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Toxicological effects of the rare earth element neodymium in *Mytilus galloprovincialis*

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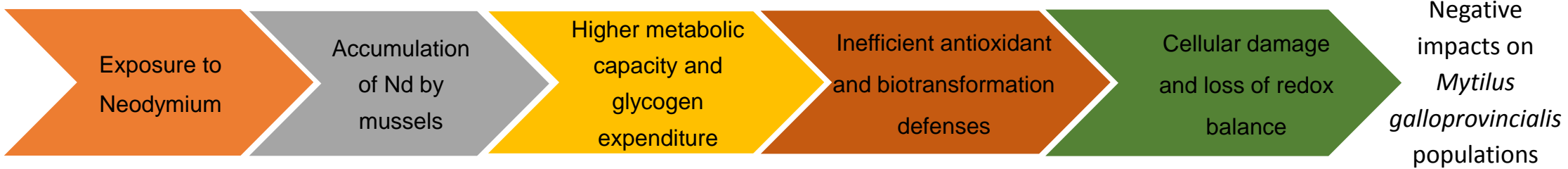
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1 Toxicological effects of the rare earth element Neodymium in

2 *Mytilus galloprovincialis*

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22 **ABSTRACT**

23 The wide range of applications of rare earth elements (REE) is leading to their
24 occurrence in worldwide aquatic environments. Among the most popular REE is
25 Neodymium (Nd), being widely used in permanent magnets, lasers, and glass additives.
26 Neodymium–iron–boron (NdFeB) magnets is the main application of Nd since they are
27 used in electric motors, hard disk drives, speakers and generators for wind turbines.
28 Recent studies have already evaluated the toxic potential of different REE, but no
29 information is available on the effects of Nd towards marine bivalves. Thus, the present
30 study evaluated the biochemical alterations caused by Nd in the mussel *Mytilus*
31 *galloprovincialis* exposed to this element for 28 days. The results obtained clearly
32 demonstrated that Nd was accumulated by mussels, leading to mussel's metabolic
33 capacity increase and GLY expenditure, in an attempt to fuel up defense mechanisms.
34 Antioxidant and biotransformation defenses were insufficient in the elimination of ROS
35 excess, resulting from the presence of Nd and increased electron transport system
36 activity, which caused cellular damages (measured by lipid peroxidation) and loss of
37 redox balance (assessed by the ratio between reduced and oxidized glutathione). The
38 results obtained clearly highlight the potential toxicity of REEs, and in particular of Nd,
39 with impacts at cellular level, which may have consequences in mussel's survival,
40 growth and reproduction, affecting mussel's population.

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43

44 **Keywords:** Rare earth elements, mussels, oxidative stress, metabolic capacity,
45 bioconcentration.

46

1. INTRODUCTION

47 **1. INTRODUCTION**

48 The extensive use of neodymium (Nd, $Z = 60$) by the industries has drawn

49 attention from the scientific community in the last years. This light rare earth element is

50 classified as one of the five most critical rare earth elements (REEs) until 2025 (U.S.

51 Department of Energy, 2011) due to its high economic importance and supply risk

52 (Batinic et al., 2017; Critical Raw Materials - European Commission Report, 2018).

53 Currently, Nd main application is related to permanent magnets (PMs) based on NdFeB,

54 whose amount is increasing every year. Neodymium magnets are the strongest type of

55 permanent magnet commercially available. It was announced that the annual use of

56 these magnets increased from 20 thousand tons in 2006 to almost 55 thousand tons in

57 2017 (Liu et al., 2019), with three electric and electronic components containing PMs:

58 hard disk drives, small electric motors and speakers. The results show that the weight

59 percentage of the PMs varies from 4 to 6% in the speakers, 2.5 to 2.8% in the hard

60 disks, and between 0.8 and 2% in some electric motors from hybrid electric vehicles

61 (Menad and Seron, 2017). Despite these low representative numbers, there were

62 generated, in 2018, 50 million tons of electric and electronic waste (UNEP et al., 2019).

63 These numbers are concerning the governments and industry analysts regarding future

64 prices and availabilities (Rabe et al., 2017) but also concerning environmental impact of

65 this element in aquatic systems. Permanent magnets are also used in wind turbines,

66 missiles, tanks, warplanes and submarines (Padhan et al., 2017). An advantage of using

67 these magnets over alternative technologies in wind turbines is that they reduce the

68 turbine's size and decreases the overall weight (Rabe et al., 2017). Even so, typically

69 wind turbines contain about 150–200 kg of Nd per megawatt of generating capacity

70 (Hatch, 2012), which means that most offshore wind turbines may require two tons of

71 this REE (Turra, 2018).

72 The increase application of Nd in high-tech processes and products leads to the
73 release of this element into the environment, mainly in the rivers and coastal areas not
74 only due to the disposal of e-waste (50 million tons in 2018) but also from the mining
75 activities which is the primary source of REEs discharge into water systems (Adeel et
76 al., 2019; UNEP et al., 2019). The concentration of Nd in waters depends on several
77 factors such as climate, geology and vegetation and its most common oxidation state is
78 Nd(III). A wide range of REEs concentrations has been detected in agricultural soils (<
79 15.9-249.1 $\mu\text{g/g}$) and in groundwater (< 3.1-146.2 $\mu\text{g/L}$) at various sites worldwide
80 (Adeel et al., 2019), with Nd being the third element most abundant (after Ce and La).
81 The concentration of REEs in soils and sediment is higher than in water resources due
82 to pH and cationic exchange capacity. This occurs because most REEs may adsorb to
83 soils and sediments through their dissolution and surface complexation reactions with
84 inorganic and organic ligands (Gwenzi et al., 2018). The concentration of Nd in the
85 environment is in the order of ng/L: 2.8 ng/L in seawater (Tai et al., 2010), 0.76 – 15
86 ng/L in rain water, 16.9 ng/L in throughfall, 58 ng/L in aqueous phase of the soil and
87 84.9 ng/L in stream water (Kabata-Pendias and Mukherjee, 2007). Despite that, its
88 concentration in surface waters is about a few $\mu\text{g/L}$, while in contaminated
89 environments it increases until hundreds of $\mu\text{g/L}$. The Terengganu River Basin, in
90 Malaysia, is an example of a surface water where concentrations of Nd between 0.0071
91 and 6.68 $\mu\text{g/L}$ were detected (Sultan and Shazili, 2009). As for the contaminated
92 environments, it was detected 771 $\mu\text{g/L}$ in streams draining from acid sulphate soils
93 during high-water flow events in autumn, in Finland (Åström, 2001). In an alluvial
94 aquifer affected by acid mine drainage (Guadamar, Spain) it was detected < 0.01 –
95 52.67 $\mu\text{g/L}$ (Olías et al., 2005). It was also found a concentration of Nd of 10.8 $\mu\text{g/L}$
96 (Khan et al., 2017) and 317 $\mu\text{g/L}$ (Migaszewski et al., 2016) in the surface water of the

97 ex-mining pit lake (Malaysia) and in a mining pit in Wisniowka (Poland), respectively.
98 In coastal areas the concentration of Nd found in the coast of Hawaii (Kona) and the
99 coast of Australia (Labrador beach), revealed concentrations of Nd in seawater of 24-32
100 $\mu\text{g/L}$ (Adeel et al., 2019).

101 The increase concentration of Nd in aquatic systems may have significant impacts
102 in the organisms inhabiting these systems. Asian continent, namely in China, shows the
103 most critical risk of REEs pollution level followed by Europe, Africa, USA and
104 Australia (Adeel et al., 2019). However, there are only few papers published regarding
105 this thematic. Wang et al. (2011) evaluated the effect of Nd on a freshwater
106 cyanobacteria, *Microcystis aeruginosa*, specifically on its growth and biochemical
107 changes. The results showed that Nd(III) concentration ≤ 1 mg/L can stimulate the
108 growth of *M. aeruginosa*; also, the content of chlorophyll a (Chl-a), soluble protein and
109 the activity of antioxidant defences increased when compared with the control.
110 However, with high concentration of Nd(III) (5.00–10.00 mg/L), the growth of *M.*
111 *aeruginosa* was inhibited while increasing the content of malondiadehyde (MDA) and
112 decreased the activity of the enzyme catalase (CAT). It was also studied the Nd effects
113 on rat (liver), and the results showed an accumulation in hepatocyte nuclei and
114 mitochondria, a decrease of superoxide dismutase (SOD) and CAT, and an increase of
115 the activity of glutathione peroxidase (GPx) and lipid peroxidation levels (Adeel et al.,
116 2019; Rim et al., 2013). Regardless of the increasing presence of Nd in oceans and
117 consequently bioaccumulation in aquatic organisms, there are only a few toxicological
118 studies on organisms published in the literature. Also, those studies evaluate the effects
119 on freshwater organisms, but Nd discharged by the industries will reach the oceans.

120 Therefore, the present study aimed to evaluate the biological effects induced by
121 Nd in the mussel species *Mytilus galloprovincialis*, through the exposure of the

122 organisms to five different concentrations (2.5, 5, 10, 20 and 40 $\mu\text{g/L}$ of Nd), during
123 twenty-eight days. For this, biomarkers related with oxidative stress, redox balance and
124 metabolic status were measured in *M. galloprovincialis*, trying to identify potential
125 impacts of Nd on this species.

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126 2. MATERIALS AND METHODS

127 2.1 Experimental conditions

128 The Mediterranean mussel *Mytilus galloprovincialis* was selected as biological
129 model for the present study. Mussels with similar size (5.7 ± 0.7 cm length; 3.0 ± 0.4 cm
130 width) were collected in September 2018, at the Ria de Aveiro lagoon (Portugal). After
131 sampling mussels were placed in aquaria for depuration and acclimation to laboratory
132 conditions for 2 weeks, during which animals were maintained under constant aeration
133 in different tanks with in artificial seawater (made by reverse osmosis water and
134 artificial salt - Tropic Marin® SEA SALT) at temperature, pH and salinity values
135 resembling the sampling site conditions (18.0 ± 1.0 °C; 8.0 ± 0.1 , 30 ± 1 , respectively).
136 Seawater was renewed daily during the first week and then every three days until the
137 end of the acclimation period. After the three first days of acclimation, mussels were fed
138 with Algamac protein plus (150,000 cells/animal) after each water renewal.

139 After acclimation organisms were distributed in different aquaria, with four
140 aquaria (containing 3L of artificial seawater each) per treatment: control (CTL, 0 µg/L
141 of Nd), 2.5, 5.0, 10, 20 and 40 µg/L of Nd (6 treatments). A total of 20 mussels were
142 used per tested concentration, with 5 mussels per replicate/aquarium (total of 120
143 mussels in total). Concentrations of Nd identified in low to highly contaminated
144 environments (see references above) were considered to select test treatments.

145 To guarantee the presence of Nd in the water medium, the stability of Nd in the
146 seawater was evaluated running a preliminary experiment in the absence of mussels. For
147 this, glass containers with 500 mL of artificial seawater were spiked with 2.5 and 40
148 µg/L of Nd (10 containers per concentration) and, during seven days (corresponding to
149 the period between water renewals along the twenty-eight days experimental assay),
150 aliquots of 5 mL were daily collected to assess concentrations of Nd in the water.

151 Results concerning the stability of Nd in seawater medium showed that, in the absence
152 of mussels, concentrations were maintained along seven days' exposure period, with
153 results showing that the values after exposure to 2.5 and 40 $\mu\text{g/L}$ of Nd were,
154 respectively, 3.5 ± 0.2 and 54 ± 4.4 $\mu\text{g/L}$ of Dy. These results clearly demonstrate the
155 stability of Nd during the seven days' exposure period, the interval used between water
156 renewal along the experimental assay, allowing to perform a twenty-eight days exposure
157 period with water renewal ever week.

158

159 During the experimental period (twenty-eight days), water medium was renewed
160 weekly and exposure conditions re-established, including Nd concentrations and
161 seawater characteristics (temperature 17 ± 1 $^{\circ}\text{C}$, pH 8.0 ± 0.1 and salinity 30 ± 1). Every
162 week, immediately after seawater renewal, water samples were collected from each
163 aquarium for Nd quantification, to compare nominal and real exposure concentrations.
164 During the exposure period, organisms were fed with Algamac protein plus (150,000
165 cells/animal) three times per week. Mortality was also daily checked, with 100% of
166 survival recorded during the experimental period. During the exposure period water
167 medium in each aquarium was continuously aerated with a photoperiod of 12h light:12h
168 dark.

169 At the end of the exposure period, organisms were frozen individually with liquid
170 nitrogen and stored at -80°C , until homogenization of each individual soft tissue using a
171 mortar and a pestle under liquid nitrogen. Each homogenized organism was divided into
172 aliquots (each with 0.5 g fresh weight, FW) for biomarkers analyses and Nd
173 quantification.

174

175 2.2 Neodymium quantification in water and tissue samples

176 To guarantee that nominal and real concentrations were similar, Nd concentrations
177 in water samples, collected every week from each aquaria immediately after water
178 contamination, were quantified using inductively coupled plasma mass spectrometry
179 (ICP-MS), on a Thermo ICP-MS X Series equipped with a Burgener nebulizer after
180 adequate sample dilution and acification to pH <2. Water samples collected daily to
181 evaluate the stability of Nd in seawater (in the absence of mussels), along seven days
182 experimental period, were analysed following the same procedure.

183 Total Nd concentrations in *M. galloprovincialis* whole soft tissues (2 individuals
184 per replicate, 8 individuals per condition) were also quantified by ICP-MS, after
185 microwave assisted acid digestion. After freeze-drying, mussel samples with 100–200
186 mg were digested in a CEM MARS 5 microwave, firstly with 2 mL of HNO₃ (70%) at
187 170 °C for 15 min, followed by a second identical microwave cycle with 0.5 mL of
188 H₂O₂ (30%). After addition of H₂O₂, the mixture was allowed to stand for 15 min so
189 that the microwave reaction was not as violent. The obtained digests were transferred
190 into 25 mL polyethylene vessels and the volume made up with ultrapure water.

191

192

193 2.3 Biochemical markers

194 The whole tissue of mussels was used for biomarkers determination. For each
195 parameter, 0.5 g FW of tissue per organism was used, with 2 individuals per replicate (8
196 individuals per condition). For each condition, metabolic capacity (electron transport
197 system activity, ETS), energy reserves (glycogen content, GLY; total protein content,
198 PROT), antioxidant and biotransformation defences (activities of superoxide dismutase,
199 SOD; catalase, CAT; glutathione peroxidase, GPx; glutathione S-transferases, GSTs),
200 cellular damage (lipid peroxidation levels, LPO) and redox balance (ratio between

201 reduced glutathione and oxidized glutathione, GSH/GSSG) markers were assessed.
202 Each sample was performed at least in duplicate, i.e., from each 0.5 g samples two sub-
203 samples were measured for each biomarker to guarantee the quality of the data. All
204 measurements were done using a microplate reader (BioTek, Synergy HT). The
205 extraction for each biomarker was performed with specific buffers: phosphate buffer for
206 SOD, CAT, GSTs, PROT and GLY; magnesium sulphate buffer for ETS;
207 trichloroacetic acid buffer for LPO and KPE buffer for GSH/GSSG. Each sample was
208 sonicated for 15 s at 4 °C and centrifuged for 25 min (or 15 min for GSH/GSSG) at
209 10,000 g (or 3,000 g for ETS) (Andrade et al., 2019; Coppola et al., 2019; De Marchi et
210 al., 2018; Freitas et al., 2019). Supernatants were stored at -80 °C.

211

212 *Metabolic capacity and energy reserves*

213 The ETS activity was measured based on King and Packard (1975) and the
214 modifications performed by De Coen and Janssen (1997). Absorbance was measured
215 during 10 min at 490 nm with intervals of 25 s and the extinction coefficient $\epsilon = 15,900$
216 $\text{M}^{-1} \text{cm}^{-1}$ was used to calculate the amount of formazan formed. Results were expressed
217 in nmol per min per g of FW.

218 For GLY quantification the sulphuric acid method was used, as described by
219 Dubois et al. (1956). Glucose standards were used (0–10 mg/ mL) to produce a
220 calibration curve. Absorbance was measured at 492 nm after incubation during 30 min
221 at room temperature. Results were expressed in mg per g FW.

222 The PROT content was determined according to the spectrophotometric Biuret
223 method (Robinson and Hogden, 1940). Bovine serum albumin (BSA) was used as
224 standard calibration curve (0–40 mg/mL). Absorbance was read at 540 nm. The results
225 were expressed in mg per g FW.

226

227 *Antioxidant defences*

228 SOD activity was determined by the Beauchamp and Fridovich (1971) method
229 after adaptations performed by Carregosa et al. (2014). The standard curve was formed
230 using SOD standards (0.25-60 U/mL). Samples' absorbance was read at 560 nm after 20
231 min of incubation at room temperature. Results were expressed in U per g FW where
232 one unit (U) represents the quantity of the enzyme that catalyzes the conversion of 1
233 μmol of substrate per min.

234 CAT activity was quantified according to the Johansson and Borg (1988) method
235 and the modifications performed by Carregosa et al. (2014). The standard curve was
236 determined using formaldehyde standards (0–150 $\mu\text{mol/L}$). Absorbance was measured
237 at 540 nm. The enzymatic activity was expressed in U per g of FW, where U represents
238 the amount of enzyme that caused the formation of 1.0 nmol formaldehyde per min at
239 25 °C.

240 GPx activity was quantified following Paglia and Valentine (1967). The
241 absorbance was measured at 340 nm in 10 sec intervals during 5 min and the enzymatic
242 activity was determined using the extinction coefficient $\epsilon = 6.22 \text{ mM}^{-1}\text{cm}^{-1}$. The results
243 were expressed as U per g FW, where U represents the amount of enzyme that caused
244 the formation of 1.0 μmol NADPH oxidized per min.

245

246 *Biotransformation defences*

247 GSTs activity was quantified following Habig et al. (1974) protocol with some
248 adaptations performed by Carregosa et al. (2014). The absorbance was measured at
249 340nm and the activity of GSTs was determined using the extinction coefficient $\epsilon = 9.6$
250 $\text{mM}^{-1}\text{cm}^{-1}$. The enzymatic activity was expressed in U per g of FW where U is defined

251 as the amount of enzyme that catalysis the formation of 1 μmol of dinitrophenyl
252 thioether per min.

253

254 *Cellular damage*

255 LPO determination was done following the method described by Ohkawa et al.
256 (1979). LPO levels were measured trough the quantification of malondialdehyde
257 (MDA), a by-product of lipid peroxidation. Absorbance was measured at 535 nm and
258 the extinction coefficient $\varepsilon = 156 \text{ mM}^{-1} \text{ cm}^{-1}$ was used to calculate LPO levels,
259 expressed in nmol of MDA formed per g of FW.

260

261 *Redox balance*

262 GSH and GSSG glutathione contents were measured at 412 nm (Rahman et al.,
263 2014) and used as standards (0–60 $\mu\text{mol/L}$) to obtain a calibration curve. Absorbance
264 was measured at 412 nm, for both assays. The results were expressed as nmol per g of
265 FW. The ratio GSH/GSSG was determined taking in account the number of thiol
266 equivalents (GSH / 2*GSSG).

267

268 2.5 Data analyses

269 Bioaccumulation factor (BCF) was calculated dividing the mean Nd concentration
270 found in mussel's tissues at the end of the experimental period by the mean value of Nd
271 found in seawater immediately after spiking (corresponding to the real exposure
272 concentration).

273 All the biochemical results (ETS, GLY, PROT, SOD, CAT, GPx, GSTs, LPO and
274 GSH/GSSG) and Nd concentrations in mussel's tissues, obtained from each tested
275 treatment, were submitted to statistical hypothesis testing using permutational analysis

276 of variance, employing the PERMANOVA+add-on in PRIMER v6 (Anderson et al.,
277 2008). The pseudo-F p-values in the PERMANOVA main tests were evaluated in terms
278 of significance and when significant ($p < 0.05$) differences were observed pairwise
279 comparisons were performed among conditions. Significant differences were identified
280 in the figures with different lowercase letters.

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3. RESULTS AND DISCUSSION

With the technological advances and economic development, the multiplicity and wide variety of applications of electrical and electronic equipment have extremely increased over the last decades. Consequently, the quantity of end-of-life products is also growing, resulting into increasing amounts of hazardous substances, including REEs. As a consequence, different authors have reported the presence of these elements in aquatic systems and inhabiting organisms (Akagi and Edanami, 2017; Adeel et al., 2019; Rim, 2016; Rim et al., 2013). Considering this, recent studies have evaluated the impacts of REE in different aquatic species, trying to identify harmful effects caused by these substances (Oral et al., 2010; Adeel et al., 2019; Henriques et al., 2019; Pinto et al., 2019; Rim, 2016; Rim et al., 2013).

The present study aimed to evaluate the accumulation and effects of Nd exposure in the mussel species *Mytilus galloprovincialis*, after a chronic exposure period.

3.1 Accumulation of Neodymium in mussel's tissues

In terms of accumulation, the results obtained showed that when exposed to environmentally relevant concentrations, Nd levels in mussel's tissues increased along the increasing exposure gradient, with significant differences among all tested conditions (Table 1). However, bioconcentration factor (BCF) showed similar values among tested conditions, indicating the efforts of mussels to prevent Nd accumulation with the increasing exposure concentrations (Table 1). The present results are in accordance with previous studies, conducted under laboratory conditions, which revealed similar responses, with accumulation of different REE by marine mussels (*M. galloprovincialis*, Henriques et al., 2019; Pinto et al., 2019), freshwater mussels (*Dreissena polymorpha*, Hanana et al., 2018), and freshwater clams (*Corbicula*

308 *fluminea*, Bonnail et al., 2017). Such findings highlight the capacity of bivalves to
309 accumulate REE, which may impair their physiological and biochemical performances.
310 Nevertheless, the present study further revealed the capacity of mussels to limit the
311 accumulation of Nd, as similar accumulation rate was observed in all tested treatments,
312 regardless the concentration of exposure. These results may indicate that, along the
313 increasing exposure gradient, mussels were able to limit the Nd accumulation by
314 reducing filtration and respiration capacity and/or were able to increase the
315 detoxification of this element. Since, higher metabolic capacity was observed in
316 contaminated mussels and no differences were observed among these treatments (except
317 for the lowest Nd concentration), the results obtained may indicate that respiration and
318 filtration rates were not decreased under the exposure of Nd. Therefore, the efforts of
319 mussels to limit Nd accumulation may result from mussel's detoxification capacity.
320 Similarly, Oliveira et al. (2017) demonstrated that *M. galloprovincialis* decreased
321 bioconcentration factor (BCF) along the increasing exposure gradient of carbamazepine
322 (CBZ). It was already showed that in the presence of pollutants bivalves may limit their
323 filtration rate to avoid their accumulation, an effort that may increase with increasing
324 exposure concentration (Almeida et al., 2015; Chen et al., 2014). In particular, Chen et
325 al. (2014) reported a decrease in the filtration rate of the clam *Corbicula fluminea* after
326 exposure to CBZ by comparison with non-contaminated clams. Also Almeida et al.
327 (2015) observed lower BCF values at the highest CBZ exposure concentration in *R.*
328 *philippinarum*.

329

330 3.2 Metabolic capacity and energy reserves

331 Except for the lowest tested concentration (2.5 µg/L), mussels exposed to Nd
332 significantly increased their metabolic capacity compared to non-contaminated

333 organisms under control condition (Figure 1A). However, except for the lowest Nd
334 concentration, higher electron transport system (ETS) values compared to control
335 organisms did not vary among tested treatments, which may explain similar BCF values
336 in concentrations higher than 2.5 $\mu\text{g/L}$. As mentioned above, as a result of similar
337 metabolic capacity among different exposure concentrations, mussels may have
338 presented similar filtration rates which led to a similar accumulation rate among
339 different tested conditions. It was already demonstrated that ETS activity may be used
340 as an indication of metabolic activity in marine macrofauna (Cammen et al., 1990), and
341 an increase in bivalves ETS activity was already identified as a protective behavior,
342 associated with the activation of defense mechanisms under pollution exposure,
343 including the increase of antioxidant and biotransformation enzymes. In particular,
344 different authors already demonstrated that marine bivalves (*Ruditapes philippinarum*
345 and *M. galloprovincialis*) increased their metabolic capacity, measured by ETS activity,
346 when in the presence of nanoparticles (multi-walled carbon nanotubes) and drugs
347 (salicylic acid) (De Marchi et al., 2018; Freitas et al., 2019). Nevertheless, several other
348 studies addressing the metabolic capacity of bivalves under pollution stress evidenced a
349 decrease on ETS activity, which was associated to a decrease in the filtration rate to
350 prevent pollutants accumulation (among others, Almeida et al., 2015; Oliveira et al.,
351 2017). Also, a previous study with other REE (Gadolinium, Gd) but testing a similar
352 concentration range (between 15 and 60 $\mu\text{g/L}$) and the same exposure period (twenty-
353 eight days), showed that the ETS activity in *M. galloprovincialis* decreased significantly
354 after the experimental period (Henriques et al., 2019). Such findings may indicate
355 higher toxicity of Nd in comparison to other pollutants, as mussels under higher stress
356 conditions may increase their metabolic capacity to fight against the stressful condition.
357 Therefore, considering the results obtained and previous studies with bivalves exposed

358 to pollutants we can hypothesize that up to certain stress levels bivalves are able to
359 decrease their metabolism for short periods of time, to avoid accumulation of pollutants
360 and reduce their toxic impacts. This strategy can seriously affect bivalves physiological
361 and biochemical performance, being tolerable for a limited period of time. Therefore, it
362 seems that at higher stressful conditions this strategy is no longer valid and organisms
363 increase their ETS activity to activate defense mechanisms, which results into higher
364 production of reactive oxygen species (ROS) by mitochondrial electron transport
365 system, with negative impacts on organism's cellular oxidative status.

366

367 As a consequence of higher metabolic activity, the results obtained showed that
368 associated with higher ETS activity contaminated organisms presented significantly
369 lower glycogen (GLY) content in comparison to control mussels (Figure 1B). These
370 findings evidence the need of mussels to use their energy reserves to fuel up defense
371 mechanisms. Previous studies conducted by Lagadic et al. (1994) already suggested that
372 energy reserves could be considered as biomarkers reflecting sublethal changes from a
373 stressful xenobiotic exposure. Also Pellerin-Massicotte et al. (1994) highlighted that
374 GLY reserves may be depleted in the presence of contaminants. Other authors identified
375 the possible use of GLY for the synthesis of lipids and/or proteins for gametogenesis
376 (Bayne et al., 1975; Parra et al., 2005). In the present study we may hypothesize that
377 GLY reserves were used in the activation of defense mechanisms. A similar response
378 was also observed by other authors, assessing the effects carbon nanotubes in the clam
379 *R. philippinarum* (De Marchi et al., 2018).

380

381 The results obtained further demonstrated that although the GLY content
382 decreased in contaminated mussels compared to control ones, the protein (PROT)

383 content was maintained regardless the Nd concentration of exposure, with no significant
384 differences among tested conditions (Figure 1C). Such results demonstrated that
385 mussels were neither using PROT as energy reserves to fuel up defense mechanisms nor
386 increasing the production of enzymes to fight against the stress caused by Nd.
387 Nevertheless, previous studies with REE showed that *M. galloprovincialis* were able to
388 increase the PROT content in the presence of an increasing concentration gradient of Gd
389 and Lanthanum (La) (Henriques et al., 2019; Pinto et al., 2019), which could be related
390 to the capacity of mussels to increase production of enzymes to fight against the stress
391 induced, indicating also that in this case mussels were experiencing a mild stress
392 condition with no need to use PROT as energy source while being able to enhance the
393 production of enzymes.

394

395 Overall, the results obtained evidenced that under Nd exposure mussels increased
396 their metabolic capacity, probably to fuel up defense mechanisms (namely antioxidant
397 enzymes activity), which was accompanied by expenditure of GLY reserves but not a
398 decrease in PROT content. It was already described that up to certain stress levels stored
399 GLY is the first source of energy used, while energy stored in lipid and PROT being
400 used at higher stress levels (Sonawane and Sonawane, 2018).

401

402 3.3 Antioxidant defenses

403 In terms of superoxide dismutase (SOD) activity, the results obtained showed no
404 significant differences among conditions except for the lowest Nd concentration where
405 the activity of this enzyme was significantly higher compared to the remaining
406 conditions (Figure 2A). In the case of glutathione peroxidase (GPx), at the highest
407 exposure concentration no significant differences were observed to control organisms,

408 while at the remaining exposure concentrations significantly lower activity was
409 recorded compared to control and the highest concentration (Figure 2B). On the
410 contrary, mussels exposed to Nd tended to increase their catalase (CAT) activity, which
411 was significantly higher at concentrations 5.0, 10 and 40 $\mu\text{g/L}$ compared to the
412 remaining conditions (Figure 2C). It is well known that when in the presence of a
413 stressful condition, including the presence of pollutants, organisms may increase the
414 production of ROS. To avoid damages caused by ROS (including lipid peroxidation,
415 protein carbonilation and DNA damage), organisms may increase the activity of
416 antioxidant enzymes. Among these enzymes are SOD, GPx and CAT that have the
417 capacity to eliminate ROS (namely, superoxide anion, hydroxyl radical, and hydrogen
418 peroxide), preventing organisms from cellular damages. Nevertheless, this response
419 normally occurs when oxidative stress is not very high or very long-during. On the other
420 hand, when exposed to extremely high stressful conditions or if the stress is persisting,
421 the proteins damage became profound and a decrease of these enzymes activity may
422 occur (either via direct oxidative damage of the enzymes molecules, or via oxidative
423 stress-altered enzymes gene expression, or both). Among others, Manduzio et al. (2004)
424 hypothesized that the over production of ROS inhibited the SOD activity in *Mytilus*
425 *edulis* collected from a polluted area. Studies conducted by Matozzo et al. (2001)
426 highlighted that the significant inhibition of SOD activity in Cu-exposed *R.*
427 *philippinarum* clams observed might be due to the oxidation of the enzyme SH groups
428 mediated by ROS, which production is increased by Cu (Halliwell and Gutteridge,
429 1984). In accordance to this, the present study may indicate that mussels were exposed
430 to high toxic conditions since the results obtained showed that *M. galloprovincialis*
431 exposed to Nd were not able to significantly increase their antioxidant defenses, namely
432 in terms of SOD and GPx activity, which was especially noticed at higher exposure

433 concentrations. This behaviour may have limited mussel's capacity to eliminate the
434 excess of ROS generated by the presence of this element. Nevertheless, one of the three
435 antioxidant enzymes analyzed seemed to be sensitive to some of the treatments, which
436 can suggest not only a complex mode of action of this element but also that not all the
437 mechanisms involved in the onset of oxidative stress due to Nd have been investigated.
438 Furthermore, the results obtained may also evidence that increased metabolic capacity
439 was not enough to significantly activate antioxidant enzymes and increased ETS activity
440 also contributed to the generation of higher ROS amount. Previous studies also showed
441 that in the presence of Gd *M. galloprovincialis* presented limited capacity to activate
442 their antioxidant enzymes, but in this case only at higher exposure concentrations (60
443 and 120 $\mu\text{g/L}$) mussels were not able to continue to increase antioxidant enzymes
444 activity (Henriques et al., 2019). Such results can, once again, corroborate the
445 hypothesis that Nd may be more toxic than Gd towards *M. galloprovincialis*, exposed to
446 a similar concentration range.

447

448 3.4 Biotransformation defenses

449 Concerning biotransformation capacity, mussels exposed to lower concentrations
450 (2.5, 5.0 and 10 $\mu\text{g/L}$) tended to increase glutathione S-transferases (GSTs) activity,
451 with significantly higher values at 5.0 and 10 $\mu\text{g/L}$ compared to the remaining
452 conditions (Figure 3). As for the antioxidant enzymes, mussels were not able to increase
453 the activity of biotransformation enzymes along the increasing exposure concentration,
454 showing limited capacity to increase GSTs activity at higher Nd concentrations (20 and
455 40 $\mu\text{g/L}$). GSTs are a superfamily of Phase II detoxification enzymes involved in the
456 detoxification of ROS and toxic xenobiotics. These enzymes are able to catalyze the
457 conjugation of the reduced form of glutathione (GSH) to xenobiotic substrates for the

458 purpose of detoxification and, therefore, in the presence of pollutants, GSTs activity is
459 induced to achieve efficient cell protection. Thus, from the results obtained, it is
460 possible to identify mussel's efforts to detoxify cells from Nd under intermediate
461 concentrations, while exposure to the highest concentrations (20 and 40 $\mu\text{g/L}$) mussels
462 were no longer able to continue to activate this defense mechanism. The decrease of
463 GSTs activity at higher concentrations may be related to diminished levels of GSH
464 susceptible of being conjugated. A similar response was observed when *M.*
465 *galloprovincialis* mussels were exposed to La and Gd, with higher activity at lower
466 concentrations and lower activity at higher concentrations (Henriques et al., 2019; Pinto
467 et al., 2019). Therefore, the results obtained in the present study as well as in previous
468 studies showed the capacity of GSTs to detoxify REE, with greater detoxification
469 capacity at lower exposure concentrations.

470

471 Overall, in what regards to defense mechanisms, the results obtained indicate high
472 toxicity of Nd, especially at higher concentrations, which may have resulted into i)
473 increase of SOD activity only at the lowest tested concentration, with inactivation of
474 this enzyme at higher concentrations; ii) inhibition of GPx in contaminated mussels; iii)
475 decrease of GSTs at higher exposure concentrations; iv) only the activation of CAT in
476 contaminated organisms.

477

478 3.4 Cellular damage

479 Levels of lipid peroxidation (LPO) significantly increased in organisms exposed
480 to Nd in comparison to control organisms, with no significant differences among 5.0,
481 10, 20 and 40 $\mu\text{g/L}$ exposure concentrations (Figure 4A). These results may result from
482 the fact that mussels were not able to efficiently activate antioxidant enzymes, resulting

483 into cellular damages in the presence of Nd. Previous studies already demonstrated that
484 the presence of pollutants results into an overproduction of ROS that, if not eliminated
485 by antioxidant enzymes, can react with lipids of the cellular membrane, causing lipid
486 peroxidation that corresponds the oxidative degradation of lipids (see for example,
487 Regoli and Giuliani, 2014). Although a high diversity of studies already showed that the
488 presence of metals, nanoparticles and drugs may originate increased LPO levels in
489 bivalves even if antioxidant enzymes were activated (see for example, Freitas et al.,
490 2019b, 2019a; McCarthy et al., 2013; Monteiro et al., 2019; Vlahogianni and
491 Valavanidis, 2007), less studies demonstrated the occurrence of LPO when bivalves are
492 exposed REE. In particular, Henriques et al. (2019) demonstrated that when *M.*
493 *galloprovincialis* were exposed to Gd LPO significantly increased in comparison to
494 control values, although antioxidant enzymes were increased, especially at intermediate
495 concentrations (30 and 60 µg/L). Hanana et al. (2017) revealed a significant increase of
496 LPO in the freshwater mussel *Dreissena polymorpha* after 28 days of exposure but only
497 when exposed to the highest concentration of La (1250 µg/L), with no significant
498 differences evidenced among the other tested concentrations (10, 50, 250 µg/L). With
499 other aquatic invertebrates, it was also demonstrated the capacity of REE to enhance the
500 production of ROS, such was in the freshwater crustacean *Daphnia magna* exposed to
501 Cerium and Erbium. LPO levels also increased in the sea urchin *Paracentrotus lividus*
502 larvae exposed to Dysprosium (Oral et al., 2017). Also Wang et al. (2011) showed that
503 the significant increasing activities of antioxidant enzyme observed in the freshwater
504 cyanobacteria *Microcystis aeruginosa* may result from overproduction of ROS due to
505 the exposure to Nd concentrations. Altogether, these findings clearly demonstrate the
506 capacity of REE to induce cellular damages.

507

508 3.5 Redox balance

509 The ratio between reduced (GSH) and oxidized (GSSG) glutathione was
510 significantly lower in mussels exposed to Nd in comparison to non-exposed ones, with
511 no significant differences among contaminated mussels (Figure 4B). Such results
512 clearly reveal that GSH content decreased while GSSG increased in contaminated
513 mussels, indicating loss of redox homeostasis in organisms exposed to Nd and the high
514 demand for GSTs. To eliminate the excess of ROS generated by a stress condition,
515 besides antioxidant enzymatic defenses, organisms further present low molecular
516 scavengers that are also to neutralize ROS by direct reaction with them, being GSH the
517 most abundant. In its reduced form, GSH, glutathione is capable of scavenging reactive
518 oxygen and nitrogen species, thereby contributing to the control of redox homeostasis.
519 Therefore, the glutathione system acts as the major redox buffer in the majority of cells
520 (Couto et al., 2016). In the presence of ROS, GSH can be oxidized into GSSG and, thus,
521 under stressful conditions the ratio GSH/GSSG tends to decrease as a result of GSSG
522 increase. Organisms which use glutathione for redox homeostasis are able to
523 synthesize reduced glutathione, but they are also characterized by their ability to recycle
524 glutathione. Glutathione reductase (GRed) is an essential enzyme that recycles oxidised
525 glutathione back to the reduced form (Couto et al., 2016). Thus, the increased content of
526 GSSG observed by lower GSH/GSSG levels in contaminated mussels indicates that
527 GRed failed to oxidize glutathione into its reduced form. The ratio GSH/GSSG is often
528 used to assess the oxidative stress of organisms exposed to pollutants (e.g. Peña-Llopis
529 et al., 2002; Almeida et al., 2015; Sellami et al., 2015; Freitas et al., 2018; 2019).
530 Similarly, to the present results, recent studies also demonstrated that in the presence of
531 REEs mussels significantly decreased in mussels exposed to La and Gd (Henriques et
532 al., 2019; Pinto et al., 2019).

533

534

535 **4. CONCLUSIONS**

536 The present findings revealed high toxicity of Nd towards *M. galloprovincialis*,
537 which showed low capacity to prevent injuries caused by this REE. After exposure,
538 mussels accumulated Nd with higher concentrations at higher exposure levels.
539 Accumulation of Nd revealed to be costly to mussels, which revealed higher metabolic
540 activity and increased expenditure of GLY content when in the presence of this element.
541 Also, after exposure to Nd, mussels showed inefficient antioxidant and
542 biotransformation strategies, leading to cellular damage and loss of redox balance
543 provoked by the excess of ROS, namely as a result of higher electron transport system
544 activity. Considering that tested concentrations resemble low to highly polluted areas,
545 the results here presented highlight the hazardous capacity of Nd towards *M.*
546 *galloprovincialis*. Toxic effects observed at individual level may result into negative
547 impacts to mussel's population as changes observed at cellular level may result into
548 impairments on organism's survival, growth, abundance and reproduction capacity.

549

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562

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FIGURE CAPTIONS

Figure 1. A: Electron transport system activity (ETS), B: Glycogen content (GLY); and C: Protein content (PROT), in *Mytilus galloprovincialis* exposed to different Neodymium concentrations (CTL-0, 2.5, 5.0, 10, 20 and 40 $\mu\text{g/L}$ of Nd). Values are mean + standard deviation. Significant differences among concentrations are represented with different letters.

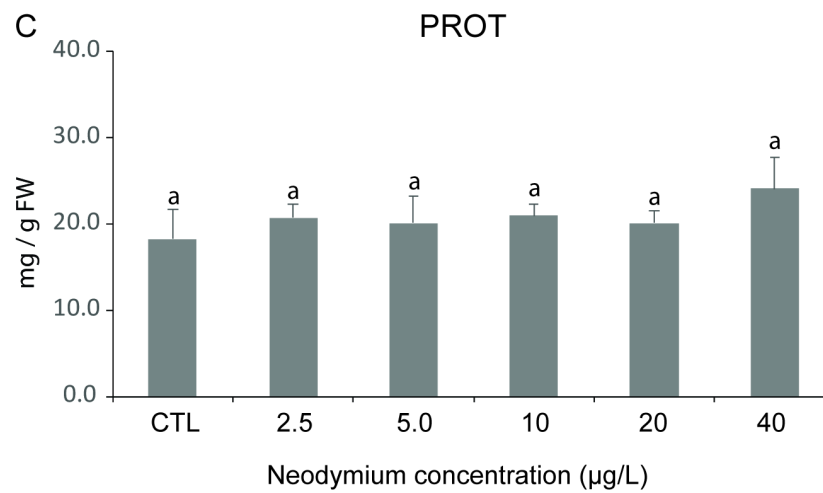
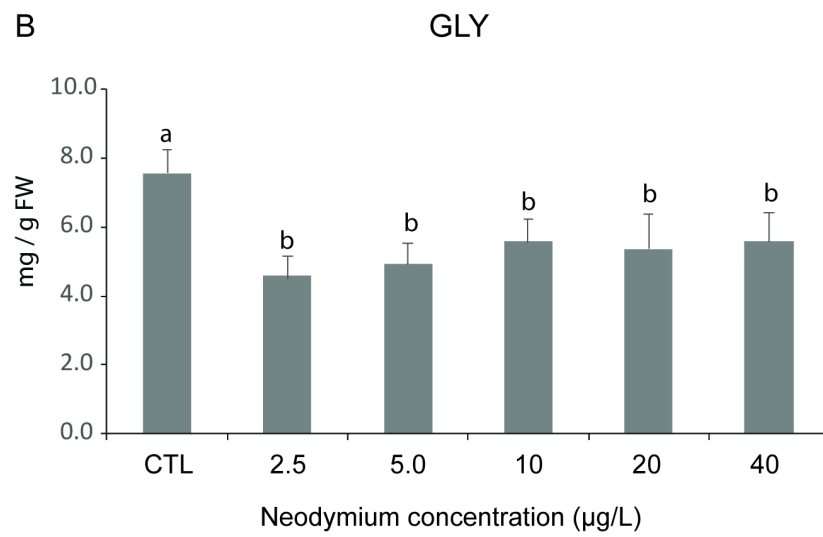
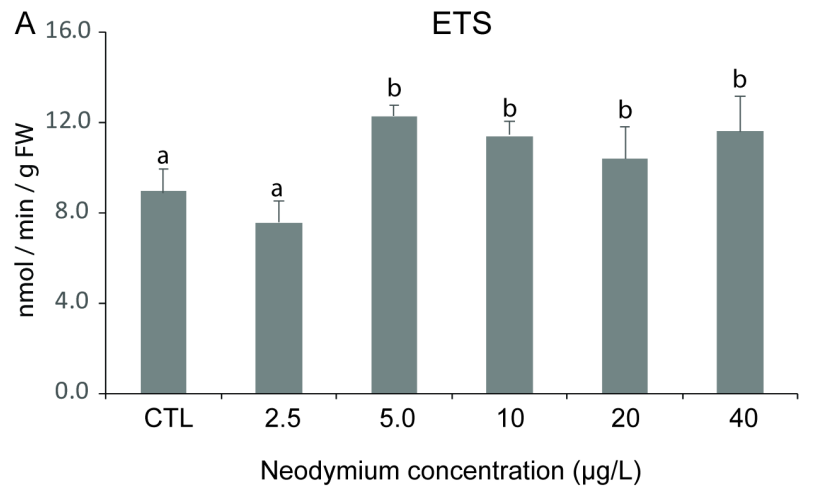
Figure 2. A: Superoxide dismutase activity (SOD); B: Glutathione peroxidase activity (GPx); and C: Catalase activity (CAT), in *Mytilus galloprovincialis* exposed to different Neodymium concentrations (CTL-0, 2.5, 5.0, 10, 20 and 40 $\mu\text{g/L}$ of Nd). Values are mean + standard deviation. Significant differences among concentrations are represented with different letters.

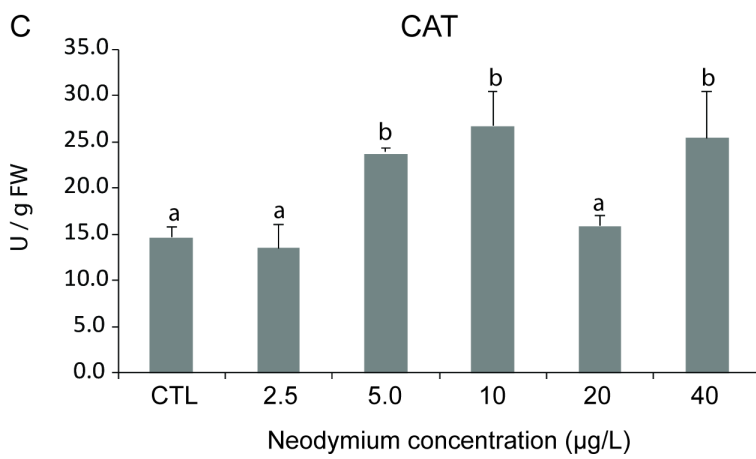
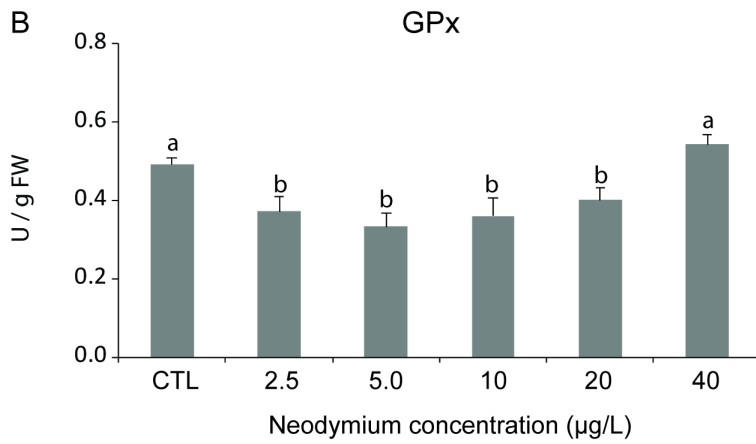
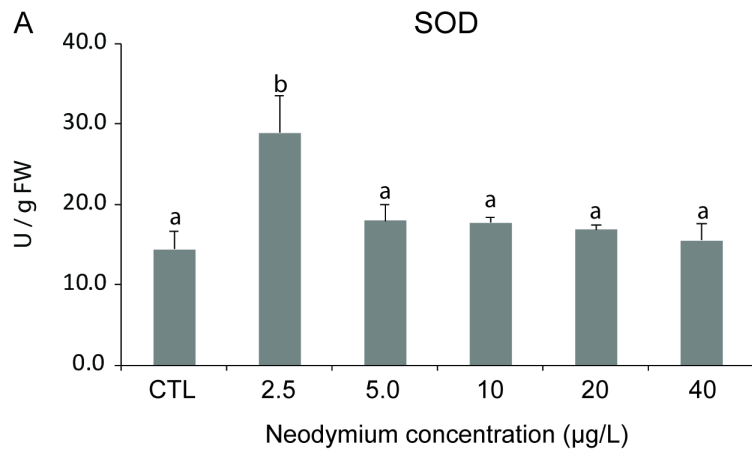
Figure 3. Glutathione S-transferases activity (GSTs), in *Mytilus galloprovincialis* exposed to different Neodymium concentrations (CTL-0, 2.5, 5.0, 10, 20 and 40 $\mu\text{g/L}$ of Nd). Values are mean + standard deviation. Significant differences among concentrations are represented with different letters.

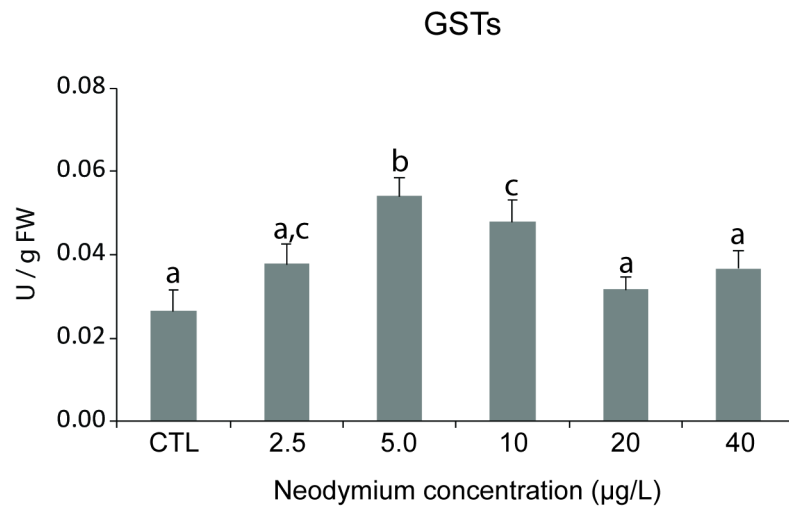
Figure 4. Lipid peroxidation levels (LPO); and B: Ratio between reduced (GSH) and oxidized (GSSG) glutathione (GSH/GSSG), in *Mytilus galloprovincialis* exposed to different Neodymium concentrations (CTL-0, 2.5, 5.0, 10, 20 and 40 $\mu\text{g/L}$ of Nd). Values are mean + standard deviation. Significant differences among concentrations are represented with different letters.

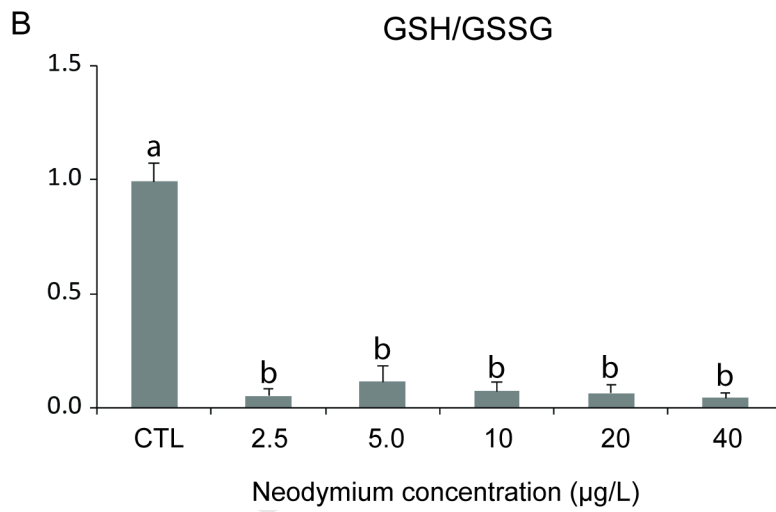
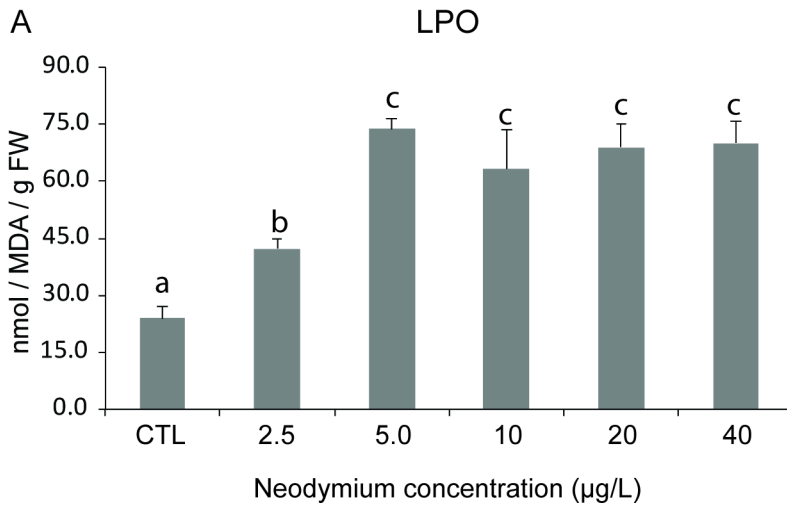
Table 1- Neodymium (Nd) mean concentrations, in water ($\mu\text{g/L}$), collected immediately after spiking at the 1st, 2nd, 3rd and 4th weeks of exposure, and in mussels tissues ($\mu\text{g/g}$ dry weight) at the end of the experimental period (28 days) from each condition (0-control, 2.5, 5.0, 10, 20, 40 $\mu\text{g/L}$ of Nd). Different letters among exposure concentrations denote statistical significance. LOQ for water samples 10 ng/L; LOQ for tissue samples 0.0025 $\mu\text{g/g}$. Bioconcentration factor (BCF) corresponds to the concentration of Nd in mussel's tissues divided by the mean values for the real exposure concentration during the four weeks of exposure.

Nd concentrations ($\mu\text{g/L}$)	Water	Mussels tissues ($\mu\text{g/g}$)	BCF
	During the four weeks	In the 4 th week	
CTL	<LOQ	0.095 \pm 0.006	---
2.5	2.6 \pm 0.32	0.136 \pm 0.016 ^a	0.05
5.0	5.3 \pm 0.30	0.26 \pm 0.013 ^b	0.05
10	10 \pm 0.3	0.435 \pm 0.003 ^c	0.04
20	22 \pm 0.8	0.982 \pm 0.008 ^d	0.05
40	43 \pm 3.0	1.72 \pm 0.03 ^e	0.04









- *Mytilus galloprovincialis* bioaccumulated Neodymium (Nd)
- Mussels exposed to Nd increased their metabolic capacity, with glycogen expenditure
- Limited antioxidant and biotransformation capacity in contaminated mussels
- Lipid peroxidation occurred in Nd contaminated mussels
- Loss of redox balance in mussels exposed to Nd

Journal Pre-proof

Rosa Freitas and Eduarda Pereira are supervisors of the students that co-authored this manuscript (Silvana Costa, Celso Cardoso, Tiago Morais, Pedro Moleiro, Ana C. Matias, Ana F. Pereira, Joana Machado, Beatriz Correia, Diana Pinheiro, Adriana Rodrigues, João Colónia). Students did the exposure assay (for 28 days under controlled conditions), performed all methods and analyses for Nd quantification and biomarkers determination.

Rosa Freitas and Eduarda Pereira gave the idea of this study to the students that accepted this challenge and performed all the analyses during their last year of their bachelor degree. Eduarda Pereira is the responsible for the laboratory where Nd quantification was done. Rosa Freitas and Amadeu Soares are the responsible persons for the labs where biomarkers were determined. Eduarda Pereira, Rosa Freitas and Amadeu Soares funded this study.

Conflict of Interest

The Authors whose names are listed immediately below certify that they have NO affiliations with or involvement in any organization or entity with any financial interest (such as honoraria; educational grants; participation in speakers' bureaus; membership, employment, consultancies, stock ownership, or other equity interest; and expert testimony or patent-licensing arrangements), or non-financial interest (such as personal or professional relationships, affiliations, knowledge or beliefs) in the subject matter or materials discussed in this manuscript.

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