

1 **Reclamation of real urban wastewater using solar Advanced Oxidation**
2 **Processes – an assessment of microbial pathogens and 74 organic**
3 **microcontaminants uptake in lettuce and radish.**

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25 **Abstract**

26 In this study, disinfection of urban wastewater (UWW) with two solar processes (H_2O_2 -
 27 20 mg/L, and photo-Fenton 10 mg/L- Fe^{2+} /20 mg/L- H_2O_2 at natural water pH) at pilot scale
 28 using a 60L-Compound Parabolic Collector reactor for irrigation of raw-eaten vegetables
 29 (lettuce and radish) has been investigated. Several microbial targets (total coliforms,
 30 *Escherichia coli*, *Salmonella spp.*, and *Enterococcus spp.*) naturally occurring in UWW and
 31 74 organic microcontaminants (OMCs) were monitored. Disinfection results showed no
 32 significant differences between both processes, showing the following inactivation resistance
 33 order: *Salmonella spp.* < *E. coli* < total coliforms < *Enterococcus spp.* Reductions of target
 34 microorganisms to concentrations below the limit of detection (LOD) was achieved in all
 35 cases with cumulative solar UV (Q_{UV}) ranged from 12 to 40 kJ/L (90 min to 5 hours). Solar
 36 photo-Fenton showed a reduction of 66% of OMCs and solar/ H_2O_2 of 56% in 4 hours
 37 treatment. Irrigation of radish and lettuce with solar treated effluents, secondary effluents and
 38 mineral water was performed for 6 and 16 weeks, respectively. The presence of bacteria was
 39 monitored in surfaces and uptake of leaves, fruit and also in soil. The bacterial concentrations
 40 detected were below the LOD in the 81.2% (lettuce) and the 87.5% (radish) of the total
 41 number of samples evaluated. Moreover, uptake of OMCs was reduced above 70% in crops
 42 irrigated with solar treated effluents in comparison with secondary effluents of UWW.

43

44 **Keywords:** Bacterial inactivation, solar photo-Fenton, uUrban wastewater, plant uptake,
 45 organic microcontaminants.

46

47 **1. Introduction**

48 The increase of water scarcity in arid zones has forced to search alternatives water sources
49 like wastewater for use in sectors like agriculture, the largest water consumer human activity,
50 turning this activity into a widespread practice [1]. The employment of treated wastewater in
51 agriculture involves important health risks, especially for raw consumption crops, due to the
52 presence of several chemical (micropollutants) and microbiological hazardous contaminants
53 [2-3]. Consequently, different agencies have established guidelines to control the microbial
54 load for agricultural irrigation meanwhile organic microcontaminants (OMCs) have not been
55 included in these regulations. OMCs refer to chemical organic substances, which have been
56 identified on water environments in the range of ng/L- μ g/L. They belong to different
57 chemical families with diverse physic-chemical characteristics and include priority
58 substances, already regulated in the EU (Directive, 2013/39/EC), such as polycyclic aromatic
59 hydrocarbons, polychlorinated biphenyls, etc. as well as contaminants of emerging concern
60 (CECs), which are unregulated, poorly characterized in terms of occurrence and have the
61 potential to cause adverse ecological or human health impacts. These include personal care
62 products, pharmaceuticals, drugs, UV-filters, transformation products (TP), etc., which in
63 many cases are still unknown [4].

64 The aforementioned guidelines are based on the control of the concentration of *E. coli* as
65 main indicator of faecal contamination. They include defined different contamination
66 categories depending on the final use of the reclaimed wastewater. For example, <1000
67 CFU/100mL for restricted irrigation according to the World Health Organization [5]; and in
68 other regulations, it is specifically described the *E. coli* concentration for raw crops irrigation
69 for unprocessed purposes, <100 CFU/100mL in Spanish legislation [6] and ISO
70 recommendation [7], and no detectable (< 1 CFU/100mL) by USEPA [8]. Recently, the

71 European Commission published a proposal to regulate wastewater reuse, which include
 72 minimum quality requirements for water reuse in agricultural irrigation [9], which reports a
 73 maximum concentration of <10 CFU/100mL for *E. coli* for the same reuse.

74 The persistence of trace levels of OMCs after a conventional secondary treatment and the
 75 presence of high microbial load (ca. 10^5 CFU/100mL) require the application of a tertiary
 76 treatment to achieve the quality levels required by the reuse guidelines. The continuous
 77 discharge of pharmaceuticals in the environment and their possible incorporation in the
 78 human food-chain-food chain represents one of the greatest threats to human health. This is
 79 also recognised as a clear route to proliferation of antimicrobial resistant microorganisms and
 80 genes in the environment [10]. Conventional tertiary treatments (O_3 , UV-C, chlorination,
 81 etc.) can achieve the microbial quality required for irrigation reuse, although they have some
 82 drawbacks that are still unsolved, as for example the low removal of OMCs, high cost of the
 83 treatments and the generation of undesired disinfection by-products (DBPs) [11].

84 Solar water treatments may represent a sustainable alternative wastewater treatment for reuse
 85 as they allow minimizing the cost of the treatment using sunlight and they also avoid
 86 the generation of DBPs during the process [12]. Solar advanced oxidation processes (AOPs)
 87 have been proven effective alternative tertiary wastewater treatment due to their high
 88 effectiveness for disinfection and decontamination based on the capacity to generate
 89 hydroxyl radicals (HO^\bullet), which are non-selective and very powerful oxidants (2.8 eV).
 90 Several studies on these solar AOPs show high OMCs degradation of OMCs and bacterial
 91 inactivation rates [12].

92 Among the different solar AOPs, photo-Fenton is one of the most investigated processes due
 93 to its simplicity and high disinfection and decontamination capability using low reagents
 94 concentrations (iron and H_2O_2). The process is based on the generation of HO^\bullet through a

95 photocatalytic cycle between iron ions, hydrogen peroxide (H₂O₂) and UV-vis radiation [13].
 96 Furthermore, solar disinfection processes based on photo-inactivation of bacteria assisted or
 97 not by H₂O₂ (Solar/H₂O₂) have also demonstrated high disinfection efficiencies [13-16].
 98 These treatments may have promising applications in highly solar irradiated areas. Previous
 99 investigations have demonstrated the capability of solar/H₂O₂ treatment to improve the
 100 microbial quality of secondary effluents. These works also demonstrated that irrigated lettuce
 101 crops with the solar/H₂O₂ treated wastewater reduced microbial contamination risk [17-18].
 102 Nevertheless, there is still a lack of comprehensive studies about the capability of these
 103 processes as tertiary treatment and on the further translocation and accumulation of OMCs
 104 and their TPs in vegetables, as a result of irrigation practices.
 105 The aim of this study was to evaluate solar/H₂O₂ and solar photo-Fenton process at natural
 106 pH for reducing the load of microorganisms and OMCs of real UWW effluents using a solar
 107 Compound Parabolic Collector (CPC) pilot reactor. Following the treatment, the microbial
 108 and chemical quality of irrigated lettuce and radish crops was assessed in an experimental
 109 greenhouse. Total coliforms (TC), *E. coli*, *Salmonella spp* and *Enterococcus spp* and 74
 110 OMCs present in real UWW were monitored. To our knowledge, this is the first time that the
 111 efficiency of solar/H₂O₂ and solar photo-Fenton at natural water pH is investigated to
 112 simultaneously reduce the microbial risk of natural occurring pathogens and the presence of
 113 OMCs in real effluents of urban wastewater. A multiresidue analytical approach has been
 114 applied to detect OMCs and some of their TPs, in both water and vegetable samples, to
 115 evaluate their potential uptake by raw-eaten crops [17-19].

116

117

118 **2. Materials and methods**

119 **2.1 Water matrixes**

120 Two types of water sources were used: i) secondary effluents from the urban wastewater
 121 (UWW) treatment plant of El Bobar (Almería, Spain). This secondary effluent was directly
 122 used as positive control for irrigation assays and also it was solar treated (as described below)
 123 and used for irrigation assays; and ii) commercial mineral water (Aguas del Marquesado S.L.,
 124 Spain) was used as negative control in the irrigation experiments.

125 Chemical characterization of UWW and mineral water is shown in Table S1 (Supplementary
 126 information). Total Organic Carbon (TOC) was analysed using a Shimadzu TOC-5050
 127 (Shimadzu, Japan). Ionic concentrations was measured with a Dionex DX-600 (Dionex,
 128 USA) IC system for anions, and with a Dionex DX-120 system for cations. Turbidity was
 129 determined using a turbidity-meter (Model 2100 N Hach, USA) and conductivity with a
 130 conductivity sensor (GLP31, CRISON, Spain).

131

132 **2.2. Microbial enumeration and OMCs quantification in water**

133 *2.2.1 Bacterial assessment*

134 Total coliforms, *E. coli*, *Salmonella spp.* and *Enterococcus spp.* naturally occurring in UWW
 135 effluents were detected and enumerated using the pour plate technique by spreading
 136 appropriate sample volumes (50, 250, or 500 μ L, depending on the bacterial load) in selective
 137 and specific agar media (Limit of Detection (LOD): 2 CFU/mL). Total coliforms and *E. coli*
 138 were simultaneously enumerated using the chromogenic ChromoCult® Coliform Agar
 139 (Merck KGaA, Germany) which permits to distinguish between total coliforms and *E. coli*
 140 by the colour of the colonies formed in the agar media, corresponding to red and violet
 141 colonies, respectively. *Salmonella spp.* was enumerated on Salmonella-Shigella agar

142 (Scharlau[®], Spain) and *Enterococcus spp.* on Slanetz Bartley agar (Scharlau[®], Spain).
 143 Colonies were counted after incubation for 24-48 h at 37°C. Concentrations of pathogens
 144 detected in secondary effluent are shown in Table S1 (Supplementary information).

145 2.2.2 Organic microcontaminants

146 74 OMCs (pharmaceuticals and some of their transformation products (TPs)) were screened
 147 in this work due to their frequent identification in UWW effluents [20-21]. The analysis was
 148 performed by a 1200 LC system (Agilent Technologies, Foster City, CA, USA) -with a XDB
 149 C18 50 x 4.6 mm and 1.8 µm particle size analytical column (Agilent Technologies, CA,
 150 USA), coupled to a hybrid quadrupole-linear ion trap-mass spectrometer (QqLIT) 5500
 151 QTRAP[®] from Sciex Instruments (Foster City, CA, USA) equipped with a TurboIon Spray[®]
 152 source featuring electrospray ionization (ESI), operating in positive (ESI+) and negative
 153 (ESI-) modes. The source settings were: ionspray voltage, 4500 V; curtain gas, 25 (arbitrary
 154 units); GS1, 50 psi; GS2, 40 psi; and temperature, 550 °C. N₂ served as nebulizer, curtain and
 155 collision gas. Samples were directly injected (10 µL) and analysed by a multi-residue
 156 analytical method previously reported [21]. The OMCs uptake was analysed in the 3
 157 vegetable matrices by a multi-residue method as reported elsewhere [22]. Briefly, it consists
 158 on the extraction of OMCs based on a modification of QuEChERS acetate sample extraction
 159 protocol followed by LC-MS/MS analysis.

160

161 2.3 Solar water treatments

162 2.3.1. Compound Parabolic Collector (CPC) reactor

163 The solar CPC photoreactor used in this work was described elsewhere [23]. It consists of
 164 two CPC mirror modules with 10 borosilicate-glass tubes per module placed on an anodized-
 165 aluminium platform tilted at 37°. CPC mirrors are made of highly reflective anodized

166 aluminium (MiroSun, Alanod, Germany), with a concentration factor of 1, 4.5 m² of total
 167 irradiated surface, and illuminated water volume is 45L over a total volume of 60L. The flow
 168 rate used was 30 L/min. pH, dissolved oxygen, and temperature were continuously monitored
 169 by several probes inserted on pipes and data were recorded by data acquisition software.

170 2.3.2. Reagents

171 Ferrous sulphate heptahydrate (FeSO₄•7H₂O, PANREAC, Spain) was used as the source of
 172 Fe²⁺ for photo-Fenton experiments. Iron concentration was measured in water samples and
 173 in soil samples from crops irrigation tests. The measurement was done according to ISO6332,
 174 with a limit of quantification (LOQ) of 0.1 mg/L [13]. Hydrogen peroxide aqueous solution
 175 was used at 30% (w/v) (Riedel-de Haën, Germany), and diluted directly into the reaction
 176 mixture. H₂O₂ concentration was measured with a spectrophotometer at 410 nm according to
 177 DIN 38409 H15, limit of quantification (LOQ) 0.1 mg/L [13].

178 2.3.3. Solar radiation measurements

179 Solar UV irradiance was measured using a global UV pyranometer (295–385 nm, Model
 180 CUV4, Kipp&Zonen, Netherlands), providing data of incident radiation in terms of W_{UV}/m²).
 181 The accumulated UV energy in the solar reactor per unit of treated water volume and time
 182 (Q_{UV} , kJ/L) was estimated to evaluate the bacterial inactivation during solar processes.
 183 Treatment time was also used to describe the effectiveness of the solar processes. Q_{UV} allows
 184 the comparison of results under different weather conditions and reactors characteristics and
 185 it is calculated by Eq. (1):

$$186 \quad Q_{uv,n} = Q_{uv,n-1} + \frac{\Delta t_n UV_{G,n} A_r}{V_t}; \quad \Delta t_n = t_n - t_{n-1} \quad \text{Eq (1)}$$

187 where $Q_{UV,n}$ and $Q_{UV,n-1}$ is the cumulative UV energy per liter (kJ/L) at times n and $n-1$; $UV_{G,n}$
 188 is the average incident radiation on the irradiated area (W/m²), Δt_n is the experimental time

189 of samples (s), A_r is the illuminated area of collector (m^2), and V_t is the total volume of water
 190 treated (L) [23].

191 2.3.4. Solar experiments

192 All experiments were conducted at Plataforma Solar de Almeria (Spain) on completely sunny
 193 days. They started at the same local time (10:30 am) and lasted 5 h in consecutive days, so
 194 that water temperature (ranged from 27.1 to 39.2°C) and solar UV irradiance (ranged from
 195 23 to 45 W/m^2) was similar for all experiments. Three solar treatments were investigated: i)
 196 solar photo-inactivation, ii) solar/ H_2O_2 with 20 mg/L of H_2O_2 and iii) solar photo-Fenton at
 197 natural water pH with 10/20 mg/L of Fe^{2+}/H_2O_2 . The reagents concentrations herein used
 198 were selected based on optimal concentrations for the same conditions as reported elsewhere
 199 [13,15,18].

200 Solar experiments were carried out as follows; the photo-reactor tank was filled with 60 L of
 201 UWW effluent. When required, the reagents were added and re-circulated in the dark for
 202 homogenisation during 15 min [23]. After that, the reactor was uncovered and 10-mL
 203 samples were taken at regular intervals during the solar experiment for bacterial, OMCs and
 204 reagents quantification. 2 batches of UWW effluents per solar treatment were sampled and
 205 monitored to obtain the average monitoring results. The average of the bacterial inactivation
 206 results is reported along with an error equal to the standard deviation. Average values of the
 207 OMCs concentrations and their degradation during these treatments are also reported (Table
 208 1).

209

210

211 **2.4. Irrigation assays**

212 *2.4.1 Experimental greenhouse*

213 The irrigation assays were performed under controlled conditions at Plataforma Solar de
214 Almeria using a 30 m²-experimental greenhouse, designed and built by Suministros D.R.
215 (Spain). It consists of 4 individual areas (7.5 m² each) equipped with temperature and
216 humidity sensors connected to Ambitrol[®] software for controlling these parameters by
217 cooling (Fisair, Spain) and heating (Gabarrón, Spain) systems and also automatic windows
218 located in the roof of each area. Averaged temperature during the experiments was 25±5 °C
219 and humidity varied daily from 50 to 90 %.

220 *2.4.2. Crops*

221 Lettuce (*Lactuca sativa*) and radish (*Raphanus sativus*) crops were selected as they are raw-
222 eaten vegetables with relative fast growing, i.e., 8-10 and 4-6 weeks from seeded to
223 harvested, respectively. Both seeds were obtained from a local provider and grown on
224 propylene pots (9x9x10 cm) filled with commercial and regular peat as substrate. According
225 to the manufacturer, it contains a N-P-K ratio (w/v) of 13-14-13 g/L, respectively, pH 7, and
226 120 mS/m of conductivity. 100 pots per each type of crop and irrigation condition were used
227 for statistical purposes.

228 *2.4.3. Crops irrigation experiments*

229 Lettuce and radish irrigation tests were done simultaneously, with similar growing conditions.
230 4 sets of 100 pots of lettuce and radish pots were planted, one per water sample evaluated
231 (negative control, positive control, solar/H₂O₂ treated effluent, and solar photo-Fenton
232 treated effluent). Each set was placed in an individual area of the greenhouse to avoid
233 potential risk of cross contamination between solar treated and secondary effluents. 2 batches
234 of solar treated effluents were collected and stored at 4°C to provide sufficient irrigation

235 water along the irrigation period. Analysis of bacterial regrowth of the selected pathogens in
 236 solar treated effluents during storage was evaluated at 24h, 48h and 72h post-treatment; no
 237 positive regrowth was found in any case. Each pot was regularly watered with 50 mL of
 238 corresponding type of water as reported elsewhere [18, 22].

239 *2.4.4. Detection of pathogens on crops and soil*

240 After the irrigation period, 15 out 100 samples/pot of each irrigation test were randomly
 241 selected and analysed to detect and quantify the pathogens on surfaces of lettuce and radish
 242 leaves, radish fruit and on soil samples following reported methodology [17-18]. Briefly it
 243 consists of the following steps:

- 244 i) Leaves of lettuce or radish: each sample weights (3 ± 0.5) g, it is cut in small ($<1\text{ cm}^2$)
 245 pieces, then mixed with 20 mL of saline solution, and homogenized in a Stomacher
 246 400 (Seward, UK) at 260 rpm for five min.
- 247 ii) Radish fruit: follows the same procedure as leaves, but with samples of (7.0 ± 0.1) g.
- 248 iii) Peat: the soil around each plant (1cm around) was collected and weighted to obtain
 249 samples of (5.0 ± 0.5) g. Then, they were mixed with 45 mL of saline solution in 150
 250 mL container and homogenized manually.

251 The enumeration of pathogens in crops and soils were performed following the pour plate
 252 counting procedure described in section 2.2 and additionally 5 mL of the generated extract
 253 samples was spread in 140mm petri dishes with the corresponding culture media. The LOD
 254 was reduced to 20 CFU/100 mL in this case.

255 *2.4.5 Quantification of OMCs on crops*

256 Composite samples of lettuces and radish (leaves and fruit) were washed with mineral water
 257 prior trituration to remove any deposition in the surface of the crop and detect only true CECs
 258 absorption [23]. After that, the extraction of the composite samples was made per triplicate.

259 Results are shown as the average of the concentrations calculated (concentrations expressed
 260 in wet weight, w.w.).

261

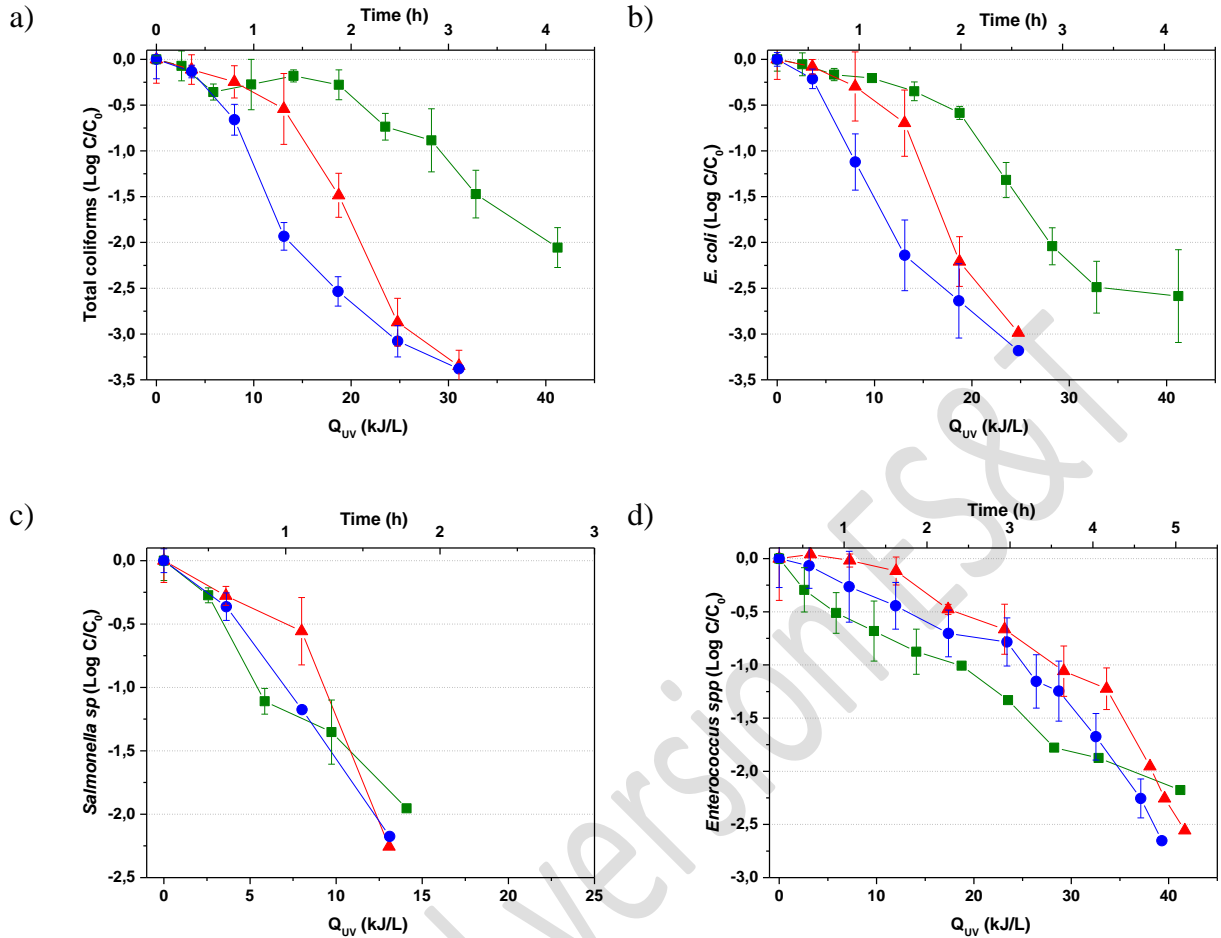
262 **3. Results and discussion**

263 **3.1 Solar treatment of UWW effluents**

264 *3.1.1 Bacterial inactivation*

265 The inactivation profile of total coliforms, *E. coli*, *Salmonella spp* and *Enterococcus spp* in
 266 effluents treated by solar processes at near neutral pH is showed in Figure 1. Water
 267 temperature never exceeded 40°C, excluding therefore the thermal effect (T>45°C) as a key
 268 parameter for bacterial inactivation in these results [24].

269 Reductions of target microorganisms to concentrations below our limit of detection ~~was~~ were
 270 attained in all cases requiring different solar cumulated UV energy and treatment times
 271 depending on the type of pathogen. As expected, the disinfection results using both solar
 272 processes were very similar and much faster than solar photo-inactivation [13-14,18].
 273 Nevertheless, in the case of *Enterococcus spp*, the solar/H₂O₂ treatment was slightly faster
 274 than for the rest of microorganisms. The inactivation order for both solar oxidation processes
 275 was: *Salmonella spp* (12 kJ/L and 90 min) > *E. coli* (23 kJ/L and 2.5 h) > total coliforms
 276 (31 kJ/L and 3 h) > *Enterococcus spp*. (39 kJ/L and 41 kJ/L or 4 for solar/H₂O₂ and photo-
 277 Fenton, respectively).



278 **Figure 1.** Inactivation profile of (a) Total coliforms, (b) *E. coli*, (c) *Salmonella sp* and (d)
 279 *Enterococcus sp.* in UWW effluents by: solar photo-inactivation (-■-), solar/H₂O₂ (20 mg/L)
 280 (-●-) and solar photo-Fenton (10/20 mg/L of Fe²⁺/H₂O₂) (-▲-).

281
 282 These results show the capability of both treatments for reducing the microbial load in urban
 283 UWW effluents. Similar results have been reported [13, 25]. The mechanisms explaining this
 284 behaviour have been already described. In brief, the bactericidal effect of small
 285 concentrations (mM range) of H₂O₂ and sunlight is mainly attributed to the oxidative effect
 286 of internal photo-Fenton reactions assisted by the internal iron and the diffused H₂O₂ inside
 287 the cell, leading to cell lethal damages [13, 25-26]. The results of photo-Fenton at natural
 288 water pH are also in agreement with previous results in UWW, where effects of oxidant

289 concentrations, pH and organic matter as key parameters for bacterial inactivation have been
290 extensively investigated and correlated with the inactivation mechanisms, which are directly
291 related to the generation of HO[•] [13,16].

292 The final H₂O₂ concentration measured was 6.13 mg/L for photo-Fenton and 7.45 mg/L for
293 solar/H₂O₂. In photo-Fenton, H₂O₂ (20 mg/L) was added at the beginning and after 60 min
294 of the experiment to balance the consumed H₂O₂ during the process. This residual
295 concentration (20 mg/L) can be considered innocuous for the crops, as their growth is not
296 compromised, as observed during the experiments. Even, the spontaneous H₂O₂ breakdown
297 in the natural water to water and oxygen will continue happening and decreasing the residual
298 H₂O₂ [18, 26-27]. Moreover, the residual presence of an oxidizing agent in the solar treated
299 effluent could be beneficial to avoiding a possible bacterial regrowth (no regrowth was
300 observed in our study) during the post-treatment water storage [28].

301

302 **3.1.2 Organic microcontaminants removal**

303 The concentration of 74 OMCs (commonly found in UWW) detected in secondary effluents
304 and solar treated effluents by solar/H₂O₂ and photo-Fenton treatment and their removal
305 percentages are shown in Table 1. The results revealed the presence (>LOQ) of 34 OMCs
306 out of the 74 investigated belonging to different classes of pharmaceutical compounds.

307 The OMCs concentration obtained in secondary effluents ranged from 10 to 6897 ng/L with
308 a total load of 14832 ng/L, highlighting the high average concentration of 4-FAA (6541 ng/L),
309 4-AAA (3590 ng/L), atenolol (681 ng/L), hydrochlorothiazide (593 ng/L) and gemfibrozil
310 (573 ng/L). These results are in line with previously reported results [21,22] and confirm the
311 common presence of OMCs, which remain in water after physical and conventional
312 biological treatments due to their high water stability and solubility [29].

313 **Table 1.** Organic microcontaminants (OMCs) detected in secondary effluents and solar
 314 treated effluents. OMCs with a100% degradation are presented in bold.
 315

Compound	Concentration range in secondary effluents (ng/L)	Average in secondary effluents (ng/L)	Degradation (%) solar/H ₂ O ₂	Degradation (%) solar photo- Fenton
4-AA	233-428	304	100	100
4-AAA	3117-4254	3590	54	61
4-FAA	6106-6897	6541	73	75
4-MAA	48-64	54	100	100
Acetaminophen	50	50	-	100
Antipyrine	294-322	308	100	100
Atenolol	626-770	681	0	6
Azithromycin	259-476	363	32	51
Caffeine	55-68	61	37	40
Carbamazepine	90-107	97	11	19
Carbamazepine epox	22-24	23	60	62
Citalopram	187-209	198	2	14
Clarithromycin	16-30	23	0	0
Diazepam	10-15	12	0	0
Famotidine	13-24	18	100	100
Fenofibric acid	26-97	66	78	89
Gemfibrozil	457-852	573	77	88
Hydrochlorothiazide	438-1015	593	83	49
Ketoprofen	40-87	66	100	100
Lincomycin	72-130	96	100	100
Mepivacaine	9-22	16	96	47
Metoclopramide	18-31	26	97	100
Metoprolol	7-17	13	0	3
Metronidazole	20-52	30	100	78
Naproxen	50-62	54	100	-
Pentoxifylline	24-90	51	0	11
Primidone	51-104	84	0	0
Propranolol	30-58	50	95	94
Ranitidine	178-484	340	100	100
Sotalol	20-37	32	100	100
Sulfamethoxazole	51-112	83	71	63
Sulfapyridine	34-76	58	91	87
Trimethoprim	25-64	45	54	47
Venlafaxine	126-287	233	0	4
Total average load (ng/L)		14832	56%	66%

316 LOQ: Limit of quantification. (-): No data.

317 *Metabolites of metamizole: 4-AA (4-Aminoantipyrine); 4-AAA (N-acetyl-4-aminoantipyrine); 4-FAA (N-formyl-4-aminoantipyrine); 4-
 318 MAA (N-methyl-4-aminoantipyrine).
 319

320 Considering the total load of the detected OMCs (14832 ng/L), solar/H₂O₂ and solar photo-
 321 Fenton showed a removal efficiency of 56 % and 66 %, respectively. Looking at specific
 322 OMCs, 10 of them were completely removed using both treatments (final concentration <
 323 LOQ); while 9 and 6 OMCs were degraded above 70% by H₂O₂/solar and solar/photo-Fenton,

324 respectively. In the case of hydrochlorothiazide, metronidazole, mepivacaine,
325 sulfamethoxazole, sulfapyridine and trimethoprim, H₂O₂/solar was more efficient than
326 photo-Fenton, while the opposite happened in the rest of the OMCs.

327 These results can be explained by the mechanisms involved in the OMCs degradation in both
328 processes. In the case of solar/H₂O₂, the partial removal of OMCs is due to the oxidation by
329 H₂O₂ in combination with solar photons. Although the generation of HO[•] under solar light
330 can be discarded, as this process requires photons with wavelengths under 290 nm, which are
331 practically inexistent in the solar spectrum at the Earth's surface [12], the generation of a
332 small amount of HO[•] cannot be completely discarded considering the chemical complexity
333 of the water matrix (secondary effluents) investigated, which may account for the OMCs
334 degradation obtained in this work.

335 Previous investigations in similar conditions confirm similar degradation rates of
336 pharmaceuticals in UWW. Ferro et al. (2015) [18] used solar/H₂O₂ for inactivation of
337 antimicrobial resistant *E. coli* and *E. faecalis*, and the removal of spiked pharmaceuticals
338 (100 µg/L). Carbamazepine, flumequine and thiabendazole were removed 36.9%, 68.3% and
339 99.9% within 5 hours of treatment, respectively. Moreira et al. (2018) [30] reported the
340 removal of spiked pharmaceuticals (100 µg/L) in UWW, 40% carbamazepine, 45%
341 sulfamethoxazole, after 5 hours, and 100% diclofenac in 3 hours of solar exposure. Higher
342 antibiotic concentrations have also removed partially in UWW, as reported by Rizzo et al.,
343 (2018) [31], where for example 71% removal of chloranphenicol was attained with a solar
344 UV dose of 1173 kJ/L. Nevertheless, the initial pharmaceutical concentration (25 000 µg/L)
345 in this work [31] is much higher than the commonly OMCs usually found in UWW samples,
346 from 0.002 to 6.9 µg/L, as shown in our results (Table 1).

347 Conversely, for solar photo-Fenton, the well-known generation of HO^{*} is responsible for a
 348 higher oxidant capacity, although limited degradation was obtained. Recent studies report on
 349 the limited efficiency of ‘mild photo-Fenton’ – i.e. mM concentrations of reagents and near
 350 neutral pH - for the degradation of OMCs (most of cases with spiked pollutants in real WW
 351 samples) [30-31]. They claim as the main reason the formation of iron sludge due to the
 352 precipitation of iron hydroxide at neutral pH. For example, Moreira et al. (2018) report
 353 degradations of only 20% for carbamazepine and sulfamethoxazole, and 100% for diclofenac
 354 [30].

355 The present contribution reports for the first time on the disinfection and degradation of 74
 356 OMCs in real UWW effluents by solar/H₂O₂ and solar photo-Fenton at pilot scale. Both solar
 357 treatments reduce more than 55% the total OMCs load determined in the effluents as well as
 358 the selected microbial pathogens below to the detection limit (2 CFU/mL). The lower
 359 chemicals consumption for solar/H₂O₂ makes this process more suitable for the reuse
 360 application under study. Further research is needed to improve the disinfection results to meet
 361 the irrigation quality criteria-restricted use (< 1/100 CFU/mL).

362

363 **3.2. Microbiological assessment of irrigated crops**

364 The microbiological results obtained for crops irrigated with mineral water (negative control)
 365 showed, as expected, negative results in all analysed samples, i.e., the absence of bacteria in
 366 lettuce, radish, radish leaves and soil samples. Table 2 shows the presence and absence of the
 367 selected pathogens in the crop samples irrigated with secondary effluents-untreated and solar
 368 treated effluents by the two solar processes.

369 The samples irrigated with secondary effluents showed the presence of all the selected
 370 bacteria in 60% of the samples analysed (leaves, fruit and soil), with concentrations ≥ 200

371 CFU/100mL. Nevertheless, the radish leaves showed complete absence of all bacteria, which
372 may be explained by the high hydrophobicity of the leaves surface, reducing therefore the
373 adhesion of bacteria. Although radish leaves are not eaten by human they can be used for
374 animal feed, therefore control of their quality may be also important. *E. coli* was detected in
375 only 33% of the analysed lettuce leaves samples, which may be attributed to the low
376 capability of survival of this bacterial strain far from their environmental conditions (water
377 or humidity and nutrients) [32]. Total absence of *Enterococcus spp.* in soil samples was also
378 observed. This can be explained by the lack of required more complex nutrients for these
379 bacteria strain [33]. Moreover, *Enterococcus spp.* has a lower survival capacity in soils after
380 watering (rainfall) compared with other gram-negative bacteria, i.e. *E. coli* [34]. These results
381 confirmed the high health risk associated with direct reuse of secondary effluents for human
382 consumed crops [2-3]. According to the guidelines for the restricted reuse of wastewater, the
383 presence of *E. coli* is, in all cases, over the permitted concentration for irrigation of
384 unprocessed raw eaten crops (< 100 CFU/100mL [6-7] and < 1 CFU/100mL [8]).

385 The microbiological quality results from the analysis of lettuce leaves, radish fruit and leaves,
386 and soil irrigated with treated effluents by solar/H₂O₂ and solar photo-Fenton show the
387 complete absence of *E. coli*, *Salmonella spp* and *Enterococcus spp* (Table 2). Regarding total
388 coliforms, a substantial reduction in lettuce leaves using both treatments was observed. Only
389 the 20% of the samples showed positive results in the case of lettuce, i.e., 3 out 15 samples,
390 with concentration of 200 CFU/100mL, and none for radish samples, including leaves and
391 fruits. On the other hand, in soil samples the detection of total coliforms showed that, in the
392 case of solar/H₂O₂, 10-11 (radish soil-lettuce soil) out of 15 samples were positive, with a
393 maximum concentration of 900 CFU/100mL; while photo-Fenton showed zero (lettuce soil)
394 or one (radish soil) out of 15. These differences observed in soil samples can be attributed to

395 the presence of a certain amount of iron in the soil due to the consecutive irrigation events.
 396 The iron concentration measured in water/soil samples used during irrigation protocol (using
 397 treated effluents by solar photo-Fenton) revealed a slightly higher concentration (0.46 mg/L)
 398 than in those irrigated by solar/H₂O₂ (< 0.1 mg/L). This iron in soil may react with the
 399 residual H₂O₂ of the solar treated effluent through Fenton and Fenton-like reactions
 400 producing a bactericidal effect. Several articles show that Fenton processes have been used
 401 in the remediation of pesticides contaminated soils including pendimethalin, DDT, diuron,
 402 2,4-dichlorophenol and pentachlorophenol [35]. However, some authors claim that Fenton
 403 applied for soil remediation is very harmful to the microbes in the soil [35].

404
 405 **Table 2.** Detection of pathogens on lettuce and radish crop irrigated with secondary effluents
 406 and solar treated effluents.

Crop sample	UWW sample	Total coliforms*	<i>E. coli</i> *	<i>Salmonella spp</i> *	<i>Enterococcus spp</i> *
Lettuce					
Leaves	secondary effluent Untreated	15/15 (400)	5/15 (200)	15/15 (200)	9/15 (200)
	Treated (H ₂ O ₂ /solar)	3/15 (200)	0/15 (<LOD)	0/15 (<LOD)	0/15 (<LOD)
	Treated (PhotoFenton)	3/15 (200)	0/15 (<LOD)	0/15 (<LOD)	0/15 (<LOD)
Soil	secondary effluent Untreated	15/15 (15500)	15/15 (1200)	15/15 (200)	0/15 (<LOD)
	Treated (H ₂ O ₂ /solar)	11/15 (900)	0/15 (<LOD)	0/15 (<LOD)	0/15 (<LOD)
	Treated (Photo-Fenton)	0/15(<LOD)	0/15 (<LOD)	0/15 (<LOD)	0/15 (<LOD)
Radish					
Leaves	secondary effluent Untreated	0/15 (<LOD)	0/15 (<LOD)	0/15 (<LOD)	0/15 (<LOD)
	Treated (H ₂ O ₂ /solar)	0/15 (<LOD)	0/15 (<LOD)	0/15 (<LOD)	0/15 (<LOD)
	Treated (PhotoFenton)	0/15 (<LOD)	0/15 (<LOD)	0/15 (<LOD)	0/15 (<LOD)
Fruits	secondary effluent Untreated	10/15 (4400)	10/15 (3900)	15/15 (200)	15/15 (23300)
	Treated (H ₂ O ₂ /solar)	1/15 (1800)	0/15 (<LOD)	0/15 (<LOD)	0/15 (<LOD)
	Treated (Photo-Fenton)	0/15 (<LOD)	0/15 (<LOD)	0/15 (<LOD)	0/15 (<LOD)
Soil	secondary effluent Untreated	12/15 (6300)	12/15 (400)	15/15 (200)	0/15 (<LOD)
	Treated (H ₂ O ₂ /solar)	10/15(400)	0/15(<LOD)	0/15 (<LOD)	0/15 (<LOD)
	Treated (Photo-Fenton)	1/15 (100)	0/15 (<LOD)	0/15 (<LOD)	0/15 (<LOD)

407 * Number of positive detected samples / Total samples analyzed, i.e., 15. Numbers in brackets show the maximum concentration in
 408 CFU/mL detected.
 409 (< LOD): below limit of detection: 20 CFU/100mL.
 410

411 These results shows the absence (with exceptions due to the plant substrate surface) of all
 412 pathogens under study when the UWW is disinfected up to the desired level (< 2CFU/mL).
 413 This is coherent with previous findings; Bichai et al (2012) [17] reported the improvement
 414 of the microbiological quality of lettuce irrigated with treated effluents by solar/H₂O₂ with
 415 consideration of natural occurring *E. coli*. Ferro et al. (2015) [18] investigated the cross
 416 contamination of lettuce by antibiotic resistant *E. coli* and *Enterococcus spp* in UWW also
 417 treated by solar/H₂O₂.

418

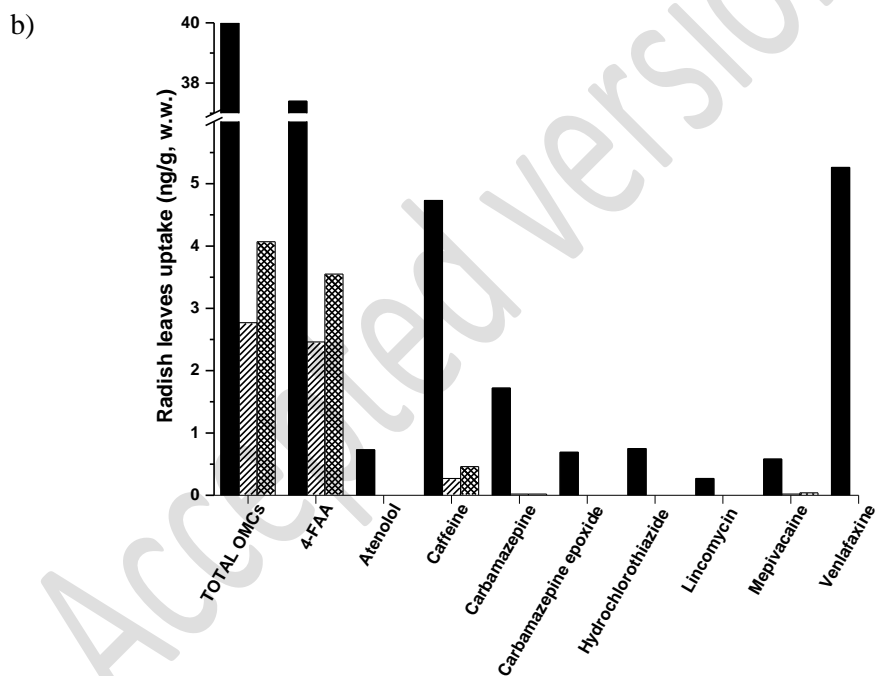
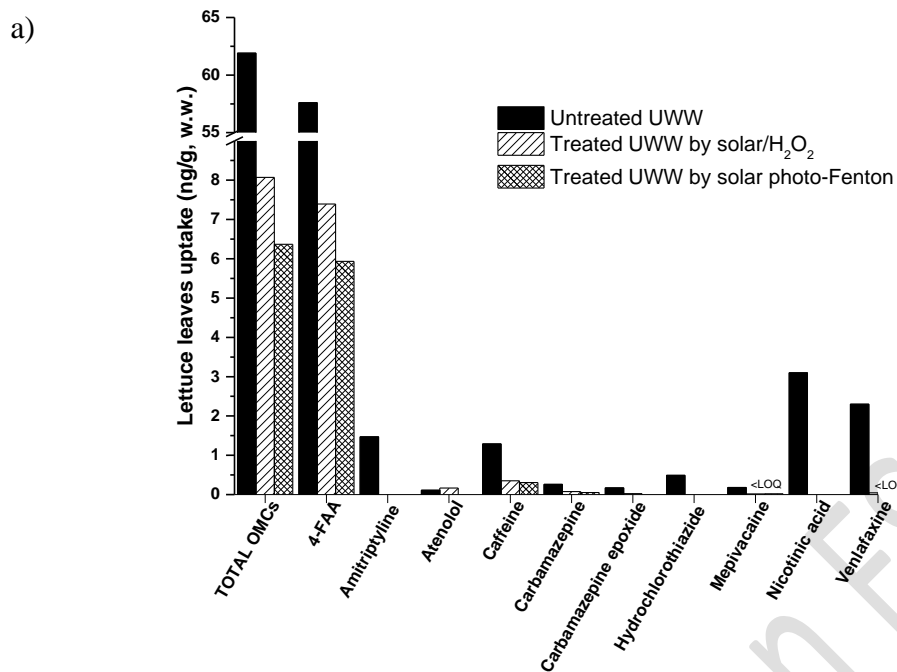
419 **3.3 Crop uptake of OMCs**

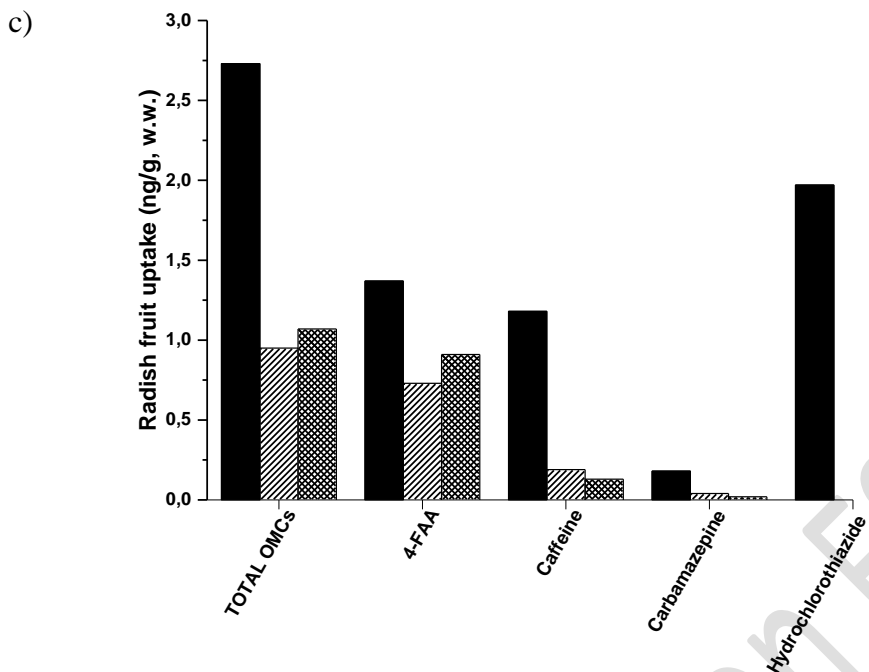
420 Figure 2 shows the OMC concentrations detected in lettuce and radish crops irrigated with
 421 solar treated effluents. The OMCs uptake in the crops irrigated with secondary effluents--
 422 same methodology- have been used for comparison purposes in Figure 2 [22]. The irrigation
 423 with secondary effluents led to the uptake of 12 OMCs in the plant samples assessed (4-AAA,
 424 4-FAA-dipyronone metabolites-, amitriptyline, atenolol, caffeine, carbamazepine,
 425 carbamazepine epoxide -carbamazepine metabolite-, hydrochlorothiazide, lincomycin,
 426 mepivacaine, nicotinic acid and venlafaxine). At harvest, the concentrations ranged from
 427 0.11 ng/g (atenolol in lettuce) to 57.6 ng/g (4-FAA in lettuce) [22].

428 The results obtained using solar treated effluents evidence the capability of the solar
 429 processes under study to reduce the amount of OMCs available for the crops uptake (Fig 2a).

430 The total OMCs uptake in lettuce leaves when irrigated with secondary effluents, 61.9 ng/g,
 431 was reduced to 8.1 ng/g and 6.4 ng/g for solar/H₂O₂ and solar photo-Fenton, respectively.

432





433 **Figure 2.** Concentration (ng/g, w.w.) of target analytes found in a) lettuces, b) radish leave
 434 and c) radish fruit irrigated with secondary effluents and solar treated effluents. Experimental
 435 data of secondary effluents obtained from [22].

436

437 The results obtained for the 10 detected OMCs uptake by lettuce crops irrigated by solar
 438 treated effluents are shown in Figure 2a. Significant level of 4-FAA (57.6 ng/g) in lettuce
 439 samples was strongly reduced to 7.4 ng/g and 5.9 ng/g when solar/H₂O₂ and solar photo-
 440 Fenton treated effluents were used, respectively. Nicotinic acid (3.1 ng/g), amitriptyline (1.5
 441 ng/g), hydrochlorothiazide (0.49 ng/g), venlafaxine (2.3 ng/g), and mepivacaine (0.18 ng/g)
 442 were reduced to levels below the LOQ when treated effluents (via both solar treatments) was
 443 used for irrigation.

444 Regarding the results of radish leaves uptake, 10 OMCs were detected and quantified (Fig.
 445 2b). The OMCs with higher concentrations detected were 4-FAA (37 ng/g), venlafaxine (5.2
 446 ng/g) and caffeine (4.7 ng/g), followed by carbamazepine (1.7 ng/g), carbamazepine epoxide,

447 hydrochlorothiazide and atenolol (0.7 ng/g), mepivacaine (0.6 ng/g) and lincomycin (0.3
 448 ng/g). Irrigation with solar treated effluents made undetectable the levels of most of them,
 449 except for the highly concentrated OMCs, i.e. 4-FAA, which decreased the uptake levels up
 450 to 2.5 (solar/H₂O₂) and 3.5 ng/g (photo-Fenton) and for caffeine below 0.5 ng/g with both
 451 treatments.

452 Figure 2c shows the accumulation of only 4 OMCs (4-FAA, caffeine, carbamazepine and
 453 hydrochlorothiazide) in radish fruit irrigated with UWW. These concentrations are lower
 454 than those found in leafy parts (lettuce leaves or radish leaves). This can be due to the fact
 455 that OMCs are translocated by the transpiration stream at the leafy parts, which normally
 456 presents a greater water flow [19]. The total OMCs uptake in radish was also reduced in a
 457 large percentage by the solar treatments: from 2.7 ng/L (secondary effluents) to 0.9 (65 % of
 458 reduction) and 1.1 (60 % of reduction) for solar/H₂O₂ and solar photo-Fenton, respectively.
 459 However, in radish fruit only one OMC; the diuretic drug hydrochlorothiazide was reduced
 460 under the limit of quantification in both cases.

461 The evaluation of the OMCs intake by the roots and their subsequent translocation to other
 462 plant organs is a difficult task in which many factors are involved. Biotic parameters, such
 463 as physiological state of the plant, surrounded micro-fauna, the crop's genotype, and other
 464 abiotic factors including the typology of soil, the organic matter present, the environmental
 465 stress and even the irrigation method can influence the process [10]. The physic-chemical
 466 properties of the microcontaminants play also an important role in this complex process. The
 467 OMCs root uptake and their translocation to aboveground parts of plants is usually evaluated
 468 taking into account compound lipophilicity ($\log K_{ow}$), pK_a values and electrical charge, which
 469 are fundamental to understand their transport capabilities. Typically, polar OMCs in neutral
 470 species ($-1 < \log K_{ow} < 5$) and cationic analytes in a wide range of plant physiology pH values

471 (~5.5 < pH < ~7.5) are more likely to be uptaken by plant roots and then transported through
 472 the vascular plant system [36]. Nevertheless, anions are more likely to be retained in cell
 473 roots due to diverse interactions such as ion trapping and, therefore, less transported [37].

474 In Table S2, the different lipophilic coefficients (log K_{ow} for neutral compounds, pH-
 475 dependent log K_{ow} , log D_{ow} , for ions), pK_a and molecular charge (soil pore solution pH = 7.5)
 476 of the identified OMCs are listed [PubChem Database (www.pubchem.ncbi.nlm.nih.gov),
 477 38]. Predominantly, moderate to strong bases ($pK_a \geq 7$) in neutral form (4-AAA, 4-FAA,
 478 amitriptyline, atenolol, caffeine, carbamazepine and carbamazepine epoxide) and weak bases
 479 ($pK_a < 6$) in their cationic or partially ionized species (hydrochlorothiazide, lincomycin,
 480 mepivacaine and venlafaxine) were found in leaves and radish roots. Only an acidic analyte
 481 in its neutral form was detected in lettuce leaves (nicotinic acid). Anionic forms of OMCs
 482 were not detected in any plant tissue. These results are in agreement with the literature, where
 483 the higher capability of neutral and cationic molecules to translocate from roots to other plant
 484 organs in comparison to anions is demonstrated [36-37].

485 The values of log K_{ow} for neutral species and log D_{ow} for cations at pH = 7.5, were from -
 486 0.62 to 4.92 (Table S2). This range covers from low to medium lipophilic values, which
 487 represents diverse affinities to lipid tissues and agrees with the reported data [36].

488 Our results revealed that leaves of lettuce and radish showed a higher uptake capacity of
 489 OMCs than radish roots, in agreement with other articles [22,36,39-40]. This behavior has
 490 been attributed to the transport properties of the OMCs by the plant transpiration-derived
 491 mass flow. Therefore the OMCs tend to accumulate at higher concentrations in leafy parts
 492 than in roots [10,40].

493 In summary, both solar water treatments have demonstrated a high purification capability to
 494 both reduce the initial load (OMCs and microbial pathogens) of UWW secondary effluents

495 as well as to reduce the presence of pathogens (> limit of guidelines) and the uptake of OMCs
496 in lettuce and radish (fruit and leaves) crops irrigated with solar treated effluents.

497

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504

505 **Supporting Information.**

506 Table S1. Physico-chemical and microbiological characterization of water matrixes used in
507 this study.

508 Table S2. Physicochemical properties of the OMCs found in real samples: log of the acid dissociation
509 constant (pKa), log of the octanol-water partition coefficient (log Kow) and log of the pH-dependent
510 octanol-water partition coefficient (log Dow) in the soil solution (pH=7.5) and predominant state

511

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