

This is a repository copy of *Insight into the distribution of pharmaceuticals in soil-water-plant systems*.

White Rose Research Online URL for this paper:

<https://eprints.whiterose.ac.uk/151016/>

Version: Accepted Version

Article:

Li, Yuanbo, Sallach, J. Brett orcid.org/0000-0003-4588-3364, Zhang, Wei et al. (2 more authors) (2019) *Insight into the distribution of pharmaceuticals in soil-water-plant systems*. *Water research*. pp. 38-46. ISSN 0043-1354

<https://doi.org/10.1016/j.watres.2018.12.039>

Reuse

This article is distributed under the terms of the Creative Commons Attribution-NonCommercial-NoDerivs (CC BY-NC-ND) licence. This licence only allows you to download this work and share it with others as long as you credit the authors, but you can't change the article in any way or use it commercially. More information and the full terms of the licence here: <https://creativecommons.org/licenses/>

Takedown

If you consider content in White Rose Research Online to be in breach of UK law, please notify us by emailing eprints@whiterose.ac.uk including the URL of the record and the reason for the withdrawal request.

1 **Insight into the Distribution of Pharmaceuticals in Soil-Water-Plant Systems**

2

3 Yuanbo Li, J. Brett Sallach, Wei Zhang, Stephen A. Boyd, and Hui Li*

4 Department of Plant, Soil and Microbial Sciences, Michigan State University, East Lansing, MI

5 48824, USA

6

7

8

9

10

11

12

13

14

15

16

17

18 * Corresponding Author:

19 Phone: 517-355-0151

20 E-mail: lihui@msu.edu

21

22 **ABSTRACT**

23 Pharmaceuticals in agricultural soils originating from irrigation with treated wastewater and land-
24 applied biosolids can enter field crops. However, little is known about the role of pore water in
25 plant uptake of pharmaceuticals from soil. In this study, the fate, uptake and distribution of fifteen
26 commonly used pharmaceuticals in soil-water-radish systems were investigated to examine the
27 relationship between the accumulation and their physicochemical processes in soils. The results
28 indicate that the distribution of pharmaceuticals between soil and pore water, as well as their
29 biodegradation, combined to govern the bioavailability of pharmaceuticals to plant uptake.
30 Fourteen out of 15 pharmaceuticals could enter radish tissues in which the accumulation ranged
31 from 2.1 to 14080 ng/g. Comparison of bioconcentration factors (BCFs) on the basis of
32 pharmaceutical concentration in bulk soil vs. in pore water implies that pharmaceuticals present in
33 soil pore water are the major bioavailable fractions to plant uptake. The pore water-based BCFs
34 exhibited a positive linear relationship with $\log D_{ow}$ for the pharmaceuticals with > 90% as neutral
35 species in soil pore water, while such relationship was not observed between bulk soil-based BCFs
36 and $\log D_{ow}$ mainly due to sorption by soil. Other than hydrophobicity, the dissociation of ionizable
37 pharmaceuticals in the soil pore water and (or) root cells may lead to the “ion-trap” effects and
38 thus influence the uptake and translocation process. The large molecular size pharmaceuticals (e.g.,
39 tylosin) manifested a minimum uptake due plausibly to the limited permeability of cell membranes.

40

41 *Keywords:* Plant uptake; Soil pore water; Bioavailability; Bioaccumulation; Translocation.

42

43 **1. Introduction**

44 Some pharmaceuticals have been recognized as chemicals of emerging concern because
45 they are widespread in the environment, and have the potential adverse effects on non-target
46 organisms and humans (e.g., endocrine disruption and preservation of antibiotic resistance)
47 (Carvalho et al., 2014; Daughton and Ternes, 1999; Kolpin et al., 2002; Madikizela et al., 2018).
48 Conventional wastewater treatment processes cannot effectively remove all pharmaceuticals from
49 the influents, leaving the pharmaceuticals in the effluents at the levels of ng/L to low µg/L (Gros
50 et al., 2010; Sui et al., 2011; Vanderford and Snyder, 2006) and in biosolids at µg/kg to low mg/kg
51 (dry weight) (Clarke and Smith, 2011; McClellan and Halden, 2010). Irrigation with the treated
52 wastewater and land application of biosolids are common agricultural practices, which lead to the
53 dissemination of a wide range of pharmaceuticals in agricultural soils with concentration up to
54 mg/kg levels (Carter et al., 2014; Durán-Alvarez et al., 2009; Kinney et al., 2006; Vazquez-Roig
55 et al., 2010). For example, land application of biosolids at a rate of 5 kg/m² caused triclosan
56 concentration of 0.77-0.95 mg/kg in the farm soils located in Bedfordshire, UK (Butler et al., 2012).
57 One major consequence of the soil contamination is that these pharmaceuticals could enter food
58 chain after plant uptake, and pose potential risks to human and animal health via dietary
59 consumption (Christou et al., 2017; de Boer et al., 2018; Sallach et al., 2015; Wu et al., 2014).
60 Paltiel et al. (2016) recently found that carbamazepine and its metabolites were detected in human
61 urine after consuming the fresh produce irrigated with treated wastewater.

62 In soil-plant systems, the amount of organic chemical uptake by plant roots depends largely
63 on sorption/desorption of contaminants in soils and their physicochemical properties. It is assumed
64 that organic chemicals present in soil pore water are readily bioavailable to plant uptake (Miller et
65 al., 2016), and soil water serves as the carrier to move the chemicals into plants. The distribution
66 of pharmaceuticals in soil-water-plant systems is commonly characterized by bioconcentration

67 factor (BCF) which is typically calculated as the ratio of pharmaceutical concentration in plant to
68 that in bulk soil (Hurtado et al., 2016; Karnjanapiboonwong et al., 2011; Pan et al., 2014; Shenker
69 et al., 2011; Wu et al., 2010). However, pharmaceuticals present in soil pore water vs sorb by soil
70 could manifest different bioavailability to plant uptake. Therefore, BCFs calculated on the basis
71 of pharmaceutical concentration in bulk soil are not comparable among the studies using different
72 soils because of the varying affinities of pharmaceuticals to soils. For instance, soil-based BCFs
73 of seven benzodiazepines in radish could vary by up to 86.0 times between two soils (Carter et al.,
74 2018). The difference of soil-based BCFs between three soils for caffeine, carbamazepine, and
75 lamotrigine in tomato or cucumber can up to 20.0, 7.8, and 245 times, respectively (Goldstein et
76 al., 2014). In fact, BCFs calculated by pharmaceutical concentration in soil pore water are believed
77 to provide more accurate information to describe the uptake process since pharmaceuticals in soil
78 pore water are directly available to plant root uptake. For example, Blaine et al. (2014) used pore
79 water-based BCFs of perfluoroalkyl acids in four vegetables grown in soils to explore the
80 relationship between bioaccumulation and chemical properties, and found the consistent results
81 with the vegetables grown in hydroponic solution but not for soil-based BCFs. Despite numerous
82 greenhouse and field studies that have been conducted, very limited studies were conducted to
83 systematically evaluate the contribution of pharmaceuticals in soil pore water to the
84 bioaccumulation in plants (Boxall et al., 2006; Carter et al., 2014).

85 Water is believed to carry xenobiotic chemicals (e.g. pharmaceuticals) to cross plant root
86 cortex by symplastic (intracellular space) and apoplastic (extracellular space) pathways
87 (McFarlane and Trapp, 1994; Trapp and Pussemier, 1991). In plant roots, Casparian strip
88 composing primarily of hydrophobic suberin and lignin functions as a water-impermeable barrier
89 to prevent water and pharmaceuticals from passing through the endodermis via the apoplastic route

90 (Naseer et al., 2012; Schreiber, 2010). Therefore, pharmaceuticals have to reenter the symplastic
91 pathway to cross cell membranes and enter the xylem. For many nonionic organic compounds, the
92 accumulation in plants is positively related to their lipophilicity as indicated by the linear
93 relationship between BCFs and octanol–water partition coefficient (K_{ow}) (Briggs et al., 1982;
94 Briggs et al., 1983). However, plant uptake of ionic compounds is determined by the combination
95 of hydrophobicity, chemical speciation and surrounding solution pH (Briggs et al., 1987; Trapp,
96 2000). Most pharmaceuticals are ionizable compounds and have low hydrophobicity (e.g., $\log D_{ow}$
97 < 2). Therefore, the relations developed for nonionic organic contaminants may not be applied to
98 the uptake of pharmaceuticals. For example, no apparent relationship was observed between \log
99 BCF and $\log D_{ow}$ (pH-adjusted K_{ow} to neutral species) for 20 pharmaceuticals (including acids,
100 bases and neutral compounds) in hydroponically grown lettuce, spinach, cucumber and pepper
101 (Wu et al., 2013). However, strong correlations were observed when the data were limited to
102 neutral pharmaceuticals. The relationship between plant uptake of ionic pharmaceuticals and their
103 physicochemical properties still remain largely unknown in the complex soil-water-plant systems,
104 where the dissociation in soil and plant at relevant pHs has been considered as an important factor
105 (Carter et al., 2014; Goldstein et al., 2014; Hyland et al., 2015; Malchi et al., 2014).

106 This study aims to investigate the fate, uptake and distribution of fifteen diverse
107 pharmaceuticals in soil-radish systems. Radish plants grew in soil with the soil moisture levels at
108 75%, 60% and 45% of maximum water holding capacity (MWHC), which represent the common
109 range of soil water contents in agricultural field. BCF values, calculated on the basis of
110 pharmaceuticals concentration in bulk soil vs. in pore water, were compared to gain more insight
111 into the bioavailable fractions of pharmaceuticals in soil to plant root uptake. The selected
112 physicochemical properties of pharmaceuticals were evaluated to examine the relation to their

113 uptake and translocation in plants. This study provided experimental results to verify and improve
114 the understanding of plant uptake of pharmaceuticals from soils in terms of pore water, which
115 delivers the useful information on the risk assessment of human exposure to pharmaceutical-
116 contaminated vegetables from the reuse of treated wastewater and biosolids in agriculture.

117

118 **2. Experimental Section**

119

120 *2.1. Chemicals and Materials*

121

122 Acetaminophen, caffeine, carbamazepine, sulfadiazine, sulfamethoxazole, lamotrigine,
123 carbadox, estrone, 17 β -estradiol, triclosan, trimethoprim, lincomycin, oxytetracycline, monensin
124 and tylosin were purchased from Sigma-Aldrich (St. Louis, MO, USA). These fifteen
125 pharmaceuticals (Table 1) were selected because they are commonly present in treated wastewater
126 and biosolids, as well as in agricultural lands. Ceramic homogenizers, octadecylsilane (C18), and
127 primary secondary amine (PSA) were purchased from Agilent Technologies (Santa Clara, CA,
128 USA). Disodium ethylenediaminetetraacetate (Na₂EDTA), formic acid, and sodium chloride
129 (NaCl) were purchased from J.T. Baker (Phillipsburg, NJ, USA). Organic solvents (HPLC grade)
130 were purchased from Fisher Scientific (Fair Lawn, NJ). Ultrapure water was produced from a
131 Milli-Q water purification system (Millipore, Billerica, MA, USA).

132 A sandy loam soil was collected from Michigan State University Research and Teaching
133 Farm located in Lansing, Michigan. The sampling site had not previously irrigated with treated
134 wastewater or amended with biosolids and manure. The soil contained 79.0 % sand, 12.6 % silt,
135 and 8.4 % clay. The soil was mixed with a portion of peat to achieve soil organic matter content

136 of 2.8 %. The soil had a cation exchange capacity of 9.2 cmol/kg. The soil was air-dried and passed
137 through 2-mm sieve before use.

138

139 2.2. Experimental Setup

140

141 A mixture of the fifteen pharmaceuticals (Table 1) in acetone was spiked to soil (1.0 kg)
142 and thoroughly mixed in a fume hood. After acetone was evaporated, the soil with spiked
143 pharmaceuticals was mixed thoroughly with 32.0 kg of clean soil using a motorized concrete mixer
144 to achieve the final concentration of 1 µg/g for each pharmaceutical. Five radish (*Raphanus sativus*)
145 seeds (Burpee & Co., Warminster, PA) were planted in each plastic pot containing 1.2 kg of the
146 soil. The pots were irrigated daily with deionized water to maintain water content at 75 %, 60 %
147 and 45 % of MWHC by monitoring the pot weights. No additional fertilizer was added. All
148 experimental pots were prepared in triplicate, including the pharmaceutical-free and radish-free
149 controls. The pots were randomly placed in a climate-controlled greenhouse under sunlight at
150 25 °C during daytime and 21 °C during night. After seed germination (at day 8), the plants were
151 thinned to two radish seedlings per pot.

152 For the radish-free controls, soil and pore water samples were collected at 4 hours after
153 irrigation at day 0, 1, 3, 7, 14, 21, 28 and 35. For the uptake experiments, radish were sampled at
154 day 28 (premature stage) and day 35 (mature stage). The radish plants were thoroughly rinsed with
155 deionized water to remove the attached soil particles, wiped with tissue paper, and separated into
156 roots (including bulb) and leaves. Plant samples were weighed, cut into small pieces, freeze-dried
157 and ground to fine powders. All samples were stored at -20 °C prior to extraction. At the same
158 time, soil samples were also collected from the plant pots at day 28 and 35 for the analysis of

159 pharmaceuticals in soil and pore water. The pH of collected pore water was measured using a
160 Fisher Scientific Accumet AB15 pH meter (Pittsburgh, PA, USA).

161

162 *2.3. Sample Extraction and Analysis*

163

164 Pharmaceuticals in radish and soil samples were extracted using a modified QuChERS
165 method (Chuang et al., 2015). In brief, dry radish (0.5 g) or soil sample (2.5 g) was placed in
166 polypropylene centrifuge tube to which 2.0 mL of 150 mg L⁻¹ of Na₂EDTA was added with two
167 pieces of ceramic homogenizers and vortexed for 1 min. The sample was then extracted with 5.0
168 mL of acetonitrile and methanol mixture (v/v = 65/35) by vigorously shaking for 3 min. Then,
169 Na₂SO₄ (2.0 g) and NaCl (1.0 g) were added and vortexed for another 2 min. The tubes were
170 centrifuged at 2990 g for 10 min, and 1.3 mL of supernatant was transferred into clean 1.5-mL
171 centrifuge tubes containing 250 mg of Na₂SO₄ and d-SPE sorbents (25 mg of C18, and 25 mg of
172 PSA). The samples were vortexed for 1 min, and centrifuged at 9240 g for 10 min. The supernatant
173 was filtered through a 0.22 µm polytetrafluoroethylene (PTFE) filter and stored at -20 °C prior to
174 analysis.

175 Soil pore water was collected immediately after soil sampling using the protocol described
176 by Carter et al. (2014). Briefly, 25 g of soil was placed in a 20-mL disposable plastic syringe with
177 a glass wool insert at the bottom. The syringes were placed in 50-mL centrifuge tubes and
178 centrifuged at 4300 g for 40 min. The collected pore water was passed through a 0.22-µm PTFE
179 membrane, and an aliquot of the pore water (0.5 mL) was diluted to 1.0 mL with methanol in clean
180 HPLC glass vials for LC-MS/MS analysis.

181 The LC-MS/MS system consisted of a Shimadzu prominence high-performance liquid
182 chromatography (Columbia, MD, USA) coupled to a Sciex 4500 triple quadrupole mass
183 spectrometer (Foster City, CA, USA), and a 50 mm × 2.1 mm Agilent C18 column (Torrance, CA,
184 USA). Detailed information about the LC-MS/MS optimized condition is provided in the
185 supporting information (SI). Multiple reaction monitoring (MRM) parameters used for the analysis
186 are listed in Table S1. Extraction efficiency and method detection limits (MDLs) of
187 pharmaceuticals are summarized in Tables S2 and S3.

188

189 2.4. Data Analysis

190

191 Soil-pore water distribution coefficient (K_d) was calculated using equation 1. BCF was
192 calculated by measured pharmaceutical concentration in soil and in pore water using equation 2
193 and 3. Translocation factor (TF) was calculated using equation 4. The first-order dissipation rate
194 constant (k) of pharmaceuticals in soil was estimated by fitting the time-dependent concentration
195 to the first-order decay model using equation 5. The dissipation half-life ($T_{1/2}$) was calculated using
196 equation 6.

$$197 \quad K_d = \frac{C_s}{C_{pw}} \quad (1)$$

$$198 \quad \text{BCF}_{\text{soil}} = \frac{C_{\text{root}}}{C_{\text{soil}}} \quad (2)$$

$$199 \quad \text{BCF}_{\text{pore water}} = \frac{C_{\text{root}}}{C_{pw}} \quad (3)$$

$$200 \quad \text{TF} = \frac{C_{\text{leaf}}}{C_{\text{root}}} \quad (4)$$

$$201 \quad C_t = C_0 e^{-kt} \quad (5)$$

$$202 \quad T_{1/2} = \frac{\ln(2)}{k} \quad (6)$$

203 where C_s is pharmaceutical concentration in soil solid phase (total amount of pharmaceuticals in
204 soil subtracted by that present in pore water) (ng/g, dry weight), C_{pw} is pharmaceutical
205 concentration in pore water (ng/mL), C_{soil} is pharmaceutical concentration in bulk soil (ng/g, dry
206 weight). C_{root} and C_{leaf} are pharmaceutical concentration in radish roots and leaves (ng/g, dry
207 weight), respectively. BCF_{soil} and $BCF_{pore\ water}$ are the BCFs calculated on the basis of
208 pharmaceutical concentration in bulk soil and in pore water, respectively. C_0 and C_t are
209 pharmaceutical concentration at the beginning of the incubation and time t (d) in soil, respectively.
210 Statistical analysis (All Pairs, Tukey HSD, $p = 0.05$) of experimental results for significant analysis
211 was carried out using SPSS 22.0 software for Windows (IBM Corp., Armonk, NY).

212

213 **3. Results and discussion**

214

215 *3.1. Plant Uptake*

216

217 Radish grew well in the soil with the three water content at 75%, 60%, and 45% of MWHC,
218 and appeared in good health. The associated radish biomass showed no significant difference at
219 the three water contents in the presence and absence of pharmaceuticals ($p > 0.05$) (Figure S1).
220 The radish plants were separated into leaves and roots, and analyzed individually for the
221 accumulated pharmaceuticals (Table S4). All the pharmaceuticals except 17 β -estradiol were
222 detected in the radishes sampled at both day 28 and 35. Pharmaceutical concentration in roots
223 ranged from 2.4 to 1774 ng/g. Carbamazepine demonstrated the highest accumulation, which was
224 up to 738 times greater than estrone (the least accumulated compound in roots) at day 35. In the
225 leaves, thirteen out of the fifteen pharmaceuticals (except 17 β -estradiol and monensin) were

226 detected with the concentration range of 2.1 to 14080 ng/g. Carbamazepine, caffeine, lamotrigine
227 and trimethoprim were measured at relatively high concentrations in leaves (462-14080 ng/g),
228 while tylosin and estrone were detected at low concentrations (2.1-12.3 ng/g). Monensin was found
229 to accumulate in the roots only, which was not detected in the leaves.

230 Pharmaceuticals accumulated in plants may be metabolized, thus further influence their
231 accumulation and distribution in plant organs. In our previous study, we found that 17 β -estradiol,
232 sulfamethoxazole, sulfadiazine, estrone, triclosan, acetaminophen, caffeine, carbadox and
233 lamotrigine were extensively metabolized in the radish plants with the mass recoveries ranging
234 from 3.0 to 32.1% after 7 days of hydroponic exposure (Li et al., 2018). In some cases, the amount
235 of formed metabolites can even be greater than the accumulated parent compounds (Goldstein et
236 al., 2014; LeFevre et al., 2017; LeFevre et al., 2015; Macherius et al., 2012a; Malchi et al., 2014;
237 Riemenschneider et al., 2017). For example, Macherius et al. (2012a) reported that the total amount
238 of eight phase-II triclosan conjugates was about 5 times that of triclosan in carrot roots after two-
239 month growth in soil.

240 The transpiration stream is considered as the main driving force for uptake and transport
241 of pharmaceuticals from soil to plants (Dodgen et al., 2015). In this study, the three common soil
242 water contents (75, 60 and 45 % of MWHC) did not have an apparent impact on the accumulation
243 of pharmaceuticals in the radish tissues. As shown in Table S4, the concentrations of all detected
244 pharmaceuticals (except caffeine) in radishes showed no significant difference ($p > 0.05$) among
245 three soil water contents. This could be due to the factor that the water present in soil at the three
246 soil water contents could be sufficient for radish growth, and the variation in three soil water
247 contents had barely impact on the water use efficiency by radishes. This is evidenced by the fact
248 that there was no significant difference in the amount of transpired water through the radish

249 growing at the three soil water contents ($p > 0.05$, Figure S2). As a result, the uptake of
250 pharmaceuticals from soil pore water could be similar, and the corresponding BCFs or TF values
251 at both harvesting times demonstrated no significant difference for all measured pharmaceuticals
252 in radish growing at the three soil water contents ($p > 0.05$, Figures S3 and S4).

253

254 3.2. Bioconcentration Factor

255

256 To better understand the accumulation potential of pharmaceuticals in radish from soil
257 system, the BCF values were calculated on the basis of pharmaceutical concentration in bulk soil
258 (BCF_{soil}) and in soil pore water ($BCF_{pore\ water}$). The BCF values were averaged across all radish
259 samples growing at the three soil water contents ($n = 9$), and are shown in Figure 1. The BCF_{pore}
260 $water$ values ranged within 0.26-45.9 mL/g, and the BCF_{soil} values were within the range of 0.02-
261 5.4 g/g. Triclosan manifested the highest $BCF_{pore\ water}$ value 45.9 mL/g at day 28, and
262 acetaminophen had the greatest BCF_{soil} 5.4 g/g at day 28. Tylosin demonstrated the lowest BCF_{soil}
263 and $BCF_{pore\ water}$ compared to the other pharmaceuticals. Other than the two sulfonamide antibiotics
264 (sulfadiazine and sulfamethoxazole), the $BCF_{pore\ water}$ values of the pharmaceuticals were 1.2-423
265 and 1.3-281 times greater than their corresponding BCF_{soil} values at day 28 and 35, respectively.
266 The greater $BCF_{pore\ water}$ than BCF_{soil} values could be due to the less bioavailability of soil-sorbed
267 pharmaceuticals to radish root uptake than pharmaceuticals in soil pore water. For the instance of
268 triclosan, the BCF_{soil} value was < 0.12 g/g at day 28, and its corresponding $BCF_{pore\ water}$ was 45.9
269 mL/g. The strong sorption of triclosan by soil ($K_d = 316.2$ mL/g) could reduce the bioavailability
270 of sorbed chemical, resulting in the substantially diminished BCF_{soil} . Different soils
271 manifest varying sorption for pharmaceuticals hence the bioavailability for plant uptake (Fu et al.,

272 2016b; Goldstein et al., 2014; Malchi et al., 2014). For example, the uptake of triclosan from soils
273 by root vegetables such as radish and carrot demonstrated a wide range of BCF_{soil} from 0.12 to 6.6
274 g/g (Carter et al., 2014; Fu et al., 2016b; Macherius et al., 2012b; Pannu et al., 2012; Prosser et al.,
275 2014). Such difference in BCF_{soil} among the soils (up to 55 times) could be attributed mainly to
276 the varying sorption by soils that lead to the different available fractions of triclosan in pore water
277 to plant uptake. The BCF_{soil} tends to be elusive as evidenced by that the chemical concentration in
278 bulk soil does not reflect its bioavailability in soil (Hung et al., 2009), and thus the BCFs of
279 pharmaceuticals with different soils give no direct insight to the efficiency of pharmaceutical
280 uptake into plants. In contrast, the concentration in pore water serves as a reasonable basis to
281 evaluate the accumulation potential of pharmaceuticals in soil-plant systems. $BCF_{pore\ water}$ could
282 also facilitate the comparison of pharmaceuticals uptake by plants from soils by minimizing the
283 effects of soil sorption. To further clarify this point, BCF_{soil} data of triclosan in radish from this
284 test and a previous study were thus compared in five different soils (Fu et al., 2016b). As illustrated
285 in Figure S5, the BCF_{soil} of triclosan in radish roots varied among five soils by up to 17.5 times
286 between soil A ($BCF_{soil} = 2.10\text{ g/g}$) and soil D (0.12 g/g), with the detailed data given in Table S6.
287 On the other hand, the corresponding $BCF_{pore\ water}$ data could also be obtained from Fu et al. 2016b,
288 by using $BCF_{pore\ water} = BCF_{soil} \times K_d$, where the soil K_d values were provided from another report
289 of the same authors using the same soils (Fu et al., 2016a). As shown in Figure S5, the triclosan
290 $BCF_{pore\ water}$ values (41.1-87.7 mL/g) varied within a factor of 2.2 among five soils (detail in Table
291 S6), which sharply narrowed the variation relative to that of the corresponding BCF_{soil} data.

292

293 *3.3. Relationship between Pharmaceutical Properties and Root Uptake*

294

295 The previous hydroponic studies suggested that the neutral form of organic chemicals
296 generally favors root uptake (Briggs et al., 1982; Tanoue et al., 2012; Trapp, 2000; Wu et al., 2013).
297 To further characterize the relationship between the uptake and pharmaceutical properties in soil-
298 plant systems, pH-adjusted octanol-water partition coefficient (D_{ow}) was used to account for the
299 lipophilicity of neutral speciation of pharmaceuticals in soil pore water (Table 1). Both soil and
300 pore water-based log BCF values are plotted against log D_{ow} for all studied pharmaceuticals. The
301 relationship of log BCF_{pore water} vs. log D_{ow} ($R^2 = 0.48$) or log BCF_{soil} vs. log D_{ow} ($R^2 = 0.04$) is
302 showed in Figures 2A and 2B. These results suggest that partitioning of pharmaceuticals in neutral
303 speciation is not the singular uptake driving force for the accumulation of the investigated
304 pharmaceuticals in roots. The ionic pharmaceutical species could also contribute to the uptake and
305 accumulation in the radish roots.

306 Root uptake of anionic chemicals could be inhibited by the negative electrical potential
307 across plant cell membranes (between -71 and -174 mV), which might repulse anionic
308 pharmaceuticals to approach the cell surfaces (Trapp, 2009). In this study, soil pore water pH value
309 was ~ 8.1 at which the majority of sulfadiazine (92.5%), sulfamethoxazole (99.6%) and monensin
310 (100%) were anionic, thus limiting their penetration into plant cells, as evidenced by a low
311 bioaccumulation in radish roots (BCF_{pore water} < 1.0 mL/g). Carbadox, acetaminophen, and estrone
312 were present in soil pore water primarily in neutral form ($> 95\%$). These chemicals could pass
313 through the cell membrane resulting in relatively higher BCF_{pore water}, 1.5 mL/g for carbadox (log
314 $D_{ow} = -1.22$), 6.6 mL/g for acetaminophen (log $D_{ow} = 0.44$), and 23.1 mL/g for estrone (log $D_{ow} =$
315 3.13).

316 Cationic chemicals could be attracted to plant root cell membranes via electrostatic
317 interaction with negatively-charged plasmalemma (Inoue et al., 1998; Trapp, 2009). Ion trapping

318 could enhance the accumulation of cationic pharmaceuticals in plants due to the alteration of
319 chemical speciation in cell organelles (Trapp, 2004). For example, trimethoprim (weak organic
320 base with $pK_a = 7.12$) existed primarily in neutral form ($> 90\%$) in soil pore water ($pH = 8.1$). The
321 neutral trimethoprim could easily enter plant root cells, and be ionized in vacuole where pH is ~ 5.5
322 (Trapp, 2000, 2009). The majority of trimethoprim is present as cationic species ($> 97\%$) in the
323 vacuole, and could be trapped in vacuole because less effective to cross the membranes. This could
324 be responsible for the observed relatively high accumulation of trimethoprim in roots ($BCF_{\text{pore water}}$
325 $= 4.3 \text{ mL/g}$) in spite of its low $\log D_{ow} = 0.86$.

326 Among the 14 pharmaceuticals measured in radish roots, acetaminophen, caffeine,
327 carbamazepine, lamotrigine, carbadox, estrone and trimethoprim were present primarily in neutral
328 form ($> 90\%$) in soil pore water. A strong positive correlation was found between $\log BCF_{\text{pore water}}$
329 and $\log D_{ow}$ ($R^2 = 0.94$) (Figure 2C), suggesting that partitioning could be the primary factor
330 affecting their accumulation in radish roots. This is consistent with the findings from the previous
331 study of uptake of neutral pharmaceuticals by vegetables from hydroponic solution (Wu et al.,
332 2013). Hyland et al. (2015b) calculated the BCFs based on the chemical concentration in irrigation
333 water, and found the positive correlation between $\log BCF$ and $\log D_{ow}$ for six pharmaceuticals
334 and three flame retardants in both lettuce and strawberry roots grown in soils ($R^2 = 0.78$). However,
335 in this study, the poor relationship was observed between $\log BCF_{\text{soil}}$ and $\log D_{ow}$ for these neutral
336 pharmaceuticals ($R^2 = 0.03$) (Figure 2D), revealing that $BCF_{\text{pore water}}$ is more appropriate to describe
337 the distribution of pharmaceuticals in soil-water-plant systems, rather than the bulk soil-based BCF.

338 In addition to charged species and hydrophobicity, the bioaccumulation of pharmaceuticals
339 could also be affected by chemical molecular weight (MW), plant physiology, metabolism,
340 exposure time, and plant growth rates, etc. For example, MW is considered as another factor that

341 associate with the membrane permeability (Kumar and Gupta, 2016; Topp et al., 1986). Studies
342 on diffusion have indicated that the compounds with MW > 500 g/mol have the restricted
343 membrane permeability (Camenisch et al., 1998), while the compounds with MW > 1000 g/mol
344 are impossible to be absorbed by cells (Sanderson et al., 2004). In this study, the MW of 15
345 pharmaceuticals is in the range of 151.16-916.10 g/mol (Table 1), among the 14 measured
346 chemicals, 12 compounds with MW < 500 g/mol and 2 compounds with MW > 500 g/mol. Ten
347 out of 12 compounds with MW < 500 g/mol (acetaminophen, caffeine, carbamazepine, lamotrigine,
348 estrone, triclosan, trimethoprim, lincomycin, and oxytetracycline) had the greater $BCF_{\text{pore water}}$
349 (1.34-45.9 mL/g) than that of the 2 compounds with MW > 500 g/mol (monensin and tylosin)
350 (0.26-1.0 mL/g). As shown in Figure 2A, tylosin with the highest MW (916.10 g/mol) showed the
351 lowest $BCF_{\text{pore water}}$ (< 0.32 mL/g) even though the most of this antibiotic presented in the pore
352 water as neutral form (~69%) with the intermediate lipophilicity ($\log D_{\text{ow}}$, 1.47). The low $BCF_{\text{pore water}}$
353 of tylosin was also reported in lettuce (< 0.1 mL/g) and carrot (< 0.54 mL/g) (Boxall et al.,
354 2006). The relative high MW of tylosin limiting its permeability across the cell membranes may
355 account for the low bioaccumulation in vegetables.

356

357 *3.4. Pharmaceuticals Translocation*

358

359 Many pharmaceuticals in plant roots can be transported upwards to aerial tissues via xylem
360 driven by transpiration stream, and the magnitude can be quantified by translocation factor (TF).
361 In this study, acetaminophen, oxytetracycline, triclosan and tylosin showed less translocation from
362 roots to leaves with TF < 1 (Figure 3A). Monensin and 17 β -estradiol were not detected in radish
363 leaves. The TF values for other nine pharmaceuticals were > 1, indicating their strong translocation

364 from roots to leaves. No apparent relationship was observed between log TF and log D_{ow} (Figure
365 3B). This is consistent with the previous hydroponic study in which no correlation between log TF
366 and log D_{ow} was found for the pharmaceuticals with log D_{ow} between -3 and 4 (Wu et al., 2013).
367 However, Tanoue et al. (2012) observed that the root-to-leaf translocation is more effective for the
368 pharmaceuticals with moderate hydrophobicity i.e. $1 < \log D_{ow} < 3$. However, our experimental
369 results showed that the hydrophilic caffeine manifested the highest translocation (TF = 36.1). The
370 relatively high transpiration stream concentration factor of caffeine could lead to the higher
371 accumulation in plant leaves (Garvin et al., 2015).

372 The poor relationship between log TF and log D_{ow} suggests that other factors could
373 influence the translocation processes such as the electrostatic interaction with ionizable
374 pharmaceuticals (e.g., ion trapping), plant physiology, molecular size, and in-plant metabolism.
375 For example, ion trapping may decrease the translocation of lamotrigine from roots to leaves.
376 Carbamazepine and lamotrigine exhibit the similar log D_{ow} (2.45 and 2.57, respectively) and MW
377 (236.27 and 256.10 g/mol, respectively). Both chemicals existed as neutral form in the pore water
378 and had the similar $BCF_{\text{pore water}}$ (~11 mL/g). However, the TF of carbamazepine (~8.0) was about
379 4 times greater than that of lamotrigine (~2.0). Lamotrigine is a weak organic base ($pK_a = 5.34$);
380 after neutral species entered the root cell, lamotrigine could be dissociated and accept protons in
381 the cell vacuole (pH ~5.5). In the vacuoles, ~41% of lamotrigine was positively charged, which
382 could be trapped in vacuoles or interact with the negatively charged cell walls. This process could
383 largely reduce the translocation of lamotrigine to radish leaves, compared to the neutral
384 carbamazepine. The less translocation of lamotrigine relative to carbamazepine was also observed
385 in sweet potato, carrot, tomato, and cucumber (Goldstein et al., 2014; Malchi et al., 2014).
386 Goldstein et al. (2018) demonstrated that the transpiration stream concentration factor of

387 carbamazepine (~0.7-0.9) in cucumber was about 4.6 times higher than that of lamotrigine (~0.1-
388 0.25), which is similar to the TF difference (~4 times) in radish between the two chemicals. The
389 TF values of intermediate hydrophobic compounds monensin ($\log D_{ow} = 1.65$) and tylosin (\log
390 $D_{ow} = 1.47$) was 0 and 0.4, respectively, which are much less than the predicted transport based on
391 the hydrophobicity (Tanoue et al., 2012). The relatively large-sized monensin and tylosin with
392 MW > 500 g/mol could limit their transport across the cell membrane and enter xylem, hence
393 decrease their accumulation in leaves (Limmer and Burken, 2014).

394

395 *3.5. Pharmaceutical Distribution in Soil and Relation to Plant Uptake*

396 The K_d , $T_{1/2}$, and residual fractions of pharmaceuticals at day 35 are reported in Table 2.
397 The tested soil demonstrated a range of sorption capacity for the fifteen pharmaceuticals as
398 indicated by the K_d values from 0.3 to 316 mL/g. The half-lives ranged from 5.0 to >35 days, and
399 the concentration profiles of pharmaceuticals in bulk soil and pore water over time are plotted in
400 Figure S6. Pharmaceutical with strong sorption to soil and/or short half-life is generally less
401 bioavailable to plant uptake. For example, 17 β -estradiol had relatively strong sorption to soil (K_d
402 = 34.9 mL/g) and a rapid degradation rate ($T_{1/2} = 7.5$ days), as a result, little 17 β -estradiol was
403 found in radish. In contrast, carbamazepine was weakly sorbed by soil ($K_d = 4.2$ mL/g) and highly
404 persistent in soil ($T_{1/2} > 35$ days, 78.4 % remained after 35 days). Therefore, a large fraction of
405 carbamazepine remained in soil and pore water, which is readily available to root uptake leading
406 to the relatively high accumulation of carbamazepine. This could partially explain the high
407 concentration of carbamazepine (e.g., up to 400 ng/g) found in various vegetables irrigated with
408 treated wastewater (Goldstein et al., 2014; Malchi et al., 2014; Paltiel et al., 2016; Wu et al., 2014).

409 Soil pore water is the media to move pharmaceuticals into plants. Pharmaceutical
410 concentration in soil pore water could be estimated using sorption coefficient (K) obtained from
411 another independent batch equilibration experiment (detailed in SI) in which $C_{\text{pore water}} = C_{\text{soil}} / K$
412 (Chiou et al., 2001). The K values used here were calculated from the slope of sorption isotherm
413 of pharmaceutical by the tested soil (Figure S7 and Table S6). To evaluate the impact of sorption
414 and desorption of pharmaceuticals in soils to the uptake by radish, the $BCF_{\text{pore water}}$ values were
415 estimated using $BCF_{\text{soil}} \times K$, and then compared to the measured $BCF_{\text{pore water}}$ (Figure 4). The
416 predicted $BCF_{\text{pore water}}$ values are within 3.0 times of difference compared to the measured BCF_{pore}
417 $_{\text{water}}$ for all the measured pharmaceuticals at both day 28 and 35, except for estrone and lincomycin
418 are within 3.0-5.5 times of difference. Recall that the measurement of sorption isotherm (to obtain
419 K values) was equilibrated for 48 hours, while plant uptake experiment was conducted for 35 days.
420 The sorption of pharmaceuticals in the pot experiments (without plant) at different sampling
421 intervals during the 35-day studies are also compared to the 48-hour sorption equilibration
422 isotherms (Figure S7). The results revealed that distribution of most pharmaceuticals between soil
423 and water in the 35-day study is similar to the sorption of 48-hour equilibration. The good
424 agreement between the measured and the predicted $BCF_{\text{pore water}}$ indicates that soil-sorbed
425 pharmaceuticals could quickly establish sorption/desorption equilibrium with soil pore water (<
426 48 hours). Therefore, pharmaceutical desorption from soil is not the limiting factor governing the
427 uptake of pharmaceuticals from soil to radish. These results provide the basis for assuming the fast
428 sorption equilibration of pharmaceuticals between soil and pore water when modeling the
429 movement of pharmaceuticals from soil to pore water for plant uptake.

430

431 **4. Conclusion**

432

433 This study reveals that pharmaceuticals could enter plant from contaminated soil, and
434 sorption and biodegradation in soil are the determinants of the bioavailable amounts of
435 pharmaceuticals to plant uptake. The comparison of the bulk soil-based BCF_{soil} and pore water-
436 based $BCF_{pore\ water}$ suggests that the pharmaceuticals in pore water represent the more effective
437 fractions for plant uptake than those sorbed by soil. In addition, the good positive correlation (R^2
438 = 0.93) between $\log BCF_{pore\ water}$ and $\log D_{ow}$ vs. the poor correlation between $\log BCF_{soil}$ and $\log D_{ow}$
439 ($R^2 = 0.03$) for neutral pharmaceuticals indicate that pharmaceuticals in pore water is more
440 appropriate for developing the relationships for predicting the accumulation in plants. We therefore
441 highly recommend $BCF_{pore\ water}$, instead of BCF_{soil} , be used to evaluate the distribution of
442 pharmaceuticals in soil-plant systems. Anionic pharmaceuticals generally showed less uptake than
443 neutral pharmaceuticals. “Ion trapping” effects may enhance the accumulation of basic
444 pharmaceuticals accumulation in the roots but limit the transport to leaves. Our results indicate
445 that pharmaceutical in pore water is an important factor to control their uptake by food crops from
446 contaminated agricultural soils. The results help to better understand the common processes of
447 pharmaceutical transport and distribution in soil-water-plant systems through land application of
448 biosolids or animal manure. However, the experiment may not well mimic the processes of plant
449 uptake of pharmaceuticals via irrigation with treated wastewater. The human dietary intake of
450 these pharmaceuticals via radishes seems to be small, as the measured amounts are much lower
451 than the amount of single medical dose (usually in the range of 10–200 mg). However, little is
452 known about the long-term risks of exposure to the mixture of pharmaceuticals and metabolites.
453 Therefore, more studies are still needed to warrant better understanding of pharmaceutical
454 transport, uptake and metabolism in soil-plant systems.

455

456 **Acknowledgement**

457 This study was supported in part by Agriculture and Food Research Initiative Competitive Grant
458 2016-67017-24514 from USDA National Institute of Food and Agriculture, and Michigan
459 AgBioResearch Project GREEN.

460

461 **Appendix A. Supporting Information**

462 Additional description is available regarding analytical methods, soil-pore water distribution
463 coefficient, pharmaceuticals concentration in plant tissues, sorption measurement using batch
464 equilibration method, radish biomass, BCF values, TF values, and water transpiration, Comparison
465 between the 48 hours sorption isotherms and long term (35 days) sorption isotherms in greenhouse
466 experiment.

467

468 **References**

469

- 470 Blaine, A.C., Rich, C.D., Sedlacko, E.M., Hundal, L.S., Kumar, K., Lau, C., Mills, M.A., Harris, K.M., Higgins,
471 C.P., 2014. Perfluoroalkyl acid distribution in various plant compartments of edible crops grown
472 in biosolids-amended soils. *Environ. Sci. Technol.* 48, 7858-7865.
- 473 Boxall, A.B., Johnson, P., Smith, E.J., Sinclair, C.J., Stutt, E., Levy, L.S., 2006. Uptake of veterinary medicines
474 from soils into plants. *J. Agric. Food. Chem.* 54, 2288-2297.
- 475 Briggs, G.G., Bromilow, R.H., Evans, A.A., 1982. Relationships between lipophilicity and root uptake and
476 translocation of non - ionised chemicals by barley. *Pestic. Sci.* 13, 495-504.
- 477 Briggs, G.G., Bromilow, R.H., Evans, A.A., Williams, M., 1983. Relationships between lipophilicity and the
478 distribution of non - ionised chemicals in barley shoots following uptake by the roots. *Pestic. Sci.*
479 14, 492-500.
- 480 Briggs, G.G., Rigitano, R.L., Bromilow, R.H., 1987. Physico - chemical factors affecting uptake by roots and
481 translocation to shoots of weak acids in barley. *Pestic. Sci.* 19, 101-112.
- 482 Butler, E., Whelan, M.J., Sakrabani, R., van Egmond, R., 2012. Fate of triclosan in field soils receiving
483 sewage sludge. *Environ. Pollut.* 167, 101-109.
- 484 Camenisch, G., Alsenz, J., van de Waterbeemd, H., Folkers, G., 1998. Estimation of permeability by passive
485 diffusion through Caco-2 cell monolayers using the drugs' lipophilicity and molecular weight. *Eur.*
486 *J. Pharm. Sci.* 6, 313-319.
- 487 Carter, L.J., Harris, E., Williams, M., Ryan, J.J., Kookana, R.S., Boxall, A.B., 2014. Fate and uptake of
488 pharmaceuticals in soil-plant systems. *J. Agric. Food. Chem.* 62, 816-825.

489 Carter, L.J., Williams, M., Martin, S., Kamaludeen, S.P., Kookana, R.S., 2018. Sorption, plant uptake and
490 metabolism of benzodiazepines. *Sci. Total. Environ.* 628, 18-25.

491 Carvalho, P.N., Basto, M.C.P., Almeida, C.M.R., Brix, H., 2014. A review of plant–pharmaceutical
492 interactions: from uptake and effects in crop plants to phytoremediation in constructed wetlands.
493 *Environ. Sci. Pollut. Res.* 21, 11729-11763.

494 Chiou, C.T., Sheng, G., Manes, M., 2001. A partition-limited model for the plant uptake of organic
495 contaminants from soil and water. *Environ. Sci. Technol.* 35, 1437-1444.

496 Christou, A., Karaolia, P., Hapeshi, E., Michael, C., Fatta-Kassinos, D., 2017. Long-term wastewater
497 irrigation of vegetables in real agricultural systems: Concentration of pharmaceuticals in soil,
498 uptake and bioaccumulation in tomato fruits and human health risk assessment. *Water Res.* 109,
499 24-34.

500 Chuang, Y.-H., Zhang, Y., Zhang, W., Boyd, S.A., Li, H., 2015. Comparison of accelerated solvent extraction
501 and quick, easy, cheap, effective, rugged and safe method for extraction and determination of
502 pharmaceuticals in vegetables. *J. Chromatogr. A* 1404, 1-9.

503 Clarke, B.O., Smith, S.R., 2011. Review of ‘emerging’ organic contaminants in biosolids and assessment of
504 international research priorities for the agricultural use of biosolids. *Environ. Int.* 37, 226-247.

505 Daughton, C.G., Ternes, T.A., 1999. Pharmaceuticals and personal care products in the environment:
506 agents of subtle change? *Environ. Health Perspect.* 107, 907-938.

507 de Boer, M.A., Hammerton, M., Slootweg, J.C., 2018. Uptake of pharmaceuticals by sorbent-amended
508 struvite fertilisers recovered from human urine and their bioaccumulation in tomato fruit. *Water*
509 *Res.* 133, 19-26.

510 Dodgen, L.K., Ueda, A., Wu, X., Parker, D.R., Gan, J., 2015. Effect of transpiration on plant accumulation
511 and translocation of PPCP/EDCs. *Environ. Pollut.* 198, 144-153.

512 Durán-Alvarez, J.C., Becerril-Bravo, E., Castro, V.S., Jiménez, B., Gibson, R., 2009. The analysis of a group
513 of acidic pharmaceuticals, carbamazepine, and potential endocrine disrupting compounds in
514 wastewater irrigated soils by gas chromatography–mass spectrometry. *Talanta* 78, 1159-1166.

515 Fu, Q., Sanganyado, E., Ye, Q., Gan, J., 2016a. Meta-analysis of biosolid effects on persistence of triclosan
516 and triclocarban in soil. *Environ. Pollut.* 210, 137-144.

517 Fu, Q., Wu, X., Ye, Q., Ernst, F., Gan, J., 2016b. Biosolids inhibit bioavailability and plant uptake of triclosan
518 and triclocarban. *Water Res.* 102, 117-124.

519 Garvin, N., Doucette, W.J., White, J.C., 2015. Investigating differences in the root to shoot transfer and
520 xylem sap solubility of organic compounds between zucchini, squash and soybean using a
521 pressure chamber method. *Chemosphere* 130, 98-102.

522 Goldstein, M., Malchi, T., Shenker, M., Chefetz, B., 2018. Pharmacokinetics in Plants: Carbamazepine and
523 Its Interactions with Lamotrigine. *Environ. Sci. Technol.* 52, 6957-6964.

524 Goldstein, M., Shenker, M., Chefetz, B., 2014. Insights into the uptake processes of wastewater-borne
525 pharmaceuticals by vegetables. *Environ. Sci. Technol.* 48, 5593-5600.

526 Gros, M., Petrović, M., Ginebreda, A., Barceló, D., 2010. Removal of pharmaceuticals during wastewater
527 treatment and environmental risk assessment using hazard indexes. *Environ. Int.* 36, 15-26.

528 Hung, H.-W., Sheng, G.D., Lin, T.-F., Su, Y., Chiou, C.T., 2009. The organic contamination level based on the
529 total soil mass is not a proper index of the soil contamination intensity. *Environ. Pollut.* 157, 2928-
530 2932.

531 Hurtado, C., Domínguez, C., Pérez-Babace, L., Cañameras, N., Comas, J., Bayona, J.M., 2016. Estimate of
532 uptake and translocation of emerging organic contaminants from irrigation water concentration
533 in lettuce grown under controlled conditions. *J. Hazard. Mater.* 305, 139-148.

534 Hyland, K.C., Blaine, A.C., Dickenson, E.R., Higgins, C.P., 2015. Accumulation of contaminants of emerging
535 concern in food crops—part 1: Edible strawberries and lettuce grown in reclaimed water. *Environ.*
536 *Toxicol. Chem.* 34, 2213-2221.

537 Inoue, J., Chamberlain, K., Bromilow, R.H., 1998. Physicochemical factors affecting the uptake by roots
538 and translocation to shoots of amine bases in barley. *Pest Manage. Sci.* 54, 8-21.

539 Karnjanapiboonwong, A., Chase, D.A., Cañas, J.E., Jackson, W.A., Maul, J.D., Morse, A.N., Anderson, T.A.,
540 2011. Uptake of 17 α -ethynylestradiol and triclosan in pinto bean, *Phaseolus vulgaris*. *Ecotoxicol.*
541 *Environ. Saf.* 74, 1336-1342.

542 Kinney, C.A., Furlong, E.T., Werner, S.L., Cahill, J.D., 2006. Presence and distribution of wastewater -
543 derived pharmaceuticals in soil irrigated with reclaimed water. *Environ. Toxicol. Chem.* 25, 317-
544 326.

545 Kolpin, D.W., Furlong, E.T., Meyer, M.T., Thurman, E.M., Zaugg, S.D., Barber, L.B., Buxton, H.T., 2002.
546 Pharmaceuticals, hormones, and other organic wastewater contaminants in US streams, 1999-
547 2000: A national reconnaissance. *Environ. Sci. Technol.* 36, 1202-1211.

548 Kumar, K., Gupta, S.C., 2016. A framework to predict uptake of trace organic compounds by plants. *J.*
549 *Environ. Qual.* 45, 555-564.

550 LeFevre, G.H., Lipsky, A., Hyland, K.C., Blaine, A.C., Higgins, C.P., Luthy, R.G., 2017. Benzotriazole (BT) and
551 BT plant metabolites in crops irrigated with recycled water. *Environ. Sci. Water Res. Technol.* 3,
552 213-223.

553 LeFevre, G.H., Müller, C.E., Li, R.J., Luthy, R.G., Sattely, E.S., 2015. Rapid phytotransformation of
554 benzotriazole generates synthetic tryptophan and auxin analogs in *Arabidopsis*. *Environ. Sci.*
555 *Technol.* 49, 10959-10968.

556 Li, Y., Chuang, Y.-H., Sallach, J.B., Zhang, W., Boyd, S.A., Li, H., 2018. Potential metabolism of
557 pharmaceuticals in radish: Comparison of in vivo and in vitro exposure. *Environ. Pollut.* 242, 962-
558 969.

559 Limmer, M.A., Burken, J.G., 2014. Plant Translocation of Organic Compounds: Molecular and
560 Physicochemical Predictors. *Environ. Sci. Technol. Lett.* 1, 156-161.

561 Macherius, A., Eggen, T., Lorenz, W., Moeder, M., Ondruschka, J., Reemtsma, T., 2012a. Metabolization
562 of the bacteriostatic agent triclosan in edible plants and its consequences for plant uptake
563 assessment. *Environ. Sci. Technol.* 46, 10797-10804.

564 Macherius, A., Eggen, T., Lorenz, W.G., Reemtsma, T., Winkler, U., Moeder, M., 2012b. Uptake of
565 galaxolide, tonalide, and triclosan by carrot, barley, and meadow fescue plants. *J. Agric. Food.*
566 *Chem.* 60, 7785-7791.

567 Madikizela, L., Ncube, S., Chimuka, L., 2018. Uptake of pharmaceuticals by plants grown under hydroponic
568 conditions and natural occurring plant species: A review. *Sci. Total Environ.* 636, 477-486.

569 Malchi, T., Maor, Y., Tadmor, G., Shenker, M., Chefetz, B., 2014. Irrigation of root vegetables with treated
570 wastewater: evaluating uptake of pharmaceuticals and the associated human health risks. *Environ.*
571 *Sci. Technol.* 48, 9325-9333.

572 McFarlane, C., Trapp, S., 1994. Plant contamination: modeling and simulation of organic chemical
573 processes. CRC Press.

574 McClellan, K., Halden, R.U., 2010. Pharmaceuticals and personal care products in archived US biosolids
575 from the 2001 EPA national sewage sludge survey. *Water Res.* 44, 658-668.

576 Miller, E.L., Nason, S.L., Karthikeyan, K., Pedersen, J.A., 2016. Root Uptake of Pharmaceuticals and
577 Personal Care Product Ingredients. *Environ. Sci. Technol.* 50, 525-541.

578 Naseer, S., Lee, Y., Lapierre, C., Franke, R., Nawrath, C., Geldner, N., 2012. Casparian strip diffusion barrier
579 in *Arabidopsis* is made of a lignin polymer without suberin. *Proc. Natl. Acad. Sci. U. S. A.* 109,
580 10101-10106.

581 Paltiel, O., Fedorova, G., Tadmor, G., Kleinstern, G., Maor, Y., Chefetz, B., 2016. Human Exposure to
582 Wastewater-Derived Pharmaceuticals in Fresh Produce: A Randomized Controlled Trial Focusing
583 on Carbamazepine. *Environ. Sci. Technol.* 50, 4476-4482.

584 Pan, M., Wong, C.K., Chu, L., 2014. Distribution of antibiotics in wastewater-irrigated soils and their
585 accumulation in vegetable crops in the Pearl River Delta, southern China. *J. Agric. Food. Chem.* 62,
586 11062-11069.

587 Pannu, M.W., Toor, G.S., O'Connor, G.A., Wilson, P.C., 2012. Toxicity and bioaccumulation of biosolids -
588 borne triclosan in food crops. *Environ. Toxicol. Chem.* 31, 2130-2137.

589 Prosser, R.S., Lissemore, L., Topp, E., Sibley, P.K., 2014. Bioaccumulation of triclosan and triclocarban in
590 plants grown in soils amended with municipal dewatered biosolids. *Environ. Toxicol. Chem.* 33,
591 975-984.

592 Riemenschneider, C., Seiwert, B., Moeder, M., Schwarz, D., Reemtsma, T., 2017. Extensive transformation
593 of the pharmaceutical carbamazepine following uptake into intact tomato plants. *Environ. Sci.*
594 *Technol.* 51, 6100-6109.

595 Sallach, J.B., Zhang, Y., Hodges, L., Snow, D., Li, X., Bartelt-Hunt, S., 2015. Concomitant uptake of
596 antimicrobials and Salmonella in soil and into lettuce following wastewater irrigation. *Environ.*
597 *Pollut.* 197, 269-277.

598 Sanderson, H., Johnson, D.J., Reitsma, T., Brain, R.A., Wilson, C.J., Solomon, K.R., 2004. Ranking and
599 prioritization of environmental risks of pharmaceuticals in surface waters. *Regul. Toxicol. Pharm.*
600 39, 158-183.

601 Schreiber, L., 2010. Transport barriers made of cutin, suberin and associated waxes. *Trends Plant Sci.* 15,
602 546-553.

603 Shenker, M., Harush, D., Ben-Ari, J., Chefetz, B., 2011. Uptake of carbamazepine by cucumber plants—a
604 case study related to irrigation with reclaimed wastewater. *Chemosphere* 82, 905-910.

605 Sui, Q., Huang, J., Deng, S., Chen, W., Yu, G., 2011. Seasonal variation in the occurrence and removal of
606 pharmaceuticals and personal care products in different biological wastewater treatment
607 processes. *Environ. Sci. Technol.* 45, 3341-3348.

608 Tanoue, R., Sato, Y., Motoyama, M., Nakagawa, S., Shinohara, R., Nomiyama, K., 2012. Plant uptake of
609 pharmaceutical chemicals detected in recycled organic manure and reclaimed wastewater. *J.*
610 *Agric. Food. Chem.* 60, 10203-10211.

611 Topp, E., Scheunert, I., Attar, A., Korte, F., 1986. Factors affecting the uptake of ¹⁴C-labeled organic
612 chemicals by plants from soil. *Ecotoxicol. Environ. Saf.* 11, 219-228.

613 Trapp, S., 2000. Modelling uptake into roots and subsequent translocation of neutral and ionisable organic
614 compounds. *Pest Manage. Sci.* 56, 767-778.

615 Trapp, S., 2004. Plant uptake and transport models for neutral and ionic chemicals. *Environ. Sci. Pollut.*
616 *Res.* 11, 33-39.

617 Trapp, S., 2009. Bioaccumulation of polar and ionizable compounds in plants, *Ecotoxicology modeling.*
618 Springer, pp. 299-353.

619 Trapp, S., Pussemier, L., 1991. Model calculations and measurements of uptake and translocation of
620 carbamates by bean plants. *Chemosphere* 22, 327-339.

621 Vanderford, B.J., Snyder, S.A., 2006. Analysis of pharmaceuticals in water by isotope dilution liquid
622 chromatography/tandem mass spectrometry. *Environ. Sci. Technol.* 40, 7312-7320.

623 Vazquez-Roig, P., Segarra, R., Blasco, C., Andreu, V., Picó, Y., 2010. Determination of pharmaceuticals in
624 soils and sediments by pressurized liquid extraction and liquid chromatography tandem mass
625 spectrometry. *J. Chromatogr. A* 1217, 2471-2483.

626 Waller, N.J., Kookana, R.S., 2009. Effect of triclosan on microbial activity in Australian soils. *Environ. Toxicol.*
627 *Chem.* 28, 65-70.

628 Wu, C., Spongberg, A.L., Witter, J.D., 2009. Adsorption and degradation of triclosan and triclocarban in
629 soils and biosolids-amended soils. *J. Agric. Food. Chem.* 57, 4900-4905.

630 Wu, C., Spongberg, A.L., Witter, J.D., Fang, M., Czajkowski, K.P., 2010. Uptake of pharmaceutical and
631 personal care products by soybean plants from soils applied with biosolids and irrigated with
632 contaminated water. *Environ. Sci. Technol.* 44, 6157-6161.
633 Wu, X., Conkle, J.L., Ernst, F., Gan, J., 2014. Treated wastewater irrigation: uptake of pharmaceutical and
634 personal care products by common vegetables under field conditions. *Environ. Sci. Technol.* 48,
635 11286-11293.
636 Wu, X., Ernst, F., Conkle, J.L., Gan, J., 2013. Comparative uptake and translocation of pharmaceutical and
637 personal care products (PPCPs) by common vegetables. *Environ. Int.* 60, 15-22.
638 Xu, J., Wu, L., Chang, A.C., 2009. Degradation and adsorption of selected pharmaceuticals and personal
639 care products (PPCPs) in agricultural soils. *Chemosphere* 77, 1299-1305.

640
641
642
643
644
645
646
647
648
649
650
651
652
653
654
655
656
657
658

Tables and Figures

659
660
661
662

663 **Table 1.** Summary of molecular weight (MW), *n*-octanol–water partition coefficient ($\log K_{ow}$), pH-
664 adjusted octanol–water partition coefficient to the basis of neutral speciation ($\log D_{ow}$), acidic
665 dissociation constant (pK_a), and fraction of neutral and ionic pharmaceutical speciation in soil pore water
666 (pH \approx 8.1).

Pharmaceutical	MW (g/mol)	$\log K_{ow}^a$	$\log D_{ow}$	pK_a	Neutral fraction (%) ⁱ	Ionic fraction (%) ⁱ
Acetaminophen	151.16	0.46	0.44	9.38 (acid) ^a	95.2	4.8 (anion)
Caffeine	194.19	-0.07	-0.07	—	100	—
Carbamazepine	236.27	2.45	2.45	2.3 (acid), 13.9 (base) ^b	100	—
Sulfadiazine	250.28	-0.09	-1.21	2.01 (base), 6.99 (acid) ^c	7.5	92.5 (anion)
Sulfamethoxazole	253.28	0.89	-1.49	1.6 (base), 5.7 (acid) ^d	0.42	99.6 (anion)
Lamotrigine	256.10	2.57 ^e	2.57	5.34 (base) ^e	99.8	0.2 (cation)

Carbadox	262.22	-1.22	-1.22	1.8 (base), 10.5 (acid) ^c	99.6	0.4 (anion)
Estrone	270.37	3.13	3.13	10.77 (acid) ^h	99.8	0.2 (anion)
17 β -estradiol	272.38	4.01	4.01	10.71 (acid) ^h	99.8	0.2 (anion)
Triclosan	289.54	4.76	4.36	7.9 (acid) ^a	39.8	60.2 (anion)
Trimethoprim	290.32	0.91	0.86	7.12 (base) ^a	90.1	9.9 (cation)
Lincomycin	406.54	0.2	0.08	7.6 (base) ^a	75.1	24.9 (cation)
Oxytetracycline	460.43	-0.9	-1.76	3.23 (acid), 7.32 (acid), 9.11 (base) ^f	13.7	86.3 (anion)
Monensin	692.87	5.43	1.65	4.3 (acid) ^a	0.02	100 (anion)
Tylosin	916.10	1.63	1.47	7.73 (base) ^g	69.1	30.9 (cation)

667

668 ^aFrom TOXNET database: <http://toxnet.nlm.nih.gov/index.html>, ^bDodgen et al.(2015), ^cChuang et al.

669 (2015), ^dTanoue et al. (2012), ^eMalchi et al. (2014), ^fSassman and Lee (2005), ^gBoxall et al. (2006), ^h

670 Lewis and Archer (1979), and ⁱcalculated on the pH-pK_a relationship.

671

672

673

674

675

676

677

678 **Table 2.** Soil-water distribution coefficient (K_d), half-life ($T_{1/2}$), and residual fractions of pharmaceuticals

679 in soil at the end of experiment (35 days). The values are the average of nine soil samples at three soil water

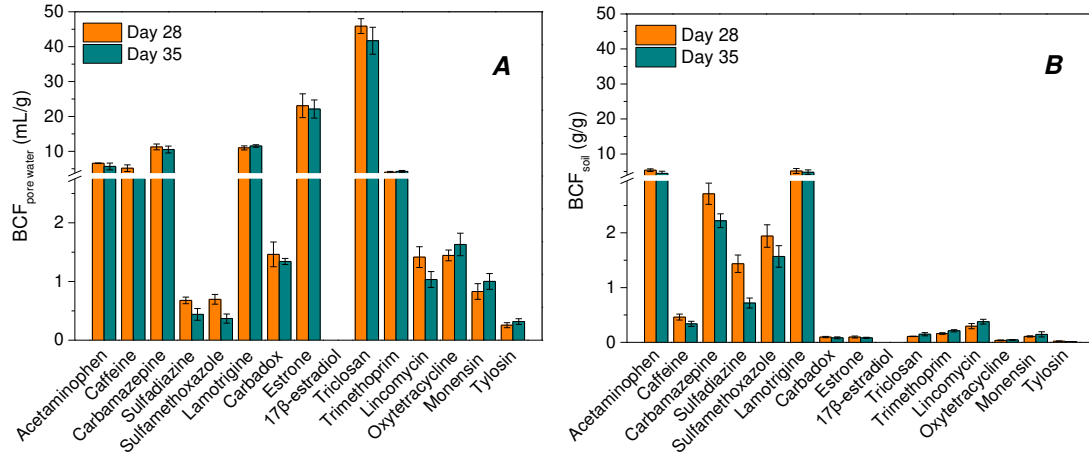
680 contents (mean \pm standard deviation, n = 9).

Pharmaceutical	K_d (mL/g) ^a	$T_{1/2}$ (days)	Residual fraction (%)
Acetaminophen	0.5 \pm 0.1	6.1 \pm 0.1	1.6 \pm 0.1
Caffeine	7.5 \pm 1.3	8.6 \pm 0.5	7.5 \pm 1.1
Carbamazepine	4.2 \pm 0.5	>35	78.4 \pm 2.4
Sulfadiazine	0.8 \pm 0.1	8.0 \pm 0.2	2.1 \pm 0.2
Sulfamethoxazole	0.3 \pm 0.0	7.2 \pm 0.4	1.5 \pm 0.2
Lamotrigine	3.0 \pm 0.2	27.8 \pm 1.1	31.5 \pm 3.1
Carbadox	15.4 \pm 0.6	>35	58.5 \pm 1.4
Estrone	116.3 \pm 20.3	10.5 \pm 0.2	9.9 \pm 0.9
17 β -estradiol	34.9 \pm 1.8	7.5 \pm 0.1	3.6 \pm 0.3

Triclosan	316.2 ± 21.9	>35	70.7 ± 2.7
Trimethoprim	25.2 ± 3.1	>35	55.0 ± 0.8
Lincomycin	1.9 ± 0.2	5.0 ± 0.2	0.6 ± 0.1
Oxytetracycline	33.6 ± 1.5	29.0 ± 1.2	33.9 ± 1.4
Monensin	3.5 ± 0.3	14.3 ± 0.5	18.5 ± 0.8
Tylosin	9.7 ± 0.3	>35	50.9 ± 2.7

681
682
683

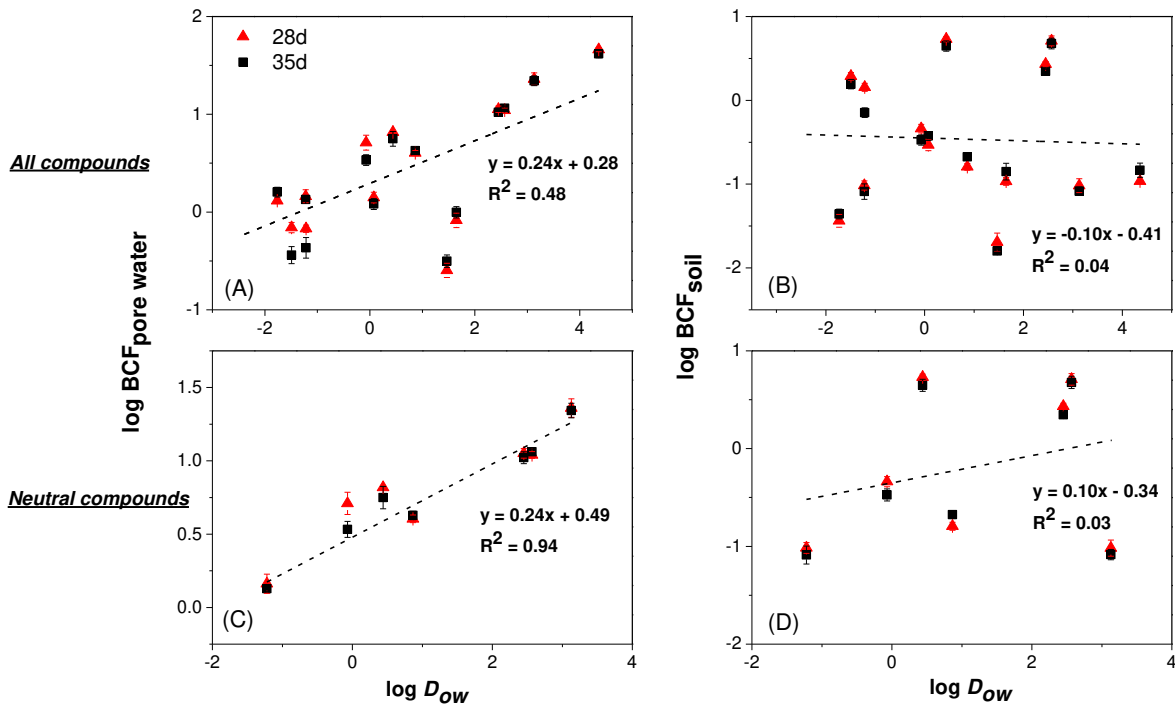
^a Average the data sampled at day 1, 3, 7, 14, 21, 28 and 35 of the radish-free controls.



684

685 **Figure 1.** Bioconcentration factors (BCF) of pharmaceuticals in radish roots on the basis of their
 686 concentrations in pore water (A) and in bulk soil (B). The values are the average of nine radish roots samples
 687 collected at the three soil water contents (one sample per pot, n = 9). The error bars represent the standard
 688 deviations.

689



690

691 **Figure 2.** Linear relationship between $\log BCF$ of radish roots and $\log D_{ow}$ for all measured pharmaceuticals

692 (A and B) and the neutral pharmaceuticals (C and D).

693

694

695

696

697

698

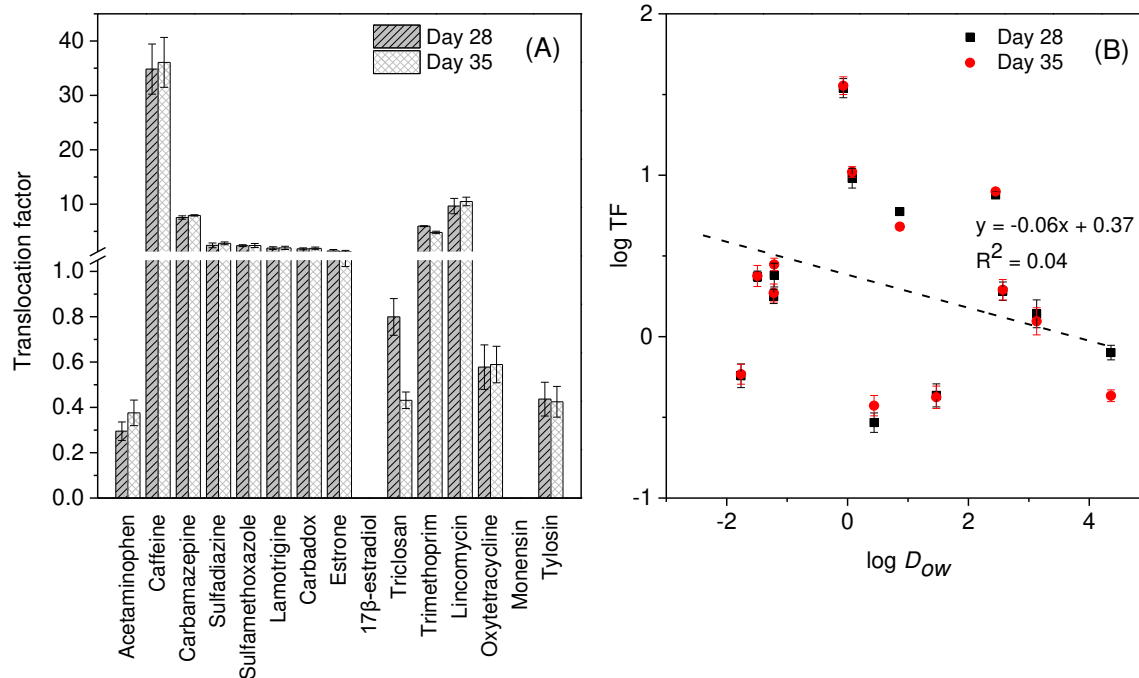
699

700

701

702

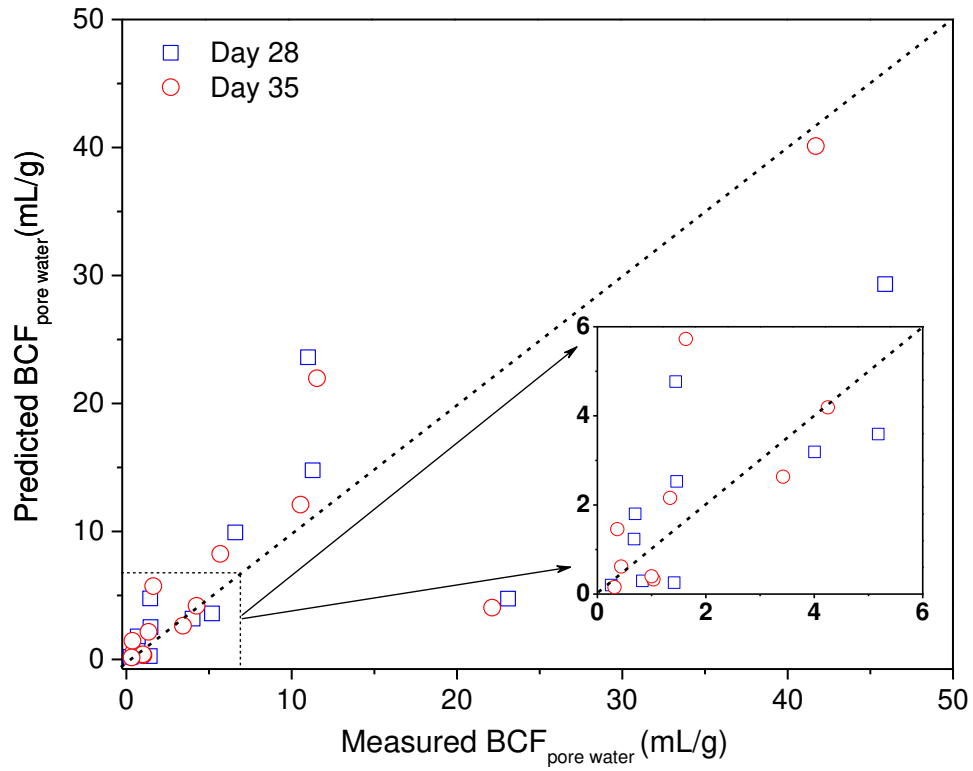
703



704 **Figure 3.** (A) Translocation factor (TF) of pharmaceuticals, and (B) relation between log TF and log D_{ow} .
 705

706 The values are the average of nine radish growing at three soil water contents. The error bars represent the
 707 standard deviation (one sample per pot, n = 9).

708
 709
 710
 711
 712
 713
 714
 715
 716
 717
 718
 719
 720
 721
 722



723

724 **Figure 4.** Comparison between predicted and measured pore water-based bioconcentration factors (BCF
 725 _{pore water}) of pharmaceuticals in radish roots.

726