Co-occurrence of polychlorinated biphenyls, cyanotoxins and trace elements in commercial fish species from a freshwater protected area (Pertusillo Lake,

Southern Italy).

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4 Abstract

A total of 79 fish samples covering nine species were collected in a preliminary investigation on a SCI (Site 5 6 of Community Importance) water reservoir (Pertusillo Lake, Southern Italy) created for drinking purpose and located in a territory used for drilling activities. Analyses for microcystins (MYCs) and cylindrospermopsins 7 8 (CYLs) presence were performed using Elisa assays, while 10 fish samples were analyzed also for trace 9 elements by atomic adsorption spectrophotometry and for polychlorinated biphenyls (PCBs) by GC-MS 10 operated in EI mode. The results showed the compresence of important cyanotoxins and industrial 11 contaminants in fish. More extended studies are needed to evaluate the combined effects of these contaminants on the lake ecosystem and ichthyic fauna, in order to establish an available risk assessment for 12 13 human population in the lake region.

Keywords: microcystins, cylindrospermopsins, trace elements, polychlorinated biphenyls, fish,
bioaccumulation, Pertusillo Lake.

16 Introduction

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In the past century, the development of industry and agriculture often caused the release or production of organic and inorganic pollutants in the environment, posing threats to wildlife and human health. Several studies have shown the presence of anthropic contaminants in inland waters of various continents, as found in lakes polluted by industries (Yu et al., 2012; Hao et al., 2013; Jia et al., 2014). A particular class of these contaminants, microcystins (MYCs), the commonest biotoxins of Cyanobacteria, are a family of more than 90 potent eptapeptide hepatotoxins (Teixeira

et al 1993, Codd 1995, Jochimsen et al 1998, Gacsi et al 2009) acting as specific inhibitors of 24 protein phosphatases (PPs) of type 1, 2A, 3 (for MC-LA; Prinsep et al 1992), 4 and 5 (Hastie et al 25 2005), and to a lesser extent of type 2B (Mackintosh et al 1990). The inhibition of PP1 and PP2A 26 results in an increased phosphorylation of proteins in liver cells, affecting several cellular processes 27 (Dawson, 1998). MYCs are responsible for liver failure and death in humans (Falconer et al., 1983; 28 29 Azevedo et al., 2002; Crux et al., 1993), wild animals, livestock and aquatic life (Sivonen and 30 Jones, 1999; Mwaura et al., 2004). Indirect evidence supporting tumour promotion of human cancer from MYCs exposure comes from the studies of Yu (1989), Ueno et al. (1996) and Zhou et al. 31 (2002) in China, Fleming et al. (2002) in Florida, and Svircev et al. (2009) in Serbia. They can 32 induce oxidative DNA damage (Zegura et al., 2003), genotoxicity (Bouaicha et al., 2005) and cause 33 the activation of proto-oncogenes c-jun, c-fos and c-myc (Li et al., 2009). In addition, MYCs from 34 contaminated lakes can percolate and contaminate groundwater proportionally to the duration of 35 toxic bloom events (Eynard et al 2000, Messineo et al 2006). Their association with primary 36 carcinogens in the aquatic environment is a problematic event. Several large scale fish death 37 38 outbreaks have been associated to massive occurrence of Cyanobacteria in waterbodies (Jewel et 39 al., 2003; Zimba et al., 2006). MYCs concentrations between 0.34 µg/kg (Magalhaes et al., 2001) 40 and 36.42 µg/kg (Bruno et al., 2009) were measured in the muscle tissue of wild or farmed fish, indicating that even the consumption of contaminated fish muscle might constitute a threat for 41 42 human health. Cylindrospermopsin (CYN), another common cyanotoxin, is, is a sulfatedguanidinium alkaloid with hepatotoxic, nephrotoxic and thymotoxic effects (Terao et al., 1994; 43 Banker et al., 1997). CYN has in vitro and in vivo mutagenic, endocrine-disrupting and 44 carcinogenic activity (Shaw et al., 2000; Shen et al., 2002; Bain et al., 2007; Young et al., 2008; 45 Zegura et al., 2011), showing neurotoxic activity in fish (Guzman-Guillen et al., 2015). Aside from 46 microcystins, other toxic substances of major concern contaminating the environment are toxic 47 48 metals, namely mercury (Hg), cadmium (Cd) and lead (Pb), and organic contaminants, including

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49	polychlorinated biphenyls (PCBs). As a consequence of their environmental persistence and
50	potential for bioaccumulation, these chemicals are widespread throughout the ecosystem, causing
51	toxic problems to all life forms. Fish, in particular, have the ability to accumulate these
52	contaminants and, often, have been employed to assess environmental contamination (Tekin-Ozan
53	and Kir, 2008). More attention should be devoted to contaminant levels in fish, especially when
54	significant alterations in industrial development can result in large pollutant releases into the
55	environment. Common carp is a good species for bioaccumulation monitoring, being bottom feeder
56	fish that do not migrate extensively, reproduce rapidly and have long life spans (up to 38 yrs.)
57	(Pérez-Fuentetaja et al., 2010).
58	Being fish an important food source and a major part of many natural food chains, the objective of
59	the present study was to investigate the presence of these specific contaminants in fish from lake
60	Pertusillo, an extended Italian reservoir part of a national park, which neighbourhood is interested
61	by intense drilling activities often accused of causing serious water and sediment pollution in the
62	lake.
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64	Materials and methods
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66	Site description
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68	Lake Pertusillo is an artificial reservoir of the Italian region Basilicata, located atlocated at the
69	conjunction of the three municipal lands of Grumento Nova, Montemurro and Spinoso towns
70	(fig.1). Created between 1957 and 1962 by damming the River Agri, its surface area is 7.5 km ² and
71	its depth reaches 90 m. The mean renewal time is six months (Calderoni and Mosello, 1978). Thick
72	and beautiful woods surround it, covering its shores; the lake is a Site of Community Importance
73	(SCI) for the preservation of natural habitats (European Commission Habitats Directive 92/43/EEC)

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and a Special Protection Zone (SPZ) (European Union Directive on the Conservation of Wild Bird 74 Directive 79/409/EEC). As part of the National Park of Val d'Agri the lake is used for angling and 75 rowing, and its waters are used for drinking and irrigation purposes by the Basilicata and Apulia 76 Regions. Lake Pertusillo is about eight kilometers distant from a center of petroleum refining and in 77 2016, during an incident, 400 oil tons were spilled from this center in the site groundwater. In 78 79 2010, 2011, 2012, 2013, 2014 and 2015 fish deaths occurred in the lake, which cause was not found. In 2010, 2011, 2012 and 2017 huge dinoflagellate blooms, covering the lake surface, 80 81 occurred in spring and winter.

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83 Sample collection

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Samplings from June, 2010 to March, 2013 and in May, 2016 and April, 2017 were carried out in 85 six stations (Rifreddo, R; Madonna Grumentina, MG; Spinoso, S; Montemurro Bridge, MB; Lake 86 Damming, LD; Masseria Crisci; MC) of the lake. Seventy nine adult fish covering 10 species and 87 88 thirty water samples were analyzed. The analyzed fish species were the zoobenthivorous species Cyprinus carpio (carp, 30 individuals), Carassius carassius (crucian carp, 10 individuals) and 89 Cyprinus carpio specularis (mirror carp, 2 individual), the carnivorous species Lepomis gibbosus 90 (pumpkinseed, 2 individuals), Perca fluviatilis (perch, 9 individuals), Scardinius erythrophtalmus 91 (rudd, 1 individual), Ictalurus melas (catfish, 1 individual), Alburnus alborella (bleak, 9 92 individuals), Squalius cephalus (chub, 5 individuals) and Micropterus salmoides (black bass, 10 93 individuals). Fish captured by angling were ice-stored and transported to the laboratory. Thirty 94 surface water samples were collected in 20 samplings by filling 1 L Pyrex glass bottles 10-20 cm 95 below the water surface from two stations (S shoreS shore and R shoreR shore). 96

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98 Fish tissue cylindrospermopsin (CYN) extraction

Cylindrospermopsin extraction from tissue samples was performed according to Saker et al. (1999, 100 2004) mod.: tissue (5 g, muscle) was homogenized in 10 ml mL 100% MeOH for 15 min. using a 101 Potter Homogenizer (Polytron), then sonicated 5 min. at 30-40 °C in an ultrasonic bath (Elgasonic 102 Swiss made, 25 kHz) at room temperature, to disrupt cells. The sample was then centrifuged for 5 103 104 min. at 5000 g and the supernatant decanted and filtered. filtered. The extraction was repeated on 105 the pellet, the sample was centrifuged and the supernatant filtered on the same filter previously used. The filter and the funnel were washed three times with little volumes of MeOH; the two 106 extracts and washings were collected together, then dried by rotavapor at 40 °C; the residue re-107 suspended in 2 mL distilled water was then stored at -30 °C until analysis. 108

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110 Fish tissue microcystin (MYC) extraction

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Five grams (wet weight) of muscle tissue from each fish was extracted. The sample was 112 homogenized in 10_mlmL MeOH for 15min. using an Ultra-Turrax T8 (IKA Werke, Staufen, 113 Germany) grinder and then sonicated for 5 min. at 30-40°C in an ultrasonic bath (Elgasonic Swiss 114 made, 25 kHz) to disrupt cell membranes. The sample was centrifuged for 5min. at 5000 g and the 115 supernatant decanted and filtered on a paper filter. The extraction was repeated on the pellet, the 116 117 sample was centrifuged, and the supernatant filtered on the same filter previously used. The filter 118 and the funnel were washed three times with little volumes of_MeOH; the two supernatants and the 119 washings were gathered, then reduced to a small volume (1-2 mlmL) by rotary evaporator (Bùchi, Switzerland) at 40°C, and diluted to 5mlmL with MeOH. One mlmL (for fish) of the extract 120 (corresponding to 1 g of tissue) were then added with 1mlmL of distilled water and loaded onto a 121 HLB SPE Waters OASIS cartridge, preconditioned with 1ml mL MeOH followed by 1ml mL of 122 123 distilled water. The column was washed with 1mlmL of 5% MeOH in distilled water. Microcystins Comment [M6]: What tissues???

Comment [M7]: What sample? The pellet??

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were eluted by 1 <u>mlmL</u> of MeOH. The MeOH eluate was dried by rotary evaporator at 40°C; the
residue, dissolved in 2 <u>mlmL</u> distilled water, was stored at -30 °C for subsequent microcystin
analysis with the EnviroGard Elisa kit.

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128 CYN and MYC analysis by ELISA assays

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Muscle tissue extracts from 17 fish caught in 2012 in MG and S stations were analyzed using the
Abraxis Cylindrospermopsin ELISA Microtiter Plate immunoassay (Abraxis Bioscience CA).

ELISA assays were performed in accordance with the manufacturer's instructions using the calibration concentrations suggested. The Abraxis immunoassay declares the detection limit is 40 ppb, with percentage coefficients of variation below 10% for standard and below 15% for samples. The final reaction solution absorbances of the kit were measured at 450 nm with an Anthos 2010 microplate spectrophotometer (Anthos – Labtech, Salzburg, Austria).

Muscle tissue extracts from 79 fish samples were analyzed using the EnviroGard Microcystins Plate 137 138 Kit (Strategic Diagnostics Inc., Newark, DE, USA), a direct competitive ELISA for quantitative detection of microcystins and nodularins (limit of quantification, LOQ = 0.1 ppb). This 139 immunoassay does not differentiate between microcystin-LR and two other microcystin variants 140 (MC-RR and MC-YR) but detects their presence to differing degrees. The concentrations at 50% 141 inhibition (50% B/Bo absorbance signal) for these compounds (ppb) are: microcystin-LR 0.31, 142 microcystin-RR 0.32, microcystin-YR 0.38. The final reaction solution absorbances of the kit were 143 measured at 450 nm with an Anthos 2010 microplate spectrophotometer (Anthos - Labtech, 144 Salzburg, Austria). The analytical method to determine microcystins in water and fish samples was 145 146 previously validated according to the decision 2002/657/CEE (De Pace et al., 2014).

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148 Sample handling and trace elements and PCB analysis

Ten specimens of Cyprinus carpio (common carp) caught from two stations (MC, LD) of Pertusillo 150 Lake in April, 2017 (figure 2,3) were analysed also for trace elements and PCBs. After sampling, 151 the specimens were stored in ice boxes with dry ice, transferred to the laboratory and immediately 152 kept in a deep freezer. Subsequently, the frozen fish samples were thawed and biometric 153 measurement were made (weight range: 868-3195 g, mean: 1296±697 g; length range: 37.0-60.0 154 cm, mean: 43.1±6.9 cm). From each specimen the muscle tissue was dissected, homogenized and 155 analyzed. The extractive analytical procedure and the instrumental conditions to determine trace 156 element concentrations have been described in detail elsewhere (Barone et al., 2013). Briefly, about 157 0.5 g of the samples were digested to a transparent solution with a mixture of HNO₃-HClO₄ (8:3) 158 for cadmium (Cd), lead (Pb), chromium (Cr), copper (Cu) and zinc (Zn) determination and with a 159 mixture of H_2SO_4 -HNO₃ (1:1) for mercury (Hg). The completely digested samples were allowed to 160 cool temperature and diluted with deionized water according to the method recommended by 161 Official Italian Agencies (G.U.R.I., 1994). The content of elements was determined by atomic 162 163 absorption spectrophotometry (Shimadzu AA 7000). Zn was analysed by flame, Cd, Pb, Cr, and Cu by using a graphite furnace (high density tube) (GFA-7000), Hg was measured by using a hydride 164 vapour generator (HVG-1) after reduction by NaBH₄. Concerning PCBs, the concentrations of 165 indicator PCBs (28, 52, 101, 138, 153 and 180) were determined using analytical procedures 166 previously described and validated (Storelli, 2014). Briefly, about 40 g of powder were mixed with 167 Na₂SO₄ and spiked with PCB 143 used as internal standard. The mixture was extracted with 168 169 hexane:acetone: acetone (9:1) and the extracts were concentrated in order to determine the fat 170 content by gravimetry. Next the extract was dissolved in hexane and cleaned by passing through 8 g 171 of acid silica (H₂SO₄, 44% w. w.), using 50 mL of a mixture of hexane/dichloromethane (1/1, v/v) for elution of the analytes. The eluate was evaporated to dryness and redissolved in 100 mL of iso-172 173 octane. For the analysis of PCBs, a Thermo Trace GC connected with a Thermo PolarisQ MS

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operated in electron impact ionization (EI) mode was equipped with a 30 m, i.d. 0.25 mm and 0.25 174 um Rtx 200 capillary column (Thermo, Austin, Texas, USA). The MS was used in the SIM mode 175 with two ions monitored for each PCBs homologue group in specific windows. One ml of the 176 cleaned extract was injected in splitless mode (injector temperature 90 °C then to 300 °C with 70 177 °C/min), splitless time 1.50 min, pulse pressure time 1.50 min, pressure pulse 25 psi. Helium was 178 used as carrier gas at constant flow (1.0 ml/min). The temperature of the Rtx 200 column was held 179 at 90 °C for 1.50 min, then increased to 180 °C at a rate of 15 °C/min, further increased to 280 °C at 180 a rate of 5 °C/min, further increased to 300 °C at a rate of 40 °C/min, held for 7 min. 181

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183 *Quality assurance*

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Reference tissue (Tort-2 Lobster Hepatopancreas, National Research Council of Canada, Ottawa, 185 186 Ontario, Canada) was treated and analysed in the same way as the samples. Results (Hg: $0.28 \pm$ 0.03; Cd: 26.2 \pm 2.4; Pb: 0.32 \pm 0.18; Cr: 0.73 \pm 0.16; Cu: 101 \pm 13; Zn: 188 \pm 12 μ g g⁻¹ dry 187 weight) were in good agreement with the certified values (Hg: 0.27 ± 0.06 ; Cd: 26.7 ± 0.60 ; Pb: 188 0.35 ± 0.13 ; Cr: 0.77 ± 0.15 ; Cu: 106 ± 10 ; Zn: 180 $\pm 6 \ \mu g \ g^{-1}$ dry weight) and the standard 189 deviations were low, proving good repeatability of the methods. The results for standard reference 190 material displayed recoveries of the elements ranging from 91 to 104% (n = 3). The limit of 191 detection (LOD) (Hg: 5; Cd: 0.12; Pb: 10; Cr: 5; Cu: 26; Zn: 24 ng g⁻¹ wet weight) was defined as 192 the concentration corresponding to three times the standard deviation of blanks, and the standards of 193 quantification (LOQs) were the following: Hg: 13; Cd: 0.30; Pb: 38; Cr: 16; Cu: 81; Zn: 87 ng g⁻¹ 194 wet weight. Two blank samples were analysed together with each sample batch. Metal 195 196 concentrations in blanks were below the detection limits in all the analyses. Blanks and calibration standard solutions were similarly analysed as the digested sample solution, and calibration curves 197 were constructed. Analyses were duplicated to check the reproducibility of the results. Relative 198

standard deviations among replicates were always less than 10%. Recovery tests were performed 199 for the investigated metals in selected samples by spiking analysed samples with aliquots of the 200 metal standards and then carrying out digestion. The recovery percentages ranged from 96 to 99%. 201 Metal concentrations are presented as $\mu g g^{-1}$ wet weight basis. For PCBs quality control was 202 performed through the analysis of procedural blanks, a duplicate sample and a standard reference 203 204 material [CRM349 for PCBs (cod liver oils) (BCR, Brussels)] within each batch of samples. The recovery percentage of the standard reference standard reference material was within the range of 205 86 and 105%. For the samples and standard reference materials, the relative standard deviations 206 (RSD) were <10% for all the detected compounds. The limit of detection (LOD) for PCBs ranged 207 from 0.02 to 0.50 ng g^{-1} on a lipid weight basis, while the limit of quantification (LOQ) varied from 208 0.20 to 1.30 ng g⁻¹ on a lipid weight basis. Appropriate standard solution was added to the samples 209 210 and recovery values were between 82 and 104%. The trace element and PCB concentrations in the samples were expressed as $\mu g g^{-1}$ and $ng g^{-1}$ wet weight, respectively. 211

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213 Statistical analysis

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Kruskal-Wallis test was conducted to verify the difference in the levels of trace metal and PCB accumulation, while simple linear regression coefficient was used to examine the correlations between PCBs and specimen length. To investigate size influence on PCB accumulation, the length of fish was chosen, because less subject to fluctuation than body weight (Diaz et al., 1994). The level of significance was set at p < 0.05.

- 220
- 221 Microscopic observations

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223 The water samples were stored in ice chests and transported to the laboratory. For microscopic

observations water subsamples were analyzed by an inverted microscope (Leitz Labovert FS)
according to Utermöhl (1931) and Lund et al. (1958), using 25 ml sedimentation chambers for
phytoplankton identification and cell density estimation.

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228 Results and discussion

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230 Trace element and PCB concentrations

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The trace element concentrations detected in the study showed Zn values ranging from 1.15 to 4.32 232 $\mu g g^{-1}$ wet weight (2.31 $\mu g g^{-1}$ wet weight), while Cu showed much lower concentrations, ranging 233 from 0.15 to 0.61 μ g g⁻¹ wet weight (0.36 μ g g⁻¹ wet weight) (p < 0.001) (fig.2). The considerable 234 difference in levels between these two metals is not unique to the species here studied, being part of 235 a general picture suggesting muscle tissue not to be considered a specific physiological site for Cu 236 (Zia and Khan, 1989). Cr levels were very low too, ranging from 0.02 to 0.05 μ g g⁻¹ wet weight 237 (0.03 μ g g⁻¹ wet weight) (p < 0.001). Among non-essential metals the highest concentrations were 238 recorded for Hg with values between 0.27 and 0.53 μ g g⁻¹ wet weight (0.40 μ g g⁻¹ wet weight), 239 followed by Pb showing levels from 0.05 to 0.28 μ g g⁻¹ wet weight (0.14 μ g g⁻¹ wet weight), while 240 Cd registered the lowest values between 0.03 and 0.05 μ g g⁻¹ wet weight (0.04 μ g g⁻¹ wet weight) (p 241 < 0.001). A comparison with data in the literature shows a wide concentration heterogeneity for all 242 243 metals studied. However, our Hg levels are very similar to those found by Stong et al. (2013) in common carp from Lake Chapala in Mexico, but very higher than those reported by Vicarova *et al.* 244 (2016) in the same species from three reservoirs in the Czech Republic. For Cd and Pb, the levels in 245 this study are in line with values reported by Yancheva et al. (2014) in muscle tissue of common 246 247 carp from Topolnitsa reservoir in Bulgaria. For essential metals, our Cr values are in good

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248	agreement with results found in the muscle tissues of common carp from the uncontaminated
249	fishponds in the Czech Republic (Čelechovská et al., 2007) and Kabul River in Pakistan (Yousafzai
250	et al., 2017). In contrast, our Zn values are lower than those reported by Yousafzai et al. (2017) and
251	by Čelechovská <i>et al.</i> (2007) in muscle tissue of common carp from the Keban Dam Lake in Turkey
252	and the fishponds in the Czech Republic, respectively. Regarding Cu concentrations, samples
253	analysed in this study showed levels of the same order of magnitude of those reported for common
254	carp from the Czech Republic (Čelechovská et al., 2007). To safeguard public health, concentration
255	standards in fish for some heavy metals have been established by the European Commission). In
256	particular, Hg, Pb and Cd limit values at 0.50, 0.30 and 0.05 μ g g ⁻¹ wet weight respectively, have
257	been fixed (Official Journal of the European Union, 2006, 2014, 2015). In this context, no analysed
258	fish sample showed concentrations exceeding the European Directive proposed limits for Pb and Cd
259	while for Hg, slightly exceeding levels were registered in two samples (0.51 and 0.53 μ g g ⁻¹ wet
260	weight).There are no European guidelines for fish consumption established as regards Cu, Zn and
261	Cr, but the UK Food Standards Committee's Report fixed Zn and Cu limits at 50 and 20 $\mu g \ g^{\text{-1}}$ wet
262	weight respectively, while the Western Australian Food and Drug Regulation List (Usero <u>et al.</u>
263	2003) fixed Cr limits at 5.5 μ g g ⁻¹ wet weight. Our detected results were always lower than these

264 **human**these human consumption limits.

265 The subset of six PCB congeners here tested were selected by the International Council for the Exploration of the Sea (ICES) as contamination indicators, due to their easy quantification 266 compared to the other non-dioxin-like PCBs, however representing all relevant degrees of 267 chlorination. The data analysis showed that PCBs 153 and 138 were the most frequently detected 268 congeners (detection in 100% of samples), while PCBs 101 and 180 were detected with 50% and 269 70% frequency, respectively, and PCBs 28 and 52 were below the detection limits in all samples 270 examined. The total concentrations of indicator PCBs were 95.8-202.5 ng g⁻¹ lipid weight, with a 271 mean value of 148.6 ng g^{-1} lipid weight. PCBs 153 and 138 with mean values of 62.6 ng g^{-1} lipid 272 11



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weight and 55.4 ng g⁻¹ lipid weight were the highest in concentration, followed by PCB 180 273 showing a mean concentration of 18.7 ng g⁻¹ lipid weight and PCB 101 exibithing the lower mean 274 value equal to 11.9 ng g⁻¹ lipid weight. The PCB bioconcentration in aquatic organisms correlates 275 with the degree of chlorination, the stereochemistry and lipophilicity (Fox et al., 1994). Generally, 276 277 congeners with a high chlorination grade are more difficult to metabolise and eliminate than less 278 chlorinated congeners. Our data well fit this general picture, being low chlorinated congeners PCBs 28 and 52 below the detection limit, PCB 101 contributing for 8%, while hexa- and 279 heptachlorinated biphenyls 138, 153 and 180 together constituted a consistent percentage of the 280 total PCB burden representing 92%. Generally, the largest and potentially oldest fish exhibit higher 281 PCB levels than younger organisms. Despite of this, no correlation between fish length and total 282 PCB concentrations was observed (R = 0.42; P > 0.05) in the present study, probably as 283 284 consequence of scarce PCB contamination in the Pertusillo basin. These PCBs have been recommended by the EU as indicators of PCB contamination because generally they represent 285 approximately half of the total ndl-like PCBs existing in food. In fact, the European Food Safety 286 287 Authority (EFSA) Scientific Panel regarding Contaminants in the Food Chain (CONTAM Panel) 288 recommends the sum of these six PCBs as an appropriate marker for risk assessment of ndl-PCBs. Regulation No. 1259/2011 of the European Union (EU) (Official Journal of the European Union, 289 2011) has set de novo maximum tolerable levels for the sum of the six indicators non-dioxin-like 290 PCBs in muscle meat of freshwater fish that, apart from some exceptions, is of 125 ng g^{-1} wet 291 weight. Our results presented on a lipid weight basis have, hence, been converted to wet weight 292 basis to conform to legal standard. According to this, the sum of six "indicator" congener 293 concentrations was below the conventional permissible consumption limit in all samples examined 294 $(1.27 \text{ ng g}^{-1} \text{ wet weight}).$ 295

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297 Microcystin and cylindrospermopsin concentration

Superficial fortnightly water samples taken from March to April 2012 and from October 2012 to 299 300 March 2013 were analyzed for phytoplankton presence. In these winter samples only 16 species were detected; the lack of summer samples, due to difficulties in carrying out regular water 301 samplings, did not allow a complete evaluation of phytoplanktonic composition. In a few summer 302 303 samples analyzed by the Basilicata Agency for Environmental Protection (ARPAB) in 2014, 9 other 304 species were detected (ARPAB, 2015). The poor presence of phytoplanktonic species detected in 305 this study may also be due to the need for column samplings and more systematic monitoring. However, even in the past the lake showed the presence of a limited number of species (29 species 306 detected, Ruggiu and Saraceni, 1978). No cyanotoxins were detected in the analyzed water samples. 307 In fish 86% of total tissue samples were positive for MYC presence, at concentration values ranging 308 from a minimum of 0.19 ng/g to a maximum of 2.01 of 2.01 ng/g b.w. (fig. 4-6). Micropterus 309 salmoides, Carassius carassius and Cyprinus carpio were the species with highest concentration 310 capacity and averages. 311

ELISA analyses showed the presence of CYN in 64% of samples, with maximum concentrations at 0.78 ng/g in muscle (fig. 7). *Cyprinus carpio* and *Perca fluviatilis* were the species with highest concentration capacity and averages. ARPAB phytoplankton analyses in summer 2014 showed the presence of *Aphanizomenon* sp., which could take account for CYN presence (Messineo <u>et_al.</u>, 2010).

In May, 2016 fifteen fish samples from four stations (2 carps from MG, 2 carps from MB, 5 chubs and 6 perchs from MC) were analyzed for MYC presence (fig. 5), showing the highest mean content (0.72 ng/g), in perchs. In the following year (April, 2017), nine samples (5 carps from MC and 4 carps from LD) showed a mean content (0.91 and 0.93 ng/g, respectively, fig.6) higher than that of 2016 carps (0.29 and 0.28 ng/g, respectively). The toxicity of microcystins in fish depends on the balance between accumulation and metabolism (Ito et al., 2002); the observed speciesFormatted: Font: Italic

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323	specific sensitivities have been interpreted as the result of anatomical, physiological and behavioral	
324	differences among the various fish orders (Tencalla and Dietrich, 1997; Fischer and Dietrich,	
325	2000): the detoxification capacities via the glutathione-S-transferase pathway are species-specific	
326	dependent, too (Cazenave et al., 2006).	Comment [M10]: Review the redaction
327	CYN accumulation in ichthyic fauna was previously investigated in crayfish (Cherax	
328	quadricarinatus), rainbow fish (Melanotaenia eachamensis) (Saker and Eaglesham, 1999),	
329	freshwater mussels (Anodonta cygnea) (Saker et al., 2004), salmonids (Salmo trutta) (Messineo et	Formatted: Font: Italic
330	al., 2010) and finfish (Berry et al. 2012).	Formatted: Font: Italic
331	The acute Tolerable Daily Intake (TDI) guideline for MC-LR, proposed by WHO in 1998 for an	
332	adult of 60 kg b.w. (0.04 µg/kg body weight/day, Chorus and Bartram, (1999) was revised by	
333	USEPA in 2006, with new proposed guidelines developed for acute and chronic risk (0.006 and	
334	0.003 microcystin µg/kg b.w./day, respectively; US Environmental Protection Agency, 2006), but	
335	no guidelines for cancerogenicity were proposed, due to the insufficient adequacy of the available	
336	studies. In the same 2006, the International Agency for Research on Cancer classified microcystin-	
337	LR as possibly carcinogenic to humans (group 2B: IARC, 2010). Case-control studies in southwest	
338	China recently confirmed the link between MYC serum levels and occurrence of hepatocellular	
339	carcinoma occurrence in humans (Zheng et al., 2017).	
340	For an adult human weighing 60 kg and ingesting 300 g serving of fish muscle, the microcystin	
341	level in 14.5 % of muscle samples analyzed from 2010 to 2012 was 1.6 -fold the recommended TDI	
342	acute value of EPA, and 36.3% of muscle samples were even 3.3 -fold the recommended chronic	

Contaminant classes like polycyclic aromatic hydrocarbons, trace elements, PCBs and microcystins 344 345 are known to produce synergistic effects on organisms: in fish heavy metals may cause enhanced toxic effects if combined (Espina et al., 1997), in vitro and in vivo studies on cyanobacterial 346 extracts, PCB 153 and fluoranthene (Bartova et al., 2011) provide evidence on synergistic effects of 347

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value.

348 tumor promotion.

In Italy, microcystin contaminations in ichthyic fauna were detected in several lakes (Bruno *et al.*, 2009; Bruno *et al.*, 2012). MYCs demonstrated to be a recurrent component among the lake Pertusillo main contaminants, being detected in fish tissue all along the duration of the study. No MYC producing cyanobacteria were found in our phytoplankton analyses but several benthic species are MYC producers, too, and an extended monitoring for phytobenthic toxic species in the sediments of the lake would be needed, to investigate the reasonthe reason why a higher presence of these toxins was detected in the cyprinid species.

Zn levels detected by ARPAB lake water monitoring in 2014 (between 5 and 83 µg/l; ARPAB,
2015) are known to increase the growth and intracellular MYC production in *Microcystis aeruginosa* cultures (Polyak *et al.*, 2013). A recent meta-analysis has also shown that persistent
organic pollutants, among which PCBs, are able to stimulate cyanobacterial growth (Harris and
Smith, 2016).

A more extended monitoring is needed to define the presence of these different contaminants in 361 362 ichthyic fauna, their role in the recurrent fish deaths in the lake, and the exposure risk of people of the lake region by consuming contaminated lake fish. As Pertusillo Lake is part of a SCI zone, the 363 PCBs content in lake fish could endanger also the fish-eating birds, through biomagnification. Lake 364 Pertusillo is mesotrophic-eutrophic (ARPAB, 2015), and several episodes of algal blooms occurred 365 in the lake during the last seven years. Organisms are usually exposed not only to isolated 366 environmental pollutants, but to chemical mixtures which individual components may be present at 367 concentrations lower than their safety threshold levels. 368

Although the concentrations of metals and PCBs detected in the analysed fish samples are not high, the presence of these different compounds in association with microcystins suggests the possibility of future cyanobacterial toxic blooms in this environment through their growth stimulation, pointing out the need for restoration programs to improve the trophic conditions of the reservoir. Moreover, Formatted: Font: Italic

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373	given the presence of the industrial activities of oil drilling in the area, further studies are needed to		
374	investigate the potential contamination of oil compounds in Pertusillo ichthyic fauna.		
375	Conclusions		
376			
377	The ichthyic fauna of Pertusillo appears to be interested by multiple contaminant concentrations.	Cor	nment [M11]: Improve the redaction
378	The MYC production may be synergistically influenced and enhanced in the aquatic environment	For	matted: Highlight
379	by some trace element concentrations.	Cor	nment [M12]: That is not a conclusion
380	Waiting for new tools to assess the overall impact of these pollutants on aquatic life and human		
381	health, the managing policy remains the exploration and implementation of cost-effective and		
382	appropriate remediation, coupled with the search for environmentally more benign products and		
383	processes, which should aim to minimize introduction of critical pollutants into the aquatic		
384	environment.	Cor	nment [M13]: That is not conclusion
385			
386			
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726 Legend

Figure 1. Study site and station coordinates.



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Sampling station Е N Masseria Crisci MC 40.28977 15.95180 40.28710 15.9527 Rifreddo R Spinoso S 40.28044 15.96638 Madonna Grumentina MG 40.29172 15.92957 Montemurro Bridge MB 40.28238 15.9825 Lake Damming LD 40.27522 15.99157

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736 Figure 3. Concentrations of six PCB indicator congeners in common carp.





Table 1. Phytoplanktonic species identified in the superficial samplings of 2012.

	Phytoplanktonic species
Cyanobacteria	Coelosphaerium kutzingianum Nageli
Diatomeae	Asterionella formosa Hassall
	Cyclotella kutzingiana Thwaites
	Cymbella sp. C. Agardh
	Fragilaria crotonensis Kitton
	Gyrosigma attenuatum (Kutzing) Rabenhorst
	Melosira italica (Ehrenberg) Kutzing
	Melosira varians C. Agardh
	Navicula ssp. Bory de Saint-Vincent
	Nitzschia acicularis (Kutzing) W. Smith
	Rhizosolenia sp. Ehrenberg
	Stephanodiscus astraea (Ehrenberg) Grunow
Chlorophyceae	Oocystis lacustris Chodat
Conjugatopnyceae	Closterium kutzingii Bredisson
\mathcal{O}	Closterium pronum Brébisson
Dinophyceae	Ceratium hirundinella (O.F.Müller) Dujardin



Figure 4. Microcystin concentration in fish muscle tissue (all the stations) during three years (2010-2012).



Figure 5. Microcystin concentration in fish samples from three lake stations (MG, MB, MC) inMay, 2016.

753 Figure 6. Microcystin concentration in fish samples from two lake stations (MC, LD) in April,

754 2017.





Figure 7. Cylindrospermopsin concentration in fish muscle tissue during 2012.

770	Figure legend
771	
772	Figure 1. Study site and sampling stations
773	
774	Figure 2. Trace element concentrations in common carp.
775	
776	Figure 3. Concentrations of six PCB indicator congener in common carp.
777	
778	Figure 4. Microcystin concentration in fish muscle tissue (all the stations) during three years (2010-
779	2012).
780	
781	Figure 5. Microcystin concentration in fish samples from three lake stations (MG, MB, MC) in
782	May, 2016.
783	
784	Figure 6. Microcystin concentration in fish samples from two lake stations (MC, LD) in April,
785	2017.
786	
787	Figure 7. Cylindrospermopsin concentration in fish muscle tissue during 2012.
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789	Table legend
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791	Table 1. Phytoplanktonic species identified in the superficial samplings of 2012.
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