu
295 et al (2003) for CB-95, CB-132 and CB-149 in human mussle, kidney and brain samples that
296 were all racemic or nearly racemic but does match that recorded by the authors for liver
297 samples which were largely non-racemic in nature. This suggest that sera may in fact reflect
298 liver profiles only, rather than whole body signatures. This has potentially important
299 implications for sera based human biomonitoring programs such as the Canadian Health
300 Measures Survey (CHMS) and the National Health and Nutrition Examination Survey

301 (NHANES) as it suggest that they may underestimate the true PCB burden and profiles of
 302 subjects. However further work would be needed to conform this theory. This data also
 303 suggests that the enantiomeric profice of the PCBs should be taken into account when
 304 assesing the toxicity of any potential exposure as the persistence and consequent effects of
 305 the enantiomers and there metabolic products may be significantly different (Kodavanti and
 306 Curras-Collazol, 2010).



307

308 **Figure 5. Enantiomer Fraction of CB-95 (Figure 5A), CB-136 (Figure 5B) and CB-149**
 309 **(Figure 5C), and their relationship with EC7 PCB concentrations and the groups**
 310 **identified by PCA. The red line on the scatter plots represents an EF of 0.5 (i.e. a**
 311 **racemic mixture), box plots display the interquartile range, median and mean (•).**

312 **4 Conclusions**

313 Identifying the source of contamination and age dating human exposure to PCBs is a highly
314 complex task. This is due to the wide range of PCB sources that humans are exposed to along
315 with different exposure pathways and processes such as volatilization, dissolution,
316 biodegradation and post uptake processes that can all alter the original PCB signature.
317 However, by considering PCB concentrations along with detailed congener specific
318 signatures it was possible to identify different sources of contamination and gain an insight
319 into both the magnitude and duration of exposure. Occupationally exposed individuals had a
320 similar PCB profile to Aroclor A1260. Individuals with prolonged exposure had depleted
321 proportions of several PCB congeners that are susceptible to metabolism (CB-95, CB-101
322 and CB-151), and elevated proportions of PCBs that are resistant to metabolism (CB-74, CB-
323 153, CB-138 and CB-180). A group of workers were also identified with a suspected
324 additional source of exposure through the inhalation of PCBs, as their sera contained elevated
325 proportions of CB-28, CB-60, CB-66, CB-74, CB-105 and CB-118.

326 Whilst there were no clear trends in enantiomer signatures with the exposure type and
327 duration identified in this study, there was a weak correlation between EFs of CB-95 and CB-
328 149 with EC7 PCB concentrations, suggesting that there may be a concentration dependent
329 element to the metabolism of these congeners in humans. The extent of enantioselective
330 metabolism or excretion in one worker (S029) was so great it resulted in the near complete
331 removal the E2 enantiomer.

332

333 **Acknowledgements**

334 The authors would like to thank; Ann Tanderitispole and Chris Gallagher (University of
335 Strathclyde) for their assistance with sample analysis and the Scottish Funding Council and

336 EPSRC Grant EP/D013739/2 for funding associated with the University of Strathclyde
337 Laboratory. Corina Brimacombe and Terry Kolic for their help with the HRMS analysis of
338 chiral PCBs. Alec Kettle (Leco) for his help and support with the research. All the volunteers
339 who gave blood used in this study. Finally, David Megson would like to thank Plymouth
340 University for funding this project as part of his PhD research.

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