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Monolithic capsule phase microextraction prior to gas chromatography-mass
spectrometry for the determination of organochlorine pesticides in
environmental water samples
Antonio Ferracane ^{a,b} , Natalia Manousi ^{b,c*} , Abuzar Kabir ^d , Kenneth G. Furton ^d , Peter
Q. Tranchida ^a , George A. Zachariadis ^c , Justyna Płotka-Wasylka ^e , Luigi Mondello ^{a,f} ,
Victoria F. Samanidou ^c , Erwin Rosenberg ^b
^a Department of Chemical, Biological, Pharmaceutical and Environmental Sciences,
University of Messina, Messina, Italy
^b Institute of Chemical Technologies and Analytics, Vienna University of Technology,
1060 Vienna, Austria
^c Laboratory of Analytical Chemistry, Department of Chemistry, Aristotle University of
Thessaloniki, Thessaloniki 54124, Greece
^d Department of Chemistry and Biochemistry, Florida International University, Miami,
FL, USA
^e Department of Analytical Chemistry, Faculty of Chemistry and BioTechMed Center,
Gdansk University of Technology, 1/12 G. Narutowicza St., 80-233 Gdansk, Poland
^f Chromaleont s.r.l., c/o Department of Chemical, Biological, Pharmaceutical and
Environmental Sciences, University of Messina, Messina, Italy
*Corresponding author: nmanousi@chem.auth.gr; Laboratory of Analytical
Chemistry, Department of Chemistry, Aristotle University of Thessaloniki, Thessaloniki
54124, Greece
Abstract: In this study, a capsule phase microextraction (CPME) protocol followed by
gas chromatography-mass spectrometry is proposed for the accurate and sensitive
monitoring of organochlorine pesticides (OCPs) in environmental water samples.
Different monolithic sol-gel encapsulated sorbents were compared and monolithic sol-
gel poly(ethylene glycol)-based sorbent incorporated into porous microextraction
capsules resulted in the highest extraction efficiency. Following the selection of the
microextraction device, the CPME conditions were optimized, while linearity, limits
detection (LODs), limits of quantification (LOQs), accuracy and precision were the
figures-of-merit measured. Under optimum conditions the LODs for the OCPs ranged

between 0.01 ng mL⁻¹ and 0.03 ng mL⁻¹. The relative standard deviations were better than 5.3% and 9.2% for intra-day and inter-day study, respectively. The relative recoveries ranged between 90.5-105.2% (intra-day) and 93.0-105.0% (inter-day), demonstrating good method trueness. Finally, the proposed protocol was implemented for the monitoring of OCPs in tap, river, pond, and lake water. The developed method presents better analytical features than existing methods.

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Keywords: Organochlorine pesticides; capsule phase microextraction; water; sample
preparation, GC-MS, monolithic sorbents

- 44
- 45
- 46 **1. Introduction**

Pesticides are used in agriculture to repel, prevent, mitigate, or destroy pests [1]. 47 48 They are classified into different types including insecticides, herbicides, nematicides, rodenticides, fungicides, molluscicides and plant growth regulators [2]. Pesticide 49 50 residues may constitute a significant source of contamination for the environment since they can be present in air, water and soil [3]. Nowadays, water resources pollution 51 52 because of the uncontrolled use of pesticides represents a significant environmental and health threat. In order to be able to design and implement strategy plans for 53 54 environmental and human health protection, the estimation of the type and amount of pesticide residues in water resources is crucial [4]. 55

Organochlorine pesticides (OCPs) are a class of pesticides used for controlling 56 vector-borne diseases (e.g., typhus and malaria) and to destroy pests due to their low 57 cost and high efficiency [5,6]. These compounds were the first major pesticide class 58 59 that was produced on large scale since the rapid growth of the pesticide industry in the 60 late 1940's [7]. Because of their chemical stability, their high persistence in the natural environment and their low biodegradability, OCPs are ubiquitous among 61 environmental, food and biological samples [3,8]. These compounds are responsible for 62 a wide variety of adverse effects including damages to the human nervous system, 63 cancer predisposition, reproductive disorders, and disruption of the cellular immune 64 system. As a result, their use has been banned by the United Nations Environment 65 Program, while they have been also listed as persistent organic pollutants by the 66 Stockholm Convention [3,9]. Moreover, OCPs have been reported to be toxic by both 67 the World Health Organization (WHO) and the Environmental Protection Agency 68

(EPA) [8]. The toxicity of these pollutants is clearly related to their chemical structure
[5]. Although the use of OCPs has been banned in most industrialized advanced
countries and their production has been terminated, their lasting and adverse influence
on aquatic biota, human health and wildlife still causes concerns regarding the
monitoring of the levels of these compounds in environmental samples [10].

74 The main difficulties of the determination of these analytes in real-world 75 samples are related to their low concentrations, the sample complexity, and the potential presence of a wide range of interfering compounds. Therefore, an extraction and 76 77 enrichment step is typically required prior to the determination of the OCPs using an instrumental analytical technique [11,12]. Conventional approaches for the extraction 78 of OCPs include liquid-liquid extraction (LLE) and solid-phase extraction (SPE) that 79 show high consumption of hazardous organic solvents, increased number of complex 80 steps and high waste generation [13]. 81

82 After the introduction of Green Analytical Chemistry (GAC) [14] that aims to provide the principles for developing of more sustainable and environmentally friendly 83 84 methodologies, SPE and LLE tend to be progressively replaced by microextraction approaches. Thus, several novel methodologies including solid phase microextraction 85 86 [15], dispersive solid-phase extraction (d-SPE) [16], magnetic solid-phase extraction 87 (MSPE) [10], fabric phase sorptive extraction (FPSE) [17], stir bar sorptive extraction (SBSE) [18], dispersive liquid-liquid extraction [19] and hollow fiber-liquid phase 88 microextraction [20] have been proposed for the accurate and sensitive monitoring of 89 90 OCPs in a wide range of samples. An interesting technique that was recently proposed for the monitoring of pesticides in environmental water samples is capsule phase 91 92 microextraction (CPME) [21].

In CPME, the analytes are extracted by appropriately designed devices that are 93 made of two permeable microporous polypropylene tubes welded together to form a 94 single, bipartite microextraction capsule. One polypropylene capillary tube contains a 95 96 sol-gel hybrid organic-inorganic sorbent, while the other capillary tube contains a cylindrical magnet that provides to the device the ability to spin when a magnetic field 97 98 is implemented [22]. The integration of sample stirring can efficiently simplify the extraction procedure to a large extent, while it prevents potential loss of analytes due to 99 their retention on external devices and it results in increased extraction yield [23]. 100 Moreover, the polypropylene membranes exhibit inherent porosity and thus the 101 capsules can be directly employed for the analysis of sample matrices containing 102

insoluble matrix interferants, debris and particulates, without any requirement of 103 sample pretreatment (i.e., filtration) [21]. CPME is based on equilibrium extraction and 104 due to the integration of the stirring mechanism, rapid extraction kinetics can be 105 obtained [24]. An important characteristic of CPME is the utilization of sol-gel 106 technology to prepare hybrid organic-inorganic porous products of various shapes, 107 sizes, and formats. Sol-gel sorbents exhibit good chemical and thermal stability, 108 selectivity, tunable porosity and high purity resulting in the fabrication of powerful 109 microextraction devices [25]. 110

111 In this work, we aimed to develop a simple and rapid method for the determination of ten OCPs in water samples by CPME combined with gas 112 chromatography-mass spectrometry (GC-MS). Various monolithic sol-gel coated 113 microextraction capsules were evaluated for their performance. Subsequently, the 114 CPME procedure was optimized to ensure that the highest possible extraction efficiency 115 is obtained. After method optimization, the CPME GC-MS protocol was validated. As 116 a final step, the validated methodology was applied for the monitoring of OCPs in 117 118 different environmental water samples.

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120 **2.** Experimental

121 **2.1. Reagents and chemicals**

Acetonitrile and methanol of LC-MS grade were obtained from Honeywell 122 (Charlotte, North Carolina, USA). HPLC grade acetone (ACE) and reagent grade NaCl 123 were obtained from Merck (Darmstadt, Germany). α -HCH (99.9%), β -HCH (98.4%), 124 γ-HCH (99.7%), alachlor (99.3%), aldrin (98.5%), p,p'-DDE (99.8%), o,p'-DDD 125 (99.9%), p,p'-DDD (99.9%) and methoxychlor (98.7%) were obtained from Supelco 126 (Bellefonte, PA, USA). o,p'-DDE (99.39%) was obtained from Dr. Ehrenstorfer GmbH 127 (Augsburg, Germany). The structures of the target analytes are summarized in Figure 128 S1. Stock solutions ($c=1000 \text{ mg } \text{L}^{-1}$) for all analytes were prepared in methanol. Multi-129 130 analyte working solutions were prepared daily with appropriate dilution in methanol.

Capsule phase microextraction devices were built using Membrana Accurel® porous capillary membranes, which were purchased from 3M Inc. (St. Paul, MN, USA). Cylindrical magnetic rods (1/4" x 1/16") were purchased from K&J Magnetics Inc. (Pipersville, PA, USA). Sol-gel synthesis materials, namely methyl trimethoxysilane (MTMS), tetramethyl orthosilicate (TMOS), poly(tetrahydrofuran) (PTHF), and polyethylene glycol 300 (PEG 300), were obtained from Sigma-Aldrich (St. Louis, MO, USA). Poly(dimethylsiloxane) (PDMS), octadecyltrimethoxysilane (C₁₈),
poly(dimethyldiphenylsiloxane) (PDMDPS) were obtained from Gelest Inc.
(Morrisville, PA, USA). Ammonium hydroxide, methylene chloride, isopropanol, and
hydrochloric acid were purchased from Fisher Scientific (Milwaukee, WI, USA).

Environmental water samples (*i.e.*, lake, river, pond and tap water) were collected in Vienna, Austria. Amber-glass vials with no headspace were used for sample collection and storage. All samples were stored at 4°C, while no sample pretreatment was required prior to the CPME GC-MS procedure.

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146 **2.2. Instrumentation**

A Shimadzu GC-2010 instrument coupled to a QP2010 Plus mass spectrometer 147 (Shimadzu, Kyoto, Japan) was used for the quantification of the OCPs. Separation was 148 achieved using Helium (99.999%) as mobile phase that was delivered at a flow rate of 149 1.00 mL min⁻¹. Constant linear velocity was employed as flow control mode. An Rtx-150 5MS (30 m \times 0.25 mm, 0.25 μ m) column (Restek Corporation, Bellefonte, PA, USA) 151 152 was used under the following oven temperature program: 100 °C initial temperature (hold time: 2.5 min), increased to 200 °C (rate: 15 °C min⁻¹), then increased to 250 °C 153 (rate: 5 °C min⁻¹) and finally increased to 300°C (rate: 6 °C min⁻¹). The run time and 154 the solvent delay were 27.5 min and 7.0 min, respectively. The injector temperature, 155 156 the ion source temperature and the interface temperature were 280°C, 220°C and 250 $^{\circ}$ C, respectively. The injection volume was 2 μ L and high-pressure injection (450 kPa) 157 took place. Finally, the OCPs were quantified at the selected ion monitoring (SIM) 158 mode. For each analyte, one target ion was used as quantifier, while two reference ions 159 were used as qualifiers. Table S1 shows the recorded m/z ratios for each analyte, as well 160 as their respective retention times. 161

162 The CPME procedure was carried out using a magnetic stirrer (Heidolph163 Instruments GmbH & CO, Schwabach, Germany).

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166 2.3. Preparation of sol-gel monolithic sorbent encapsulated CPME devices

167 CPME devices with built-in magnet and encapsulated sol-gel PDMS, sol-gel 168 C18, sol-gel PEG 300, sol-gel PTHF and sol-gel PDMDPS monolithic sorbent beds 169 were prepared using a simple protocol which is illustrated in Figure 1.



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Figure 1. Preparation of the sol-gel monolithic sorbent encapsulated CPME devices

Environmental samples can be collected in large volumes and the target analytes are often at very low concentration levels. As such, a higher sorbent loading is needed to accomplish higher method sensitivity. Thus, CPME media of 3 cm length were used. The synthetical route for the fabrication of the microextraction capsules is described in Supplementary Material.

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179 **2.4 CPME procedure**

Initially, the CPME capsules were immersed into a vial containing 2 mL of
MeOH mixture for 5 min to remove impurities that remained from the material
preparation. Accordingly, the capsules were rinsed with water.

Following the activation step, the pre-treated CPME capsules were immersed 183 into 20 mL of sample solution containing 5% w/v NaCl and adsorption was carried out 184 185 within 50 min under stirring at 800 rpm. Following this step, the supernatant was discarded, and the capsule was rinsed with water and dried using lint free tissue. The 186 adsorbed OCPs were eluted by placing the CPME media into Eppendorf tubes 187 containing 250 µL acetone. Finally, the capsules were recovered, and eluent filtration 188 was performed using 0.22 µm polytetrafluoroethylene (Frisenette ApS, Knebel, 189 Denmark). Then, 2 µL was analyzed by GC-MS system. Figure 2 shows a 190 representative chromatogram of a blank river water sample (a) and a spiked river water 191 sample (b) subjected to the herein described protocol. 192

The monolithic sol-gel PEG 300 CPME device used was immersed in the initial mixture of ACN: MeOH (50:50 v/v) for 5 min. The utilization of the initial solvent mixture meets the requirements of GAC regarding material recycling and reusing [26]. Following this procedure, no carry-over effects were observed. After washing, the clean capsules were left to dry at ambient temperature, and they were stored in airtight sealed vials.



Figure 2. Chromatogram of (a) a blank river water sample and (b) a spiked river
water sample (c=20 ng mL⁻¹) after their sample preparation using the monolithic solgel PEG 300 CPME device

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2043. 3. Results and discussion

205 **3.1 Characterization of the CPME media**

The characterization of the CPME media by scanning electron microscopy and Fourier transform infrared spectroscopy has been previously conducted [27], with the aim of investigating the functional makeup of the sol-gel sorbents and the surface morphology of the sol-gel sorbent.

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211 **3.2 Optimization of CPME method**

212 **3.2.1 Selection of sol-gel coated microextraction capsule**

Initially, five monolithic sol-gel coated capsules were evaluated to assess the performance of different sorbents and to select the material with the highest affinity towards the target analytes. The examined sorbents were sol-gel PDMS, sol-gel C₁₈, sol-gel PEG 300, sol-gel PTHF and sol-gel PDMDPS. As shown, in Figure S2, sol-gel PEG 300 showed the best extraction performance towards the majority of the examined analytes. Higher extraction efficiency was observed for aldrin, o,p'-DDE and p,p'-DDE, only in the case of sol-gel PDMS. However, this sorbent exhibited significantly lower extraction efficiency for α -HCH, β -HCH and γ -HCH and thus, further experiments were conducted using sol-gel PEG 300 as a compromise for all the analytes.

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3.2.2 Optimization of the adsorption step

To ensure high extraction efficiency of the OCPs from the water samples, the experimental parameters that influence the performance of the adsorption step (*i.e.*, the stirring rate, the sample volume, the extraction time, and the salt content) were examined under a univariate approach. A spiked water sample solution ($c=10 \text{ ng mL}^{-1}$) was used in the optimization study. Table S2 summarizes the experimental conditions before and after the optimization study.

231 The effect of stirring rate was primarily investigated because it affects the analyte diffusion. For this purpose, four different stirring rates (i.e., 0, 400, 800, and 232 233 1000 rpm) were studied. According to the mass transfer theory, sample agitation is important to assist the movement of the analytes to the sol-gel sorbent surface with a 234 reduction in the thickness of the boundary layer in order to shorten the thermodynamic 235 equilibrium time [17]. As shown in Figure S3, sample stirring is critical for the 236 adsorption of the OCPs. Under no stirring (i.e., 0 rpm) negligible adsorption was 237 achieved. The performance of the CPME method increased at 800 rpm for most analytes 238 239 and it remained constant up to 1000 rpm. Thus, a stirring rate of 800 rpm was chosen.

Accordingly, the sample amount was studied using three different volumes, *i.e.*, 10 mL, 20 mL and 50 mL (Figure S4). A reduction of the extraction recovery was observed by increasing the sample volume from 10 mL to 50 mL. However, the utilization of 10 mL of sample results in lower preconcentration factors (PF) and thus in reduced method sensitivity, Therefore, an amount of 20 mL was used in the CPME method as a compromise between the extraction efficiency and the overall method sensitivity.

The extraction time is an significant factor in equilibrium-based techniques [28]. The extraction time was studied from 10 to 60 min (Figure 3). Equilibrium was achieved at 40 min for aldrin, alachlor, o,p'-DDE, p,p'-DDE, o,p'-DDD and p,p'-DDD while 50 min were required for α-HCH, β-HCH, γ-HCH and methoxychlor. Thus, an
extraction time of 50 min was chosen taking into consideration all the analytes.



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Figure 3. Evaluation of the effect of extraction times. Sample volume: 20 mL, salt content: 0% w/v NaCl, eluent: acetonitrile, stirring rate: 800 rpm, volume of eluent: 500 μ L, desorption time: 5 min.

As a final step, the effect of salt addition was studied by adding variable 257 concentrations of NaCl (i.e., 0-20% w/v). An increase in the extraction recovery was 258 259 observed for α -HCH, β -HCH, γ -HCH by enhancing the NaCl content up to 5% w/v (Figure S5). This phenomenon can be attributed to the salting-out effect, based on 260 261 which, the addition of salt results in a reduction of the analyte solubility in the sample solution, favoring its interaction with the sorptive phase. However, a further increase 262 from 5% w/v to 20% w/v had a negative impact on the extraction performance for o.p'-263 DDD, p,p'-DDD and methoxychlor, probably due to the reduced mass transfer of the 264 OCPs which can be attributed to the enhancement of sample density [21]. Thus, a NaCl 265 content of 5% w/v was chosen for further experiments. 266

3.2.3 Optimization of the elution step

The main parameters that influence the performance of the elution step (*i.e.*, 270 type/volume of eluent and elution time) were also investigated and optimized. Different 271 solvents were examined for their performance to elute the adsorbed analytes from the 272 CPME device. During method optimization, ACN was used as eluent, however its use 273 is not recommended from an environmental aspect [29]. For this purpose, MeOH and 274 ACE were also examined, since these solvents are "preferred" organic solvents 275 according to the Pfizer solvent selection guide [30]. Although the usage of MeOH 276 277 resulted in decreased elution efficiency, ACE exhibited similar performance as ACN (Figure S6). Thus, ACE was chosen as eluent taking into consideration the principles 278 of GAC regarding the utilization of more environmentally-friendly chemicals [14]. 279 Accordingly, the usage of different aliquots (i.e., 250 µL, 500 µL and 1000 µL) of ACE 280 was evaluated, aiming to reduce the quantity of chemicals used in this study. In 281 282 principle, it is desirable to use the lowest possible quantity of organic solvent to ensure low waste generation and low cost [14]. Meanwhile, the quantity of the solvent must be 283 284 enough for the complete analyte elution and for avoiding potential carry over effects. As shown in Figure S7, an aliquot of 250 µL of ACE was sufficient for the elution of 285 286 OCPs. Lower solvent amounts were not studied to ensure complete immersion of the capsule in the eluent and to ensure sufficient contact between the eluent and the sol-gel 287 sorbent. Thus, further experiments were carried out using this quantity of solvent. 288 Finally, different elution times were studied to ensure the development of a rapid 289 method with high sample throughput. A time span of 2 min was sufficient (Figure 4) 290 for the elution of the OCPs from the CPME device. 291



Figure 4. Evaluation of the effect of different elution times. Sample volume: 20 mL,
salt content: 5% *w/v*, extraction time: 50 min, stirring rate: 800 rpm, eluent: acetone,
volume of eluent: 500 μL.

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298 **3.3. Figures-of-merit**

In order to assess the linearity of the proposed methodology, spiked samples 299 300 were subjected to the optimum extraction protocol and linear regression analysis was performed by plotting the peak area of each analyte versus its respective concentration. 301 302 For each OCP pesticide, the determination coefficient, the slope, and the intercept of the regression lines were calculated. In Table 1, the regression analysis results are 303 presented. The coefficients of determination were 0.9939-0.9993 indicating good 304 method linearity. The lowest point of the calibration curve for each pesticide that had a 305 signal-to-noise ratio higher than 10 was considered to be the limit of quantification 306 (LOQ) and the limit of detection (LOD) was the concentration that corresponded to a 307 signal-to-noise ratio of 3 [31]. For the studied analytes, the LOD values were 0.01-0.03 308 ng mL⁻¹ and the LOQ values were 0.02-0.10 ng mL⁻¹. 309

Accordingly, the preconcentration factor (PF), the enhancement factor (EF) and the extraction recovery (ER%) were calculated [32]. PFs were calculated as ratio of the 312 sample volume (20 mL) compared to the eluent amount (250 µL). Thus, the theoretical PF for each analyte is 80. The EF values of each analyte were calculated as the ratio of 313 the slope derived from the calibration curve of the respective analyte prior and after the 314 CPME method. Finally, the ER% values were calculated by dividing the EF with the 315 theoretical PF *100. As shown in Table 1, the EF values for each analyte were in the 316 range 11.5-59.9. Finally, the ER% values were obtained by dividing the EF values by 317 the PF values and multiplying with 100. The ER% values of the OCPs were in the range 318 of 14.4-74.8% 319

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Table 1. Figures-of-merit for the proposed CPME GC-MS protocol

OCD	Degregation Analysia	D2	Linear range	LOD ¹	LOQ ²	ER ³	EE4	
UCP	Regression Analysis	K-	(ng mL ⁻¹)	(ng mL ⁻¹)	(ng mL ⁻¹)	(%)	EL.	
α-HCH	y = 4884.6x + 765.51	0.9993	0.05-50.0	0.02	0.05	74.8	59.9	
β-ΗCΗ	y = 3808.1x + 2789.5	0.9975	0.10-50.0	0.03	0.10	41.8	52.9	
γ-HCH	y = 3722.5x + 550.9	0.9991	0.10-50.0	0.03	0.10	70.7	59.4	
Alachlor	y = 14054x - 1809.6	0.9989	0.05-50.0	0.02	0.05	56.5	45.2	
Aldrin	y = 2887x + 2850.8	0.9939	0.05-50.0	0.01	0.02	18.4	14.7	
o,p'-DDE	y = 10197x + 1204.2	0.9986	0.02-20.0	0.01	0.02	20.8	16.6	
p,p'-DDE	y = 6404.8x + 1222	0.9970	0.02-20.0	0.01	0.02	14.4	11.5	
o,p'-DDD	y = 5885.3x + 3022.2	0.9949	0.02-50.0	0.01	0.02	38.6	30.9	
p,p'-DDD	y = 20404x + 5194.9	0.9973	0.02-50.0	0.01	0.02	53.5	42.8	
Methoxychlor	y = 33202x - 11634	0.9989	0.05-50.0	0.02	0.05	56.6	45.3	

322 ¹LOD: Limit of detection

323 2 LOQ: Limit of quantification

324 ³ER: Extraction recovery

325 ⁴EF: Enhancement factor

Method accuracy and method precision was investigated by analyzing spiked samples (*i.e.*, c=2.00 and 10.00 ng mL⁻¹). For the intra-day studies, five replicate measurements (n=5) of each spiked concentration level were conducted in the same day, while for the inter-day studies triplicate analysis of each spiked concentration level were performed on four consecutive days ($n=3 \times 4$) [33]. Method accuracy was expressed in terms of relative recovery (RR%) between the found and added concentration of each pesticide. Method precision was expressed in terms of relative standard deviation (RSD%). The results of the assessment of method trueness and method precision are summarized in Table 2. As it can be observed, the RR% values for intra-day study were between 90.5% and 105.2% and the RSD% values were less than 5.3% for all analytes. As for the inter-day study, the RR% values were 93.0% and 105.0% and the RSD% values were less than 9.2%, indicating good method trueness and precision.

Table 2. Intra-day (n=5) and inter-day ($n=4 \times 3$) performance studies of the CPME GC-MS method

	Added	Intra	-Day (n=5)		Inter-Day ($n=4 \times 3$)			
Analyte	(ng mL ⁻¹)	Found	RSD% ¹	DD 0/2	Found	RSD%	DD 0/	
	(ing int)	(ng mL ⁻¹)		KK 70 ⁻	(ng mL ⁻¹)		KK 70	
	2.00	1.99 ± 0.04	1.8	99.5	2.09 ± 0.14	6.7	104.5	
и-псп	10.00	9.74 ± 0.21	2.1	97.4	9.31 ± 0.56	6.0	93.1	
виси	2.00	1.95 ± 0.07	3.6	97.5	2.03 ± 0.08	3.9	101.5	
р-псп	10.00	9.78 ± 0.24	2.4	97.8	9.30 ± 0.45	4.8	93.0	
	2.00	2.03 ± 0.06	3.2	101.5	2.06 ± 0.09	4.2	103.0	
ү-псп	10.00	9.78 ± 0.30	3.1	97.8	9.36 ± 0.54	5.8	93.6	
Alashlar	2.00	1.96 ± 0.07	3.5	98.0	2.02 ± 0.11	5.3	101.1	
Alacilloi	10.00	9.66 ± 0.30	3.1	96.6	9.61 ± 0.35	3.6	96.1	
Aldrin	2.00	1.81 ± 0.06	3.3	90.5	1.95 ± 0.09	4.8	97.5	
Aluliii	10.00	9.88 ± 0.28	2.9	98.8	9.51 ± 0.58	6.1	95.1	
o n DDE	2.00	1.82 ± 0.08	4.5	91.0	1.94 ± 0.07	3.7	97.0	
0,р-DDE	10.00	9.85 ± 0.24	2.4	98.5	9.62 ± 0.47	4.9	96.2	
n n DDE	2.00	1.94 ± 0.08	4.2	97.0	2.10 ± 0.15	7.2	105.0	
р,р-DDE	10.00	10.11 ± 0.14	1.4	101.1	9.79 ± 0.42	4.3	97.9	
	2.00	1.99 ± 0.07	3.5	99.5	2.00 ± 0.13	6.3	100.0	
0,р-иии	10.00	10.00 ± 0.16	1.6	100.0	9.88 ± 0.67	6.7	98.8	
	2.00	2.03 ± 0.05	2.7	101.5	1.96 ± 0.14	7.4	98.0	
р,р-иии	10.00	9.97 ± 0.14	1.4	99.7	9.84 ± 0.32	3.2	98.4	
athowyahlar	2.00	2.10 ± 0.06	2.7	105.0	2.02 ± 0.06	3.1	101.0	
suloxychior	10.00	9.42 ± 0.50	5.3	94.2	9.84 ± 0.90	9.2	98.4	
	Analyte α-HCH β-HCH γ-HCH Alachlor ο,p-DDE ρ,p-DDE ο,p-DDD ρ,p-DDD εthoxychlor	AnalyteAdded (ng nL-1) α -BHCH2.00 β -HCH2.00 β -HCH2.00 γ -HCH2.00 γ -HCH2.00 β -DDE2.00 β -DDE3.00 β -DE3.00 β -DE3.00	AnalyeeAddeed (rg mL ⁻¹)AnalyeeFound(rg mL ⁻¹)(rg mL ⁻¹)α-HCH2.009.74 ± 0.21β-HCH2.009.74 ± 0.21β-HCH2.009.78 ± 0.07β-HCH2.009.78 ± 0.07β-HCH2.009.78 ± 0.07β-HCH2.009.78 ± 0.03β-HCH2.009.78 ± 0.03β-HCH10.009.66 ± 0.03β-HCH2.001.81 ± 0.06β-HCH2.001.81 ± 0.06β-DDE2.001.82 ± 0.08β-PDDE1.0009.85 ± 0.24β-PDDE2.001.91 ± 0.14β-PDDE2.001.91 ± 0.14β-PDDE2.002.03 ± 0.05β-PDDE2.009.97 ± 0.14β-PDDE2.002.10 ± 0.06β-PDDE2.002.10 ± 0.06β-PDDE2.002.10 ± 0.06β-PDDE2.002.10 ± 0.06β-PDDE2.002.10 ± 0.06β-PDDE2.003.97 ± 0.14β-PDDE2.003.97 ± 0.14β-PDDE2.003.97 ± 0.14β-PODE2.003.97 ± 0.14β-PODE2.003.97 ± 0.14β-PODE2.003.97 ± 0.14β-PODE2.003.91 ± 0.06β-PODE3.003.91 ± 0.06β-PODE3.003.91 ± 0.06β-PODE3.003.91 ± 0.06β-PODE3.003.91 ± 0.06β-PODE3.91 ± 0.063.91 ± 0.06β-PODE	AnalyaAdded (ag mL1)FoundRSD%1FoundFoundRSD%1α-HCH2.001.99±0.041.8β-HCH2.009.74±0.212.1β-HCH2.009.78±0.073.6β-HCH2.009.78±0.242.4β-HCH2.009.78±0.243.1β-HCH2.009.78±0.303.1β-HCH2.001.96±0.073.5β-HCH1.009.66±0.303.1β-HCH1.009.88±0.283.1β-HCH1.009.88±0.283.1β-HCH1.009.88±0.283.1β-HCH1.009.85±0.243.3β-HCH1.009.85±0.243.5β-PDDE1.001.94±0.084.2β-PDDD1.001.01±0.141.4β-PDDD2.001.99±0.073.5β-PDDD2.002.03±0.052.7β-PDDD2.009.97±0.141.4β-PDDD3.009.97±0.141.4β-PDDD2.009.97±0.141.4β-PDDD3.009.97±0.141.4β-PODD3.003.13.1β-PODD2.003.13.1β-PODD3.003.13.1β-PODD3.003.13.1β-PODD3.003.13.1β-PODD3.003.13.1β-PODD3.003.13.1β-PODD3.003.13.1β-PODD3.	AnalyseAddea (ag mL1)FoundRSD%1FoundRSD	Analya (nmm)Intra-Jum (nmm)Intra-Jum (nmm)FoundFoundRPMFound(nmm)(nmm)(nmm)(nmm)\alpha-CH2.001.99±0.041.899.52.09±0.14\alpha-CH10.009.74±0.212.197.49.31±0.56\beta-CH1.95±0.073.697.52.03±0.08\beta-CH9.78±0.242.497.89.30±0.45\beta-CH2.03±0.063.2101.52.06±0.09\beta-CH9.78±0.203.197.80.36±0.54\beta-CH1.96±0.073.598.02.02±0.11\beta-CH1.96±0.073.59.61±0.350.61±0.35\beta-CH1.96±0.073.19.65±0.249.65±0.249.65±0.24\beta-CH1.81±0.063.390.51.95±0.07\beta-CH1.82±0.84.59.101.94±0.03\beta-CH1.94±0.084.29.101.94±0.14\beta-CH1.94±0.084.29.011.94±0.14\beta-CH1.94±0.084.29.011.94±0.14\beta-CH1.94±0.031.64101.19.79±0.42\beta-CH1.94±0.031.641.019.79±0.42\beta-CH1.94±0.031.641.019.84±0.32\beta-CH1.94±0.031.641.019.84±0.32\beta-CH1.94±0.031.641.019.84±0.32\beta-CH1.94±0.031.641.019.84±0.32\beta-CH <td< td=""><td>AnalyceAdded (ng mL')Intra-Jur (n=5)Intra-Jur (n=4 × 3)RanalyceFoundRSD%1Reffer (ng mL')FoundRSD%1α-HCH2.001.99 ± 0.041.89.052.09 ± 0.146.7β-HCH10.009.74 ± 0.212.1197.49.31 ± 0.566.0β-HCH2.001.95 ± 0.073.697.52.03 ± 0.083.9β-HCH2.009.78 ± 0.242.4497.89.30 ± 0.454.8γ-HCH2.009.78 ± 0.303.197.89.36 ± 0.545.8β-HCH1.0009.78 ± 0.303.197.89.36 ± 0.545.8β-HCH1.0009.66 ± 0.303.196.69.61 ± 0.353.6β-HCH1.0009.66 ± 0.303.196.69.61 ± 0.353.6β-HCH1.0009.88 ± 0.282.998.89.51 ± 0.586.1β-HCH1.0009.88 ± 0.282.99.89.51 ± 0.586.1β-PDDE2.001.81 ± 0.063.390.51.95 ± 0.094.8β-PDDE2.001.94 ± 0.084.297.02.10 ± 0.157.2β-PDDE2.001.91 ± 0.141.44101.19.79 ± 0.424.3β-PDDD2.001.99 ± 0.073.59.52.00 ± 0.136.1β-PDDD2.001.99 ± 0.151.96 ± 0.147.47.4β-PDDD2.002.01 ± 0.062.7101.51.96 ± 0.147.4β-PDDD<t< td=""></t<></td></td<>	AnalyceAdded (ng mL')Intra-Jur (n=5)Intra-Jur (n=4 × 3)RanalyceFoundRSD%1Reffer (ng mL')FoundRSD%1α-HCH2.001.99 ± 0.041.89.052.09 ± 0.146.7β-HCH10.009.74 ± 0.212.1197.49.31 ± 0.566.0β-HCH2.001.95 ± 0.073.697.52.03 ± 0.083.9β-HCH2.009.78 ± 0.242.4497.89.30 ± 0.454.8γ-HCH2.009.78 ± 0.303.197.89.36 ± 0.545.8β-HCH1.0009.78 ± 0.303.197.89.36 ± 0.545.8β-HCH1.0009.66 ± 0.303.196.69.61 ± 0.353.6β-HCH1.0009.66 ± 0.303.196.69.61 ± 0.353.6β-HCH1.0009.88 ± 0.282.998.89.51 ± 0.586.1β-HCH1.0009.88 ± 0.282.99.89.51 ± 0.586.1β-PDDE2.001.81 ± 0.063.390.51.95 ± 0.094.8β-PDDE2.001.94 ± 0.084.297.02.10 ± 0.157.2β-PDDE2.001.91 ± 0.141.44101.19.79 ± 0.424.3β-PDDD2.001.99 ± 0.073.59.52.00 ± 0.136.1β-PDDD2.001.99 ± 0.151.96 ± 0.147.47.4β-PDDD2.002.01 ± 0.062.7101.51.96 ± 0.147.4β-PDDD <t< td=""></t<>	

- ¹RSD: Relative standard deviation
- 344 2 RR: Relative recovery
- 345
- 346

347 3.4. Reusability of the sol-gel PEG 300 CPME media

In a further step, the reusability of the sol-gel PEG 300 CPME media was 348 349 studied to provide a more comprehensive assessment regarding the performance of the proposed method. In green sample preparation, the utilization of reusable materials over 350 351 those of disposable nature is of high importance to promote the reduction of waste aiming to develop more environmentally-friendly and low-cost methods [34]. For this 352 study, one capsule was used for 20 repeated extraction cycles using a spiked sample 353 (c= 10 ng mL⁻¹) and the criterion of a reduction of \geq 10% of the recovery compared to 354 the initial recovery was set. As shown in Figure S8, the performance of the CPME 355 device was unaffected after 20 consecutive extraction cycles. Thus, the capsules are 356 357 reusable for at least 20 times.

358

359 **3.5.** Evaluation of method's green character and comparison with other 360 approaches

The herein developed method was compared with previously reported methodsfor the extraction of OCPs, as shown in Table 3.

			time(min)	Liuciit	reconstitution	KDD %0°	(ng mL ⁻¹)	Ref.
GC-MS	15	No	120/15	1.5 mL of toluene: acetonitrile (20:80 v/v)	Required	<14.8	0.02-0.38	[18]
GC-MS	100	No	-	12 mL of dichloromethane	Required	<6.69	0.007- 0.126	[35]
GC-ECD	20	No	10/15	5 mL of ethyl acetate	Required	<5.6	0.025- 0.049	[36]
GC-ECD	10	Required	40/7	dichloromethane	Required	<9.88	0.0076- 0.10	[37]
GC-MS	5	Required	20/5	4 mL of acetonitrile	Required	<8.5	0.07-1.03	[38]
GC-µECD	10	Required	10/2	250 μL of acetonitrile	No	<7.3	0.4-4.1 x 10 ⁻³	[39]
GC-MS	20	No	50/2	250 μ L of acetone	No	<5.3 (intra-day) <9.2 (inter-day)	0.01-0.03	This study
	GC-MS GC-ECD GC-ECD GC-MS GC-µECD GC-MS	GC-MS 15 GC-MS 100 GC-ECD 20 GC-ECD 10 GC-MS 5 GC-µECD 10 GC-MS 20	GC-MS 15 No GC-MS 100 No GC-ECD 20 No GC-ECD 10 Required GC-MS 5 Required GC-MS 10 Required GC-MS 20 No	GC-MS 15 No 120/15 GC-MS 100 No - GC-ECD 20 No 10/15 GC-ECD 10 Required 40/7 GC-MS 5 Required 20/5 GC-µECD 10 Required 10/2 GC-MS 20 No 50/2	GC-MS15No120/15acetonitrile (20:80 v/v)GC-MS100No-12 mL of dichloromethaneGC-ECD20No10/155 mL of ethyl acetateGC-ECD10Required40/7dichloromethaneGC-MS5Required20/54 mL of acetonitrileGC- μ ECD10Required10/2250 μ L of acetonitrileGC-MS20No50/2250 μ L of acetone	GC-MS15No120/15acetonitrile (20:80 v/v)RequiredGC-MS100No-12 mL of dichloromethaneRequiredGC-ECD20No10/155 mL of ethyl acetateRequiredGC-ECD10Required40/7dichloromethaneRequiredGC-MS5Required20/54 mL of acetonitrileRequiredGC-MS10Required10/2250 µL of acetonitrileNoGC-MS20No50/2250 µL of acetoneNo	GC-MS15No120/15acetonitrile (20:80 v/v)Required<14.8GC-MS100No- $\begin{array}{c} 12 \text{mL of} \\ dichloromethane \end{array}$ Required<6.69	GC-MS 15 No 120/15 acetonitrile (20:80 v/v) Required <14.8 0.02-0.38 GC-MS 100 No - 12 mL of dichloromethane Required <6.69 0.007-

Table 3. Comparison of the proposed method with other methodologies.

366	¹ HS-SBSE: Headspace-stir bar sorptive extraction, DMIP: Dummy template molecularly imprinted polymer, SWCNTs: single-walled
367	carbon nanotubes, M-M-ZIF-67: zeolitic imidazolate framework based on magnetic multi-walled carbon nanotubes, RGO/Fe ₃ O ₄ @Au:
368	reduced graphene oxide/ Fe ₃ O ₄ @gold nanocomposite,
369	² GC-ECD: Gas chromatography-electron capture detector, GC-µECD: Gas chromatography- micro electron capture detector
370	³ RSD: Relative standard deviation
371	⁴ LODs: Limits of detection

Ś

The sample amount used during the extraction procedure was higher than the 373 sample amount used in refs. [18,37–39], similar to the sample amount used in ref. [36] 374 and lower than the sample amount used in ref. [35]. The overall extraction time was 375 comparable with the extraction time required in ref. [37], higher than the extraction 376 time required in refs. [36,38,39] but lower than the extraction time required in ref. [18]. 377 378 Moreover, the sensitivity of the proposed method (in terms of LOD values) was comparable to those of refs. [18,35–37], higher than the sensitivity of ref. [38] but lower 379 380 than the sensitivity of ref. [39].

381 A significant advantage of the proposed procedure is the utilization of a small 382 amount of organic solvent as eluent. In this work, elution is performed using 250 μ L of acetone, while the organic solvent requirements in most of the other studies are above 383 1.5 mL (i.e., refs. [18,36,38] and they range up to 12 mL (i.e., ref. [35]). Only in ref. 384 [39] the same amount (*i.e.*, 250 µL) of organic solvent is employed; however 385 386 acetonitrile is used which is not recommendable from an environmental perspective [29]. On the other hand, acetone is considered to be a "preferred" solvent, as already 387 388 discussed in section 3.2.3 [30]. The proposed method does not require acetonitrile (used in ref. [18] and [38]), toluene (used in ref. [18]) and chlorinated solvents (used in refs. 389 390 [35] and [37]) which are more hazardous chemicals. Thus, the proposed method meets the requirements of GAC [14] regarding the low consumption of organic solvents and 391 392 the replacement of chemicals with less hazardous ones.

Another advantage of CPME is that it overcomes the need for sample filtration 393 prior to the extraction procedure and the need for sample evaporation following the 394 extraction procedure. Sample filtration was required in refs. [37–39], while sample 395 evaporation and reconstitution were required in refs. [18,35–38]. In principle, a multi-396 step sample preparation procedure may result in reduced precision and accuracy, while 397 it can be time-consuming and demand high expenditures of chemicals and energy [34]. 398 Thus, the reduction of sample preparation steps is a significant factor towards the 399 development of greener methods. An additional benefit of CPME is the increased 400 simplicity of the method, because the microextraction capsules can be removed from 401 the eluent and the sample solution using tweezers. Thus, they overcome the need of 402 magnetic separation that is necessary in MSPE [38,39] and centrifugation that is 403 necessary in d-SPE [36] processes. 404

405 A usefool tool to make the greenness of an analytical method visible and 406 comparable are ComplesGAPI pictograms [40]. In this tool, the environmental 407 friendliness of an analytical method is assessed by evaluating the sample preparation procedure, the instrumentation used for the analytical determination, the reagents and 408 chemicals used and the overall method type. Moreover, the hexagonal field of the 409 ComplexGAPI pictogram examines the impact of the yield and conditions, chemicals, 410 instrumentation purification and workup used during the manufacturing of extraction 411 materials which in this case includes the CPME device. Figure 5 depicts the 412 ComplexGAPI pictogram that corresponds to the herein developed CPME GC-MS 413 method, as well as to two of the other existing methodologies for the monitoring of 414 415 OPCs in water samples found in the literature. With regard to the preparation of the microextraction capsules most of the assessment criteria are met (green colour). The 416 synthesis was characterized by a high process yield and reduced waste generation, as 417 well as a low E-factor. As for the extraction, the proposed scheme is characterized by 418 low chemical consumption and waste generation since microextraction is used. Future 419 420 recommendations towards the reduction of the environmental impact of the herein developed method include the utilization of more environmentally-friendly chemicals 421 422 (*i.e.*, deep eutectic solvents, DESs) instead of conventional organic solvents.

DMIP-SPE-GC-MS [35] d-SPE-GC-MS [36] CPME-GC-MS

423

Figure 5. ComplexGAPI pictogram of the developed method (right), compared to other
selected methods (left and middle) [35, 36]

426

429 **3.6.** Analysis of real-world water samples

Following method development and measurement of the figures-of-merit of the 430 proposed method, water samples of different origin (*i.e.*, tap, river, pond and lake water) 431 were analysed. Each sample was spiked at two different concentration levels (i.e., 432 $c=2.00 \text{ ng mL}^{-1}$ and 10.00 ng mL⁻¹) to investigate the applicability of the proposed 433 methodology to different water samples. As shown in Table 4, the relative recoveries 434 435 in the examined spiked levels ranged between 80.1-112.5% indicating good method applicability of the proposed scheme in different environmental water samples. The 436 437 absence of interferences in the blank samples shows that the proposed method is characterized by specificity, while no contamination occurred during sample analysis. 438

	AddedLake water 1La		Lake wat	Lake water 2Pond water			River v	vater	Tap water		
Analyte	(ng	Found		Found	RR%	Found	RR%	Found	RR%	Found	RR%
	mL ⁻¹)	(ng mL ⁻¹)		(ng mL ⁻¹)	1117,0	(ng mL ⁻¹)	1111/0	(ng mL ⁻¹)		(ng mL ⁻¹)	
a-HCH	0	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<>	-	<lod< td=""><td>-</td></lod<>	-
u-mem	2.00	2.01±0.04	100.3	1.82 ± 0.04	91.0	1.71 ± 0.01	85.5	2.22±0.09	111.0	1.98 ± 0.02	99.1
	10.00	10.16±0.06	101.6	9.57±0.18	95.7	10.22 ± 0.38	102.2	9.89±0.16	98.9	9.92±0.18	99.2
	0	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<>	-	<lod< td=""><td>-</td></lod<>	-
β-НСН	2.00	1.90 ± 0.02	95.2	1.80 ± 0.03	90.2	1.75 ± 0.02	87.4	1.77±0.03	88.6	1.92 ± 0.02	96.1
2	10.00	10.52±0.12	105.2	9.28±0.34	92.8	10.38 ± 0.22	103.8	9.97±0.22	99.7	10.12±0.25	101.2
	0	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<>	-	<lod< td=""><td>-</td></lod<>	-
ү-НСН	2.00	1.97±0.02	98.3	1.85 ± 0.07	92.4	2.06 ± 0.02	103.0	2.21±0.04	110.5	1.91 ± 0.04	95.4
)	10.00	9.71±0.09	97.1	9.25±0.07	92.3	9.77±0.28	97.7	9.88±0.17	98.8	9.23±0.27	92.3
- 5 5	0	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<>	-	<lod< td=""><td>-</td></lod<>	-
Alachlor	2.00	1.64±0.05	82.0	1.96±0.05	98.1	1.95 ± 0.05	97.4	2.03±0.01	101.3	2.01±0.03	100.5
)	10.00	9.74±0.44	97.4	9.14±0.28	91.4	9.89±0.30	98.9	9.58±0.60	95.8	10.73±0.24	107.3
	0	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<>	-	<lod< td=""><td>-</td></lod<>	-
Aldrin	2.00	1.62 ± 0.02	81.2	2.01±0.01	100.6	1.93±0.06	96.4	1.97 ± 0.02	98.6	1.98 ± 0.03	99.0
	10.00	10.51±0.04	105.1	10.25±0.15	102.5	8.90±0.37	89.0	9.65±0.01	96.5	10.25±0.27	102.5

440 **Table 4.** Analysis of environmental water samples through CPME GC-MS.

Added		Lake water 1		Lake water 2		Pond water	Pond water		River water		water
Analyte	(ng	Found	RR% ¹	Found	RR%	Found	RR%	Found	RR%	Found	RR%
	mL ⁻¹)	(ng mL ⁻¹)		(ng mL ⁻¹)		(ng mL ⁻¹)		$(ng mL^{-1})$		(ng mL ⁻¹)	
	0	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<>	-	<lod< td=""><td>-</td></lod<>	-
o,p-DDE	2.00	1.72 ± 0.01	86.0	1.91±0.06	95.5	$1.60{\pm}0.01$	80.1	1.76±0.06	87.8	2.10±0.11	105.1
	10.00	10.98 ± 0.05	109.8	9.95±0.07	99.5	10.05 ± 0.19	100.5	9.51±0.13	95.1	10.25±0.27	102.5
p,p-DDE	0	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<>	-	<lod< td=""><td>-</td></lod<>	-
	2.00	1.97 ± 0.01	98.7	1.75 ± 0.05	87.6	$1.70{\pm}0.07$	85.2	1.74 ± 0.01	87.1	2.15±0.05	107.3
	10.00	9.53±0.14	95.3	9.68±0.17	96.8	8.87±0.45	88.7	9.25±0.20	92.5	10.87±0.15	108.7
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o,p-DDD	2.00	1.85 ± 0.01	92.7	2.11±0.03	105.3	2.00±0.09	99.8	2.06±0.02	102.8	2.07 ± 0.05	103.4
	10.00	10.34±0.15	103.4	11.17±0.16	111.7	8.91±0.20	89.1	8.51±0.07	85.1	11.25±0.16	112.5
EO	0	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<>	-	<lod< td=""><td>-</td></lod<>	-
p,p-DDD	2.00	1.81 ± 0.04	90.4	1.93 ± 0.02	96.4	1.81 ± 0.10	90.6	1.94 ± 0.03	97.2	1.98 ± 0.06	98.8
	10.00	8.79±0.02	87.9	8.24±0.30	82.4	8.04±0.02	80.4	8.61±0.01	86.1	10.72±0.16	107.2
thoxychlor	0	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<></td></lod<>	-	<lod< td=""><td>-</td><td><lod< td=""><td>-</td></lod<></td></lod<>	-	<lod< td=""><td>-</td></lod<>	-
	2.00	$1.70\pm0.0.03$	85.1	1.82 ± 0.03	90.9	$1.74{\pm}0.07$	87.1	1.93±0.05	96.5	1.96 ± 0.02	98.2
	10.00	9.15±0.90	91.5	8.43±0.82	84.3	8.89±1.05	88.9	8.79±0.10	87.9	9.94±0.81	99.4

441 ¹RR: Relative recovery

442 **4.** Conclusions

In this work, CPME was used for the extraction of OCPs in environmental water 443 samples. Among the examined sorbents, the monolithic sol-gel poly(ethylene glycol)-444 based CPME device resulted in the highest extraction efficiency. Under optimum 445 sample preparation conditions, the proposed method showed good linearity, accuracy, 446 precision, and sensitivity. Moreover, the capsules could be reused for at least 20 times. 447 The proposed scheme exhibits multiple benefits including handling simplicity, rapid 448 extraction kinetics, and low consumption of organic solvents. CPME efficiently 449 450 overcomes the need for sample pretreatment (*i.e.*, filtration) prior to sample preparation, 451 while it also reduces the need for sample manipulation (e.g., evaporation/reconstitution) after the sample preparation. Moreover, the integration of stirring mechanism results in 452 453 a less error prone and more powerful extraction device, that prevents potential loss of analytes due to their retention on external devices (e.g., magnetic rods). 454

455

456 Declaration of Competing Interest

- 457 The authors declare no conflict of interest
- 458
- 459 **References**
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Monolithic capsule phase microextraction prior to gas chromatography-mass spectrometry for the determination of organochlorine pesticides in environmental water samples

Antonio Ferracane^{a,b}, Natalia Manousi^{b,c*}, Abuzar Kabir^d, Kenneth G. Furton^d, Peter Q. Tranchida^a, George A. Zachariadis^c, Justyna Płotka-Wasylka^e, Luigi Mondello^{a,f}, Victoria F. Samanidou^c, Erwin Rosenberg^b

^a Department of Chemical, Biological, Pharmaceutical and Environmental Sciences, University of Messina, Messina, Italy

^b Institute of Chemical Technologies and Analytics, Vienna University of Technology, 1060 Vienna, Austria

^c Laboratory of Analytical Chemistry, Department of Chemistry, Aristotle University of Thessaloniki, Thessaloniki 54124, Greece

^d Department of Chemistry and Biochemistry, Florida International University, Miami, FL, USA

^e Department of Analytical Chemistry, Faculty of Chemistry and BioTechMed Center, Gdansk University of Technology, 1/12 G. Narutowicza St., 80-233 Gdansk, Poland ^f Chromaleont s.r.l., c/o Department of Chemical, Biological, Pharmaceutical and Environmental Sciences, University of Messina, Messina, Italy

*Corresponding author: nmanousi@chem.auth.gr; Laboratory of Analytical Chemistry, Department of Chemistry, Aristotle University of Thessaloniki, Thessaloniki 54124, Greece

Preparation of the capsule phase microextraction devices

First, Accurel® polypropylene S6/2 porous membranes were cut into 3 cm pieces. The porous capillary membranes were then cleaned with methylene chloride for 30 min under sonication and subsequently, air dried at room temperature for 30 min. A cylindrical magnet (3/4" x 1/16") was used in each of the CPME device. The magnet was inserted into an empty polypropylene capsule. Subsequently, one empty capsule and a capsule containing the magnet were fused together by the ends using an impulse heat-sealing machine. As a result, both the capsules were connected to each other by their ends. The CPME devices were then ready for the sol-gel sorbent coating process.

Sol solutions for in situ creation of sol-gel PDMS/sol-gel PEG 300/ sol-gel C₁₈/sol-gel PDMDPS monolithic sorbent bed within the lumen of the empty propylene capsule were prepared by the sequential addition of tetramethyl orthosilicate (TMOS), methyl trimethoxysilane (MTMS), polymer, isopropanol, hydrochloric acid catalyst and deionized water at a molar ratio: 1: 1: 0.2: 30: 0.04: 8, respectively in a 50mL amber glass reaction container. The mixture was kept at room temperature for 12 h so that acidic hydrolysis of the sol-gel precursors moves towards completion. Subsequently, the sol solution was centrifuged and the supernant particle free fraction was transferred into a wide-mouth glass reaction vessel. Successively, NH₄OH (1 M) was added to the solution in droplets at a molar ratio between TMOS and NH₄OH at 1: 0.10 with continuous stirring to achieve a homogeneous mixture. The process allows creation of batch of 30 units of CPME devices at once that can be even expanded by extending the size of the reaction vessel. A batch of 30 CPME devices were submerged into the sol solution and then at the reaction vessel containing the submerged CPME devices was sonicated for 5 min to remove air bubbles from the system. The gelation of the sol solution begins with the addition of the base catalyst and the sol solution turns into solid gel in 1 h at room temperature. The sol solution formed a solid monolithic bed within the lumen of the capsules and a mesh-like network on the surface of the porous polypropylene capsules and inside the pores of its thick walls. The CPME devices were then subjected to aging and thermal conditioning at 50°C for 24 h. The CPME devices were subsequently cleaned by scrubbing the sol-gel sorbent from their outer surface and rinsing with a mixture of methanol: methylene chloride (50:50 v/v) under sonication for 30 min. The monolithic bed of the sol-gel sorbent was disintegrated into fine microparticles by

sonication. The CPME devices were then dried in an oven at 50°C. Finally, the CPME devices were ready for the analyte extraction.



Figure S1. Chemical structures of the target analytes



Figure S2. Evaluation of the effect of different monolithic sol-gel coated microextraction capsules. Sample volume: 20 mL, extraction time: 30 min, stirring rate: 800 rpm, salt content: 0% w/v NaCl, eluent: methanol, volume of eluent: 1000 μ L, desorption time: 5 min.



Figure S3. Evaluation of the effects of different stirring rates on extraction recovery. Sample volume: 20 mL, extraction time: 30 min, salt content: 0% w/v NaCl, eluent: acetonitrile, volume of eluent: 500 µL, desorption time: 5 min.



Figure S4. Evaluation of the effect of different sample volumes on extraction recovery. Extraction time: 30 min, salt content: 0% w/v NaCl, eluent: acetonitrile, stirring rate: 800 rpm, volume of eluent: 500 µL, desorption time: 5 min.



Figure S5. Evaluation of the effect of salt content on extraction recovery. Sample volume: 20 mL, extraction time: 50 min, eluent: acetonitrile, stirring rate: 800 rpm, volume of eluent: 500 μ L, desorption time: 5 min.



Figure S6. Evaluation of the effect of different eluents. Sample volume: 20 mL, salt content: 5% w/v, extraction time: 50 min, stirring rate: 800 rpm, volume of eluent: 500 μ L, desorption time: 5 min.



Figure S7. Evaluation of the effect of eluent volume on extraction yield. Sample volume: 20 mL, salt content: 5% w/v, extraction time: 50 min, stirring rate: 800 rpm, eluent: acetone, desorption time: 5 min.



Figure S8. Results of the reusability study of sol-gel PEG 300 microextraction capsules

Analyta	Retention time	Target Ion	Reference Ions
Analyte	(min)	(m / z)	(m/z)
	10.45	191	109
u-mem	10.45	101	219
β-ΗCΗ	11.00	191	109
	11.00	101	219
ү-НСН	11.80	181	109
	11.00	101	219
Alachlor	12.46 160	160	146
Alacinoi		100	188
Aldrin	13.46	66	91
	15.16	00	263
o n'-DDF	15.02	246	318
o,p DDL	15.02	240	176
n n'-DDF	15.87	246	318
p,p DDL	15.07	210	176
o n'-DDD	16 14	235	165
o,p DDD	10.11	200	199
n n'-DDD	17 11	235	165
p,p DDD	17.11	255	199
Methoxychlor	19 98	227	153
	17.70	,	212

Table S1. Retention times and m/z values used for the OCPs.

Variable In		itial value	Interval studied	Optimum/selected value
Adsorption step				
Sample amount (mL)		20	10-50	20
Stirring rate (rpm)		800	0-1200	800
Extraction time (min)		30	10-60	50
Salt content (% w/v)		0	0-20	5
Elution step				
Eluent ¹		ACN	ACN, ACE, MeO	H ACE
Volume of eluent (µL)		500	250-1000	250
Elution time (min)		5	2-20	2
¹ ACN:	Acetonitrile,	ACE:	Acetone,	MeOH: Methanol

Table S2. Initial parameters, interval studies and optimum/selected values for the CPME

 method using monolithic sol-gel PEG 300 microextraction capsules.

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