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1 Perfluoroalkylated acids in the eggs of great tits (*Parus major*) near a
2 fluorochemical plant in Flanders, Belgium

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19 **Abstract**

20 Perfluoroalkyl acids (PFAAS) are highly persistent substances which have been detected in wildlife around
21 the world, including birds. Although bird eggs have often been used to determine and monitor PFAAs
22 levels in the marine environment, this has rarely been done in the terrestrial environment. In the present
23 study we examined the concentrations and composition profile of 12 PFAAs (4 perfluoroalkyl sulfonic
24 acids (PFSA) and 8 perfluoroalkyl carboxylic acids (PFCA) in the eggs of great tits (*Parus major*) collected
25 at a fluorochemical plant and in three other areas, representing a gradient in distance from the pollution
26 source (from 1 to 70km), in Antwerp, Belgium.

27 The PFSA concentrations measured at the site of the fluorochemical plant were among the highest ever
28 reported in eggs with median concentrations of 10380 ng/g (extrapolated), 99.3 ng/g and 47.7 ng/g for
29 PFOS, PFHxS and PFDS respectively. Furthermore, the median concentration of 19.8 ng/g for PFOA was
30 also among the highest ever reported in bird eggs. Although these concentrations decreased sharply with
31 distance from the fluorochemical plant, levels found in the adjacent sites were still high compared to what
32 has been reported in literature. Moreover, based on what is known in literature, it is likely that these
33 concentrations may cause toxicological effects. PFOS was the dominant contributor to the PFSA and PFAAs
34 (63.4 – 97.6 %) profile at each site, whereas for PFCA this was PFOA at the plant site and the nearest
35 locations (41.0 – 52.8 %) but PFDoA (37.7 %) at the farthest location.

36 Although there is some evidence that PFAAs concentrations close to the plant site are decreasing in
37 comparison with earlier measurements, which may be due to the phase out of PFOS, more research is
38 necessary to understand the extent of the toxicological effects in the vicinity of this PFAAs hotspot.

39 **Keywords:** Perfluoroalkyl acids, PFAAs, Birds, Eggs, Belgium, Great tit

40

41 **Capsule**

42 Levels of perfluorinated compounds in passerine bird eggs were very high at a perfluorochemical hotspot
43 but decreased sharply with distance.

44

45

46

Introduction

47 Perfluoroalkyl acids (PFAAs) have been produced for more than 50 years. The strength and stability of the
48 C-F binding in combination with the hydrophobic and lipophobic character of PFAAs lead to unique
49 physicochemical properties. PFAAs applications include fire-fighting foams, fast food packaging and
50 surface coatings for carpets. (Buck et al., 2011; Kissa, 2001). PFAAs are highly persistent and may enter
51 the environment either directly or indirectly from environmental degradation of precursors (Buck et al.,
52 2011; Prevedouros et al., 2006). The widespread use of PFAAs has resulted in a global presence in the
53 environment, wildlife and even humans as described in many studies (e.g. Butt et al., 2010; D'Hollander
54 et al., 2010; Giesy & Kannan, 2001, 2002; Houde et al., 2006; Miller et al., 2015).

55 The attention of regulatory agencies and researchers has focused on long chain perfluoroalkyl carboxylic
56 acids (PFCAs) and perfluoroalkyl sulfonic acids (PFSAs), because of their higher bioaccumulative potential
57 compared to their short chain analogues (Buck et al., 2011). They are particularly interested in the two
58 most widely known ones: PFOA ($C_7F_{15}COOH$) and PFOS ($C_8F_{17}SO_3H$).

59 PFOS, PFOA and related compounds have been phased out by 3M, the major global manufacturer, in 2002,
60 due to their persistence, potential health effects and global distribution. Furthermore, PFOS was included
61 in the Stockholm Convention on Persistent Organic Pollutants (POPs) in 2009. These measures, in most
62 cases, appear to be reducing PFOS environmental levels while levels of other PFAAs are still rising (Ahrens
63 et al., 2011; Filipovic et al., 2015; Miller et al., 2015).

64 Bird eggs have been used in multiple studies to monitor PFAAs levels in many regions of the world (e.g.
65 Gebbink & Letcher, 2012; Giesy & Kannan, 2001; Holmström et al., 2005; Miller et al., 2015; Yoo et al.,
66 2008). However, the majority of these studies have been performed on aquatic birds, whereas data on
67 terrestrial birds, especially passerine birds, remain scarce (Ahrens et al., 2011; Custer et al., 2012;
68 Holmström et al., 2010; Rüdél et al., 2011; Yoo et al., 2008).

69 Previous studies conducted near a fluorochemical plant in Antwerp, Belgium, revealed the highest PFOS
70 levels ever found in wildlife (Dauwe et al., 2007; D'Hollander et al., 2014; Hoff et al., 2005; Lopez Antia et
71 al., 2017). Liver PFOS levels measured in great tits (*Parus major*) and blue tits (*Cyanistes caeruleus*) from
72 this area were higher than those measured in top predators in other regions worldwide, and were also
73 above the benchmark concentrations for the possible risk levels of avian species (Dauwe et al., 2007).
74 Furthermore, PFOS levels in eggs were among the highest ever reported in bird eggs worldwide (Lopez
75 Antia et al., 2017). These studies conducted nearby the fluorochemical plant in Antwerp have
76 demonstrated that PFOS levels measured in wildlife decreased significantly at relatively short distances
77 from the plant site (from 3 to 10 km) on the one hand, and that levels found at these distances are still
78 very high on the other hand. Monitoring PFAAs levels and composition profile in this hot spot and its
79 surroundings is therefore extremely important.

80 In the present study concentrations of multiple PFAAs were measured in eggs of a terrestrial songbird,
81 the great tit, at a fluorochemical plant in Antwerp. Additionally eggs from three other areas were analyzed,
82 representing a gradient in distance from the pollution source. It is important to compare the levels and
83 composition profile of PFAAs along this distance gradient to better understand the environmental
84 dynamics of PFAAs. Moreover, the outcome of the present study can be used for further monitoring
85 studies, to investigate temporal changes in PFAAs concentrations using 1) minimally invasive sampling,
86 namely eggs (Furness, 1993), and 2) a species that has demonstrated to be useful as sentinel species for
87 local contamination of Persistent Organic Pollutants (Dauwe et al., 2003, 2007; Van den Steen et al., 2006,
88 2009). Finally, detected levels were used to assess the potential risk to birds based on the current
89 toxicological benchmark levels.

90

91 **Materials and methods**

92 *Study species and Sample collection*

93 Great tits, insectivorous songbirds, are increasingly being used in biomonitoring studies because they
94 readily nest in man-made nestboxes, are abundant and can even be attracted to polluted areas (Eens et
95 al., 1999; Eeva & Lehikoinen, 1995, 1996; Eeva et al., 1998; Dauwe et al., 1999, 2004, 2005; Van den Steen
96 et al., 2006).

97 During the winter of 2011, nestboxes were placed at four sampling sites. Three locations were situated in
98 the vicinity of a perfluorochemical plant (3M) in Antwerp, Belgium. These locations were the
99 perfluorochemical plant itself (32 nestboxes), Vlietbos (1km SE from the plant site; 23 nestboxes) and Rot-
100 Middenvijver (shortly Rot; 2.3km ESE from the plant site; 16 nestboxes). As a reference site, Tessenderlo-
101 Ham (20 nestboxes), approximately 70 km ESE from the plant site was selected, as it is an area without a
102 known perfluorochemical point source in the direct environment.

103 Nestboxes were checked weekly or daily just before laying to be able to determine the laying date and
104 clutch size. At each site one egg per clutch was collected randomly by hand from 10 to 12 different
105 nestboxes before the incubation had started (early April).

106 *Chemical analysis*

107 The used abbreviations of PFAAs are according to Buck et al. (2011). The target analytes included 4 PFSA
108 (PFBS, PFHxS, PFOS and PFDS) and 8 PFCAs (PFBA, PFHxA, PFOA, PFNA, PFDA, PFDoA, PFTrA and PFTeA).

109 The isotopically mass-labelled internal standards (ISTDs) comprised [1,2-¹³C₂]PFHxA, ¹³C₈-PFOA, ¹³C₉-PFNA,
110 [1,2,3,4,5,6-¹³C₆]PFDA, [1,2,3,4,5,6,7-¹³C₇]PFUdA, [1,2,3,4,5,6,7-¹³C₇]PFDoA, ¹⁸O₂-PFHxS and ¹³C₈-PFOS and
111 were purchased by Wellington Laboratories (Guelph, Canada). HPLC-grade Acetonitrile (ACN) and water
112 (Acros Organics, New Jersey, USA) were used.

113 *Sample extraction*

114 After removal of the shell, the content of the egg was homogenized with an Ultra Turrax mixer (T25,
115 Staufen, Germany) in a polypropylene (PP) tube and divided into two parts of approximately 0.5g.

116 The extraction procedure was based on a method described by Powley et al. (2005) with minor
117 modifications. Samples were spiked with an internal standard mixture (ISTD, 80 μL , 125 $\text{pg}/\mu\text{L}$), containing
118 125 $\text{pg}/\mu\text{L}$ of each ISTD and mixed thoroughly. Hereafter 10 mL acetonitrile was added, samples were
119 sonicated (3x10 min) and left overnight at room temperature on a shaking plate. After centrifugation (4°C,
120 10 min, 2400 rpm, Eppendorf centrifuge 5804R), the supernatant was transferred to a 15 mL PP tube and
121 reduced to approximately 0.5 mL by using a rotational-vacuum-concentrator at 20°C (Martin Christ, RVC
122 2-25, Osterode am Harz, Germany). The concentrated extract and 2 times 250 μL acetonitrile, which was
123 used to rinse the tubes, were transferred to a PP micro centrifuge tube containing 50 mg graphitized
124 carbon powder (Supelclean ENVI-Carb, Sigma-Aldrich, Belgium) and 70 μL glacial acetic acid merely to
125 eliminate pigments. These tubes were vortex-mixed during at least one minute and centrifuged (4°C, 10
126 min, 10 000 rpm, Eppendorf centrifuge 5415R). The cleaned-up supernatants were stored at -20°C until
127 analysis. Before analyses, 70 μL of extract was diluted with 130 μL 2mM aqueous ammonium acetate and
128 filtrated through an Ion Chromatography Acrodisc 13mm Syringe Filter with 0.2 μm Supor (PES)
129 Membrane (Leuven, Belgium) attached into a PP auto-injector vial.

130 *UPLC-TQD analysis*

131 We analyzed PFAAs by UPLC coupled tandem ES(-) mass spectrometry (ACQUITY, TQD, Waters, Milford,
132 MA, USA) using an ACQUITY BEH C18 column (2.1 X 50 mm; 1.7 μm , Waters, USA), mobile phase: 0.1 %
133 formic acid in water(A), 0.1 % formic acid in acetonitrile(B), solvent gradient: from 65% A to 0 % A in 3.4
134 min and back to 65%A at 4.7 min, flow rate: 450 $\mu\text{L}/\text{min}$, injection volume: 10 μL . To retain any PFAA
135 contamination originating from the system, we inserted an ACQUITY BEH C18 pre-column (2.1 \times 30 mm;
136 1.7 μm , Waters, USA) between the solvent mixer and the injector. Identification and quantification was

137 based on multiple reaction monitoring (MRM) of the following diagnostic transitions: 213 → 169 (PFBA),
138 313 → 296 (PFHxA), 315 → 270 (¹³C₂-PFHxA), 413 → 369 (PFOA), 421 → 376 (¹³C₈-PFOA), 463 → 419
139 (PFNA), 472 → 427 (¹³C₉-PFNA), 513 → 469 (PFDA), 519 → 474 (¹³C₆-PFDA), 613 → 569 (PFDoA), 613 →
140 319 (PFDoA), 615 → 169 (¹³C₇PFDoA), 615 → 570 (¹³C₇PFDoA), 570 → 525 (¹³C₇PFUdA), 663 → 619 (PFTrA),
141 713 → 669 (PFTeA), 713 → 369 (PFTeA), 299 → 99 (PFBS), 399 → 99 (PFHxS), 403 → 103 (¹⁸O₂-PFHxS),
142 599 → 80 (PFDS), 499 → 80 (PFOS), 499 → 99 (PFOS) and 507 → 80 (¹³C₈-PFOS).

143 *Calibration*

144 Non-labelled standards of all the target analytes were used to construct ten-level calibration curves ($r^2 >$
145 0.99) covering the entire linear range (0.0125 till 16 ng/mL) in HPLC-grade ACN and water. Labeled internal
146 standards were added to each calibration point in the same amount as in samples. Each PFAA was
147 quantified using the corresponding internal standard with the exception of PFBS, PFDS, PFTrA and PFTeA
148 of which no labelled standards were available. PFBS and PFDS were quantified using ¹⁸O₂-PFHxS and ¹³C₄-
149 PFOS respectively, whereas for both PFTrA and PFTeA, ¹³C₂-PFDoA was used. The internal standards
150 allowed us to correct for matrix effects and recovery losses for the corresponding compounds.

151 *Quality assurance.*

152 One procedural blank per 10 samples was analyzed as quality control. Minor levels of contamination (<0.4
153 pg/μL) of PFOA and PFOS were subtracted from the correspondent concentrations found in the samples.
154 For PFOA and PFOS, the quality of the applied method was evaluated by 3 laboratories on spiked egg
155 samples; a triplicate analysis of a sample, spiked with linear (61.7 ng/g and 63.2 ng/g for PFOA and PFOS
156 respectively) or branched (32.2 ng/g and 32.0 ng/g for PFOA and PFOS respectively) isomers of PFOS and
157 PFOA, was performed in each laboratory (Table S1). No significant differences were detected between the
158 laboratories. For the spiked samples, an accuracy of 93 – 107% was achieved. The precision of the applied
159 method varied between 2-4% (Table S1). The limit of quantifications (LOQs), corresponding to a signal-to-

160 noise ratio 10, ranged from 0.02 ng/g to 1 ng/g for PFBS, PFHxS, PFOS, PFDS, PFOA, PFDoA and PFTTrA. Due
161 to some high noise levels the LOQs for PFBA, PFHxA, PFNA, PFDA and PFTTeA are considerably higher and
162 ranged from 1.4 ng/g to 4.3 ng/g. Individual LOQs are displayed in Table 1. For all samples, of which
163 concentrations were within the linear range of the calibration curve, recoveries of the ISTDs were
164 calculated. The samples were corrected for recoveries, which were between 92% and 110%. At two
165 locations, some PFOS concentrations were outside the linear range of the calibration curve and therefore
166 the samples were 10 to 800 times diluted. As a consequence, the internal standards were no longer visible
167 and therefore a correction based on the recoveries was extrapolated.

168 *Statistical analysis.*

169 Statistical analyses were performed using SPSS 23. Samples with a bad recovery were excluded from the
170 analyses. PFAAs concentrations were log transformed to obtain a normal distribution.

171 Differences in concentrations between the different sampling locations were evaluated in two ways. First
172 of all, we performed a one way ANOVA using Least Significant Difference (LSD) test for Post-hoc analysis
173 for PFAAs found in all samples, i.e. PFOS and PFOA. Secondly, for PFAAs with at least one value above the
174 LOQ (i.e. PFDoA, PFTTrA, PFHxS and PFDS), we used a reverse Kaplan Meier (KM) analysis and a Mantel-
175 Cox test for pairwise comparisons among sampling sites. This analysis is commonly used for survival
176 analysis of left censored data (Gillespie et al., 2010) and has been proven useful to cope with levels below
177 the LOQ (Jaspers et al., 2013). Details about how to perform this analysis with SPSS are provided in
178 Gillespie et al. (2010). As reverse KM is a nonparametric analysis we used untransformed data to perform
179 the analysis.

180 To study correlations between the individual compound concentrations, and between the Σ PFSAAs and
181 Σ PFCAAs in each study site Spearman rank correlation analyses were performed.

182 For each site the composition profiles were determined by calculating the proportions of individual
183 compounds to the total concentrations of PFAAs, PFSAs and PFCAs in each egg and then averaging the
184 percentages for all the eggs at a site. For this calculation, values below the LOQ were replaced with a value
185 of LOQ/2 (Bervoets et al., 2004; Custer et al., 2000).

186 **Results**

187 *PFAAs levels*

188 An overview of median levels, ranges and detection frequencies of PFAAs in the eggs is given in Table 1.
189 Some PFOS levels at 3M and Vlietbos exceeded the linear range of the calibration curve and were thus
190 higher than 16 ng/mL. Although these levels were already very high, the extrapolated levels have been
191 used in this study.

192 PFOS, PFOA, PFDoA and PFTrA were detected at all the sampling sites. PFHxS and PFDS were only detected
193 at 2 sampling sites (at the plant site and 1km away from the plant site, at Vlietbos). PFDA and PFNA were
194 only detected at the plant site. PFBS, PFBA, PFHxA and PFTeA were not detected in any of the samples at
195 any of the sites. The overall detection frequencies of the analyzed PFAAs decreased in following order:
196 both PFOS and PFOA were detected in all the samples (100%), followed by PFDoA (60%), PFTrA (56%),
197 PFHxS (38%), PFDS (33%), PFDA (16%) and PFNA (11%). The detection frequencies should be interpreted
198 with caution as there were relatively large differences between the LOQs.

199 Significant differences between sampling sites in PFOS ($F_{3,44}=114.15$, $p<0.001$) and PFOA ($F_{3,44}=77.14$,
200 $p<0.001$) concentrations were observed. Post hoc test revealed that levels were significantly higher at the
201 plant site compared to Vlietbos, 't Rot and Tessenderlo (all $p<0.001$). For PFOS, significant differences
202 were found also between Vlietbos and Tessenderlo (all $p<0.001$) and between Rot and Tessenderlo (all
203 $p<0.001$) but not between Vlietbos and Rot. Concentrations of PFOA were significantly higher at the plant
204 than all of the other sites (all $p<0.001$). For PFDoA and PFTrA levels were significantly higher at the plant

205 site compared with all the other sampling sites (all $\chi^2 \geq 24.79$, all $p < 0.05$) but no significant differences
206 existed among the other sampling sites. Finally, significantly higher levels of PFHxS ($\chi^2 = 24.7$, $p < 0.001$) and
207 PFDS ($\chi^2 = 19.9$, all $p < 0.001$) were found at the plant site compared to Vlietbos.

208 Figure 1 shows the PFAAs concentrations in function of the distance from the pollution source (the center
209 of the plant site is considered to be the pollution source (0m).

210 *PFAAs profile*

211 For all the sampling sites PFOS was the dominant contributor to the Σ PFSA (Figure S1) and to the Σ PFAAs
212 as its contribution to the Σ PFAAs ranged between 97.6 ± 0.3 % (mean \pm SE) at the plant site and 63.4 ± 6.4
213 % at Tessenderlo. For Σ PFCA, the major compound was PFOA at the plant site (43.2 ± 6.5 %), Vlietbos
214 (52.8 ± 4.3 %) and Rot (41 ± 5.6 %), but not at Tessenderlo where it accounted for the 27.0 ± 7.5 % and
215 where PFDoA and PFTrA represented 37.7 ± 6.5 % and 35.2 ± 5.2 % respectively (Figure 2).

216

217 *Correlations*

218 All correlations found among PFAAs at the different sampling sites are summarized in Table 2. Significant
219 correlations were observed mostly at the plant site (14 significant correlations), followed by Vlietbos (13),
220 Rot (3) and Tessenderlo (where no significant correlations were observed). All significant correlations
221 were positive. At the plant site PFOS, PFDS, PFDoA and PFTrA were all correlated with each other, whereas
222 PFOA, PFHxS, PFNA and PFDA were also correlated with each other. However, PFNA was also correlated
223 with PFDoA. At Vlietbos PFOS levels were correlated with levels of all other compounds. Although many
224 of these compounds were also related with each other, no correlations were observed between PFTrA
225 and PFOA, PFHxS and PFDS. At Rot PFOS was only correlated with PFDoA, which was also correlated with
226 PFTrA. Overall PFCA levels (Σ PFCA) were correlated with overall PFSA levels (Σ PFSA) at the plant site,
227 Vlietbos and Rot (Figure 3).

228

229 **Discussion**

230 *PFAAs levels*

231 At the plant site, the observed concentrations of the detected PFSAs (PFOS, PFHxS and PFDS) were among
232 the highest ever reported in bird eggs with median concentrations of 10380 ng/g, 99.3 ng/g and 47.7 ng/g
233 respectively. The median PFOA concentration (19.8 ng/g) was also among the highest ever reported in
234 bird eggs.

235 Compared to a study in 2006 on PFOS in eggs and blood of great tit, northern lapwing and Mediterranean
236 gull, near the same fluorochemical plant (Lopez Antia et al., 2017), ranges of PFOS levels at Vlietbos were
237 approximately 6.5 times lower in the present study. However, the highest concentration reported in
238 northern lapwing (90m from the fluorochemical plant) was 1.5 times lower than the highest concentration
239 in great tit at the fluorochemical plant in the present study, but 4 times higher than the median PFOS level
240 at the plant.

241 To be able to make comparisons among species, some examples of PFAAs concentrations found previously
242 in terrestrial and marine bird eggs are shown in Table 3.

243 To the best of our knowledge only four papers on PFAAs in passerine birds have been published. Until
244 now, the highest concentrations in passerine birds were found by Yoo et al. (2008) in parrot bill
245 (*Paradoxornis webbiana*) eggs collected around the shores of a lake in Korea that receives wastewaters
246 from an industrial complex. Mean PFOS concentrations detected in that study (314 ng/g) were slightly
247 higher than the one found in Vlietbos but more than 60 times lower than the one found at the plant site
248 in the current study. Interestingly, Σ PFCAs concentrations measured in the study in Korea are much higher

249 than the one found in the present study, mainly due to PFNA (40 ng/g) and PFDA (114.2 ng/g)
250 concentrations, suggesting a different type of contamination between both places.

251 The highest mean PFOS concentration detected at the plant site in the present study is more than 18 times
252 higher than the highest mean concentration reported in Great Blue Heron (*Ardea herodias*) eggs (1014
253 ng/g) in 1993 in the Mississippi river, in a colony located approximately 20 km from a 3M fluorochemical
254 plant site (Custer et al., 2010). In the same study, concentrations of the other reported PFAAs (PFDS,
255 PFHxS, PFDA, PFNA, and PFOA) were also, at least two times, lower than the ones we measured in the
256 present study. It must be considered that PFOS was still produced in the plant when the Great Blue Heron
257 eggs were collected. Mean concentrations found in the same Great Blue Heron colony in 2010 (465 ng/g;
258 Custer et al., 2013) remain among the highest ever found, however they are 41 times lower than the one
259 reported at the fluorochemical plant in the present study. Mean PFOS (109 ng/g), PFHxS (0.52 ng/g) and
260 PFOA (0.9 ng/g) concentrations found in free range chicken eggs (*Gallus gallus*) collected at a distance of
261 less than 500 m from a fluorochemical manufacturing plant in China (Wang et al., 2010), were lower than
262 those found in the present study in Vlietbos and very similar to the ones found in Rot, located at 1 and 2.3
263 km from the plant site respectively. In fact, PFOS and PFOA levels found 1km away from the
264 fluorochemical plant in China (9.8 and 0.15 ng/g respectively) were lower than those found in our
265 reference site, Tessenderlo, 70 km away from the plant site. We must consider that when the study of
266 Wang et al. (2010) was performed the plant in China still produced PFOS and related compounds. A study
267 on PFOS concentrations in chicken eggs in Belgium also showed high concentrations of PFOS (highest
268 mean concentration of 3500 ng/g) in the vicinity of the same fluorochemical plant as in the present study
269 (D'Hollander et al., 2011). However, this mean concentration is similar to the lower PFOS levels at the
270 plant site in the present study.

271 The present study site contains one of the highest PFOS levels ever reported in wildlife worldwide. This
272 was confirmed by previous studies performed in the surroundings of this hot-spot in Antwerp in which

273 PFOS levels measured in wood mice (*Apodemus sylvaticus*) livers (D'Hollander et al., 2014; Hoff et al.,
274 2004), great and blue tit nestlings livers (Hoff et al., 2005) and great tit blood and plasma (Dauwe et al.,
275 2007) were all the highest ever reported in these matrices in wildlife.

276 We found a steep decrease in concentrations of all the detected compounds with distance from the plant
277 site. However, significant differences in concentrations were not detected between the sites 1km or
278 2.3km away from the plant and only for PFOS and PFOA significantly lower concentrations were found at
279 the reference site (70 km away from the plant site) comparing with the two other points outside the plant.
280 This decrease with distance from the pollution source was also observed for PFOS concentrations in the
281 aforementioned studies conducted in this area (Dauwe et al., 2007; D'Hollander et al., 2014; Hoff et al.,
282 2004; Hoff et al., 2005). In these studies, as in ours, despite the decrease in concentrations with distance
283 (between 3 and 10km away from the plant site), levels found in the furthest sites remained high
284 comparing with the literature. Also in China, in the surroundings of a perfluorochemical manufacturing
285 facility, a decreasing trend of PFOA, PFOS and PFHxS concentrations in soils, water, and chicken eggs with
286 increasing distance from the plant was observed (Wang et al., 2010).

287 Within each site, the variability between the concentrations of the individual PFAAs of the different
288 nestboxes was considerably high. The highest variability was observed at the plant site where the
289 minimum and maximum concentrations varied up to 20 times. The variability of the PFSA concentrations
290 was higher compared to those observed for the PFCAs. These differences could be indicating variations in
291 concentrations in the laying females. For example, higher PFOS levels were found in young great tits
292 (<one-year old) than in old ones (>1 year-old) in a study performed in the same study area than ours
293 (Dauwe et al., 2007). Unfortunately we know neither the age of the laying females in the present study,
294 nor the origin of these birds (locally born versus immigrant females). Therefore, we do not know the
295 degree of prior exposure. On the other hand, this variability found at the plant site may also be due to
296 variations in egg concentrations within the clutches. Variations within the clutch have been demonstrated

297 in a study about PFOS levels in eggs of tree swallow, in a PFAAs contaminated area in Minnesota, where
298 a 4-fold difference between the highest and lowest concentration within a clutch was found (Custer et al.,
299 2012). Moreover, a study of PFAAs concentrations in entire clutches of Audouins' gulls demonstrated that
300 PFOS concentrations decreased with the laying order of the eggs (Vicente et al., 2015). Unfortunately, in
301 the present study, the eggs were randomly collected before incubation so we could not evaluate the effect
302 of the laying order.

303 *PFAAs profile*

304 Our results showed that PFOS is the major contributor to the total PFAAs. This is in agreement with the
305 literature on PFAAs in bird eggs (e.g. Ahrens et al., 2011; Custer et al., 2012; Nordén et al., 2013; Rüdél et
306 al., 2011). Regarding the PFCAs composition profile, PFOA, followed by PFDoA, is the major contributor at
307 the plant site, Vlietbos and Rot whereas at PFDoA and PFTrA are the major contributors at Tessenderlo.
308 Moreover, a trend can be observed for PFOA to reduce and PFTrA to increase their relative concentrations
309 with the increase in distance (Figure 2). In the plant site and surrounding areas the PFCAs composition
310 profile could be explained by the influence of a direct contamination source, where PFOA is the main
311 product (Prevedouros et al., 2006), whereas 70km away from the plant site, in Tessenderlo, the
312 composition profile could be explained by the atmospheric and biological degradation of the volatile
313 polyfluorinated precursor compounds (fluorotelomer alcohols; FTOH), and the fact that long chain
314 fluorinated compounds (PFTrA and PFDoA) are more bioaccumulative than shorter chain ones (PFOA),
315 (Armitage et al., 2009; Conder et al., 2008; Ellis et al., 2004; Houde et al., 2006). The composition profile
316 found in Tessenderlo is similar to the ones found in eggs of tawny owl in Norway (Ahrens et al., 2011) and
317 peregrine falcon in Sweden (Holmström et al., 2010), where PFTrA and PFUndA were the major
318 contributors to Σ PFCAs.

319 *Toxicological implications*

320 The toxicological and biological effects of PFAAs on avian species are not well characterized but several
321 laboratory studies have verified developmental toxicity (Cassone et al., 2012; Jiang et al., 2012; Molina et
322 al., 2006; O'Brien et al., 2009a,b ; Pinkas et al., 2010). Furthermore, negative effects on the
323 neuroendocrine system (Cassone et al., 2012; Smits & Nain, 2013; Vongphachan et al., 2011) and histology
324 (Molina et al., 2006) have been suggested. Most of these studies focus on the effects of PFOS and PFOA
325 while information on other PFAAs is limited. Additionally, there is a considerable variation in the effect
326 concentrations. For example, *in ovo* exposure to PFOS in chicken eggs determined an LD50 based on
327 hatchability of 4.9 µg/g (Molina et al., 2006) whereas O'Brien et al. (2009a) established it as 93 µg/g. These
328 levels are in the same order of magnitude as the levels found in the present study. However, most of the
329 effects on PFAAs in the laboratories have been established after egg injection which strongly differs from
330 the exposure route of the eggs in the present study.

331 Regarding PFOS, both laboratory and field studies are present. Molina et al. (2006) indicated that PFOS
332 caused a significant reduced hatchability of the chicken embryo after *in ovo* exposure at a dose as low as
333 0.1 µg/g egg. Pathological changes in the liver, including bile duct hyperplasia, periportal inflammation
334 and necrosis were observed at a dose of 1.0 µg/g after *in ovo* exposure. Peden-Adams et al. (2009)
335 observed increased spleen mass, increased lysozyme activity, suppressed total sheep red blood cell -
336 specific immunoglobulin production, shorter right wings and increased frequency and severity of brain
337 symmetry in chickens at *in ovo* exposure level of 1 µg/g. Newsted et al. (2005) derived a predicted-no-
338 effect concentration of 1 µg/ml PFOS in egg yolk based on chronic and acute dietary exposures of northern
339 bobwhite quail (*Colinus virginianus*) and mallard (*Anas platyrhynchos*). According to these values, the
340 PFOS concentrations observed at the plant site may cause physiological and neurological effects on great
341 tits if we assume equal sensitivity between species. Moreover, in a PFAAs contaminated area in east
342 central Minnesota, USA, reduced hatching success was associated with PFOS concentrations as low as 150
343 ng/g in eggs of tree swallow (*Tachycineta bicolor*) (Custer et al., 2012; Custer et al., 2014). If great tit have

344 the same sensitivity as tree swallow, the current PFOS contamination at all the sampling locations, except
345 Tessenderlo, would result in reduced hatchability.

346 The available studies on the toxicity of other PFAAs to birds are limited. The toxic effects of PFOA, PFUdA
347 and PFDS on hatching success and liver mRNA expression in chicken embryos after *in ovo* exposure were
348 evaluated by O'Brien et al. (2009b). Even at the highest exposed group of 10 µg/g these PFAAs did not
349 influence the hatching success. Furthermore, Smits and Nain (2013) evaluated the immunotoxicity of
350 subchronic exposure to PFOA via drinking water in Japanese quail and they found that although birds
351 exposed to the highest dose (10 µg/g) presented a reduced T cell immune response. This reduced
352 response did not translate into an increased disease susceptibility. However, they also found that the
353 highest dose of PFOA reduced thyroid hormone levels and increased the growth rate of exposed Japanese
354 quail (*Coturnix coturnix japonica*).

355 Reduced hatching success and a decrease in tarsus length and embryo mass have been observed in
356 chickens that were exposed *in ovo* to PFHxS concentrations up to 38 µg/g (Cassone et al., 2012).
357 Furthermore, a reduction in plasma thyroid hormone levels was observed at concentrations up to 0.89
358 µg/g (Cassone et al., 2012), a level about 5 times higher than the one found at the plant site in the present
359 study.

360 We have to consider that while most of these toxicological studies were focussed on the effects of a single
361 compound, free-living animals such as the great tits in the present study are exposed to a mixture of PFAAs
362 and other contaminants in combination with natural stressors and therefore more research on
363 toxicological effects under real conditions is urgently needed.

364 **Conclusion**

365 Even though PFOS concentrations have been decreased since the phase out in 2002, the PFAAs
366 concentrations, especially these of PFHxS, PFOS, PFDS and PFOA, in the eggs of great tit at the plant site
367 in 2011 were still among the highest ever reported in wild birds. Furthermore, levels in adjacent sites
368 decreased with distance from the fluorochemical plant, but remained high compared to what has been
369 reported in literature. It is therefore expected that concentrations have decreased further since the
370 present study, although this remains to be tested.

371 More research on PFAAs toxicological effects along with studies on other bird species and biota (to cover
372 the entire food chain) is needed to understand the extent of the problem in this PFAAs contamination hot
373 spot and its surroundings.

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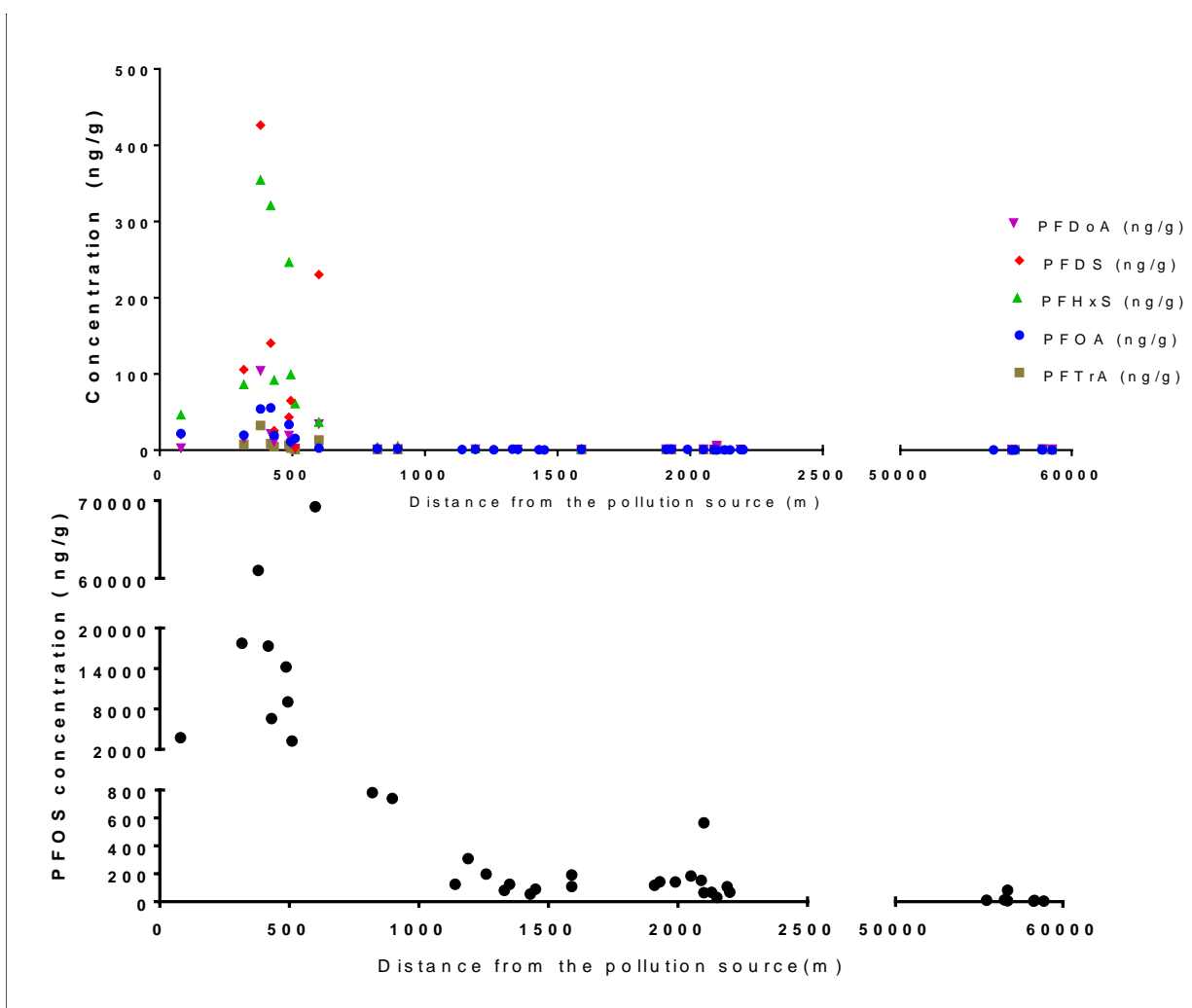
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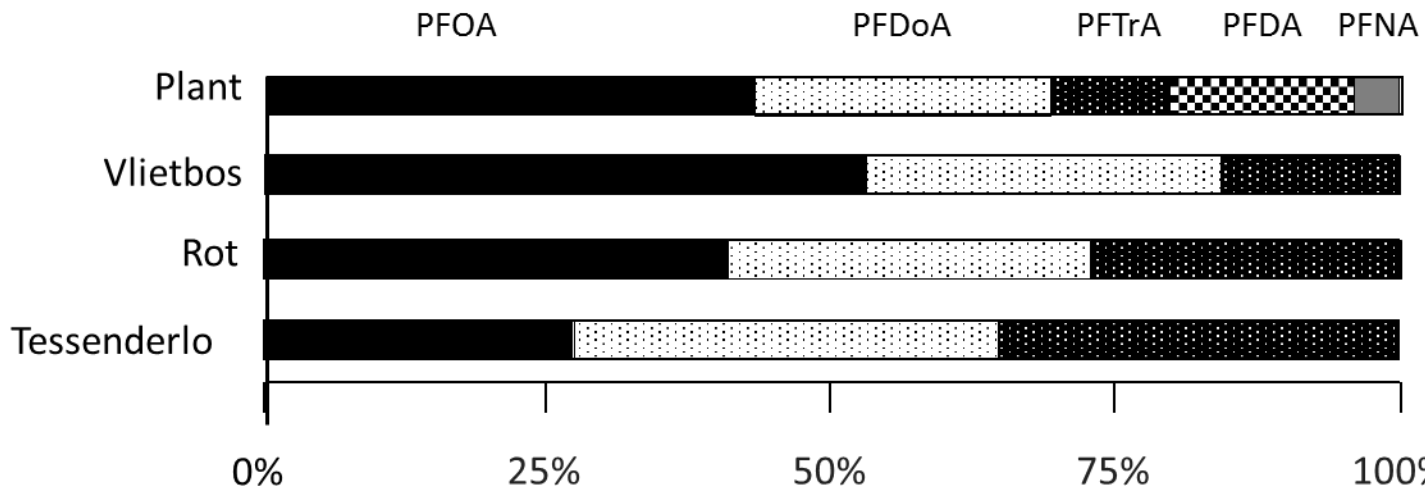
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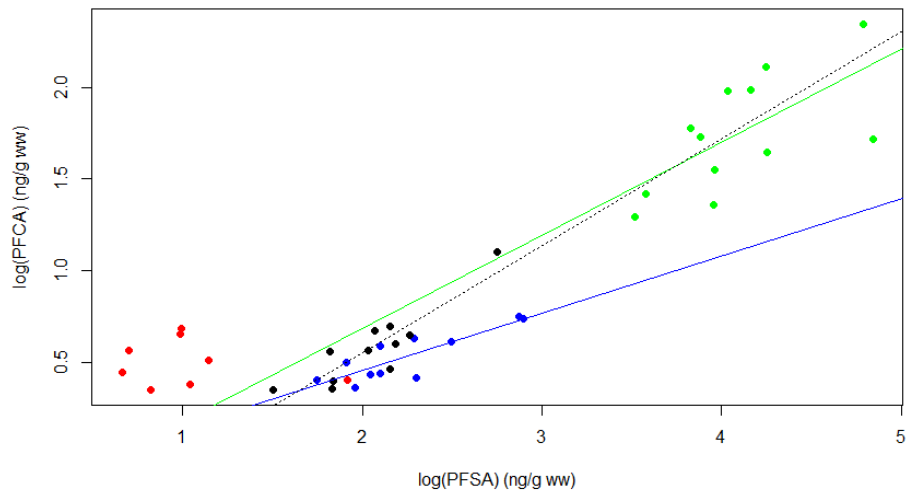


542
543 *Figure 1. a) PFAAs concentrations along the distance gradient from the pollution source. The center of the fluorochemical plant is*
544 *considered to be the pollution source (0m). b) PFOS concentrations along the distance gradient from the pollution source.*



546

547 *Figure 2. Composition profile of PFCA in eggs of great tit at the four sampling sites; a fluorochemical plant and at three sites*
 548 *with increasing distance from the plant site: 1 km (Vlietbos), 2.3 km (Rot) and 70 km (Tessenderlo, reference site).*



549

550 *Figure 3. Correlations between \sum PFSA and \sum PFCA concentrations amongst sites. Green = 3M, Blue = Vlietbos, Black =*
 551 *Middenvijver-Rot and Red = Tessenderlo.*

552 Table 1. Individual limits of quantification (LOQ: ng/g, determined as 10 times the signal to noise ratio), median and mean
 553 concentrations (ng/g ww), range (ng/g ww) and detection frequencies (Freq) of PFAAs in eggs of great tit at the four sampling
 554 sites: a perfluorochemical plant and at three sites with an increasing distance from the plant site (i.e. 1 km Vlietbos, 2.3 km Rot
 555 and 70 km Tessenderlo). ND = not detected. Different letters indicate significant differences ($p \leq 0.05$) between sampling sites in
 556 each compound concentration and in each compound prevalence respectively.

		PFCAs							PFASs				
		PFOA	PFDA	PFNA	PFD _o A	PFT _r A	PFH _x A	PFBA	PFT _e A	PFOS	PFH _x S	PFDS	PFBS
LOQ		0.02	1.4	1.8	0.32	0.38	2.9	4.3	2.1	0.02	0.45	0.2	1.1
Plant (n = 11)	Median	19.8A	12.0	<LOQ	13.7A	5.6A	ND	ND	ND	10380A	99.3A	47.7A	ND
	Mean	26.9	12.3	4.2	22.0	7.9	ND	ND	ND	20122	162.3	100.8	ND
	Range	2.7 - 56.3	<LOQ - 37.2	<LOQ - 20.5	2.0 - 103.9	<LOQ - 32.3				3237 - 69218	36.9 - 354.6	<LOQ - 426.3	
	Freq	100	58.3	41.6	100	91.7				100	100	91.6	
Vlietbos (n = 11)	Median	0.9B	<LOQ	<LOQ	1.0B	<LOQB	ND	ND	ND	125B	<LOQB	<LOQB	ND
	Mean	1.0	<LOQ	<LOQ	0.7	0.4	ND	ND	ND	254	1.6	0.7	ND
	Range	0.3 - 1.9	/	/	<LOQ - 1.50	<LOQ - 0.9				55.1 - 782	<LOQ - 5.6	<LOQ - 2.9	
	Freq	100			50	25				100	50	25	
Rot (n=11)	Median	0.8B	<LOQ	<LOQ	<LOQB	<LOQB	ND	ND	ND	107.1B	<LOQ	<LOQ	ND
	Mean	0.7	<LOQ	<LOQ	1.0	0.8	ND	ND	ND	133.2	<LOQ	<LOQ	ND
	Range	0.3 - 1.3	/	/	<LOQ - 6.0	<LOQ - 4.8				4.3 - 565.3	/	/	
	Freq	100			54.5	54.5				100			
Tessenderlo (n = 8)	Median	0.3B	<LOQ	<LOQ	0.6B	0.5B	ND	ND	ND	9.4C	<LOQ	<LOQ	ND
	Mean	0.3	<LOQ	<LOQ	0.8	0.6	ND	ND	ND	17.6	<LOQ	<LOQ	ND
	Range	0.1 - 0.8	/	/	<LOQ - 1.9	<LOQ - 1.6				4.3 - 82.2	/	/	
	Freq	100			40	50				100			

557

558

		Plant site (n = 11)		Vlietbos (n = 11)		Rot (n = 11)		Tessenderlo (n = 8)	
		p-value	Rho	p-value	Rho	p-value	Rho	p-value	Rho
PFOS	PFOA	0.654	0.155	0.016	0.702	0.818	0.082	0.233	-0.7
	PFHxS	0.451	0.255	0.008	0.745	/	/	/	/
	PFDS	<0.001	0.891	0.005	0.776	/	/	/	/
	PFNA	0.296	0.347	/	/	/	/	/	/
	PFDA	0.614	0.172	/	/	/	/	/	/
	PFDoA	<0.001	0.927	<0.001	0.898	0.009	0.739	0.858	-0.112
	PFTTrA	<0.001	0.900	0.015	0.707	0.097	0.525	0.614	0.308
PFOA	PFHxS	0.031	0.664	0.038	0.630	/	/	/	/
	PFDS	0.435	0.264	0.024	0.671	/	/	/	/
	PFNA	0.013	0.719	/	/	/	/	/	/
	PFDA	0.031	0.648	/	/	/	/	/	/
	PFDoA	0.341	0.318	0.015	0.706	0.620	0.169	0.718	-0.224
	PFTTrA	0.755	0.109	0.062	0.579	0.481	0.238	0.219	-0.667
PFHxS	PFDS	0.503	0.227	0.009	0.744	/	/	/	/
	PFNA	0.007	0.758	/	/	/	/	/	/
	PFDA	0.022	0.677	/	/	/	/	/	/
	PFDoA	0.214	0.409	0.038	0.629	/	/	/	/
	PFTTrA	0.341	0.318	0.052	0.597	/	/	/	/
PFDS	PFNA	0.197	0.421	/	/	/	/	/	/
	PFDA	0.427	0.267	/	/	/	/	/	/
	PFDoA	0.003	0.827	0.002	0.825	/	/	/	/
	PFTTrA	0.010	0.755	0.089	0.536	/	/	/	/
PFNA	PFDA	<0.001	0.858	/	/	/	/	/	/
	PFDoA	0.038	0.630	/	/	/	/	/	/
	PFTTrA	0.113	0.506	/	/	/	/	/	/

560

561 *Table 2 (continued). Correlations found between different PFAAs in the different sampling sites. Values in bold are significant*
 562 *correlations.*

		Plant site (n = 11)		Vlietbos (n = 11)		Rot (n = 11)		Tessenderlo (n = 8)	
		p-value	Rho	p-value	Rho	p-value	Rho	p-value	Rho
PFDA	PFD _o A	0.210	0.410	/	/	/	/	/	/
	PFT _r A	0.462	0.248	/	/	/	/	/	/
PFD _o A	PFT _r A	<0.001	0.945	0.009	0.740	0.038	0.641	0.199	0.688
∑PFSA	∑PFCA	0.034	0.6	<0.001	0.882	0.006	0.791	0.683	0.3

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565 Table 3. Median PFAAs concentrations in bird eggs (ng/g ww) published in the literature. All studies on levels in terrestrial birds
 566 and four studies with higher levels in sea birds have been included. *Geometric means; ** median levels; *** range; NP=levels
 567 were measured but were not provided; NA = not assessed; § Levels found in active fluorochemical plant † Levels found in a
 568 fluorochemical plant not used since 2002.

Species	Country	Year	PFHxS	PFOS	PFDS	PFOA	PFNA	PFDA	PFDoA	PFTra	PFTeA	Publication
<i>Corvus frugilegus</i> **	Germany	2009	<LOQ	5.3	NA	0.5	2.1	0.8	NA	NA	NA	Rüdel et al., 2011
<i>Paradoxornis webbiana</i>	Korea	2006	1.3	314.1	1.1	0.8	40	114.2	25.6	NA	NA	Yoo et al. 2008
<i>Strix aluco</i> *	Norway	1986-2009	0.05	10.9	0.06	<LOQ	<LOQ	0.20	0.12	0.36	NA	Ahrens et al.2011
<i>Falco peregrinus</i>	Sweden	2006	0.80	83	0.66	<LOD	1.6	3.1	3.2	7.3	2.7	Holmström et al. 2010
<i>Tachycineta bicolor</i>	Minnesota (USA)	2008-2009	NP	141	NA	<LOD	NP	5.51	NP	NA	NA	Custer et al. 2012
<i>Tachycineta bicolor</i> *	Minnesota (USA)	2007-2011	0.95	270	NA	18.7	3.10	5.47	1.96	NA	NA	Custer et al. 2014
<i>Gallus gallus</i> §	China	2009	0.52	109	NA	0.9	<LOD	<LOD	<LOD	NA	NA	Wang et al. 2010
<i>Gallus gallus</i> †	China	2009	0.24	85.2	NA	0.76	<LOD	<LOD	<LOD	NA	NA	Wang et al. 2010
<i>Ardea herodias</i> *	Minnesota (USA)	1993	1.5	940	33	<LOD	0.9	3.6	3.7	NA	NA	Custer et al. 2010
<i>Ardea herodias</i> *	Minnesota (USA)	2010-2011	0.65	342	8	0.6	2.55	22	12.9	NA	NA	Custer et al. 2013
<i>Phalacrocorax auritus</i>	San Francisco (USA)	2009	LOD-25.2	483.7	NA	ND-24.3	13.4	13.8	7.08	NA	NA	Sedlak and Greig 2012
<i>Phalacrocorax carbo</i> **	Sweden	2007-2009	2.5	552	2.06	4.05	20.7	44.8	23.9	23.7	4.08	Nordén et al.2013
<i>Phalacrocorax carbo</i> **	Germany	2009	2.8	400	NA	1.1	2.7	10.4	1.0	NA	NA	Rüdel et al., 2011
<i>Parus major</i> ***	Belgium	2006	NA	19 – 5635	NA	NA	NA	NA	NA	NA	NA	Lopez Antia et al., 2017
<i>Vanellus Vanellus</i> ***	Belgium	2006	NA	143 - 46182	NA	NA	NA	NA	NA	NA	NA	Lopez Antia et al., 2017
<i>Ichthyaetus melanocephalus</i> ***	Belgium	2006	NA	150 – 916	NA	NA	NA	NA	NA	NA	NA	Lopez Antia et al., 2017

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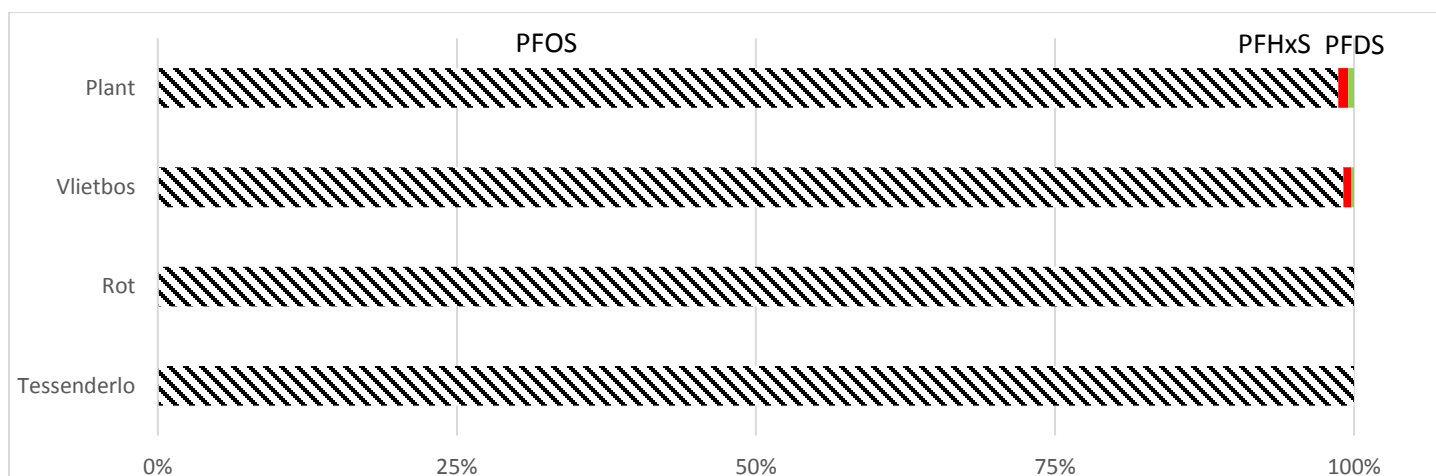
571 **Appendix**

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573 *Table S1. Results obtained for spiked egg samples in the inter-laboratory study between SGS, University of Antwerp and the*
 574 *University of Amsterdam.*

Spiked isomers	Trial	SGS		University of Antwerp		University of Amsterdam		theoretical values	
		PFOA (ng/g)	PFOS (ng/g)	PFOA (ng/g)	PFOS (ng/g)	PFOA (ng/g)	PFOS (ng/g)	PFOA (ng/g)	PFOS (ng/g)
Linear	1	55.6	59	57.0	57.4	56	66	61.7	63.2
	2	50.1	62.8	60.3	57.1	57	67		
	3	53.7	59.2	61.4	60.9	55	68		
Branched	1	28.5	30.1	34.5	34.9	28	30	32.2	32
	2	28.4	29.5	32.1	33.5	26	28		
	3	29.1	30.5	33.4	34.8	28	29		

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576 *Figure S1. Composition profile of Σ PFSA in eggs of great tit at the four sampling sites; a fluorochemical plant and at three sites*
 577 *with increasing distance from the plant site: 1 km (Vlietbos), 2.3 km (Rot) and 70 km (Tessenderlo, reference site).*
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