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The interactions of Fullerene C₆₀ and Benzo(α)pyrene influence their bioavailability and toxicity to zebrafish embryos

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14 Abstract

15 This study aimed to assess the toxicological consequences related to the interaction of fullerene 16 nanoparticles (C₆₀) and Benzo(α)pyrene (B(α)P) on zebrafish embryos, which were exposed to C₆₀ 17 and B(α)P alone and to C₆₀ doped with B(α)P. The uptake of pollutants into their tissues and intra-18 cellular localization were investigated by immunofluorescence and electron microscopy. A set of 19 biomarkers of genotoxicity and oxidative stress, as well as functional proteomics analysis were applied to assess the toxic effects due to C_{60} interaction with $B(\alpha)P$. The carrier role of C_{60} for 20 21 $B(\alpha)P$ was observed, however adsorption on C_{60} did not affect the accumulation and localization of 22 $B(\alpha)P$ in the embryos. Instead, C_{60} doped with $B(\alpha)P$ resulted more prone to sedimentation and less 23 bioavailable for the embryos compared to C₆₀ alone. As for toxicity, our results suggested that C₆₀ 24 alone elicited oxidative stress in embryos and a down-regulation of proteins involved in energetic metabolism. The C₆₀ + B(α)P induced cellular response mechanisms similar to B(α)P alone, but 25 26 generating greater cellular damages in the exposed embryos.

27 Capsule

28 Once C_{60} nanoparticles and $B(\alpha)P$ meet in water, they reciprocally affect their bioavailability and, by

29 consequence, their toxicity to organisms.

30 *Keywords*: Fullerene nanoparticles; *Danio rerio*; oxidative stress; proteomics; trojan horse effect

31 **1. Introduction**

32 Fullerenes are carbonaceous nanoparticles (NPs) broadly used in several applications including 33 targeted drug delivery systems, lubricants, energy devices, catalysis, surfaces for antiwear 34 applications, cosmetics and sporting goods, due to their outstanding chemico-physical properties 35 (Yadav and Kumar 2008; Mousavi et al., 2017). As a result of the growing production of fullerenes 36 - and especially C₆₀- there is rising concern regarding their presence and impacts on the natural 37 ecosystems. Indeed, fullerenes nanoparticles have been detected in many environmental matrices as 38 atmospheric aerosol (Sanchis et al., 2012) waters (Farrè et al., 2010; Pakarinen et al., 2013; 39 Astefanei et al., 2014), sediments (Sanchis et al., 2015) and soils (Carboni et al., 2016). Therefore, 40 it is extremely important to assess their interactions and toxic effects on wildlife, with particular 41 emphasis on aquatic environments, which act as ultimate sinks for NPs. In fact, the same properties 42 that render fullerenes a unique and innovative material can trigger deleterious effects to natural 43 biocenosis.

44 The toxicity of C_{60} has been described in bacteria (Freitas Cordiero et al., 2014), crustaceans 45 (Klaper et al., 2009), bivalves (Canesi et al., 2010; Al-Subiai et al., 2012), chironomids (Waissi et 46 al., 2017) and fishes (Ferreira et al., 2012; Gorrochategui et al., 2017) as well as the potential for 47 trophic transfer (Fortner et al., 2010; Chen et al., 2014). Besides its inherent toxicity, C₆₀ has also 48 exceptional sorption capacity towards hydrophobic chemicals (Hu et al., 2014; Velzeboer et al., 49 2014) that may significantly affect their bioavailability, bioconcentration and toxicity. Although 50 some studies showed the ability of C_{60} to sequester diverse contaminants and to reduce their toxicity 51 (Yang et al., 2010; Park et al., 2011), it can conversely act as carrier for organic pollutants 52 enhancing their biological effects on the organisms (Baun etal., 2008; Al-Subiai et al., 2012; 53 Ferreira et al., 2014; Seke et al., 2017; Li et al., 2017). Therefore, the release of C₆₀ into the aquatic 54 environment in the presence of toxic chemicals may pose a further risk for ecosystems, with hardly 55 predictable effects.

56 This study aimed to assess the interactive effects of fullerene NPs (C_{60}) and Benzo(α)pyrene 57 $(B(\alpha)P)$ on zebrafish (*Danio rerio*) embryos. Specifically, we doped C_{60} with $B(\alpha)P$ ($C_{60}+B(\alpha)P$) 58 from now on) and compared the effect on zebrafish embryos exposed to the two contaminants 59 singly administered and to the $C_{60} + B(\alpha)P$ complex. This experimental plan allowed to assess the 60 accumulation and toxicity only of the $B(\alpha)P$ adsorbed on C_{60} without any interference of free 61 hydrocarbon. A thorough evaluation of chemico-physical interactions between the two pollutants 62 has been performed, and the uptake and distribution of C_{60} and $B(\alpha)P$ were shown through 63 advanced microscopy techniques. To evaluate whether $C_{60} + B(\alpha)P$ affects different molecular pathways compared to the two singly administered pollutants, a suite of biomarkers was applied. 64 65 The activity of proteins involved in the detoxification and antioxidant response, namely glutathione-S-transferase (GST), catalase (CAT) and superoxide dismutase (SOD), was measured, while the 66 67 oxidative damage was assessed by the measurement of protein carbonylation (PCC). The 68 genotoxicity was assessed by the application of single gel cell electrophoresis (SCGE) assay, DNA diffusion assay and Micronucleus test. Proteomics analysis was also performed to evaluate changes 69 70 of embryos proteome profile, and suggesting possible mechanisms of action of the pollutants, both 71 alone or in combination.

72

73 **2. Materials and Methods**

74 2.1 Materials

75 C_{60} (CAS number: 99685-96-8) and all reagents used for chemical and biomarker analyses were 76 purchased by Sigma-Aldrich (Steinheim, Germany). B(α)P powder (CAS number: 50-32-8) was 77 supplied by Dr. Ehrenstorfer, (Augsburg, Germany).

78

- 79 2.2 C₆₀ characterization
- 80 The bulk C_{60} and C_{60} + B(α)P were observed at Zeiss LEO 912ab Energy Filtering TEM operating
- at 100 kV, at a magnification of 25-50,000x using a CCD-BM/1 K system.

Dynamic Light Scattering (DLS) was used to measure the hydrodynamic diameter (size distribution) and the charges at the surface (ζ-potential) of water suspended C_{60} nanoparticles. The measurements were performed on a Malvern Zetasizer Nano ZS instrument (Malvern instruments, UK) equipped with a device for the ζ-potential measurement, employing a solid state He-Ne laser (633 nm) as a light source and recovering the scattered light at an angle of 173° (Series software – version 7.02 – Particular Sciences, UK).

88

89 2.3 B(α)P sorption on C₆₀

90 Two aliquots of C₆₀ (200 mg/L) were suspended in 200 mL of MilliQ water and stirred for 15 days 91 at 20 °C. The first aliquot was doped with 1 mg/L B(α)P dissolved in dimethylsulhpoxide (DMSO), 92 while the second portion was not contaminated. The two suspensions were stirred for 5 days at 20 93 °C in the dark. They were then centrifuged at 3,000 x g for 30 min. The precipitated C_{60} with $B(\alpha)P$ 94 adsorbed were completely dried in an oven at 40 °C and used for embryos exposures. For 95 measuring the $B(\alpha)P$ fraction remaining solubilized in water, the supernatants were treated with 96 toluene (1:5 v/v, respectively) leaving the mixture under stirring for 90 min at 20 °C in order to 97 extract the free B(α)P solubilized in water by liquid/liquid extraction. The amount of B(α)P in 98 toluene solutions was measured by fluorescence detection, as described in Supplementary 99 Materials. The same technique was used to measure the amount of $B(\alpha)P$ adsorbed on C₆₀ NPs. In 100 this case, the samples were prepared by dissolving 5 mg of dried C_{60} + B(α)P in 200 mL toluene. A 101 sample containing 5 mg of untreated C_{60} in 200 mL toluene served as a blank.

102

103 2.4 Preparation of C_{60} suspensions and hydrodynamic behaviour in exposure media

104 The C₆₀ and C₆₀ + B(α)P were suspended and equilibrated at concentration of 20 mg/L, in zebrafish

105 water (ZFW) for 10 days by stirring in the dark at 20 °C. DLS analysis was performed as described

above to determine both hydrodynamic diameters and surface charges (ζ potentials) of each sample.

To evaluate the sedimentation process, the UV-vis absorbance spectra of the two suspensions were
 aquired on an Agilent model 8543 spectrophotometer at room temperature.

109

110 2.5 Zebrafish embryo exposure

Adult zebrafish of the AB strain were bred in the fish facility of the Department of Biosciences (University of Milan), to obtain 1-cell stage embryos. Our facility is strictly compliant with the Italian legislation (Legislative Decree No. 116/92) concerning animal welfare, as also certified by the authorization released by the Milan municipality (Art. 10 of Legislative Decree No. 116, dated 27.1.1992). Animal procedures were carried out in conformity with the relevant guidelines and regulations.

117 To avoid any physical interference with the uptake of C_{60} , removal of chorion with pronase (0.5 118 mg/mL) was performed at 24 h post-fertilization (hpf), immediately prior to the exposure. Embryos were then exposed to B(α)P (8 μ g/L), C₆₀ (20 mg/L) and to C₆₀ + B(α)P in Petri dishes in a total 119 120 volume of 4 mL. B(α)P concentrations were defined based on the effective B(α)P sorption on C₆₀ 121 (20 mg/L) measured by emission spectra (see results). A preliminary range-finding assured that 122 concentrations of C_{60} and $B(\alpha)P$ were not able to produce mortality or any morphological embryos 123 alteration. Control embryos were exposed to zebrafish water (ZFW) and to vehicle (0.08% DMSO) 124 only. The exposure proceeded until 96 hpf under semistatic conditions, renewing the exposure 125 solutions every 24 h in new vessels. To prevent embryos pigmentation for $B(\alpha)P$ visualization in 126 tissues, ZFW was added with 0.003% 1-phenyl 2-thiourea (PTU). For biochemical analyses and 127 proteomics, embryos were stored at -80 °C until processing. For advanced microscopy and genotoxicity assessment, embryos were immediately processed at the end of the exposure as 128 129 described below. Experiments were run at least 3 times for each analysis.

130

131 2.6 Electron microscopy

132 A detailed description of ultrastructural analysis procedures is reported in Binelli et al. (2017). Ten 133 embryos from each experimental group were fixed in a mixture containing 4% paraformaldehyde 134 and 2.5% glutaraldehyde in 0.1 M sodium cacodylate buffered solution (pH 7.4). Then the embryos 135 were postfixed in 1% OsO₄, dehydrated in a graded ethanol series and infiltrated in Araldite-Epon. 136 Ultrathin sections of about 70 nm were obtained by Ultracut E microtome (Reichert, Austria). Counterstain of sections was not performed to avoid interference with C₆₀ visualization. Digital 137 138 images were acquired using a CCD-BM/1K system, and image elaboration was performed using the 139 ESI vision software AnalySIS (Soft Imaging Systems, Muenster, Germany).

140

141 2.7 Immunohistochemistry

Details of the procedure are described in Binelli et al. (2017). Briefly, embryos were fixed in paraformaldehyde (4%) and cryo-protected. For immunofluorescence, cryostat sections (10 μm) were incubated with primary antibody anti-polycyclic aromatic hydrocarbons (anti-PAHs 1/100 in PBS, Santa Cruz) and exposed to secondary antibody (Alexa Fluor 488 goat anti-mouse 1:200, Thermo Fisher Scientific). Finally, samples were mounted in PBS/glycerol (1:2 v/v) with DNAbinding dye 40-60-diamidino2-phenylindole (DAPI). Sections were observed by a confocal microscope Leica SP2 microscope equipped with He/Kr and Ar laser (Leica, Wetzlar, Germany).

149

150 2.8 Accumulation of C_{60} in zebrafish embryos

The quantification of C_{60} accumulated in zebrafish embryos was performed according to the method described by Waissi et al. (2017), with slight modifications. Embryos were exposed in triplicate to C_{60} and $C_{60} + B(\alpha)P$ and to ZFW only (N = 160 for each treatment). After 96 hpf, embryos were collected and washed with MilliQ water, then homogenized in 1 mL of 2% NaCl solution. Toluene (1 mL) was added, the solution vortexed and transferred in an ultrasonic bath for 15 min. After sedimentation, the toluene fraction was collected and C_{60} absorbance was measured at 335 nm using a Jenway spectrophotometer (Stone, UK). The baseline absorbance detected in controls was subtracted to absorbance of C_{60} samples. The C_{60} concentration was determined based on standard curve (0.01-10 µg/mL r² = 0.9997).

160

161 2.9 Biomarkers analyses

A detailed description of biomarkers analysis is reported in Supplementary Materials. The GST,
SOD and CAT activities were analysed on homogenates obtained from pools of 60 embryos for
each treatment. PCC was evaluated on homogenates obtained from pools of 80 embryos.

Genotoxicity was performed on cells dissociated from a pool of 10 zebrafish embryos (three pools per treatment) according to the methods described in Parolini et al. (2017). Briefly, cell viability was assessed by the trypan blue dye exclusion method. The percentage of DNA in the comet tail and the ratio between migration length and comet head diameter (LDR) were used as endpoints of primary genetic damages. The apoptotic and necrotic cell frequency and the frequency of micronuclei (MN‰) were measured as fixed genetic damage.

171

172 2.10 Functional proteomics

173 The analysis was performed on pools of 90 embryos for each experimental group. A detailed 174 description of the procedure is reported by Binelli et al. (2017). Briefly, 200 µg of protein for each 175 group were precipitated using a chloroform/methanol/water mixture (4:1:3 v/v). Proteins 176 resuspended in rehydratation (denaturing) buffer were loaded in 18 cm pH 3-10 non-linear gradient 177 IPG strips (GE Healthcare, Milan, Italy) and IEF was performed on the Ettan IPGphor II system 178 (GE Healthcare, USA). Protein separation in the second dimension was performed on 12% 179 acrylamide gel in an Ettan DALTsix electrophoretic unit (GE Healthcare, UK). Gels were died by 180 silver stain (ProteoSilver Plus Silver Stain kit; Sigma Aldrich, Milan, Italy), according to producer 181 instructions. Gel images were analyzed by the ImageMaster 2D Platinum software (Amersham 182 Biosciences, USA). Significant protein differences were investigated comparing gels from controls 183 group (ZFW, DMSO) with those from treatment groups. Spots were statistically evaluated in terms

184 of the mean relative volume (vol.%) using Student's t-test for unpaired samples taking p<0.05 as 185 significant treshold. A further criterion for differential regulation was employed such as minimum 2-folds change cut-off relative to controls. Significantly modified protein spots were excised from 186 187 gels, destained, dehydrated with acetonitrile and digested with trypsine (Sigma Aldrich, Milan, 188 identified MALDI **TOF-TOF** (matrix-assisted Italy). The proteins were by laser 189 desorption/ionization time of flight) mass spectrometry analysis; the Peptide Mass Fingerprinting (PMF) was performed using an Ultraflex III MALDI-TOF/TOF mass spectrometer (Bruker 190 191 Daltonics, Billerica, MA, United States). Spectra were analyzed by the Flex Analysis software 192 v.3.0. Mascot (Matrix Science Ltd., London, UK, http://www.matrixscience.com). On-line-193 available software was used for PMF search in NCBInr or Swiss-Prot/TrEMBL databases with 194 taxonomy set for Danio rerio.

195

196 2.11 Statistical analysis

Biomarker data were investigated through one-way analysis of variance (ANOVA) after checking for normality and homoscedasticity, taking p<0.05 as significance cut-off. The ANOVA was followed by the Duncan's *post-hoc* test to investigate significant differences between exposure groups. The analyses were performed using the STATISTICA 7.0 software package.

201

202 **3. Results and Discussion**

203 3.1 C₆₀ characterization

The bulk C_{60} consisted mostly of isometric NPs, with mean diameter estimated by TEM micrographs of about 35.6 ± 10.9 nm and few submicrometric aggregates. A similar structure was observed also for $C_{60} + B(\alpha)P$ (Fig. S1). The DLS analysis of C_{60} suspension in MilliQ water (1 mg/mL) showed the presence of a homogeneous population of NPs aggregates with hydrodynamic radius of 519 ± 169 nm, and polydispersion index (PDI) of 0.39. The stability of the suspension was 209 confirmed by the highly negative surface charge value (-36 \pm 1 mV) derived from a ζ -potential 210 measurement.

211 In ZFW, the Z average measured for C_{60} suspension (20 mg/L) was 899 ± 97 nm with PDI of 0.376, 212 indicating the presence of homogeneous population of NPs (trace red in Fig. 1A). On the contrary, 213 the suspension of $C_{60} + B(\alpha)P$ showed the presence of two aggregate populations, centered at 767 ± 214 44 nm and 189 ± 13 nm (trace blue in Fig. 1A) showing also the presence of smaller aggregates. 215 Nevertheless, the counts per second of scattered light in this second set of measurements resulted 216 very poor, indicating that a very quick sedimentation phenomenon occurred. Indeed, the suspension 217 just after the DLS measurement pointed out evident sediment at the bottom of the cuvette (Fig. S2). 218 Moreover, UV-Vis analysis highlighted also a significant sedimentation of $C_{60} + B(\alpha)P$ with respect 219 to C_{60} alone, leading to a decrease of concentration of the suspended NPs in ZFW (Fig. 1B). These results suggested that once contaminated with $B(\alpha)P$, C_{60} NPs were more prone to aggregation and 220 221 to be easily settled out of suspension.

Finally, the ζ potential value of C₆₀ alone was -23.4 ± 0.2 mV, and a similar value was measured for C₆₀ combined with B(α)P, equal to -21.6 ± 0.3 mV.

224

225 3.2 B(α)P sorption on C₆₀

226 A small fraction (2.9 %) of the administered $B(\alpha)P$ was recovered in the water phase after 5 days of contamination, while a marked amount of B(α)P (38%) was adsorbed on C₆₀, equal to 378 µg/g. 227 Based on these results, the concentration of $B(\alpha)P$ corresponding to suspensions containing 20 228 mg/L of C₆₀ was set to 8 μ g/L. These results confirmed the sorption capacity of C₆₀ towards B(α)P 229 230 in the water media as reported for other PAHs (Baun et al., 2008; Hu et al., 2014). Such findings 231 highlighted that C_{60} could alter significantly the fate and transport of B(α)P in the aquatic 232 ecosystems. Moreover our results suggest that as $B(\alpha)P$ is relatively stable and can move in the 233 atmosphere for a long time, it can bind to atmospheric NP such as C₆₀, and be subsequently 234 introduced in the water environment. The high sedimentation rate observed for the complex C60 +

B(α)P strongly suggests that a relevant fraction of the hydrocarbon adsorbed on C₆₀ could reach and accumulate in the sediments.

237

238 3.3 Accumulation of C₆₀

239 The measurement of C_{60} body burden showed a higher accumulation in embryos exposed to C_{60} alone (16.32 \pm 6.45 ng/embryo, corresponding to three-fold increase) than C₆₀ contaminated with 240 241 $B(\alpha)P(3.90 \pm 3.39 \text{ ng/embryo})$. Therefore, the observed increase of C₆₀ sedimentation due to $B(\alpha)P$ 242 adsorption reduced the NPs bioavailability and uptake by the embryos. This result highlight that, in 243 natural conditions, the presence of contaminants in water could significantly influence the 244 environmental fate of C₆₀, potentially enhancing its distribution in sediments. According with our result, a previous study on zebrafish embryos showed that the combination of C_{60} and Hg^{2+} 245 246 increased NPs size and sedimentation, resulting in a lower accumulation of NPs in embryos, 247 compared to the C₆₀ alone (Henry et al., 2013). Similarly, two other studies on nano-TiO₂ in 248 combination with metals have suggested that chemico-physical interactions (e.g. adsorption) 249 between NPs and contaminants can significantly alter their accumulation in organisms (Pavagadhi 250 et al., 2014; Fan et al., 2016).

TEM observations showed a microvilli-mediated internalization of C_{60} NPs in enterocytes mediated (Fig. 2A). The adsorption of $B(\alpha)P$ on C_{60} did not modify this behavior, as also doped NPs entered enterocytes, (Fig. 2B). TEM observations confirmed the ability of C_{60} to pass through the gill cell membranes and accumulate in the epithelium cells (Fig. 2C).

255 3.4 B(α)P accumulation

The B(α)P fluorescence signal was detected in gills (Fig. 3C, D) and in the gastrointestinal tract (Fig. 3F, G) of embryos exposed to the hydrocarbon. A similar pattern was revealed also in C₆₀ + B(α)P exposed embryos, showing that C₆₀ can act as carrier for the adsorbed B(α)P. Confocal observations suggested that the B(α)P adsorbed on C₆₀ enters the organism mostly through the

260 gastrointestinal tract, where it can be released and transferred to other compartments as described 261 for different carbon nanomaterials (CNMs; Wang et al. 2011; Su et al., 2013; Seke et al., 2017). 262 Nevertheless, the mechanisms determining the release of contaminants from CNMs and distribution 263 in the organism are still barely understood, and might vary depending on the CNM. For instance, in 264 our recent study we showed that the $B(\alpha)P$ sorbed on carbon nanopowder (CNPW) was taken up by 265 zebrafish embryos and it followed the physical contaminant distribution rather than its natural 266 accumulation (Binelli et al., 2017). On the contrary, the immunoistochemistry analysis showed that 267 the adsorption on C_{60} did not affect the embryo $B(\alpha)P$ distribution. It is known that fullerene 268 structures, as well as other allotropic carbon-based materials like carbon nanotubes (CNT), interact 269 with the aromatic moieties of many different molecules by π - π interactions (Lu et al., 2006). Yet the 270 sorption of aromatic hydrocarbons by C₆₀ has been calculated and compared with the ability of 271 CNTs to absorb these small molecules (Huffer et al., 2017), concluding that the sorption by CNTs is 272 stronger than that by C₆₀ and may be attributable, among others, to the smaller surface area of the 273 fullerene aggregates in water with respect to the ones of other CNMs (Yang et al., 2006). Therefore, 274 the small difference in the biodistribution of $B(\alpha)P$ when administered alone or associated to C_{60} , 275 could be due to a faster equilibrium release from this material with respect to the dissociation from 276 other carbon-based materials in the physiological environment as the gastrointestinal fluids.

277

278 3.5 Effects of C₆₀

No mortality or teratogenic effects have been recorded in zebrafish embryos exposed to contaminants alone or in combination. Concerning the oxidative stress biomarkers, exposure to $B(\alpha)P$ determined a significant inhibition of CAT activity compared to DMSO (p <0.0001), but did not affect SOD activity. A significant increase of SOD (p = 0.0004) and CAT (p = 0.01) activities was observed in embryos exposed to C₆₀ alone. On the other hand, the activity of the two enzymes was restored to control levels following C₆₀ + B(α)P exposure, resulting significantly lower in respect to C₆₀ alone (p = 0.0003 for SOD and p = 0.0013 for CAT) (Fig. 4A,B). The measurement of protein carbonylation showed an increase in carbonyl content exclusively in embryos exposed to C_{60} alone compared to controls (p = 0.019; Fig. 4D).

288 The increase of SOD and CAT activity confirmed the ability of C₆₀ to induce antioxidant response, 289 as already pointed out in other studies performed on several aquatic models (Usenko et al., 2008; 290 Klaper et al., 2009; Ferreira et al., 2012; Waissi et al., 2017; Lv et al., 2017). The oxidative stress 291 generated by C₆₀ was also confirmed by the increase of protein carbonylation -marker of oxidative 292 damage- observed in embryos exposed to C₆₀ alone. Indeed, the carbonylation of proteins involved 293 in various cellular mechanisms has been described, as consequence of the oxidative stress generated 294 following exposure to NPs (Driessen et al., 2015). On the contrary, there was no evidence of 295 oxidative damage in co-exposure, and a significant reduction of the activity of antioxidant enzymes 296 was observed in comparison to the single pollutant. This result is likely related to the lower accumulation of C_{60} observed in embryos exposed to the $B(\alpha)P$ doped NPs in respect to C_{60} alone, 297 298 therefore unable to induce a measurable cellular response.

299 A significant increase of GST activity was observed in embryos exposed to $B(\alpha)P$ (p = 0.012) and 300 C_{60} (p = 0.0002 vs control) administered alone. GST is involved in phase II of 301 metabolism/detoxification catalyzing the conjugation of glutathione to several environmental 302 pollutants and oxidative stress by-products (van der Oost et al., 2003). The induction of GST 303 activity upon exposure to C₆₀ alone confirmed the active role of this enzyme in the cellular response 304 to the NPs, as observed in previous studies (Usenko et al., 2008; Klaper et al., 2009). On the 305 contrary, it is intriguing that $C_{60} + B(\alpha)P$ determined a significant reduction of GST activity with 306 respect to control (p = 0.0002) (Fig. 3C). This result agrees with previous observations concerning 307 CNPW contaminated with $B(\alpha)P$ (Della Torre et al., 2017). The same effect was also reported in a 308 study on zebrafish hepatocytes exposed to C_{60} and $B(\alpha)P$ (Ferreira et al., 2014), suggesting a 309 specific inhibition of this enzyme, which may be due to a physical interaction with the doped NPs. 310 However, the results obtained so far do not allow the identification of the mechanism underlying 311 this inhibition.

Concerning genotoxicity, the exposure to $B(\alpha)P$ alone increased significantly the DNA % in the comet tail (p < 0.0001) as well as the LDR (p = 0.033) in respect to DMSO (Fig. 4F,G), but did not induce cell necrosis and occurrence of MN (Fig. 4H,I). Our results confirmed the genotoxic potential of $B(\alpha)P$, which triggered the onset of DNA damage; its weak effect is likely related to the short exposure time (72 h), which may have been not sufficient to determine fixed genetic damages, as normally expected after exposure to $B(\alpha)P$ (Parolini et al., 2017).

The C₆₀ administered alone did not cause any primary or fixed DNA damage compared to controls. The results highlighted the absence of genotoxic effects by C₆₀ in agreement with previous studies showing the low genotoxic potential of C₆₀ (Jacobsen et al., 2008) and the inability to generate primary damage to biological systems both *in vitro* and *in vivo* (Shinoara et al., 2009; Matsuda et al., 2011; Ema et al., 2012).

323 Conversely, in embryos exposed to $C_{60} + B(\alpha)P$, a significantly higher frequency of necrotic cells 324 was found compared to control (p < 0.0001) and C_{60} administered alone (p = 0.0012) (Fig. 4H). 325 Exposure to $C_{60} + B(\alpha)P$ also resulted in the increase of MN occurrence compared to controls (p = 326 0.047), even if the MN frequency of this group did not exceed 5 ‰ (Fig. 4I). An extremely low 327 frequency of apoptotic cells was found in all exposure conditions (<2% data not shown). The results highlighted that the adsorption of $B(\alpha)P$ increased the cellular damage with respect to the C₆₀ alone. 328 329 Two possible hypotheses could explain this effect: the first one suggests that when the pollutants 330 are administered in co-exposure $(C_{60} + B(\alpha)P)$ they induce an increase of cell disruption without 331 direct interacting with the DNA. Alternatively, the reduction of GST activity elicited by the two 332 contaminants administered together might reduce the detoxifying capacity of the embryos, thus 333 enhancing the genotoxic effects of $B(\alpha)P$. In support of the latter hypothesis, a higher cell death and 334 genotoxicity, together with the inhibition of GST activity, were observed upon exposure to CNPW 335 and $C_{60} + B(\alpha)P$, in previous studies (Ferreira et al., 2014; Della Torre et al., 2017).

336

337 3.6 Functional proteomics

The application of proteomics allowed the identification of molecular events involved in the responses to pollutants alone and in co-exposure. About 2,000 different spots in each analyzed gel were visualized: 220 spots were in common between DMSO and B(α)P, 235 between control and C₆₀, and 173 between control and C₆₀+B(α)P (Fig. S3). A significant variation in terms of volume percentage for 28, 50 and 21 spots was measured for the three treatments, respectively. The final cut-off (\geq 2-folds) revealed 23, 34 and 14 varied spots for the treatments in comparison to controls.

The exposure to $B(\alpha)P$ up-regulated 12 different proteins and down-regulated the remaining 11 with respect to DMSO (Fig. S4). The C₆₀ administered alone down-regulated 31 different proteins and overregulated 3 of them (Fig. S4). The co-exposure C₆₀ + B(α)P induced a significant overregulation of 13 proteins and down-regulation of 1 protein only (Fig. S4). Mass spectrometry analysis allowed the identification of 12 changed proteins in embryos exposed to B(α)P, 27 proteins for C₆₀ exposure and 5 varied proteins for the co-exposure (Tab. 1,S1,S2,S3).

350 Going deeper, a variation in the amount of vitellogenin cleavage products (Vtg1,5,7) was observed 351 in all the three exposure groups. Vtgs are glycophospholipoproteins, which constitute the yolk-352 proteins precursors in all oviparous species, including Teleosts. Vtgs are synthesized in the liver of 353 female and incorporated into oocytes where, following a proteolytic cleavage, provide essential 354 nutrients for the embryos (Byrne et al., 1989). Therefore, Vtgs proteolitic cleavage processes are 355 fundamental for the proper embryo development and the evaluation of Vtgs profiles, both at gene 356 and protein level, is considered a useful tool for highlighting toxic effects due to various types of 357 environmental pollutants (Muncke and Eggen 2006; Gundel et al., 2007; 2012; Hanish et al., 2010; 358 Ponnodurai et al., 2012; Hao et al., 2013). The C_{60} -induced down-regulation of Vtgs is in line with 359 the effects on zebrafish embryos following exposure to Quantum Dots (Petushkova et al., 2015) and 360 on adults of Daphnia magna exposed to Ag-NPs (Rainville et al., 2014). In this latter study, a 361 significant reduction of Vtg-like proteins was observed together with an increase in protein 362 oxidation. Therefore the down-regulation of Vtgs could be due to the protein oxidation processes as 363 a consequence of the oxidative stress generated by C_{60} .

In addition to nutritional function, Vtgs have a protective role towards different stressors (Sun and Zhang, 2015). Particularly, the involvement of Vtg1-like proteins in the DNA repair mechanism has been demonstrated in zebrafish embryos (Lai et al., 2006). Therefore, the over-regulation of Vtg1 observed in embryos exposed to B(α)P and to C₆₀ + B(α)P might suggest the induction of a protective mechanism involved, for instance, in the genotoxic damage repair processes.

Another protein engaged in the lipid metabolism and in metabolic processes is the 369 370 ApolipoproteinA-I (Apoa1b), which was modulated in embryos exposed to $B(\alpha)P$ and C_{60} 371 administered singly, albeit in opposite way. The alteration of Apoalb due to modification oxidation for instance- might trigger cytotoxic and degenerative effects, therefore promoting the 372 373 onset of circulatory alterations (Park and Cho, 2011; Filipe et al., 2013). Indeed, a recent study 374 highlighted that the exposure of zebrafish embryos to particulate matter_{2.5} (PM_{2.5}) could enhance the 375 occurrence of cardiovascular toxicity through the proteolitic degradation of lipoproteins (Kim et al., 376 2015). The down-regulation of Apoa1b suggested a similar mechanism also for C_{60} . Conversely, the 377 over-regulation of Apoa1b measured in embryos exposed to $B(\alpha)P$ paralleled the increase of Vtg1. 378 Indeed, Apoab1 in fact also owns anti-inflammatory and antioxidant properties (Filipe et al., 2013), 379 which could contribute to the protective response of the embryos towards this pollutant.

The exposure to both contaminants, administered alone and in combination, affected the betahemoglobin (BE1), a protein assigned to oxygen transport. The modulation of BE1 in fish is usually related to environmental stress conditions, such as modification of temperature, salinity and hypoxia (Eissa e Wang, 2016), but also to the exposure to environmental pollutants (Duarte et al., 2010; Narra 2016). The observed alteration of BE1 content might affect oxygen supply, thereby compromising the development and survival of the embryos.

The exposure to C_{60} induced the down-regulation of several kinases such as muscle creatine kinase A (Ckma), creatine kinase M-type isoform X1 (Ckmb) and nucleoside diphosphate kinase B (Nme2b.2). These proteins are involved in cellular signaling, growth and differentiation, as well as energetic metabolism (Tanimura et al., 2014). The alteration of kinases levels might induce the onset of negative effects on embryos. Indeed, the loss of Nme2b.2 induced severe vasculature
 malformations (Feng et al., 2014) and cardiomyopathy (Hippe et al., 2009) in zebrafish embryos.

The exposure to $B(\alpha)P$ determined a down-regulation of Type I cytokeratin enveloping layer (Cyt1). Cytokeratins are structural proteins involved in the formation of intermediate filaments of epithelial cells and in the mainteinance of cell integrity and adhesion in tissues, promoting resistance to mechanical stress (Padhi et al., 2006). The down-regulation of several keratins (Krt4, Krt5 e Krt8) has been already observed in zebrafish embryos exposed to $B(\alpha)P$ (20 µg/L) (Binelli et al., 2017), supporting the hypothesis that this chemical is able to affect the functionality of structural proteins.

Another down-regulated protein in embryos exposed to $B(\alpha)P$ is the Fatty Acid Binding Protein7 (Fabp7), a chaperonine responsible for cellular fatty acid transport (Furuhashi and Hotamisligil, 2013). At the embryonic level, Fabp7 plays a key role in the development of the central nervous system and affects proper development of the visual system (Liu et al., 2004). Therefore, $B(\alpha)P$ might alter the development and function of the nervous system and the visual apparatus through the down-regulation of Fabp7, as previously suggested (He et al., 2012; Binelli et al., 2017).

405 Overall, proteomic analysis confirmed the different mechanism of action of single contaminants and 406 their combination. Results suggested that the oxidative stress generated by C_{60} triggers a general 407 reduction of the metabolic activity in the embryos, confirming recent findings of Lv and coauthors 408 (2017), who suggested that the toxicity of C_{60} in *D. magna* might be correlated with oxidative stress 409 and reduction of energy acquisition. On the contrary, $B(\alpha)P$ alone and adsorbed on C_{60} up-regulated 410 proteins involved in the homeostatic response to cellular stress.

411

412 4. Conclusions

413 The present study showed how the adsorption of $B(\alpha)P$ by C_{60} altered the hydrodynamic behavior 414 of the NPs, consequently reducing their bioavailability and intake by the embryos. We showed that the B(α)P adsorbed on C₆₀ is bioavailable and accumulated in embryos. The integration of data obtained through biomarkers and functional proteomics suggests that B(α)P alone and C₆₀ + B(α)P affect similar cellular mechanisms, with the latter triggering severer cellular damages.

418 Our results highlight that, in the natural environment complex chemico-physical-biological 419 interactions arise, possibly determining unexpected ecotoxicological consequences for the 420 organisms.

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424

425 Figure Captions

Figure 1 C₆₀ characterization. DLS profile of C₆₀ alone and contaminated with B(α)P, in zebrafish water (A). Particle size distribution of 3 measurements of 10 runs is given by numbers. UV-Vis spectra of C₆₀ suspensions (B).

Figure 2. C_{60} observation in embryos. TEM images showing C_{60} indicated by arrows in gut (A) of embryos exposed to C_{60} and in gut (B) and gills (C) of $C_{60} + B(\alpha)P$. n = nucleus, m = mitochondrion, mv = microvilli, l = lumen.

432 **Figure 3**. **B**(α)**P accumulation**. Cryostate sections showing the uptake of B(α)P (in red) in gills 433 (C,D,E) and digestive apparatus (F,G,H) of zebrafish embryos. Controls (A-B), B(α)P (C,F), C₆₀ +

434 $B(\alpha)P(D-G)$ and $C_{60}(E,H)$. DNA (nuclei) is marked in blue (DAPI coloration). SB = swim bladder,

435 L = gut lumen, Y = yolk sac.

436 Figure 4. Effects on biomarkers. Effects on the activity (mean \pm SEM) of SOD (A) CAT (B),

437 GST (C); protein carbonylation (D); and genotoxic effects as DNA strand breaks (E), LDRs (F),

438 occurrence of necrotic cells (G) and MN (H) measured in zebrafish embryos (96 hpf) (n = 3; pool of

439 3 independent experiments). Different letters correspond to values significantly different (one-way
440 ANOVA, Duncan's *post-hoc* test, p < 0.05).

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Spot ^a	Fold change (↓/↑) ^b	Protein identification	NCBInr Accession number	Molecular function ^c
B	(a)P vs DMSO			
2	2.3↓	Type I cytokeratin. enveloping layer (Cyt1)	AAH65653.1	Structural molecule activity
6	3.2↑	Apolipoprotein A-I precursor (Apoa1b)	NP 001093614.2	Lipid transport
7	3.6↑	Hemoglobin beta embryonic-1.1 (BE1)	NP 932339.1	Oxygen transport
9	2.5↑	Hemoglobin beta embryonic-1.1 (BE1)	NP_932339.1	Oxygen transport
11	2.0.	Fatty acid binding protein 7, brain, a (Fabp7a)	NP 571680.1	Lipid transport
14	2.9↑	Vitellogenin 1 (Vtg 1)	AAH94995.1	Lipid transport
19	2.8↑	Hemoglobin beta embryonic-1,1 (BE1)	NP 932339.1	Oxygen transport
20	2.5↑	Vitellogenin 1 (Vtg 1)	AAH94995.1	Lipid transport
21	2.5↑	Hemoglobin beta embryonic-1.1 (BE1)	NP 932339.1	Oxygen transport
27	3.61	Vitellogenin 1 (Vtg 1)	AAH94995.1	Lipid transport
30	2.0.	Vitellogenin 5 (Vtg 5)	AAH97081.1	Lipid transport
	C ₆₀ vs Ctrl			
1	4.0↓	Vitellogenin 1 (Vtg 1)	AAH94995.1	Lipid transport
5	3.5↓	Vitellogenin 1 (Vtg 1)	AAH94995.1	Lipid transport
6	2.9↓	Vitellogenin 1 (Vtg 1)	AAH94995.1	Lipid transport
7	3.0↓	muscle creatine kinase a (Ckma)	NP_571007.2	Kinase activity
				ATP binding
9	3.4↓	Vitellogenin 1 (Vtg 1)	AAH94995.1	Lipid transport
10	2.1↓	Vitellogenin 1 (Vtg 1)	AAH94995.1	Lipid transport
11	2.0↓	Apolipoprotein A-I precursor (Apoa1b)	NP_001093614.2	Lipid transport
12	3.0↓	Vitellogenin 1 (Vtg 1)	AAK94945.1	Lipid transport
13	2.3↓	Vitellogenin 1 precursor (Vtg 1)	NP_001038362.3	Lipid transport
16	3.7↓	Vitellogenin 1 (Vtg 1)	AAI39514.1	Lipid transport
17	2.7↓	Hemoglobin beta embryonic-1.1 (BE1)	NP_932339.1	Oxygen transport
20	3.2↓	Hemoglobin beta embryonic-1.1 (BE1)	NP_932339.1	Oxygen transport
21	3.2↓	Vitellogenin 1 (Vtg 1)	AAK94945.1	Lipid transport
22	8.1↓	Vitellogenin 1 (Vtg 1)	AAH94995.1	Lipid transport
24	7.2↓	Vitellogenin 7 (Vtg7)	AAW56971.1	Lipid transport
26	2.8↓	muscle-specific creatine kinase (Ckma)	AAK64515.1	Kinase activity ATP binding
27	2.8↓	muscle-specific creatine kinase (Ckma)	AAK64515.1	Kinase activity ATP binding
32	3.1↓	Vitellogenin 1 (Vtg 1)	AAH94995.1	Lipid transport
33	4.9↓	Hemoglobin beta embryonic-1.1 (BE1)	NP_932339.1	Oxygen transport
35	2.6↓	Vitellogenin 1 (Vtg 1)	AAH94995.1	Lipid transport
38	3.0↓	creatine kinase M-type isoform X1 (Ckmb)	XP_005157650.1	Kinase activity ATP binding
43	3.0↓	nucleoside diphosphate kinase B (Nme2b.2)	NP_571002.1	ATP binding
46	2.3↓	Vitellogenin 1 (Vtg 1)	AAK94945.1	Lipid transport
47	3.3↓	Hemoglobin beta embryonic-1.1 (BE1)	NP_932339.1	Oxygen transport
52	2.3↓	Vitellogenin 1 precursor (Vtg 1)	NP_001038362.3	Lipid transport
53	2.7↓	nucleoside diphosphate kinase B (Nme2b.2)	NP_571002.1	ATP binding
54	3.0↓	Vitellogenin 1 (Vtg 1)	AAK94945.1	Lipid transport
C ₆₀	+ B(α)P vs Ctrl			
1	1 1 1	Vitalloganin 1 (Vtg 1)	A A HO4005 1	Linid transport
1	4.1 ∕ 1↑	Vitallogenin 1 (Vtg 1)	AA1194993.1 AA120514 1	Lipid transport
ט ד	4.1 2 0↑	Vitallogenin 1 (Vtg 1)	AA139314.1 AA120514.1	Lipid transport
/ 0	2.7 2.41	Hemoglohin beta ombruonia 11 (PE1)	MD 022220 1	Ovvgen transport
0	∠. 4 2.9↑	Vitalloganin 1 (Vta 1)	A A LIO A005 1	Linid transport
12	2.8	vitenogenin I (vig I)	AAN94993.1	Lipid transport

Table1 Proteins modified in zebrafish embryos upon exposure to C_{60} and $B(\alpha)P$ singly and in combination

^a ID number of spot on 2-DE map; ^b fold change increase (↑) or decrease (↓) in terms of relative spot volume (%V) in comparison with control (ZFW or DMSO);

^c from <u>www.uniprot.org</u> site.

Supporting information

The interactions of Fullerene C_{60} and $Benzo(\alpha)$ pyrene influence their bioavailability and toxicity to zebrafish embryos

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1. Methods

1.1 Fluorescence detection of $B(\alpha)P$ adsorption on C_{60}

Excitation and emission spectra were obtained with an Edinburgh FLS980 spectrofluorometer equipped with a 450 W xenon arc lamp. The spectra were corrected for source intensity (lamp and grating) and emission spectral response (detector and grating) by standard correction curves. The emission spectra of four toluene solutions of B(α)P at different concentrations (0.10, 1.0, 10, 100 mg/L) added with C₆₀ fullerene (25 mg/L) - to take into account possible energy transfer phenomena occurring between B(α)P and C₆₀- were analyzed, and a calibration curve obtained by plotting the intensity maxima of the peaks ($\lambda_{exc} = 360$ nm, $\lambda_{em} = 430$ nm). Then, the B(α)P concentration was measured on three independent samples of C₆₀ stock suspension contaminated with B(α)P (1 mg/L) in water, dried, and the aggregates suspended in toluene and analysed for the quantification of the B(α)P adsorbed portion (detection limit ≥ 0.05 mg/L).

1.2 Biomarkers analysis

Embryos were homogenized using a pestle in 100 mM potassium phosphate buffer (KCl 100 mM, EDTA 1 mM, protease inhibitors 1:100 v/v, dithiothreitol 1 mM pH 7.4). The homogenates were centrifuged at 15,000 x g for 10 minutes at 4 °C. The GST activity was measured by adding reduced glutathione (1 mM) in 100 mM phosphate buffer (pH 7.4) and using CDNB (1mM) as substrate. The reaction was monitored for 1 min at 340 nm. The CAT activity was determined by measuring the consumption of H_2O_2 (50 mM) in 100 mM potassium phosphate buffer (pH 7) at 240 nm. The SOD activity was determined by measuring the degree of inhibition of cytochrome c (10 μ M) reduction by the superoxide anion generated by the xanthine oxidase (1.87 mU/mL)/hypoxanthine (50 μ M) reaction at 550 nm. The activity is given as SOD units (1 SOD unit=50% inhibition of the xanthine oxidase reaction). Protein carbonyls were derivatized with 2,4-dinitrophenylhydrazine (DNPH) (10 mM in 2M HCl). Proteins were then precipitated and the pellet washed and resuspended in guanidine hydrochloride (6 M). The absorbance of protein-hydrozone was measured at 370 nm (Mecocci et al., 1999). The total protein content of each sample was measured according to the Bradford (1976) method using bovine serum albumin as standard.

Concerning biomarkers of genotoxicity, first we confirmed that the mean cell viability of dissociated cells from embryos exposed to B(α)P and C₆₀ alone and combined was always higher than the threshold value (70%) suggested for the application of the genotoxicity tests (Kirkland et al., 2007). The alkaline (pH > 13) Single Cell Gel Electrophoresis (SCGE) assay was performed according to the method described in Koshmel et al. (2008). One hundred cells per slide (n = 9; three slides per each pool) were analyzed using the Comet Score[®] image analysis software. The apoptotic cell frequency (%) was assessed analyzing three hundred cells per slide (n=6; two slides per each pool). The frequency of micronuclei (MN‰) was calculated on 400 cells/slide (n=6; two slides per each pool) according to Pavlica et al. (2000).

2. Results



Fig. S1 TEM images of bulk $C_{60}(A)$ and $C_{60} + B(\alpha)P(B)$.



Fig. S2 Sedimentation of suspensions of C_{60} alone compared to $C_{60} + B(\alpha)P$ in zebrafish water.



Fig. S3 Representative 2DE gels of zebrafish (96 hpf) exposed to $B(\alpha)P$ and C_{60} alone and C_{60} + $B(\alpha)P$. Red circles highlight proteins identified through mass spectrometry analysis.



Fig. S4 Protein spots differentially expressed in zebrafish (96 hpf) exposed to $B(\alpha)P$, C_{60} and $C_{60} + B(\alpha)P$, with respect to controls. The *y*-axis represents the fold change (in terms of relative spot volume, % V) of the protein spots, where a positive value indicates an increase in abundance and a negative value indicates a decrease in abundance.



Fig. S5 Venn diagram shows common protein spots between treatments (blue=down-regulation; red=up-regulation).

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	Eald shangs		NCBIng		Theoretical	Experimen	IT LASED	л эсат сп т сэч	1112
Spot ^a	roid change (↓/↑) ^b	Protein identification	Accession number ^e	Molecular function	pI/MW (kDa) ^d	tal pl/MW (kDa) ^e	Sequence coverage (%) ^g	N°. of matched peptides ^h	Scor
2	2.3↓	Type I cytokeratin enveloping layer (Cyt1) <i>[Danio rerio]</i>	AAH65653.1	Structural molecule activity	46.53/5.13	41.36/4.7	15	6/11	89
6	3.2↑	Apolipoprotein A-I precursor (Apoa1b) [Danio rerio]	NP_001093614.2	Lipid transport	30.18/6.05	30.56/5.1	28	7/11	78
7	3.6↑	Hemoglobin beta embryonic-1.1 (BE1) [Danio rerio]	NP_932339.1	Oxygen transport	16.27/6.89	13.19/5.7	29	4/4	84
6	2.5↑	Hemoglobin beta embryonic-1.1 (BE1) [Danio rerio]	NP_932339.1	Oxygen transport	16.27/6.89	13.19/7.3	29	4/6	73
11	2.01	Fatty acid binding protein 7, brain, a (Fabp7a) [Danio rerio]	NP_571680.1	Lipid transport	14.97/5.43	15.09/5.3	40	5/16	70
14	2.9↑	Vitellogenin 1 (Vtg 1) [<u>Danio rerio]</u>	AAH94995.1	Lipid transport	36.58/ 9.23	27.40/5.5	30	7/10	109
19	2.8↑	Hemoglobin beta embryonic-1.1 (BE1) [Danio rerio]	NP_932339.1	Oxygen transport	16.27/6.89	13.19/6.9	29	4/4	84
20	2.5↑	Vitellogenin 1 (Vtg 1) [<u>Danio rerio]</u>	AAH94995.1	Lipid transport	36.58/ 9.23	27.40/5.3	24	7/10	101
21	2.5↑	Hemoglobin beta embryonic-1.1 (BE1) [Danio rerio]	NP_932339.1	Oxygen transport	16.27/6.89	13.19/6.7	29	4/4	84
27	3.6↑	Vitellogenin 1 (Vtg 1) [<u>Danio rerio]</u>	AAH94995.1	Lipid transport	36.58/ 9.23	28.10/9.2	26	7/21	72
30	2.01	Vitellogenin 5 (Vtg 5) [Danio rerio]	AAH97081.1	Lipid transport	150/8.77	15.09/5.2	12	12/32	67

Table S1 Proteins modified in zebrafish embryos (96 hpf) exposed to $B(\alpha)P$ 8 µg/L respect to DMSO

from www.uniprot.org site;

Table S2 Proteins modified in zebrafish embryos (96 hpf) exposed to C₆₀ 20 mg/L respect to Ctrl

^d Predicted p*I* and MW according to protein sequence; ^e Experimentally determined p*I* and MW;

^f Results obtained by Peptide Mass Fingerprinting analysis;
 ^g Percentage of sequence coverage of matched peptides in the identified proteins;
 ^h number of matched peptide/total number of peptide searched;

ⁱ probabilistic score sorted by the software (protein scores greater than 60 were indicated as significant. p<0.05. by the program)

52	47	46	43	38	35	33	32	27	26	24	22	21	20	17	
2.3↓	3.3↓	2.3↓	3.0↓	3.0Ų	2.6↓	4.9↓	3.1↓	2.8J	2.8↓	7.2↓	8.1↓	3.2↓	3.2↓	2.7↓	
Vitellogenin 1 precursor (Vtg 1)	Hemoglobin beta embryonic-1.1 (BE1) [Danio rerio]	Vitellogenin 1 (Vtg 1) [<u>Danio rerio]</u>	nucleoside diphosphate kinase B (Nme2b.2)	creatine kinase M- type isoform X1 (Ckmb) [<i>Danio rerio</i>]	Vitellogenin 1 (Vtg 1) [Danio rerio]	Hemoglobin beta embryonic-1.1 (BE1) [Danio rerio]	Vitellogenin 1 (Vtg 1) [Danio rerio]	muscle-specific creatine kinase (Ckma) [<i>Danio rerio</i>]	muscle-specific creatine kinase (Ckma) [<i>Danio rerio</i>]	Vitellogenin 7 (Vtg7) <u>[Danio rerio]</u>	Vitellogenin 1 (Vtg 1) [Danio rerio]	Vitellogenin 1 (Vtg 1) [<u>Danio rerio]</u>	Hemoglobin beta embryonic-1.1 (BE1) [Danio rerio]	Hemoglobin beta embryonic-1.1 (BE1) <i>[Danio rerio]</i>	(Vtg 1) [<u>Danio rerio]</u>
NP_001038362.3	NP_932339.1	AAK94945.1	NP_571002.1	XP_005157650.1	AAH94995.1	NP_932339.1	AAH94995.1	AAK64515.1	AAK64515.1	AAW56971.1	AAH94995.1	AAK94945.1	NP_932339.1	NP_932339.1	
Lipid transport	Oxygen transport	Lipid transport	ATP binding	Kinase activity ATP binding	Lipid transport	Oxygen transport	Lipid transport	Kinase activity ATP binding	Kinase activity ATP binding	Lipid transport	Lipid transport	Lipid transport	Oxygen transport	Oxygen transport	
150/8.74	16.27/6.89	150/ 8.68	17.23/6.75	43.11/6.29	36.58/9.23	16.27/6.89	36.58/ 9.23	43.03/6.32	43.03/6.32	24.49/ 8.37	36.58/9.23	150/ 8.68	16.27/6.89	16.27/6.89	
30.68/7	14.87/6.5	16.01/6.2	18.56/6.7	48.50/6.3	27.67/7.3	14.87/6.9	27.67/5.8	45.05/6.3	45.05/6.3	28.50/ 7.9	27.26/8.3	16.25/7	13.19/6.7	14.87/6.5	
12	43	9	56	18	27	34	38	27	26	49	36	10	29	29	
12/34	5/10	9/14	8/12	6/7	7/18	5/9	10/17	9/16	9/13	12/21	10/21	11/21	4/4	4/9	
63	83	79	117	84	78	84	131	103	113	182	117	81	84	64	

																		I
^a ID nu ^b fold c ^c from	12	∞	Γ	5	-	opor	Snota		able S3	^h numt ⁱ proba	g Perce	f P eeul	^d Predi	° from	^b fold c	Ĵ	54	53
mber of sp hange incr www.unipr	2.8↑	2.4↑	2.9↑	4.1↑	4.1↑	(1/1) €	Fold		Proteins	er of matcl bilistic sco	ntage of se	rimentally of the obtained	cted p <i>I</i> and	www.unipr	hange incr		3.0	2.7J
ot on 2-DE map; ease (†) or decrease (1) ot.org site;	Vitellogenin 1 (Vtg 1) [Danio rerio]	Hemoglobin beta embryonic-1.1 (BE1) [Danio rerio]	Vitellogenin 1 (Vtg 1) [Danio rerio]	Vitellogenin 1 (Vtg 1) [Danio rerio]	Vitellogenin 1 (Vtg 1) [Danio rerio]	identification	Protein		modified in zebra	ned peptide/total numl re sorted by the softw:	quence coverage of m	tetermined pl and MV	MW according to pro	ot.org site;	ease (↑) or decrease ((Vtg 1) [Danio rerio]	diphosphate kinase B (Nme2b.2) Vitellogenin 1	nucleoside
) in terms of relative spot	AAH94995.1	NP_932339.1	AAI39514.1	AAI39514.1	AAH94995.1	Accession number ^e	NCBInr		fish embryos (96 hpf)	per of peptide searched; are (protein scores greater)	atched peptides in the ider	V; remrinting analysis:	otein sequence;) in terms of relative spot		AAK94945.1	NP 571002.1
volume (%V) in com	Lipid transport	Oxygen transport	Lipid transport	Lipid transport	Lipid transport	function	Molecular		exposed to C ₆₀ 20	than 60 were indicate	tified proteins;				volume (%V) in com		Lipid transport	ATP binding
parison with c	36.58/9.23	16.27/6.89	117/ 9.07	117/ 9.07	36.58/ 9.23	(kDa) ^d	n1/MW) mg/L + B	d as significar					parison with c		150/ 8.68	17.23/6.75
ontrol (ZFW).	27.40/5.3	13.19/6.7	27.40/5.5	30.7/6	27.40/5.2	(kDa)°	Experimental		(α)P 8 μg/L res	1t. p<0.05. by the p					ontrol (ZFW).		16.25/5.8	18.56/6.2
	24	29	30	10	30	Sequence coverage (%) ^g		Mase	pect to Ctrl	orogram)							∞ :	37
	7/10	4/4	7/10	7/12	9/18	N°. of matched peptides ^h		ot search resu									8/12	4/6
	101	84	109	93	112	Score		ılts ^r									73	71

^d Predicted p*I* and MW according to protein sequence;
^e Experimentally determined p*I* and MW;
^f Results obtained by Peptide Mass Fingerprinting analysis;
^g Percentage of sequence coverage of matched peptides in the identified proteins;
^h number of matched peptide/total number of peptide searched;
ⁱ probabilistic score sorted by the software (protein scores greater than 60 were indicated as significant. p<0.05. by the program)