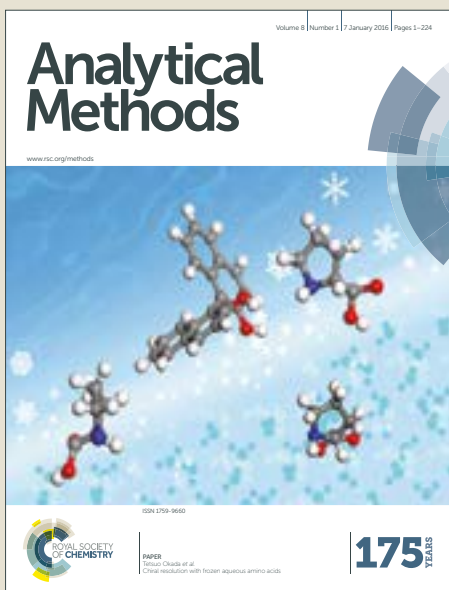


# Analytical Methods

Accepted Manuscript

This article can be cited before page numbers have been issued, to do this please use: P. N. Carvalho, Y. Zhang, T. Lyu, C. Arias, K. Bester and H. Brix, *Anal. Methods*, 2018, DOI: 10.1039/C8AY00393A.



This is an Accepted Manuscript, which has been through the Royal Society of Chemistry peer review process and has been accepted for publication.

Accepted Manuscripts are published online shortly after acceptance, before technical editing, formatting and proof reading. Using this free service, authors can make their results available to the community, in citable form, before we publish the edited article. We will replace this Accepted Manuscript with the edited and formatted Advance Article as soon as it is available.

You can find more information about Accepted Manuscripts in the [author guidelines](#).

Please note that technical editing may introduce minor changes to the text and/or graphics, which may alter content. The journal's standard [Terms & Conditions](#) and the ethical guidelines, outlined in our [author and reviewer resource centre](#), still apply. In no event shall the Royal Society of Chemistry be held responsible for any errors or omissions in this Accepted Manuscript or any consequences arising from the use of any information it contains.

# Methodologies for the analysis of pesticides and pharmaceuticals in sediments and plant tissue

Pedro N. Carvalho<sup>1\*†</sup>, Yang Zhang<sup>1,2‡</sup>, Tao Lyu<sup>1§</sup>, Carlos A. Arias<sup>1</sup>, Kai Bester<sup>3</sup>, Hans Brix<sup>1</sup>

<sup>1</sup>Department of Bioscience, Aarhus University, Ole Worms Allé 1, Building 1135, 8000 Aarhus C, Denmark

<sup>2</sup>College of Life Science, South China Normal University, Guangzhou 510631, PR China

<sup>3</sup>Department of Environmental Science, Aarhus University, Frederiksborgvej 399, 4000 Roskilde, Denmark

\* Corresponding author:

Tel.: Tel.: +45 87158462

E-mail address: [pedro.carvalho@envs.au.dk](mailto:pedro.carvalho@envs.au.dk)

ORCID: 0000-0002-7131-9102

## Abstract

Eco-technologies that utilize natural processes involving wetland vegetation, soil and their associated microbial assemblages are increasingly used for the removal of contaminants of emerging concern (CECs) from polluted water. However, information on removal processes in these systems is not always available, possibly due to the lack of simple and robust methodologies for analysis of CECs in complex matrices such as sediment and plant tissue. The aim of the present study was to use a simple and fast procedure based on ultrasonic extraction (USE) and reduced clean-up procedures to analyse 8 pesticides and 9 pharmaceuticals by high-performance liquid chromatography (HPLC) coupled with diode array detector.

The established methods demonstrated suitable sensitivity and reliability, and proved fit-for-purpose in quantifying multiple classes of pesticides and pharmaceuticals. For sediments, extraction with methanol/acetone (95:5, v/v) followed by a simple evaporation to dryness and redissolution (water:methanol 50:50) provided acceptable recovery (50 - 101%) and RSD < 14%. The complex matrix of plant samples posed specific problems resulting in individualized approaches for pesticides and pharmaceuticals in the final optimized conditions. Pesticides were extracted with *n*-hexane followed by saponification (KOH), pH

<sup>†</sup> Current address: Department of Environmental Science, Aarhus University, Frederiksborgvej 399, 4000 Roskilde, Denmark

<sup>‡</sup> Current address: School of environmental Science and Engineering, Southern University of Science and Technology, Shenzhen 518055, PR China

<sup>§</sup> Current address: School of Animal, Rural and Environmental Sciences, Nottingham Trent University, Nottinghamshire NG25 0QF, UK

1  
2  
3  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60

adjustment and solid-phase extraction; while pharmaceuticals were extracted with methanol:acetone (95:5), supernatant cleaned with activated carbon, evaporated to dryness and redissolved (water:methanol 50:50) prior to HPLC injection. Final method characteristics, with a few exceptions, showed acceptable recovery (> 64%) with RSD < 22% determined using different types of wetland plants.

The methodology has been successfully applied in different studies on the fate of emerging contaminants in water treatment eco-technology systems.

**Keywords:** emerging contaminants; biological sample; environmental matrix; constructed wetlands; water treatment

## 1. Introduction

Emerging contaminants are a new class or classes of unregulated chemicals previously known to be present in the environment but showing new documented environmental impacts [1]. Many of these emerging contaminants are detected in the aquatic environment at low  $\mu\text{g/L}$  to  $\text{ng/L}$  levels, including trace organic pollutants [2], referred to as contaminants of emerging concern (CECs). Examples of CECs are pharmaceuticals, personal care products, plasticizers, surfactants and biocides that are discharged to the environment as a consequence of human activity.

Major sources of the discharge of most of these CECs into the environment are usually the wastewater treatment plants (WWTPs) [3]. Discharge of CEC with unknown potential adverse effects and/or bioaccumulation into the environment may pose a risk to humans considering their uptake either via the food chain or via drinking water [4]. Therefore, there is an increasing interest in the development of more efficient wastewater treatment technologies capable of dealing with CECs [5]. Among these, eco-technologies such as constructed wetland systems (CWs) or phytoremediation engineered systems, that utilize natural processes involving wetland vegetation,

1  
2  
3  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60

66 soil and their associated microbial assemblages to treat polluted water, have  
67 been pursued.

68 Studies along the last decade have shown that these eco-technologies are  
69 able to degrade CECs [6]. However, in spite of promising results [7], detailed  
70 information on the removal processes is lacking. In fact, analysis of  
71 sediment/substrate and plant tissues samples is crucial to be able to perform  
72 flow studies and total mass balances in wastewater treatment systems<sup>[8, 9]</sup>. In  
73 several of the applied studies on CWS, sediments and plant levels have not been  
74 studied, or when studied, the methodology used is not always sufficiently  
75 described. Sediment is already considered a complex matrix with different  
76 organic and inorganic fractions as well as biomass, and humic compounds. Plants  
77 present even greater challenges in terms of matrix interferences due to their  
78 high contents of pigments and fatty or waxy materials [10]. In addition to the  
79 compounds/matrix interactions, the large variety of CECs combined with the  
80 normally very low concentrations of the target compounds pose difficult  
81 challenges to their detection and analysis [11]. There is a clear need for simple but  
82 reliable and robust methodologies concerning CECs analysis in sediment and  
83 plant tissue.

84 The analytical procedures usually comprise three steps, which are  
85 followed by detection and data analysis: i) sampling, ii) compound extraction  
86 and iii) clean-up of the extract that contains the compound [12]. In general, solid  
87 samples will go through a series of steps for preservation (freezing, lyophilizing,  
88 chemical drying) followed by homogenization (blending, chopping, grinding,  
89 milling, etc.). Homogenization with a mortar and pestle is one of most common  
90 procedures for sediment [13]. Considering the analytical procedures for the  
91 determination of CECs in crop plants a recent review by Matamoros, Calderon-  
92 Preciado [14] has covered the major achievements and drawbacks. Several  
93 extraction techniques have been tested for both sediment and plant tissue  
94 samples, including accelerated solvent extraction (ASE) also called pressurized  
95 liquid extraction (PLE), ultrasonic extraction (USE), sea sand disruption method  
96 (SSDM), microwave assisted extraction (MAE), "Quick, Easy, Cheap, Effective,  
97 Rugged, and Safe" method (QuEChERS), and matrix solid-phase dispersion  
98 (MSPD) in combination with pressurized fluid extraction (PFE) [10, 15, 16]. Classical  
99 Soxhlet extractions have been phased out for techniques allowing for higher  
100 throughput such as PLE, USE and QuEChERS. Independently of the extraction

technique used, these primary extracts of multi-residue methods need to be cleaned up before final measurements. In the early days liquid-liquid partitioning (LLP) between an aqueous and organic solvents (such as acetone or dichloromethane), at modulated pH was often performed for pesticide analysis [10, 16, 17], followed by laborious and extensive procedures for condensation, particles removal, gel permeation chromatography (GPC) more commonly referred to as size exclusion chromatography (SEC) and polarity fractionation previous to chromatographic analysis [18]. More recently a typical approach after the extraction of solid samples is the use of solid-phase extraction (SPE), where several different adsorbents can be used and solvents use reduced. SPE and *n*-hexane washing for sample clean-up methods, however, either lack good sensitivity or have considered just a few target analytes [17]. While research on pesticides has historically been more important due to the need for monitoring their levels in food matrices, interest in the analysis of pharmaceuticals in environmental samples has recently risen [14]; therefore very little information on clean-up applications focused on pharmaceuticals analysis is available [19]. The clean-up steps are important to reduce co-extracted compounds that may compromise the chromatographic run avoiding further laborious and/or expensive quantification procedure such as the use of matrix matched [20] or standard addition calibrations and surrogate and internal standards (often isotopically labelled compounds).

In spite of the different available extraction techniques for sediment and plant extracts, recoveries reported are generally variable [14, 21]. On the other hand, several published articles focused on environmental studies, due to different final aims, only briefly report the methodology used without a complete description of optimization and/or validation details. Plant matrices present added difficulties as lipids and pigments which can interfere with analytical procedures are co-extracted with the analytes, resulting in critical ion-enhancement or ion-suppression during MS analysis in HPLC-MS [22]. Therefore, development of simple clean-up steps is important. Simple and fast, but reliable analytical methods are necessary to monitor and control the distribution of CECs in different environmental matrices.

133 In this work a method for the analysis of triclosan and pesticides  
134 (referred further as pesticides group) and pharmaceuticals (Table S1) in  
135 sediment and plant tissue samples was developed. The compounds selected are

1 known to be present in wastewater and comprise different families and chemical  
2 characteristics (molecular weight and  $\log K_{ow}$ ). Ultrasonic extraction (USE) was  
3 selected due to the wide availability of the equipment and its easy operation.  
4 Following extraction, the need for a simple clean-up procedure prior to sample  
5 analysis was evaluated. The compounds were analysed by high-performance  
6 liquid chromatography (HPLC) coupled with a diode array detector (DAD).  
7  
8  
9  
10  
11  
12  
13  
14

## 144 **2. Experimental**

### 145 2.1 Material and Reagents

146 Methanol, acetone and *n*-hexane (SupraSolv ®) and formic acid (98 %,  
147 reagent ACS) were purchased from Merck (Darmstadt, Germany). High purity  
148 grade triclosan (by Dr. Ehrenstorfer GmbH Augsburg, Germany) and the  
149 analytical standards of the pesticides carbendazim, benzoisothiazolinone,  
150 imazalil, terbutryn, diuron, and mecoprop were supplied by Sigma-Aldrich  
151 (Schnelldorf, Germany) and tebuconazole by Dr. Ehrenstorfer GmbH (Augsburg,  
152 Germany). High purity grade analytical standards of the pharmaceuticals  
153 iopamidol, iohexol, iomeprol, iopromide, propranolol and diclofenac were  
154 supplied by Dr. Ehrenstorfer GmbH (Augsburg, Germany) and carbamazepine,  
155 naproxen and ibuprofen by Sigma-Aldrich (Schnelldorf, Germany). Other  
156 solvents and reagents used were analytical grade. Water used in this study was  
157 ultrapure water ( $18.2 \text{ M}\Omega \text{ cm}^{-1}$ , Milli-Q plus system).

158 Individual standard solutions of each pharmaceutical and pesticide (1000  
159  $\text{mg L}^{-1}$ ) were prepared in methanol. A standard working solution of the mixture  
160 of all compounds in methanol, at a concentration of  $60 \text{ mg L}^{-1}$ , was prepared  
161 weekly. This solution was used to prepare daily calibration standard solutions in  
162 Milli-Q water and for the sample (sediment and plant tissue) spiking. All  
163 standard solutions were kept at  $5 \text{ }^{\circ}\text{C}$  in a refrigerator (light protected from  
164 photo-degradation).

165 For decontamination purposes all plastic and glassware used were rinsed  
166 with soap, water, deionized water, soaked overnight in 4.5 % (v/v) hydrochloric  
167 acid (technical -30% purity, VWR BDH Prolabo), rinsed with deionized water  
168 again and dried at  $60 \text{ }^{\circ}\text{C}$ . Procedural blanks were used to control material  
169 cleanliness.

170

1  
2  
3  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60171  
172 **2.2. Sample collection and preparation**173       Samples were selected in order to provide real environmental matrices  
174 for method development and performance check. Sediment (anaerobic, TOC 3%-  
175 7%) and plant tissue samples (*Typha latifolia* and *Berula erecta*) were both  
176 collected in a stormwater pond designed for urban-runoff treatment near  
177 Skoldhoejvej, Aarhus, Denmark.178       Plants were cleaned with deionized water and the plant material divided  
179 into roots and leaves. The sediment and plant tissue were frozen at -4 °C and  
180 subsequently lyophilized (Christ Alpha 1-4 LSC Freeze Dryer, Martin Christ  
181 Gefriertrocknungsanlagen GmbH, Osterode, Germany). Before proceeding to the  
182 extraction, the lyophilized plant material was finely ground (< 2 mm) using a  
183 rotor mill (Retsch KG, Haan, Germany), while the sediment material was  
184 homogenized with mortar and pestle and sieved (particle size < 2 mm).185       Spiked samples were prepared by addition of a methanolic standard  
186 solution mixture of either pesticides or pharmaceuticals (representing an added  
187 volume of 0.5 mL) to the lyophilized and ground samples (0.2 g for plant tissue  
188 and 2 g for sediment) into a glass vial (20 mL) per individual sample for future  
189 extraction. The mixture was shaken and let to dry overnight in the hood, light  
190 protected. The target levels for method optimization and validation ranged  
191 between 0.5 to 5 µg g<sup>-1</sup> dry sediment and 0.5 to 100 µg g<sup>-1</sup> dry plant material of the  
192 individual compounds, as observed before<sup>[15, 23, 24]</sup>. The pesticides and  
193 pharmaceuticals studies were performed in separate batches.194       Method optimization and further characterization was carried out using  
195 spiked samples of both sediment and plant material. Once real sediment and  
196 plant material were used for spiking, non-spiked samples were also analysed to  
197 control background levels. All results further presented along both optimization  
198 and method validation report means and standard deviation of at least 3  
199 replicates.200  
201202 **2.3. Sample extraction**203       Optimization of the sample extraction was performed using ultrasonic  
204 solvent extraction (USE). The first parameter to be tested was the selection of  
205 extraction solvent. For that, six different solvents methanol, *n*-hexane,

1  
206 dichloromethane, methanol:formic acid (96:4, v:v), methanol:acetone (95:5, v:v)  
3 and acetonitrile:formic acid (99:1, v:v) were tested keeping a fixed solvent  
4 volume (10 mL) and a fixed sample mass, 0.2 g for plant material and 2 g for  
5 sediment. Each spiked sample was mixed with the different solvents and further  
6 placed in an ultrasonic bath (Metason 120, Struers, Denmark) for 30 min.  
7  
8

9  
10 After extraction, the samples were centrifuged (3000 rpm for 10 min;  
11 Sigma 3-18K Centrifuge, Laborzentrifugen GmbH, Osterode, Germany) and  
12 supernatants collected. For direct analysis, the supernatants were filtered  
13 through nylon filter (0.45  $\mu$ m) (Frissenette, Knebel, Denmark), while for pre-  
14 concentration the supernatants were evaporated to dryness under a nitrogen  
15 stream at 35°C, further dissolved in 1.0 mL of methanol and filtered through  
16 nylon filters 0.45  $\mu$ m. All extracts analysis was processed by HPLC-DAD (see  
17 section 2.5). Filters were previously tested in terms of blanks as well as sorption,  
18 to ensure that the filtration step would not affect the results.  
19  
20

21 In the optimized operating conditions, for both pesticides and  
22 pharmaceuticals, 2 g of sediment samples were extracted with 10 mL of  
23 methanol/acetone (95:5, v/v) for 30 min in the ultrasonic bath. The resulting  
24 samples were centrifuged and the supernatant evaporated to dryness. Residues  
25 were dissolved in 1 mL of methanol and subsequently the solution was filtered  
26 and injected into the HPLC system. No clean-up procedures were required for the  
27 sediment extracts.  
28  
29

30 Regarding plant material, in the optimized operating conditions, for  
31 pesticides, 0.2 g of plant tissue samples were extracted with 10 mL of *n*-hexane  
32 for 30 min in the ultrasonic bath. For pharmaceuticals, 0.2 g of plant tissue  
33 samples were extracted with 10 mL of methanol/acetone (95:5, v/v) for 30 min  
34 in the ultrasonic bath. Optimization of the clean-up for plant tissue extracts for  
35 pesticides and pharmaceuticals is further discussed in section 2.4.  
36  
37

#### 38 235 2.4. Clean-up procedure

39 Extracts obtained by USE generally require an additional clean-up step,  
40 such as solid-phase extraction (SPE) which is one of the most common  
41 techniques [25]. In the present study a clean-up based on reversed phase  
42 approach using Phenomenex Strata-X SPE columns (200 mg / 6 mL) and a  
43 normal phase approach using a Supelclean™ LC-Florisil® (1 g / 6 mL) were  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60

1  
241 tested. SPE cartridges were processed accordingly to the technical indications  
3 (described in the SI).  
4

5 SPE eluted samples were then evaporated to dryness under a nitrogen  
6 stream at 35°C and the residues dissolved in 1.0 mL of methanol prior to HPLC  
7 injection.  
8

9  
10 Plants pigments, mainly chlorophylls and carotene, are highly  
11 hydrophobic and co-extracted together with the micropollutants. A  
12 saponification step with KOH suggested by Dugay, Herrenknecht [26] to improve  
13 PAHs recovery from plant material was investigated. For that, 5 mL of KOH  
14 solution 1 mol L<sup>-1</sup> (methanol:water (4:1, v/v)) was used to dissolve dried  
15 residues (after extraction solvent evaporation) and the obtained solution further  
16 sonicated for additional 30 min.  
17

18 In the optimized clean-up conditions, plant slurry samples for pesticide  
19 analysis were centrifuged and the supernatant evaporated to dryness.  
20 Afterwards, saponification was performed by dissolving the residues in 5 mL of  
21 KOH solution (methanol:water (4:1, v/v)) and sonicating the sample for 30 min.  
22 Then, samples were filtered, diluted with MilliQ water (MeOH content < 5%),  
23 acidified to pH 5.5 (HCl addition) and further processed through SPE (Strata-X)  
24 prior to HPLC analysis.  
25

26 For pharmaceuticals, in the clean-up step optimized conditions, plant  
27 slurry samples were centrifuged, pellet discarded and the supernatant passed to  
28 a clean vial to which 0.25 g of activated charcoal was added and the solution  
29 sonicated for 30 min. After an additional centrifugation, supernatants were  
30 filtered, evaporated to dryness and the residues were then dissolved in 1.0 mL of  
31 methanol prior to HPLC analysis.  
32  
33

34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60

### 2.5. High performance liquid chromatography conditions

269 Analytes separation was performed using a HPLC Thermo Scientific  
270 Dionex UltiMate 3000 equipment with automatic sampler, column oven and  
271 diode array detector (DAD). The analytes were separated on a Synergy 4 $\mu$  Polar  
272 80 Å column (150 mm  $\times$  2.0 mm ID) using a linear gradient program with two  
273 eluents, water (0.2% formic acid) and methanol (0.2% formic acid). The linear  
274 gradient program used was: 100 % of eluent A (water), keeping isocratic  
275 conditions for 2 min, followed by a 2 min linear gradient to 35 % of eluent A (65

1  
276 % of eluent B (methanol)), followed by a second slower 9 min linear gradient to  
3 0 % of eluent A which was held afterwards for 3 min. Finally, initial conditions  
4 (100 % of eluent A) were reached again in 1 min, with a re-equilibration time of  
5 3 min to restore the column. Flow rate gradient started with  $0.25 \text{ mL min}^{-1}$ ,  
6 maintained for 16 min, followed by a 1 min linear gradient to  $0.3 \text{ mL min}^{-1}$ , which  
7 was held for 1 min and another linear gradient along 1 min back to the initial  
8  $0.25 \text{ mL min}^{-1}$ . The two groups of micropollutants (i.e., a) pesticides plus  
9 triclosan and b) pharmaceuticals were quantified separately using a 6 points  
10 external calibration. The Chromeleon® 7.1 software (Thermo Scientific,  
11 Germany) was used for data integration of chromatograms. The sample injection  
12 volume was set at  $10 \mu\text{L}$ , sampler temperature at  $8 \text{ }^\circ\text{C}$ , column oven at  $20 \text{ }^\circ\text{C}$  and  
13 the detector signal was acquired simultaneously in 3 channels, for quantitation  
14 at 220 nm and 240 nm, and a 3D-field in the  $\lambda$  range 190 to 800 nm (bunch width  
15 of 5 nm). These two wavelengths provide a suitable compromise to obtain  
16 acceptable sensitivity for the detection of all compounds. The instrument (HPLC-  
17 DAD) basic analytical figures of merit (LOD, LOQ, linearity and RSD) are  
18 presented in Table S2.  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60

### 2.6 Analysis of Real Samples

296 The here described optimized and validated methodology has been  
297 efficiently applied by the authors on different works focused on the removal of  
298 micropollutants from water through the use of constructed wetland systems.  
299 Plant samples from an uptake study in spiked hydroponic medium ( $10 \text{ mg L}^{-1}$   
300 level) where both the above and below ground tissues were analysed, as well as  
301 for the quantification of the accumulated amount of micropollutants in the  
302 substrate of constructed wetland bed mesocosms along a 9 months trial. Fully  
303 described experimental setups can be found elsewhere [27, 28].  
304  
305

### 2.7. Statistical analysis

307 Statistically significant differences between samples were evaluated  
308 through Student's t-test ( $p$ -value cut-off: 0.05).  
309  
310

1  
2 311 **3. Results and discussion**  
3 312 3.1 Extraction optimization  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60

The solvents tested were chosen based on typical applications for extraction of solid matrices for a variety of organic contaminants. Ultrasonic extraction (USE) was chosen due to its fast and easy to use approach, besides being attractive because the equipment necessary is widely available and the extraction can be done using a reasonably small amount of sample (0.1 – 2 g) and volume of solvent (5 – 25 mL) [25]. Furthermore, this method has a short extraction time compared to those of classical liquid extraction methods.

320

321

322 Sediment samples

Recovery percentages obtained for both pesticides and pharmaceuticals in spiked sediment extracts with the different solvents (methanol, *n*-hexane, dichloromethane, methanol:formic acid (96:4, v:v), methanol:acetone (95:5, , v:v), acetonitrile:formic acid (99:1, v:v)) were compared in order to identify the best solvent/mixture to be further optimized (Figure 1). In general, methanol or methanol mixtures presented better recoveries, although some low recoveries were observed for the pesticides carbendazim, BIT, imazalil and for the iodinated X ray contrast agents. A careful look on methanol-based extracts showed higher recovery efficiency for mixture with either formic acid or acetone. Once the recoveries for methanolic extracts were very similar among themselves, the next step to choose the best solvent passed by visually study the quality of the different chromatograms. The interpretation of the signal to noise ratio based on chemical noise (Typical chromatogram shown in Figure S1) was used to evaluate chemical background effects and interferences, and also the reproducibility of the two most promising mixtures.

An extraction with methanol:aqueous formic acid resulted in higher chemical background noise than acetone. For the pesticides, triclosan and tebuconazole were affected by the background noise resulting in recovery rates exceeding 100%. On the other hand, with acetone good recoveries were obtained for all pesticides except BIT and carbendazim. For pharmaceuticals, the mixture methanol:acetone also provided better resolved peaks. The final decision was in favour of methanol:acetone (95:5, v:v) for both pesticides and pharmaceuticals as a compromise for lower recoveries but having chromatograms with less

1  
2  
3  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60

346 background noise, less interference peaks and well defined target compound  
347 peaks.

348 The introduction of a condensation/evaporation step is a common  
349 practice along extraction procedures, typically due to solvents change or as a  
350 pre-concentration step. Thus, differences in recovery using methanol:acetone  
351 (95:5, v:v) were also accessed with direct analysis of the extract or using a pre-  
352 concentration step by drying and redissolution (in water:methanol 50:50, v:v) in  
353 order to achieve a 10x concentration factor, Table 1. For pesticides, there were  
354 no differences in the recovery (carbendazim, BIT, mecoprop) or there was a  
355 significant negative effect on the recoveries (imazalil, terbutryn, diuron and  
356 triclosan) and a significant increase in the recovery of tebuconazole. Due to the  
357 significant decrease of triclosan recovery, the use of the concentration step needs  
358 to be carefully evaluated depending on the target analytes of most interest for  
359 specific studies. However, for pharmaceuticals drying and redissolving improved  
360 significantly the recovery rate of the iodinated pharmaceuticals, without impact  
361 on the other compounds. The evaporation step resulted in precipitation of  
362 particles that were not redissolved by the mixture water:methanol (50:50).  
363 These particles most probably worked as a sink for the more hydrophobic  
364 compounds present in the extract. This co-precipitation explains both the  
365 reduced recovery for some moderately hydrophobic target compounds ( $\log K_{ow}$   
366 2.67 – 4.66) and the decrease in background noise in the chromatogram.  
367 Therefore, there was increased S/N of the target peaks rather than a true  
368 recovery improvement.

369 Once sample extracts resulted in clean chromatograms and similar or  
370 better recoveries than the existing techniques (PLE, MAE) [29-31], the use of  
371 sequential extraction (commonly used) or further extract clean-up were not  
372 considered in order to ensure a fast and simple method.

373

374

375 Plant samples

376 For the optimization stage, only leaf material was used. As leaf extracts  
377 were expected to show higher backgrounds, they were not analysed directly, but  
378 only after the evaporation to dryness and a redissolution (in water:methanol  
379 50:50) step. The recovery percentages of the pharmaceuticals and pesticides  
380 were evaluated for the most promising solvent/mixture (methanol, *n*-hexane,

1  
2  
3  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60

381 dichloromethane, methanol:formic acid (96:4, v:v), methanol:acetone (95:5, v:v),  
382 acetonitrile:formic acid (99:1, v:v)) (Figure 2).

383 Main results considering both pesticides and pharmaceuticals are that  
384 either some compounds show low recovery efficiencies (< 50%) or recoveries  
385 are higher than 120% as a consequence of high background influence on results  
386 (typical chromatogram shown in Figure S2). For pesticides, independently of the  
387 solvent used, the chemical background noise in the first part of the  
388 chromatographic run resulted in poor recovery for carbendazim,  
389 benzoisothiazoline and imazalil. As for the sediments, x-ray contrast agents had  
390 lower recoveries also in plant extracts, while the propranolol peak was  
391 overlapping with the background noise. Additional solvents (acetone, ethanol)  
392 and mixtures of solvents in different proportions (dichloromethane:methanol, *n*-  
393 hexane:acetic acid) were tested without noticeable improvements (results not  
394 shown) to reduce the background influence while providing acceptable recovery  
395 rates. Therefore, optimization of a clean-up step was further pursued.

396 A commonly used technique for environmental samples clean-up is the  
397 employment of Florisil in the form of SPE cartridges, for a variety of organic  
398 contaminants such as organochlorine pesticides or PAHs. For the pesticides  
399 included in this study clean-up by Florisil presented a general improvement in  
400 the results by reducing the matrix effect considerably. However, the extracts still  
401 contained too much background to analyse carbendazim and benzoisothiazoline.  
402 Regarding the Florisil step in itself, benzoisothiazoline and mecoprop also  
403 showed reproducibility problems that could not be overcome by optimizing the  
404 elution solvent. For pharmaceuticals, the Florisil SPE step results (not shown)  
405 revealed the occurrence of strong sorption to the sorbent, not only of the  
406 chemicals responsible for the background but also the target compounds. The  
407 obtained extracts provided chromatograms with reduced background, but low  
408 recoveries. Possibly there were problems eluting the target analytes. Therefore,  
409 the use of Florisil SPE cartridges was further discarded.

410 The next option chosen for both pesticides and pharmaceuticals was a  
411 typical reverse phase SPE approach for water samples. For that, extracts (after  
412 drying) were re-dissolved in water and processed in polymeric SPE orthogonal  
413 to the separation column (i.e., Strata-X cartridges) as water samples. Although  
414 the improvement in the chromatographic run was noticeable as for Florisil  
415 cartridges, it was still not enough to eliminate the chromatogram background,

1  
2  
3  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60  
masking the results mainly for carbendazim and benzoisothiazoline (pesticides) and the x-ray contrast agents (pharmaceuticals). Use of SPE in these conditions would not ensure the quantification of all the compounds.

1  
2  
3  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60  
Therefore, a less commonly used but promising approach for sample clean-up tested was pigments saponification [26]. Chlorophylls and carotenes, are present in high concentrations in plants and will interfere in the analysis because they are extracted into the organic solvent. The saponification step addresses a base hydrolysis (at pH 13) of chlorophylls by cleavage of the two-ester bonds present in the chlorophylls. Nevertheless, it does not affect carotenes in the solution. Results revealed an improvement in the background removal showing clear chromatograms. For pesticides, the introduction of this saponification step resulted in less background and consequently in improved recovery (Figure 3) for the first pesticides of the run (early retention times) for all solvents, especially carbendazim and imazalil, and in general less co-eluted peaks with the target compounds. In fact, at this stage, *n*-hexane extraction followed by the saponification step was the most effective choice considering the amount of compounds and acceptable recoveries obtained. However, for pharmaceuticals, saponification was not as promising as for the pesticides (results not shown). Although showing chromatograms with less background, it was still not enough to reduce the interferences with the x-ray contrasts agents, as well as the last compound of the chromatographic run, diclofenac.

1  
2  
3  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60  
For the clean-up step, the use of less commonly applied materials was further considered. Activated carbon<sup>[32]</sup>, Sephadex LH-20® or LRA (Lipid Removal Agent) media® have been previously employed on environmental samples for clean-up procedures [33]. Preliminary tests using methanolic plant extracts (5 mL) spiked with the target compounds, mixed with the different materials (0.25 g) in an ultrasonic bath for 30 min, revealed (results not shown) a general improvement in the chromatogram background, after the analysis of the supernatant. Especially for activated carbon, the typical green colour of the plant extracts was completely removed. Nevertheless, for pesticides this also resulted in strong sorption of the pesticides to the activated carbon causing lower recoveries. For the other tested materials, LRA and Sephadex, the improvement in the chromatograms were still not sufficient to completely remove the background. For the pharmaceuticals, activated carbon was the most promising material, especially because it allowed the quantification of some of

1  
2 the x-ray contrast agent compounds. Further tests were performed by adding the  
3 activated carbon to the extracts obtained with the six solvents under screening  
4 (Figure 3). Although allowing an acceptable analysis of the x-ray contrasts  
5 agents, it resulted in lower recovery efficiency than previously observed with for  
6 instance SPE for the remaining compounds, especially naproxen and diclofenac.  
7  
8

9  
10 Considering the advantages and disadvantages of the previously tested  
11 steps, different procedural lines were further considered in order to clean-up the  
12 plant extracts. For pesticides, *n*-hexane at 100% was chosen as the most  
13 promising solvent for the extraction, and further efforts were placed in  
14 optimizing the saponification procedure, instead of working on improving the  
15 elution from activated carbon. For pharmaceuticals, activated carbon was  
16 considered to be more promising than the saponification step for improved  
17 recoveries of the iodinated compounds.  
18  
19

20 Final procedures establishment for pesticides was conducted by checking  
21 the pH influence in the SPE after the saponification step. The crude extract after  
22 evaporation to dryness was re-dissolved in methanolic KOH solution,  
23 ultrasonicated for 30 min, then the pH adjusted with hydrochloric acid (no  
24 adjustment, 2, 4, 5.5, 7) and further processed by SPE. A general improvement in  
25 recovery, except for imazalil, was observed when the pH of the KOH solution was  
26 adjusted to 5.5 before the SPE step, by comparison with no adjustment (Table 2).  
27  
28

29 Regarding pharmaceuticals, the last optimization step was to check which  
30 of the most promising solvents (Figure 3), methanol or methanol:acetone  
31 mixture (95:5, v:v) followed by the activated carbon clean-up step would provide  
32 the best and most reproducible results (Table 2). There were no significant  
33 differences in recovery between solvents, nevertheless the methanol:acetone  
34 mixture was chosen as it provided the highest recovery values. It should be  
35 noted that some of the recovery values obtained after the optimized clean-up  
36 step are lower than the methanolic (solution obtained by direct extracts  
37 evaporation to dryness and redissolution) extracts analysis. However, the  
38 existence of background noise on the extract analysis raises doubts on the  
39 reliability of this method when used as a routine for a high number of samples. In  
40 the present work, the choice of a multi approach overcomes individual best  
41 recoveries optimization for all compounds. Therefore, extraction with  
42 methanol:acetone mixture (95:5, v:v) followed by the activated carbon clean-up  
43  
44

1  
2  
3  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60  
485 was selected for the improvement in the reliability of iodinated compounds  
486 analysis compromising recovery efficiency of diclofenac and naproxen.

487

488 The final optimized procedures selected were (Figure 4):  
489 a) for sediments, samples were extracted with methanol:acetone (95:5, v/v) in  
490 an ultrasonic bath for both pesticides and pharmaceuticals. The extract was  
491 evaporated to dryness and dissolved in methanol prior to HPLC injection;  
492 b) for plant tissue, pesticides were extracted with *n*-hexane followed by  
493 saponification (KOH), pH adjustment and SPE (Strata X) steps; while  
494 pharmaceuticals were analysed after extraction with methanol:acetone (95:5,  
495 v:v), supernatant cleaning with activated carbon and drying and re-dissolving in  
496 methanol/water prior to HPLC injection.

497

498

### 499 3.2 Method characteristics and testing

500 Precision, limits of detection (LOD) and quantification (LOQ), were  
501 assessed for the final method. The HPLC instrument LOD and LOQ were  
502 determined based on the signal-to-noise ratio (S/N) of 3 and 10, respectively,  
503 and further confirmed by injection of decreasing concentrations of standards  
504 (Table S2). The overall methodology limits were calculated based on samples  
505 mass used for extraction and further confirmed by assessing S/N of spiked  
506 matrix extracts in the calculated limits range. Overall methodology precision was  
507 based on extracts analysis.

508

#### 509 Sediment samples

510 In sediment, the LODs and LOQs were calculated considering the  
511 extraction of 2 g of sediment sample. LODs ranged from 5 to 100 ng g<sup>-1</sup> for the  
512 pesticides and 15 to 50 ng g<sup>-1</sup> for the pharmaceuticals, while LOQ ranged from 25  
513 to 250 ng g<sup>-1</sup> for the pesticides and 50 to 150 ng g<sup>-1</sup> for the pharmaceuticals  
514 (Table 3). The characteristics of the method are consistent with the analysis of  
515 different organic contaminants in sediments using different extraction  
516 procedures (Table S3). The LODs for sediment samples were higher than those  
517 obtained for pesticides in sediment samples by LC-MS/MS (0.01 – 17 ng g<sup>-1</sup>) [13, 15,  
518 29, 34] or GC-MS (0.01 to 2 ng g<sup>-1</sup>). For example, a direct comparison of specific  
519 compounds across studies showed that the present LODs for terbutryn and

1 diuron (5 ng g<sup>-1</sup>), mecoprop and tebuconazole (50 ng g<sup>-1</sup>), and triclosan (40 ng g<sup>-</sup>  
2 1) were higher than those reported for PLE-LL-LC-HRMS/MS (0.05, 0.31, 0.4, 0.24  
3 and 0.89 ng g<sup>-1</sup>, respectively)<sup>[13]</sup> and PLE-SPE-LC-MS/MS (diuron 0.06 and  
4 mecoprop 4.17 ng g<sup>-1</sup>)<sup>[40]</sup>. For pharmaceuticals, the present LODs for sediment  
5 samples were higher than those obtained by LC-MS/MS (0.01 – 10 ng g<sup>-1</sup>)<sup>[13, 15, 21,</sup>  
6 <sup>35, 36]</sup> or GC-MS (0.3 – 6 ng g<sup>-1</sup>)<sup>[30, 37]</sup> and similar to pharmaceuticals  
7 determination in sediments by DAD (LOD < 167 ng g<sup>-1</sup><sup>[11]</sup> and LOQ of 1 -187 ng g<sup>-</sup>  
8 <sup>1</sup><sup>[38]</sup>). For example, the comparison for propanolol showed that the present LOD  
9 (15 ng g<sup>-1</sup>) was higher than that reported for USE-SPE-HPLC-DAD/FL (2 ng g<sup>-</sup>  
10 <sup>1</sup>)<sup>[38]</sup>, USE-SPE-LC-MS/MS (0.9 ng g<sup>-1</sup>)<sup>[15]</sup> and PLE-LL-LC-HRMS/MS (0.03 ng g<sup>-</sup>  
11 <sup>1</sup>)<sup>[13]</sup>. Main differences in LOD performance are related to the use of more  
12 powerful detector such as MS or MS/MS, and less to the extraction techniques.

13 The overall precision of the methodology was determined based on the  
14 intermediate precision (i.e., replicates analysed by HPLC-DAD on various  
15 working days) of the extraction recovery of 6 spiked sediment samples, including  
16 both 0.5 and 5 µg g<sup>-1</sup> level. This precision, reported as a relative standard  
17 deviation (RSD), was lower than 14 % (except for benzoisothiazoline 30%)  
18 (Table 3). Overall methodology recoveries (Table 3) ranged between 50 to 98%  
19 for the pharmaceuticals and 53 to 101% for the pesticides. For the  
20 pharmaceuticals, naproxen, and for the pesticides, benzoisothiazoline and  
21 triclosan, were the more affected compounds by the background noise resulting  
22 in poorer recoveries. Nevertheless, the obtained results are similar to previous  
23 published methodologies (Table S3) for sediment analysis of pesticides using  
24 simple solid-liquid extraction (40-125%)<sup>[39]</sup>, PLE followed by SPE (67 –  
25 118%)<sup>[40]</sup>, USE followed by SPE (68 – 102%)<sup>[15]</sup>, QuEChERS (46 – 102%)<sup>[34]</sup> or  
26 even MAE (81 – 112%)<sup>[37, 41]</sup>. For example, a direct comparison for carbendazim  
27 across studies showed that the present recovery (79%) is similar or higher to  
28 that reported for QuEChERS-LC-MS (61-80%)<sup>[34]</sup> and SLE-LC-MS (68%)<sup>[39]</sup>.  
29 Similarly, the current methodology recovery for pharmaceuticals is higher than  
30 obtained by Wagil, Maszkowska<sup>[35]</sup> (98 – 103%) and in the same range of  
31 previous works using MAE (25 – 81%)<sup>[11, 30]</sup> PLE (< 57 – 139 %)<sup>[13]</sup> or even USE  
32 followed by SPE (< 10 – 343%)<sup>[15, 21, 36, 38]</sup> (Table S3). For example, a comparison  
33 for carbamazepine showed that the present recovery (98%) was higher than that  
34 reported for USE-SPE-HPLC-DAD/FL (95%)<sup>[38]</sup>, MAE-HPLC-DAD (78%)<sup>[11]</sup> and  
35 PLE-LL-LC-HRMS/MS (72%)<sup>[13]</sup>.

1  
2  
3  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60555  
556        Plant samples

557        For plant tissue, LODs and LOQs were calculated considering the  
558 extraction of 0.2 g of sample. Values ranged from 0.05 to 1  $\mu\text{g g}^{-1}$  for LOD and  
559 from 0.25 to 2.5  $\mu\text{g g}^{-1}$  for LOQ for both the pesticides and the pharmaceuticals  
560 (Table 4). The overall methodology limits were higher than those obtained for  
561 pesticides in plant samples (Table S4) by LC-MS/MS (LOD of 3  $\text{ng g}^{-1}$  [29], LOQ of  
562 10  $\text{ng g}^{-1}$  [42, 43]) and GC-MS/MS (LOQ of 10  $\text{ng g}^{-1}$  [44]). For example, a direct  
563 comparison for tebuconazole across studies showed that the present LOQ (2  $\mu\text{g}$   
564  $\text{g}^{-1}$ ) was higher than that reported for dispersive-SPE-LC-MS/MS (100  $\text{ng g}^{-1}$ )<sup>[42]</sup>.  
565 For pharmaceuticals, the present limits for plant material were higher than those  
566 (Table S4) by LC-MS (LOD 2 – 13  $\text{ng g}^{-1}$ )<sup>[14]</sup>, LC-MS/MS (LOD of 0.5 to 8  $\text{ng g}^{-1}$ <sup>[14,</sup>  
567 <sup>45]</sup>) or GC-MS (7 – 58  $\text{ng g}^{-1}$ )<sup>[14]</sup>. For example, a comparison for carbamazepine  
568 showed that the present LOD (0.25  $\mu\text{g g}^{-1}$ ) was higher than that reported for  
569 buffer extraction followed by SPE-GC-MS (10-20  $\text{ng g}^{-1}$ )<sup>[14]</sup>, PLE-SPE-GC-MS (19  
570  $\text{ng g}^{-1}$ )<sup>[14]</sup>, QuEChERS-LC-MS/MS (0.7  $\text{ng g}^{-1}$ )<sup>[45]</sup> or PLE-SPE-LC-MS (0.17  $\text{ng g}^{-1}$ )<sup>[14]</sup>. Again, the main differences in LOD performance are related to the use in  
572 other works of a powerful detector such MS and less to the extraction and clean-  
573 up technique.

574        For the optimized conditions, recoveries (Table 4) ranged between 9 to  
575 99% for the pharmaceuticals and 56 to 103% for the pesticides. The proposed  
576 methodology is not appropriate for iopamidol (25 %), propranolol (31%),  
577 naproxen (9%) and diclofenac (46%) quantification in plant tissue samples. The  
578 recoveries of the remaining pharmaceuticals were above 65%. For the pesticides,  
579 acceptable recoveries for this type of matrix (above 75%) were determined with  
580 the exception of benzoisothiazoline (56% recovery). The obtained recoveries are  
581 generally similar or higher than those previous published (Table S4) for  
582 pesticides in plant tissue samples using dispersive-SPE (72 – 104%)<sup>[42]</sup>, solid-  
583 liquid extraction followed by salting out and SPE steps (10 – 120%)<sup>[44]</sup> and  
584 QuEChERS (80 - 136%)<sup>[43]</sup>. For example, a comparison for tebuconazole across  
585 studies showed that the present recovery (92%) was similar to the one reported  
586 for dispersive-SPE-LC-MS/MS (94%)<sup>[42]</sup>. Similarly, the current methodology  
587 recovery for pharmaceuticals are generally similar or higher than obtained using  
588 buffer extraction followed by SPE (15 – 98%)<sup>[14]</sup>, USE followed by SPE (73 –  
589 192%)<sup>[14]</sup>, PLE with<sup>[14]</sup> or without<sup>[45]</sup> SPE (46 – 176%) and QuEChERS (70 –

1  
2 590 119%)<sup>[45]</sup> (Table S4). For example, a comparison for carbamazepine showed  
3 591 that the present recovery (82%) was higher than that reported for buffer  
4 592 extraction followed by SPE-GC-MS (15-61%)<sup>[14]</sup>, PLE-SPE-GC-MS (75%)<sup>[14]</sup>, and  
5 593 similar or lower than QuEChERS-LC-MS/MS (84-96)<sup>[45]</sup> and PLE-SPE-LC-MS  
6 594 (110)<sup>[14]</sup>.  
7  
8

9  
10 595 The overall precision of the methodology was determined as the  
11 596 intermediate precision (i.e., replicates analysed by HPLC-DAD on various  
12 597 working days) of the extraction of different spiked plant tissue (*Typha* and  
13 598 *Berula* n=2) parts (leaves n=3 and roots n=3), including both 2.5 and 5  $\mu\text{g g}^{-1}$   
14 599 level. This precision, reported as a relative standard deviation (RSD), was lower  
15 600 than 21% (except for iopromide, 38%). These results suggest good method  
16 601 repeatability, even considering different type of plant tissue (leafs and roots). In  
17 602 fact, in previous works the RSD for pharmaceuticals has been considered matrix-  
18 603 dependent<sup>[10]</sup>. The RSDs presently obtained (6-38%) is within the range  
19 604 previously found for pesticides and pharmaceuticals determination in plant  
20 605 tissue<sup>[43-45]</sup>.  
21  
22

23 606 The use of the standard addition method could improve the overall  
24 607 quality of the proposed methodology for both sediment and plant analysis.  
25 608 However, that would have negative impact on simplicity and sample throughput.  
26 609 Since the objective was to establish a reliable but fast and simple method, the  
27 610 standard addition methodology was disregarded in the present study. Another  
28 611 option especially interesting for MS detectors would be the use of stable isotope  
29 612 labelled internal and/or surrogate standards, although Zhou, Ying<sup>[36]</sup> showed  
30 613 that even the addition of internal standards does not always overcome the  
31 614 matrix effects obtained for sediment samples. The sensitivity of HPLC-MS/MS is  
32 615 very dependent on the chemical ionisation procedure that is conditioned by the  
33 616 sample, the analyte, the eluent and the ion source design<sup>[22]</sup>. The use of matrix-  
34 617 matched calibration can be an interesting approach to minimize the matrix  
35 618 effects<sup>[20]</sup>. However, to match the matrix of the calibration standards with all  
36 619 individual plant samples (i.e., standard addition technique) can result in  
37 620 extended number of injections and consequently instrument time. Therefore, for  
38 621 MS detectors the use of internal standards is preferred over matrix-matched  
39 622 calibration<sup>[46]</sup>. Application of the methodology should be accompanied by  
40 623 recovery tests on the specific substrate to ensure a proper quality assurance and  
41 624 control.  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60

1  
2  
3  
4  
5  
6  
625

626

627 3.3 Application to real samples628        The optimized and validated methodology has subsequently been used in  
629 different studies focused on removal of micropollutants from water by  
630 constructed wetland mesocosm systems. As an example of the method  
631 applicability, the quantification of the total accumulation of imazalil in a  
632 constructed wetland mesocosms substrate/sediment continuously run over 9  
633 months under various hydraulic loading rates and imazalil concentrations of  
634 both 10 and 100  $\mu\text{g L}^{-1}$  (Figure S3) [28], as well as ibuprofen accumulation in  
635 plant tissue (roots and leaves) after exposure to an initial spike of 10  $\text{mg L}^{-1}$  in  
636 the hydroponic media (Figure S4) [27]. In a recent work by the authors, studying  
637 an initial exposure of *Phragmites australis* to 10  $\mu\text{g L}^{-1}$  of imazalil in hydroponic  
638 solution, plant extracts obtained with the present methodology were successfully  
639 analysed by HPLC-MS/MS for quantification of imazalil enantiomers and  
640 screened for transformation products with success [47]. The intra-equipment  
641 deviation for control samples ( $n \geq 8$ ) analysed by both HPLC-DAD and HPLC-  
642 MS/MS were below 15% for the quantification of imazalil in plant tissue [47].643        The validated methodologies proved fit-for-purpose in quantifying  
644 multiple classes of pesticides and pharmaceuticals in complex matrices.  
645 However, a broader application of the current methodology should be  
646 approached carefully. The use of a non-selective (non-confirmatory) DAD  
647 detector is only recommended when dealing with systems studied under  
648 controlled conditions. The application to field-samples should always be coupled  
649 with a confirmation step, or in alternative, the current extraction and clean-up  
650 steps can also be coupled with LC-MS. Nevertheless, as discussed before, the  
651 coupling to LC-MS, needs to be validated prior to full application specialty to  
652 assess matrix-effects and ion suppression in the detector. The range of  
653 compounds studied was broad and the methods may be applied for other  
654 compounds from the same family, chemical properties. But such application will  
655 always require a validation step.656        The proposed USE methodology is a fast, easily accessible and effective  
657 alternative to the most advanced PLE or MAE methods (Table S3 and S4). Sample  
658 preparation time will be grossly similar across platforms. However, USE  
659 (presently, 24 samples in 30 min) and MAE (typically 24 samples in 40 min)

allow the simultaneous extraction of samples being faster than PLE, which implies a sequential process (typically 20 min per cell, resulting in 24 samples in 8 hours). USE extraction is done in disposable glass vials, while MAE and PLE require additional clean-up and decontamination of the Teflon vessels or cells after use. PLE and MAE require an additional programming of an extraction/sequence procedure. Therefore, sample throughput is larger for USE. It should be noted that as drawback, USE does not have any automated control over the extraction process, as can be achieved by MAE and PLE. The difference in cost and accessibility to a simple ultrasonic bath that can be used for USE and the more advanced and dedicated equipment for MAE or PLE with the respective dedicated consumables is distinct.

671

672

#### 673 **4. Conclusions**

674 The here established USE methods with the different optimized clean-up  
675 and pre-concentration steps coupled to HPLC-DAD analysis demonstrated  
676 suitable sensitivity and reliability, and proved fit-for-purpose in quantifying  
677 multiple classes of pesticides and pharmaceuticals in complex matrices such as  
678 sediment and plant tissue. For sediments, an acceptable extraction efficiency (50  
679 - 101%) and RSD < 14% (except for benzoisothiazoline) were achieved without  
680 performing any clean-up step. The complex matrix of plant tissues poses specific  
681 problems, especially for improving the methodology recoveries. Thus, the final  
682 optimized method implies individualized approaches for the extraction of  
683 pesticides and pharmaceuticals. The established final method shows in general  
684 an acceptable extraction efficiency (> 46%) (except for iopamidol, propranolol  
685 and naproxen) with RSD < 21% (except iopromide) for different type of wetland  
686 plant tissues.

687 Compared with the existing methods in the literature, the proposed USE  
688 methodology is a fast, easily accessible, and effective alternative to PLE or MAE  
689 for extracting emerging contaminants from sediment and plant tissue samples.  
690 The methodology was successfully applied in different studies on the fate of  
691 pesticides and pharmaceuticals in water treatment eco-technology systems.

692

693

#### 694 **Acknowledgments**

1 695 Aarhus University Research Foundation (AUFF) funded Center for  
2 Advanced Water Purification. The PhD fellowships of Tao Lv and Yang Zhang  
3 were supported by the China Scholarship Council (CSC).  
4  
5 698  
6  
7 699 **Conflict of Interest:** The authors declare that they have no conflict of interest.  
8  
9 700  
10 701  
11  
12 702 **References**  
13  
14 703

13  
14 704 [1] J. P. Bellenger, H. Cabana. Emerging contaminants: A scientific challenge  
15 without borders Preface. *Sci Total Environ.* **2014**, *487*,  
16 [2] K. E. Murray, S. M. Thomas, A. A. Bodour. Prioritizing research for trace  
17 pollutants and emerging contaminants in the freshwater environment.  
18 *Environmental Pollution.* **2010**, *158*,  
19 [3] T. A. Ternes, A. Joss, H. Siegrist. Scrutinizing pharmaceuticals and personal  
20 care products in wastewater treatment. *Environmental Science and Technology.*  
21 **2004**, *38*,  
22 [4] T. Ternes. The occurrence of micropollutants in the aquatic environment: a  
23 new challenge for water management. *Water Sci Technol.* **2007**, *55*,  
24 [5] J. Diamond, K. Munkittrick, K. E. Kapo, J. Flippin. A framework for  
25 screening sites at risk from contaminants of emerging concern. *Environmental  
Toxicology and Chemistry.* **2015**, *34*,  
26 [6] J. García. Advances in pollutant removal processes and fate in natural and  
27 constructed wetlands. *Ecological Engineering.* **2011**, *37*,  
28 [7] P. Verlicchi, E. Zambello. How efficient are constructed wetlands in  
29 removing pharmaceuticals from untreated and treated urban wastewaters? A  
30 review. *Sci Total Environ.* **2014**, *470–471*,  
31 [8] T. A. Ternes, M. Bonerz, N. Herrmann, D. Löffler, E. Keller, B. B. Lacida, et  
32 al. Determination of pharmaceuticals, iodinated contrast media and musk  
33 fragrances in sludge by LC tandem MS and GC/MS. *Journal of Chromatography A.*  
34 **2005**, *1067*,  
35 [9] Z. F. Chen, G. G. Ying, Y. S. Liu, Q. Q. Zhang, J. L. Zhao, S. S. Liu, et al.  
36 Triclosan as a surrogate for household biocides: An investigation into biocides in  
37 aquatic environments of a highly urbanized region. *Water Res.* **2014**, *58*,  
38 [10] X. Wu, J. L. Conkle, J. Gan. Multi-residue determination of pharmaceutical  
39 and personal care products in vegetables. *Journal of chromatography A.* **2012**,  
40 *1254*,  
41 [11] L. Sanchez-Prado, C. Garcia-Jares, M. Llompart. Microwave-assisted  
42 extraction: Application to the determination of emerging pollutants in solid  
43 samples. *J Chromatogr A.* **2010**, *1217*,  
44 [12] D. M. Pavlovic, S. Babic, A. J. M. Horvat, M. Kastelan-Macan. Sample  
45 preparation in analysis of pharmaceuticals. *Trac-Trend Anal Chem.* **2007**, *26*,  
46 [13] A. C. Chiaia-Hernandez, M. Krauss, J. Hollender. Screening of lake  
47 sediments for emerging contaminants by liquid chromatography atmospheric  
48 pressure photoionization and electrospray ionization coupled to high resolution  
49 mass spectrometry. *Environmental science & technology.* **2013**, *47*,

1  
2  
3  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60

[14] V. Matamoros, D. Calderon-Preciado, C. Dominguez, J. M. Bayona. Analytical procedures for the determination of emerging organic contaminants in plant material: a review. *Analytica chimica acta*. **2012**, *722*, [15] H. Darwano, S. V. Duy, S. Sauve. A new protocol for the analysis of pharmaceuticals, pesticides, and hormones in sediments and suspended particulate matter from rivers and municipal wastewaters. *Archives of environmental contamination and toxicology*. **2014**, *66*, [16] M. C. Bruzzoniti, L. Checchini, R. M. De Carlo, S. Orlandini, L. Rivoira, M. Del Bubba. QuEChERS sample preparation for the determination of pesticides and other organic residues in environmental matrices: a critical review. *Analytical and bioanalytical chemistry*. **2014**, *406*, [17] S.-B. Consuelo, L. T. José, A. Beatriz. in *Analysis of Pesticides in Food and Environmental Samples* 2008, (CRC Press). [18] B. Gilbert-López, J. F. García-Reyes, A. Molina-Díaz. Sample treatment and determination of pesticide residues in fatty vegetable matrices: A review. *Talanta*. **2009**, *79*, [19] K. M. Dimpe, P. N. Nomngongo. Current sample preparation methodologies for analysis of emerging pollutants in different environmental matrices. *TrAC Trends in Analytical Chemistry*. **2016**, *82*, [20] J. M. Montiel-León, S. V. Duy, G. Munoz, M. Amyot, S. Sauvé. Evaluation of on-line concentration coupled to liquid chromatography tandem mass spectrometry for the quantification of neonicotinoids and fipronil in surface water and tap water. *Analytical and bioanalytical chemistry*. **2018**, *410*, [21] B. Albero, C. Sánchez-Brunete, A. I. García-Valcárcel, R. A. Pérez, J. L. Tadeo. Ultrasound-assisted extraction of emerging contaminants from environmental samples. *TrAC Trends in Analytical Chemistry*. **2015**, *71*, [22] K. Bester. Quantification with HPLC-MS/MS for environmental issues: quality assurance and quality assessment. *Analytical and bioanalytical chemistry*. **2008**, *391*, [23] P. N. Carvalho, M. C. Basto, C. M. Almeida, H. Brix. A review of plant-pharmaceutical interactions: from uptake and effects in crop plants to phytoremediation in constructed wetlands. *Environmental science and pollution research international*. **2014**, *21*, [24] M. Shenker, D. Harush, J. Ben-Ari, B. Chefetz. Uptake of carbamazepine by cucumber plants – A case study related to irrigation with reclaimed wastewater. *Chemosphere*. **2011**, *82*, [25] W. W. Buchberger. Current approaches to trace analysis of pharmaceuticals and personal care products in the environment. *J Chromatogr A*. **2011**, *1218*, [26] A. Dugay, C. Herrenknecht, M. Czok, F. Guyon, N. Pages. New procedure for selective extraction of polycyclic aromatic hydrocarbons in plants for gas chromatographic-mass spectrometric analysis. *Journal of chromatography A*. **2002**, *958*, [27] Y. Zhang, T. Lv, P. N. Carvalho, C. A. Arias, Z. Chen, H. Brix. Removal of the pharmaceuticals ibuprofen and iohexol by four wetland plant species in hydroponic culture: plant uptake and microbial degradation. *Environmental Science and Pollution Research*. **2016**, *23*, [28] T. Lv, Y. Zhang, L. Zhang, P. N. Carvalho, C. A. Arias, H. Brix. Removal of the pesticides imazalil and tebuconazole in saturated constructed wetland mesocosms. *Water Res*. **2016**, *91*, [29] E. Maillard, G. Imfeld. Pesticide mass budget in a stormwater wetland. *Environmental science & technology*. **2014**, *48*,

1  
2  
3  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60

793 [30] J. Kumirska, N. Migowska, M. Caban, P. Lukaszewicz, P. Stepnowski.  
794 Simultaneous determination of non-steroidal anti-inflammatory drugs and  
795 oestrogenic hormones in environmental solid samples. *Sci Total Environ.* **2015**,  
796 508,  
797 [31] S. Babić, D. Mutavdžić Pavlović. in *Comprehensive Analytical Chemistry*  
798 Eds. Mira Petrović DB, Sandra P)2013, pp. 129-67 (Elsevier).  
799 [32] H. Dabrowska, L. Dabrowski, M. Biziuk, J. Gaca, J. Namiesnik. Solid-phase  
800 extraction clean-up of soil and sediment extracts for the determination of various  
801 types of pollutants in a single run. *Journal of Chromatography A.* **2003**, 1003,  
802 [33] S. W. C. Chung, B. L. S. Chen. Determination of organochlorine pesticide  
803 residues in fatty foods: A critical review on the analytical methods and their  
804 testing capabilities. *Journal of Chromatography A.* **2011**, 1218,  
805 [34] M. Kvicalova, P. Doubravova, R. Jobanek, M. Jokesova, V. Ocenaskova, H.  
806 Sussenbekova, et al. Application of Different Extraction Methods for the  
807 Determination of Selected Pesticide Residues in Sediments. *Bulletin of*  
808 *Environmental Contamination and Toxicology.* **2012**, 89,  
809 [35] M. Wagil, J. Maszkowska, A. Bialk-Bielinska, P. Stepnowski, J. Kumirska. A  
810 comprehensive approach to the determination of two benzimidazoles in  
811 environmental samples. *Chemosphere.* **2015**, 119 Suppl,  
812 [36] L.-J. Zhou, G.-G. Ying, S. Liu, J.-L. Zhao, F. Chen, R.-Q. Zhang, et al.  
813 Simultaneous determination of human and veterinary antibiotics in various  
814 environmental matrices by rapid resolution liquid chromatography-electrospray  
815 ionization tandem mass spectrometry. *J Chromatogr A.* **2012**, 1244,  
816 [37] P. N. Carvalho, P. N. Rodrigues, F. Alves, R. Evangelista, M. C. Basto, M. T.  
817 Vasconcelos. An expeditious method for the determination of organochlorine  
818 pesticides residues in estuarine sediments using microwave assisted pre-  
819 extraction and automated headspace solid-phase microextraction coupled to gas  
820 chromatography-mass spectrometry. *Talanta.* **2008**, 76,  
821 [38] J. Martín, J. L. Santos, I. Aparicio, E. Alonso. Multi-residue method for the  
822 analysis of pharmaceutical compounds in sewage sludge, compost and sediments  
823 by sonication-assisted extraction and LC determination. *J Sep Sci.* **2010**, 33,  
824 [39] A. Lazartigues, C. Fratta, R. Baudot, L. Wiest, C. Feidt, M. Thomas, et al.  
825 Multiresidue method for the determination of 13 pesticides in three  
826 environmental matrices: water, sediments and fish muscle. *Talanta.* **2011**, 85,  
827 [40] M. Kock-Schulmeyer, M. Olmos, M. L. de Alda, D. Barcelo. Development of a  
828 multiresidue method for analysis of pesticides in sediments based on isotope  
829 dilution and liquid chromatography-electrospray-tandem mass spectrometry.  
830 *Journal of Chromatography A.* **2013**, 1305,  
831 [41] L. Wu, M. Hu, Z. Li, Y. Song, C. Yu, H. Zhang, et al. Dynamic microwave-  
832 assisted extraction combined with continuous-flow microextraction for  
833 determination of pesticides in vegetables. *Food Chemistry.* **2016**, 192,  
834 [42] S. Walorczyk, D. Drożdżyński, R. Kierzek. Determination of pesticide  
835 residues in samples of green minor crops by gas chromatography and ultra  
836 performance liquid chromatography coupled to tandem quadrupole mass  
837 spectrometry. *Talanta.* **2015**, 132,  
838 [43] A. Abad-Fuentes, E. Ceballos-Alcantarilla, J. V. Mercader, C. Agulló, A.  
839 Abad-Somovilla, F. A. Esteve-Turrillas. Determination of succinate-  
840 dehydrogenase-inhibitor fungicide residues in fruits and vegetables by liquid  
841 chromatography-tandem mass spectrometry. *Analytical and bioanalytical*  
842 *chemistry.* **2015**, 407,

1  
2  
3  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60

843 [44] S. Saito-Shida, S. Nemoto, R. Teshima. Multiresidue determination of  
844 pesticides in tea by gas chromatography-tandem mass spectrometry. *Journal of*  
845 *Environmental Science and Health, Part B*. **2015**, *50*,  
846 [45] Y. H. Chuang, Y. Zhang, W. Zhang, S. A. Boyd, H. Li. Comparison of  
847 accelerated solvent extraction and quick, easy, cheap, effective, rugged and safe  
848 method for extraction and determination of pharmaceuticals in vegetables.  
849 *Journal of chromatography A*. **2015**, *1404*,  
850 [46] A. K. Hewavitharana. Matrix matching in liquid chromatography-mass  
851 spectrometry with stable isotope labelled internal standards—Is it necessary?  
852 *Journal of Chromatography A*. **2011**, *1218*,  
853 [47] T. Lv, P. N. Carvalho, M. E. Casas, U. E. Bollmann, C. A. Arias, H. Brix, et al.  
854 Enantioselective uptake, translocation and degradation of the chiral pesticides  
855 tebuconazole and imazalil by *Phragmites australis*. *Environmental pollution*.  
856 **2017**, *229*,

857