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# 1 High levels of fluoroalkyl substances and potential disruption of thyroid

## 2 hormones in three gull species from South Western France

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#### 14 Abstract

Per- and Poly-fluoroalkyl substances (PFAS) raised increasing concerns over the past years due to 15 their persistence and global distribution. Understanding their occurrence in the environment and 16 17 their disruptive effect on the physiology of humans and wildlife remains a major challenge in ecotoxicological studies. Here, we investigate the occurrence of several carboxylic and sulfonic 18 PFAS in 105 individuals of three seabird species (27 Great black-backed gull Larus marinus; 44 19 20 Lesser black-backed gull Larus fuscus graellsii; and 34 European herring gull Larus argentatus) from South western France. We further estimated the relationship between plasma 21 concentrations of PFAS and i) the body condition of the birds and ii) plasma concentrations of 22 23 thyroid hormone triiodothyronine (TT3). We found that great and lesser black-backed gulls from South Western France are exposed to PFAS levels comparable to highly contaminated species 24 25 from other geographical areas, although major emission sources (i.e. related to industrial 26 activities) are absent in the region. We additionally found that PFAS are negatively associated with the body condition of the birds in two of the studied species, and that these results are sex-27 28 dependent. Finally, we found positive associations between exposure to PFAS and TT3 in the great 29 black-backed gull, suggesting a potential disrupting mechanism of PFAS exposure. Although only 30 three years of data have been collected, we investigated PFAS trend over the study period, and found that great black-backed gulls document an increasing trend of plasma PFAS concentration 31 from 2016 to 2018. Because PFAS might have detrimental effects on birds, French seabird 32 populations should be monitored since an increase of PFAS exposure may impact on population 33 viability both in the short- and long-term. 34

#### 35 Introduction

Per- and Poly-fluoroalkyl substances (PFAS) raised increasing concerns over the past years due to 36 their persistence and global distribution. Because of their high thermal and chemical stability, 37 38 these synthetic substances have found an application in the manufacturing industry, mostly used 39 as surfactants and additives (Buck et al., 2011), and have been widely produced over the past 50 40 years (Wang et al., 2017). Being extremely persistent in the environment, and due to their long-41 range transport via atmospheric and oceanic currents, they have been detected worldwide (Giesy and Kannan, 2001). Several studies have found PFAS to accumulate into living organisms (including 42 invertebrates, fishes, amphibians, mammals, and birds) and to biomagnify through food webs 43 44 (Kannan et al., 2005; Kelly et al., 2009; Simonnet-Laprade et al., 2019), and to date, PFAS exposure represents a global threat to human health and wildlife (Sunderland et al., 2019). Documenting 45 46 their occurrence in the environment and understanding their disruptive effect on the physiology of 47 humans and wildlife remains a major challenge.

48 Seabirds are long-lived apex predators generally exposed to high levels of environmental 49 contaminants (Elliott and Elliott, 2013; Furness and Camphuysen, 1997), thus they prove particularly valuable to investigate PFAS accumulation in marine food webs especially in northern 50 areas. High levels of PFOS have been found in plasma samples of several seabird species from the 51 Arctic including ivory gulls Pagophila eburnea (average concentration of 31 ng/g, Lucia et al., 52 2017); glaucous gulls Larus hyperboreus (average concentration of 47 ng/g, Melnes et al., 2017; 53 and of 134 ng/g, Verreault et al., 2005); in black-legged kittiwakes from Svalbard (average 54 concentration of 10.2 ng/g, Tartu et al., 2014); and in European shags Phalacrocorax aristotelis 55 from Isle of May in Scotland (average concentration of 251 ng/g in females and 163 ng/g in males, 56 57 Carravieri et al. 2020). High levels of PFOS were also found in egg samples of the European shag 58 and common eider Somateria mollissima from Norway (average concentration of 36.8 ng/g and

59 37.4 ng/g, respectively, Herzke et al., 2009); in whole blood of the endangered lesser black backed gull Larus fuscus from Norway (average concentration of 33.5 ng/g, Bustnes et al., 2008a); and in 60 several other seabird species. However, much work has been devoted to seabirds from the Arctic 61 62 regions (i.e. considered a sink for environmental contaminants; Barrie et al., 1992; Braune et al., 63 2014; Wong et al., 2018), or in highly contaminated areas (e.g. China; Xie et al., 2013), while fewer 64 studies have focused on areas with not-known sources of PFAS (i.e. Antarctica, Munoz et al., 2017b). In France, most studies examining PFAS occurrence and exposure in aquatic ecosystems 65 66 focused on water, sediments, invertebrates, and fishes (Couderc et al., 2015; Fernandes et al., 67 2018; Munoz et al., 2019; Simonnet-Laprade et al., 2019). However, to the extent of our 68 knowledge, no studies have been carried out on top predators including birds in this area, which 69 may be exposed to concentrations of concern. It is therefore crucial to investigate PFAS exposure 70 in French seabirds to document PFAS occurrence in marine biota and to provide early warning of its effects on their health status. 71

Over the past few years, there has been an increased body of evidence showing that PFAS 72 73 may i) impact on adipogenesis thus with body condition (Tartu et al., 2014), and ii) disrupt several 74 physiological traits of seabirds. For instance, previous work found that PFAS exposure is associated 75 with lower levels of the stress hormone corticosterone (Tartu et al., 2014), higher oxidative stress 76 (Costantini et al., 2019), longer telomeres (Blévin et al., 2017a; Sebastiano et al., 2020), and a higher metabolic rate (Blévin et al., 2017b). Further studies found PFAS to be associated with 77 higher levels of the parental hormone prolactin and altered incubation behaviours (Blévin et al., 78 79 2020), lower hatching success (Tartu et al., 2014), and a higher survival rate (Sebastiano et al., 80 2020). Specifically, one way through which PFAS may impact on organism function is by disrupting 81 hormonal mechanisms. Previous work provided evidence that PFAS have a strong affinity for 82 proteins and are known to bind to the thyroid hormone transport protein transthyretin (Ren et al.,

83 2016; Weiss et al., 2009). In birds, the hypothalamic-pituitary-thyroid (HPT) axis controls the 84 secretion of the thyroid hormone thyroxine (T4), which is then converted to triiodothyronine (T3), the active form of T4 (McNabb, 2007). Although Blévin et al. (2017b) found no association 85 between PFAS exposure and thyroid hormones in adult black-legged kittiwakes Rissa tridactyla, 86 Braune et al. (2011) found a significant positive correlation between total triiodothyronine (TT3) 87 88 levels and hepatic concentrations of PFAS in northern fulmars. Nøst et al. (2012) also found a positive association between PFAS levels and total thyroxine (TT4) in black-legged kittiwake and 89 90 northern fulmar chicks, suggesting that PFAS may potentially act through an endocrine disrupting mechanism. More recently, Melnes et al. (2017) found that PFAS were positively associated with 91 free triiodothyronine (FT3) in the glaucous gull. To date, further work is needed to understand the 92 93 relationship between PFAS exposure and thyroid functioning in birds, especially considering that in 94 birds, T3 and T4 are involved in a multitude of physiological pathways (McNabb, 2007). A disruption of thyroid hormone levels may be detrimental to development, behaviour, and 95 reproduction (McNabb, 2007). 96

97 The Lilleau des Niges Natural Reserve is an important site for breeding, wintering, and migration of several bird species. It is located north of Ile de Ré, an island off the west coast of 98 99 France, in front of La Rochelle, in the Bay of Biscay. By hosting several seabird species during the 100 breeding season, this island offers a unique opportunity to investigate the occurrence of PFAS in a 101 French seabird community. Although most previous studies have been carried out on a single species (Blévin et al., 2017b; Costantini et al., 2019; Melnes et al., 2017; Tartu et al., 2014), 102 investigating several species simultaneously and from the same geographical area can help to 103 104 better understand the mechanisms of exposure to PFAS and the potential physiological 105 consequences of PFAS contamination. For instance, the Herring gull Larus argentatus, the lesser black-backed gull Larus fuscus graellsii, and the great black-backed gull Larus marinus, which 106

107 breed sympatrically on the island, are characterized by different foraging and migratory strategies, thus potentially exposed to different concentrations of PFAS. The aims of this study were to i) 108 assess to which extent French seabirds are contaminated by PFAS; ii) investigate the relationship 109 between exposure to PFAS and body condition; and iii) determine the association between 110 111 exposure to PFAS and plasma thyroid hormone T3 concentration in the three above mentioned 112 seabirds from Ile de Ré. Data on PFAS occurrence and their potential adverse effects in seabirds 113 from France are not yet available. To date, we are not aware of known point sources of PFAS in 114 the region. But considering that diverse important rivers may discharge PFAS near the study area (Simonnet-Laprade et al., 2019), and that PFAS may reach and accumulate in remote areas due to 115 their long-range oceanic and atmospheric transport (Munoz et al., 2019), we expect comparable 116 117 PFAS concentrations with seabirds from the Arctic. In addition, if PFAS have a stimulating effect on 118 thyroid hormone production as found in previous work (DeWitt, 2015; Liu et al., 2011; Nøst et al., 2012), we expect a positive association between PFAS and the concentration of thyroid hormones. 119 Furthermore, although some PFAS are listed as POPs by the Stockholm Convention and their 120 121 production has subsequently been reduced over the past years, studies investigating temporal 122 trends of PFAS in tissues of wildlife are limited. Although our data have been solely collected over 123 three years of study, we further aim to describe the temporal variation in blood concentration of 124 PFAS from 2016 to 2018 in local seabirds.

#### 125 Materials and Methods

#### 126 Sampling

Field work was performed in 2016, 2017, and 2018 at the Lilleau des Niges Natural Reserve (46° 127 13' 53" N, -1° 30' 22" W), managed by the Ligue pour la Protection des Oiseaux (LPO) located on 128 the North side of Ile de Ré, France, as a part of a monitoring program for PFAS in the region. A 129 total of 108 breeding adult birds from three species were captured during the incubation stage on 130 131 their nests using a nest trap. Because out of the 108 observations, three were coming from the 132 same individuals sampled at different years, one or the other observation was randomly excluded to perform statistical analyses. Therefore, the final dataset included a total of 105 birds (European 133 herring gull, n=9 in 2016, n=16 in 2017, and n=9 in 2018; lesser black-backed gull, n=11 in 2016, 134 135 n=17 in 2017, and n=16 in 2018; great black-backed gull, n=9 in 2016, n=7 in 2017, and n=11 in 2018). After capture, 2mL of blood was collected from the alar vein using a heparinized syringe 136 137 and a 25 gauge needle. Blood was kept in a cold container and centrifuged for 10 min at 8,000 x g at 20 °C at the laboratory within a few hours after collection; plasma and red blood cells were kept 138 139 frozen at -20 °C until laboratory analyses. Skull and tarsus were measured with an accuracy of 0.1 mm using a caliper. Wing length was also measured with an accuracy of 1 mm using a ruler, and 140 141 birds were weighted to the nearest 5 g using a Pesola spring balance. Birds were sexed from red blood cells by polymerase chain reaction amplification (PCR) of part of two highly conserved genes 142 143 (CHD) of sexual chromosomes. Briefly, DNA was extracted from erythrocytes and the sex was 144 determined by molecular sexing based on polymerase chain reaction (PCR) amplification of the CHD gene as described in Fridolfsson and Ellegren (1999). Amplification was performed in 20µl 145 final volume with a Eppendorf Mastercycler using 0.5 U Taq DNA polymerase, 200µM dNTPs, 146 10mM Tris-HCl pH 8.3, 50mM KCl, 1.5mM MgCl2 and 0.4 $\mu$ M of primers 2550F (5'-147 148 GTTACTGATTCGTCTACGAGA-3') and 2718R (5'-ATTGAAATGATCCAGTGCTTG-3'). Female birds may

149 deposit a significant amount of PFAS into their eggs. Therefore, to minimize the variation due to PFAS deposition in eggs, we have only sampled individuals having either two (17/105, 16% of 150 birds) or three eggs (87/105, 83% of birds) except one sampled females that only laid one egg 151 152 (1/105, 1%). Preliminary statistical analyses were carried out to test whether females with two 153 (8/52, 15% of females) or three eggs (43/52, 83%) had different concentrations of PFAS. However, 154 linear models showed that for any PFAS, concentrations were similar between females that laid 155 two or three eggs (all t<1.14, all P>0.26), thus clutch size was not further included in the statistical 156 analyses.

#### 157 **PFAS analyses**

A total of 14 PFAS were analysed in each plasma sample, including eight carboxylates: branched(Br-PFOA) and linear-perfluorooctanoate (L-PFOA), perfluorononanoate (PFNA),

160 perfluorodecanoate (PFDA), perfluoroundecanoate (PFUnDA), perfluorododecanoate (PFDoDA),

161 perfluorotridecanoate (PFTrDA), perfluorotetradecanoate (PFTeDA); and six sulfonates:

perfluorohexanesulfonate (PFHxS), branched- (Br-PFHpS) and linear-perfluoroheptasulfonate (L-162 PFHpS), branched perfluoroctanesulfonate (Br-PFOS), linear perfluoroctanesulfonate (L-PFOS), and 163 perfluorooctanesulfonamide (FOSA). Analytical standards of native PFAS along with a series of 10 164 <sup>13</sup>C, <sup>18</sup>O or D mass-labelled internal standards used for quantification purposes were supplied by 165 Wellington laboratories. All reagents were analytical grade or equivalent (see Munoz et al. (2017b) 166 167 for full details). Briefly, in a 2 mL polypropylene Eppendorf tubes, a 25 μL aliquot of plasma was 168 weighed (~25 mg) and internal standards (ISs) were subsequently added under gravimetric control ( $\sim$ 15 mg of a 1 pg/µL IS mixture prepared in methanol). Following protein precipitation 169 with 100 µL of acetonitrile (ACN), extracts were centrifuged for 10 min at 24,000 x g at 20 °C. The 170 171 supernatant was then transferred to 2 mL polypropylene centrifuge tubes (0.22 µm nylon filter).

172 After centrifugation for 3 min at 7,000 x g at 20 °C, extracts were transferred to 2 mL auto sampler

173 glass vials and diluted with 675 µL of HPLC-water. Extracts were briefly vortexed and then processed using an Agilent Technologies (Massy, France) on-line SPE platform which comprises a 174 standard auto sampler (1260 Infinity ALS), a quaternary pump (1260 Infinity Quaternary Pump VL), 175 176 a switch valve (1200 2 Position/6 Port Valve) and an on-line SPE column support (1200 6 Position 177 Selection Valve), which were all automatically controlled via the Acquisition module of the Agilent 178 Mass Hunter software as previously done (Munoz et al., 2017b). HPLC-water aliquots were run 179 between each seabird plasma sample to eliminate any cross-contamination. Note that on-line 180 extraction was performed with Waters Oasis HLB on-Line SPE columns (2  $\times$  10 mm, dp= 25-35  $\mu$ m) while analyte separation was carried out using an Agilent  $C_{18}$  Poroshell analytical column (2.1 × 181 182 100 mm, 2.7 μm).

#### 183 **Quality assurance/Quality control (QA/QC)**

When analytes were detected in blanks, blank correction was performed and a limit of reporting 184 185 (LOR) was defined as three times the maximum blank signal divided by the average mass of plasma used for analysis. A limit of detection (LOD) was also defined as the concentration yielding 186 a signal to noise ratio of 3 in spiked plasma samples. Because laboratory analyses were performed 187 in different years, a unique left-censoring threshold was set for each analyte, i.e. the maximum 188 between LORs and LODs, all years combined. For those PFAS with concentrations below this 189 threshold in less than 30% of samples, left-censored data were arbitrarily replaced with ½ x LOR or 190 191 LOD to enable statistical analyses. Therefore, ten PFAS (PFNA, PFDA, PFUnDA, PFDoDA, PFTrDA, 192 PFTeDA, PFHxS, L-PFHpS, Br-PFOS, and L-PFOS) could be further investigated (i.e. other analytes were excluded from statistical analyses). LORs, LODs and detection frequencies are presented in 193 the supplementary information (Table S1). For each sample batch (20 samples), several QA/QC 194 195 points were assessed by analyzing: i) two procedural blanks consisting of 25 µL of HPLC-water that 196 went through the entire analytical procedure; ii) one human serum standard reference material

197 (NIST SRM 1957, trueness assessment); iii) replicate spiked chicken plasma samples (target analytes added jointly with mass-labelled ISs at the beginning of the preparation procedure at 2 198 ng/g each, accuracy assessment); and iv) HPLC-water samples spiked at 2 ng/g, accuracy 199 assessment) as previously described (Munoz et al. 2017b). Procedural blanks showed very limited 200 201 contamination. The analysis of NIST SRM 1957 gave satisfactory results, i.e. within the specified 202 uncertainty interval. For those compounds with a reference concentration (i.e. PFHpA, PFOA, PFNA, PFDA, PFUnDA, PFHxS and PFOS), levels deviated between 2 and 20% from the reference 203 204 concentration (except for FOSA, which deviated 36%).

#### 205 Thyroid hormone analyses

206 TT3 was determined by radioimmunoassay. Briefly, 25 µL of plasma was incubated for 24h at 4 °C 207 with a known concentration (10000 cpm) of T3 marked with the radioisotope lodine-125 (T3-<sup>125</sup>I, 208 Perkin Elmer, US, reference: NEX110X100UC) and an antibody Ab (polyclonal rabbit antiserum, 209 Sigma-Aldrich, US, reference: T-2777). Because *Ab* is available in a limited concentration, T3 and 210 T3-<sup>125</sup>I compete for *Ab*, to which they bind. Therefore, after incubation, there is a bound fraction (T3 and T3-<sup>125</sup>I bound to Ab) and a free fraction (T3 and T3-<sup>125</sup>I unbound to Ab), which are 211 separated by adding a sheep anti-rabbit antibody (whole anti-serum anti rabbit IgG produced in 212 213 sheep), incubated for 12h at 4 °C followed by centrifugation at 4,300 x g at 18-20°C for 45 min. The bound fraction is then counted with a wizard 2 gamma counter (Perkin Elmer, US). Pooled plasma 214 215 of diverse gull samples were serially diluted and produced a dose-response curve parallel to the T3 216 standard curve. The lowest TT3 detectable concentration was 0.07 ng/ml (LOD). Samples below this limit (n=3) were replaced with a value equal to ½ x LOD to enable statistical analyses. All 217 samples were run in duplicates. Samples that had a coefficient of variation above 15% and could 218 219 not be done in triplicates due to low plasma volume were not included in statistical analyses 220 (n=11). An additional measurement of TT3 was excluded from statistical analyses since it was

considered an outlier (the measurement exceeded the mean ± 3 times the standard deviation and
was highly influential in statistical analyses). Therefore, for a total of 58 samples (n=6 in 2016 and
n=12 in 2017 for the European herring gull; n=10 in 2016 and n=14 in 2017 in lesser black-backed
gull; n=8 in 2016 and n=8 in 2017 in great black-backed gull), both TT3 and PFAS data were
available. The intra-assay coefficient of variation was 9.73%, while the inter-assay coefficient of
variation amounted to 15.13%.

#### 227 Statistical analyses

228 After PFAS data were log-transformed to reduce the influence of extreme values (see below), linear models were used to investigate differences in PFAS concentrations among species and 229 230 between genders (in samples collected from 2016 to 2018). In each model, each PFAS was 231 considered as a dependent variable while the factors Species, Sex, and their interaction were 232 considered as predictors. To simultaneously investigate species specific temporal trends in PFAS 233 exposure from 2016 to 2018, the Year, the factor Species, and their interaction were also included 234 in the model as explanatory variables. We used a similar model to test the difference in body condition between sexes. Briefly, the body condition has been calculated using the body mass 235 236 adjusted by a linear body measurement (i.e. skull length) using the formula described in Peig and Green (2009). 237

Linear models were additionally used to study the association between TT3 and PFAS (in samples collected in 2016 and 2017). In these models, a three-way interaction between *PFAS*, the factors *Species* and *Sex*, was used to investigate sex-related responses to PFAS exposure. These models additionally included the *Year* (as a factor, to control for the temporal variation in TT3 and PFAS), and *body condition* (as a covariate, to control for the individual condition of birds). A similar model

was built to test the association between the body condition and PFAS in all samples collectedfrom 2016 to 2018.

245 All PFAS concentrations (except PFTeDA, which was normally distributed and assumptions listed below were respected without data transformation), were log-transformed when testing for time 246 trends and when testing for inter-species and between sex differences. All PFAS concentrations 247 were log-transformed when testing for the association between either TT3 or body condition and 248 249 PFAS. Data transformation was done to meet model assumptions as homoscedasticity and 250 normality of residuals, further confirmed by visually inspecting Q-Q plots. All data transformation and violation of models' assumptions are reported throughout the manuscript. Statistical 251 significance was set to  $\alpha$ =0.05 and 95% confidence intervals were used during data processing and 252 data visualization. All statistical analyses were performed using R version 3.5.2. 253

#### 255 Results

PFAS used in statistical analyses were detected in all samples (Table S1) and their concentrations 256 are summarized in Table 1 and Figure 1. PFOS was the most abundant, followed by the odd-chain 257 258 carboxylates PFTrDA and PFUnDA. Linear models showed statistically significant differences among species for all carboxylic and sulfonic PFAS (all F>5.15, all P<0.01), and all statistical outputs 259 and post-hoc differences can be found in Table S2. Among carboxylates, PFNA levels were higher 260 261 in herring gulls and lesser black-backed gulls than great black-backed gulls (both P<0.05, Figure 1), PFDA levels were higher in great black-backed gulls than herring gulls (P<0.01, Figure 1), PFUnA, 262 PFDoDA, PFTrDA, and PFTeDA levels were higher in great and lesser black-backed gulls than 263 264 herring gulls (all P<0.01, Figure 1), and PFTrDA levels were also higher in great than lesser blackbacked gulls (P<0.05, Figure 1). Among sulfonic acids, PFHxS was higher in both lesser black-265 266 backed and herring gulls than great black-backed gulls (both P<0.01, Figure 1), PFHpS and L-PFOS 267 were highest in lesser black-backed gulls (all P<0.001, Figure 1), and Br-PFOS was higher in lesser than great black-backed gulls (P<0.05, Figure 1). 268

All carboxylates showed significantly higher concentrations in males than in females for all species
(all P<0.05, except for PFNA in the herring gull, for which P=0.07; Table 1). Among sulfonates,</li>
PFHxS showed similar concentrations between females and males in all species (all P>0.16, Table
1), while PFHpS, Br- and L-PFOS showed significantly higher concentrations in males than females
(all P<0.05, Table 1). In all three species, there was no difference in the body condition between</li>
males and females (all t<1.27, all P>0.80). Finally, TT3 levels were similar between sexes in all
three species (t=2.76, P=0.08).

In great black backed gulls, TT3 was positively associated with PFUnDA, PFDoDA, PFTrDA, PFTeDA
 and Br-PFOS in females (all t> 2.10, all P≤0.04; Figure 2a-d, 2f, Table S3), while TT3 was negatively

associated with PFHxS in males (t=-2.69, P=0.01; Figure 2e, Table S3). There was no association
between TT3 and any PFAS in herring gulls and lesser black-backed gulls (all t≤0.79, all P≥0.43,
Table S3).

281 In great black backed gull females, increasing levels of PFNA, PFDA, PFHxS, PFHpS, Br- and L-PFOS

were associated with a reduced body condition (all t $\leq$ -2.19, all p $\leq$ 0.03; Figure S1, S2, Table S4),

283 while increasing levels of PFNA and PFDA were associated with a reduced body condition (both t≤-

284 2.18, both p=0.03; Figure S1, Table S4) in lesser black-backed gull males, but not females.

285 In great black backed gulls, there was a significant or marginally-significant increase in all PFAS

from 2016 to 2018 (all t≥1.93 and all P≤0.056; Figure 3 and 4, Table S5), while lesser black backed

287 gulls showed an increase in PFTeDA and PFHxS, and a marginally significant decrease in PFUnDA

288 levels from 2016 to 2018 (all t≥1.98, all P≤0.05; Figure 3, 4, Table S5). All PFAS showed similar

concentrations among years in the herring gulls except for PFTrDA, which increased from 2016 to

290 2018 (t=2.30, P=0.02; Figure 3, Table S5). Body condition did not change in any of the species from

one year to another (all t<1.14, all P>0.26). TT3 levels remained similar between 2016 and 2017 in

all three species (all t<1.88, all P>0.42).

#### 294 Discussion

Our study is the first to provide evidence that although not-known point sources of emission are 295 296 present in the region, several PFAS were detected in seabird species from South Western France. 297 Great and lesser black-backed gulls show that both plasma carboxylate and sulfonate 298 concentrations are comparable to highly contaminated seabird species from Arctic regions, while 299 herring gulls are exposed to relatively lower levels of PFAS. We found that PFAS are negatively 300 associated with the body condition of the birds. Furthermore, TT3 levels were associated with 301 several PFAS in a contrasted manner between sexes in the great black-backed gull, suggesting a 302 potential disrupting mechanism of PFAS exposure. Finally, the great black-backed gulls 303 documented an increasing trend of plasma PFAS concentration from 2016 to 2018.

304 Our results show that among carboxylates, PFUnDA and PFTrDA are the most abundant 305 congener in all three species, a pattern that is commonly found in seabird species (Bustnes et al., 306 2008b; Melnes et al., 2017; Tartu et al., 2014). Because of the strong winds and oceanic currents 307 that characterize the Atlantic Ocean, and considering that, to the best of our knowledge, there are no point sources of PFAS in the region, perfluorinated compounds should occur at a lower 308 concentration than in the Mediterranean and the Arctic regions, considered as sinks for pollutants 309 310 (Danovaro, 2003; Wong et al., 2018). However, the levels of carboxylates found in this study (ranging from a median of 0.6 ng/g of PFTeDA in Herring gulls to 5.8 ng/g of PFTrDA in great black-311 312 backed gulls) are similar to those reported for glaucous gulls (ranging from a median of 0.1 ng/g of 313 PFOA to 3.8 ng/g of PFUnDA, Melnes et al., 2017) and lesser black-backed gulls (ranging from a median of 0.2 ng/g of PFDoDA to 5.9 ng/g of PFDA, Bustnes et al., 2008a) from Arctic regions, but 314 lower than those found in other species (e.g. in black-legged kittiwakes from Svalbard; ranging 315 316 from a mean of 1.0 ng/g of PFNA to 18.2 ng/g of PFTrDA, Tartu et al., 2014). One possible 317 explanation may be related to the continental input of PFAS through the Gironde, Loire, and

318 Charente estuaries (Munoz et al., 2019; Munoz et al., 2017a; Munschy et al., 2019), which may have contributed to the observed concentrations. Except for PFNA (which levels were higher in 319 herring gulls and lesser black-backed gulls than great black-backed gulls), most carboxylates were 320 higher in lesser and great black backed gulls than herring gulls. Although we cannot exclude that 321 322 these three species differ in their ability to excrete PFAS from their body, our results suggest that 323 the differences in exposure likely depend on the trophic niche occupied by the species. For 324 instance, a recent study on PFAS in six seabird species from the Arctic regions showed that 325 predatory birds (e.g. Great skua Stercorarius skua, 44.8 ng/g of sumPFAS) showed the highest contaminant load compared with species from a lower trophic level (e.g. Common eider Somateria 326 mollissima, 1.3 ng/g of sumPFAS; Haarr et al. 2018). Great black-backed gulls feed on higher 327 trophic level preys and mainly forage along the shore (Maynard and Davoren, 2020), while lesser 328 329 black-backed and herring gulls are known to have a generalist diet which also includes food items from both terrestrial and marine origin (Corman et al., 2016; Maynard and Davoren, 2020). 330

331 Among sulfonates, L-PFOS was the most abundant, followed by Br-PFOS, PFHxS, and PFHpS 332 and all occurred at very high concentrations. For instance, PFHxS ranged from a median value of 1.2 ng/g in great black-backed gulls to 2.2 ng/g in lesser black-backed gulls, while other studies 333 from highly contaminated areas reported lower PFHxS plasma concentrations (a median below 1 334 ng/g in lesser black-backed gulls, Bustnes et al., 2008b; a median below 0.7 in glaucous gulls, 335 Melnes et al., 2017; all samples below 0.2 ng/g in black-legged kittiwakes, Tartu et al., 2014). 336 Furthermore, L-PFOS ranged from a median of 11.6 ng/g in herring gull females to a median of 337 54.7 ng/g in lesser black-backed gull males. Thus, lesser black-backed gulls in our study showed 338 339 very high sulfonate levels, even higher than Norwegian populations (a median of 40 ng/g of PFOS 340 in males, Bustnes et al., 2008b), and in this species, sulfonate levels are significantly higher than those observed in great black-backed gulls. In addition, plasma concentrations of most PFAS 341

342 showed significantly higher levels in males than in females, but this difference was not related to the body condition of the birds, with males showing a similar body condition than females. 343 Because females transfer contaminants in the eggs, it is thus possible that females have lower 344 345 levels of circulating PFAS in plasma. However, previous work pointed out contrasting results between PFAS in eggs and plasma, suggesting that the extent of PFAS transfer to the eggs may 346 347 significantly vary among the studied species (Bustnes et al., 2008a; Herzke et al., 2009; Verreault 348 et al., 2005). Given that in this study we did not analyse PFAS levels in eggs, it is not possible to 349 clarify whether the species differ in terms of PFAS excreted in eggs. Although Verreault et al. (2006) found that the contaminant content in glaucous gull eggs fluctuated irrespectively of the 350 laying order, other work suggests that the majority of PFAS are found in the first or the first two 351 352 eggs, while negligible concentrations of PFAS are found in the third egg, as previously shown in 353 Audouin's gulls Larus audouinii (Vicente et al., 2015). Our results showed that females that laid three eggs had similar PFAS concentrations than females that laid two eggs, therefore our results 354 should not be affected by the difference in PFAS deposition in eggs. 355

356 Because of the higher bioaccumulative properties and biomagnification and of longer-chain PFAS (Boisvert et al., 2019; Simonnet-Laprade et al., 2019), these compounds tend to occur at 357 higher concentrations in wildlife tissues (Conder et al., 2008; Muir et al., 2019; Muir and de Wit, 358 2010). Therefore, individuals feeding at a higher trophic position are likely to be exposed to higher 359 concentrations of long-chained carboxylates. Previous work also showed that longer chained PFAS 360 are more likely to induce adverse health effects in seabirds compared to shorter chained PFAS. For 361 instance, negative associations between PFAS and baseline corticosterone in black-legged 362 363 kittiwakes were only found for PFTrDA and PFTeDA (Tartu et al., 2014), while such association was 364 not found for shorter-chain PFAS. Additional work on the same bird population found higher 365 protein oxidative damage in those birds having higher concentrations of PFDoDA, PFTrDA and

PFTeDA (Costantini et al., 2019). Similarly, previous work found a positive association between
 PFTrDA and metabolic rate in the same species (Blévin et al., 2017b). These results were further
 corroborated by experimental work on rat *Rattus sp.* cell cultures, showing that the cytotoxicity of
 PFAS increases with increasing carbon chain length (Berntsen et al., 2017), and that comparing
 molecules with a similar chain length, a sulfonate functional group may lead to greater toxicity
 than a carboxyl group (Berntsen et al., 2017).

372 Our results are thus of particular interest as they suggest that birds feeding at a higher trophic position and showing a more marine diet (i.e. great black-backed gulls) should also be 373 exposed to greater toxicological risks from carboxylates, while birds with a more generalist diet 374 (i.e. lesser black-backed gulls) can be exposed to higher levels of sulfonates. This is likely the 375 reason why we found associations between TT3 and PFAS only in great black-backed gulls, while 376 377 no significant associations have been found in herring gulls and lesser black-backed gulls. 378 Interestingly, our results were dependent on the sex of the birds. Indeed, although we found that PFHxS was associated with TT3 in great black-backed gull males, all other associations were found 379 380 in females only. Despite the lower concentration of circulating plasma PFAS levels, females can deposit PFAS in eggs therefore we cannot be certain that females were exposed to lower PFAS 381 382 concentrations than males, and further work including other tissues (e.g. liver or muscle) would clarify whether PFAS intake differs between the sexes of sampled birds. Despite the absolute 383 concentrations to which they are exposed, a possible explanation for the results in females may 384 rely on the fact that incubation can be extremely costly for female birds (Hanssen et al., 2005), 385 thus they may be more susceptible to PFAS exposure. Previous work in birds suggest a modulation 386 387 of thyroid function induced by exposure to various environmental contaminants. Smits et al. 388 (2002) reported decreased TT3 levels in American kestrels Falco sparverius experimentally exposed to PCBs, while Verreault et al. (2004) reported a decrease in T4:T3 ratio in the glaucous 389

390 gull. Similarly, exposure to organochlorines was associated with reduced TT3 in kittiwakes (Blévin et al., 2017b), and with reduced T3 and T4 in glaucous gulls (Melnes et al., 2017; Verreault et al., 391 2004). However, specifically related to PFAS, further work on seabirds found a positive association 392 with thyroid functioning (i.e. between PFOS and FT3 in glaucous gulls, Melnes et al., 2017; 393 between several PFAS and TT4 in black-legged kittiwakes and northern fulmars, Nøst et al., 2012). 394 395 Although being conducted in fish, an experimental approach showed that exposure to PFOS in zebra fish (Danio rerio) led to increased thyroid hormones secretion (Liu et al., 2011). Thus, our 396 397 results on great black backed gulls are in line with previous studies. In this study, increasing TT3 levels in this species were found with increasing concentrations of longer chain PFAS (PFUnDA, 398 PFDoDA, PFTrDA, and PFTeDA) and Br-PFOS. This suggests that despite carboxylates and 399 400 sulfonates are functionally different, their effect on TT3 is similar. However, this was not the case 401 for PFHxS, which showed a decrease in TT3 levels with increasing concentrations. PFHxS is highly toxic and causes thyroid disruption by lowering thyroid hormone levels in rats (Ramhøj et al., 402 2020), but it remains unclear why this effect was only found in great black-backed gull males. 403 404 Indeed, females showed similar levels than males, and the other species exhibited higher PFHxS 405 concentrations than those found in great black backed gulls, thus this result would strongly benefit 406 from experimental support. Because the avian thyroid gland secretes almost exclusively T4 (Darras et al., 2006), most T3 is derived from the deiodination of T4 (Darras et al., 2006). A possible 407 408 explanation is that in great black-backed gulls, exposure to PFHxS may negatively impact either the transport of T4 (by reducing the activity of serum binding proteins) or deiodination processes. It is 409 410 therefore strongly warranted to supplement in vitro experiments to verify the effect of PFHxS on 411 T4 transformation. Our results do not provide evidence for a causal relationship PFAS exposure 412 and circulating thyroid hormones. But the contrasting results found between sexes strongly call for

further work to experimentally investigate the effect of PFAS exposure on thyroid functioning ofbirds.

Sex-related differences were also found while investigating the relationship between 415 exposure to PFAS and the birds' body condition. In female great black-backed gulls, we found that 416 increasing concentrations of PFNA, PFDA, and all four sulfonates were negatively associated with 417 418 body condition, while in lesser black-backed gulls, a similar negative relationship between PFNA, 419 PFDA, and body condition was only found in males. Previous work in humans and various animal models found that exposure to certain PFAS is suspected to disrupt fatty acids metabolism and 420 promote adipogenesis (Cheng et al., 2016; Wan et al., 2012; Xu et al., 2016; Yeung et al., 2007). 421 422 More specifically, these changes in lipid content are related to the capacity of PFAS to alter the expression of genes involved in the metabolism of lipids and fatty acids (Jacobsen et al., 2018; 423 424 Wan et al., 2012). To date, work on the effect of PFAS on lipid metabolism and body condition in 425 birds remain extremely limited. The negative association we found is in contrast with a previous 426 study on black-legged kittiwakes, showing that PFNA was positively associated with body condition 427 in males (Tartu et al., 2014). One possible explanation for the negative relationship found in females may be related to the ability to deposit PFAS into the eggs. For instance, great black-428 backed gull females in a better body condition may be more efficient in eliminating PFAS through 429 430 egg-deposition, although evidences to support this statement are lacking. However, this would not 431 explain why in a closely related species (i.e. lesser black-backed gull) a similar association has been found in males. Our study results are novel but emphasize the need to experimentally investigate 432 433 the potential association between exposure to PFAS and body condition in birds.

Finally, not only does our work provide evidence of high PFAS levels in seabirds from
metropolitan France, but our results clearly suggest increasing blood concentration of most PFAS
over a relatively short period of time (i.e. from 2016 to 2018) in great and lesser black-backed

437 gulls. A previous study showed that birds caught later on over the breeding season had lower 438 concentration of PFAS (Bustnes et al., 2008a). However, all birds included in this study were sampled during the same period of the year (difference of a few days from one year to another), 439 and, more importantly, all birds were sampled while incubating eggs, thus during the same 440 reproductive state, which should not affect the results. Additionally, we cannot exclude that we 441 442 unintentionally captured older birds in more recent years (assuming that PFAS levels increase with age in these species). However, considering that all individuals included in this study were adults, 443 444 and assuming that PFAS concentrations in birds reach a steady level relatively early in life as previously shown for organochlorines (Bustnes et al., 2003), these trends should not be affected 445 by the age of the bird. Information on temporal trends of PFAS in birds' tissues in recent years are 446 447 scarce and do not exhibit any overall trend (Jouanneau et al., 2020; Land et al., 2018; Muir et al., 448 2019; Sun et al., 2019). Thus, although only three years of data could be included in the present study, our results provide valuable information on PFAS trends in South western France. Further 449 450 work including several years of study is strongly warranted to corroborate these findings.

451

#### 452 **Conclusions**

Our study provides the first evidence of the presence of high levels of PFAS in seabirds from South western France. Despite some PFAS showed similar levels of other seabird species, L-PFOS and some other PFAS showed either comparable or higher levels than highly contaminated seabird species, and may therefore pose a threat to long-lived seabirds. This hypothesis is further corroborated by our results showing an association between PFAS and the level of the thyroid hormone TT3. Similarly, we provide evidence that PFAS may interfere with lipid accumulation and body condition in birds, and we call for further work to experimentally test this hypothesis. Work

on species-specific mechanisms of contaminant excretion and susceptibility to PFAS exposure
would prove useful to understand the consequences of PFAS exposure in different species. Our
results also document an increase in blood PFAS concentrations over time, particularly in great
black-backed gulls, suggesting that PFAS concentrations may also be increasing in the investigated
species. Because PFAS have detrimental effects on birds, these and other seabird populations
should be monitored as an increase of PFAS exposure may impact on population viability both in
the short- and long-term.

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480 PFNA PFDA PFUNDA PFDoDA PFTrDA PFTeDA PFT

483 <0.01, and <0.001, respectively.



Species 💽 Herring gull 🔺 Lesser black-backed gull 🔳 Great black-backed gull



**Figure 2:** Relationship between the concentration of the thyroid hormone TT3 (expressed as ng/mL) and log-transformed carboxylic (C-11 to C-14, panel **a** to **d**, respectively), and sulfonic PFAS (PFHxS and Br-PFOS, panel **e** and **f**, respectively) of the three seabird species from IIe de Re. The solid line represents the trend while the grey area represents 95% confidence intervals. Shapes of data points and three different grades of grey are used to distinguish the three species as explained in the figure legend. Only significant trends are shown. Data refer to the period 2016-2017 for which both TT3 and PFAS were available (n=58).





492 Figure 3: Error bar plots mean ± standard error of carboxylate concentrations in 2016, 2017, and 2018 in
493 the seabird species from IIe de Re. The shapes and the three different grades of grey are used to distinguish
494 the three species as explained in the figure legend. Only significant trends are shown. Dashed lines indicate
495 a trend close to significance.



Figure 4: Error bar plots mean ± standard error of sulfonate concentrations in 2016, 2017, and 2018 in the
 seabird species from IIe de Re. The shapes and the three different grades of grey are used to distinguish the
 three species as explained in the figure legend. Only significant trends are shown. Dashed lines indicate a

502 trend close to significance.

Table 1: PFAS concentration (ng/g of ww) in females and males of the three seabird species from IIe de Re.
 P-values refer to the difference in PFAS between females and males. Significant P-values are bolded. SE

506 refers to standard errors.

Herring gull Larus argentatus								
	Females (n=18)		Males (n=16)					
	mean ± SE	median (range)	mean ± SE	median (range)	P-value			
PFNA	$1.16 \pm 0.11$	1.06 (0.36, 2.16)	$1.68 \pm 0.17$	1.63 (0.67, 2.85)	0.07			
PFDA	$1.16 \pm 0.19$	0.94 (0.38, 4.01)	$1.98 \pm 0.30$	1.79 (0.69, 6.04)	0.01			
PFUnDA	$1.23 \pm 0.17$	1.01 (0.47, 3.45)	$2.43 \pm 0.34$	2.17 (0.61, 6.49)	<0.001			
PFDoDA	0.95 ± 0.16	0.9 (0.3, 3.22)	$1.82 \pm 0.32$	1.56 (0.55, 6.17)	<0.001			
PFTrDA	1.76 ± 0.27	1.42 (0.55, 5.25)	3.03 ± 0.29	3.05 (1.04, 4.81)	<0.001			
PFTeDA	$0.61 \pm 0.07$	0.61 (0.25, 1.23)	$1.23 \pm 0.16$	1.03 (0.33, 2.8)	0.04			
PFHxS	2.26 ± 0.21	2.16 (0.88, 3.5)	2.29 ± 0.36	1.89 (0.79 <i>,</i> 6.92)	0.99			
PFHpS	0.27 ± 0.02	0.25 (0.11, 0.5)	$0.49 \pm 0.09$	0.38 (0.11, 1.63)	0.045			
Br-PFOS	2.03 ± 0.34	1.71 (0.86, 6.8)	3.95 ± 0.45	3.75 (1.07, 7.9)	<0.001			
L-PFOS	13.82 ± 2.0	11.64 (5.57, 37.04)	30.44 ± 4.31	28.44 (6.6, 62.52)	<0.001			

### Lesser black-backed gull Larus fuscus graellsii

	Females (n=20)		Males (n=24)				
PFNA	1.14 ± 0.09	1.11 (0.46, 1.83)	2.04 ± 0.1	2.06 (0.95, 3.36)	<0.001		
PFDA	$1.1 \pm 0.08$	1.15 (0.54, 1.64)	2.63 ± 0.15	2.58 (1.17, 4.16)	<0.001		
PFUnDA	$1.88 \pm 0.16$	1.69 (0.66, 3.28)	5 ± 0.29	4.89 (1.98, 8.7)	<0.001		
PFDoDA	$1.02 \pm 0.07$	1.02 (0.39, 1.87)	3.06 ± 0.2	2.86 (1.75, 5.52)	<0.001		
PFTrDA	$1.97 \pm 0.12$	2.06 (0.66, 2.91)	6.38 ± 0.44	5.74 (3.53, 11.66)	<0.001		
PFTeDA	0.66 ± 0.07	0.61 (0.13, 1.39)	2.23 ± 0.2	1.87 (1.1, 4.35)	<0.001		
PFHxS	$2.18 \pm 0.25$	1.95 (1.12, 6.01)	2.56 ± 0.26	2.24 (1.08, 5.56)	0.94		
PFHpS	0.37 ± 0.03	0.35 (0.16, 0.82)	0.79 ± 0.07	0.69 (0.34, 1.84)	<0.001		
Br-PFOS	$1.86 \pm 0.19$	1.96 (0.58, 3.15)	5.06 ± 0.5	4.41 (2.38, 13.67)	<0.001		
L-PFOS	21.09 ± 2.06	22.76 (5.74, 34.49)	60.23 ± 4.44	54.68 (26.51, 119.69)	<0.001		

# Great black-backed gull Larus marinus

	Females (n=14)		Males (n=13)				
PFNA	0.59 ± 0.06	0.59 (0.24, 0.95)	$1.94 \pm 0.29$	1.72 (0.89, 4.85)	<0.001		
PFDA	$1.41 \pm 0.23$	1.13 (0.32, 3.55)	3.83 ± 0.58	3.34 (1.52, 7.2)	<0.001		
PFUnDA	2.42 ± 0.35	2.23 (0.49, 4.93)	4.67 ± 0.65	4.08 (2.62, 11.46)	0.01		
PFDoDA	$1.69 \pm 0.24$	1.63 (0.32, 3.12)	2.79 ± 0.29	2.37 (1.68, 5)	0.04		
PFTrDA	4.04 ± 0.72	3.55 (0.68, 9.61)	6.87 ± 0.83	5.79 (3.85, 14.39)	0.02		
PFTeDA	0.96 ± 0.17	0.92 (0.13, 2.4)	1.94 ± 0.32	1.42 (0.91, 4.17)	0.03		
PFHxS	$1.2 \pm 0.11$	1.16 (0.62, 2.06)	2.01 ± 0.32	1.58 (0.93, 5.33)	0.16		
PFHpS	$0.2 \pm 0.03$	0.15 (0.09, 0.47)	0.63 ± 0.21	0.42 (0.22, 3.01)	<0.001		
Br-PFOS	$1.32 \pm 0.17$	1.1 (0.39 <i>,</i> 2.53)	4.21 ± 0.89	3.21 (1.85, 13.01)	<0.001		
L-PFOS	13.45 ± 1.52	10.53 (5.18, 25.92)	46.55 ± 13.74	27.62 (16.71, 194.72)	<0.001		

#### 508 References

- Barrie L, Gregor D, Hargrave BT, Lake R, Muir D, Shearer R, et al. Arctic Contaminants: Sources, Occurrence
   and Pathways. Science of the total environment 1992; 122: 1-74.
- Berntsen HF, Bjørklund CG, Audinot J-N, Hofer T, Verhaegen S, Lentzen E, et al. Time-dependent effects of
   perfluorinated compounds on viability in cerebellar granule neurons: Dependence on carbon chain
   length and functional group attached. NeuroToxicology 2017; 63: 70-83.
- Blévin P, Angelier F, Tartu S, Bustamante P, Herzke D, Moe B, et al. Perfluorinated substances and
   telomeres in an Arctic seabird: Cross-sectional and longitudinal approaches. Environmental
   Pollution 2017a; 230: 360-367.
- Blévin P, Shaffer SA, Bustamante P, Angelier F, Picard B, Herzke D, et al. Contaminants, prolactin and
   parental care in an Arctic seabird: Contrasted associations of perfluoroalkyl substances and
   organochlorine compounds with egg-turning behavior. General and Comparative Endocrinology
   2020; 291: 113420.
- Blévin P, Tartu S, Ellis HI, Chastel O, Bustamante P, Parenteau C, et al. Contaminants and energy
   expenditure in an Arctic seabird: Organochlorine pesticides and perfluoroalkyl substances are
   associated with metabolic rate in a contrasted manner. Environmental Research 2017b; 157: 118 126.
- Boisvert G, Sonne C, Rigét FF, Dietz R, Letcher RJ. Bioaccumulation and biomagnification of perfluoroalkyl
   acids and precursors in East Greenland polar bears and their ringed seal prey. Environmental
   Pollution 2019; 252: 1335-1343.
- Braune BM, Gaston AJ, Letcher RJ, Grant Gilchrist H, Mallory ML, Provencher JF. A geographical comparison
   of chlorinated, brominated and fluorinated compounds in seabirds breeding in the eastern
   Canadian Arctic. Environmental Research 2014; 134: 46-56.
- Braune BM, Trudeau S, Jeffrey DA, Mallory ML. Biomarker responses associated with halogenated organic
   contaminants in northern fulmars (*Fulmarus glacialis*) breeding in the Canadian Arctic.
   Environmental Pollution 2011; 159: 2891-2898.
- Buck RC, Franklin J, Berger U, Conder JM, Cousins IT, de Voogt P, et al. Perfluoroalkyl and polyfluoroalkyl
   substances in the environment: terminology, classification, and origins. Integrated environmental
   assessment and management 2011; 7: 513-541.
- Bustnes JO, Bakken V, Skaare JU, Erikstad KE. Age and accumulation of persistent organochlorines: A study
   of arctic-breeding glaucous gulls (*Larus hyperboreus*). Environmental Toxicology and Chemistry
   2003; 22: 2173-2179.
- Bustnes JO, Borgå K, Erikstad KE, Lorentsen SH, Herzke D. Perfluorinated, brominated, and chlorinated
   contaminants in a population of lesser black-backed gulls (*Larus fuscus*). Environmental Toxicology
   and Chemistry 2008a; 27: 1383-92.
- Bustnes JO, Erikstad KE, Lorentsen S-H, Herzke D. Perfluorinated and chlorinated pollutants as predictors of
   demographic parameters in an endangered seabird. Environmental Pollution 2008b; 156: 417-424.
- Carravieri A, Burthe SJ, de la Vega C, Yonehara Y, Daunt F, Newell MA, Jeffreys RM, Lawlor AJ, Hunt A, Shore
   RF, Pereira MG, Green JA. Interactions between Environmental Contaminants and Gastrointestinal
   Parasites: Novel Insights from an Integrative Approach in a Marine Predator. Environmental Science
   & Technology 2020; 54: 8938-8948.
- Cheng J, Lv S, Nie S, Liu J, Tong S, Kang N, et al. Chronic perfluorooctane sulfonate (PFOS) exposure induces
   hepatic steatosis in zebrafish. Aquatic Toxicology 2016; 176: 45-52.
- Conder JM, Hoke RA, Wolf Wd, Russell MH, Buck RC. Are PFCAs Bioaccumulative? A Critical Review and
   Comparison with Regulatory Criteria and Persistent Lipophilic Compounds. Environmental Science
   & Technology 2008; 42: 995-1003.
- Corman A-M, Mendel B, Voigt CC, Garthe S. Varying foraging patterns in response to competition? A
   multicolony approach in a generalist seabird. Ecology and Evolution 2016; 6: 974-986.
- Costantini D, Blevin P, Herzke D, Moe B, Gabrielsen GW, Bustnes JO, et al. Higher plasma oxidative damage
   and lower plasma antioxidant defences in an Arctic seabird exposed to longer perfluoroalkyl acids.
   Environmental Research 2019; 168: 278-285.

- Couderc M, Poirier L, Zalouk-Vergnoux A, Kamari A, Blanchet-Letrouve I, Marchand P, et al. Occurrence of
   POPs and other persistent organic contaminants in the European eel (*Anguilla anguilla*) from the
   Loire estuary, France. Science of the Total Environment 2015; 505: 199-215.
- Danovaro R. Pollution threats in the Mediterranean Sea: An overview. Chemistry and Ecology 2003; 19: 15 32.
- Darras VM, Verhoelst CH, Reyns GE, Kühn ER, Van der Geyten S. Thyroid hormone deiodination in birds.
   Thyroid 2006; 16: 25-35.
- 566 DeWitt J. Toxicological Effects of Perfluoroalkyl and Polyfluoroalkyl Substances, 2015. Springers press.

567 Elliott JE, Elliott KH. Tracking Marine Pollution. Science 2013; 340: 556.

- Fernandes AR, Mortimer D, Holmes M, Rose M, Zhihua L, Huang X, et al. Occurrence and spatial distribution
   of chemical contaminants in edible fish species collected from UK and proximate marine waters.
   Environment International 2018; 114: 219-230.
- Fridolfsson A-K, Ellegren H. A Simple and Universal Method for Molecular Sexing of Non-Ratite Birds.
   Journal of Avian Biology 1999; 30: 116-121.
- Furness RW, Camphuysen K. Seabirds as monitors of the marine environment. ICES Journal of Marine
   Science 1997; 54: 726-737.
- Giesy JP, Kannan K. Global distribution of perfluorooctane sulfonate in wildlife. Environmental Sciences &
   Technology 2001; 35: 1339-42.
- Haarr A, Hylland K, Eckbo N, Gabrielsen GW, Herzke D, Bustnes JO, Blévin P, Chastel O, Moe B, Hanssen SA,
   Sagerup K, Borgå K. DNA damage in Arctic seabirds: Baseline, sensitivity to a genotoxic stressor, and
   association with organohalogen contaminants. Environmental Toxicology and Chemistry 2018; 37:
   1084-1091.
- Hanssen SA, Hasselquist D, Folstad I, Erikstad KE. Cost of reproduction in a long-lived bird: incubation effort
   reduces immune function and future reproduction. Proceedings. Biological sciences 2005; 272:
   1039-1046.
- Herzke D, Nygård T, Berger U, Huber S, Røv N. Perfluorinated and other persistent halogenated organic
   compounds in European shag (*Phalacrocorax aristotelis*) and common eider (*Somateria mollissima*)
   from Norway: A suburban to remote pollutant gradient. Science of The Total Environment 2009;
   408: 340-348.
- Jacobsen AV, Nordén M, Engwall M, Scherbak N. Effects of perfluorooctane sulfonate on genes controlling
   hepatic fatty acid metabolism in livers of chicken embryos. Environmental Science and Pollution
   Research 2018; 25: 23074-23081.
- Jouanneau W, Bårdsen B-J, Herzke D, Johnsen TV, Eulaers I, Bustnes JO. Spatiotemporal Analysis of
   Perfluoroalkyl Substances in White-Tailed Eagle (*Haliaeetus albicilla*) Nestlings from Northern
   Norway—A Ten-Year Study. Environmental Science & Technology 2020; 54: 5011-5020.
- Kannan K, Tao L, Sinclair E, Pastva SD, Jude DJ, Giesy JP. Perfluorinated compounds in aquatic organisms at
   various trophic levels in a Great Lakes food chain. Archives of Environmental Contamination and
   Toxicology 2005; 48: 559-66.
- Kelly BC, Ikonomou MG, Blair JD, Surridge B, Hoover D, Grace R, et al. Perfluoroalkyl Contaminants in an
   Arctic Marine Food Web: Trophic Magnification and Wildlife Exposure. Environmental Science &
   Technology 2009; 43: 4037-4043.
- Land M, de Wit CA, Bignert A, Cousins IT, Herzke D, Johansson JH, et al. What is the effect of phasing out
   long-chain per- and polyfluoroalkyl substances on the concentrations of perfluoroalkyl acids and
   their precursors in the environment? A systematic review. Environmental Evidence 2018; 7: 4.
- Liu Y, Wang J, Fang X, Zhang H, Dai J. The thyroid-disrupting effects of long-term perfluorononanoate
   exposure on zebrafish (*Danio rerio*). Ecotoxicology 2011; 20: 47-55.
- Lucia M, Strøm H, Bustamante P, Herzke D, Gabrielsen GW. Contamination of ivory gulls (*Pagophila eburnea*) at four colonies in Svalbard in relation to their trophic behaviour. Polar Biology 2017; 40:
   917-929.
- Maynard L, Davoren G. Inter-colony and interspecific differences in the isotopic niche of two sympatric gull
   species in Newfoundland. Marine Ornithology 2020; 48: 103-109.

- McNabb FMA. The Hypothalamic-Pituitary-Thyroid (HPT) Axis in Birds and Its Role in Bird Development and
   Reproduction. Critical Reviews in Toxicology 2007; 37: 163-193.
   Melnes M, Gabrielsen GW, Herzke D, Sagerup K, Jenssen BM. Dissimilar effects of organohalogenated
- 613 compounds on thyroid hormones in glaucous gulls. Environtal Research 2017; 158: 350-357.
   614 Muir DCG, Bossi R, Carlsson P, Evans M, De Silva A, Halsall C, et al. Levels and trends of poly- and
   615 perfluoroalkyl substances in the Arctic environment An update. Emerging Contaminants 2019; 5:
   616 240-271.
- Muir DCG, de Wit CA. Trends of legacy and new persistent organic pollutants in the circumpolar arctic:
   Overview, conclusions, and recommendations. Science of The Total Environment 2010; 408: 3044 3051.
- Munoz G, Budzinski H, Babut M, Lobry J, Selleslagh J, Tapie N, et al. Temporal variations of perfluoroalkyl
   substances partitioning between surface water, suspended sediment, and biota in a macrotidal
   estuary. Chemosphere 2019; 233: 319-326.
- Munoz G, Budzinski H, Labadie P. Influence of Environmental Factors on the Fate of Legacy and Emerging
   Per- and Polyfluoroalkyl Substances along the Salinity/Turbidity Gradient of a Macrotidal Estuary.
   Environmental Science & Technology 2017a; 51: 12347-12357.
- Munoz G, Labadie P, Geneste E, Pardon P, Tartu S, Chastel O, et al. Biomonitoring of fluoroalkylated
   substances in Antarctica seabird plasma: Development and validation of a fast and rugged method
   using on-line concentration liquid chromatography tandem mass spectrometry. J Chromatogr A
   2017b; 1513: 107-117.
- Munschy C, N. B, Pollono C, Aminot Y. Perfluoroalkyl substances (PFASs) in the marine environment: Spatial
   distribution and temporal profile shifts in shellfish from French coasts. Chemosphere 2019; 228:
   640-648.
- Nøst TH, Helgason LB, Harju M, Heimstad ES, Gabrielsen GW, Jenssen BM. Halogenated organic
   contaminants and their correlations with circulating thyroid hormones in developing Arctic
   seabirds. Science of The Total Environment 2012; 414: 248-256.
- Peig J, Green AJ. New perspectives for estimating body condition from mass/length data: the scaled mass
   index as an alternative method. Oikos 2009; 118: 1883-1891.
- Ramhøj L, Hass U, Gilbert ME, Wood C, Svingen T, Usai D, et al. Evaluating thyroid hormone disruption:
   investigations of long-term neurodevelopmental effects in rats after perinatal exposure to
   perfluorohexane sulfonate (PFHxS). Scientific Reports 2020; 10: 2672.
- Ren X-M, Qin W-P, Cao L-Y, Zhang J, Yang Y, Wan B, et al. Binding interactions of perfluoroalkyl substances
   with thyroid hormone transport proteins and potential toxicological implications. Toxicology 2016;
   366-367: 32-42.
- Sebastiano M, Angelier F, Blevin P, Ribout C, Sagerup K, Descamps S, et al. Exposure to PFAS is associated
   with telomere length dynamics and demographic responses of an arctic top predator.
   Environmental Science & Technology 2020 ; 54: 10217–10226
- Simonnet-Laprade C, Budzinski H, Maciejewski K, Le Menach K, Santos R, Alliot F, et al. Biomagnification of
   perfluoroalkyl acids (PFAAs) in the food web of an urban river: assessment of the trophic transfer of
   targeted and unknown precursors and implications. Environmental Science: Processes & Impacts
   2019; 21: 1864-1874.
- 651Smits JE, Fernie KJ, Bortolotti GR, Marchant TA. Thyroid hormone suppression and cell-mediated652immunomodulation in American kestrels (*Falco sparverius*) exposed to PCBs. Archives of653Environmental Contamination and Toxicology 2002; 43: 338-44.
- Sun J, Bossi R, Bustnes JO, Helander B, Boertmann D, Dietz R, et al. White-Tailed Eagle (*Haliaeetus albicilla*)
   Body Feathers Document Spatiotemporal Trends of Perfluoroalkyl Substances in the Northern
   Environment. Environmental Science & Technology 2019.
- Sunderland EM, Hu XC, Dassuncao C, Tokranov AK, Wagner CC, Allen JG. A review of the pathways of
   human exposure to poly- and perfluoroalkyl substances (PFASs) and present understanding of
   health effects. Journal of exposure science & environmental epidemiology 2019; 29: 131-147.

- Tartu S, Gabrielsen GW, Blévin P, Ellis H, Bustnes JO, Herzke D, et al. Endocrine and Fitness Correlates of
   Long-Chain Perfluorinated Carboxylates Exposure in Arctic Breeding Black-Legged Kittiwakes.
   Environmental Science & Technology 2014; 48: 13504-13510.
- Verreault J, Houde M, Gabrielsen GW, Berger U, Haukås M, Letcher RJ, et al. Perfluorinated Alkyl
   Substances in Plasma, Liver, Brain, and Eggs of Glaucous Gulls (*Larus hyperboreus*) from the
   Norwegian Arctic. Environmental Science & Technology 2005; 39: 7439-7445.
- Verreault J, Skaare JU, Jenssen BM, Gabrielsen GW. Effects of orgonochlorine contaminants on thyroid
   hormone levels in arctic breeding glaucous gulls, *Larus hyperboreus*. Environmental Health
   Perspectives 2004; 112: 532-537.
- Verreault J, Villa RA, Gabrielsen GW, Skaare JU, Letcher RJ. Maternal transfer of organohalogen
   contaminants and metabolites to eggs of Arctic-breeding glaucous gulls. Environmental Pollution
   2006; 144: 1053-1060.
- Vicente J, Sanpera C, García-Tarrasón M, Pérez A, Lacorte S. Perfluoroalkyl and polyfluoroalkyl substances in
   entire clutches of Audouin's gulls from the ebro delta. Chemosphere 2015; 119: S62-S68.
- Wan HT, Zhao YG, Wei X, Hui KY, Giesy JP, Wong CKC. PFOS-induced hepatic steatosis, the mechanistic
   actions on β-oxidation and lipid transport. Biochimica et Biophysica Acta (BBA) General Subjects
   2012; 1820: 1092-1101.
- Wang Z, DeWitt JC, Higgins CP, Cousins IT. A Never-Ending Story of Per- and Polyfluoroalkyl Substances
   (PFASs)?. Environmental Science & Technology 2017; 51: 2508-2518.
- Weiss JM, Andersson PL, Lamoree MH, Leonards PEG, van Leeuwen SPJ, Hamers T. Competitive Binding of
   Poly- and Perfluorinated Compounds to the Thyroid Hormone Transport Protein Transthyretin.
   Toxicological Sciences 2009; 109: 206-216.
- Wong F, Shoeib M, Katsoyiannis A, Eckhardt S, Stohl A, Bohlin-Nizzetto P, et al. Assessing temporal trends
   and source regions of per- and polyfluoroalkyl substances (PFASs) in air under the Arctic Monitoring
   and Assessment Programme (AMAP). Atmospheric Environment 2018; 172: 65-73.
- 685Xie S, Wang T, Liu S, Jones KC, Sweetman AJ, Lu Y. Industrial source identification and emission estimation686of perfluorooctane sulfonate in China. Environment International 2013; 52: 1-8.
- Ku J, Shimpi P, Armstrong L, Salter D, Slitt AL. PFOS induces adipogenesis and glucose uptake in association
   with activation of Nrf2 signaling pathway. Toxicology and applied pharmacology 2016; 290: 21-30.
- Yeung LWY, Guruge KS, Yamanaka N, Miyazaki S, Lam PKS. Differential expression of chicken hepatic genes
   responsive to PFOA and PFOS. Toxicology 2007; 237: 111-125.
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# Ile de Ré (South western France)

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30

20

10

0

L- PFOS (ng/g)

# **T**3) hormones nyroid

# Larus marinus

PFAS