

The data underlying this study are openly available in RiUAL (Repository of the University of Almeria) at DOI [10.1021/acs.jafc.9b01656](https://doi.org/10.1021/acs.jafc.9b01656)

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Organic microcontaminants in tomato crops irrigated with reclaimed water grown under field conditions: occurrence, uptake and health risk assessment

Journal:	<i>Journal of Agricultural and Food Chemistry</i>
Manuscript ID	jf-2019-01656r.R1
Manuscript Type:	Article
Date Submitted by the Author:	n/a
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1 **Organic microcontaminants in tomato crops irrigated with reclaimed**
2 **water grown under field conditions: occurrence, uptake and health**
3 **risk assessment**

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15

16 ABSTRACT

17 In many regions reuse of reclaimed water (RW) is a necessity for irrigation. Presence of
18 organic microcontaminants (OMCs) in RW and their translocation to plants may
19 represent a risk of human exposure. Nevertheless, information available about real field
20 crops is scarce and focused on a limited number of compounds. The novelty of this
21 work relies on the application of a wider-scope analytical approach based on a multi-
22 analyte target analysis (60 compounds) and a suspect screening (1300 compounds). This
23 methodology was applied to real field-grown tomato crops irrigated with RW. The
24 study revealed the presence of 17 OMCs in leaves (0.04 - 32 ng g⁻¹), and 8 in fruits
25 (0.01 - 1.1 ng g⁻¹); 5 of them not reported before in real field samples. A health-risk
26 assessment, based on the toxicological threshold concern (TTC) concept, showed that
27 RW irrigation applied under the conditions given do not pose any threat to humans.

28

29

30 KEY WORDS

31 Organic microcontaminants

32 Plant uptake

33 Reclaimed water reuse

34 Health risk assessment

35 LC-MS target/suspect analysis

36

37 INTRODUCTION

38 The lack of fresh water resources for agriculture in arid and semiarid regions is a
39 worldwide problem that needs to be addressed in the 21st century. Factors such as
40 climate change and increasing population have led to severe droughts in areas where
41 intensive agriculture is the main economic activity. The reuse of reclaimed water (RW)
42 for agriculture irrigation seems to be an excellent approach to deal with water scarcity,¹⁻
43 ⁵ since it not only promotes efficient water usage, but also has other advantages such as
44 reducing the application of fertilizers and avoiding the discharge of waste into natural
45 water bodies, thus contributing towards the preservation of the environment.⁶

46 In Europe, the Mediterranean area is heavily influenced by low and irregular rainfall, a
47 fact that has worsened water shortages leading to a lower water supply for agricultural
48 purposes mainly during peak water demand periods. Nowadays, countries such as
49 Cyprus, France, Greece, Italy, Portugal and Spain, have adopted regulations regarding
50 the reuse of RW for crop irrigation due to the increasing application of this practice.⁷
51 So much so, in Spain, the 10.8% of the RW is reused, being the 71% of it destined to
52 agriculture.⁷ In most cases, the national regulations include specific threshold values for
53 either microbiological (e.g. *E. coli*, intestinal nematodes) and physical-chemical
54 parameters (e.g. total suspended solids, turbidity) for any restricted use,⁸ being more
55 strict for agricultural uses. The European Commission has recently launched a proposal
56 for a regulation on minimum requirements for water reuse, which includes
57 recommendations based on a health and environmental risk management framework for
58 future water reuse legislation.⁹ Again, only microbiological and physical-chemical
59 parameters have been considered. However, in the last decade, the presence of organic
60 microcontaminants (OMCs) in RW, which are not completely removed during the
61 treatments,¹⁰ have been pointed out as a potential risk. It has been demonstrated that

62 intensive use of RW in agriculture leads to their accumulation in agricultural soils^{11,12}
63 and their subsequent uptake by plant roots, in some cases being able to translocate to
64 aerial parts of plants such as leaves and fruits through the vascular plant system.^{2,13-15}
65 However, some knowledge gaps and the lack of reliable data still prevent to make
66 definite conclusions about their risk posed to humans and the environment.

67 Numerous studies have shown translocation of OMCs to edible parts of crops in
68 simulated or controlled conditions.¹³⁻²¹ Nevertheless, little is known about their
69 occurrence and accumulation in real field crops exposed to RW irrigation for long time
70 periods. Recently, Picó et al.¹⁴ have evaluated the accumulation of OMCs in agricultural
71 soils and crops irrigated with treated wastewater, finding up to 6 pharmaceuticals in
72 different crops as cabbage, green beans or eggplants. Also Riemenschneider et al.⁵
73 reported the translocation in real field samples of 12 micropollutants and metabolites to
74 different plant organs such as roots, stems, leaves and fruits of 10 different vegetables
75 irrigated with river water mixed with effluent from a wastewater treatment plant
76 (WWTP). In another study, Wu et al.² monitored the accumulation of 19 OMCs in 8
77 vegetables irrigated with RW showing a detection frequency of 64% at concentrations
78 in the range of 0.01-3.87 ng g⁻¹, dry weight, (d.w.).

79 However, most of the reported studies analyze a low number of compounds or are
80 focused on certain pharmaceutical classes. In order to obtain a comprehensive
81 evaluation of the impact of OMCs in the food chain, it is necessary to apply multi-
82 analyte/class methodologies able to provide qualitative information for a wide range of
83 compounds, given the large number of OMCs reported in RW. Therefore, in addition to
84 wider target methods, non-target screening methodologies based on high resolution
85 mass spectrometry (HRMS) should be applied, leading to the identification of
86 substances outside the limited scope of the target analysis.^{12,14} This approach should

87 contribute towards improving data available regarding the occurrence/accumulation of
88 OMCs in final products intended for human consumption to ensure safe use of RWW in
89 terms of health risk assessment.

90 Finally, the reported accumulation of OMCs in crops is in general low and no risk for
91 public health is expected to be associated to the until now, few known individual
92 compounds in crops grown under real field conditions.^{22,23} However, further work needs
93 to be carried out to assess the risk of not previously evaluated compounds that are
94 present in the edible tissues of plants grown under long-term and continuous exposition
95 to these microcontaminants.²⁴ This data will be also valuable to study the risk associated
96 with mixtures of OMCs in end-products in future works.

97 The goal of this work was to increase the current information about the translocation
98 of OMCs derived from reuse by providing reliable data on their occurrence and fate in
99 real tomato crops (leaves and fruits) after long-term exposure to RW irrigation practices
100 under field conditions. Field-grown tomato plants were cultivated in agricultural soils
101 previously analyzed¹² and irrigated with RW for more than 10 years without soil
102 substitution. With this aim in mind, a combined strategy based on a multi-analyte target
103 analysis (including 60 compounds considered as contaminants of emerging concern)
104 together with a suspect screening methodology (covering a list of 1300 potential
105 contaminants) was applied. A simple and quick QuEChERS-based method was used for
106 sample preparation and liquid chromatography coupled to low and high resolution mass
107 spectrometry, were selected. A health-risk assessment approach was also applied to
108 evaluate human exposure of the RW-derived OMCs in tomato fruits.

109

110 **MATERIALS AND METHODS**

111 **Chemicals and Reagents.** A total of 60 OMCs (mainly pharmaceuticals from a
112 variety of therapeutic classes) (Table S1) were analyzed due to their frequent
113 identification in WWTP effluents.¹⁰ All reference standards (purity > 98%) were
114 acquired from Sigma-Aldrich (Steinheim, Germany). Methanol (MeOH), acetonitrile
115 (ACN), water, formic acid and acetic acid (LC-MS grade) were obtained from Sigma-
116 Aldrich. Ultrapure water for LC-MS/MS analysis was produced using a Milli-Q water
117 purification system from Millipore (Darmstadt, Germany). For the QuEChERS
118 extraction method, anhydrous magnesium sulfate (MgSO₄) and sodium acetate
119 (NaOAc) were purchased from Sigma Aldrich (all purity > 98%). Octadecyl-silyl-
120 modified silica gel (C18) and primary-secondary amine (PSA) were acquired from
121 Supelco (Bellefonte, PA, USA).

122 Stock standard solutions of each compound were prepared at 1000-2000 mg L⁻¹ in
123 MeOH. Multi-compound working solutions were prepared at a concentration of 10 mg
124 L⁻¹ in MeOH by diluting the individual stock solutions. All standard solutions were
125 stored in amber glass vials at -20°C. Matrix matched calibration solutions were daily
126 prepared and used for quantification purposes. Two surrogate standards,
127 carbamazepine-d₁₀ and ¹³C-caffeine, were used to check the extraction efficiency.

128 **Sample Collection.** To study the occurrence and distribution of OMCs in the plant
129 system, three greenhouses were selected (GH1, GH2 and GH3; intensive production;
130 13000–25000 m²), in which two different tomato varieties, ramyle (GH1, GH2) and
131 retinto (GH3) were grown. A fourth greenhouse dedicated to the experimental soilless
132 culture (SP1) of the cherry tomato variety, which was grown in pots filled with perlite
133 substrate, was also included in the study. All greenhouses were located in Almeria
134 province (Spain) and had been irrigated with RW for no less than ten years without soil
135 replacement. The RW was provided by a regeneration plant facility which treats

136 municipal wastewater secondary effluents by filtration (sand and anthracite filters) and
137 chlorination (NaClO). Treated water fulfilled the requirements of water quality
138 according to the Spanish regulation for water reuse.⁸ Drip irrigation was employed in
139 all greenhouses. Four sampling events during the commercial tomato campaign took
140 place from January (full plant growth) to May 2016 (when all tomato plants are
141 removed to prepare the next crop). In each sampling event, tomatoes at a mature stage
142 of growth and leaves of tomato plant samples (500 g in each case) of similar size were
143 taken from different parts of the greenhouse following a W sampling route. The
144 subsamples were chopped and mixed to form a homogeneous composite sample and
145 were kept in the dark at -20°C until their analysis. Three replicates of each sample were
146 extracted for quantification purposes. RW was analyzed coinciding with the first
147 sampling of tomato fruits and leaves.

148 **Sample Extraction.** The extraction of OMCs in tomato fruit and leaves was carried
149 out by a modification of the QuEChERS acetate extraction method previously published
150 by our group.¹³ Briefly, a portion of 10 g of plant material were placed into a 50-mL
151 polypropylene centrifuge tube. After that, 10 mL of 1% acetic acid in ACN and 20 μ L
152 of the extraction quality control solution were added to the sample and the tube was
153 shaken for 5 min and centrifuged at 3500 rpm (2054xg) for 5 min. Following the
154 extraction procedure, a clean-up step was carried out. An aliquot of 5 mL of the upper
155 organic layer was transferred to a 15-mL centrifuge tube containing 750 mg of
156 anhydrous MgSO₄, 125 mg of primary-secondary amine (PSA) and 125 mg of C18.
157 Then the tube was shaken for 30 s in a vortex and centrifuged at 3500 rpm for 5 min.
158 Following this, the extract (4 mL) was transferred to screw-cap vials adding 10 μ L of
159 ACN at 1% of formic acid per milliliter of extract. Prior to injection into the LC-

160 MS/MS system, 100 μL of the extract was evaporated and reconstituted in 100 μL of
161 ACN:H₂O (10:90, v/v).

162 **Liquid Chromatography-Mass Spectrometry.** *LC-MS/MS Target Analysis.* The
163 HPLC system (Agilent Series 1200, Agilent Technologies, Palo Alto, CA, USA)
164 consisted of a binary pump, a degasser and an autosampler. Chromatographic separation
165 was accomplished using a XDB C18 (50 x 4.6 mm, 1.8 μm particle size) column
166 (Agilent Technologies). Mobile phases were 0.1% formic acid in MilliQ water (solvent
167 A) and ACN (solvent B). The gradient used ranged from 10% to 100% of solvent B:
168 initially it was kept at 10% for 1 min, increased from 10% to 50% over 3 min and from
169 50% to 100% over 10 min; kept at 100% for 4 min and finally returned to its initial
170 conditions. The total analysis run time was 18 min. The injection volume was 10 μL and
171 the flow rate was set to 0.4 mL min⁻¹. The column outlet system was connected to a
172 hybrid triple quadrupole-linear ion trap-mass spectrometer 5500 QTRAP® (Sciex
173 Instruments, Foster City, CA, USA) equipped with an ESI source (TurboIon Spray)
174 operating with positive and negative polarities. The ionization settings used were:
175 ionspray voltage, 5000 V; curtain gas, 25 (arbitrary units); GS1, 50 psi, GS2, 40 psi; and
176 a temperature, 500 °C. Nitrogen was used as a nebulizer, curtain and collision gas. The
177 multiple reaction monitoring (MRM) mode was chosen for the analysis of the target
178 compounds. To increase the sensitivity for the acquisition performance, the schedule
179 MRM™ algorithm was applied with a retention time window of 40 s per transition. The
180 optimal mass spectrometric parameters for each compound are summarized in Table S2.
181 Sciex Analyst version 1.6.2 software was applied for data acquisition and processing,
182 and MultiQuant 3.0.1 software for data quantification.

183 *LC-QTOF-MS/MS Suspect screening analysis.* Chromatographic separation was
184 performed using a HPLC (Agilent 1260 Infinity system) equipped with a Poroshell 120

185 EC-C18 (50 x 4.6 mm, 2.7 μm particle size) analytical column (Agilent Technologies).
186 0.1% formic acid in ultrapure water (solvent A) and ACN (solvent B) were used as
187 mobile phases. The injection volume was 20 μL and the flow rate was 0.5 mL min^{-1} .
188 The gradient used ranged from 10% to 100% of solvent B: initially it was kept constant
189 at 10% for 2 min, then increased linearly from 10% to 100% for 9 min and finally it
190 remained constant for 4 min before being returning to initial conditions. The total
191 analysis run time was 22 min. The LC system was coupled to a QTOF mass analyzer
192 Triple TOF 5600+ (Sciex Instruments), with a DuoSprayTM ion source consisting of an
193 electrospray (ESI) interface for sample injection and an atmospheric-pressure chemical
194 ionization interface (APCI) for calibrant delivery. Samples were analyzed in ESI+ and
195 ESI- modes. The ESI source parameters were: ionspray voltage, 4500 V; curtain gas,
196 25 (arbitrary units); GS1, 60 psi; GS2, 60 psi; and temperature, 575°C. Nitrogen served
197 as a nebulizer, curtain and collision gas. The equipment worked via TOF MS survey
198 scan (resolving power of 30000) with an accumulation time of 250 ms followed by four
199 IDA (Information Dependent Acquisition) TOF MS/MS scans with an accumulation
200 time of 100 ms. The IDA feature allows the performance of MS/MS acquisitions
201 simultaneously with the MS acquisition. The m/z range was from 100 to 2000. IDA
202 criteria considered dynamic background subtraction. Collision energy of 30 eV with a \pm
203 15 eV spread was applied for MS/MS fragmentation. Diverse Sciex software (Analyst
204 TF 1.5, PeakViewTM 2.2 and MasterView 1.1) was used to record and process LC-
205 QTOF-MS/MS data. A suspect list containing 1300 OMCs commonly found in WWTP
206 effluents was made before sample processing. The settings considered for a final
207 confirmation of the compounds were: a) a mass accuracy error for the precursor ion < 5
208 ppm; b) an isotope ratio difference < 10%; c) a MS/MS spectra fit \geq 80% when the
209 acquired spectra was compared with the MS/MS spectra of the standard; and d) a

210 difference of ± 0.1 min in the retention time (RT) when it was compared with the
211 standard in matrix.

212 **Method Validation.** Concerning the quantitative method for tomato fruits and leaves,
213 the present methodology was validated assessing trueness (in terms of recoveries),
214 precision (expressed as relative standard deviation, RSD), linearity and limits of
215 quantification (LOQs). For method validation, tomato leaves and fruits not irrigated
216 with RW were used as blank matrices. Triplicate analyses of samples spiked at 0.5 ng g^{-1}
217 were used to calculate the recoveries. Satisfactory mean recovery values were
218 considered in the range 70-120% with an associated precision RSDs $\leq 20\%$. The
219 linearity was studied by matrix-matched standard calibration curves at six concentration
220 levels ranging from 0.01 to 10 ng g^{-1} . Linearity was considered as acceptable when the
221 determination coefficients (R^2) were ≥ 0.990 . The LOQs were set as the lowest
222 acceptable concentration in the matrix-matched calibration curve which yielded the
223 signal-to-noise (S/N) ratio closer to 10 for the quantification transition (SRM 1). The
224 quantification of the analytes present in the samples was carried out by matrix-matched
225 calibration curves of all validated compounds. OMCs quantified in real samples
226 fulfilled the requirements for recoveries, precision and linearity (Table S3).

227 Regarding RW, the sample collected was analyzed per triplicate by direct injection
228 following the methodology reported elsewhere,¹⁰ which was previously validated for the
229 analysis of 115 OMCs in WWTP effluents.

230 **Health-risk Assessment.** The health risk associated with presence of OMCs in tomato
231 fruits was estimated using the threshold of toxicological concern (TTC) approach. This
232 is useful for assessing the risk involved with substances present in food at low
233 concentrations and for which toxicity data is still scarce.²⁵ TTC has previously been
234 applied to the risk assessment of OMCs in crops.^{3,20} In this study, an average body

235 weight of 70 kg for adults and 12 kg for toddlers was considered for the estimation of
236 daily consumption. The TTC values and compounds classification were estimated based
237 on the well-known Cramer classification tree. The Cramer method mainly utilizes
238 chemical structures and evaluates the total human intake to establish priorities for
239 testing.²⁶ This protocol considers a number of factors related to the presence of the
240 chemical component under study or the frequency of ingestion, including: a) different
241 metabolic pathways for either activation or deactivation of the chemicals under study; b)
242 partial presence of a target substance in a variety of standard foods and their
243 metabolites; c) toxicity data for each substance; and d) the level of exposure to humans
244 via oral ingestion. This information is then managed to obtain the TTC value of each
245 compound in terms of $\mu\text{g}\cdot\text{ng kg}^{-1}$ body weight (b.w.) day^{-1} .²⁷ For the OMCs
246 translocation that were not reported before, we considered as minimum tolerated
247 exposure of each OMC as equals to the TCC value given for the parent compound.²⁸⁻³⁰

248 TCC values and compound classification were estimated using ToxTree software
249 (ToxTree v.3.1.0, by JRC Computational Toxicology and Modeling and developed by
250 Ideaconult Ltd, Sofia, Bulgaria). The TTC values for all the compounds under study
251 were determined for the highest CEC concentrations of all greenhouses (SP1, GH1,
252 GH2, GH3) obtained in each sampling event (S1 - S4). Statistical analysis of all the
253 samples and repeated measurements in pairs ($p < 0.05$) were performed using ANOVA
254 analysis.

255

256 **RESULTS AND DISCUSSION**

257 **Method validation results.** The proposed methodology was validated in tomato fruits
258 and leaves of tomato plant for a total of 60 OMCs. The validation results are presented
259 in Table S3. A total of 48 out of 60 compounds (80%) in tomato fruit showed

260 acceptable mean recoveries, within the range 70-120% with a $RSD \leq 20\%$. Lower
261 recoveries, in the 47-69% range, were obtained for 9 compounds. They were also
262 accepted because of the good repeatability obtained, $RSD \leq 20\%$, in all cases. These
263 results are in line with those obtained in a previous work for the same compounds in
264 other vegetable matrices such as lettuce, radish and strawberry.¹³ However, the number
265 of successfully recovered OMCs in leaves was lower than in fruits (31 compounds,
266 52%), probably due to the complexity of this matrix, with a high content of chlorophylls
267 and pigments that may affect OMCs extraction and analysis. 14 additional compounds
268 yielded consistent ($RSD \leq 20\%$) recovery rates within the range 37-66%. However,
269 fenofibrate, gemfibrozil, loratadine, mefenamic acid, nalidixic acid and sulfathiazole
270 gave recovery values lower than 30% and two compounds, sulfapyridine and
271 clotrimazole were not recovered at all.

272 Concerning linearity, all compounds presented R^2 values higher than 0.991. LOQs
273 ranged from 0.01 to 2 ng g⁻¹; showing more than the 50% of the analytes LOQs below
274 0.1 ng g⁻¹ in both commodities. All OMCs quantified in the real samples analyzed
275 fulfilled the requirements for recoveries, precision and linearity.

276 **OMCs in Irrigation Water.** An analysis of the irrigation water was carried out at the
277 beginning of the study to obtain an overview of the potential exposure of the crops to
278 the tested OMCs. As can be observed in Table S4, up to 51 OMCs could be identified at
279 concentration values ranging from 15 to 14424 ng L⁻¹. The metabolites of dipyrone, 4-
280 FAA and 4-AAA (14424 and 5396 ng L⁻¹, respectively), the diuretics
281 hydrochlorothiazide and furosemide (2758 and 1694 ng L⁻¹, respectively), and the beta-
282 blocker atenolol (1279 ng L⁻¹), were detected at the highest concentrations. It was
283 expected that OMC concentrations in RW would vary throughout the study. However,
284 overall these results are in line with previous monitoring studies carried out on urban

285 WWTP effluents from Almeria^{10,13} and can be considered as representative of the
286 type/concentration of compounds usually present in the RW. The presence of 35 of
287 these compounds has also been previously reported by our group in soil and soilless
288 perlite substrate samples taken from the greenhouses monitored, which show average
289 concentrations in the range 0.14 - 99 ng g⁻¹, d.w. (Table S4). Although the presence of
290 OMCs in irrigation water and soils cannot be directly related to their occurrence in plant
291 tissues due to the diverse factors involved in plant uptake, it can be assumed that their
292 availability to be taken up by roots and translocate to edible parts is feasible when RW
293 is used in irrigation.

294 **Occurrence of OMCs in Tomato Plant Leaves.** Greater knowledge about the
295 occurrence of OMCs in vegetables irrigated with RW under field conditions is key to
296 evaluating the quality of crops and determining potential consequences of reusing RW
297 in agriculture irrigation. Moreover, the analysis of non-edible parts of the tomato crop,
298 such as leaves, which may be used as sustenance for livestock feeding, is also important
299 since it could represent another pathway for human exposure to OMCs. In this study,
300 OMCs were monitored in real samples of tomato and tomato plant leaves to evaluate
301 their distribution throughout the plant-fruit system.

302 The average concentration levels of OMCs found in leaf samples during the four
303 sampling events are shown in Table 1. Up to 17 compounds were detected in leaves
304 with average concentrations ranging from 0.04 to 32 ng g⁻¹ wet weight (w.w.). The
305 compounds that eventually reached the higher concentrations were the metabolites of
306 dypirone, 4-AAA and 4-FAA (11 and 32 ng g⁻¹, respectively), the anticonvulsant drug
307 carbamazepine (8.9 ng g⁻¹), its metabolite carbamazepine epoxide (8.1 ng g⁻¹) and the
308 antidepressant venlafaxine (4.0 ng g⁻¹). Regarding the frequency of detection, only 7
309 OMCs were found in all samples, namely caffeine, paraxanthine, carbamazepine,

310 carbamazepine epoxide, hydrochlorothiazide, mepivacaine and venlafaxine; evidencing
311 their higher capability of uptake and translocation within the plant. Nevertheless, their
312 concentrations did not increase during the sampling period; a fact that could
313 demonstrate stable accumulation despite constant irrigation with RW. Another group of
314 OMCs were detected at very low concentrations (<LOQ) and/or showed low frequency
315 of detection. This was the case for acetaminophen, antipyrine, diazepam, propranolol
316 and the antibiotic trimethoprim.

317 In addition to the target analysis, samples were retrospectively analyzed by the
318 acquired LC-QTOF-MS/MS sample information. The strategy allowed the identification
319 of 3 additional OMCs: flecainide, lidocaine and tramadol (Figures S1-S3). These
320 compounds were also found in the irrigation water and soil samples (Table S4).¹²
321 Almost all of them were identified in every sampling event, showing uptake from soil to
322 leaf plant tissues. As the methodology could not be validated for these analytes,
323 estimated concentration values had to be calculated (Table 1).

324 In general, no significant differences considering concentration levels were found
325 between the different tomato plants produced in the greenhouses. This suggests there is
326 no correlation between plant uptake and the tomato plant variety.

327

328 **Table 1. Average OMC concentrations (ng g⁻¹, w.w.) quantified in tomato plant leaves**

Compound	SP1 ^a				GH1 ^b				GH2				GH3			
	S1 ^c	S2	S3	S4	S1	S2	S3	S4	S1	S2	S3	S4	S1	S2	S3	S4
4-AAA	<LOQ ^d	-	0.5	<LOQ	<LOQ	-	-	<LOQ	0.4	-	<LOQ	<LOQ	<LOQ	<LOQ	12	<LOQ
4-FAA	8	n.d. ^e	32	13	n.d.	n.d.	7	5	<LOQ	n.d.	4	3	<LOQ	3	4	2
Acetaminophen	n.d.	n.d.	<LOQ	2	n.d.	n.d.	n.d.	2	n.d.	n.d.	n.d.	3	n.d.	n.d.	n.d.	3
Antipyrine	<LOQ	<LOQ	1	<LOQ	<LOQ	<LOQ	0.7	n.d.	<LOQ	<LOQ	0.6	n.d.	<LOQ	<LOQ	<LOQ	n.d.
Caffeine	0.5	1	0.7	0.5	0.5	1	0.4	<LOQ	0.4	1	0.5	<LOQ	0.5	1	0.5	<LOQ
Carbamazepine	5	5	5	2	3	6	5	2	2	9	3	6	6	4	7	4
Carbamazepine epox	3	3	2	3	3	2	0.7	4	2	2	0.5	8	4	2	1	8
Flecainide^f	n.d.	2	n.d.	0.9	n.d.	2	4	4	2	4	4	4	3	2	3	4
Diazepam	<LOQ	<LOQ	0.06	0.04	<LOQ	<LOQ	<LOQ	0.01	<LOQ	<LOQ	n.d.	<LOQ	<LOQ	n.d.	<LOQ	<LOQ
Hydrochlorothiazide	1	1	1	0.6	1	1	2	1	0.9	1	0.9	0.6	1	0.6	1	0.6
Lidocaine^f	1	2	10	8	3	8	6	11	3	5	6	6	7	6	4	8
Mepivacaine	0.6	0.5	0.5	0.3	0.8	0.7	0.8	0.9	0.6	1	0.6	1	1	0.6	0.3	0.8
Paraxanthine	0.2	0.6	0.4	0.3	0.2	0.6	<LOQ	<LOQ	0.2	0.5	0.3	0.2	0.2	0.3	<LOQ	<LOQ
Propranolol	<LOQ	<LOQ	<LOQ	<LOQ	0.3	<LOQ	<LOQ	<LOQ	<LOQ	<LOQ	<LOQ	<LOQ	0.4	<LOQ	<LOQ	<LOQ
Tramadol^f	1	2	1	4	0.8	2	2	3	0.6	3	2	3	0.2	3	3	3
Trimethoprim	n.d.	n.d.	2	n.d.	n.d.	n.d.	0.7	n.d.	n.d.	n.d.	2	n.d.	n.d.	n.d.	n.d.	n.d.
Venlafaxine	2	1	2	0.7	2	2	2	3	2	4	3	4	4	2	2	4

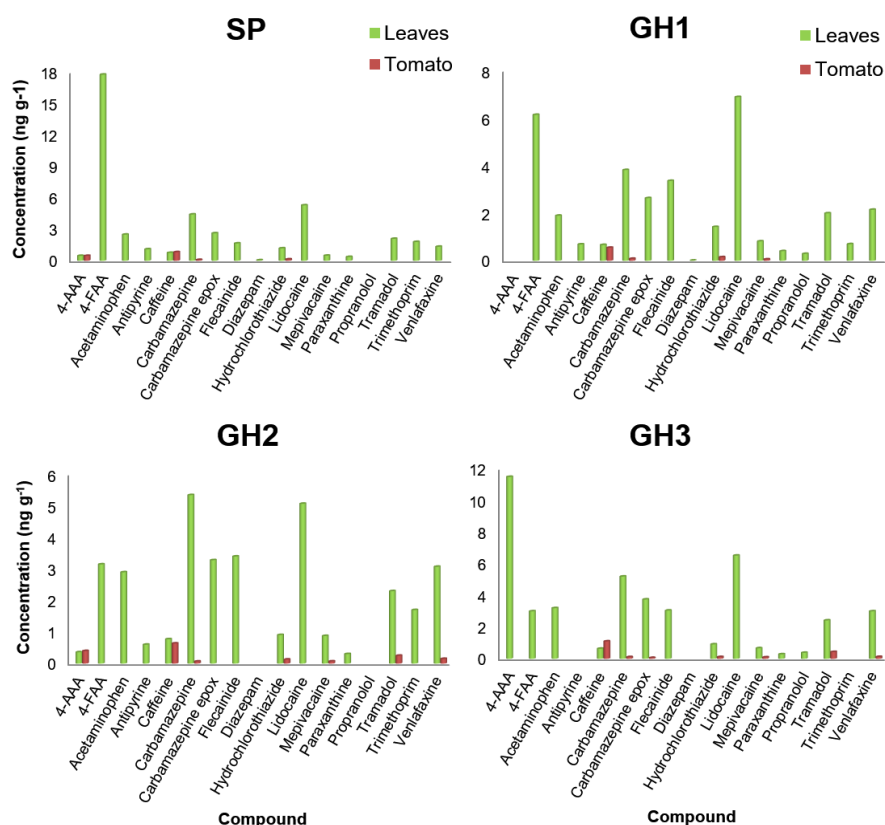
329 ^aSP: soiless perlite culture; ^bGH: greenhouse; ^cS: sampling event; ^d<LOQ: concentration below the limit of quantification; ^en.d.: not detected; ^fEstimated OMC concentrations
330 quantified by LC-QTOF-MS/MS

331

332 Results obtained in the field study concerning translocation of selected OMCs via
333 plant roots to other plant tissues, confirm previous results reported in studies under
334 controlled conditions. For instance, Martínez-Piernas et al.¹³ reported the accumulation
335 of diverse analytes such as 4-AAA, 4-FAA, caffeine, carbamazepine, carbamazepine
336 epoxide, hydrochlorothiazide, lincomycin, mepivacaine and venlafaxine, among others,
337 in lettuce and leaves of radish when RW was used as irrigation water. Wu et al.¹⁶
338 compared the concentrations found for a group of OMCs such as acetaminophen,
339 caffeine, carbamazepine and diazepam in roots and leaves of lettuce, spinach, cucumber
340 and pepper irrigated with spiked water. The metabolism and plant uptake of diazepam
341 has also been evaluated by Carter et al.¹⁷ in radish and silverbeet cultivated with spiked
342 soil. The antibiotic trimethoprim has been reported by Dodgen et al.¹⁸ as being
343 translocated to lettuce, carrot and tomato leaves in an experiment carried out under
344 controlled conditions of temperature and humidity. Other studies have investigated the
345 impact of soil composition in OMCs' plant uptake in leafy crops when they were
346 cultivated with spiked water, observing correlations between soil characteristics and
347 root uptake.^{19,20}

348 However, very few studies have analyzed real field samples exposed to OMCs. Wu et
349 al.² described the translocation of caffeine and carbamazepine within the different plant
350 organs in vegetables irrigated with RW and cultivated under field conditions. In
351 addition, Riemenschneider et al.⁵ observed the accumulation of caffeine,
352 carbamazepine, carbamazepine epoxide and hydrochlorothiazide in different vegetables
353 and agricultural plant tissues. In another study, levels of lincomycin were reported up to
354 20 $\mu\text{g kg}^{-1}$ (d.w.) in leafy vegetables such as rape, celery and coriander grown in soil
355 amended with manure.³¹

356 **Occurrence of OMCs in Tomato Fruit.** Concentrations of OMCs were found to a
 357 lesser extent in tomato fruits, these generally being 10 times lower in fruit compared to
 358 leaves (Figure 1). A total of 12 OMCs were detected in tomato samples. However, only
 359 8 compounds could be quantified in at least one sample (Table 2). In general, the
 360 compounds that showed higher frequencies of detection and concentrations in leaves
 361 were also present in tomatoes, showing mobility through the plant transpiration stream
 362 up to fruits. The highest concentration was observed for caffeine (1.1 ng g⁻¹), followed
 363 by the metabolite 4-AAA (0.4 ng g⁻¹), carbamazepine (0.23 ng g⁻¹), hydrochlorothiazide
 364 (0.15 ng g⁻¹), venlafaxine (0.15 ng g⁻¹), mepivacaine (0.09 ng g⁻¹) and carbamazepine
 365 epoxide (0.07 ng g⁻¹). 4-FAA, acetaminophen, acetanilide and paraxanthine were
 366 identified at concentrations below the LOQ in at least one sample.



367
 368 **Figure 1. Average OMC concentrations found in leaves (green) and tomatoes (red)**
 369 **in each sampling site during the four sampling events.**

370

371 The retrospective analysis of tomato fruit samples revealed the presence of a
372 previously detected analyte in leaves by the same approach: tramadol (Figure S3). It is
373 an opioid analgesic generally used for moderate and severe pain. Tramadol was found in
374 tomatoes from two different greenhouses. Estimated concentrations of this compound
375 are shown in Table 2.

376 No remarkable differences were found between the concentrations observed for cherry
377 (SP1), ramyle (GH1, GH2) or retinto (GH3) tomato varieties. This fact evidences that
378 despite the higher size of the last two types and the different agricultural practices
379 (soilless culture for cherry and real soils for the rest), OMC accumulation was similar in
380 all cases.

381

382 **Table 2. Average OMC concentrations (ng g⁻¹, w.w.) quantified in tomato fruit samples**

Compound	SP1 ^a				GH1 ^b				GH2				GH3			
	S1 ^c	S2	S3	S4	S1	S2	S3	S4	S1	S2	S3	S4	S1	S2	S3	S4
4-AAA	-	<LOQ ^d	-	0.4	-	-	-	<LOQ	<LOQ	-	-	0.4	-	-	-	-
Caffeine	<LOQ	0.4	<LOQ	<LOQ	<LOQ	0.3	0.8	<LOQ	<LOQ	0.8	0.5	<LOQ	<LOQ	1	<LOQ	<LOQ
Carbamazepine	0.2	0.01	0.01	<LOQ	0.2	0.01	0.03	0.1	0.05	<LOQ	0.06	0.1	0.05	0.07	0.1	0.2
Carbamazepine epox	<LOQ	<LOQ	<LOQ	-	<LOQ	<LOQ	<LOQ	<LOQ	<LOQ	<LOQ	<LOQ	<LOQ	0.05	<LOQ	<LOQ	0.07
Hydrochlorothiazide	-	-	0.1	0.1	-	-	<LOQ	0.2	-	<LOQ	<LOQ	0.1	-	0.1	<LOQ	0.1
Mepivacaine	<LOQ	-	-	-	<LOQ	-	<LOQ	0.06	<LOQ	-	<LOQ	0.07	-	<LOQ	<LOQ	0.1
Tramadol^a	-	-	-	-	-	-	-	-	-	0.2	-	-	-	0.1	-	0.7
Venlafaxine	<LOQ	-	<LOQ	-	<LOQ	-	<LOQ	<LOQ	<LOQ	<LOQ	0.1	<LOQ	<LOQ	<LOQ	<LOQ	0.1

383 ^aSP: soiless perlite culture; ^bGH: greenhouse; ^cS: sampling event; ^d<LOQ: concentration below the limit of quantification; ^eestimated OMC concentrations quantified by LC-
384 QTOF-MS/MS

385

386 As was observed in leaves, caffeine and carbamazepine were detected in all
387 greenhouses in every sampling event. Their plant uptake and translocation to the edible
388 parts of vegetables is well described in literature.^{2,13,32} Some studies have already
389 reported them in tomato crops cultivated under field and controlled conditions.^{5,21} Also
390 metabolites such as carbamazepine epoxide have been identified in tomatoes when
391 plants were irrigated with water spiked with carbamazepine under experimental
392 conditions.³³ Hydrochlorothiazide has been reported in other vegetable tissues such as
393 roots and leaves of parsley cultivated under field conditions⁵ evidencing its high
394 capability of translocation through the plant system. OMCs such as 4-AAA, 4-FAA,
395 mepivacaine and venlafaxine, which were quantified in leaves, were also translocated to
396 fruits and identified in certain sampling events. This group of OMCs has been found in
397 the edible parts of lettuce and radish cultivated under controlled conditions submitted to
398 RW irrigation.¹³ Considering that 4-AAA and 4-FAA have exhibited toxicity,³⁴ it is
399 important to monitor their occurrence and to evaluate their repercussions on human
400 exposure.

401 To our knowledge, 4-AAA, mepivacaine, paraxanthine, tramadol and venlafaxine have
402 not previously been identified either in plant tissues or edible parts of real field samples,
403 which highlights the importance of applying wide-scope analytical methods for the
404 evaluation of reuse of RWW in agriculture under different conditions and crops, and the
405 potential of HRMS for the identification of non reported substances in environmental
406 analysis. These results contribute to cover the gap of knowledge regarding the possible
407 OMCs that can be present in edible parts of crops. This will help future studies dealing
408 with the evaluation of the environmental and human risks associated with mixtures of
409 analytes.

410 **Accumulation in Plant Tissues and Properties of Compounds.** It is well-known that
411 OMCs' uptake by roots is accessible for those compounds that are dissolved in the
412 solution of the soil pore water. In general, neutral and cationic species in the soil
413 solution are susceptible to uptake by roots and subsequently translocate to the
414 aboveground parts of plants by the transpiration stream.^{16,20,35} On the other hand, anions
415 are considered less transported to aerial parts due to their accumulation in cell roots by
416 mechanisms such as ion-trapping.³⁵ The translocation of OMCs from roots to other
417 plant organs is possible due to their capability of moving through the transpiration
418 streams. This mobility depends on diverse analyte physical-chemical properties such as
419 lipophilicity (K_{ow}), pK_a or the type of crop, among others.^{3,36}

420 The results found in this study revealed that the OMC concentration values detected in
421 tomato leaves were significantly higher (up to ten times in some cases, Table 1) than
422 those found in tomatoes (Table 2). This behavior has been already reported in several
423 studies.^{2,13,20,36} This issue is explained by the greater water flow to leaves, leading to
424 higher accumulation of OMCs in leafy parts than in fruits.

425 In Table S5, the diverse lipophilic coefficients ($\log K_{ow}$ for neutral compounds and
426 $\log D_{ow}$ for ions), pK_a and molecular charge (soil pore solution $pH = 7.5$) of the OMCs
427 identified in this work are shown. In general, moderate to strong bases ($pK_a \geq 7$), in its
428 cationic species or partially ionized (flecainide, hydrochlorothiazide, lidocaine,
429 mepivacaine, propranolol, trimethoprim and venlafaxine); weak bases ($pK_a < 6$) in
430 neutral form (4-AAA, 4-FAA, antipyrine, caffeine, carbamazepine, carbamazepine
431 epoxide, diazepam and paraxanthine) and a very weak acid ($pK_a > 7.5$) in its neutral
432 form (acetaminophen) were detected. The fact that these analytes are neutral or cations
433 for a wide range of pH values explains their good distribution through the transpiration
434 streams ($\sim 5.5 < pH < \sim 7.5$), being able to cross membranes, reaching leaves and fruits.²⁰

435 Although some compounds were partially ionized, they were translocated via the
436 transpiration-derived mass flow subsequently being found in leaves, and in case of
437 mepivacaine and hydrochlorothiazide, in both leaves and fruits. No OMC in anionic
438 form was detected in either leaves or tomato. This is in accordance with the
439 aforementioned reasons about the expected low translocation of anions through the
440 vascular system, making its distribution less possible through plant streams.

441

442

443 As shown in Table S5, $\log K_{ow}$ and $\log D_{ow}$ of the OMCs identified, ranged from low
444 to medium lipophilic values ($-0.63 < \log K_{ow}, D_{ow} < 3.08$), demonstrating that the
445 OMCs observed have different affinities to lipid tissues. According to Miller et al.³⁶
446 non-ionized compounds with $-1 < \log K_{ow} < 5$ are expected to translocate from roots to
447 other plant tissues, which is consistent with the results obtained for all the neutral
448 compounds identified in this study (Table S5).

449 **Human Exposure and Health-risk Assessment Analysis.** Tomato is one of the most
450 important crops around the world, with global production currently around 130 million
451 tons, of which 88 million is destined for the fresh market and 42 million processed.
452 Considering the intensive consumption of tomato worldwide, the evaluation of human
453 OMC exposure when RW is used as irrigation is of particular interest, even more when
454 this assessment focuses on real samples submitted to long-time RW irrigation.

455 In this study, an assessment of human exposure for each analyte quantified in samples
456 was carried out by the estimation of the daily tomato consumption required to reach
457 TTC levels in adults (average 70 kg) and toddlers (12 kg). All daily intakes were
458 calculated taking into account the worst-case scenario possible. To this aim, a single

459 sampling event with the highest value for TTC estimations was taking into account to
460 provide the most conservative considerations out of this study.

461 Regarding toxicological effects, substances were classified as follows. ‘Class I’ for
462 chemicals with simple structure and known metabolic pathways leading to innocuous
463 end products showing a low order of oral toxicity. Class II contains substances that are
464 intermediate. Very few compounds are included in this category, which is not very well
465 characterized and even questionable.³⁷ They have less innocuous structures than those in
466 Class I but they do not contain potentially toxic structural features. Class III contains
467 substances with complex chemical structures that provide no strong initial presumption
468 of safety and indeed may produce a significant toxicity effect, some of them being
469 genotoxic compounds. Examples of Class III are a number of pharmaceuticals and other
470 common used stimulants including, carbamazepine, caffeine, bezafibrate, clofibrac acid,
471 ketoprofen, naproxen, and metoprolol.²⁰ TTC levels of these pharmaceuticals typically
472 reach values of around $1500 \text{ ng kg}^{-1} \text{ b.w. day}^{-1}$, while the TTC for genotoxic chemicals
473 is only $2.5 \text{ ng kg}^{-1} \text{ b.w. day}^{-1}$ or $0.15 \text{ } \mu\text{g person}^{-1} \text{ day}^{-1}$.^{27,37} Nevertheless, it is important
474 to remark that the consumption of a substance above the estimated TTC level would not
475 imply that there is a toxicological risk. It may even point out a demand for specific
476 toxicity analysis of the compound.

477 Some analytes quantified in tomato samples in this study are classified in Cramer
478 Class III (4-AAA, caffeine, carbamazepine, hydrochlorothiazide and mepivacaine).
479 Regular TTC values for these substances range from 1500 to $1800 \text{ ng kg}^{-1} \text{ b.w. day}^{-1}$.²⁹
480 Venlafaxine and tramadol are categorized as chronic toxic, being their TTC value
481 commonly set in $240 \text{ ng kg}^{-1} \text{ b.w. day}^{-1}$.³⁰ Carbamazepine epoxide has potential
482 genotoxic carcinogenicity. Therefore, TTC reported values are between 1.5 and 2.5 ng
483 $\text{kg}^{-1} \text{ b.w. day}^{-1}$.²⁸

484 As can be observed in Table 3, the OMC concentrations found require an adult and
485 toddler consumption of tens to hundreds of kg to reach the TTC values in most cases.
486 Considering a reasonable tomato daily consumption (according to FAP the average is
487 0.13 kg of tomatoes per adult per day,³⁸ depending on the dietary habits and country),
488 these results do not bring along a health risk for the consumers.

489 As carbamazepine epoxide exhibit genotoxic carcinogenicity, it presented the lowest
490 daily consumption of tomatoes per toddler and adult (400 g and 2.5 kg, respectively) to
491 reach the TTC, despite its low concentration in the samples. These results are in
492 agreement with the low amount intake of carrots to reach the estimated TTC reported by
493 Malchi et al.²⁰

494 The results of this presented study clearly indicate that the estimated TTC values do
495 not pose a health risk for any of the substances at the concentrations found. This
496 contributes towards the safe usage of RW for tomato irrigation under the conditions
497 presented even when the worst conditions were taking into account.

498 Nevertheless, more studies are needed including the evaluation of exposure to other
499 hazards such as synergistic effects due to the addition of concentrations, mixtures of
500 compounds and the formation of metabolites and transformation products that may be
501 more toxic than the original compound. More information about OMCs identified in
502 real crop samples, agricultural procedures and the consideration of sensitive population
503 groups, should also be evaluated to conclude that reuse of RW in agriculture is a safe
504 approach.

505

506

507 **Table 3. Health-risk assessment based on TTC levels of the OMCs quantified in**
 508 **tomato samples.**

Sampling	S1 ^a	S2	S3	S4
Maximum OMC concentration (ng g⁻¹, w.w.) detected in tomato samples				
4-AAA	<LOQ ^b	<LOQ	<LOQ	0.4
Caffeine	<LOQ	1	0.8	<LOQ
Carbamazepine	0.2	0.07	0.1	0.2
Carbamazepine epoxide	0.05	<LOQ	<LOQ	0.07
Hydrochlorothiazide	<LOQ	<LOQ	0.1	0.2
Mepivacaine	<LOQ	<LOQ	<LOQ	0.1
Tramadol	<LOQ	0.2	<LOQ	0.7
Venlafaxine	<LOQ	<LOQ	0.1	0.1
Daily consumption of tomatoes (kg) per adult (70 kg) to reach the TTC values				
4-AAA ^c	-	-	-	315
Caffeine ^c	-	114	150	-
Carbamazepine ^c	548	1800	1260	600
Carbamazepine epoxide ^d	3.5	-	-	2.5
Hydrochlorothiazide ^c	-	-	840	840
Mepivacaine ^c	-	-	-	1400
Tramadol ^d	-	67	-	22
Venlafaxine ^d	-	-	112	140
Daily consumption of tomatoes (kg) per toddler (12 kg) to reach the TTC values				
4-AAA ^c	-	-	-	54
Caffeine ^c	-	20	26	-
Carbamazepine ^c	94	309	216	103
Carbamazepine epoxide ^d	0.6	-	-	0.4
Hydrochlorothiazide ^c	-	-	144	144
Mepivacaine ^c	-	-	-	240
Tramadol ^d	-	67	-	4
Venlafaxine ^d	-	-	19	24

509 ^aS: sampling event; ^b<LOQ: concentration below the limit of quantification; ^ccompound classified
 510 according to Munro *et al.* 1996;²⁹ ^dcompound classified according to Houeto *et al.* 2012.²⁸

511

512

513 ABBREVIATIONS USED

514 4-AAA: 4-acetyl-aminoantipyrine

515 4-FAA: 4-formyl-aminoantipyrine

516 APCI: Atmospheric pressure chemical ionization

517 ESI: Electrospray

518 GH: Greenhouse

519 IDA: Information dependent acquisition

520 LOQ: Limit of quantification

521 OMCs: Organic microcontaminants

522 PSA: Primary-secondary amine

523 RSD: Relative standard deviation

524 RW: Reclaimed water

525 SP: Soilless perlite culture

526 TTC: Toxicological threshold concern

527

528 **ASSOCIATED CONTENT**

529 **Supporting Information**

530 Information about experimental details: list of target analytes, LC-MS/MS details;
531 analytical method validation in tomato fruit and leaves information, OMCs found in
532 RW and agricultural soils, physical-chemical properties of compounds detected in
533 samples and extracted ion chromatograms and MS/MS spectra of the identified
534 compounds by suspect screening strategy (PDF).

535

536 **ACKNOWLEDGMENTS**

537 The authors would like to acknowledge the ALICE project (GA: 734560) “AcceLerate
538 Innovation in urban wastewater management for Cli-mate changeE” supported by the
539 H2020-MSCA-2016. A.B. Martínez-Piernas gratefully acknowledges the Cooperation
540 agreement between the University of Almería and PSA-CIEMAT regarding financial
541 support for her PhD scholarship. P. Plaza-Bolaños acknowledges the University of
542 Almeria for her PhD research contract (Hipatia Program).

543

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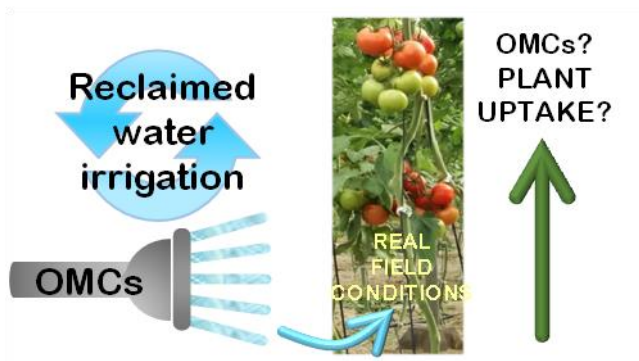
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