

1 Enriching *Artemia* nauplii with selenium from different sources and interactions with essential
2 fatty acid incorporation

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20 **Abstract**

21 The production of high-quality marine fish fry is limited by the low survival observed during
22 the larval phase, which is often attributed to dietary deficiencies of the diets at first feeding.
23 Despite progress made with live feed (i.e. rotifers, *Artemia*), enrichments in essential fatty acids
24 for marine fish larvae, little is known on the micronutrient requirements such as selenium (Se).
25 Se is a critical component of several enzymes maintaining important biological functions such
26 as cellular oxidation, and therefore plays a key role in oxidative and stress status of marine
27 larvae. The levels of Se found in the larvae's natural diet (i.e. copepods) is generally higher
28 than those of the enriched live preys used in hatcheries. This study aimed at establishing a
29 protocol to enrich *Artemia* nauplii with Se using different inorganic (sodium selenite) and
30 organic (selenoyeast). Results indicated that the use of dissolved sodium selenite, an alternative
31 inorganic and cheaper form of Se, did not increase the levels of Se in the nauplii. However, the
32 use of selenoyeast (Sel-Plex) confirmed that it is possible to enrich the nauplii with targeted
33 levels of Se, since this process followed a dose-response pattern with Se enrichment ranging
34 from 1.7 to 12.4 mg kg⁻¹. In addition, the supplementation of Sel-Plex to the regular enrichment
35 product did not impact on lipids and fatty acids enrichment irrespective of the dose dispensed.
36 Overall, this study contributes to the refinement of the live prey enrichment protocols that are
37 critical to the success of marine finfish larviculture protocols.

38 **Keywords:** *Artemia*; Essential fatty acids; Selenium; Sel-Plex; Sodium selenite.

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43 **1. Introduction**

44 Marine finfish larviculture has remarkably improved during the last three decades but it still
45 remains one of the most challenging research areas and commercial bottleneck in marine fish
46 aquaculture. Despite significant advances in the hatcheries' operational procedures, survival
47 rates during the larval phases usually remains low with at best 30 % in well-established species
48 such as gilthead seabream (*Sparus aurata*) (Atalah et al., 2011) and European seabass
49 (*Dicentrarchus labrax*) (Villamizar et al., 2009), and <10 % in most other species such as
50 California yellowtail (*Seriola lalandi*) (Hawkyard et al., 2016), Atlantic halibut (*Hippoglossus*
51 *hippoglossus*) (Bjornsdottir et al., 2009) and ballan wrasse (*Labrus bergylta*) (Øie et al., 2015).
52 Causes accounting for the low survival in marine larviculture are multifactorial but nutrition
53 and feeding have been often regarded as major causal factors (Hamre et al., 2013). While it is
54 fairly well-understood which nutrients are essential in marine fish larvae diets (NRC, 2011),
55 quantitative requirements are largely unknown for most fish species and nutrients. This is
56 mainly due, among other reasons, to the inherent difficulty to manipulate and reliably control
57 dietary content of individual nutrient within live preys (i.e. rotifers, *Artemia*) to match marine
58 finfish larvae requirements (Støttrup, 2000; Conceição et al., 2010; Dhont et al., 2013).
59 Nutrients can be incorporated into live preys through the so-called "bioencapsulation" or
60 "enrichment" protocols (Sorgeloos et al., 2001), however, targeting a recommended level is
61 challenging due primarily to the metabolic activity of live preys towards the enrichment
62 products (Navarro et al., 1999; Reis et al., 2017) and the inherent variability associated with
63 the live prey enrichment process (Navarro et al., 1999; Hamre, 2016). Furthermore, the
64 nauplii's gut can only contain a defined amount of nutrient thus increasing the level of one
65 nutrient (i.e. lipid) can result in the decrease of another nutrient's level (e.g. micromineral).

66 The brine shrimp *Artemia* sp. is the most commonly used live feed in marine finfish
67 hatcheries (Conceição et al., 2010; Dhont et al., 2013). Unlike copepods, the natural preys of

68 marine fish larvae (Hunter, 1980), *Artemia* have a suboptimal nutritional profile which does
69 not satisfy the requirements of marine fish larvae. It is well-established that *Artemia* is deficient
70 in certain essential lipids for larval stages of fish, particularly phospholipids and long-chain
71 polyunsaturated fatty acids (LC-PUFA) (i.e. arachidonic acid, ARA; eicosapentaenoic acid,
72 EPA; docosahexaenoic acid, DHA) (Navarro et al., 1999; Monroig et al., 2003; Dhont et al.,
73 2013). Along with essential lipids, successful *Artemia* enrichment strategies have also been
74 tested and established for other essential nutrients such as methionine (Tonheim et al., 2000),
75 vitamins A, C and E (Monroig et al., 2007; Adloo, 2012), minerals including cobalt (Fehér et
76 al., 2013), iodine (Moren et al., 2006), manganese (Nguyen et al., 2008) and selenium (Se)
77 (Hamre et al., 2008a,b; Penglase et al., 2011). Importantly, Se is regarded as an essential trace
78 element for virtually all animal species (Hefnawy and Tórtora-Pérez, 2010) and it is a major
79 component of the glutathione peroxidase (GPx), which is involved in the regulation of the
80 antioxidant status in finfish by reducing hydrogen peroxide and hydroperoxides to their base
81 constituents (Lall, 2003; Pacitti et al., 2015). Moreover, Se has also been shown to play a
82 protective role by reducing oxidative stress caused by heavy metals such as copper (Cu),
83 resulting in enhanced immune response in fish (Lin and Shiau, 2007). Studies have shown that
84 dietary deficiencies of Se can result in reduced growth in channel catfish *Ictalurus punctatus*
85 (Wang and Lovell, 1997) and increased mortality in rainbow trout *Oncorhynchus mykiss*
86 (Hilton et al., 1980). The NRC (2011) recommendations for Se dietary requirements in fish
87 vary across species but in average, Se dietary requirements for juvenile and adult stages are
88 $\sim 0.35 \text{ mg kg}^{-1}$ (NRC, 2011). Importantly, Se requirements in marine fish larvae are largely
89 unknown despite playing a crucial role as an antioxidant and subsequently reducing stress
90 levels in fragile marine larvae (Saleh et al., 2014). Se levels in enriched *Artemia* nauplii are
91 usually around 2 mg kg^{-1} (Ribeiro et al., 2012a,b), which is at the lower end of wild copepods
92 Se content (i.e. between 2 to 5 mg kg^{-1}) (Hamre et al., 2008b; Mæhre et al., 2013). This may

93 therefore not satisfy the dietary requirements of marine fish larvae (Solbakken et al., 2002;
94 NRC, 2011; Hamre et al., 2013). A study performed in Atlantic cod (*Gadus morhua*) showed
95 that larvae fed rotifers enriched with Se exhibited a higher survival compared to larvae fed the
96 control rotifers (Penglase et al., 2010). In addition, the inclusion of Se in marine fish larval
97 diets has been suggested to potentially offset the high oxidation risk from diets boosted in LC-
98 PUFA (Saleh et al., 2014).

99 Bioavailability of Se, as per other minerals (NRC, 2011), is greatly dependent upon its
100 form when accumulated in the diet (Pacitti et al., 2015). One of the most common forms of
101 dietary Se is sodium selenite (Na_2SeO_2 , hereafter referred to as Na-Se), a highly water-soluble
102 inorganic compound. An alternative Se source is selenoyeast (Se-yeast), an organic source of
103 Se produced by exposing yeast (*Saccharomyces cerevisiae*) to Na-Se (Suhajda et al., 2000),
104 which results in an accumulation of selenomethionine (Se-Met). The latter (organic) form of
105 Se is regarded to be more bioavailable to organisms, thus explaining why it is broadly used as
106 a livestock feed additive (Wang and Lovell, 1997; Rayman, 2004; Thiry et al., 2012).
107 Nevertheless, inappropriate dietary Se levels can induce toxicity effects. Se dietary toxicity
108 levels has been reported in rainbow trout (*Onchorynchus mykiss*) at 13 mg Se kg^{-1} when
109 presented as Na-Se (Hilton et al., 1980) and at levels of 20 mg Se kg^{-1} when presented as Se-
110 Met in hybrid striped bass (*Morone chrysops* \times *M. saxatilis*) (Jaramillo et al., 2009). In both
111 studies, Se toxicities resulted in growth impairment and elevated mortality.

112 Importantly, with a few exceptions (e.g. Penglase et al., 2010), most studies mentioned
113 above were conducted in fish juveniles and information on Se requirements and toxicity levels
114 in fish larvae remain scarce. The present study aimed to compare the effectiveness of different
115 Se enrichment protocols for *Artemia* nauplii using different Se sources and determine how
116 these impact on essential fatty acids enrichment using a commercially available enrichment

117 product. The effects of varying doses of Se-yeast were first tested then the efficiency of
118 different Se sources were studied on *Artemia* nauplii content in Se and essential fatty acids.

119 **2. Materials and Methods**

120 *2.1.Artemia hatching and culture*

121 *Artemia* cysts GSL (EG, Inve, Belgium) were decapsulated and hatched according to Sorgeloos
122 et al. (2001). Newly hatched *Artemia* nauplii were subsequently used for the enrichment
123 experiments. Enrichments were carried in 1-litre Imhoff cones using glass pipes to strongly
124 aerate the cones from the bottom. The cones were subsequently placed in a 28 °C water bath.
125 After the 24 h hatching process, enriched nauplii were collected on a 100 µm sieve, rinsed with
126 freshwater, and distributed in each cone at 300 nauplii ml⁻¹ for further enrichment. Artificial
127 seawater (32 ppt) (Instant Ocean, Virginia, USA), disinfected with Pyceze® (0.05 ml l⁻¹), was
128 dispensed in each 1-litre Imhoff cone to provide a final volume of 800 ml after the addition of
129 the enrichment products (see below). The enrichment experiments were run under constant
130 illumination of about 47 klx at the surface of the water.

131 *2.2.Experiment 1: Effects of varying doses of Se-yeast on Se content of Artemia nauplii.*

132 Newly hatched *Artemia* were first enriched with Sel-Plex (SP) (Alltech, Kentucky, USA),
133 during 4 h at different concentrations: 0 mg l⁻¹ (Treatment SP0), 12 mg l⁻¹ (Treatment SP12),
134 24 mg l⁻¹ (Treatment SP24) and 36 mg l⁻¹ (Treatment SP36). Each SP dose was tested in
135 triplicated 1-litre Imhoff cones (i.e. 4 treatments x 3 replicates). After 4 h, Larviva Multigain
136 (MG) (BioMar, Denmark) was added to each of the 12 cones at 0.6 g l⁻¹ for a further 24 h. At
137 the end of the MG enrichment period, nauplii were collected on a 100 µm sieve, thoroughly
138 rinsed with freshwater to remove the excess of enrichment product, and gently dried on
139 absorbent paper before transferring them into universal sample tubes. The samples were frozen
140 at -20 °C before being freeze-dried and stored at -20 °C for further analysis.

141 2.3. *Experiment 2: Effects of different sources of Se on Artemia nauplii enrichment*
142 *efficiency.*

143 Experiment 2 investigated the efficiency of two sources of Se to increase the content of Se in
144 *Artemia nauplii*. Four enrichment diets were tested in triplicate: no addition of Se sources
145 (Treatment SP0); SP at 12 mg l⁻¹ for 4 h (Treatment SP12); sodium selenite Na-Se (Sigma
146 Aldrich, UK) at the Se equivalent dose (i.e. 24 µg l⁻¹) of Treatment SP12 for 4 h (Treatment
147 NS); soya lecithin emulsion (0.6 g l⁻¹) (Optima Health, UK) containing Na-Se (24 µg l⁻¹) for 4
148 h (Treatment SL+NS). Treatments SP12, NS and SL+NS contained the same effective dose of
149 Se (24 µg l⁻¹) irrespectively of the form under which Se was presented. For treatment SL+NS,
150 we used a soya lecithin emulsion, potentially creating lipid vesicles (liposomes) that can
151 encapsulate dissolved Se, as this has been proven to be a good strategy to deliver water-soluble
152 nutrients in *Artemia nauplii* (Monroig et al., 2007). The hereby tested sodium selenite was an
153 inorganic, water soluble compound with ≥ 95.0 % purity. During the first 4 h, nothing was
154 added to enrichment medium of Treatment SP0, whereas the other treatments were enriched
155 with Se as indicated above. After the 4 h, Larviva Multigain (0.6 g l⁻¹) was added to all
156 treatments for a further 24 h. Sample collection and storage was done as described in
157 Experiment 1.

158 2.4. *Nutritional analysis*

159 Total lipids (TL) from the enrichment products MG, SL and SP (Table 1), as well as freeze-
160 dried *Artemia* samples collected from Experiments 1 and 2, were extracted according to Folch
161 et al. (1957), with modifications as described by Monroig et al. (2006). Fatty acid methyl esters
162 (FAME) from TL were prepared, extracted and purified according to Christie (2003).
163 Identification and quantification of FAME were carried out using a gas chromatograph coupled
164 with flame ionisation detection as previously described (Houston et al. 2017).

165 Se concentration was determined after digestion of Sel-Plex, enrichment products and
166 freeze-dried *Artemia* samples in AristAR nitric acid (VWR International, Pennsylvania, USA)
167 in a microwave MARSXpress (CEM, North Carolina, USA) for 40 min (20 min ramping to
168 120 °C and 20 min holding that temperature). Digests were transferred into a volumetric flask
169 and made up into x 25 dilutions with distilled water. Samples were analysed by Inductively
170 Coupled Plasma Mass Spectrometry (ICP-MS, Thermo Scientific Model X Series 2,
171 Massachusetts, USA) (Smedley et al., 2016).

172 2.5. Statistical analysis

173 All enrichment treatments in both experiments were carried out in triplicate cones ($n = 3$).
174 Biological and analytical data are expressed as means \pm standard deviation (SD). Percentage
175 data were transformed using the arcsine square root function prior to statistical analysis.
176 Difference among treatments for total lipids and fatty acids were analysed by one-way ANOVA
177 followed by a Tukey post-hoc multiple comparison test at a significance level of $P \leq 0.05$ (IBM
178 SPSS Statistics 23, New-York (state), USA).

179

180 3. Results

181 3.1. Experiment 1: Effects of varying doses of Se-yeast on Se content of *Artemia* nauplii.

182 The analysis of the unenriched *Artemia* nauplii revealed the presence of Se ($1.83 \pm 0.18 \text{ mg kg}^{-1}$
183 DW). In the experimental groups, Se concentration in *Artemia* nauplii increased linearly with
184 increasing levels of selenoyeast Sel-Plex (Fig. 1). The equation of the linear regression was:

$$185 \quad [\text{Se}]_{\text{Artemia}} = 3.79 \times [\text{SelPlex}]_{\text{enrichment}} - 2.42 \quad (r^2 = 0.97)$$

186 where $[\text{Se}]_{\text{Artemia}}$ is the Se content in the enriched *Artemia* (mg kg^{-1} DW) and $[\text{SelPlex}]_{\text{enrichment}}$
187 is the dose of Sel-Plex used to prepare enrichment (mg l^{-1}). The Se content in *Artemia* nauplii

188 varied from $1.7 \pm 0.1 \text{ mg kg}^{-1}$ (Treatment SP0), defined as the basal level of Se found in the
189 non-enriched nauplii, to $12.4 \pm 1.0 \text{ mg kg}^{-1}$ (Treatment SP36), with significant differences
190 between treatments ($P < 0.05$) (Fig. 1). Sel-Plex is a yeast-based product that contains lipids
191 (Table 1). However, the dose of Sel-Plex used in Experiment 1 (from 0 to 36 mg l^{-1}) did not
192 affect the TL contents of enriched *Artemia* nauplii ($P > 0.05$) (Table 2), which ranged between
193 $209.5 \pm 9.0 \text{ mg g}^{-1}$ (Treatment SP0) and $220.4 \pm 22.5 \text{ mg g}^{-1}$ (Treatment SP12). Similarly, no
194 statistical differences were observed in the levels of ARA, EPA and DHA, nor DHA/EPA ratios
195 in *Artemia* nauplii enriched with varying doses of Se (Table 2).

196 3.2. Experiment 2: effects of different sources of Se on *Artemia* nauplii enrichment efficiency

197 In Experiment 2, Se levels of *Artemia* from Treatments SP0 and SP12 (1.7 ± 0.1 and 4.3 ± 0.4
198 mg kg^{-1} , respectively) were consistent with results from Experiment 1 (1.70 and 4.17 mg kg^{-1} ,
199 respectively). Moreover, *Artemia* nauplii from NS and SL+NS treatments contained Se levels
200 of 1.7 ± 0.0 and $1.7 \pm 0.1 \text{ mg kg}^{-1}$, respectively, very similar to those of the control *Artemia*
201 (Treatment SP0).

202 The different enrichment regimes resulted in variations in the TL content and fatty acid
203 profiles of *Artemia* nauplii (Table 3). While nauplii from Treatments SP0, SP12 and NS
204 showed similar TL contents ($P > 0.05$), nauplii from Treatment SL+NS exhibited significantly
205 higher TL content (Table 3). In terms of fatty acid profiles, the nauplii enriched with SL+NS
206 showed a significantly higher n-6 fatty acid content (Table 3), largely due to the contribution
207 of 18:2n-6 ($16.2 \pm 0.9 \%$) present in the soybean lecithin (Table 1). Consistently, the n-3/n-6
208 ratio was significantly lower in SL+NS nauplii (1.7 ± 0.1) compared to that of the other
209 treatments (~ 3.0). ARA levels were significantly lower in the nauplii SL+NS ($1.7 \pm 0.2 \%$)
210 than in the other treatments. The EPA contents of the SP0 nauplii were overall the highest (4.9
211 $\pm 0.2 \%$), but the difference was only significant when compared to the SL+NS nauplii ($3.7 \pm$
212 0.1%). DHA contents were significantly higher in SP0 and SP12 nauplii (13.8 ± 0.7 and 13.2

213 ± 0.5 %, respectively) compared to SL+NS (10.8 ± 0.3 %). DHA/EPA ratios were not
214 significantly different between treatments.

215

216 **4. Discussion**

217 Nutritional deficiencies play an important role in explaining elevated mortalities in marine fish
218 during the early larval stages (Hamre et al., 2013), themselves resulting from a knowledge gap
219 in our understanding of the digestive development and physiology of marine larvae (Rønnestad
220 et al., 2013). Se is an essential trace element required for a variety of biological functions
221 throughout the entire fish life-cycle including early larval stages (Hamre et al., 2008a; Ribeiro
222 et al., 2012a). Copepods, natural preys of marine finfish larvae, contain relatively high levels
223 of Se compared to live preys and it is therefore critical to develop Se enrichment protocols to
224 guarantee that live preys, otherwise deficient in Se, provide adequate Se levels to meet larvae
225 requirements. Importantly, Se enrichment must be achieved along with the provision of
226 essential fatty acids, micronutrients which are typically encapsulated into live preys with
227 commercial products with very low or, even non-existing, levels of Se.

228 The effects of varying doses of selenoyeast Sel-Plex, a commercial Se additive used for
229 animal feed, were first tested on the levels of Se and essential fatty acids of *Artemia* nauplii.
230 The control treatment in Exp. 1 (Treatment SP0) without Sel-Plex resulted in *Artemia*
231 containing 1.7 ± 0.1 mg Se kg⁻¹, similar to previously published results (Ribeiro et al., 2012b)
232 despite using a different enrichment protocol (i.e. 3 h instead of 20 h in the current study) and
233 enrichment product (i.e. DHA Selco instead of MG in the current study). In the control
234 treatment, Se concentrations of the nauplii reflected the background level of Se present in
235 unenriched *Artemia*, mainly provided by the enrichment product MG, which contained 2.2 mg
236 Se kg⁻¹. Interestingly, the provision of selenoyeast Sel-Plex into the enrichment medium
237 resulted in increased levels of Se in *Artemia* nauplii. The relationship between Sel-Plex and

238 *Artemia* Se was linear thus enabling us to predict Se content in *Artemia* nauplii when enriched
239 with a given Sel-Plex dose. SP12 nauplii showed Se contents ($4.2 \pm 0.1 \text{ mg kg}^{-1}$) within the
240 upper range of Se concentration reported in wild copepods (i.e. 5 mg kg^{-1}) (Hamre et al.,
241 2008b), whereas SP24 and SP36 nauplii, enriched with Sel-Plex doses of 12 and 24 mg l^{-1} ,
242 respectively, contained Se above 10 mg kg^{-1} . Thus, SP12 enrichment treatment resulted in a
243 diet with Se levels well below the potential dietary toxicity threshold observed in rainbow trout
244 and hybrid striped bass ($> 10 \text{ mg kg}^{-1}$; Hilton et al., 1980; Jaramillo et al., 2009). However,
245 SP24 and particularly SP36 treatments resulted in nauplii Se contents that could cause toxicity
246 for fish larvae. When compared to other published studies, the efficiency of Se enrichment
247 obtained in Experiment 1 differs from those reported by Ribeiro et al. (2012a). In the latter
248 study, a Sel-Plex dose of 0.6 mg l^{-1} resulted in a Se content in *Artemia* of $3.11 \pm 0.27 \text{ mg kg}^{-1}$
249 while in our study, a similar Se content in *Artemia* was obtained using 12 mg l^{-1} of Sel-Plex
250 (4.17 mg kg^{-1}). While the reasons explaining such discrepancy remain unknown, Ribeiro's
251 study lacks details on the enrichment protocol used (e.g. rinsing *Artemia* prior sampling,
252 selenium content of Sel-Plex), which could help explain the differences observed in the
253 efficiency of Se incorporation in nauplii. In addition, the lack of data on the fatty acid
254 composition of the enriched nauplii in Ribeiro et al. (2012a) does not allow us to clarify
255 whether the high Se incorporation correlated with a concomitant increase in essential fatty acids
256 within *Artemia* nauplii.

257 Results from Exp. 1 clearly showed that simultaneous delivery of Se and essential fatty
258 acids is possible under our enrichment protocol. Nauplii from SP12 treatment contained, in
259 addition to $4.2 \text{ mg Se kg}^{-1}$, markedly higher levels of essential fatty acids such as DHA (18.0
260 $\pm 1.1 \%$). This data is consistent with an *Artemia* enrichment study that showed the high
261 efficiency of Larviva Multigain at supplying DHA, in which levels of $21.8 \pm 0.7 \%$ DHA post-
262 enrichment were obtained (Cavrois-Rogacki et al., 2019). Furthermore, none of the treatments

263 with Sel-Plex significantly affected the levels of essential fatty acids (i.e. ARA, EPA and DHA)
264 in the *Artemia* compared to the control, despite the potential dilution effect derived from the
265 inclusion of LC-PUFA free lipids from yeast (Santomartino et al., 2017). Therefore, these
266 results showed that Sel-Plex can be successfully used to enrich *Artemia* in Se while preserving
267 the essential fatty acid contents achieved using commercial enrichment products. Our analyses
268 suggested that Sel-Plex does contain traces of EPA and DHA, although their low levels (<0.2
269 %) do not appear to have a major contribution to the essential fatty acids of nauplii.

270 Although it is known that organic Se (e.g. selenomethionine) may be more easily
271 absorbed by living animals compared to inorganic forms (e.g. sodium selenite) (Wang and
272 Lovell, 1997; Izquierdo et al., 2017), high cost of the former can constitute a barrier to their
273 use at a large commercial scale. Cheaper inorganic source of Se have previously been tested
274 on fish larvae fed Se-enriched rotifers (Hamre et al., 2008a) and thus represent potential
275 alternatives. This was investigated in Exp. 2, in which NS and SL+NS treatments consisted of
276 a Se dose of 24 $\mu\text{g l}^{-1}$ (equivalent to Se contained in SP12 treatment with Sel-Plex) supplied as
277 dissolved sodium selenite. Results indicated that neither NS nor SL+NS treatment appeared to
278 be effective ways to enhance Se contents in *Artemia* nauplii since they did not differ from those
279 of control nauplii. While previous studies reported on the low efficiency of delivering dissolved
280 materials into *Artemia* (Tonheim et al., 2000; Monroig et al., 2007), it was somewhat
281 unexpected that delivering the same dose of Se encapsulated into phospholipid vesicles
282 (SL+NS) did not result in any increased Se enrichment efficiency despite *Artemia* being
283 adapted to filtrate discrete particles. The reasons for such a result are unknown but it is
284 reasonable to believe that lipid vesicles produced with the soya lecithin source used in the
285 present study were leaky and did not retain the dissolved Se in the inner aqueous phase of the
286 vesicle. Other highly purified sources of phospholipids, particularly when constituted of more
287 saturated fatty acyl chains, have proven to produce relatively stable vesicles with good

288 efficiency in delivering water soluble compounds into live preys (Hontoria et al., 1994;
289 Monroig et al., 2003, 2007). Thus, in spite of low Se incorporation into *Artemia* nauplii,
290 incorporation of soya lecithin was observed as evidenced by the increased levels of linoleic
291 acid (18:2n-6), its most abundant fatty acid, and the corresponding reduced percentage of other
292 fatty acid contents including EPA, DHA and ARA. The importance of essential fatty acids for
293 marine fish larvae nutrition has been extensively reviewed (Izquierdo, 1996; Tocher 2010,
294 2015) and it is therefore crucial that the enrichment of live preys with micronutrients is not
295 detrimental to the fatty acids' levels of live feed.

296 One important aspect of live prey enrichment is its reproducibility and predictability,
297 which is essential in commercial hatcheries to ensure a constant daily production of high-
298 quality live preys. Producing enriched *Artemia* with consistent levels of essential nutrients such
299 as LC-PUFA can be challenging as shown in previous studies (Navarro et al., 1999; Monroig
300 et al., 2006). Importantly, the results from the present study suggests that enrichment of
301 *Artemia* nauplii with selenoyeast Sel-Plex is highly reproducible, according to the consistent
302 levels of Se found in nauplii from the SP12 treatment in two independent experiments (Exp. 1
303 to 2). While this proves the reproducibility of the method on a small-scale enrichment system,
304 further trials are necessary to confirm the reproducibility at commercial scale.

305 The study showed that it is possible to enrich *Artemia* with targeted levels of Se using
306 selenoyeast Sel-Plex. Enriching *Artemia* nauplii with 12 mg of Sel-Plex per litre for 4 h prior
307 to a 24 h enrichment with LC-PUFA rich commercial diets produces *Artemia* with Se contents
308 similar to those found in the natural preys (wild zooplankton) of marine fish larvae and high
309 levels of essential fatty acids. The use of inorganic Se was not an effective strategy to enrich
310 *Artemia* nauplii even when it was delivered through phospholipid vesicles. In the case of soya
311 lecithin, the use of low-quality liposomes in the experiment could be a potential cause and

312 should be deeper investigated. Ultimately, these results can be implemented to the *Artemia*
313 enrichment protocols of any marine fish species.

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474 **Table 1.** Contents of Selenium (Se), total lipids, and selected fatty acids in enrichment products
 475 used in Experiments 1 and 2.

	Larviva Multigain	Soya lecithin	Sel-Plex
Se (mg kg ⁻¹)	2.2	ND	2000
Total lipids (mg g ⁻¹)	397.4	808.6	13.4
<i>Fatty acids (% of total)</i>			
14:0	6.1	0.1	0.6
15:0	0.4	ND	0.4
16:0	32.3	19.0	15.3
18:0	0.9	4.2	5.2
Saturates	40.1	24.2	22.2
16:1n-9	0.3	ND	ND
16:1n-7	0.1	0.8	25.1
18:1n-9	1.9	10.2	28.5
18:1n-7	ND	2.0	ND
Monounsaturates	2.3	13.4	54.3
18:2n-6	2.2	54.1	14.7
18:3n-6	0.2	ND	0.1
20:4n-6	1.2	ND	0.1
22:5n-6	14.4	ND	ND
Total n-6	18.4	54.3	15.0
18:3n-3	0.3	6.9	3.9
18:4n-3	0.3	0.1	0.2
20:3n-3	0.1	ND	0.1
20:4n-3	0.7	ND	ND
20:5n-3	0.8	ND	0.1
22:5n-3	0.3	ND	ND
22:6n-3	36.6	ND	0.2
Total n-3	39.0	7.0	4.3

DW, dry weight; ND, not detected.

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482 **Table 2.** Total lipids and selected fatty acids of *Artemia* nauplii from Experiment 1 enriched
 483 for 4 h with different doses of selenoyeast Sel-Plex (SP0: 0 mg l⁻¹; SP12: 12 mg l⁻¹; SP24: 24
 484 mg l⁻¹; SP36: 36 mg l⁻¹) followed by a 24 h enrichment with Larviva Multigain (0.6 g l⁻¹). Data
 485 are expressed as means ± standard deviations (*n* = 3). Differences among treatments were
 486 analysed by a one-way ANOVA followed by a Tukey post-hoc test (*P* ≤ 0.05).

Treatment	SP0	SP12	SP24	SP36
Total lipids (mg g ⁻¹ DW)	209.5 ± 9.0	220.4 ± 22.5	212.5 ± 8.7	211.9 ± 8.9
<i>Fatty acids (% of total)</i>				
14:0	1.6 ± 0.1	1.5 ± 0.0	1.7 ± 0.3	1.7 ± 0.3
15:0	0.2 ± 0.0	0.2 ± 0.0	0.2 ± 0.0	0.2 ± 0.0
16:0	14.6 ± 0.9	14.2 ± 0.3	16 ± 1.9	16.8 ± 3.0
18:0	4.4 ± 0.1	4.3 ± 0.3	4.6 ± 0.3	5.1 ± 1.2
Saturates	21.2 ± 1.1	20.6 ± 0.6	22.8 ± 2.5	24.3 ± 4.6
16:1n-9	0.5 ± 0.0	0.4 ± 0.1	0.5 ± 0.0	0.5 ± 0.2
16:1n-7	1.3 ± 0.1	1.2 ± 0.1	1.5 ± 0.0	1.5 ± 0.6
18:1n-9	13.4 ± 0.4	12.7 ± 0.9	14.6 ± 0.1	15.7 ± 4.8
18:1n-7	4.0 ± 0.1	3.9 ± 0.2	4.3 ± 0.0	4.7 ± 1.3
Monounsaturates	19.7 ± 0.6	18.8 ± 1.4	21.3 ± 0.1	23.0 ± 7.0
18:2n-6	4.7 ± 0.2	4.6 ± 0.1	4.8 ± 0.3	4.8 ± 0.3
18:3n-6	0.4 ± 0.0	0.4 ± 0.0	0.4 ± 0.0	0.4 ± 0.0
20:4n-6	2.6 ± 0.1	2.8 ± 0.3	2.1 ± 0.2	2.1 ± 1.5
22:5n-6	6.7 ± 0.2	7.0 ± 0.6	5.6 ± 0.5	4.9 ± 3.0
Total n-6	14.8 ± 0.3	15.4 ± 0.9	13.3 ± 0.5	12.7 ± 4.3
18:3n-3	16.9 ± 0.6	16.7 ± 0.3	18.5 ± 2.8	18.6 ± 2.2
18:4n-3	2.5 ± 0.1	2.4 ± 0.1	2.9 ± 0.6	2.8 ± 0.7
20:3n-3	0.8 ± 0.0	0.8 ± 0.0	0.8 ± 0.1	0.8 ± 0.0
20:4n-3	0.9 ± 0.0	0.9 ± 0.0	0.9 ± 0.1	0.8 ± 0.2
20:5n-3	5.4 ± 0.2	5.9 ± 0.6	4.4 ± 0.4	4.3 ± 2.9
22:5n-3	0.4 ± 0.0	0.3 ± 0.2	0.2 ± 0.2	0.2 ± 0.3
22:6n-3	17.3 ± 0.4	18.0 ± 1.1	14.5 ± 1.3	12.2 ± 6.9
Total n-3	44.1 ± 0.8	45.0 ± 1.2	42.2 ± 2.5	39.7 ± 7.4
n-3/n-6	3.0 ± 0.1	2.9 ± 0.1	3.2 ± 0.2	3.2 ± 0.5
DHA/EPA	3.2 ± 0.0	3.1 ± 0.2	3.3 ± 0.1	3.0 ± 0.4
Total FA (mg g ⁻¹ DW)	137.6 ± 11.0	136.0 ± 7.4	129.1 ± 2.8	116.5 ± 25.2

DW: dry weight; FA: fatty acids.

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490 **Table 3.** Total lipids and selected fatty acids in the *Artemia* nauplii from Experiment 2 treated
 491 with different enrichment diets (MG: Larviva Multigain; SP: Sel-Plex; NS: sodium selenite;
 492 SL: soya lecithin emulsion). Data are expressed as means \pm standard deviations ($n = 3$).
 493 Differences in fatty acid contents among treatments were analysed by a one-way ANOVA
 494 followed by a Tukey post-hoc test ($P \leq 0.05$). Variables that do not share the same superscript
 495 letter within a row are significantly different from each other.

Treatment	SP0	SP12	NS	SL+NS
Total lipids (mg g ⁻¹ DW)	177.1 \pm 8.6 ^a	180.1 \pm 2.4 ^a	184.7 \pm 2.9 ^a	213.9 \pm 7.8 ^b
<i>Fatty acids (% of total)</i>				
14:0	1.3 \pm 0.0 ^{ab}	1.3 \pm 0.1 ^{ab}	1.5 \pm 0.0 ^b	1.2 \pm 0.1 ^a
15:0	0.2 \pm 0.0	0.2 \pm 0.0	0.2 \pm 0.0	0.2 \pm 0.0
16:0	13.1 \pm 0.5	13.0 \pm 0.2	14.8 \pm 2.1	13.0 \pm 0.3
18:0	4.3 \pm 0.2	4.2 \pm 0.1	4.7 \pm 0.6	4.2 \pm 0.1
Saturates	19.3 \pm 0.7	18.9 \pm 0.3	21.6 \pm 2.8	19.0 \pm 0.3
16:1n-9	0.6 \pm 0.0	0.6 \pm 0.0	0.4 \pm 0.2	0.5 \pm 0.0
16:1n-7	1.1 \pm 0.9	1.6 \pm 0.0	1.5 \pm 0.1	1.3 \pm 0.1
18:1n-9	14.8 \pm 0.2 ^{ab}	15.1 \pm 0.4 ^b	14.9 \pm 0.5 ^{ab}	13.7 \pm 0.6 ^a
18:1n-7	4.4 \pm 0.1 ^b	4.5 \pm 0.1 ^b	4.5 \pm 0.2 ^b	3.8 \pm 0.1 ^a
Monounsaturates	21.3 \pm 0.6 ^b	22.1 \pm 0.4 ^b	21.7 \pm 0.7 ^b	19.7 \pm 0.7 ^a
18:2n-6	5.2 \pm 0.2 ^a	5.3 \pm 0.1 ^a	5.2 \pm 0.1 ^a	16.2 \pm 0.9 ^b
18:3n-6	0.4 \pm 0.1	0.4 \pm 0.0	0.4 \pm 0.0	0.4 \pm 0.0
20:4n-6	2.3 \pm 0.2 ^b	2.2 \pm 0.1 ^b	2.3 \pm 0.1 ^b	1.7 \pm 0.2 ^a
22:5n-6	5.3 \pm 0.1 ^b	4.9 \pm 0.2 ^b	4.8 \pm 0.4 ^{ab}	4.3 \pm 0.1 ^a
Total n-6	13.7 \pm 0.0 ^a	13.2 \pm 0.2 ^a	13.1 \pm 0.4 ^a	23.0 \pm 0.8 ^b
18:3n-3	21.4 \pm 0.4 ^b	22.1 \pm 0.5 ^b	21.4 \pm 1.1 ^b	19.3 \pm 0.6 ^a
18:4n-3	3.3 \pm 0.1 ^b	3.4 \pm 0.1 ^b	3.3 \pm 0.1 ^b	2.7 \pm 0.2 ^a
20:3n-3	0.8 \pm 0.0 ^b	0.8 \pm 0.0 ^b	0.8 \pm 0.0 ^b	0.7 \pm 0.0 ^a
20:4n-3	0.9 \pm 0.0 ^b	0.9 \pm 0.0 ^b	0.9 \pm 0.0 ^b	0.7 \pm 0.0 ^a
20:5n-3	4.9 \pm 0.2 ^b	4.7 \pm 0.1 ^{ab}	4.5 \pm 0.7 ^{ab}	3.7 \pm 0.1 ^a
22:5n-3	0.2 \pm 0.2	0.2 \pm 0.2	0.3 \pm 0.0	0.1 \pm 0.1
22:6n-3	13.8 \pm 0.7 ^b	13.2 \pm 0.5 ^b	12.0 \pm 1.4 ^{ab}	10.8 \pm 0.3 ^a
Total n-3	45.4 \pm 1.0 ^b	45.3 \pm 0.1 ^b	43.1 \pm 2.6 ^b	38.0 \pm 0.7 ^a
n-3/n-6	3.3 \pm 0.1 ^b	3.4 \pm 0.1 ^b	3.3 \pm 0.2 ^b	1.7 \pm 0.1 ^a
DHA/EPA	2.8 \pm 0.3	2.8 \pm 0.1	2.7 \pm 0.2	2.9 \pm 0.1
Total FA (mg g ⁻¹ DW)	122.1 \pm 7.0 ^a	122.7 \pm 1.3 ^a	121.4 \pm 2.4 ^a	155.8 \pm 12.1 ^b

DW: dry weight; FA: fatty acids.

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498 **Figure 1.** Selenium (Se) concentration of *Artemia* nauplii ($\mu\text{g Se g}^{-1}$ dry weight, “DW”) from
499 Experiment 1 enriched 4 h with different dose of selenoyeast Sel-Plex (SP0: 0 mg l^{-1} ; SP12: 12
500 mg l^{-1} ; SP24: 24 mg l^{-1} ; SP36: 36 mg l^{-1}) followed by a 24 h enrichment with Larviva Multigain
501 (0.6 g l^{-1}). Data are expressed as means \pm standard deviations ($n = 3$). Differences in Se contents
502 among treatments were analysed by a one-way ANOVA followed by a Tukey post-hoc test ($P \leq$
503 0.05). Treatments with different superscripts are significantly different from each other.

504 **Figure 2.** Selenium (Se) concentration of *Artemia* nauplii ($\mu\text{g Se g}^{-1}$ dry weight, “DW”) from
505 Experiment 2 enriched with different treatments. SP0: SP (0 mg l^{-1} , 4 h); SP12: SP (12 mg l^{-1} ,
506 4 h); NS: NS (24 $\mu\text{g l}^{-1}$, 4 h); SL+NS: SL emulsion (0.6 mg l^{-1}) + NS (24 $\mu\text{g l}^{-1}$ emulsion). All
507 treatments were followed by a second enrichment with MG (0.6 g l^{-1} , 24 h) Data are expressed
508 as means \pm standard deviations ($n = 3$). Differences in fatty acid contents among treatments
509 were analysed by a one-way ANOVA followed by a Tukey post-hoc test ($P \leq 0.05$). Variables
510 that do not share the same superscript letter within a row are significantly different from each
511 other.

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513 **Figure 1**

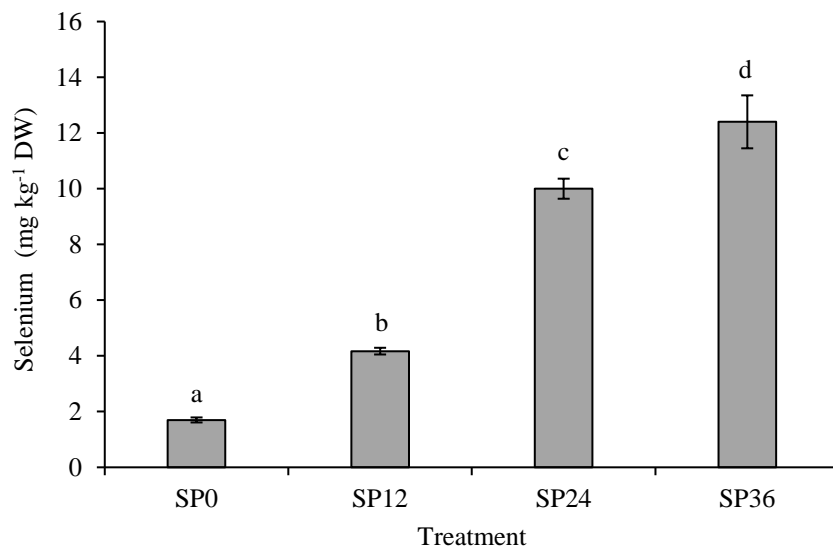


Figure 2