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# <sup>1</sup> Perfluoroalkyl Acid Uptake in Lettuce (*Lactuca sativa*) and Strawberry <sup>2</sup> (*Fragaria ananassa*) Irrigated with Reclaimed Water

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8 Supporting Information

ABSTRACT: Using reclaimed water to irrigate food crops presents an 9 exposure pathway for persistent organic contaminants such as perfluoroalkyl 10 acids (PFAAs) to enter the human food chain. This greenhouse study used 11 reclaimed water augmented with varying concentrations (0.2-40  $\mu$ g/L) of 12 PFAAs, including perfluorocarboxylates  $(C_3F_7COO^- \text{ to } C_8F_{17}COO^-)$  and 13 perfluorosulfonates (C<sub>4</sub>F<sub>9</sub>SO<sub>2</sub>O<sup>-</sup>, C<sub>6</sub>F<sub>13</sub>SO<sub>2</sub>O<sup>-</sup>, C<sub>8</sub>F<sub>17</sub>SO<sub>2</sub>O<sup>-</sup>), to investigate 14 potential uptake and concentration-response trends in lettuce (Lactuca sativa) 15 16 and strawberry (Fragaria ananassa). In addition, studies were conducted to evaluate the role of soil organic carbon concentrations on plant uptake of 17 PFAAs. PFAA concentrations in lettuce leaves and strawberry fruit were 18 measured for each incremental aqueous PFAA concentration applied. PFAA 19 plant concentrations increased linearly with the aqueous concentration for all 20 PFAAs, with PFCAs bioaccumulating to a greater degree than PFSAs in the 2.1 edible portions of the tested plants. Chain-length-dependency trends were 22



evident in both lettuce shoot and strawberry fruit, with decreasing concentrations associated with increasing chain length. 23 Perfluorobutanoate (PFBA) and perfluoropentanoate (PFPeA), both short-chain PFAAs (<8 carbon chain length), accumulated 24 the most compared with other PFAAs tested in the edible parts of both lettuce and strawberry. PFAA concentrations in 25 strawberry root and shoot were also measured at selected PFAA aqueous concentrations (0.4, 4, and 40  $\mu$ g/L). Short-chain 26 27 perfluorocarboxylates were the dominant fraction in the strawberry fruit and shoot compartments, whereas a more even distribution of all PFAAs appears in the root compartment. Lettuce grown in soils with varying organic carbon contents (0.4%, 28 2%, 6%) was used to assess the impact of organic carbon sorption on PFAA bioaccumulation. The lettuce grown in soil with the 29 6% organic carbon content had the lowest bioaccumulation of PFAAs. Bioaccumulation factors for lettuce were correlated to 30 carbon chain length of PFAAs, showing approximately a 0.4 to 0.6 log decrease per CF<sub>2</sub> group. This study confirms that PFAAs 31 can enter and bioaccumulate in food crops irrigated with reclaimed water. Bioaccumulation potential depends on analyte 32 functional group and chain length, concentration in the reclaimed water, and organic carbon content of the soil. 33

## 34 INTRODUCTION

<sup>35</sup> Perfluoroalkyl acids (PFAAs) are ubiquitous synthetic chem-<sup>36</sup> icals that are widely used in both consumer and industrial <sup>37</sup> settings and that have attracted much attention with regard to <sup>38</sup> their persistent, accumulative, and toxic nature.<sup>1,2</sup> The extensive <sup>39</sup> use of PFAAs in consumer products means that municipal <sup>40</sup> wastewaters are a collection vehicle for the compounds. In <sup>41</sup> addition, the prominence of PFAAs in some manufacturing <sup>42</sup> processes can lead to high levels in industrial wastewaters that <sup>43</sup> also flow to wastewater treatment plants (WWTPs).<sup>3</sup> Most <sup>44</sup> conventional WWTPs are ineffective at removing PFAAs<sup>3,4</sup> <sup>45</sup> and, thus, may represent significant sources of PFAA releases <sup>46</sup> into the environment.<sup>5</sup> Unlike many organic contaminants, the <sup>47</sup> dual hydrophobic/lipophobic nature of PFAAs enables the <sup>48</sup> compounds to reside in significant quantities in both the <sup>49</sup> aqueous and sludge effluent streams of WWTPs.<sup>4,6</sup> The aqueous effluent stream of a WWTP is, in general, <sup>50</sup> returned to the surrounding aquatic environment; however, <sup>51</sup> growing water scarcity is driving alternative uses of treated <sup>52</sup> wastewater. In particular, interest in the use of recycled or <sup>53</sup> reclaimed water, which typically consists of municipal waste- <sup>54</sup> water (treated to remove pathogens, organic matter, and <sup>55</sup> nutrients), for agricultural purposes is growing and is likely to <sup>56</sup> continue in the future.<sup>7</sup> Reclaimed water has been safely used <sup>57</sup> for many years in the U.S. for the irrigation of nonfood crops<sup>8</sup> <sup>58</sup> and, on a more limited scale, for food crops eaten raw (e.g., in <sup>59</sup> the Salinas Valley, CA). Recently, however, concerns have been <sup>60</sup>

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<sup>61</sup> raised regarding the presence of chemicals of emerging concern <sup>62</sup> in reclaimed water.<sup>9</sup>

Although the U.S. Environmental Protection Agency has 4 published guidelines for water reuse, no federal regulations 5 govern water reclamation and reuse in the U.S., and thus, 6 regulations or guidelines have been developed at the state 7 level.<sup>8</sup> This nonunified approach has resulted in differing 8 standards among states that have developed reuse criteria. 9 Existing water reuse regulations for food crop irrigation in each 70 state vary according to crop type and irrigation method, but are 71 principally directed at health protection from microbial 72 pathogens and do not typically include requirements addressing 73 organic contaminants.<sup>10</sup>

The potential risks associated with bioaccumulation of 74 75 organic contaminants are most easily studied in edible crops 76 eaten raw since processing and cooking can add confounding 77 factors, such as chemical transformation, chemical volatilization, 78 or additional contamination by cookware or packaging.<sup>11</sup> 79 Unfortunately, although data on the occurrence of many 80 contaminants in reclaimed water are plentiful,<sup>12-14</sup> limited data 81 exist on the potential for uptake of PFAAs from reclaimed 82 water into edible plants. To date, human health risk 83 assessments are based on plant uptake models primarily 84 developed for neutral organic chemicals in general and are 85 limited to crop-specific data.<sup>15,16</sup> PFAAs exhibit surfactant 86 behavior, and thus, octanol-water partitioning coefficients used 87 in traditional bioaccumulation modeling are not applicable to 88 this class of compounds; instead, PFAA chain length is a better 89 proxy for hydrophobicity.<sup>17</sup>

A few studies have demonstrated the potential for crop 90 91 uptake of pharmaceuticals applied via real or simulated 92 wastewater;<sup>18-21</sup> however, the behavior of the contaminants 93 studied is very different from that of PFAAs, particularly in 94 regard to charge. Felizeter et al.<sup>22</sup> reported uptake of PFAAs in 95 lettuce plants via hydroponic solution, with higher concen-96 trations of the short-chain (<8 carbon chain length) PFAAs 97 accumulating in the leaves. However, fundamental differences between hydroponic and solid media experiments as well as 98 99 differences in water quality between nutrient solutions and 100 actual reclaimed water prevent direct applicability of these data 101 to crops irrigated with reclaimed water and grown in soil. Blaine 102 et al.<sup>23</sup> examined lettuce uptake of PFAAs from biosolids-103 amended soils and also found preferential short-chain 104 accumulation in the lettuce leaves, as well, although since the 105 bioavailability of PFAAs for uptake may vary considerably 106 depending on the uptake matrix, these data may also have 107 limited applicability to crop uptake of PFAAs via reclaimed 108 water.

This study was conducted to examine the uptake of PFAAs in 109 110 lettuce (Lactuca sativa "Multy") and strawberry (Fragaria ananassa "Albion") via reclaimed water under conditions 111 112 representative of current agricultural practices. Experiments were carried out using reclaimed water augmented with varying 113 concentrations of PFAAs. The intent of this research was to 114 elucidate PFAA accumulation potential in response to varying 115 116 concentrations of PFAAs in reclaimed irrigation water. Lettuce 117 and strawberry crops were chosen to represent typical food 118 crops grown in the U.S. using reclaimed water. In addition, 119 lettuce grown in soils with varying organic carbon (OC) 120 content was used to assess the impact of OC sorption on PFAA 121 bioaccumulation given the propensity of PFAAs to sorb to 122 OC.<sup>17</sup>

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## MATERIALS AND METHODS

Chemicals. All calibration standards and stable isotopes 124 were acquired from Wellington Laboratories (Guelph, ON, 125 Canada) and prepared using established protocols.<sup>23</sup> Specific 126 PFAAs used in this study include perfluorobutanoate (PFBA), 127 perfluoropentanoate (PFPeA), perfluorohexanoate (PFHxA), 128 perfluoroheptanoate (PFHpA), perfluorooctanoate (PFOA), 129 perfluorononanoate (PFNA), perfluorobutanesulfonate 130 (PFBS), perfluorohexanesulfonate (PFHxS), and perfluorooc- 131 tanesulfonate (PFOS); PFAAs and corresponding surrogate 132 standards are listed in Table S1. Spiking solutions for dosing 133 experiments were prepared from individual standards pur- 134 chased from Sigma-Aldrich (St. Louis, MO). High purity 135 Chromasolv dichloromethane, HPLC-grade methanol, and all 136 other reagent grade solvents were acquired through Sigma- 137 Aldrich. Water for extractions was obtained from a Milli-Q 138 system (Millipore, Billerica, MA), whereas HPLC-grade water 139 was utilized for liquid chromatography tandem mass spectrom- 140 etry (LC-MS/MS) analysis. Extraction cleanup was facilitated 141 with Chromabond diamino from Macherey-Nagel Inc. 142 (Bethlehem, PA) and Supelclean ENVI-Carb from Sigma- 143 Aldrich. 144

**Greenhouse Study.** Plant uptake experiments were 145 conducted in a climate controlled greenhouse with two food 146 crops, leaf lettuce (*L. sativa* "Multy") and strawberry (*F.* 147 *ananassa* "Albion"). These selected cultivars are similar to 148 cultivars currently grown in the western U.S. using reclaimed 149 water. Five replicate plants were grown for each set of 150 experimental conditions. Pots (15 cm diameter) were randomly 151 arranged to account for any spatial variations in light and 152 temperature within the greenhouse. Day temperatures ranged 153 from 18 to 21 °C, and night temperatures ranged from 10 to 13 154 °C. Full spectrum, metal halide, and high pressure sodium 155 supplemental lighting (Plantmax 1000W bulbs) was also 156 supplied to achieve 16 h of daylight to mimic field conditions. 157 Additional information regarding plant propagation and 158 cultivation can be found in the Supporting Information (SI). 159

Reclaimed water was supplied by the Mines Park pilot-scale 160 Sequencing Batch/Membrane Bioreactor. This test site at 161 Colorado School of Mines treats raw sewage from a student 162 apartment complex (a graduate housing community of  $\sim$ 400 163 individuals); a full description of the site can be found 164 elsewhere.<sup>24</sup> Although the effluent from Mines Park was not 165 specifically tailored to meet the reclaimed water regulations of 166 any particular region, the water represented a steady and 167 realistic source of reclaimed water for the experiments (SI 168 Table S3). Concentration-dependent accumulation was exam- 169 ined by spiking the reclaimed water with eight levels of PFAAs 170  $(0.2, 0.4, 1, 2, 4, 10, 20, 40 \,\mu g/L)$  in addition to using ambient 171 reclaimed water and tap water (control). This concentration 172 range was chosen to give a range of values starting with ambient 173 concentrations found in the Mines Park reclaimed water 174 (~0.02  $\mu$ g/L), bracketing typical WWTP effluent concen- 175 trations  $(0.2-4 \ \mu g/L)$ <sup>5</sup> and reaching concentrations repre- 176 sentative of contaminated groundwater  $(10-40 \ \mu g/L)$ .<sup>25,26</sup> To 177 assess the accuracy of the actual irrigation solutions as opposed 178 to the nominal concentrations, aliquots from each solution 179 were analyzed. The lowest recoveries were for PFNA (36%) 180 and PFOS (23%), the strongest sorbing analytes, most likely as 181 a result of losses onto the walls of the watering containers. The 182 linearity of the aqueous concentrations remained fairly constant 183 for each analyte, with each applied concentration being 1.5-3 184



Figure 1. Concentrations of PFCAs (a) and PFSAs (b) in lettuce leaves versus measured aqueous concentration of PFAAs. Means and standard errors (n = 5) are shown.

185 times the next lowest concentration. Both the tap water control 186 and ambient reclaimed water had trace detections (<25 ng/L) 187 of PFHxA, PFOA, and PFOS. Measured concentrations of 188 PFAAs (SI Table S2) were used in all calculations. Plants 189 received PFAAs via hand watering three times a week, 100 mL 190 of solution per lettuce plant and 200 mL of solution per 191 strawberry plant. Additional information on the reclaimed water 192 quality is found in SI Table S3.

Because soil organic matter can significantly impact the 193 <sup>194</sup> bioavailability of PFAAs,<sup>27</sup> a prepared sandy soil mix (3:1 sand/ topsoil by mass) with only 0.4% OC was used to represent a 195 'worst-case" scenario in terms of bioavailability. Plant essential 196 197 nutrients were supplied by mixing a single application of slow release Osmocote (nitrogen-phosphorus-potassium: 19-6-198 12) into the media ( $\sim$ 5 g/plant). To more specifically test the 199 impacts of OC on PFAA uptake, lettuce was grown in two 2.00 additional soils with varying OC content (2%, 6%) at a single 201 202 PFAA concentration (10  $\mu$ g/L). Details on the soils utilized are provided in the SI. 203

Edible portions of lettuce and strawberry plants were harvested at maturity. In addition, after sufficient strawberry fruit biomass was harvested, whole strawberry plants were collected and separated into root and shoot portions. All plant material was frozen (-20 °C) in PFAA-free plastic bags prior to analysis. Additional details concerning soil and produce sampling are found in the SI.

Sample Extraction and Data Analysis. Homogenized 211 212 plant samples (0.5-2 g) were prepared and extracted using the 213 protocol from Blaine et al.<sup>23</sup> Lettuce shoots from all 214 experimental replicate pots (n = 3-5) were extracted 215 independently, and concentrations were averaged. Ripe 216 strawberry fruit from the replicate plants at each aqueous applied concentration were composited to achieve adequate 217 biomass for extraction, resulting in composited averages of 218 219 analytical triplicate measurements. Strawberry shoot and root 220 experimental replicates for three aqueous concentrations (0.4, 4, 40  $\mu$ g/L) were extracted separately to enable an estimation 221 222 of interpot variability (since field replicates of fruit were 223 composited); concentrations in replicate plants (n = 3-5) were 224 averaged to obtain sample values. All results for plants are 225 presented in terms of a dry weight basis. Aqueous sample 226 analyses were completed per established methods.<sup>28</sup>

227 Samples were analyzed with isotope dilution using LC-MS/ 228 MS under conditions outlined in previous work.<sup>23</sup> Briefly, chromatography was performed using a Shimadzu LC-20AD 229 unit (Kyoto, Japan) by injecting samples onto a Gemini C18 230 column with a  $3-\mu$ m particle size (Phenomenex, Torrance, CA). 231 In addition, two transitions for each PFAA were observed using 232 an ABSCIEX 3200 (ABSCIEX, Ontario) with negative 233 electrospray ionization operating in scheduled multiple reaction 234 mode. Quantitation of LC–MS/MS data was accomplished 235 using Analyst software. 236

Quality Assurance and Control. All of the strawberry 237 fruit, as well as approximately 20% of all other samples, were 238 extracted and analyzed in triplicate. The relative standard 239 deviation for all analytical replicates averaged <25%. One 240 laboratory blank with surrogate standard and one double blank 241 without surrogate standard were prepared for each batch of 242 samples. Limits of quantitation (LOQ) for plant material 243 ranged from 0.07 to 29 ng/g; LOQs were determined by the 244 lowest calibration standard calculated to be within 30% of its 245 actual value and were analyte-, matrix-, and run-dependent. 246 LOQs were also required to be at least twice as high as the 247 highest concentration in the corresponding blanks and have 248 signal-to-noise ratios >30. If a minimum of three pot replicates 249 were above the LOQ, an average value was calculated for that 250 treatment; otherwise, the value was reported as <LOQ. To 251 account for any losses during the extraction process, each 252 analyte contained an internal surrogate. In line with previous 253 work analyzing PFAAs in plant tissues,<sup>29</sup> surrogate recovery for 254 the samples averaged 43% for root tissues, 33% for shoot 255 tissues, and 45% for fruit tissues across all analytes. Statistical 256 analysis including all calculations of regression equations was 257 completed using OriginPro 9.0. 2.58

**Bioaccumulation Metrics.** Bioaccumulation factors 259 (BAFs) for lettuce leaves (at the 10  $\mu$ g/L applied 260 concentration) and, more specifically, fruit-to-soil concentra- 261 tion factors<sup>29</sup> (FCFs) for strawberry fruit (at the 0.4, 10, and 40 262  $\mu$ g/L applied concentrations) were calculated for each PFAA 263 that had concentrations in the plant tissues above the LOQ. To 264 enable comparisons to previous studies examining PFAA 265 bioaccumulation from soils, the aqueous concentration ( $C_w$ ) 266 was first converted to an estimated soil concentration ( $C_s$ ) 267 using the respective solid–water partitioning coefficient ( $K_d$ ) 268 for each soil and analyte (eq 1).

$$C_{\rm s}\left(\frac{\rm ng}{\rm kg}\right) = C_{\rm w}\left(\frac{\rm ng}{\rm L}\right) \times K_{\rm d}\left(\frac{\rm L}{\rm kg}\right) \tag{1}_{270}$$

271 Although it remains unclear as to whether equilibrium 272 conditions were present, single point  $K_d$  values were used in the 273 absence of soil pore water concentrations; moreover, previously 274 measured isotherms for PFAAs were fairly linear (Freundlich *n* 275 values ~ 0.9–1) providing validity for this estimation 276 method.<sup>17</sup> More information concerning the determination of 277  $K_d$  values can be found in the SI. Concentration factors were 278 then calculated as in previous work<sup>23,29</sup> by dividing the 279 concentration of chemical in the respective plant tissue on a 280 dry weight basis by the concentration of chemical in the soil. In 281 addition, intercompartmental concentration factors (fruit to 282 shoot and shoot to root) were calculated as in previous work<sup>29</sup> 283 for strawberry plants grown at the 0.4, 4, and 40  $\mu$ g/L applied 284 concentrations.

#### 285 **RESULTS AND DISCUSSION**

Concentration-Dependency Trends. All PFAAs meas-286 287 ured in lettuce leaves showed predominately linear concen-288 tration-response relationships (Figure 1), suggesting passive 289 transport through the plant.<sup>30</sup> The slopes of the nonlog linear 290 regressions for each analyte (SI Figure S1) imply preferential uptake by the short-chain PFAAs. In general, PFCAs 291 292 accumulated in much greater quantities than the perfluorosul-293 fonates (PFSAs) with concentrations in lettuce leaves receiving 294 the highest application of PFAAs reaching 25  $\mu$ g/g for PFBA. Conversely, PFCA accumulation was <LOQ in the lettuce 2.95 treated with control tap water and ambient reclaimed water (SI 296 Table S6). 297

Short-chain PFCA accumulation in strawberry fruit was also in fruit receiving the highest application of PFAAs were >10  $\mu g/g$  for PFBA and PFPeA (SI Table S7). PFHxA accumulation in strawberry fruit was not measured above the the S03 LOQ except for the four highest aqueous doses. PFHpA, except for the four highest aqueous doses. PFHpA, out PFOA, and PFNA concentrations were all <LOQ. Linear soc concentration-response relationships, similar to those obsof served in lettuce, are shown in Figure 2, and the nonlog linear regressions are displayed in SI Figure S2. Of the PFSAs, only pFBS concentrations were above LOQ, and accumulation in som the strawberry fruit was minimal (<56 ng/g) compared with the som PFCAs (SI Table S7). The lack of PFSA accumulation in the



**Figure 2.** Concentrations of PFCAs in strawberry fruit versus measured aqueous concentration of PFAAs. Means of composited berries are shown with analytical standard deviation (n = 3).

fruit compartment is consistent with previous findings in 311 tomato and pea fruit.<sup>29</sup> The observed bias against accumulation 312 of long-chain PFAAs in the fruit compartment may suggest that 313 other specific transport mechanisms exist for long-chain PFAAs. 314 Wen et al.<sup>31</sup> studied uptake of PFOA and PFOS by maize and 315 found indications of potential active transport for PFOA, partial 316 aquaporin (water channel) transport for PFOS, and limited 317 carrier-mediated transport through different anion channels for 318 both analytes. Although strawberry plants are very different 319 from maize, similar mechanisms may contribute to the lack of 320 accumulation of long-chain PFAAs in strawberry fruit. 321

Chain Length Trends. As evidenced in Figures 1 and 2, 322 concentrations in both lettuce leaves and strawberry fruit 323 decreased as PFAA chain length increased. To further illustrate 324 this trend, PFAA concentrations for a single aqueous applied 325 concentration (10  $\mu$ g/L) in both lettuce and strawberry are 326 provided in Figure 3. In lettuce leaves, PFAA concentrations 327 f3 spanned more than an order of magnitude from PFBA to 328 PFNA, a gain of 5 carbons, and also more than an order of 329 magnitude from PFBS to PFOS, a gain of 4 carbons. In 330 strawberry fruit, PFAA concentrations spanned more than an 331 order of magnitude from PFBA to PFHxA, a gain of only 2 332 carbons, further evidencing the disparity of accumulation 333 potential between short- and long-chain PFCAs. This 334 preferential accumulation of short chain carboxylates in plants 335 is consistent with previous findings.<sup>29</sup> 336

Strawberry Plant Compartments. Nonedible portions of 337 strawberry plants were analyzed to assess interpot variability 338 (22%) and help elucidate bioaccumulation trends within the 339 plant. At the highest aqueous concentration applied (40  $\mu$ g/L), 340 strawberry root concentrations were greatest for PFHxA (5450 341 ng/g; SI Table S8), strawberry shoot concentrations were 342 greatest for PFBA (3900 ng/g; SI Table S8), and strawberry 343 fruit concentrations were greatest for PFPeA (11 500 ng/g; SI 344 Table S7). Moreover, the concentrations of both PFBA and 345 PFPeA in the fruit were more than twice that of any analyte 346 that accumulated in the root or shoot compartments. The 347 distribution of PFAAs in each plant compartment (root, shoot, 348 and fruit) for a representative aqueous concentration (4  $\mu$ g/L) <sup>349</sup> is shown in Figure 4. Of the three compartments in the 350 f4 strawberry plant, the root compartment had the greatest 351 accumulation of PFAAs (2840 ng/g), and the distribution of 352 PFAAs in the root compartment was fairly evenly spread, 353 confirming the lack of selectivity of analytes in the root 354 compartment described by Blaine et al.<sup>29</sup> The shoot compart- 355 ment had the lowest total accumulation of PFAAs (705 ng/g) 356 of the three compartments, and the accumulation was 357 dominated by the short-chain analytes, PFBA, PFBS, and 358 PFPeA. The fruit compartment had almost as much total 359 accumulation (2520 ng/g) as the root compartment; however, 360 the distribution of PFAAs was highly skewed toward the short- 361 chain PFCAs, and no PFSAs were present in the fruit. Because 362 plants were grown with limited irrigation (i.e., not hydroponi- 363 cally) to represent field conditions, a high percentage of the 364 water taken up was most likely used for fruit development;<sup>32,33</sup> 365 the relatively low amount of water transpired versus the water 366 used for fruit development could explain the higher 367 accumulation of PFAAs in the fruit versus the shoot. 368

The mass distribution between plant compartments for each 369 analyte can be estimated by multiplying typical dry weights for 370 each compartment (1.3 g for root, 4 g for shoot, 3 g for fruit) 371 by the concentration of each analyte in the respective 372 compartment. The dominant fractions (>65%) of PFBA and 373



**Figure 3.** Concentrations of PFAAs in lettuce leaves (a) and strawberry fruit (b) for the aqueous applied PFAA concentration of 10  $\mu$ g/L. Mean and standard error for lettuce (n = 5) are shown. Means of composited berries are shown with analytical standard deviation (n = 3).



Figure 4. Distribution of PFAAs in strawberry root, shoot, and fruit compartments for the applied aqueous PFAA concentration of 4  $\mu$ g/L.

<sup>374</sup> PFPeA resided in the fruit compartment, whereas the dominant <sup>375</sup> fractions (>70%) of the long-chain PFAAs accumulated in the <sup>376</sup> root compartment (SI Figure S3). Similar trends concerning <sup>377</sup> mass distribution within plant compartments were observed for <sup>378</sup> both lower (0.4  $\mu$ g/L) and higher (40  $\mu$ g/L) aqueous <sup>379</sup> concentrations.

FCF values (converted to a soil basis for the 10  $\mu$ g/L applied 380 381 aqueous concentration) for strawberry ranged from the 200s for PFBA and PFPeA to 35 for PFHxA (SI Table S9). Although 382 the data are limited, when plotted versus carbon chain length, 383 the overall average decrease of FCF per CF<sub>2</sub> group was ~0.3 384 log units (Figure 5). These results are similar to the findings of 385 Blaine et al.<sup>29</sup> for tomato and pea fruit, as shown alongside the 386 strawberry fruit data in Figure 5. The differences in FCF values 387 for strawberry as compared to tomato and pea could be due to 388 differences in plant morphology or to the delivery of PFAAs via 389 390 irrigation water versus biosolids. At the highest aqueous <sub>391</sub> concentration (40  $\mu$ g/L), the FCF versus chain length trend 392 is also ~0.3 log units, indicating consistency in accumulation

f5

trends at varying concentrations (SI Figure S4). Intercompart- <sup>393</sup> mental factors for strawberry plotted versus PFAA chain length <sup>394</sup> (SI Figure S5) showed a decrease of 0.2 log units from fruit to <sup>395</sup> shoot per CF<sub>2</sub> group and 0.3 log units from shoot to root per <sup>396</sup> CF<sub>2</sub> group. These factors also correspond well to the <sup>397</sup> intercompartmental factors calculated for tomato and pea fruits <sup>398</sup> from previous work<sup>29</sup> (SI Figure S5). <sup>399</sup>

**Lettuce–Soil Organic Carbon Study.** To assess the  $_{400}$  impact of soil sorption on plant uptake of PFAAs, lettuce was  $_{401}$  grown in soils with differing OC contents and compared with  $_{402}$  the lettuce grown in the sand–soil mix. Lettuce grown in the  $_{403}$  two additional soil treatments (2% and 6% OC content) at the  $_{404}$  10  $\mu$ g/L applied concentration accumulated similar (in general  $_{405}$  within a factor of 3) PFAA concentrations to that grown in the  $_{403}$  soil mix (SI Table S10). Lettuce in all three soils had the  $_{407}$  highest concentrations of PFBA and PFPeA. Concentrations  $_{408}$  ranged from approximately 15  $\mu$ g/g of PFBA to 47 ng/g of  $_{409}$  PFNA in the 2% OC soil and from almost 5  $\mu$ g/g of PFBA to  $_{410}$  21 ng/g of PFNA in the 6% OC soil (SI Table S10).



**Figure 5.** Correlations for PFCAs between log fruit—soil concentration factors and carbon chain length in strawberry, tomato, and pea. Strawberry values from this study were from the 10  $\mu$ g/L applied aqueous concentration; tomato and pea values were from a previous study.<sup>29</sup> Means and standard errors are shown (n = 3 to 5). Linear regressions with slopes, intercepts, and associated error values are shown.

412 Lettuce BAFs for all three soil treatments at the 10  $\mu$ g/L 413 applied concentration varied widely, spanning more than 2 414 orders of magnitude within each treatment. The lettuce grown 415 in the 6% OC soil had the smallest BAF values for all PFAAs, 416 presumably because of the sorption in the media (SI Table 417 S11). All BAFs, with the exception of PFNA and PFOS in the 418 6% OC soil, were ~1, indicating the accumulation of PFAAs in 419 the lettuce. PFNA and PFOS have the highest  $K_d$  values of the 420 PFAAs in this study (SI Table S4), so it follows well that they 421 would exhibit minimal bioavailability in the highest OC content 422 soil.

f6

A linear relationship between log BAF values and carbon 423 chain length is shown in Figure 6 for lettuce grown in each soil 424 treatment. For each increase in carbon chain length, the BAF 425 426 decreased ~0.4–0.6 log units. In addition, lettuce grown in two 427 different PFAA contaminated biosolids-amended soils (2.2% 428 and 6.3% OC) from a previous study<sup>23</sup> are plotted alongside 429 the values from the present study for comparison. The slopes of 430 all lines are somewhat similar (-0.31 to -0.70), with the slopes 431 of biosolids-grown lettuce being slightly flatter than the lettuce 432 grown with aqueous-applied PFAAs. This difference could 433 indicate that the mobility of the PFAAs supplied by aqueous 434 application allows immediate plant uptake prior to significant 435 sorption in the soil. Figure 6 assumes that the irrigation 436 solution is representative of pore water and that the water-soil 437 system had reached equilibrium prior to plant uptake. In reality, 438 however, equilibrium may not have been reached, thus 439 increasing the bioavailability of the PFAAs applied in the 440 reclaimed water. The lettuce slopes from the present study 441 represented in Figure 6 may therefore be artificially steep 442 compared with equilibrium conditions; however, they may be 443 more representative of actual field conditions. Regardless, 444 greater bioaccumulation overall is seen in the lettuce plants 445 from the present study, suggesting that the mobility and



**Figure 6.** Correlations for PFCAs between log BAFs and carbon chain length in lettuce. Log BAFs from lettuce grown in soils with varying OC content (0.4%, 2%, 6%) at the 10  $\mu$ g/L applied concentration are shown alongside values from lettuce grown in biosolids-amended soils in a previous study.<sup>23</sup> Means and standard errors are shown (n = 3-5). Linear regressions with slopes, intercepts, and associated error values are shown.

bioavailability of PFAAs is greater when delivered via irrigation 446 water as compared with biosolids-amended soil. 447

Implications. The results of this study are novel and 448 important because it is the first of its kind to examine PFAA 449 accumulation in food crops simulating, at least in part, a real- 450 world field scenario by using authentic reclaimed water as the 451 delivery medium. Certainly, real-world conditions such as the 452 dilution of the contaminant concentrations crops are exposed 453 to as a result of precipitation, perennial application of reclaimed 454 water (potentially leading to a higher exposure concentration 455 due to contaminant accumulation in the soil), and other 456 numerous water management factors further complicate the 457 picture for real-world extrapolation. Regardless, the data 458 presented herein show clearly that PFAAs can be taken up 459 and accumulated into food crops grown in soil and irrigated 460 with reclaimed water, suggesting the potential for human 461 exposure if irrigation water contains PFAAs. At typical WWTP 462 effluent concentrations of PFAAs  $(0.02-4 \ \mu g/L)$ ,<sup>5</sup> values 463 reached up to 0.2  $\mu$ g/g for PFOA and 3  $\mu$ g/g for PFBA in 464 lettuce and up to 2  $\mu$ g/g for PFBA in strawberry fruit. In 465 addition, at higher aqueous concentrations, more representative 466 of contaminated surface or ground waters  $(10-40 \ \mu g/L)$ ,<sup>25</sup> 467 concentrations of PFAAs accumulated up to 1  $\mu$ g/g for PFOA 468 and 25  $\mu$ g/g for PFBA in lettuce and up to and 11  $\mu$ g/g for 469 PFBA in strawberry fruit. The proposed subchronic reference 470 dose for PFOA according to the U.S. EPA is 0.2  $\mu$ g/kg-day;<sup>34</sup> 471 for an average 70 kg adult, the maximum daily allowance of 472 PFOA would then be 14  $\mu$ g/day. If a person were to consume 473 lettuce irrigated with contaminated water (40  $\mu$ g/L of PFOA, 474 well above the provisional health advisory level for drinking 475 water of 0.4  $\mu g/L^{35}$ ), then presumably, less than half of a small 476 head of lettuce (126 g on a wet weight basis) would be enough 477 to reach the daily maximum for PFOA. Concentrations of 478 short-chain PFAAs in the lettuce would be even higher; 479 however, substantial toxicological data are lacking for short- 480 chain PFAAs. Although the levels of PFAAs in lettuce and 481

482 strawberries irrigated with typical WWTP effluent would be 483 expected to be considerably lower, concerns may arise if the 484 WWTP received significant inputs from fluorochemical-using 485 industries.

The concentration-dependent response for all PFAAs in 486 487 lettuce and for short-chain PFAAs in strawberry fruit implies 488 that PFAA accumulation in this range of aqueous concen-489 trations does not plateau, and thus, the uptake potential for 490 crops grown with highly contaminated water (e.g., surface water 491 or groundwater near industry) is great. Long-chain PFCAs and 492 PFSAs, however, do not readily accumulate in high quantities in 493 strawberry fruit, regardless of increased aqueous concentration, 494 and therefore, indications are that fruit may not be a major 495 route of exposure for long-chain PFAAs. In general, 496 bioaccumulation patterns observed in this study are consistent <sup>497</sup> with literature<sup>22,23</sup> showing greater uptake and accumulation for 498 PFCAs over PFSAs and for short-chain PFAAs over long-chain 499 PFAAs. These plant compartment accumulation trends are 500 important with respect to assessing potential human exposure 501 through consumption. As industry trends shift toward the 502 manufacture of short-chain PFAAs, increased concentrations of 503 shorter PFAAs in WWTP effluents can be expected.

If the current use of reclaimed water for food crops is to be sustained or increased, concerns about the potential contamsof ination of food products must be fully addressed through careful scientific study, evaluation, and communication with the som public. Future research is warranted by this potential exposure som route to humans. More work is needed to understand PFAA sit transport mechanisms in additional crops. Investigations of sit crop uptake using a broader suite of PFAAs present in sit reclaimed water, including potential precursors of PFAAs, are sit also needed to expand the body of knowledge on this emerging sit topic of concern.

# 515 ASSOCIATED CONTENT

#### 516 **Supporting Information**

517 Additional details are available regarding PFAA standards, 518 reclaimed water quality, soil characteristics, greenhouse experi-519 ment details, PFAA concentrations in lettuce (shoots) and 520 strawberry (fruits, shoots, and roots), linear regressions for 521 concentration-dependency study, mass distribution of PFAAs in 522 plant compartments, plots of strawberry intercompartmental 523 factors versus carbon chain length, and lettuce concentration 524 and BAF values for organic carbon study. This material is 525 available free of charge via the Internet at http://pubs.acs.org.

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529 Notes

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