

Sub-lethal effects induced by morphine to the freshwater biological model *Dreissena polymorpha*

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ABSTRACT

Opioids are considered as emerging contaminants in aquatic ecosystems, mainly due to their large illicit consume worldwide. Morphine (MOR) is the main opiate and it was commonly found at measurable concentrations in freshwaters. Even though its occurrence is well documented, just limited information is available regarding its hazard to non-target organisms. The aim of this study was of the evaluation of sub-lethal effects induced by MOR to the freshwater bivalve *Dreissena polymorpha*. We exposed mussels to two MOR concentrations (0.05 µg/L and 0.5 µg/L) for 14 days and we investigated the sub-lethal effects by a suite of biomarkers. The Neutral Red Retention Assay (NRRA) was used as a test of cytotoxicity, while the oxidative stress was evaluated by the activity of antioxidant and detoxifying enzymes, namely catalase (CAT), superoxide dismutase (SOD), glutathione peroxidase (GPx) and glutathione-S-transferase (GST), and by measuring the levels of lipid peroxidation (LPO) and protein carbonylation (PCC). The genetic damage was assessed by the Single Cell Gel Electrophoresis (SCGE) assay, the DNA diffusion assay and the micronucleus test (MN test). Finally, the filtration rate of *D. polymorpha* was evaluated in order to investigate possible physiological effects. Both tested concentrations reduced the lysosome membrane stability of bivalves, but only the highest MOR concentration induced significant changes in the activity of antioxidant enzymes (SOD, CAT and GPx) and increase in lipid peroxidation levels. Slight increase in primary DNA fragmentation was noticed, while no fixed genetic damage and alterations of the filtering rate were found.

Keywords: Morphine, Biomarkers, *Dreissena polymorpha*

1. INTRODUCTION

The scientific community has recently shown a growing interest for problems related to the presence of new contaminants such as pharmaceuticals and personal care products (PPCPs) and illicit drugs (Pal *et al.*, 2013) in the aquatic environment. Once used, these chemicals and their metabolites enter the sewage waters through urines and feces (Ternes, 1998; Zuccato *et al.*, 2000;

38 Heberer, 2002; Castiglioni *et al.*, 2006). The wastewater treatment plants (WWTPs) are built to
39 remove the organic matter and nutrients and are not suitable for the removal of most PPCPs and
40 drugs of abuse (Reungoat *et al.*, 2011; Pal *et al.*, 2013). Many monitoring studies showed
41 measurable concentrations (in the ng/L- μ g/L range) of several PPCPs and illicit drugs in both
42 European and US WWTP effluents and surface waters (Fent *et al.*, 2006; Santos *et al.*, 2010).
43 Among pharmaceuticals, analgesics are topical pain relievers and can be divided into two groups:
44 non-opioids (non-narcotic analgesics) and opioids (narcotic analgesic). The first family reduces pain
45 and inflammation interfering with the synthesis of prostaglandin hormones (Julien 1997), while the
46 latter group causes a muscular relaxation interacting with specific opioid receptor (MOR), a class of
47 G-protein-coupled receptors (Suzuki and Misawa, 1997). Considering their pharmacological
48 features, opioids are used as pharmaceuticals in human medicine, but also as drugs of abuse. The
49 latest World Drug Report (UNODC, 2013) has estimated that about 16.5 million people worldwide,
50 accounting for the 0.4% of the population aged 15-64, have used opiates as drugs of abuse at least
51 once in 2012. These chemicals are not the most common illicit drugs used worldwide, since
52 cannabis (3.9 % of the global population) and amphetamines (0.7 % of the global population)
53 showed a higher use prevalence (UNODC, 2013). However, their use trend remains stable over the
54 last years, with high prevalence in South-Western and Central Asia, Eastern and South-Eastern
55 Europe and North America (UNODC, 2013). Opiates are opium derivatives, a substance extracted
56 from *Papaver somniferum* and *Papaver setigerum*, historically prescribed for the care of cough,
57 anemia and diarrhea (Nicholson, 2003). Opium contains many active alkaloid compounds, mainly
58 morphine (MOR). In humans, MOR acts on the nervous system by binding to opioid receptors,
59 reducing pain and smoothing muscle contraction (Zhu *et al.*, 2005). MOR is metabolized for 87%
60 from hepatic carboxylase, and the main metabolite is represented by morphine-3 β -D-glucuronide
61 (75%, Baselt *et al.*, 2004). In addition, MOR is a metabolite of heroin, which has low affinity for
62 opioid receptors and only when it is converted into MOR (4%), 6-acetylmorphine (1,3%) and
63 morphine-3 β -D-glucuronide (38%) performs a pharmacological action (Baselt *et al.*, 2004; Maurer
64 *et al.*, 2006). For many decades, studies on vertebrates were focused on the pharmacological effects
65 of exogenous MOR and exogenous morphine-like compounds, but after the discovery of the
66 binding of MOR with opioid receptors, endogenous opioids have been identified (Lord *et al.*, 1977).
67 The presence of endogenous MOR is not a prerogative of vertebrates, as shown by studies
68 performed on different species of invertebrates (Stefano *et al.*, 2000). For instance, it is known that
69 mussels have opioid receptors in their nervous system (Stefano and Scharrer, 1996). The MOR-
70 opioid receptors interaction in these bivalves involves a release of dopamine (Zhu *et al.*, 2005), a
71 crucial neurotransmitter involved in oogenesis. Despite the abovementioned evidences, very few

72 studies have been carried out on aquatic organisms to evaluate the effects of MOR towards non-
73 target organisms. Mantione and co-workers (2002) showed that nitric oxide (NO) is released by the
74 pedal ganglia in *Mytilus edulis*, after the stimulation by the interaction between opioid receptors and
75 MOR metabolites. Other studies performed on microglia and immunocytes of *Mytilus edulis*
76 suggested an immunosuppressive activity of MOR (Stefano 1989; Stefano *et al.*, 1993), similar to
77 that described in humans (Stefano *et al.*, 1994; Makman *et al.*, 1995). A recent investigation by
78 Gagné and co-workers (2010) showed the neurochemical consequence of MOR exposure to the
79 freshwater bivalve *Elliptio complanata*. After injections of increasing MOR doses in the adductor
80 muscle (0.07; 0.15 and 0.75 mg/g wet weight), reductions in levels of serotonin and
81 acetylcholinesterase (AChE), as well as increases in dopamine and γ -aminobutyric acid (GABA)
82 levels, were noticed. Similar effects were obtained in the same mussel species exposed to a WWTP
83 effluent extract, in which MOR was detected at 0.1 $\mu\text{g/L}$ concentration (Gagné *et al.*, 2004).
84 However, to date no one investigation was performed to study neither the MOR cyto-genotoxicity
85 nor the involvement of oxidative stress in the mechanism of action of this drug towards non-target
86 organisms. Considering that MOR is frequently detected in European surface waters with an
87 average concentration of 50-55 ng/L (Karolak *et al.*, 2010; Terzic *et al.*, 2010; Jurado *et al.*, 2012;
88 Martinez Bueno *et al.*, 2011; Rosa Boleda *et al.*, 2011), and that the continual input of this drug can
89 lead to the exposure for the entire life-cycle of aquatic organisms, the investigation of its potential
90 sub-lethal toxicity is new and pivotal in freshwater ecotoxicology. The aim of this study was to
91 investigate the effects of MOR on the zebra mussels *Dreissena polymorpha*, using an *in vivo* multi-
92 biomarkers approach. Thanks to its physio-ecological features, this bivalve species is commonly
93 used in ecotoxicology, showing a good sensitivity to different emerging aquatic pollutants (Binelli
94 *et al.*, 2009a,b; Parolini *et al.*, 2010; Parolini and Binelli 2011; 2012), including illicit drugs
95 (Parolini *et al.*, 2013; Parolini and Binelli, 2013; Parolini and Binelli, 2014). Moreover, this filter-
96 feeding species has a great filtration rate (mean=200 mL/h/mussels) and it is more prone than other
97 biological models to introduce the aquatic pollutants into the organism, pointing out rapidly their
98 potential toxic effects. We exposed *D. polymorpha* specimens for 14 days to two low MOR
99 concentrations: 0.05 $\mu\text{g/L}$ and 0.5 $\mu\text{g/L}$. The end-points of twelve different biomarkers were
100 measured to assess MOR sub-lethal effects. Cytotoxicity was evaluated on hemocytes by the
101 Neutral Red Retention Assay (NRRA), while the activity of antioxidant and detoxifying enzymes,
102 namely catalase (CAT), superoxide dismutase (SOD), glutathione peroxidase (GPx) and glutathione
103 S-transferase (GST), as well as the lipid peroxidation (LPO) and the protein carbonyl content (PCC)
104 were applied as indices of oxidative stress in mussel homogenates. Primary (DNA strand breaks)
105 and fixed (apoptotic and micronucleated cell frequency) genetic damage was investigated on *D.*

106 *polymorpha* hemocytes by the Single Cell Gel Electrophoresis (SCGE) assay, the DNA diffusion
107 assay and the micronucleus test (MN test), respectively. Finally, the filtration rate was evaluated as
108 physiological biomarker.

109

110 2. MATERIALS AND METHODS

111 The MOR standard (CAS number 57-27-2) was purchased from Alltech-Applied Science (State
112 College, PA, USA), while all the reagents used for biomarker analyses were purchased from Sigma-
113 Aldrich (Steinheim, Germany). We diluted the methanol stock solution (1 g/L) to 10 mg/L in
114 ultrapure water (working solution), which was then used to obtain the MOR concentration in
115 experimental aquaria.

116

117 2.1 Experimental design

118 *D. polymorpha* specimens were collected in September 2012 by a scuba diver at a depth of 4-6 m in
119 Lake Lugano (Northern Italy), which is considered a reference site due to its low drug pollution
120 (Zuccato *et al.*, 2008). The mussels were gently cut off from the rocks, quickly transferred to the
121 laboratory in bags filled with lake water and placed in 15 L glass-holding aquaria filled with tap and
122 lake water (50:50 v/v) to avoid a drastic change in the chemical composition of the water and to
123 guarantee a food supply for the mussels during the first 24 h of acclimation. Mussel (n=60), having
124 the same shell length (15±4 mm), were placed within 5 L beakers filled with 4 L of tap and
125 deionized water (50:50 v/v), previously de-chlorinated by aeration, under a natural photoperiod with
126 constant temperature (20±1 °C), pH (7.5) and oxygenation (>90% of saturation). In order to avoid
127 the so-called tank effect, we prepare three beakers *per* treatment, including control. The bivalves
128 were fed daily with lyophilized algae *Spirulina* spp., and the water was regularly renewed every two
129 days for 2 weeks to gradually purify the mollusks by any possible pollutants that had previously
130 accumulated in their soft tissues. Only specimens that were able to re-form their byssus were used
131 in the experiments. Mussel viability was checked daily by the Trypan blue exclusion method and
132 was 93±2%, whereas biomarker baseline levels were checked weekly. Mussels were exposed to
133 MOR concentrations only when target biomarker levels were comparable with baseline ones
134 obtained in our previous laboratory studies (Parolini *et al.* 2010; 2011a,b; 2013; Parolini and Binelli
135 2013). Exposure assays were performed under semi-static conditions for 14 days. Control and
136 exposure beakers were processed at the same time and the whole water volume (4 L) was renewed
137 on a daily basis. Mussels were exposed to 0.05 µg/L (0.17 nM, Low) and 0.5 µg/L (1.7 nM, High)
138 of MOR. The first concentration was similar to the levels found in European surface waters (Pal *et*
139 *al.*, 2013), while the second one was the same tested in previous studies investigating the toxicity of

140 cocaine metabolites, benzoylecgonine (BE; Parolini *et al.*, 2013), ecgonine methyl ester (EME;
141 Parolini and Binelli, 2013), and Δ -9-tetrahydrocannabinol (Δ -9-THC; Parolini and Binelli, 2014) in
142 order to allow a comparison among drug toxicity administered at the same dose. Exact volumes of
143 working solution (10 ± 0.6 mg/L) were added daily to the exposure aquaria until reaching the
144 selected concentrations. Specimens were fed 2 h before the daily change of water and chemicals to
145 avoid the adherence of the drugs to food particles and to prevent the reduction of their
146 bioavailability. Every 3 days, 8 specimens were randomly collected from each tank (24 specimens
147 *per* treatment) to evaluate MOR-induced sub-lethal effects. Hemolymph was withdrawn by 10
148 bivalves and cyto-genotoxicity was evaluated on hemocytes. After the withdrawal, the soft tissue of
149 mussels was immediately frozen in liquid nitrogen and stored at -80 °C until LPO and PCC
150 analyses. Lastly, the soft tissue of the other 14 specimens was frozen in liquid nitrogen and stored at
151 -80 °C until the enzymatic activity was measured. Simultaneously, 10 zebra mussels were placed in
152 other control and exposure 500 mL beakers (three replicates *per* treatment), maintained at the same
153 condition described above and exposed at the same concentrations to assess the variation in
154 filtration rate due to MOR treatments.

155

156 2.2 Evaluation of MOR concentrations

157 In order to guarantee the reliability of the experimental design, the MOR concentration in both
158 working solution and exposure beakers were measured. At each time of biomarker analysis, water
159 was sampled 1 h after the contamination from both the three control and exposure beakers and
160 integrated in a unique sample (100 mL) per treatment. Water samples were spiked with 0.2 μ g/L of
161 MOR-D₃ as internal recovery standard. The concentration of the MOR was checked in LC-MS/MS
162 by using a HCT Ultra (Bruker, Germany) using a Phenomenex Luna PFP (2 x 50 mm-5 μ m) column
163 after purification and concentration by SPE (HLB 1 cm³, Waters). After cartridge activation (2 mL
164 methanol and 3 mL of water), 5 mL of each sample with internal standard were load on SPE and
165 then resuspended with 50 μ L of water. 20 μ L of each sample were then injected and analyzed in LC-
166 MS/MS. MOR quantification in water was performed by a calibration curve (0.025 - 1 μ g/L;
167 $R^2=0.99$) and internal standard recoveries were $>90\%$. The analysis of MOR concentration in
168 working solution, control and exposure beakers was performed in triplicate.

169

170 2.3 Biomarkers of cytotoxicity

171 The NRRA was performed to assess cytotoxicity following the method proposed by Lowe and Pipe
172 (1994) and applied on mussel hemocytes. Slides were examined systematically thereafter at 15 min
173 intervals to determine at what point in time there was evidence of dye loss from the lysosomes to

174 the cytosol. Tests finished when dye loss was evident in at least 50% of the hemocytes. The mean
175 retention time was then calculated from five replicates.

176

177 2.4. Oxidative stress biomarkers

178 The activity of SOD, CAT, GPx, and GST was measured in triplicate (n=3) in the cytosolic fraction
179 extracted from a pool of three whole mussels (≈ 0.3 g fresh weight) homogenized in 100 mM
180 phosphate buffer (pH 7.4; KCl 100 mM, EDTA 1 mM) with dithiothreitol (DTT, 100 mM) using a
181 Potter homogenizer. Specific protease inhibitors (1:10) were also added to the buffer:
182 phenanthroline (Phe, 10 mM) and trypsin inhibitor (Try, 10 mg/mL). The homogenate was
183 centrifuged at 15.000 g for 1 hour at 4 °C. The sample was held in ice and immediately processed
184 for the determination of protein and enzymatic activities. The total protein content of each sample
185 was determined according to the Bradford method (1976) using bovine serum albumin as a
186 standard. Enzymatic activities were determined spectrophotometrically as described by Orbea *et al.*
187 (2002). Briefly, the CAT activity was determined by measuring the consumption of H₂O₂ at 240 nm
188 using 50 mM of H₂O₂ substrate in 67 mM potassium phosphate buffer (pH 7). The SOD activity
189 was determined by measuring the degree of inhibition of cytochrome c (10 μ M) reduction at 550
190 nm by the superoxide anion generated by the xanthine oxidase (1.87 mU/mL)/hypoxanthine (50
191 μ M) reaction. The activity is given in SOD units (1 SOD unit=50% inhibition of the xanthine
192 oxidase reaction). The GPx activity was measured by monitoring the consumption of NADPH at
193 340 nm using 0.2 mM H₂O₂ substrate in 50 mM potassium phosphate buffer (pH 7) containing
194 additional glutathione (2 mM), sodium azide (NaN₃; 1 mM), glutathione reductase (2 U/mL), and
195 NADPH (120 μ M). Lastly, the GST activity was measured by adding reduced glutathione (1 mM)
196 and 1-chloro-2,4 dinitrobenzene in phosphate buffer (pH 7.4) to the cytosolic fraction; the resulting
197 reaction was monitored for 1 min at 340 nm. LPO and PCC were measured in triplicate (n=3) from
198 a pool of three whole mussels (≈ 0.3 g fresh weight) homogenized in 50 mM phosphate buffer (pH
199 7.4; KCl 100 mM, EDTA 1 mM) containing 1 mM DTT and 1 mM PMSF using a Potter
200 homogenizer. LPO level was assayed by the determination of thiobarbituric acid-reactive substances
201 (TBARS) according to Ohkawa (1979). The absorbance was read at 532 nm after removal of any
202 fluctuated material by centrifugation. The amount of thiobarbituric acid reactive substances
203 (TBARS) formed was calculated by using an extinction coefficient of 1.56×10^5 M/cm and
204 expressed as nmol TBARS formed/g fresh weight. For carbonyl quantification the reaction with
205 2,4-dinitrophenylhydrazine (DNPH) was employed according to Mecocci *et al.* (1999). The
206 carbonyl content was calculated from the absorbance measurement at 370 nm with the use of molar
207 absorption coefficient of 22 000 mol/cm and expressed as nmol/(mg protein).

208 2.5 Genotoxicity biomarkers

209 Since methods and procedures of genotoxicity biomarkers applied in this study were described in
210 detail by Parolini *et al.* (2010), only a brief description of the followed techniques was reported
211 here. The alkaline (pH>13) SCGE assay was performed on hemocytes according to the method
212 adapted for the zebra mussel by Buschini *et al.* (2003). Fifty cells *per* slide were analyzed using an
213 image analysis system (Comet Score[®]), for a total of 500 analyzed cells per specimen (n=10). Two
214 SCGE assay end-points were evaluated: the ratio between migration length and comet head
215 diameter (LDR) and the percentage of DNA in tail. The apoptotic cell frequency was evaluated
216 through the protocol described by Singh (2000). Two hundred cells per slide were analyzed for a
217 total of 1000 cells per sample (n=5). The MN test was performed according to the method of
218 Pavlica *et al.* (2000). Four hundred cells were counted per each slide (n=10) for a total of 4000
219 cells/treatment. Micronuclei were identified by the criteria proposed by Kirsch-Volders *et al.*
220 (2000), and the MN frequency was calculated (MN‰).

221

222 2.6 Filtration rate

223 Mussel filtration rate was measured according to a procedure adapted from Faria *et al.* (2009) and
224 Palais *et al.* (2012) and based on the loss of neutral red dye particles from the water column as a
225 result of mussel filtration activity (Coughlan, 1969). 10 zebra mussels per beaker (three replicates
226 per treatment) were placed in 500 mL and were exposed to 0.05 µg/L and 0.5 µg/L MOR
227 concentrations. Every 3 days for 14 days a 250 µg/L neutral red solution was added in the beakers.
228 After 15 min of acclimation, bivalve were allowed to filter for 3 h in the dark at 20 °C. Dye particle
229 concentration in the test beakers was then measured spectrophotometrically (340 nm) at the
230 beginning and at the end of the experiment, using a standard curve for neutral red solution. The
231 filtration rate (f), expressed in mL of water *per* individual and *per* hour (mL/ind/h), was calculated
232 using the following formula:

$$233 \quad f = [V / (n * t)] * \log (C_0 / C_t)$$

234

235 where V is the volume (mL) of the dye solution in the beaker, n the number of mussels, t the
236 duration of the filtration period (h), C₀ and C_t the initial and final dye particle concentrations in the
237 beaker (Coughlan, 1969).

238

239 2.7 Statistical analysis

240 Data normality and homoscedasticity were verified using the Shapiro-Wilk and Levene's tests,
241 respectively. To identify dose/effect and time/effect relationships a two-way analysis of variance

242 (ANOVA) was performed using time and MOR concentrations as variables, while biomarker end-
243 points served as cases. The ANOVA was followed by a Fisher LSD post-hoc test to evaluate
244 significant differences (* $p < 0.05$; ** $p < 0.01$) between treated samples and related controls (time to
245 time), as well as among exposures. All statistical analyses were performed using the STATISTICA
246 7.0 software package.

247

248 3. RESULTS

249 3.1 MOR concentration in exposure tanks

250 The MOR concentration in the control tanks was $< 0.025 \mu\text{g/L}$, which is the limit of detection of the
251 used equipment. In the exposure tanks, MOR level was close to the nominal values ($0.05 \mu\text{g/L}$ and
252 $0.5 \mu\text{g/L}$). We found an average value of $0.045 \pm 0.005 \mu\text{g/L}$ for the lower concentration and a value
253 of $0.35 \pm 0.01 \mu\text{g/L}$ for the higher tested concentration, accounting for the 90% and 70% of the
254 nominal values, respectively. Considering that the coefficient of variation of the method was $\pm 20\%$,
255 our analyses confirmed the reliability of the whole experimental design.

256 3.2 Baseline levels of applied biomarkers

257 During the 14-day experiment, very low mortality was observed in the control (0.6%) and exposure
258 ($< 3\%$) tanks. Baseline levels of cyto-genotoxic and oxidative stress biomarkers were similar to
259 those obtained in our previous laboratory studies (Binelli *et al.*, 2009a,b; Parolini *et al.* 2010;
260 2011a,b; 2013; Parolini and Binelli 2011; 2013). The baseline filtration rate of zebra mussel
261 specimens ranged between 0.74 ± 0.29 and $1.92 \pm 0.17 \text{ mL/individual/h}$, according to values measured
262 in the same species with a similar method by Palais *et al.* (2012).

263

264 3.3 Biomarkers of cytotoxicity

265 The NRRA showed a significant destabilization of hemocytes lysosome membranes (Figure 1)
266 according to time- ($F=27.87$, $p < 0.01$) and concentration-dependent ($F=48.56$, $p < 0.01$) relationships.
267 Both the MOR concentrations were able to significantly increase generic cellular stress in mollusks:
268 $0.05 \mu\text{g/L}$ treatment caused a significant decrease ($p < 0.01$) of NRRT starting to 11 days of
269 exposure, while at $0.5 \mu\text{g/L}$ a significant destabilization ($p < 0.01$) of the lysosomal membranes was
270 notice as early as 7 days of exposure.

271

272 3.4 Biomarkers of oxidative stress

273 The activity of antioxidant enzymes (SOD, CAT and GPx) and detoxification enzymes (GST)
274 showed some significant changes compared to controls during the exposure tests (Figure 2A, B, C,
275 D). Although the activity of GST was not significantly altered ($p > 0.05$) after exposure to the two

276 MOR tested concentrations (Figure 2A), the activity of SOD showed a significant time-dependent
277 ($F=3.07$, $p<0.05$) and dose-dependent ($F=11.54$, $p<0.01$) inhibition already after 4 days of exposure
278 to 0.5 $\mu\text{g/L}$ (Figure 2B), reaching at the end of the test values lower than 40% compared to controls.
279 Regarding the GPx, a significant time- ($F=4.64$, $p<0.01$) and concentration-dependent ($F=10.46$,
280 $p<0.01$) increase was noticed after 7 days of exposure at the highest MOR concentration (Figure
281 2C). Accordingly, CAT showed significant activity increase ($p<0.05$) compared to baseline values
282 (Figure 2D) at the end of the exposure to both the treatments. Finally, significant time-dependent
283 ($F=5.37$; $p<0.01$) and concentration-dependent ($F=4.36$; $p<0.05$) differences were found in the lipid
284 peroxidation levels (Figure 3A) with an increase of 15% compared to controls at the end of
285 exposure. In contrast, no significant differences ($p>0.05$) of protein carbonylation compared to
286 baseline levels were found, except for a single value obtained after 11 days at the lowest treatment
287 (Figure 3B).

288

289 *3.5 Biomarkers of genotoxicity and filtration rate*

290 Even if no significant increase in LDR parameter was found (data non shown), significant increase
291 in DNA fragmentation was noticed at the end of the 0.5 $\mu\text{g/L}$ exposure, as pointed out by the raise
292 of percentage of DNA in the hemocyte comet tail with respect to control ($F= 2.57$, $p<0.05$; Figure
293 4A). No significant increase ($p>0.05$) in frequencies of apoptotic (Figure 4B) and micronucleated
294 cells (Figure 4C), whose levels were similar to the baseline ones throughout the test, even if a
295 significant time-dependent relationship ($F=2.91$, $p<0.05$) was noticed for the latter end-point.
296 Lastly, the filtration rate (ranged between 1.43 ± 0.25 and 3.11 ± 0.28 mL/individual/h for 0.05 $\mu\text{g/L}$
297 MOR concentration and ranged between 1.19 ± 0.17 and 3.05 ± 0.66 for 0.5 $\mu\text{g/L}$) followed a bell-
298 shaped curves for both MOR tested concentrations, with no significant changes ($p>0.05$) compared
299 to controls (data not shown).

300

301 4. DISCUSSION

302 *4.1 Sub-lethal effects of MOR*

303 Although the lowest MOR concentration did not induced any significant variation ($p>0.05$) for all
304 the investigated end-points, the exposure to 0.5 $\mu\text{g/L}$ MOR caused a notable cytotoxicity to *D.*
305 *polymorpha* specimens, as pointed out by the significant ($p<0.01$) time- and concentration
306 dependent decrease of NRRT (Figure 1). This showed a progressive aggravation of the bivalve
307 health status, suggesting that bivalves suffer a situation of general cellular stress, which could be
308 linked to the induction of oxidative stress (Lowe *et al.* 1995). The destabilization of lysosome
309 membranes in aquatic organisms could be caused by the production of reactive oxygen species

310 (ROS) following exposure to pollutants (Regoli *et al.*, 1998). The ROS are mainly produced as side-
311 products of oxygen metabolism and among these the biotransformation of xenobiotics is an
312 oxidative process in which the production of ROS and the formation of more polar (reactive)
313 intermediates occur (Gagné *et al.*, 2010). In vertebrates, MOR is biotransformed by cytochrome
314 P450 3A4 and 2C19, which involves oxidative N-dealkylation, hydroxylation and conjugation to
315 glucuronide (Charney *et al.*, 2001). Even if no information regarding the biotransformation of MOR
316 in the zebra mussel is currently available, the observed trends of antioxidant enzymes suggested that
317 this drug could induce the production of ROS. Variation in antioxidant levels, in fact, indirectly
318 supply information on the changes of pollutant-induced reactive oxygen species (ROS) levels in
319 different aquatic organisms (Viarengo *et al.* 2007), including *D. polymorpha* (Parolini *et al.*, 2010;
320 2013). The inhibition of SOD activity (Figure 2B) suggested an increase of superoxide anion in
321 bivalves (O^{2-} ; Verlecar *et al.*, 2008), as found in our previous study exposing zebra mussel to
322 cocaine metabolites (Parolini *et al.*, 2013; Parolini and Binelli, 2013). Since dismutation of O^{2-} leads
323 to the production of hydrogen peroxide, this particular trend could be due to a phenomenon of
324 product inhibition, according to a negative feedback mechanism (Vlahogianni and Valavanidis,
325 2007). The inhibition of SOD should therefore indicate both an accumulation of O^{2-} and an
326 overproduction of H_2O_2 , which could be also produced through the spontaneous conversion of
327 superoxide anion mediated by non-enzymatic pathways (Gwoździński *et al.*, 2010). The significant
328 time-dependent trend in the levels of GPx and CAT (Figure 2C and D) confirmed that MOR was
329 able to increase the levels of H_2O_2 , whose toxicity seems to be counterbalanced by the antioxidant
330 shield of bivalves. However, it is important to note that the accumulation of superoxide radical
331 caused by SOD inhibition, combined with the increase of H_2O_2 caused by the activation of CAT and
332 GPx, could lead to the formation of hydroxyl radicals through the Haber-Weiss reaction, with the
333 consequent increase in the levels of lipid peroxidation and protein carbonylation (Verlecar *et al.*,
334 2008). The analysis of lipid peroxidation levels and protein carbonyl content just partially
335 confirmed this hypothesis, since we observed a significant ($p<0.01$) increase in the levels of lipid
336 peroxidation (Figure 3A), despite no variations in protein carbonylation were noticed (Figure 3B).
337 Accordingly, MOR treatments caused negligible genotoxic effects to zebra mussels, since just slight
338 significant ($p<0.01$) increase of DNA fragmentation was found at the end of exposure to 0.5 $\mu\text{g/L}$
339 (Figure 4A). Although several studies showed that the increase in DNA fragmentation is one of the
340 main factor leading to the onset of fixed genetic damage in *D. polymorpha* specimens (Binelli *et al.*,
341 2009a,b; Parolini and Binelli, 2012), no significant ($p>0.05$) increases in frequency of apoptotic
342 cells and MN were found (Figure 4B and 4C). Lastly, even if our data showed that low MOR
343 concentrations could alter antioxidant status of the zebra mussel leading to low oxidative damage,

344 no physiological effect was noticed, as pointed out by the lack of significant ($p>0.05$) alteration of
345 bivalve filtration rate (data not shown). Despite of the moderate MOR-induced adverse effects
346 found in the zebra mussel, the potential toxicity of this illicit drug cannot be neglected since it could
347 cause other deleterious effects that further studies should have to investigate. For instance, being a
348 psychotropic substance, MOR could act as neurotoxic compound, as pointed out by a recent *in vivo*
349 study on the freshwater bivalve *Elliptio complanata* in which the exposure to three MOR
350 concentrations (0.07; 0.15 and 0.75 mg/g wet weight) induced decreases in serotonin and AChE,
351 and increases in dopamine and GABA levels, suggesting the induction of a relaxation state in
352 mussels (Gagnè *et al.*, 2010).

353

354 4.2 Comparison of MOR toxicity with the effects of other illicit drugs

355 Considering the moderate sub-lethal effects of MOR to treated zebra mussel, the comparison
356 between its toxicity and that of previously analyzed illicit drugs, namely benzoylecgonine (BE;
357 Parolini *et al.*, 2013), ecgonine methyl ester (EME; Parolini and Binelli, 2013) and Δ -9-
358 tetrahydrocannabinol (Δ -9-THC; Parolini and Binelli, 2014), should allow the drawing of a toxicity
359 scale to individuate the most dangerous compound towards our biological model, laying the bases
360 for further in-depth investigations. Although it is well-known that the assessment of several
361 biomarkers is the best approach to the understanding of adverse effects and mechanism of action of
362 pollutants on organism (Viarengo *et al.*, 2007; Sforzini *et al.*, 2011), the simple examination of the
363 simultaneous changes of dissimilar biological parameters is considered insufficient to rank the
364 hazard of different pollutants because of the wide variability in biomarker responses. For example,
365 by comparing present data and those from our experiments on the sub-lethal effects induce by other
366 illicit drugs (Parolini *et al.*, 2013; Parolini and Binelli, 2013; 2014) we can note a remarkable
367 variability in most of investigated end-points, depending on tested compound and probably due to
368 dissimilarities in their mechanism of action, which prevents from ranking their toxicity. In fact,
369 even though all psychotropic substances were able to induce significant destabilization of lysosome
370 membranes, substantial differences in the activity of antioxidant/detoxification enzymes, as well as
371 in oxidative and genetic damage were found. For this reason, the application of procedures able to
372 integrate the biomarker responses within a simple synthetic index could help to minimize the
373 variation of responses, allowing to draw an accurate scale of toxicity. To compare the sub-lethal
374 toxicity of tested illicit drugs we integrated the whole biomarker dataset obtained at 0.5 μ g/L for
375 MOR into a synthetic index called Biomarker Response Index (BRI), previously described by
376 Parolini *et al.* (2013). We excluded from integration analysis the results from filtration rate since it
377 is not assessed in previous studies. Briefly, since changes in each specific biomarkers follow

378 different trends (increasing, decreasing or bell-shaped curves, Hagger *et al.*, 2010), we calculated
379 the percentage of alteration level (AL) of each biomarker *per* exposure time compared to the
380 correspondent control. To calculate the BRI, we attributed a specific score to each obtained AL
381 value according to Parolini *et al.* (2013) and each biomarker was then weighted in relation to its
382 level of biological organization (Hagger *et al.*, 2010). Finally, we compared the BRI value obtained
383 for MOR with those calculated for BE and EME (Parolini and Binelli, 2013), and Δ -9-THC
384 (Parolini and Binelli, 2014), deriving the following toxicity scale:

385

386

$$\Delta\text{-9-THC} > \text{BE} \approx \text{EME} \gg \text{MOR}$$

387

388 The toxicity of Δ -9-THC (BRI=8.78; Parolini and Binelli, 2014) seems to be slightly higher than
389 that of BE (BRI=8.22) and EME (BRI=8.06; Parolini and Binelli 2013), while MOR (BRI=6.17)
390 showed the lowest value among the tested illicit drugs, suggesting its possible lowest hazard
391 towards the zebra mussel for the measured end-points at least.

392

393 5. CONCLUSION

394 Our findings showed that MOR exposure could induce moderate adverse effects to this freshwater
395 bivalve species, highlighting its possible hazard to freshwater communities. Even if current
396 environmental MOR levels seem not cause any deleterious effect to bivalves, 14-day treatment to
397 0.5 $\mu\text{g/L}$ concentration affected the oxidative status of bivalves and induced slight oxidative damage
398 to cellular macromolecules. Although our findings suggest that oxidative stress seems to be
399 involved in the mechanism of action of MOR in zebra mussel, further studies using powerful
400 techniques, as well as the analysis of different end-points (i.e. neurotoxicity parameters), should be
401 necessary to confirm it. Although the integrated MOR toxicity obtained at 0.5 $\mu\text{g/L}$ treatment
402 resulted lower compared to that of other common illicit drugs previously tested at the same
403 experimental conditions, its environmental hazard cannot be underestimated. In fact, considering
404 that in the real environment organisms are exposed to MOR concentrations for their whole life span,
405 and its levels could increase due to the stable use of opiates worldwide, MOR effects could be more
406 deleterious with respect to those highlighted in the present study. For this reason, further
407 investigations should be necessary to enhance knowledge on MOR sub-lethal effects and its
408 mechanism of action in non-target aquatic organisms.

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566

567 FIGURES:

568 Fig. 1: Assessment of lysosomal membrane stability (Neutral Red Retention Time-mean \pm SEM)
569 found in the hemocytes of treated bivalves (n=5). Asterisks indicate significant differences between
570 the treated and the corresponding controls (two-way ANOVA, Fisher LSD post-hoc test, *p<0.05,
571 **p<0.01).

572

573 Fig. 2: Mean values (\pm SEM) of the activity of glutathione-S-transferase (GST, A), superoxide
574 dismutase (SOD, B), glutathione peroxidase (GPx, C) and catalase (CAT, D), measured in the
575 bivalves (n=3, pool of 3 individuals) exposed to both MOR concentrations. The significant
576 differences (two-way ANOVA, Fisher LSD post-hoc test, *p<0.05; **p<0.01) refer to the
577 comparison of exposed with the corresponding baseline value.

578

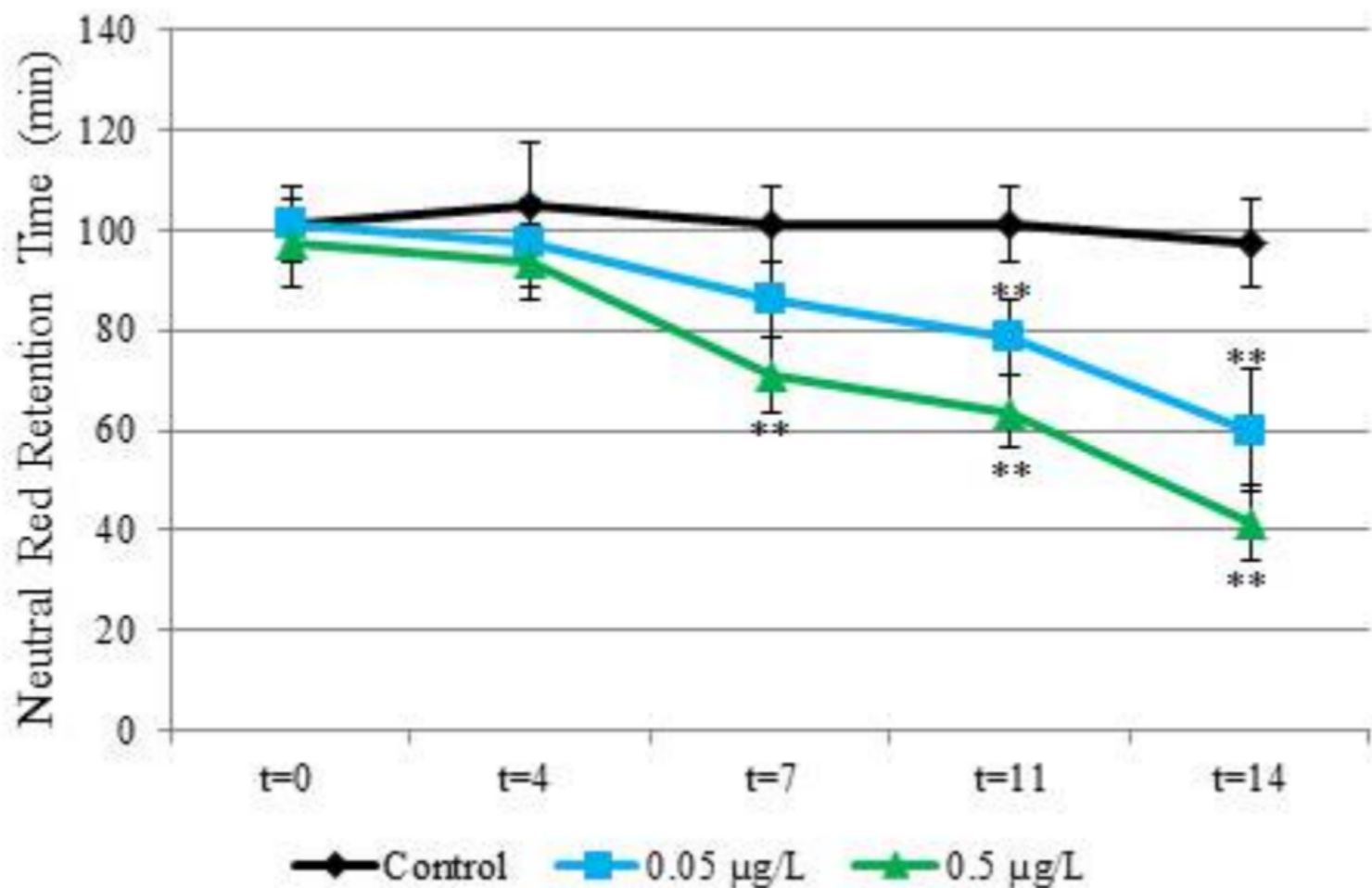
579 Fig. 3: Mean values (\pm SEM) of lipid peroxidation levels (LPO, A) and protein carbonylation (PCC,
580 B) found in the bivalves (n=3, pool of 3 individuals) exposed to both MOR concentrations. The
581 significant differences (two-way ANOVA, Fisher LSD post-hoc test, * $p < 0.05$, ** $p < 0.01$) relate to
582 the comparison between the exposed and corresponding baseline value.

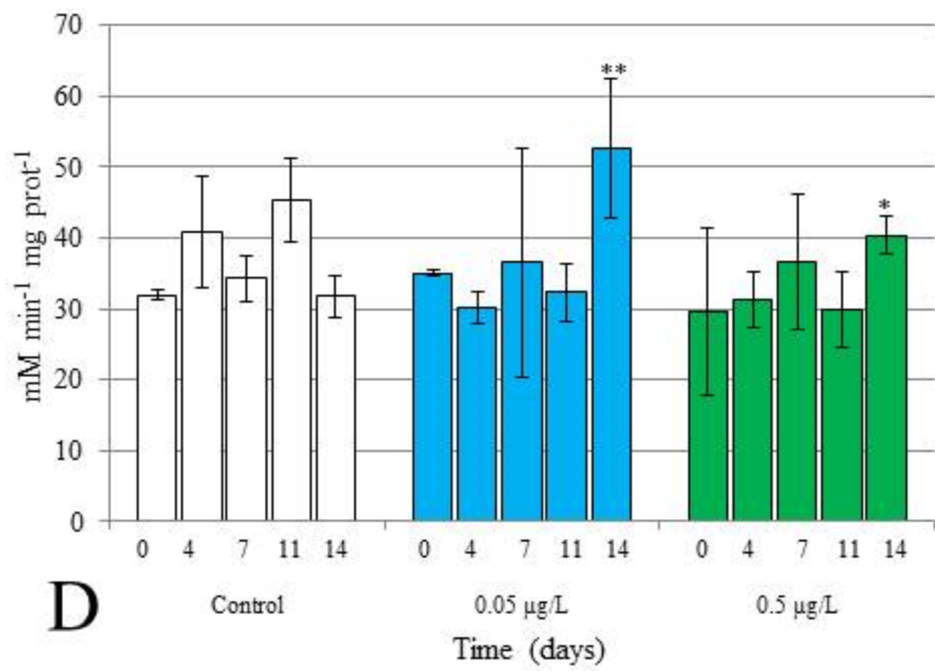
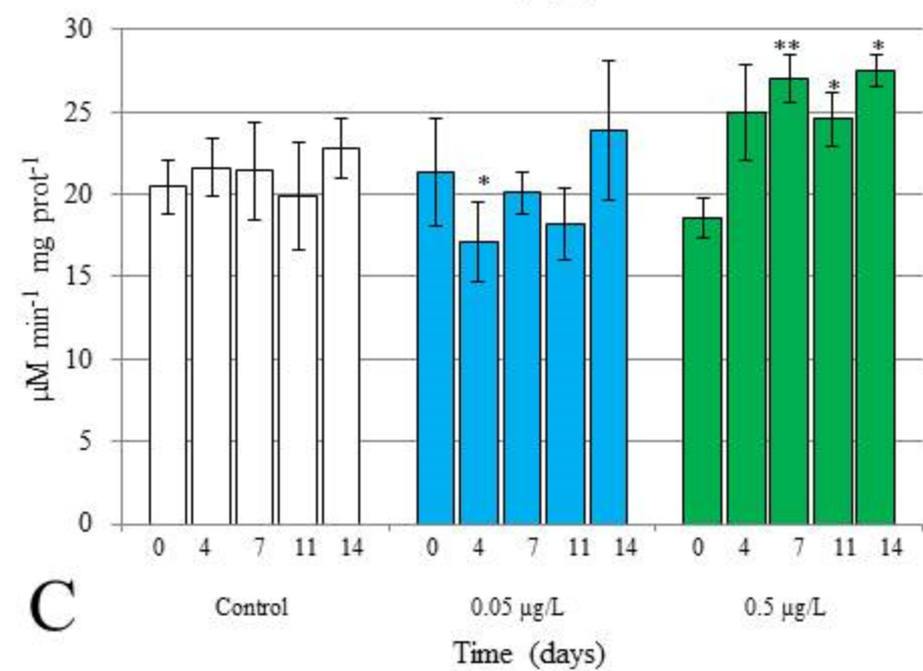
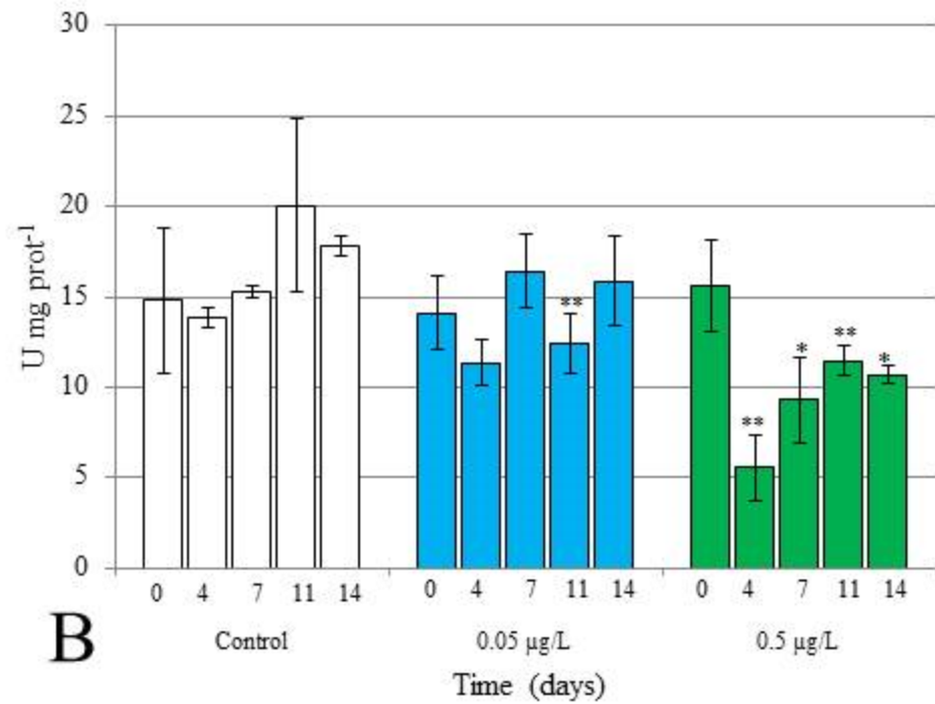
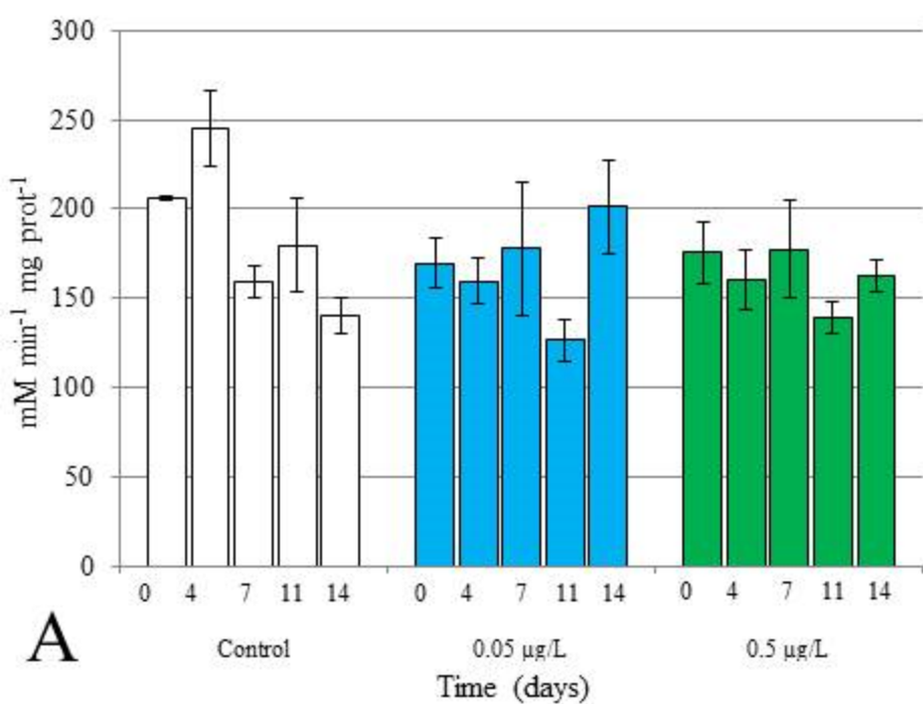
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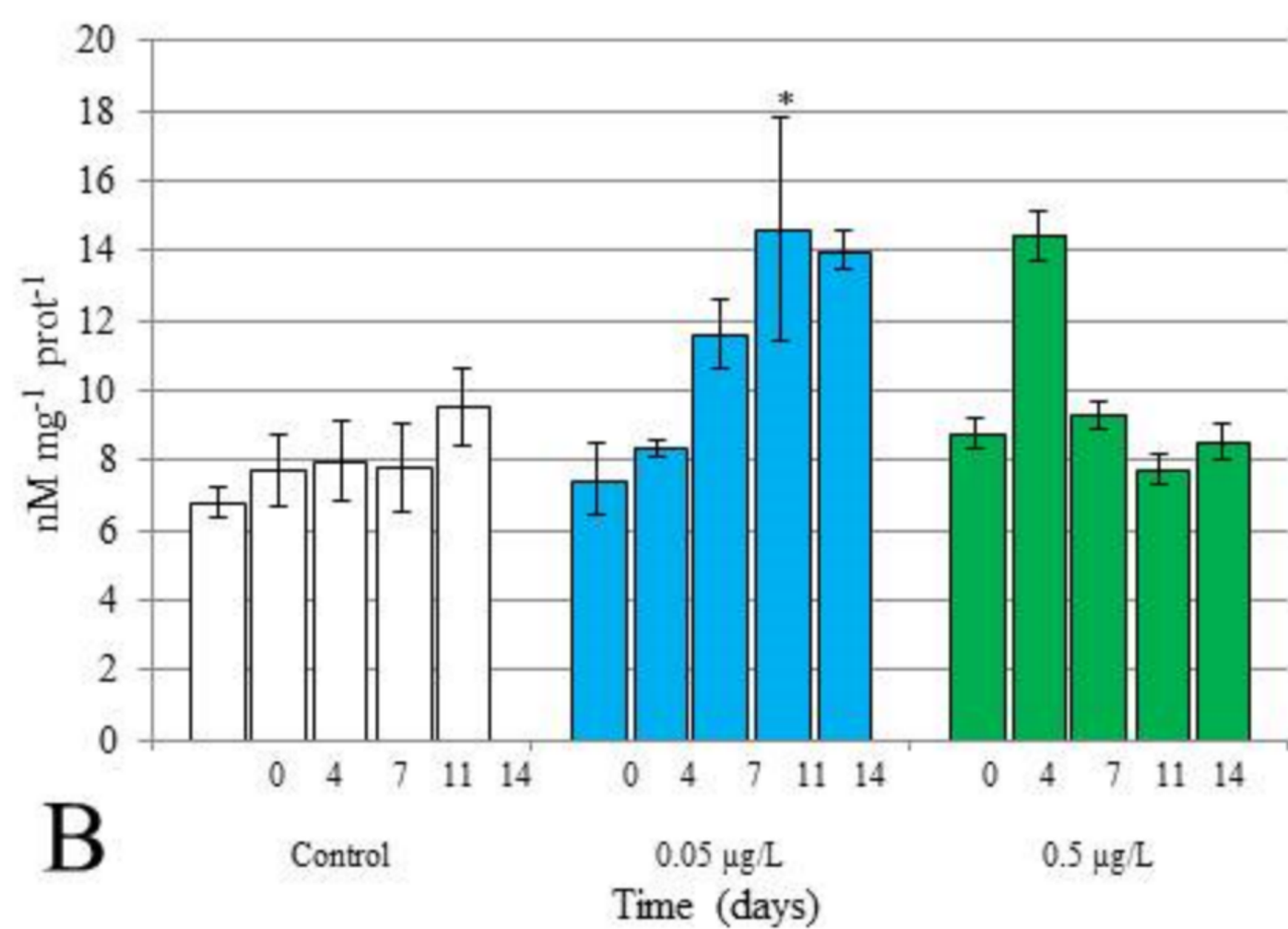
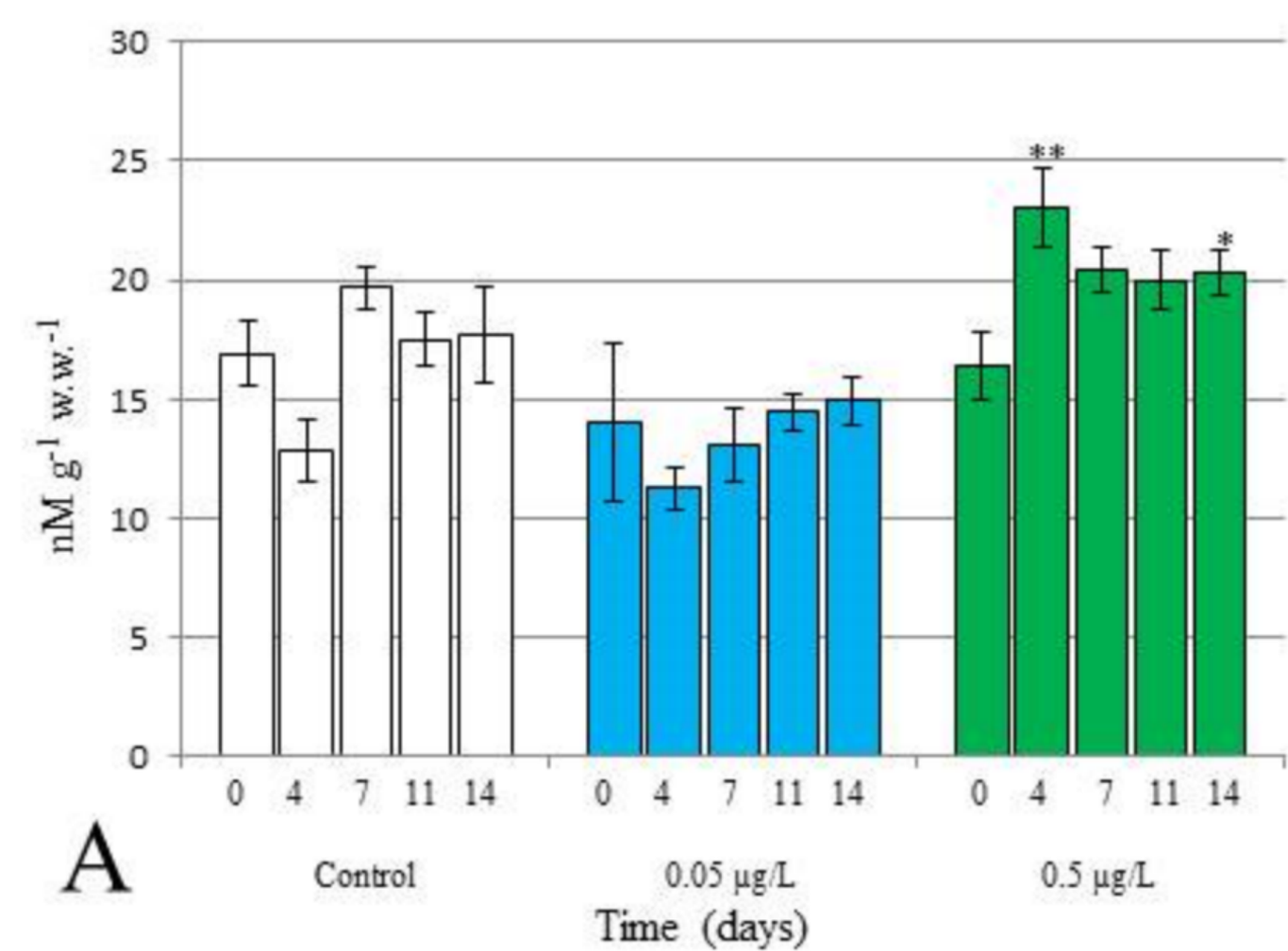
584 Fig. 4: DNA percentage in the comet tails (A; mean \pm SEM) of the bivalve hemocytes (n=8) exposed
585 to both MOR concentrations. The significance (two-way ANOVA, Fisher LSD post-hoc test,
586 ** $p < 0.01$) refers to the comparison between the exposed and controls. Mean (\pm SEM) apoptotic
587 cells frequency (%) (B) and micronucleated frequency (%MN; C) showed no significant
588 differences compared to the corresponding controls (two-way ANOVA, $p > 0.05$).

589

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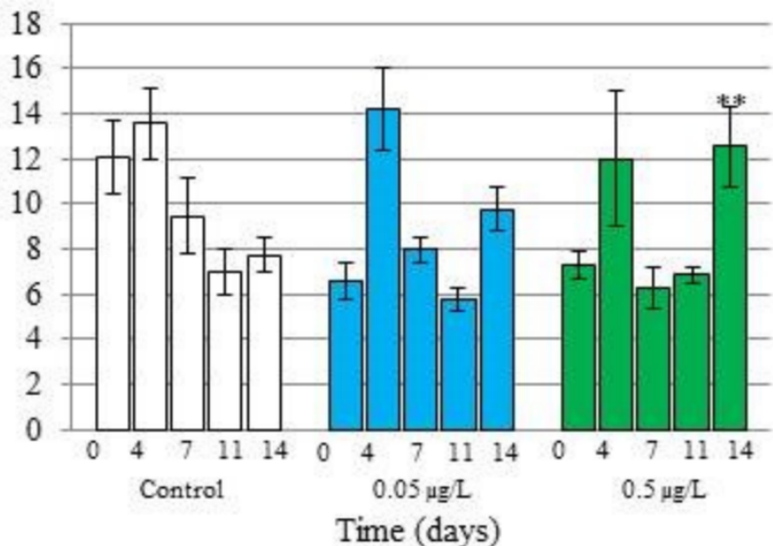




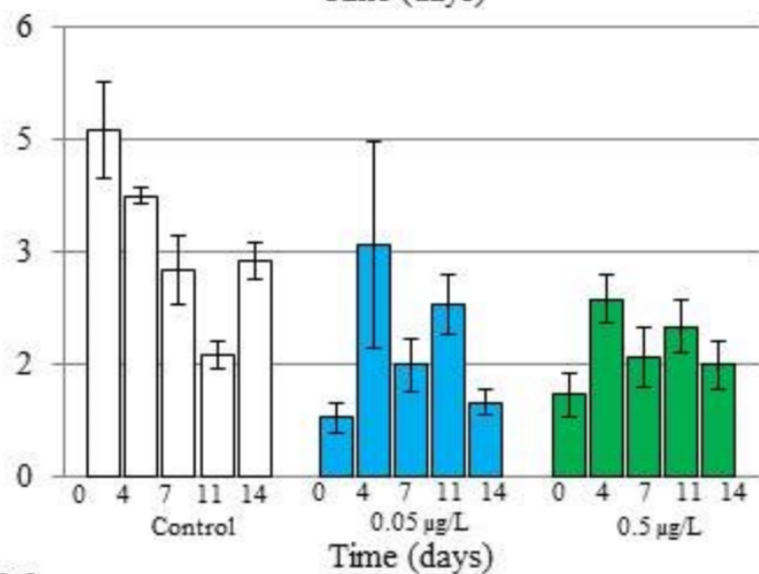


A

Mean of the % DNA (in the comet tail)

**B**

Apoptotic cell frequency (%)

**C**

Frequency of micronuclei (%)

