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1 Potential metabolism of pharmaceuticals in radish: Comparison of *in vivo* and

2 *in vitro* exposure

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8 ABSTRACT

9 Metabolism of pharmaceuticals in plants is important to evaluate their fate and accumulation in vegetables, and subsequently the risks to human health. However, limited knowledge is available 10 11 to evaluate the metabolism of pharmaceuticals in plants due to the lack of appropriate research 12 approaches. In this study, radish was selected as a model plant to investigate the metabolism of 13 pharmaceuticals in intact plants (in vivo) growing in hydroponic solution and in plant tissue enzyme extracts (in vitro). For caffeine, six phase-I demethylation metabolites identified in the 14 15 intact radish plant were also found in the plant enzyme extracts. After 7 days of in vivo exposure, 16 the amount of the identified metabolites was about 5.4 times greater than the parent caffeine in 17 radish roots. Furthermore, the metabolism potential of fifteen pharmaceuticals in radish was 18 evaluated on the basis of mass balance. After 7 days of hydroponic exposure, oxytetracycline, 19 trimethoprim, carbamazepine, lincomycin, monensin and tylosin manifested relatively less extent of metabolism with the mass recoveries ranging from 52.3 to 78.2%. In contrast, 17 β -estradiol, 20 21 sulfamethoxazole, sulfadiazine, estrone, triclosan, acetaminophen, caffeine, carbadox and lamotrigine underwent extensive metabolism with only 3.0 to 32.1% of the parent compound 22 recovered. In the *in vitro* system, 17 β-estradiol, estrone, triclosan, oxytetracycline, 23 acetaminophen, sulfadiazine and sulfamethoxazole were readily metabolized in radish root 24

enzyme extracts with 1.8 to 34.0% remaining after 96-h exposure. While in the leaf enzyme extracts, only triclosan was rapidly metabolized with 49.2% remaining, and others pharmaceuticals were $\geq 60\%$, indicating that the varying extents of metabolism occurred in different plant parts. This study highlights the importance of pharmaceutical metabolism in plants, and suggests that plant tissue enzyme extracts could serve as an alternative tool to assess pharmaceutical metabolism in plants.

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Capsule: Similar metabolism patterns were observed for rapidly metabolized pharmaceuticals in
both *in vivo* (radish tissue enzyme extracts) and *in vivo* (the intact plant) exposure.

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35 *Keywords:* Plant metabolism; Plant uptake; Pharmaceuticals; Plant tissue enzyme extracts.

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37 **1. Introduction**

38 Reclaimed water reuse in agricultural irrigation has been increasingly practiced to alleviate the burden of water scarcity, especially in arid and semiarid regions (Bischel et al., 2011; 39 Sato et al., 2013). Biosolids and animal manures are also commonly applied to agricultural lands 40 41 for their fertility values, and as a convenient disposal approach as well (Clarke and Smith, 2011; Cogger et al., 2013; Kumar et al., 2005). These practices also release pharmaceuticals to the 42 environment, resulting in their ubiquitous presence in soils with concentrations at ng kg⁻¹ to μ g 43 kg⁻¹ levels (Chen et al., 2014; Durán-Alvarez et al., 2009; Kinney et al., 2006; Vazquez-Roig et 44 45 al., 2010). These pharmaceuticals could enter vegetables from soils via root uptake, which serves 46 as the starting point in the food chain of human and animal dietary consumption. It has been well 47 documented that vegetables could accumulate a range of pharmaceuticals from soils (Malchi et

al., 2014; Prosser and Sibley, 2015; Tanoue et al., 2012; Wu et al., 2013). However, only a few
attempts have been made to elucidate the metabolism of pharmaceuticals in plants. In addition to
uptake and translocation, metabolic transformation of pharmaceuticals is also an important
process influencing their residue levels in plants and their potential risk of food safety.

52 Pharmaceuticals could be metabolized in plants by multiple pathways and form a variety 53 of transformation products (Macherius et al., 2012; Miller et al., 2016; Riemenschneider et al., 2017). In general, the metabolic processes involve three phases of reactions. Pharmaceuticals 54 55 could be mediated by plant enzymes and form more polar and water-soluble products via phase I reactions (oxidation, reduction and hydrolysis). In phase II reactions, pharmaceuticals and their 56 phase-I metabolites could be conjugated to endogenous plant biomolecules such as amino acids, 57 sugars and glutathione, and form relatively larger-sized molecules. Compared to their parent 58 compounds these conjugates generally demonstrate an enhanced water solubility and higher 59 mobility in plants. These conjugates formed by phase II reactions could be sequestered in plant 60 61 vacuoles or cell walls by phase III reactions (He et al., 2017; Macherius et al., 2012; Sandermann, 1992). The metabolic transformations of some pharmaceuticals in plants have been studied such 62 as triclosan (Macherius et al., 2012), benzotriazole (LeFevre et al., 2015), ibuprofen (He et al., 63 64 2017), diclofenac (Huber et al., 2012), iopromide (Cui et al., 2017) and carbamazepine (Riemenschneider et al., 2017). Carbamazepine was moderately metabolized in tomato after 35 65 66 days of exposure, 21 phase I and II transformation products were identified, and their total amount was equivalent to ~ 45% of carbamazepine accumulated in plants (Riemenschneider et 67 al., 2017). Among these metabolites, the 10,11-epoxycarbamazepine manifested even greater 68 69 toxic potency than carbamazepine (Tomson et al., 1990). In many cases, the amount of formed 70 metabolites exceeded that of the parent compounds remaining in plants. For instance, the sum of 71 eight phase II triclosan conjugates was 5 times greater than triclosan in carrots during two-month growth in greenhouse (Macherius et al., 2012). In another study, three major metabolites of 72 73 benzotriazole were found to be >1.5 times more than benzotriazole after 8-day exposure in hydroponic Arabidopsis (LeFevre et al., 2015). Besides the intact plants, the metabolism of 74 pharmaceuticals was also studied using plant cell cultures (Fu et al., 2017; Huber et al., 2009; 75 76 Macherius et al., 2012; Marsik et al., 2017; Sauvêtre et al., 2018; Wu et al., 2016). Triclosan, naproxen, diclofenac, ibuprofen, gemfibrozil, sulfamethoxazole and atorvastatin were found to 77 rapidly disappeared in carrot cell cultures with 0.4 to 47.3% remaining after 90 h of exposure 78 79 (Wu et al., 2016). The lack of assessment of metabolism could underestimate the total accumulation of pharmaceuticals and metabolites in plants and the potential risk related to the 80 consumption of contaminated agricultural products. Paltiel et al. (2016) recently reported that 81 human consumption of fresh produce irrigated with reclaimed water could lead to the appearance 82 of bioactive metabolites of carbamazepine in their urine. 83

84 *In vivo* exposure (intact plants) is an effective way to examine the fate and metabolism of pharmaceuticals in plant-water (or -soil) systems. However, the intact plant experiments 85 86 conducted in hydroponic solution or soil are usually labor-intensive and time-consuming. The 87 presence of microorganisms and root exudates also influence pharmaceutical metabolism, which cannot be excluded from the in vivo studies (Yu et al., 2013; Zhang et al., 2016). The uptake and 88 89 transport processes could further complicate the metabolism of pharmaceuticals in plants by 90 governing the input and output of parent compounds/metabolites in a specific plant part. In 91 plants, enzyme-mediated transformation has been considered as a major process to the metabolism of xenobiotics in plants (Huang et al., 2013; Van Eerd et al., 2003). Plant enzyme 92 93 extracts have been used in the *in vitro* system to investigate the enzyme-mediated metabolism of 94 polybrominated diphenyl ethers (Huang et al., 2013), estrogens (Card et al., 2013) and metformin 95 (Cui and Schröder, 2016). These studies showed that the metabolites from the reactions with 96 plant enzyme extracts were similar to the products formed in the intact plants. Therefore, plant 97 enzyme extracts may serve as an alternative system to provide a simple and quick approach to 98 evaluate the metabolism of pharmaceuticals in plants, which is minimally affected by plant 99 microbial, uptake and transport processes.

The objective of this study was to compare the pharmaceutical metabolism in radish 100 101 using both *in vitro* and *in vivo* experiments. Radish was selected because this root vegetable is 102 usually consumed in raw, and might represent the worst case of exposure scenarios. Caffeine was selected as one representative pharmaceutical for further evaluation of the formed metabolites in 103 both intact radish and radish tissue enzyme extracts because it is one of the most frequently 104 detected pharmaceuticals in vegetables (Goldstein et al., 2014; Malchi et al., 2014; Wu et al., 105 2014; Wu et al., 2013). In addition, we expanded the experiments to fifteen commonly-used 106 107 pharmaceuticals with a wide range of physicochemical properties to assess their metabolism potential in radish. This study provides new knowledge on the metabolic transformation of 108 pharmaceuticals in plants. The *in vitro* exposure method could serve as an alternative to quickly 109 110 screen the metabolism potential of pharmaceuticals in plants.

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112 **2. Materials and methods**

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114 2.1. Chemicals and regents

Fifteen pharmaceuticals were purchased from Sigma-Aldrich (St. Louis, MO, USA)
including acetaminophen, caffeine, carbamazepine, sulfadiazine, sulfamethoxazole, lamotrigine,

117 carbadox, estrone, 17 β-estradiol, triclosan, trimethoprim, lincomycin, oxytetracycline, monensin and tylosin. These pharmaceuticals were selected because they are frequently detected in the 118 environment, and demonstrate a wide range of physicochemical properties (Table S1). Caffeine 119 metabolites xanthine, 3-methylxanthine, 7-methylxanthine, theophylline, paraxanthine and 120 theobromine were purchased from Sigma-Aldrich (St. Louis, MO, USA). Ceramic homogenizers, 121 122 octadecylsilane (C18), primary secondary amine (PSA), and graphitized carbon black (GCB) Technologies (Santa Clara, CA, USA). Disodium 123 purchased from Agilent were ethylenediaminetetraacetate (Na₂EDTA), formic acid, glacial acetic acid, and sodium chloride 124 125 (NaCl) were purchased from J.T. Baker (Phillipsburg, NJ, USA). All organic solvents were of HPLC grade from Fisher (Fair Lawn, NJ, USA). Water used in this study was produced from a 126 Milli-Q water purification system (Billerica, MA, USA). 127

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129 2.2. In vivo hydroponic experiment

Radish seeds (Raphanus sativus, Burpee & Co., Warminster, PA) were germinated on 130 moistened filter paper for 4 days. The seedlings were then transferred to a plastic container with 131 10 L of nutrient solution (pH 6.5) (Hydrodynamics International, Lansing, MI, USA). After 14 132 133 days of growth, the radish plants (~10 cm in height) were used in the exposure experiments. Each radish plant was carefully transferred to a 250-mL glass jar containing 230 mL of nutrient 134 135 solution with spiked pharmaceuticals. This exposure experiment was conducted with two treatments (1) the radish was exposed to 200 ng mL⁻¹ of caffeine in nutrient solution; (2) the 136 radish was exposed to a mixture of fifteen pharmaceuticals with the initial concentration of 100 137 ng mL⁻¹ for each compound. The solution was continuously aerated via Teflon tubing connected 138 139 to a fusion air pump. The jars were wrapped with aluminum foil to prevent the potential

photolysis of pharmaceuticals and to minimize algae growth. Pharmaceutical-free controls 140 (radish only) and pharmaceutical solution controls (without radish) were also conducted in the 141 experiments. All controls and exposure experiments were performed in triplicate. During the 142 experiment, nutrient solution (without pharmaceuticals) was added daily to each jar to 143 compensate the water loss from plant transpiration. The glass jars were placed in a controlled 144 145 growth chamber at 20 ± 2 °C. The light cycle was set up as 10 h of fluorescent light every day with the intensity of 150 mmol/m²/s followed by 14 h of darkness. For the treatment with only 146 147 caffeine, radish roots, leaves and solution were sampled at day 1, 3 and 7. For the exposure of the mixture of pharmaceuticals, aqueous solution, radish roots and leaves were collected after 7 days 148 of growth. The radish samples were thoroughly rinsed with deionized water, wiped with tissue 149 paper, weighed, chopped into small pieces, freeze-dried and ground to fine powders. All samples 150 were stored in freezer at -20 °C prior to the extraction of pharmaceuticals. 151

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153 2.3. In vitro reactions with radish enzyme extracts

Pharmaceutical metabolism was also examined in vitro using the crude enzyme extracts 154 from radish tissues. Radish enzyme extracts were prepared using the method reported by Card et 155 156 al. (Card et al., 2013). In brief, fresh radish tissues were separated into roots and leaves, chopped into small pieces, quickly frozen, ground into powders in liquid nitrogen, and then extracted with 157 158 50 mM potassium phosphate buffer (pH 7.0) at the mass to solution ratio of 1: 2 (g: mL) in ice 159 for 30 min. The homogenate was centrifuged at 8000 g for 40 min at 4 °C. The supernatant of tissue enzyme extracts was collected, passed through a 0.22-µm cellulose ester membrane 160 (Millipore, Cork, Ireland), and immediately stored in an ice bath prior to use. 161

The *in vitro* exposure experiment was performed by mixing 200 ng mL⁻¹ of caffeine or a 162 mixture of fifteen pharmaceuticals at 100 ng mL⁻¹ for each chemical with the radish enzyme 163 extracts. The reactant mixtures were incubated in 1.5-mL centrifuge tubes in a water bath (25 °C), 164 and the reaction was quenched by adding 50 µL of glacial acetic acid. The experiments were 165 conducted in triplicate, along with enzyme extracts-free controls. The resultant solution was 166 167 sampled at the time intervals of 0, 4, 10, 24, 48, 72 and 96 h. An aliquot of the solution (0.5 mL) was diluted to 1.0 mL with methanol and subject to pharmaceutical analysis. The proteins 168 concentration in radish enzyme extracts was used to estimate the activity of the enzyme mixture, 169 170 which was measured during the experiment using Bradford assay (Bradford, 1976). The proteins concentration in the radish enzyme extracts demonstrated relatively stable during 96 h of 171 reaction (Figure S1). 172

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174 2.4. Analysis of pharmaceuticals

Pharmaceuticals in radish root and leaf samples were extracted using a modified 175 QuEChERS method (Chuang et al., 2015). The detailed extraction procedure is given in the 176 supporting information (SI). Hydroponic solution was filtered through a 0.22-um cellulose ester 177 178 membrane, and an aliquot of the filtrate (0.5 mL) was diluted to 1.0 mL with methanol and injected directly into a LC-MS/MS system. The LC-MS/MS system consisted of a Shimadzu 179 prominence high-performance liquid chromatography (Columbia, MD, USA) coupled to a Sciex 180 181 4500 triple quadrupole mass spectrometer (Foster City, CA, USA), and a 50 mm \times 2.1 mm Agilent C18 column (Torrance, CA, USA). The optimized conditions and quality assurance of 182 183 the LC-MS/MS is detailed in SI. Multiple reaction monitoring (MRM) parameters are listed in Tables S2 and S3. The corresponding matrix recoveries and method detection limits (MDLs) forthe studied pharmaceuticals are summarized in Tables S4-S7.

The metabolism of pharmaceuticals in radish enzyme extracts was fit to the first-order kinetic model $C_i = C_0 e^{-kt}$, where C_i and C_0 is pharmaceutical concentration at sampling time *t* (h) and the beginning time, respectively, and *k* is the rate constant (h⁻¹). The dissipation half-life ($T_{1/2}$) was calculated as $T_{1/2} = \ln 2/k$.

190

191 **3. Results and discussion**

192 *3.1. Biotransformation of caffeine in hydroponic system*

The mass distribution of caffeine in radish leaves, roots and hydroponic solution is shown 193 in Figure S2. On the basis of the initial amount of caffeine added to the solution, 11.8, 48.9 and 194 82.6% disappeared in the system after 1, 3 and 7 days of exposure, respectively. The quick 195 depletion of caffeine in the solution, compared to the unplanted controls (< 5% of loss after 7 196 days), suggests that caffeine could be taken up by radish and undergoes relatively rapid 197 transformation in the hydroponic system. Six demethylation metabolites from phase I reactions 198 were identified and quantified using their authentic standards including xanthine, 3-199 200 methylxanthine, 7-methylxanthine, theophylline, paraxanthine and theobromine. All six metabolites were detected in both radish roots and leaves. None of the metabolites was found in 201 202 the radish-free controls indicating that the metabolism is plant-associated. Only a small amount 203 of xanthine was found in the hydroponic solution with radishes at day 3 and 7, and no other metabolites were found in the solution (Table S8). The presence of xanthine in the solution could 204 205 be attributed to the excretion from plants and/or caffeine transformation by root exudates.

206 The molar fractions of the six metabolites increased with exposure time in both roots and leaves (Figure 1). In the radish roots, the parent compound caffeine gradually decreased from 207 67.1% at day 1 to 49.8% at day 3, and to 15.6% at day 7, indicating that caffeine was 208 substantially metabolized. The product xanthine (loss of three -CH₃ groups from caffeine) was 209 the most abundant metabolite with the molar fraction increasing from 16.7% at day 1 to 58.5% at 210 211 day 7. Theobromine (loss of one $-CH_3$ group) was the major intermediate with the fraction increasing from 9.0% at day 1 to 13.2% at day 7. This time-dependent transformation pattern 212 213 was also observed for 3- and 7-methylxanthine with their molar fractions up to 5.3% at the end 214 of experiment. Theophylline and paraxanthine (loss of one -CH₃ group) were the least abundant metabolites with the molar fractions < 3.8% (day 3); their fractions increased between day 1 and 215 3 and then decreased from day 3 to 7. Compared with the metabolism occurring in radish roots, 216 caffeine experienced less transformation in the leaves (Figure 1). The parent compound caffeine 217 was predominant with the molar fraction of 94.6% at day 1, 77.5% at day 3, and 68.1% at day 7. 218 219 Xanthine was the most abundant metabolite at day 1 (2.2%) and day 3 (9.6%), and theobromine was most abundant at day 7 (9.9%). Given that these phase I reaction products all contained the 220 moiety of xanthine that maintains the bioactivity of the parent compound (Goth and Cleaver, 221 222 1976), the extensive biotransformation of caffeine in radish, particularly in the edible root fraction, could lead to an underestimate of the risk associated with food safety if only 223 224 considering the presence of the parent compound.

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226 *3.2. Transformation of caffeine in radish enzyme extracts*

The results of the *in vivo* hydroponic experiment provided the information on the metabolism of caffeine in the intact radish. However, the uptake and translocation of caffeine in

229 radish, as well as radish growth, could add more confounding impacts to elucidating the metabolism process. The in vitro exposure experiments with radish enzyme extracts offer an 230 alternative to investigate the metabolism of pharmaceuticals in plants. The amount of caffeine 231 and its metabolites formed in the radish tissue enzyme extracts as a function of time is shown in 232 Figure 2. During 96-h exposure, the loss of caffeine in root enzyme extracts was 29.9%, which 233 234 was approximately 2 times that in leaf enzyme extracts (15.4%) (Figure 2). Meanwhile, the total amount of metabolites formed by demethylation reactions in root enzyme extracts was 235 236 approximately 2 times that in leaf enzyme extracts, suggesting that a larger extent of 237 biotransformation occurred in radish roots. This is consistent with the results of the in vivo hydroponic experiment in which the larger fraction of metabolites was formed in radish roots 238 than in leaves. The amount of the six demethylation metabolites continued to increase over time 239 in both root and leaf enzyme extracts, except for xanthine in the leaf enzyme extracts which 240 decreased slightly from 72 h to 96 h. The amount of metabolites formed in both tissue enzyme 241 242 extracts at 96 h of exposure ranked in the order of xanthine > theobromine > 3-methylxanthine \approx 7-methylxanthine > theophylline \approx paraxanthine (Figure 2), which is also consistent with the 243 ranking of demethylation metabolites found in the in vivo intact radish. In the in vivo hydroponic 244 experiment, caffeine accumulation in roots and leaves varied with time due to the continuous 245 246 water movement into radish; therefore, it is not feasible to compare the absolute amount of the formed metabolites between the *in vitro* and *in vivo* experiments. However, the consistence in the 247 248 types of metabolites and their rankings in magnitude in both radish roots and leaves suggests that the *in vitro* exposure to plant tissue enzyme extracts could provide a convenient alternative to 249 evaluate the metabolism of pharmaceuticals in a specific plant compartment (Card et al., 2013; 250 251 Cui and Schröder, 2016).

252 In both *in vivo* and *in vitro* studies, phase I demethylation reaction is identified as a metabolism pathway for caffeine in radish (Figure 3). The possible transformation route follows 253 stepwise N-demethylation reactions via the loss of one -CH₃ group to form paraxanthine, 254 theobromine and theophylline, loss of the second –CH₃ group to form 7- and 3-methylxanthine, 255 and loss of all three $-CH_3$ groups to form the end product xanthine. The initial N-demethylation 256 257 reaction (loss of one $-CH_3$ group) is common in mammalians and insects, which is believed to be mediated by cytochrome P450 enzyme systems (i.e. CYP1A2 isoenzyme) (Berthou et al., 1991; 258 259 Berthou et al., 1992; Coelho et al., 2015; Kot and Daniel, 2008). The N-demethylation reaction 260 of antidiabetic agent metformin was also observed in Typha latifolia (Cui and Schröder, 2016). P450 enzymes are also commonly present in plants, and catalyze many metabolic reactions of 261 xenobiotics including some pharmaceuticals and herbicides (Fonne-Pfister et al., 1988; Huber et 262 al., 2009; Siminszky, 2006; Thies et al., 1996). Caffeine is believed to undergo similar enzyme-263 mediated transformation in radish. In mammalians paraxanthine represents the large fraction of 264 265 the primary metabolites (70-80%) during the first step of demethylation (Berthou et al., 1991; Kalow and Tang, 1993; Kot and Daniel, 2008). However, in this study theobromine was the most 266 abundant intermediate in radish. 267

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269 *3.3. Mass distribution of pharmaceuticals in hydroponic system*

Pharmaceuticals have been frequently detected in vegetables grown in fields amended with biosolids or irrigated with reclaimed water (Christou et al., 2017; Malchi et al., 2014; Wu et al., 2014); however, their metabolism in plants still remains largely unknown. To obtain more basic information on the metabolism of pharmaceuticals in vegetables, we further assessed metabolism potential of fifteen pharmaceuticals in radish on the basis of mass balance using the hydroponic experiment. The mass recoveries of pharmaceuticals in the radish-free controls ranged from 92.0 to 108.2% indicating that pharmaceuticals in nutrient solution were relatively stable, with the only exception of oxytetracycline (14.5%) (Table S9). The substantial loss of oxytetracycline could be due to the rapid hydrolysis in the nutrient solution at pH 6.5 (near neutral), which is the most favorable condition for the hydrolysis of oxytetracycline (Xuan et al., 2009).

The mass distribution of the fifteen pharmaceuticals in solution, roots and leaves after 7-d 281 exposure is presented in Figure 4. The mass recoveries varied between 3.0 and 78.2% of the 282 initial pharmaceutical input, with the relatively high recoveries for tylosin (78.2%), monensin 283 lincomycin (74.3%), carbamazepine (68.4%), 284 (76.4%), trimethoprim (58.6%) and oxytetracycline (52.3%). Among these six pharmaceuticals, the majority of trimethoprim 285 (36.7%), lincomycin (69.4%), monensin (54.5%), and tylosin (43.8%) remained in the nutrient 286 solution, whereas large fractions of oxytetracycline (27.1%) and carbamazepine (40.8%) were 287 288 accumulated in radish plants. The metabolism of carbamazepine in vegetables such as sweet potato, carrot, tomato, and cucumber could produce 10,11-epoxycarbamazepine and 10,11-289 dihydroxycarbamazepine (Goldstein et al., 2014; Malchi et al., 2014). Riemenschneider et al. 290 291 (2017) found 21 phase-I and II transformation products derived from carbamazepine in tomato after 35 days of exposure, and accounted for $\sim 45\%$ of carbamazepine uptake into plant. The 292 293 recoveries were relatively low for other nine pharmaceuticals ranging from 3.0 to 32.1%, only 294 3.0-4.0% for 17 β-estradiol, sulfamethoxazole, sulfadiazine and estrone, 11.7-18.4% for triclosan, acetaminophen and caffeine, 20.7 and 32.1% for carbadox and lamotrigine. Similar metabolism 295 296 potential of triclosan was also observed in carrot by Macherius et al. (2012), who identified 8 297 triclosan conjugation products and their total amounts was about 5 times more than the parent

compound triclosan. The total recovered mass was < 80% for all tested pharmaceuticals after 7 days of exposure, and the metabolism in radish is expected to be responsible for such discrepancies between the initial input and the remaining amount. The potential risk originating from the metabolites are still unclear, particularly for those quickly and intensively metabolized pharmaceuticals such as estrogens and sulfonamide antibiotics.

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4 *3.4. Metabolism of pharmaceuticals in radish enzyme extracts*

The metabolism potential of the fifteen pharmaceuticals in radish root and leaf enzyme 305 extracts was examined by measuring their dissipation during 96 h of exposure (Figure 5). In root 306 enzyme extracts the tested pharmaceuticals demonstrated a wide range of variation in mass 307 recoveries from 1.8 to 98.1% of the initial dosage (Figure 5). Lincomycin, monensin, tylosin, 308 carbadox, carbamazepine, lamotrigine, trimethoprim and caffeine showed relatively low to 309 minimal transformation with the mass recoveries ranging between 72.3 and 98.1%. 310 311 Carbamazepine and trimethoprim demonstrated high stability in an *in vitro* study, where their concentration remained nearly unchanged in carrot cell cultures during the 90 h of exposure, and 312 < 5% of carbamazepine was transformed after 22 days of incubation (Wu et al., 2016). In another 313 314 lately in vitro study, about 5% of carbamazepine was transformed in horseradish hairy root cell cultures after 6 days of exposure (Sauvêtre et al., 2018). Sulfamethoxazole, sulfadiazine, 315 acetaminophen, oxytetracycline, triclosan, estrone and 17 ß-estradiol manifested a rapid 316 317 metabolism with the recovered amount of < 34.0% of the initially spiked dosage. The fast 318 metabolism of estrogens, acetaminophen, sulfamethoxazole and triclosan is consistent with 319 previous in vitro studies (Card et al., 2013; Huber et al., 2009; Macherius et al., 2012; Wu et al., 320 2016). Three conjugates (acetaminophen–glucoside, acetaminophen-glutathione, and cysteine

321 conjugate) of acetaminophen in the cell cultures of A. rusticana were identified after 6 h of exposure, and their total amounts was about 4.6 times that of acetaminophen (Huber et al., 2009). 322 Wu et al. (2016) found that about 55.5 and 91.7% of triclosan and sulfamethoxazole disappeared 323 in carrot cell cultures within 90 h. Macherius et al. (2012) reported that ~95% of triclosan was 324 quickly metabolized within 24 h in carrot cell cultures. In this study, the rapidly depleted 325 326 pharmaceuticals were found for sulfamethoxazole, sulfadiazine, acetaminophen, oxytetracycline, triclosan, estrone and 17 β -estradiol in both intact radish and root enzyme extracts. However, the 327 328 highly metabolized carbadox and caffeine in intact radish showed less depletion in the enzyme 329 extracts.

The extent of metabolism of pharmaceuticals in leaf enzyme extracts was less than that in 330 root extracts with the exception of carbadox and lamotrigine (Figure 5). Among the 331 pharmaceuticals studied, triclosan demonstrated the highest transformation in leaf enzyme 332 extracts (50.8%); however, the remaining fraction was still \sim 5 times greater than that in root 333 enzyme extracts (9.5%). Sulfadiazine, carbadox and sulfamethoxazole were moderately 334 metabolized in leaf enzyme extracts with the remaining fractions of 59.4%, 61.4% and 72.1%, 335 respectively. For the remaining ten pharmaceuticals, the majority of the initially spiked 336 337 pharmaceuticals still existed in the leaf enzyme extracts (79.2-98.8%) (Figure 5). The different metabolism rates of pharmaceuticals between root vs leaf enzyme extracts could be related to the 338 339 types and amount of extracted enzymes, and their metabolic activities in different radish tissues. 340 The changes in metabolism magnitude could alter the distribution and accumulation of pharmaceuticals in different plant parts, especially in the scenario that pharmaceuticals enter 341 342 plants by root uptake. For instance, only 1.8% of 17 β -estradiol remained in root enzyme extracts 343 after 96-h exposure, while > 90% of 17 β -estradiol was still present in leaf enzyme extracts

(Figure 5). In the *in vivo* experiment, 17 β-estradiol concentration in the leaves was 16 ng g⁻¹ which was >10 times greater than that in roots (1.4 ng g⁻¹). The results from the *in vitro* experiments clearly support the rapid metabolism of 17 β-estradiol in radish roots, which is believed to lead to the corresponding lower concentration in roots in the *in vivo* experiments. Pharmaceuticals in different plant parts could have varying metabolism rates such as carbamazepine in sweet potato, carrot, and tomato (Malchi et al., 2014; Riemenschneider et al., 2017) and iopromide in cattail (Cui et al., 2017).

The rapid metabolism of sulfamethoxazole, sulfadiazine, acetaminophen, oxytetracycline, 351 352 triclosan, estrone and 17 β-estradiol in radish root enzyme extracts was fit to the first-order kinetic model, and the fittings are shown as curves in Figure 6 with $R^2 = 0.94-0.97$ (Table S10). 353 The corresponding half-life was estimated as 17 β -estradiol (15.4 h) < estrone (17.3 h) < 354 triclosan (30.1 h) < acetaminophen (33.0 h) < oxytetracycline (36.5 h) < sulfadiazine (49.5 h) < 355 sulfamethoxazole (63.0 h). The rapid dissipation of triclosan and sulfamethoxazole were also 356 357 found in the carrot cell cultures with half-life of ~ 0.2 h and ~ 7.7 h, respectively (Wu et al., 2016). Compared with the results of carrot cell cultures, triclosan and sulfamethoxazole both exhibited a 358 much slower dissipation rate in the radish enzyme extracts. Triclosan demonstrated a faster 359 360 dissipation rate than sulfamethoxazole in both in vitro systems. The high removal of these pharmaceuticals in root enzyme extracts is consistent with their low mass recoveries in the *in* 361 362 vivo study (Figure 4), suggesting that these biotransformation reactions in root enzyme extracts 363 could also take place in radish roots in the in vivo experiments. Application of plant tissue enzyme extracts to react with pharmaceuticals could serve as an appropriate approach for rapid 364 365 evaluation of their metabolism potential in plants. This particularly simplifies the evaluation of

plant compartment-dependent metabolism, and could also eliminate the impact of uptake andtranslocation to metabolism process in the intact plant experiments.

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369 4. Conclusion

The deficient knowledge on metabolism of pharmaceuticals in plants limits the accurate 370 371 risk assessment of pharmaceuticals accumulated in vegetables because of the lack of appropriate experimental or technique protocols for identification and quantification of metabolites. In this 372 373 study, the comparison in caffeine metabolism in radish between *in vivo* and *in vitro* exposure 374 experiments demonstrates that the demethylation metabolites identified in radish tissue enzyme extracts (in vitro study) were similar to those formed in the intact radish roots and leaves in the in 375 vivo experiment. Similar metabolism patterns (dissipation of parent compounds) were observed 376 for rapidly metabolized pharmaceuticals including 17 β-estradiol, estrone, acetaminophen, 377 triclosan, oxytetracycline and sulfonamides in both radish tissue enzyme extracts and in the 378 379 intact plant. These results suggest that the metabolic reactions with plant tissue enzyme extracts could be used as an appropriate approach for rapid examination of metabolism potential of 380 pharmaceuticals in a specific plant tissue, with the minimal impact from uptake and translocation. 381 382 For those extensively metabolized pharmaceuticals such as 17 β -estradiol, acetaminophen, triclosan and sulfonamides, the measurement of the parent compounds in plants is not sufficient 383 384 to assess their accumulation and potential risks since the formed metabolites might still carry the 385 bioactive moieties and pose similar adverse impacts to plant and human health. Further investigation of the metabolism of these pharmaceuticals in plants and their related 386 387 bioactivity/toxicity are needed to warrant the safe use of reclaimed water for irrigation and land 388 application of biosolids and animal manures.

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395 Appendix A. Supporting Information

396 Additional details are provide in the Supporting Information including physicochemical

397 properties of pharmaceuticals, analytical methods, mass recovery of pharmaceuticals in the

398 hydroponic controls, protein concentration in enzyme extracts, mass distribution of caffeine in

399 hydroponic-radish system, dissipation curves of pharmaceuticals in radish enzyme extracts, and

400 representative chromatograms of pharmaceuticals.

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