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- 1 Sea turtles across the North Pacific are exposed to perfluoroalkyl substances
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21 Abstract

- 22 Perfluorinated alkyl substances (PFASs) are global, persistent, and toxic contaminants.
- 23 We assessed PFAS concentrations in green (Chelonia mydas) and hawksbill (Eretmochelys
- 24 *imbricata*) turtles from the North Pacific. Fifteen compounds were quantified via liquid

chromatography tandem mass spectrometry from 62 green turtle and 6 hawksbill plasma samples 25 26 from Hawai'i, Palmyra Atoll, and the Northern Marianas Islands. Plasma from 14 green turtles 27 severely afflicted with fibropapillomatosis, and eggs from 12 Hawaiian hawksbill nests from 7 females were analyzed. Perfluorooctane sulfonate (PFOS) predominated in green turtle plasma; 28 29 perfluorononanoic acid (PFNA) predominated in hawksbill tissues. Concentrations were greater in hawksbill than green turtle plasma (p<0.05), related to trophic differences. Green turtle plasma 30 31 PFOS concentrations related to human populations from highest to lowest: Hawai'i, Marianas, 32 Palmyra. Influence on fibropapillomatosis was not evident. PFASs were maternally transferred 33 to hawksbill eggs, with decreasing concentrations with distance from airports and with clutch 34 order from one female. A risk assessment of PFOS showed concern for immunosuppression in Kailua green turtles and alarming concern for hawksbill developmental toxicity. 35 36 Perfluoroundecanoic (PFUnA) and perfluorotridecanoic (PFTriA) acid levels were correlated 37 with reduced emergence success (p<0.05). Studies should further examine toxicity of PFASs on 38 sea turtle development. 39 **40 Capsule:** 41 Endangered or threatened sea turtle species at remote Pacific islands contain perfluoroalkyl 42 substances in blood and offload them to eggs, at concentrations known to be detrimental in birds. 43 44 Keywords: Perfluorinated contaminants, marine turtles, Pacific islands, reptile, maternal transfer

46 Introduction

47 Perfluoroalkyl substances (PFASs) are a chemical family used in many industrial and 48 commercial products (Lau et al. 2007). Their chemical makeup typically consists of a fluorinated 49 carbon backbone (4 to 14 in length) and a charged functional moiety, such as a carboxylate. The 50 carbon-fluorine bond is incredibly stable, which gives PFASs extreme persistence and both 51 hydrophobic and lipophobic properties (Buck et al. 2011). Since their invention in the 1930s and 52 commercialization in the 1950s, PFASs have been incorporated into every-day consumer 53 products including coatings, fabrics, grease-proof papers, soil repellents, and aqueous film-54 forming foam fire suppressants (AFFFs) (Moody and Field 2000; Buck et al. 2011; Ahrens and 55 Bundschuh 2014; Interstate Technology Regulatory Council 2020). AFFFs used for firefighting 56 and training at military bases and airports are large point sources of PFAS contamination 57 (Schultz et al. 2004). Their nearly non-biodegradable properties, along with global (Yamashita et 58 al. 2008) render PFASs highly persistent pollutants in the environment worldwide(Lau et al. 59 2007) with detectable baseline concentrations in the most remote regions (Young et al. 2007). 60 PFASs are prominent contaminants in wildlife and human tissues, and can result in toxicological effects, including immune, developmental, hepatic, and endocrine disruptions (Lau 61 62 et al. 2007; DeWitt 2015). In 2001, 3M Corporation, a major producer of perfluorooctane 63 sulfonate (PFOS), began phasing out production of this chemistry. In 2006, eight manufacturers 64 committed to reduce perfluorooctanoate (PFOA) production by 2015 through a US EPA 65 Stewardship program (US EPA 2018). Additional policies have focused on eliminating further production of PFASs in Europe and Asia, but PFASs remain prominent in the environment 66 (McCarthy 2017). 67

68 Because sea turtles are rare and protected, understanding their exposure to toxic 69 contaminants is important for the recovery of these species (NMFS and US FWS 1998b, a). 70 Moreover, certain life stages of sea turtles can indicate the contaminant levels in their respective 71 foraging grounds, because they integrate those contaminants over several years (O'Connell et al. 72 2010; Keller 2013). The generalized long life history of green (Chelonia mydas) and hawksbill 73 (Eretmochelys imbricata) sea turtles includes an ontogenetic switch that separates two juvenile 74 phases (Bolten 2003). The youngest turtles forage in open ocean habitats, and transition to 75 foraging in nearshore benthic habitats as older juveniles where they grow into subadults and 76 adults. As adults, they migrate from foraging grounds to inter-nesting habitat where they do not 77 eat much. The green turtle is the lowest trophic level sea turtle species (Bjorndal 1997). They transition from omnivorous pelagic juveniles (Parker et al. 2011) to herbivory in the neritic 78 79 stages (Arthur and Balazs 2008; Summers et al. 2017). Green turtles are highly migratory, across 80 entire ocean basins, but have strong site fidelity to foraging grounds within a life stage (Summers 81 et al. 2017; Shimada et al. 2020). Hawksbill sea turtles are less migratory (Gaos et al. 2020b) and 82 continue to feed at higher trophic levels after their ontogenetic switch to eating primarily sponges 83 in the neritic phase (Bjorndal 1997; Summers et al. 2017). Six studies have reported PFAS concentrations in tissues of six sea turtle species (Keller 84

et al. 2005; O'Connell et al. 2010; Keller et al. 2012; Guerranti et al. 2013; Guerranti et al. 2014;

86 Pasanisi et al. 2016), but none from turtles across the North Pacific. Most analyzed plasma

87 (Keller et al. 2005; O'Connell et al. 2010; Keller et al. 2012), and none have analyzed eggs.

88 Differences in concentrations among five species provide evidence that PFASs biomagnify in the

89 turtles' distinct food webs and that the major route of exposure is diet (Keller et al. 2012). For

90 example, PFAS concentrations are greater in carnivorous loggerhead (*Caretta caretta*) sea turtles
91 than herbivorous green turtles from the same neritic habitat (Keller et al. 2012).

92 The toxicological effects of PFASs on sea turtles are still largely undetermined. 93 Immunosuppressive contaminants have long been a suspected contributing influence in a 94 debilitating, tumor-forming, viral disease, fibropapillomatosis (FP) (Herbst and Klein 1995; Keller 2013), but to date, no convincing causal relationship between exposure to toxicants and 95 96 development of FP has been made. For instance, when levels of other types of persistent organic 97 pollutants (POPs) were measured in the plasma of many of the same turtles in the current study, 98 the levels did not relate to severity of FP in Hawaiian green turtles. PFASs were not included in 99 that analysis (Keller et al. 2014a). Furthermore, maternal offloading of POPs occurs during yolk 100 deposition (Keller 2013; Munoz and Vermeiren 2020), but transfer of PFASs to eggs and effects 101 on hatching success and embryonic development are unknown.

102 In this study, we measured concentrations of 15 PFASs in the plasma of neritic-phase 103 green and hawksbill sea turtles across three North Pacific regions: the Main Hawaiian Islands 104 (MHI), the Commonwealth of the Northern Marianas Islands (CNMI), and Palmyra Atoll. Eggs 105 from excavated hawksbill nests from MHI were also analyzed We hypothesized that (1) PFOS 106 will predominate like in most other studies, (2) higher trophic positioned, spongivorous 107 hawksbills will have greater plasma PFAS concentrations than herbivorous green turtles, (3) 108 PFAS concentrations in turtles will correlate to human population or proximity to military bases 109 or airports (and in the lack of local point sources, remote Palmyra Atoll will represent globally 110 diffuse contamination levels transported by air and ocean currents), (4) PFASs will be detected 111 in the sampled hawksbill eggs, revealing maternal offloading, (5) PFAS concentrations in eggs

will negatively correlate to nest success variables, and (6) eggs from successive clutches fromthe same female within a nesting season will have stable PFAS concentrations.

114

115 2. Materials and Methods

116 2.1 Study Sites

117 Samples were collected on beaches and coastal waters from three Pacific Island regions: 118 the Main Hawaiian Islands (MHI), the Commonwealth of the Northern Marianas Islands 119 (CNMI), and Palmyra Atoll (Figure 1). The MHIs have roughly 1.4 million people, three 120 international airports, and 11 military bases (US Census Bureau 2019a). CNMI has about 57,000 121 people (United Nations Dept. of Economic and Social Affairs 2019), three international airports, 122 and no active military bases, but a historically large military presence from World War II. 123 Palmyra Atoll is a remote U.S. National Wildlife Refuge with less than 20 staff, no commercial 124 airport, and a decommissioned military base.

125 2.2 Sample Collection

126 Cryogenically archived sea turtle samples were selected from the NIST Biorepository's 127 Biological and Environmental Monitoring and Archival of Sea Turtle tissues project (BEMAST). 128 Sample collection and processing methods are detailed in Keller et al. (2014b). Sample metadata 129 are provided (Tables S1, S2). Plasma samples were selected to investigate spatial, species, and 130 health status differences and included 62 samples from 61 free-ranging green sea turtles from 131 across three study sites (Table 1, Figure 1): 39 from MHI (n=13 from three sites), 12 from CNMI 132 (n=6 from Saipan and Tinian), and 10 from Palmyra Atoll (n=2 from each segment of the refuge, 133 with one turtle sampled twice over two years). Plasma from 14 severely tumored, MHI-stranded 134 turtles were also analyzed. All MHI plasma samples came from the same turtles that were

135	analyzed previously for other POPs (Keller et al. 2014a), representing four groups of green
136	turtles with varying prevalence of FP: Kiholo Bay (0% FP), Kailua Bay (low to moderate FP 0%
137	to 5%), Kapoho Bay (higher FP 35%) and tumored-stranded (100% FP). All six hawksbill
138	plasma samples available in the BEMAST inventory from CNMI (n=4, two each from Saipan
139	and Tinian) and Palmyra (n=2) were included. All 12 hawksbill nests available in BEMAST
140	inventory at the time were included. They were from beaches on Maui, Hawai'i, and Kauai
141	(Figure 1) in 2012 and represent seven nesting females (Table S2). Unhatched eggs were
142	collected from each nest upon excavation after hatchling emergence. Egg contents from at most
143	three unhatched eggs per nest were pooled, homogenized, aliquoted, and cryogenically archived.
144	The number of eggs laid, days of incubation, hatching success, and emergence success were
145	calculated per nest according to Miller (1997) to inspect the fitness of each nest in relation to
146	PFAS concentrations. The percent of eggs laid that developed to at least stage 23 of
147	development, as defined by Miller et al. (2017) was also calculated (Table S2).
148	2.3 PFAS Measurements
149	Detailed methods are provided in Supplemental Information. Samples were measured
150	using methods modified from Keller et al. (2012). Concentrations of 15 PFASs were quantified
151	using the internal standard approach (Table S3). Field blanks were prepared with the same lot
152	number of blood collection supplies (Table S4). Samples were extracted via sonication with
153	acetonitrile and purified on 250 mg Envi-carb columns (Supelco, Bellefonte, Pennsylvania).
154	Methanol extracts (20 μ L) were injected on an Agilent Zorbax Eclipse Plus C18 column using a
155	liquid chromatograph (Agilent 1100 HPLC, Palo Alto, CA) negative electrospray ionization
156	tandem mass spectrometer (API 4000, Applied Biosystems-MDS Sciex, Foster City, CA). The
157	mobile phase consisted of methanol and 20 mmol/L ammonium acetate in water. Two transitions

158 for PFOS were monitoring $(499 \rightarrow 99 \text{ and } 499 \rightarrow 80)$ to ensure bile acids did not interfere.

159 Concentrations are totals of linear and branched isomers. Reporting limits (RLs) were

160 determined according to O'Connell et al. (2010). The mass fractions of PFASs measured in NIST

161 SRM 1957 Organic Contaminants in Non-Fortified Human Serum and SRM 1947 Lake

162 Michigan Fish Tissue were within the uncertainty of the reference values (Table S5).

163 2.4 Data Analyses

164 R software package, NADA, was used whenever possible when samples were <RL as 165 suggested by Helsel (2005), otherwise JMP (SAS Institute Inc. Cary, NC) was used with values 166 <RL substituted with half the RL. Significance was determined as p<0.05. Data was assessed for 167 assumptions of normality and equal variances. Non-parametric ANOVAs in NADA tested 168 differences among sites for green turtle plasma PFAS concentrations, followed by Steel Dwass 169 non-parametric multiple comparison tests using JMP. Wilcoxon t-tests in JMP compared 170 hawksbill plasma PFAS concentrations between Palmyra and CNMI, except PFUnA required 171 NADA's Wilcoxon t-test. Non-parametric ANOVAs in NADA tested differences among sites 172 and FP incidence (grouped as in Keller et al. (2014a)) for green turtle plasma PFOS 173 concentrations, followed by Steel Dwass non-parametric multiple comparison tests using JMP. 174 Species differences were assessed with CNMI data. 175 Amount of PFASs offloaded into each nest was estimated by multiplying the total 176 number of eggs laid by the average grams of each pooled egg contents from that nest, and by the 177 measured PFAS concentration (ng/g). The nanograms of PFASs offloaded were then compared 178 across clutches from one nesting season of a mother using linear regressions. PFAS 179 concentrations in the eggs were compared to nest success variables using Kendall's tau 180 correlations in NADA.

181	To assess toxicological risk, estimated margins of safety (EMOS) were calculated as
182	described in Keller et al. (2012), as the ratio of the average plasma, serum, or egg PFOS
183	concentration in laboratory-exposed animals in the lowest observed adverse effect level
184	(LOAEL) dose group to either the average or maximum sea turtle plasma or egg PFOS
185	concentration. EMOS below 100 were considered of concern (Faustman and Omenn 1996).
186	

187 3. Results and Discussion

188 3.1 Plasma PFAS Concentrations

PFAS mass fractions (hereafter called concentrations) in ng/g wet mass for each turtle
plasma sample are provided (Table S6). Summary statistics for all possible groupings of Pacific
turtles are compared to turtles along the Eastern U.S. (Table S7).

192 3.1.1 Green Turtle Plasma PFAS Concentrations

193 PFOS comprised >96% of the Σ PFAS concentrations in green turtle plasma regardless of 194 capture location (Table 1; Figures 2 and S1), similar to previous reports for green, leatherback, 195 loggerhead, and Kemp's ridley turtles from the Eastern U.S. (O'Connell et al. 2010; Keller et al. 196 2012). PFOS concentrations in green turtle plasma were significantly different among the Pacific 197 Island regions and increased with human population (Figure 3a). This trend continues when 198 including previously published PFOS concentrations in green turtles from Core Sound, North 199 Carolina (mean=2.41 ng/g; (Keller et al. 2012) Figure 3a), a watershed with $\approx 6,390,000$ people 200 (O'Connell et al. 2010). The MHI with 1,400,000 people had the next highest mean PFOS 201 concentration at 1.14 ng/g and significantly greater than CNMI (57,000 people) at 0.524 ng/g 202 and Palmyra (20 people) at 0.155 ng/g. This relationship between PFAS plasma concentration 203 and human population corroborates previous relationships for sea turtle contaminant exposure

204 (O'Connell et al. 2010; Alava et al. 2011), is expected for these man-made chemicals, and
205 suggests that local or regional sources add to the diffuse, globally distributed PFAS levels in
206 nearshore marine organisms.

207 Green turtles from the MHIs were categorized into four groups based on their capture 208 location and incidence and severity of FP using the same grouping as Keller et al. (2014a). These 209 were No FP (Kiholo), Low FP (Kailua), High FP (Kapoho), and FP stranded. The FP stranded 210 turtles were alive, severely afflicted with FP tumors, and exceedingly emaciated because of the 211 disease. Significant differences in PFOS plasma concentrations were noted among the groups 212 (p=5e-9), but not in a dose-dependent manner if PFOS was contributing to FP (Figure S2). The 213 low FP Kailua group (mean=2.45 ng/g), from the most urbanized area sampled, had significantly 214 higher concentrations than the other three groups and was relatively similar to those in Core 215 Sound, NC (2.41 ng/g) (Table 1, Table S7) (Keller et al. 2012). This is likely due to O'ahu being 216 a heavily populated island (974,563 people), and hosting numerous military bases, airports, and 217 other point-sources (US Census Bureau 2019a). The Kailua sampling site is adjacent to Marine 218 Corps Base Hawai'i, with an active airstrip and firefighting training sites. Additionally, the 219 Kawainui watershed empties into Kailua Bay at this sampling site and contains 19.1% urban 220 development, while Kapoho and Kiholo watersheds had 7.8% and 0% urban development, 221 respectively (Parham et al. 2008a, b, c). Given that the MHIs are one of the most isolated 222 archipelagos in the world, approximately 4,000 km away from the nearest continent (National 223 Academy of Sciences 2004), air and ocean circulation transport a baseline level of globally 224 diffuse PFASs to Hawaiian waters. The differences among turtle groups here indicate that local 225 sources compound global sources of PFASs in Hawai'i.

226 The lack of a dose-dependent relationship between PFASs and FP is similar to results for 227 other POPs that were measured in these same turtles (Keller et al. 2014a) in which Kailua turtles 228 with low FP incidence had higher concentrations than the other two locations. What differs 229 between the two studies is the relative difference in the FP stranded turtles. For protein-230 associating PFOS, the FP stranded turtles had levels that were close to average among the MHI turtles (Table 1). For lipophilic POPs, the FP stranded turtles had elevated levels compared to the 231 232 other MHI turtles (Keller et al. 2014a). This can be explained by the different distribution of 233 these compounds during weight loss and lipid mobilization. The lipophilic POPs flooded into the 234 blood of the emaciated FP stranded group upon weight loss. With no lipid to associate within the 235 thin turtles, those POPs continued to circulate in the blood at high levels. Conversely, PFOS 236 preferentially associates with proteins in the blood and liver. Upon weight loss, protein 237 concentrations in these tissues do not change as much as lipid levels. Thus, PFOS remains 238 associated with proteins, even in emaciated animals, resulting in less mobilization of PFOS into 239 the blood upon weight loss.

Green and hawksbill turtles inhabit the coral reefs of CNMI (Summers et al. 2017), a U.S. Commonwealth comprised of 14 islands. They face several threats, including illegal harvest and marine debris entanglement (Summers et al. 2018). Sampling sites were on the archipelago's two most populated islands, Saipan with 48,200 residents and Tinian with 3,056 residents (Department of Commerce Central Statistics Division 2017). Green turtles from Saipan had significantly greater plasma PFOS and Σ PFAS concentrations than from Tinian (p=0.02); (Table 1), corroborating other human population trends.

Palmyra Atoll is extremely remote in the Central Pacific at 5.9° N 162.1° W, with a
history of extensive human disturbance during World War II in the 1940s when construction of a

U.S. naval base necessitated the remodeling of the small islets (Collen et al. 2009). Construction
of >100 buildings and two airstrips occurred between 1940 and 1944. Dredging in the West
Lagoon, as well as causeway construction across multiple reef flats, roughly doubled the land
area and resulted in major hydrodynamic changes in the lagoon (Collen et al. 2009). By 1945,
most military activity had ceased and personnel were evacuated, which prompted many natural
changes in the atoll's topography (Collen et al. 2009). Since 2001, the atoll has been protected as
a U.S. Fish and Wildlife Refuge.

256 No uses or disposal of PFASs are known on Palmyra Atoll. AFFFs were invented 15 257 years after the U.S. Navy abandoned the atoll (Moody and Field 2000). A cursory spatial 258 comparison using green turtle plasma PFAS concentrations showed no evidence of a significant 259 point source on the atoll (see Supplemental Information). The comparison was confounded by 260 life stage with the greatest PFAS burden in a young green turtle that recently recruited from the 261 pelagic carnivorous stage into the nearshore herbivorous stage. New recruits favor the eastern region of Palmyra (Sterling et al. 2013), and may carry a greater PFAS burden to Palmyra, which 262 263 dilutes with growth thereafter (Keller 2013). The current study suggests that remote Palmyra 264 Atoll is a control reference site with little to no local point sources of PFASs, reflecting baseline 265 globally diffuse concentrations. The first study to measure chemical contaminants in Palmyra sea 266 turtles found elevated aluminum and iron McFadden et al. (2014).

267

268 3.1.2 Hawksbill Plasma PFAS Concentrations

Perfluorononanoic acid (PFNA) predominated in hawksbill plasma from CNMI (n=4) and Palmyra (n=2), with means of 1.08 ng/g and 3.40 ng/g, respectively (Figures 2 and 3b). The same PFAS profile was seen in hawksbills from Juno Beach, FL (a typo in in Keller et al. (2012) should read 17.3 ng/g for mean PFNA). It is interesting that hawksbill turtles from such distant regions have similar PFAS profiles (PFNA predominating), while green turtles from these
disparate regions show PFOS dominating. The reasons for the species differences in PFAS
profiles are difficult to explain, especially since marine mammals stranded in Hawai'i show still
another profile (perfluoroundecanoic acid (PFUnA) dominated (Kurtz et al. 2019)). Differences
may be explained by elimination mechanisms, prey selection, or migratory pathways.

278 CNMI and Palmyra hawksbill plasma had lesser PFAS concentrations than Juno Beach 279 hawksbills, in line with the drastic differences in human population (Figure 3b). This finding is 280 congruent with the green turtle spatial differences described above and those in loggerhead 281 turtles from the U.S. East coast (O'Connell et al. 2010). Together these findings support the idea 282 that PFAS levels, biomagnifying in marine organisms inhabiting developed nearshore regions, 283 are influenced more from local or regional land-based sources than diffuse sources from global 284 air and ocean currents. Juno Beach, FL, is within the St. Lucie-Loxahatchee Watershed which 285 has 6,400,000 human residents (South Florida Water Management District 2009; US Census Bureau 2019b, c). Surprisingly, Palmyra hawksbills (only 20 human residents) had a greater 286 287 mean concentration of PFNA, PFUnA, and Σ PFASs than those in CNMI, but low sample size 288 prevented statistical analyses. The reasons for the observed spatial differences could include prey 289 selection, migratory pathways, or sources of contamination.

290

291 3.1.3 Species Comparisons of Plasma PFAS Concentrations

292 The two hawksbill turtles from Palmyra were both captured in the eastern region of the 293 atoll, and had PFAS concentrations one order of magnitude greater than the Palmyra green turtle 294 mean (Tables 1 and S6). In CNMI, ∑PFAS concentrations in hawksbill plasma was significantly 295 greater than green turtles (Figure S3). These results corroborate previous results showing Eastern 296 U.S. hawksbills had greater plasma PFOS levels than three other species, including green turtles (Keller et al. 2012). Similarly, in Australia and Japan, hawksbills had greater concentrations of
POPs (PFASs were not tested) than green turtles (Hermanussen et al. 2008; Malarvannan et al.
2011). The trophic position of the two species may explain greater biomagnification of PFASs in
hawksbills. Omnivorous hawksbills primarily prey on sponges, but also forage on other
invertebrates and marine vegetation, placing them higher on the food web than the herbivorous
stage of green turtles studied (Bjorndal 1997).

303

304 3.2 Egg PFAS Concentrations

305 Eleven PFAS compounds were detected in at least one hawksbill nest (Tables 1 and S2). 306 Other POPs are known to be deposited into eggs from mother sea turtles during egg production, 307 rather than crossing the eggshell in the nest environment (Keller 2013; Munoz and Vermeiren 308 2020). Therefore, POP concentrations in eggs represent the mother's exposure from her foraging 309 grounds (Alava et al. 2011). It is expected that PFASs deposit into eggs in a similar fashion as 310 maternal offloading has been documented in other egg-laying species (Wilson et al. 2020). The 311 presence of PFASs in hawksbill eggs reveals, for the first time, offloading from females to their 312 eggs and, to our knowledge, is the first report of PFAS offloading in any reptile species. These 313 data fill an important gap in understanding exposure of adult female hawksbills and their 314 developing embryos to PFASs.

 \sum PFAS concentrations were greatest in the nest laid in Wailua, Kauai, and lowest in the four nests from three mothers in Pohue, Hawai'i (Figure S4). Nest PFAS concentrations were negatively, significantly (p=0.030) correlated with the distance over water from the nests to the nearest international airport (Figure S4). This preliminary finding may be explained by airports and military bases being some of the largest point sources of PFASs due to firefighting training 320 (Schultz et al. 2004; Houtz et al. 2016). The relationship was driven by the Wailua nest, and a
321 larger sample size should be analyzed with future satellite tracks of females.

322 PFAS profiles in eggs when averaged across the seven mothers were dominated by PFOS 323 (28.5% of *SPFASs*), PFUnA (24.7%), and PFNA (23.8%) (Figures 2 and S1). More 324 interestingly, the eggs from different mothers displayed drastic differences in PFAS profiles (Table S2, Figure S5). The nest laid in Wailua, Kauai, had the highest predominance of PFOS 325 326 (88%) and PFTA (3.5%) and the lowest PFNA contribution (0.9%), a profile that reflects older, 327 phased-out formulations of AFFFs (Place and Field 2012). The four nests laid by three females 328 in Pohue, Hawai'i had an intermediate profile with PFUnA dominating, followed by either PFOS 329 (16% to 31%) or PFDA. The five nests laid by one female (ID 19591-04) on Maui were shifted 330 towards PFNA > PFOS (22.6%) > PFUnA > PFDA. Finally, the two nests from likely different 331 females on Apua and Kamehame beaches on Hawai'i had a profile most different from the Kauai 332 nest, dominated by PFNA > PFUnA > PFDA, with PFOS comprising only 6%. The extreme 333 spectrum of PFOS contributions, from 88.3% in Kauai to 6% in Hawai'i, suggests that adult 334 female Hawaiian hawksbill turtles forage in distinct areas that have different exposure profiles. 335 This suggestion is partially explained by understanding that POP concentrations in sea turtle 336 eggs originate from the mother's diet from her foraging grounds (Alava et al. 2011; Keller 2013). 337 Therefore, the interpretation of the extreme spectrum of PFOS contribution must be placed in 338 context of hawksbill migration and foraging selection, as described next. 339 Though hawksbill turtles are capable of long-distance migrations (>2,000 km) (Vaughan 340 and Spring 1980), both foraging juveniles and nesting adults exhibit natal philopatry with 341 relatively small foraging ranges (Gaos et al. 2017; Wood et al. 2017; Gaos et al. 2018; Gaos et al. 342 2020b). Hawaiian hawksbills have recently been recognized as a genetically distinct

343 management unit, with little connectivity to other populations (Gaos et al. 2020a). Satellite tracks 344 of nine Hawaiian hawksbills show that their nesting and foraging grounds are close, on the same 345 or neighboring island (Parker et al. 2009). One Maui nester (ID 19591-04) migrated to O'ahu to 346 forage in 2004 (Parker et al. 2009). Four years later, she nested again in the same region on Maui 347 and her post-nesting track (King et al. unpublished data) showed the same path to the same O'ahu foraging site (Figure S6). Four more years later, she laid five nests on Maui that were 348 349 sampled in the current study. This strong site fidelity to foraging grounds, relatively near the 350 nesting beaches, with potentially different local point sources across the Main Hawaiian Islands 351 may explain the extreme differences in PFAS profiles observed between the nests laid farthest 352 from each other: Kauai vs. Kamehame. Future studies could test this theory by measuring PFAS 353 concentrations and profiles in marine environmental samples, such as sediment or prey in known 354 hawksbill foraging grounds.

355 Of the 12 nests sampled in 2012, five were laid during the same nesting season (22 days to 25 days apart) by Turtle 19591-04 in Makena, Maui. Two other nests were laid by another 356 357 female (ID 112) 43 days apart at Pohue, Hawai'i; it is likely this turtle laid at least one more nest 358 between these dates but it was not sampled. Two additional Hawai'i Island nests came from 359 turtles with unique IDs, and three nests were laid by unknown females (Table S2). The amounts 360 of PFASs in nests of Turtle 19591-04 significantly declined with clutch order (Figure 4), 361 indicating maternally offloaded contaminant concentrations may be a function of time within a 362 nesting season. However, a decline in transferred PFAS amounts was not apparent in the two 363 nests from Turtle 112.

When sea turtles arrive at their nesting site, they are equipped with all lipid-rich follicles
ready to become the yolk of all eggs to be laid that nesting season (Miller 1997). The follicles for

366 a single nest transit the oviduct where they are fertilized and surrounded by protein-rich albumen 367 (Miller 1997). After laying this clutch, hawksbills prepare the next clutch which is laid 14 to 25 368 days later. Because PFASs associate with serum albumin and fatty acid binding proteins (Ahrens 369 and Bundschuh 2014) rather than lipids, they should deposit more in egg albumen than follicles. 370 Since the albumen is deposited just before each nest is laid, and because in general mother turtles fast during nesting (Miller 1997; Hays et al. 2002; Guirlet et al. 2008; Guirlet et al. 2010), the 371 372 mother's body could have less PFASs to transfer through albumen into successive clutches. 373 Theoretically, females would offload a greater portion of her body burden of PFASs into the first 374 clutch. Turtle 19591-04 offloaded a total of 367 µg of PFASs into these five clutches in one year, 375 with approximately 25% of that into the first clutch and 12% in the fifth clutch (Table S8). 376 Previously, a decrease in Σ PCBs, Σ HCHs and Σ DDTs yolk concentrations from successive 377 leatherback clutches suggested that reproductive lipid investment into eggs decreases as the 378 maternal lipid stores decrease (Guirlet et al. 2010). The current findings suggest that PFASs are 379 offloaded through albumen, and that Turtle 19591-04 was fasting during this nesting season 380 while Turtle ID 112 may have been foraging. These interpretations are supported by changes in 381 chicken egg PFAS concentrations during and after exposure of hens to PFASs in drinking water 382 (Wilson et al. 2020).

Nest success variables were examined for relationships with PFAS concentrations (n=11 nests). Only two significant correlations were observed (p<0.05). Emergence success was negatively correlated with concentrations of two contaminants: PFUnA and perfluorotridecanoic acid (PFTriA) (Figure 5). Few studies exist on developmental effects of PFASs, and no toxicology studies are available for reptiles. In chickens, hatching success was significantly reduced to 61.4 % by a 100 ng/g injection of PFOS into eggs compared to 85.7 % in controls 389 (Molina et al. 2006). In tree swallow (Tachycineta bicolor) nests, hatching success was 390 significantly, negatively correlated with PFOS concentrations, and complete nest failure was 391 observed in three nests with concentrations at or above 150 ng/g (Custer et al. 2012). While we 392 saw no negative correlation between egg burdens of PFOS and hatching success, PFASs may 393 have more insidious reproductive effects for hawksbill turtles. These novel results indicate a 394 potential consequence of PFAS maternal offloading to embryonic sea turtles, and are supported 395 by our risk assessment. Future studies should address the developmental effects of PFASs in turtles. It is possible that the correlations in this study between emergence success and PFASs are 396 397 confounded by the decline in PFASs in successive clutches, but to the authors' knowledge, no 398 studies have examined whether emergence success increases in successive clutches of sea turtles. 399 3.3 Risk Assessment

Using surrogate species is the only available option for sea turtle toxicology risk 400 401 assessments, but because reptiles can be more sensitive than other taxa (Weir et al. 2010) a wider 402 margin of safety (<100) should be used. Keller et al. (2012) estimated margins of safety for 403 toxicological effects of PFOS based on plasma concentrations in five species of sea turtles along 404 the Eastern U.S. All five species had margins of safety <100, which indicate a risk of at least 405 immunosuppressive effects. Using the same method for PFOS plasma concentrations, average 406 green turtles from Kailua Bay and the maximum green and hawksbill turtles from CNMI were at 407 risk of immunosuppression (Table 2). Likewise for eggs, the average hawksbill nest had a 408 margin of safety of only six, indicating heightened risk of reduced hatching success (Table 2). 409 More concerningly, the maximum nest from Kauai was nearly equal to the PFOS concentrations 410 (no margin of safety) that cause reduced hatching success in chickens and tree swallows (Molina 411 et al. 2006; Custer et al. 2012). This represents the first PFAS risk assessment for embryonic

stage sea turtles and suggests that sea turtles inhabiting regions close to military bases andairports, even on remote islands, could be at risk of PFAS toxicity.

414 4. Conclusion

415 Reptiles are significantly under-studied in toxicology, particularly for PFASs (Reiner and 416 Place 2015). This is the first report of plasma PFAS concentrations in Pacific sea turtles, plus the 417 first to document maternal offloading of PFASs into eggs of any sea turtle species. The results 418 reveal contamination patterns similar to those documented along the Eastern U.S., with PFOS 419 predominating in green turtles, hawksbills accumulating greater levels than green turtles, and 420 PFAS concentrations being related to human population and specific land uses. The PFAS 421 concentrations in Pacific turtles are generally less than those along the Eastern U.S., which can 422 be attributed to the remoteness and smaller human populations of the islands studied. Across the 423 study sites, islands with greater population densities and closer proximity to military bases and 424 airports rendered greater PFAS concentrations in turtles. Incidence and severity of FP did not 425 relate to PFAS concentrations, so the search continues for environmental stressors that may 426 contribute to this viral disease. No prior study has analyzed sea turtle tissues concurrently with 427 prey items for PFAS levels; future studies on this would improve our understanding of trophic 428 transfer. The PFAS egg concentrations are novel for reptilian species and show maternal 429 offloading is strongest in the first clutch of a season and egg concentrations were highest in nests 430 laid nearest international airports. Two contaminants (PFUnA and PFTriA) were related to 431 reduced emergence success of hatchlings, which aligns with the risk assessment showing 432 hawksbill egg PFOS concentrations are concerningly near concentrations causing developmental 433 toxicity in birds. Future studies should address the toxicological impacts that PFASs may have 434 on sea turtle development.

435

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695 Table 1. Samples sizes and mass fractions (ng/g wet mass) of predominant perfluoroalkyl

696 substances (PFASs) in plasma of North Pacific green and hawksbill sea turtles.

697

	Green turtle	Plasma	Kapoho (live captured)	2011-2012	39.2 - 83.4	13	0.407	0.796	0.734	0.196 - 2.19	100	0.407	0.796	0.734	0.196-2.19	100	
	Green turtle	Plasma	Kailua (live captured)	2011-2012	42.9 - 80.3	13	2.11	2.45	1.32	0.865 - 4.52	100	2.11	2.45	1.32	0.865-4.52	100	
	Green turtle	Plasma	MHI FP stranded	2011-2012	35.5 - 70.2	14	0.242	0.902	1.04	<0.054 - 2.77	85.7	0.242	0.816	0.912	<0.054-2.52	85.7	
	Green turtle	Plasma	CNMI	2013		12	1.74	0.562	1.12	0.141 - 4.11	100	0.634	0.524	0.987	0.141-3.65	100	
	Green turtle	Plasma	Saipan	2013	46.3 - 85.6	6	0.295	0.926	1.56	0.175 - 4.11	100	0.295	0.850	1.37	0.175-3.65	100	
	Green turtle	Plasma	Tinian	2013	47.7 - 63.7	6	0.204	0.199	0.045	0.141 - 0.270	100	0.204	0.199	0.045	0.141-0.27	100	
	Green turtle	Plasma	Palmyra Atoll	2012-2013	44.3 - 87.6	10	0.155	0.171	0.051	<0.085 - 0.265	80	0.155	0.171	0.051	<0.085-0.265	80	
	Green turtle	Plasma	Core Sound, NC, USA (Keller et al., 2012	2006	within 25-70	10	2.92	3.39	1.74	0.871 - 6.09	100	1.12	2.41	2.37	0.871 - 3.87	100	
	Hawksbill turtle	Plasma	CNMI	2013	35.1 - 48.3	4	1.74	1.99	0.951	1.19 - 3.29	100	0.634	0.654	0.285	0.334-1.01	100	0.972
	Hawksbill turtle	Plasma	Palmyra	2013	54.4 - 55.5	2		4.18	N/A	3.373 - 4.978	100		0.300	N/A	0.253-0.348	100	
	Hawksbill turtle	Plasma	Juno Beach, FL, USA (Keller et al., 2012)	2006	within 25-70	5	33.5	44.1	23.5	24.8 - 79.0	100	11.9	11.9	6.27	5.45 - 21.2	100	17.0
8	Hawksbill turtle	Egg contents	MHI	2012	unknown	7*	31.0	35.0	34.1	5.46 - 106	100	2.92	16.3	34.2	0.862 - 93.8	100	1.28

- 699 Abbreviations: Commonwealth of the Northern Marianas Islands (CNMI), Main Hawaiian
- 700 Islands (MHI), North Carolina (NC), Florida (FL), % > R = percent of samples
- above the reporting limit.
- 702 *Eleven nests from seven mothers were analyzed. Multiple nests from the same mom were
- 703 averaged before the summary statistics shown here were calculated for all seven moms.
- 704 NC and FL data taken from Keller et al. (2012).

706 Table 2. Estimated margins of safety (EMOS) for average/maximum sea turtle exposure to

707 perfluorooctane sulfonate based on the lowest adverse effect levels in lab-exposed animals.

708

		LOEAL (ng/g or ng/mL)	85000	1500	2290	1000	č
		LOEAL in	serum	serum	serum	serum	se
		LOEAL in	rat/mouse	rat	rat	zebrafish	m
			Bjork et al 2008;				Peden-A
		Reference	Lau et al 2003	Currane et al 2008	Wanget al 2011	Zhang et al 2011	2
Species	Tissue	Turtle Grouping		Estimate	ed margin of safety f	or average / maxim	um turtle
Green turtle	Plasma	MHI Live captured, no FP stranded	74561 / 18805	1316 / 332	2009 / 507	877 / 221	80*
Green turtle	Plasma	Kiholo (live captured)	464481 / 151786	8197 / 2679	12514 / 4089	5464 / 1786	500
Green turtle	Plasma	Kapoho (live captured)	106784 / 38813	1884 / 685	2877 / 1046	1256 / 457	115
Green turtle	Plasma	Kailua (live captured)	34694 / 18805	612 / 332	935/507	408/221	37*
Green turtle	Plasma	MHI FP stranded	104167 / 33730	1838 / 595	2806 / 909	1225/397	112
Green turtle	Plasma	CNMI	162214 / 23288	2863 / 411	4370 / 627	1908 / 274	175
Green turtle	Plasma	Palmyra Atoll	497076 / 320755	8772 / 5660	13392 / 8642	5848 / 3774	53!
Hawksbill turtle	Plasma	CNMI	129969 / 84158	2294 / 1485	3502 / 2267	1529/990	140
Haw ksbill turtle	Plasma	Palmyra	283333 / 244253	5000 / 4310	7633 / 6580	3333 / 2874	305
Haw ksbill turtle	Egg contents	мні					

710

709

* concern of risk when EMOS <100, ** heightened concern of risk when EMOS <10.



- 713 Figure 1. Green and hawksbill sea turtle sampling sites and sample sizes. Dots show capture sites
- of free-ranging green (green) and hawksbill (purple) turtles and nest excavation sites of
- 715 hawksbill nests (yellow). Fourteen stranded green turtles from MHI severely afflicted with FP
- 716 plasma are not mapped.



717

Figure 2. PFAS profiles detected in Pacific Island sea turtles. MHI = Main Hawaiian Islands,
CNMI = Commonwealth of the Northern Marianas Islands. *All green turtles from MHI, CNMI
and Palmyra were combined excluding the MHI FP-stranded green turtles. Perfluoroalkyl
sulfonates (blue shades) are visually distinguished from perfluoroalkyl carboxylates (other
colors).







726 the Northern Marianas Islands (CNMI), Main Hawaiian Islands (MHI), North Carolina (NC),

- 727 Florida (FL). Sample sizes were 10, 12, 39, and 10 green turtles, and 2, 4, and 5 hawksbills,
- 728 respectively. NC and FL PFOS data are from Keller et al. (2012). Population data are from

- 729 O'Connell et al. (2010) for NC, and summed for counties in the St. Lucie-Loxahatchee
- 730 Watershed (US Census Bureau 2019b, c) for FL. Stranded green turtles with severe
- fibropapillomatosis were excluded from the MHI data. The asterisk indicates a difference in
- 732 PFOS concentration from other Pacific sites p < 0.05).



Figure 4. PFAS quantities (ng) estimated in the entire clutch of eggs from five clutches laid by
hawksbill turtle 19591-04 on Maui. Linear regression statistics for ng vs. clutch order are shown
for each compound.





738 Figure 5. Significant correlations between perfluoroundecanoate (PFUnA) (A) and

739 perfluorotridecanoate (PFTriA) (B) concentrations (ng/g) in eggs from 11 hawksbill sea turtle

740 nests from the Main Hawaiian Islands and emergence success.

Table 1. Samples sizes and mass fractions (ng/g wet mass) of predominant perfluoroalkyl substances (PFASs) in plasma of North Pacific green and

hawksbill sea turtles.

							ТОТ	TAL PFA	ASs	1			PFOS			PFNA					
				SCL range					Min-					Min-					Min-		
Species	Tissue	Grouping	Year	(cm)	n	Median	Mean	SD	Max	%>RL	Median	Mean	SD	Max	%>RL	Median	Mean	SD	Max	%>RL	
Green		MHI Live captured, no FP	2011-	35.5 -					<0.063			£		<0.063-					<0.188		
turtle	Plasma	stranded	2012	83.4	39	0.447	1.14	1.29	- 4.52	92.3	0.447	2.14	1.29	4.52	92.3				<1.18	0	
Green turtle	Plasma	Kiholo (live captured)	2011- 2012	43.7 - 65.1	13	0.126	0.183	0.121	<0.063 - 0.560	76.9	Q 126	0.183	0.121	<0.063- 0.560	76.9				<0.188 - <1.03	0	
Green turtle	Plasma	Kapoho (live captured)	2011- 2012	39.2 - 83.4	13	0.407	0.796	0.734	0.97 - 2.13	100	0.407	0.796	0.734	0.196- 2.19	100				<0.239 - <0.893	0	
Green turtle	Plasma	Kailua (live captured)	2011- 2012	42.9 - 80.3	13	2.11	2.45	1.52	0.865 - 4.52	100	2.11	2.45	1.32	0.865- 4.52	100				<0.233 - <1.18	0	
Green turtle	Plasma	MHI FP stranded	2011- 2012	35.5 - 70.2	14	0.242	0.902	1.04	<0.054 - 2.77	85.7	0.242	0.816	0.912	<0.054- 2.52	85.7				<0.162 - <1.21	0	
Green turtle	Plasma	CNMI	2013		12	1.74	0.562	1.12	0.141 - 4.11	100	0.634	0.524	0.987	0.141- 3.65	100				<0.301 - <0.377	0	
Green turtle	Plasma	Saipan	2013	46.3 - 85.6	6	0.295	0.926	1.56	0.175 - 4.11	100	0.295	0.850	1.37	0.175- 3.65	100				<0.326 - <0.377	0	
Green turtle	Plasma	Tinian	2013	47.7 - 63.7	6	0.204	0.199	0.045	0.141 - 0.270	100	0.204	0.199	0.045	0.141- 0.27	100				<0.301 - <0.369	0	
Green turtle	Plasma	Palmyra Atoll	2012- 2013	44.3 - 87.6	10	0.155	0.171	0.051	<0.085 - 0.265	80.0	0.155	0.171	0.051	<0.085- 0.265	80.0				<0.273 - <8.70	0	
Green turtle	Plasma	Core Sound,	2006	25 - 70	10	2.92	3.39	1.74	0.871 - 6.09	100	1.12	2.41	2.37	0.871 - 3.87	100				<0.070 -	10	

		NC, USA (Keller et al., 2012)																	0.182	
Hawksbill				35.1 -					1.19 -					0.334-					0.582	
turtle	Plasma	CNMI	2013	48.3	4	1.74	1.99	0.951	3.29	100	0.634	0.654	0.285	1.01	100	0.972	1.08	0.516	- 1.79	100
Hawksbill turtle	Plasma	Palmyra	2013	54.4 - 55.5	2		4.18	N/A	3.373 - 4.978	100		0.300	N/A	0.253- 0.348	100		3.40	N/A	2.90 - 3.90	100
Hawksbill turtle	Plasma	Juno Beach, FL, USA (Keller et al., 2012)	2006	25 - 70	5	33.5	44.1	23.5	24.8 - 79.0	100	11.9	11.9	6.27	5.45 - 21.2	100	17.0	17.3	1.21	3.87 - 30.8	100
Hawksbill	Egg								5.46 -					0.862 -					0.756	
turtle	contents	MHI	2012	unknown	7*	31.0	35.0	34.1	106	100	2.92	16.3	34.2	93.8	100	1.28	7.44	8.90	- 22.8	100

Abbreviations: Commonwealth of the Northern Marianas Islands (CNMI), Main Hawaiian Islands (MHI), North Carolina (NC), Florida (FL), % >R

= percent of samples

above the reporting limit.

*Eleven nests from seven mothers were analyzed. Multiple nests from the same mom were averaged before the summary statistics shown here were

calculated for all seven moms.

NC and FL data taken from Keller et al. (2012).

Table 2. Estimated margins of safety (EMOS) for average/maximum sea turtle exposure to perfluorooctane sulfonate based on the lowest adverse

effect levels in lab-exposed animals.

			Neonate mortality,		Altered	Altered	Decreased T-cell dependent IgM	Decreased
			altered liver histology	Increased	thyroid	development of	antibody response	hatching
		Adverse effect	and gene expression	liver weight	hormones	motor neurons	(immunosuppression)	success
		LOEAL (ng/g or						
		ng/mL)	85000	1500	2290	1000	91.5	100
		LOEAL in	serum	serum	serum	serum	serum	eggs
		LOEAL in	rat/mouse	rat	rat	zebrafish	mouse	chicken
			Bjork et al 2008; Lau	Currane et	Wang et al	ن ک		Molina et al
		Reference	et al 2003	al 2008	2011	Zhang et al 2011	Peden-Adams et al 2008	2006
		Turtle				\mathcal{O}		
Species	Tissue	Grouping		Estir	nated marg	safety for average /	' maximum turtle	
		MHI Live			0	•		
Green		captured, no FP						
turtle	Plasma	stranded	74561 / 18805	1316 / 332	2009 / 507	877 / 221	80* / 20*	
Green		Kiholo (live						
turtle	Plasma	captured)	464481 / 151786	8197 / 2	12514 / 4089	5464 / 1786	500 / 163	
Green		Kapoho (live						
turtle	Plasma	captured)	106784 / 38813	1884/685	2877 / 1046	1256 / 457	115 / 42*	
Green		Kailua (live		\sim				
turtle	Plasma	captured)	34694 / 18805	612 / 332	935 / 507	408 / 221	37* / 20*	
Green		MHI FP	J					
turtle	Plasma	stranded	104167 / 33730	1838 / 595	2806 / 909	1225 / 397	112 / 36*	
Green								
turtle	Plasma	CNMI	162214 / 23288	2863 / 411	4370 / 627	1908 / 274	175 / 25*	
Green								
turtle	Plasma	Palmyra Atoll	497076 / 320755	8772 / 5660	13392 / 8642	5848 / 3774	535/ 345	
Hawksbill								
turtle	Plasma	CNMI	129969 / 84158	2294 / 1485	3502 / 2267	1529 / 990	140 / 91*	
Hawksbill								
turtle	Plasma	Palmyra	283333 / 244253	5000 / 4310	7633 / 6580	3333 / 2874	305 / 263	
Hawksbill	Egg							
turtle	contents	MHI						6**/1**

* concern of risk when EMOS <100, ** heightened concern of risk when EMOS <10.

Highlights

- Perfluorinated alkyl substances were detected in sea turtles from the North Pacific
- PFAS levels in sea turtles were correlated with human population and land use
- Maternal offloading of PFASs was detected in unhatched hawksbill eggs from Hawaii
- Levels in eggs were near those which have caused developmental toxicity in birds
- No correlation was found between fibropapillomatosis (FP) and PFAS concentrations

1	Supplemental Information for
2	Sea turtles across the North Pacific are exposed to perfluoroalkyl substances
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35 Detailed methods of laboratory analysis and data analysis

36 PFAS Measurements

Each plasma and egg sample was measured for concentrations of 15 PFASs according to 37 methods modified from (Keller et al. 2012). The calibration series consisted of NIST Reference 38 39 Materials 8446 Perfluorinated Carboxylic Acids and Perfluorooctane Sulfonamide in Methanol 40 and 8447 Perfluorinated Sulfonic Acids in Methanol. The internal standard (IS) solution 41 consisted of 11 mass-labeled PFASs purchased from Wellington Laboratories (Guelph Ontario, 42 Canada), RTI International (Research Triangle Park, NC, USA) and Cambridge Isotope Labs 43 (Tewksbury, MA, USA) gravimetrically combined in methanol. Compound names, 44 abbreviations, and IS choices are in Table S3. As part of the BEMAST protocols, three field blanks were prepared by pulling Millipore water stored in a polyethylene bottle through the same 45 lot number of blood collection needles, tubes, pipets, and storage vials. Control materials were 46 47 five replicates of SRM 1957 Organic Contaminants in Non-Fortified Human Serum (≈1 g each) and three replicates of SRM 1947 Lake Michigan Fish Tissue (\approx 1 g each). Plasma samples and 48 49 field blanks (≈ 1 g each) archived in plastic cryovials were thawed, mixed, transferred to glass 50 culture tubes gravimetrically, and 0.6 mL of IS solution was added gravimetrically to provide ≈ 3 ng of each mass-labeled compound to each sample. The calibration curve consisted of six tubes 51 with gravimetric dilutions from ≈ 0.05 ng to 275 ng of each compound. The IS solution and 1 g of 52 53 Millipore water were added to calibrants, and to three laboratory blanks. Tubes were vortexed, 54 and equilibrated at room temperature for 1.5 h or longer. Samples were extracted with 4 mL of 55 acetonitrile and 30 min of sonication. The supernatant, after centrifuging the tubes at 157 rad/s 56 (1500 rpm) for 5 min, was transferred to clean glass tubes. SRM 1957 replicates were processed 57 the same way.

58	Egg samples archived in plastic cryovials were thawed (≈ 1 g each), transferred to glass
59	tubes gravimetrically, centrifuged at 157 rad/s for 5 min, and IS solution (0.6 mL) was added
60	gravimetrically to provide \approx 3 ng of each mass-labeled compound to each sample. Tubes were
61	vortexed for 20 s and allowed to equilibrate for 1.5 h or longer. Samples were extracted with 6
62	mL of acetonitrile and 20 s of vortex and 30 min of sonication. The supernatant, after
63	centrifuging the tubes at 157 rad/s for 5 min, was transferred to clean glass tubes. Samples were
64	extracted a second time with 6 mL acetonitrile, and extracts were combined with the first
65	extraction. After freezing, the extracts were no longer cloudy. The precipitate was discarded after
66	centrifugation at 157 rad/s for 5 min and transferring the supernatant to a clean culture tube.
67	SRM 1947 replicates were processed the same way.
68	Extracts were solvent exchanged to methanol using nitrogen evaporation being careful to
69	not evaporate to dryness. They were purified using 250 mg Envi-carb columns (Supelco,
70	Bellefonte, Pennsylvania) by loading the 3 mL sample, a 1 mL sample rinse of methanol, and 6
71	mL of methanol. The extracts were concentrated under nitrogen, and transferred to glass
72	autosampler vials for a final volume of 0.5 mL in methanol. Samples were analyzed using a
73	liquid chromatograph (Agilent 1100 HPLC, Palo Alto, CA) interfaced to a negative electrospray
74	ionization tandem mass spectrometer (API 4000, Applied Biosystems-MDS Sciex, Foster City,
75	CA). Samples (20 μ L) were injected onto an analytical column (Agilent Zorbax Eclipse Plus
76	C18, 150 mm x 2.1 mm x 5 μ m). The solvent gradient at a flow rate of 0.25 mL/min (all
77	mixtures expressed in volume fractions) started at 50 $\%$ methanol and 50 $\%$ 20 mmol/L
78	ammonium acetate in water and increased to 75% methanol by 20 min, held for 5 min, and then
79	increased to 95 % methanol by 35 min, held for 5 min, before reverting back to original
80	conditions at 40.5 min with a 9.5 min hold time. The MS/MS method was divided into six

81 windows and two of the most abundant transitions for each PFAS were monitored. The internal 82 standard approach was used to quantify each compound amount. Amounts of each analyte were 83 calculated using the slope and y-intercept of at least a three point calibration curve that bracketed 84 the peak area ratios observed in the samples. Concentrations were determined by dividing the 85 calculated analyte mass by the extracted sample mass. Reporting limits (RLs) were determined 86 using the highest calculated RL of two methods described in O'Connell et al. (2010). 87 Concentrations are totals of linear and branched isomers.

88 Data Analyses

89 JMP statistical software (SAS Institute Inc. Cary, NC) was used when all samples were >RLs, and on occasion when required, with values <RL substituted with half the RL. When 90 91 some samples were <RLs, every attempt was made to use the R software package, NADA, as suggested by Helsel (2005). Means, medians and standard deviations were calculated with either 92 Kaplan-Meier or regression on order statistical models. Significance was determined as p < 0.05. 93 94 Data normality and homoscedasticity were tested with Shapiro-Wilk W tests and O'Brien or 95 Bartlett tests, respectively. If assumptions were violated even with transformed data, non-96 parametric tests were used. The cendiff function (non-parametric ANOVA) in R NADA was 97 used to test significant differences among sites for green sea turtle plasma PFAS concentrations, 98 followed by the Steel Dwass non-parametric multiple comparison test using JMP. The Wilcoxon 99 t-test in JMP was used to compare hawksbill plasma PFOS, PFNA, and Σ PFAS plasma 100 concentrations between Palmyra Atoll and CNMI, while the cendiff function was necessary for 101 PFUnA. Species differences were assessed only within CNMI data. PFOS and Σ PFAS plasma 102 concentrations between species could be compared using Wilcoxon tests in JMP, and PFNA and 103 PFUnA were tested with cendiff function. MHI green turtle plasma samples were grouped

according to location and FP incidence (same as in Keller et al. (2014)). Differences among
groups in PFOS concentrations was assessed with the cendiff function in R NADA, followed by
Steel-Dwass multiple comparison tests in JMP.

Amount of PFASs offloaded into each nest was estimated by multiplying the total number eggs laid by the average grams of each pooled egg contents from that nest, and by the measured PFAS concentration (in ng/g). The nanograms of PFASs offloaded were then compared across clutches laid within one nesting season from a single mother using linear regressions. PFAS concentrations in the eggs were compared to nest success variables using the cenken function in R NADA, a version of Kendall's tau correlation.

To assess toxicological risk, estimated margins of safety (EMOS) were calculated as described in Keller et al. (2012), as the ratio of the average plasma, serum, or egg PFOS concentration in laboratory-exposed animals in the lowest observed adverse effect level (LOAEL) dose group to either the average or maximum sea turtle plasma or egg PFOS concentration. EMOS below 100 were considered of concern (Faustman and Omenn 1996).

118 Palmyra Atoll Spatial Comparison

119 If local sources released PFASs at Palmyra Atoll, which are unexpected, they could be 120 from runoff/sewage from the scientific field station or non-military use of AFFFs on the airstrips 121 from Cooper Island in the northwestern portion of the atoll. Surface currents nearshore and 122 within the lagoon flow westward (Gardner et al. 2011), so we hypothesized that turtles sampled 123 from the lagoon or west would have greater PFAS concentrations. Sample sizes prevented 124 statistical testing, but a cursory comparison was attempted of green turtle PFAS concentrations 125 on the four sides and within the lagoon to investigate the possibility of a point source on the 126 atoll. Σ PFASs in green turtles were greatest in a turtle (MT13W003) from the east (0.265 ng/g)

and least in the turtle captured twice (MT12W064 and MT13W014, adult female) in the lagoon 127 (<0.088 ng/g and 0.155 ng/g), refuting the hypothesis. No evidence of a significant PFAS point 128 129 source on Palmyra Atoll was observed. The small difference in PFAS concentrations between the 130 eastern and lagoon green turtles is likely related to life stage. The green turtle in the east was 131 only 44.3 cm straight carapace length, indicating recent recruitment to Palmyra. The lagoon 132 green turtle was an adult female and was captured twice over two years, suggesting residency for 133 at least a year. Green turtles undergo an ontogenetic shift from a carnivorous pelagic juvenile 134 phase to the herbivorous benthic juvenile phase when they recruit into nearshore habitats 135 (Bjorndal 1997). Because of their higher trophic status in the pelagic phase, green turtles may 136 recruit into Palmyra carrying a greater PFAS burden, which dilutes with growth through the next benthic herbivorous phase (Keller 2013). The lesser concentrations in the adult female in the 137 lagoon may have resulted from her ability to offload PFASs to eggs. The eastern region of 138 139 Palmyra is known habitat for the smallest, recently recruited green turtles from the pelagic-phase (Sterling et al. 2013) and green turtles in this region may enter with greater PFAS concentrations. 140 141 Spatial trends across the atoll are confounded by these life history traits. The data in the current 142 study suggest that remote Palmyra Atoll is a control reference site for PFAS with little to no 143 local point sources. McFadden et al. (2014) was the first study to measure chemical contaminants 144 (heavy metals) in Palmyra sea turtles and found elevated aluminum and iron.







154 CNMI and Palmyra were combined, excluding the MHI FP-stranded green turtles.

Figure S2. Mean and one standard deviation of PFOS concentrations (ng/g) in plasma of four
green turtle groups from MHI related to fibropapillomatosis (FP) and urban development
percentage of each watershed. Mean PFOS concentrations increased with watershed
urbanization, with the highest being in Kailua, Hawai'i, rather than with FP incidence. FP
incidence is noted by more red colored bars. Black circles indicate % urbanization for each
Hawaiian watershed (Parham et al. 2008a; b; c). Different letters above bars indicate a significant
difference between groups (p<0.05).



163

Figure S3. Mean and one standard deviation PFAS concentrations (ng/g) in green and hawksbill
sea turtle plasma from CNMI. Patterned bars indicate that means fell below the detection

166 limits. Sample sizes are 12 and 4. * indicates difference between species (p<0.05).



167

168 Figure S4. Σ PFAS concentrations (ng/g wet mass) in hawksbill eggs in relation to the

169 approximate distance (km) between the nests and the nearest international airport. When females

170 laid multiple clutches, the average $\sum PFAS$ concentrations of all clutches from that female were

171 used. Pearson correlation statistics are shown.





173 174 Figure S5. PFAS profiles in egg contents of nests from individual hawksbill sea turtles laid in

different Hawaiian Island locations. 175

176 a.



180 Figure S6. Hawksbill sea turtle 19591-04's a) post-nesting migrations, satellite tracked in 2004

- 182 (2009)) and again in 2008 (yellow arrows) from Makena, Maui, to foraging grounds near
- 183 Kahuku, Oahu, and b) dive behavior profile for the 2008 track.

^{181 (}white dots, black line and arrows; base map modified from Figure 3C within Parker et al.

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Declaration of interests

x The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

□The authors declare the following financial interests/personal relationships which may be considered as potential competing interests: