

1 **Evaluation of different feeding protocols for larvae of Atlantic bluefin**
2 **tuna (*Thunnus thynnus*, L.)**
3

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18 Accepted refereed manuscript of:

Betancor M, Ortega A, de la Gándara F, Varela JL, Tocher DR & Mourente G (2019) Evaluation of different
19 feeding protocols for larvae of Atlantic bluefin tuna (*Thunnus thynnus*, L.). *Aquaculture*, 505, pp. 523-538.

DOI: <https://doi.org/10.1016/j.aquaculture.2019.02.063>

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21 **Abstract**

22 Mass mortality is still one of the main constraints in larval rearing of Atlantic bluefin tuna (*Thunnus*
23 *thynnus* L.; ABT). Early data related to the feeding sequence of ABT larvae suggested that mortality
24 observed during the first stages of life could be partly due to nutritional deficiencies. Previous studies
25 demonstrated that copepods appeared to be a superior live prey compared to rotifers during the first
26 two weeks of life. Our overarching aim was to evaluate different feeding strategies during first
27 feeding of ABT larvae from a performance, compositional and molecular perspective. In order to do
28 so, two groups of ABT larvae were fed with either copepod (*Acartia tonsa*; C) nauplii or rotifers
29 (*Brachionus rotundiformis*; R) enriched with Algamac 3050[®] from mouth opening to 13 days after
30 hatching (dah). After this, the group C-larvae was fed either *Artemia* enriched with Algamac 3050[®]
31 (CA), *Acartia* nauplii and copepodites (CC) or sea bream (*Sparus aurata*) yolk-sac larvae (CY), while
32 the R group were fed on *Artemia* enriched with Algamac 3050[®] (RA) up to 18 dah. At 13 dah, larvae
33 fed copepods (C) had grown better than those fed enriched rotifers (R) although there were no
34 significant differences in survival. ABT larvae fed R accumulated highest eicosapentaenoate (EPA)
35 but lowest docosahexaenoate (DHA) and total n-3 long-chain polyunsaturated fatty acids (LC-PUFA)
36 than C-fed larvae, reflecting the dietary contents. There was no activation in the expression of the
37 enzymes involved in EPA and DHA biosynthesis. However, the different live prey showed regulation
38 of transcription factor, digestive enzyme, lipid metabolism and oxidative stress genes. At 18 dah,
39 larvae fed CY and CA treatments were largest in size, with larvae fed RA displaying the lowest
40 growth, with no significant differences in survival among the dietary treatments. The highest DHA
41 contents were found in ABT larvae fed CC and CY, whereas the lowest contents were found in RA-
42 fed larvae. Indeed, larvae fed RA showed the highest level of the intermediate product n-3
43 docosapentaenoate, which could reflect increased activity of the biosynthetic pathway although this
44 was not supported by gene expression data.

45 **Keywords:** Bluefin tuna, larvae, rotifers, copepods, *Artemia*, yolk sac larvae, lipid metabolism,

46 digestibility, antioxidant status, gene expression.

47 *Abbreviations:* ABT, Atlantic bluefin tuna; *aco*, acyl coA oxidase; *alp*, alkaline phosphatase; *amy*,
48 amylase; *anpep*, amino peptidase; ARA, arachidonic acid (20:4n-6); *ball1*, bile salt activated lipase 1;
49 *bal2*, bile salt activated lipase 2; *cat*, catalase; *cpt1*, carnitine palmitoyl transferase I; dah, days after
50 hatch; DHA, docosahexaenoic acid (22:6n-3); DPA, docosapentaenoic acid (n-3 or n-6); EPA,
51 eicosapentaenoic acid (20:5n-3); *elovl5*, fatty acyl elongase 5; *fabp2*, fatty acid binding protein 2
52 (intestinal); *fabp4*, fatty acid binding protein 4 (adipocyte); *fabp7*, fatty acid binding protein 7 (brain-
53 type); *fads2d6*, delta-6 fatty acyl desaturase; *fas*, fatty acid synthase; *gpx1*, glutathione peroxidase 1;
54 *gpx4*, glutathione peroxidase 4; *hmgcl*, 3-hydroxy-3-methylglutaryl-CoA lyase; LA, linoleic acid
55 (18:2n-6); LNA, α -linolenic acid (18:3n-3); LC-PUFA, long-chain polyunsaturated fatty acid; LPC,
56 lyso phosphatidylcholine; *lpl*, lipoprotein lipase; *lxr*, liver X receptor; *myhc*, myosin heavy chain;
57 PBT, Pacific bluefin tuna; *pl*, pancreatic lipase; *pla2*, phospholipase A₂; *ppara*, peroxisome
58 proliferator-activated receptor alpha; *ppar γ* , peroxisome proliferator-activated receptor gamma; *rxr*,
59 retinoid X receptor; *sod*, superoxide dismutase; *srebp1*, sterol regulatory element-binding protein 1;
60 *srebp2*, sterol regulatory element-binding protein 2; TF, transcription factor; *tropo*, tropomyosin;
61 *tryp*, trypsin.

62

63 1. Introduction

64 Regardless of the efforts and progress that have been made in the Mediterranean area in the
65 larval rearing of Atlantic bluefin tuna (ABT; *Thunnus thynnus*), a number of challenges and issues
66 continue to restrict the hatchery production of ABT fingerlings in commercial quantities with early
67 “mass mortality” during first feeding being a common occurrence. (De la Gándara et al., 2016; Van
68 Beijnen, 2017). Aside from improvement of zootechnical aspects of the culture system focussed on
69 broodstock management and egg and larval production and quality, the refinement of the live food
70 trophic chain (prey size, sequence and nutritional quality) is necessary until a reliable artificial
71 commercial diet is fully developed. Currently, ABT larval first feeding is attained using either
72 enriched rotifers (preferably *Brachionus rotundiformis*) and/or copepod nauplii (*Acartia tonsa*) as
73 initial live prey (De la Gandara et al., 2010; 2016; Betancor et al., 2017a,b). Previous studies have
74 shown that copepods were better live prey for first feeding ABT based on growth and survival data
75 (Betancor et al., 2017a). In addition, broodstock nutrition was also identified as a possible factor
76 explaining differences in growth performance and lipid metabolism observed between larvae from
77 different year classes (Betancor et al., 2018).

78 The metabolism and deposition of lipids is a complex process in fish, involving multiple
79 pathways such as lipogenesis, β -oxidation and biosynthesis of long-chain polyunsaturated fatty acids
80 (LC-PUFA) (Ayisi et al., 2018). In a recent trial where ABT larvae were fed on rotifers enriched with
81 five commercial products or copepods nauplii, responses in lipid gene expression, which are a
82 consequence of differences in dietary lipid contents and fatty acid compositions, appeared to indicate
83 that lipid levels provided by enriched rotifers exceeded ABT requirements (Betancor et al., 2017b).
84 This study corroborated that copepods were a superior live prey when compared to rotifers in first
85 feeding ABT larvae, as indicated by the higher growth achieved by copepod-fed larvae, possibly
86 reflecting the higher protein content of the copepods (Betancor et al., 2017b). However, the mass
87 production of *Acartia* copepods is labourious and costly in comparison to the production of enriched

88 rotifers or *Artemia*, which explains why copepods are not widely used as a live prey in hatcheries for
89 any species of marine fish.

90 In the present study, further feeding trials were performed to expand our knowledge of early
91 nutrition and nutritional requirements of first feeding ABT larvae. Our overarching aim was to
92 evaluate different strategies during first feeding (up to weaning onto an artificial, formulated diet) of
93 ABT larvae from a performance, compositional and molecular perspective. In order to do so, two
94 groups of ABT larvae were fed with either copepod (*Acartia tonsa*; C) nauplii, or rotifers (*Brachionus*
95 *rotundiformis*; R) enriched with Algamac 3050[®] from mouth opening to 13 dah. After this, the C-
96 larvae group was fed either *Artemia* enriched with Algamac 3050[®] (CA), *Acartia* nauplii and
97 copepodites (CC) or sea bream (*Sparus aurata*) yolk-sac larvae (CY), while the R-larvae group were
98 fed on *Artemia* enriched with Algamac 3050[®] (RA) up to 18 dah. Growth performance,
99 developmental indices and survival were determined. Additionally, the expression of genes related to
100 lipid metabolism (transcription factors, fatty acid metabolism and lipid homeostasis), antioxidant
101 enzymes, myogenesis and digestive enzymes was carried out in order to assess the impacts of the
102 different feeding protocols on larval metabolism and physiology.

103

104 **2. Materials and Methods**

105 *2.1 Atlantic bluefin tuna larvae rearing conditions*

106 The ABT eggs used in this study were obtained in June 2017 from ABT broodstock fish
107 maintained in captivity in a floating net cage located at El Gorguel, off the Cartagena coast, SE Spain.
108 Captive-reared ABT broodstock fish spawned naturally and spontaneously, and floating eggs were
109 collected inside the cage by means of a net of 500 µm mesh screen size. A 1.5 m polyvinyl sheet was
110 also placed around the inside of the cage to avoid eggs drifting away from the cage (or into the cage)
111 by means of currents and/or waves. Collected eggs were transported in a 500 L plastic tank supplied
112 with pure oxygen to the Spanish Institute of Oceanography (IEO) Planta Experimental de Cultivos

113 Marinos (Puerto de Mazarrón, Murcia, Spain) aquaculture facilities and placed in 100 L tanks with
114 gentle oxygenation and flow-through sterilized seawater. After 1 h, aeration and water flow were
115 stopped to separate buoyant (viable) from non-buoyant (non-viable) eggs. After washing and
116 counting, the fertilized eggs were incubated in 1400 L cylindrical tanks at a density of 8.5 eggs.L⁻¹.
117 Incubation was carried out at a water temperature 23 - 25 °C, 37 ‰ salinity, dissolved oxygen 6.5
118 mg.L⁻¹ and continuous photoperiod, with a light intensity of 1000 lux. An upwelling flow-through
119 with gentle aeration was employed in order to maintain oxygen levels near to saturation. Larvae
120 hatched approximately 32 h after fertilization, with a hatching rate of almost 90 %, and were fed with
121 enriched rotifers (R-larvae) or copepod nauplii (C-larvae) from 2 dah to 13 dah. A mixture of the
122 microalgae *Isochrysis* sp. (T-Iso) and *Chlorella* (V12 DHA-enriched, Pacific Trading Co., Japan)
123 were added to tanks at a density of 2 - 3 x10⁵ cells. mL⁻¹ as green water. During the feeding trial,
124 photoperiod was maintained at 14 h / 10 h light/dark (light intensity about 500 lux), temperature
125 ranged between 23 - 25 °C and daily water renewal was 100 - 200 % tank volume.day⁻¹. Incoming
126 seawater was filtered at 10 µm and UV sterilized. An upwelling current was created to avoid larvae
127 sinking (mainly at night) and to maintain oxygen level (Ortega, 2015; De la Gándara et al., 2016;
128 Betancor et al., 2017a,b). The rearing conditions for feeding trials are summarized in Supplementary
129 Table 1.

130

131 2.2 Dietary trial 1 with live prey, rotifers and copepod nauplii, from first feeding up to 13 dah

132 In the first trial, ABT larvae at a density 8.5 larvae.L⁻¹ were fed the two different live prey in
133 quadruplicate (4 tank replicates/treatment) from mouth opening (2 dah) up to 13 dah (Supplementary
134 Table 1). Larvae were fed either with rotifer *Brachionus rotundiformis* enriched with Algamac 3050®
135 (treatment R) or *Acartia tonsa* copepod nauplii (treatment C). To maintain constant live prey
136 concentration of 10 rotifer.mL⁻¹ or copepod nauplii/copepodite.mL⁻¹ within each experimental tank,
137 three water samples (10 mL) from each tank were counted twice per day before supplying new feed
138 (Ortega, 2015; De la Gándara et al., 2016; Betancor et al., 2017a,b).

139

140 2.3 Dietary trial 2 from 13 to 18 dah

141 After sampling ABT larvae at 13 dah, the remaining larvae were redistributed into four 1,400
142 L tanks at a density of 0.43 larvae.L⁻¹ (Supplementary Table 1). The dietary treatments in single
143 replicate tanks were established as follows: group C-larvae (ABT larvae that had been previously fed
144 on *A. tonsa* nauplii) were given three different feeds, being either *Artemia* enriched with Algamac
145 3050[®] at 5 metanauplii.mL⁻¹ (treatment CA), *Acartia* nauplii and copepodites at 10
146 nauplii/copepodites.mL⁻¹ (treatment CC) or gilthead sea bream (*Sparus aurata*) yolk-sac larvae at 5
147 larvae.mL⁻¹ (treatment CY). The R group (ABT larvae that had been fed on rotifer *Brachionus*
148 *rotundiformis* enriched with Algamac 3050[®]) were passed on to being fed only *Artemia* enriched
149 with Algamac 3050[®] (treatment RA) at 5 metanauplii.mL⁻¹ to represent a reference treatment
150 reflecting the common commercial protocol for marine fish larvae. All four groups were fed these
151 treatments from 13 dah to 18 dah when larvae were again sampled.

152

153 2.4 Rotifer culture and enrichment protocol

154 S-type rotifers *B. rotundiformis* were continuously cultured with commercial DHA-enriched
155 algal paste (Chlorella V-12; Chlorella Industry, Kyushu, Japan), at a concentration of 3 mL Chlorella
156 paste per 10⁶ rotifers per day, in four 2,000 L cylindro-conical tanks supplied with filtered and
157 sterilized sea water at 24 – 26 °C, 38 ‰ salinity, dissolved oxygen at saturation level and 24 h
158 continual illumination. Enrichment treatment consisted of the commercial product, Algamac 3050[®].
159 Additionally, rotifers were supplemented with taurine (0.5 g per 10⁶ rotifers), organic Se (Selplex[®]
160 Alltech Spain SL; 3.0 mg per 10⁶ rotifers), and vitamin E, as dl- α tocopheryl acetate (Lutavit E50;
161 BASF; 0.9 mg per 10⁶ rotifers), 18 h before the enrichment treatment. The enrichment protocols
162 were performed in 100 L cylindro-conical tanks at a density of 1000 rotifers.mL⁻¹, adding a dose of
163 the enrichment product of 0.3 g plus 3 mg Selplex[®] and 0.9 mg vitamin E per 10⁶ rotifers over a
164 period of 6 h for Algamac 3050[®] according to manufacturer's recommendations.

165

166 *2.5 Cultivation of the copepod Acartia tonsa*

167 The copepods (*A. tonsa*) were cultivated in 4,000 L cylindrical tanks with seawater of 34 ‰
168 salinity at 20 °C and were continuously fed with algae *Rhodomonas baltica* at a concentration not
169 below 3×10^4 cells.mL⁻¹. *Acartia* eggs were obtained every day with a harvesting arm to collect the
170 eggs deposited on the flat bottom of the tanks. The eggs were washed thoroughly and stored in flasks
171 at 2 °C. Egg harvest started 3 months in advance of the ABT feeding trial and continued until the end
172 of the trial. The water in the flasks was renewed every 2 weeks and the number of eggs counted. The
173 copepod eggs were incubated in 100 L tanks at a maximum density of 150 eggs.mL⁻¹. From 2 days
174 after hatch the nauplii and copepodites were fed *ad libitum* with a mixture of *R. baltica* and *Isochrysis*
175 *galbana* clone T-Iso and, before harvesting, the nauplii/copepodite density in the tanks was estimated,
176 harvested with a siphon, concentrated in a 60 µm sieve and then transferred to the ABT larval tanks.

177

178 *2.6 Artemia metanauplii enrichment and gilthead sea bream yolk sac larvae*

179 *Artemia* cysts EG type were decapsulated and enriched with Algamac 3050[®] over 12 h before
180 being fed to ABT larvae. Recently hatched nauplii were disinfected with bronopol (100 ppm;
181 Pyceze[®]; Novartis). Enriched *Artemia* metanauplii were deposited in the tanks at a density of 0.1-
182 0.2 metanauplii.mL⁻¹. Before reproduction, gilthead sea bream broodstock were fed Vitalis CAL and
183 Vitalis REPRO (Skretting[®]) at a rate of 0.6 – 0.8 % of broodstock biomass. Gilthead sea bream 1 dah
184 yolk sac larvae were added to tanks at a density of 5-10 larvae.L⁻¹.

185

186 *2.7 Sampling for biometrical, biochemical and molecular analysis*

187 Thirty and twenty randomly caught ABT larvae per replicate treatment were anaesthetized
188 (0.02 % 2-phenoxyethanol, Sigma, Spain) in Trial 1 and 2, respectively, total length measured and
189 individual larvae photographed. The tank was considered the experimental unit in Trial 1 (n = 4),
190 where individual fish were utilized in Trial 2 (n = 20). The developmental stage was assessed by

191 counting the number of ABT larva which had attained full flexion of the notochord by the end of the
192 feeding trials (13 and 18 dah) in each replicate set of samples. Individual larvae dry mass was
193 determined on a precision balance after maintaining samples at 110 °C for 24 h and cooling *in vacuo*
194 for 1 h before weighing. Final survival (%) was calculated by counting individual live larvae at the
195 beginning and end of the trial.

196 In trial 1, three samples of 15 larvae per sample of 13 dah larvae were collected per tank: i)
197 two samples/tank were placed in 1 ml of RNAlater[®] (Sigma, Madrid, Spain) for RNA extraction and
198 molecular analysis (n = 8), and ii) one sample/tank was frozen in liquid N₂ and stored at -80 °C for
199 biochemical analysis (n = 4). In trial 2, nine samples of 15 larvae of 18 dah larvae were collected per
200 tank with 6 samples/tank used for molecular analysis (n = 6) and 3 samples/tank used for biochemical
201 analyses, respectively (n = 3). Triplicate samples of enriched rotifers, copepods (*Acartia*), enriched
202 *Artemia metanauplii*, and 1 dah sea bream yolk sac larvae were filtered and washed, excess water
203 drained and blotted with filter paper, immediately frozen in liquid N₂ and stored at -80 °C prior to
204 analysis. All procedures were carried out according to the current national and EU legislation on the
205 handling of experimental animals.

206

207 2.8 Biochemical analysis.

208 2.8.1 Proximate gross composition

209 Proximate compositions of live feeds (protein and lipid) were determined according to
210 standard procedures (AOAC, 2000). Three technical replicates of feeds (single batch production)
211 were freeze-dried prior to analyses. Ash content determined after incineration at 600 °C for 16 h.
212 Crude protein was measured by determining nitrogen content (N × 6.25) using automated Kjeldahl
213 analysis (Tecator Kjeltex Auto 1030 analyser, Foss, Warrington, UK), and crude lipid content
214 determined gravimetrically after Soxhlet lipid extraction (Tecator Soxtec system 2050 Auto
215 Extraction apparatus). Carbohydrate content was calculated as 100 – (percentages of protein + lipid

216 based on dry weight). The gross energy content was calculated from gross composition data and using
217 values of 5.65, 9.45 and 4.20 kcal.g⁻¹ for protein, lipid and carbohydrates, respectively (Henken *et al.*
218 1986).

219

220 *2.8.2. Preparation of hydrolysates, derivatisation, UPLC analysis of taurine and total amino acid*
221 *content of live prey and ABT larvae, and calculation of amino acid sufficiency index*

222 A Waters AccQ-Tag Ultra Method[®] was used in determining taurine and amino acids contents
223 in samples of enriched rotifers *B. rotundiformis* and 13 dah ABT larvae. Hydrolysis and derivatization
224 were carried out according to manufacturer's instructions, and amino acid analysis (including taurine)
225 was performed using a Waters H-Class UPLC fitted with an ACQUITY BEH Phenyl 1.7μ UPLC
226 column (AAA for H-Class System Guide, Waters Corporation 2012). An essential amino acid
227 sufficiency index (Sla.a.) was calculated by assuming the total amino acid profile/content of 1 dah ABT
228 yolk sac larvae as an indicator of ABT larvae amino acid requirements. The index was calculated by
229 dividing the content of a determined amino acid in a live prey by the content of the same amino acid in
230 1 dah ABT larvae and multiplying the result by 100.

231
$$\text{Sla.a.} = ([\text{a.a. prey}] / [\text{a.a. 1 dah ABT larvae}]) \times 100$$

232 Values of a Sla.a. equal or above 100 will indicate sufficient dietary amount of that amino acid
233 in the live prey, whereas values below 100 show potential insufficiency for that amino acid (Oser, 1959).

234

235 *2.8.3. Total lipid, lipid class composition and fatty acid analysis*

236 Total lipid of live feeds (enriched rotifers, copepods, enriched *Artemia* and gilthead sea bream
237 yolk sac larvae) and ABT larvae fed the different dietary regimes was extracted from triplicate pooled
238 samples according to the method of Folch et al. (1957). Approximately 200 mg samples of feeds/ABT

239 larvae were placed in 10 mL of ice-cold chloroform/methanol (2:1, by vol) and homogenized with an
240 Ultra-Turrax tissue disrupter (Fisher Scientific, Loughborough, U.K.). The non-lipid and lipid layers
241 were separated by addition of 5 mL of 0.88 % (w/v) KCl and allowed to separate on ice for 1 h. The
242 upper aqueous layer was aspirated and the lower organic layer dried under oxygen-free nitrogen. The
243 lipid content was determined gravimetrically after drying overnight in a vacuum desiccator.

244 Lipid class composition was determined by high-performance thin-layer chromatography
245 (HPTLC) using 10 x 10 cm plates (VWR, Lutterworth, England). Approximately 1 µg of total lipid
246 was applied as a single spot, and the plates developed in methyl acetate/isopropanol/
247 chloroform/methanol/0.25 % aqueous KCl (25:25:25:10:9, by vol.) to two-thirds up the plate. After
248 drying for 20 min, the plate was fully developed with isohexane/diethyl ether/acetic acid (85:15:1, by
249 vol.). The lipid classes were visualized by charring at 160 °C for 15 min after spraying with 3 % (w/v)
250 aqueous cupric acetate containing 8 % (v/v) phosphoric acid and quantified by densitometry using a
251 CAMAG-3 TLC scanner (version Firmware 1.14.16). Scanned images were recorded automatically
252 and analyzed using winCATS Planar Chromatography Manager software (version 1.2.0) (Henderson
253 and Tocher, 1992).

254 Fatty acid methyl esters (FAME) from the extracted total lipids were prepared by acid-
255 catalyzed transesterification at 50 °C for 16 h according to the method of Christie (1993). Methyl
256 esters were separated and quantified by gas-liquid chromatography (Agilent Technologies 7890B GC
257 System) using a 30 m x 0.32 mm i.d. fused silica capillary column (SUPELCOWAX™-10, Supelco
258 Inc., Bellefonte, USA) and on-column injection at 50 °C. Hydrogen was used as carrier gas and
259 temperature programming was from 50 °C to 150 °C at 40 °C per min and then to 230 °C at 2.0 °C
260 per min. Individual methyl esters were identified by comparison with known standards and by
261 reference to published data (Ackman, 1980; Tocher and Harvie, 1988). Data were collected and
262 processed using Agilent Technologies Openlab CDS Chemstation for Windows (version A.02.05.21).

263

264 *2.8.4. Determination of alpha-tocopherol (vitamin E) content*

265 Alpha-tocopherol concentrations in enriched rotifers and *Artemia* as well as copepods were
266 determined using high-pressure liquid chromatography (HPLC) with UV detection. Samples were
267 weighed, homogenized in pyrogallol, and saponified as described by McMurray et al. (1980) and
268 Cowey et al. (1981). HPLC analysis was performed using a 150 x 4.60 mm, reverse-phase Luna 5 μ m
269 C18 column (Phenomenox, CA, USA). The mobile phase was 98 % methanol pumped at 1.0 mL.min⁻¹.
270 The effluent from the column was monitored at a wavelength of 293 nm and quantification achieved
271 by comparison with alpha-tocopherol (Sigma-Aldrich) as external standard.

272

273 2.8.5. *Selenium determination*

274 Total selenium concentration was measured in feeds according to the method established by
275 Betancor et al. (2012). Dried samples were weighed in three replicates of between 0.04 and 0.1 g and
276 digested in a microwave digester (MarsXpress, CEM, USA) with 5 % of 69 % pure nitric acid in
277 three steps as follows; 21° C to 190° C for 10 min at 800 W, then 190° C for 20 min at 800 W, and
278 finally a 30 min cooling period. The digested solution was poured into a 10 mL volumetric flask and
279 made up to volume with distilled water. A total of 0.4 mL of this solution was added to 10 mL tubes,
280 10 μ L of internal standard (Gallium and Scandium, 10 ppm, BDH, UK) included and 0.2 mL of
281 methanol added. The tube was made up to volume with distilled water and total selenium was
282 measured in a reaction cell by Inductively Coupled Plasma Mass Spectrometry (Thermo Scientific,
283 XSeries2 ICP-MS, USA), using argon and hydrogen as carrier gas.

284

285 2.8.6. *Stable isotope analysis*

286 Triplicate samples of rotifer *B. rotundiformis* enriched with Algamac 3050[®], *A. tonsa* copepod
287 nauplii and 13 dah ABT larvae fed on rotifers and/or copepods were frozen in liquid nitrogen and
288 subsequently lyophilized. Then, samples were ground to powder by pestle and mortar, packed into
289 tin capsules to be analyzed for isotopic relative abundance (δ) of ¹⁵N and ¹³C, carbon (%) and nitrogen

290 (%) . Prior to $\delta^{13}\text{C}$ analysis, samples with high lipid content (C:N ratio > 3.5; see Post et al., 2007)
291 were subjected to total lipid extraction by chloroform/methanol (2:1, by volume) (Varela et al., 2012;
292 2013). The relative abundances of ^{13}C and ^{15}N ($\delta^{13}\text{C}$ and $\delta^{15}\text{N}$, respectively) were measured by a
293 continuous gas flow system using a Thermo Finnigan Elementary Analyzer Flash EA1112 coupled
294 to a Finnigan MAT Delta Plus mass spectrometer. All carbon and nitrogen isotope data are reported
295 in δ notation according to the following equation: $\delta X = [(R_{\text{sample}}/R_{\text{standard}}) - 1] \times 1000$, where
296 X is ^{13}C or ^{15}N and R is the ratio $^{13}\text{C}/^{12}\text{C}$ or $^{15}\text{N}/^{14}\text{N}$ (Peterson and Fry, 1987). Standard materials are
297 Vienna Pee Dee belemnite for carbon and atmospheric N_2 for nitrogen and expressed as parts per
298 thousand (‰) relative to standards (Peterson and Fry, 1987). The isotopic enrichment (Δ) of the
299 predator in relation to its prey was calculated as: $\Delta^{15}\text{N} = \delta^{15}\text{N}_{\text{predator}} - \delta^{15}\text{N}_{\text{prey}}$ and $\Delta^{13}\text{C} = \delta^{13}\text{C}_{\text{predator}} -$
300 $\delta^{13}\text{C}_{\text{prey}}$, respectively.

301

302 2.9. Tissue RNA extraction and cDNA synthesis

303 Samples of pooled larvae (approximately 100 mg) [2 samples per tank in Trial 1 (n = 8) and
304 6 samples per tank in trial 2 (n = 6)] were placed in RNAlater[®] (Sigma–Aldrich, Dorset, UK) and
305 frozen at -20 °C prior to total RNA extraction. Samples were homogenized in 1 mL of TriReagent[®]
306 (Sigma-Aldrich) RNA extraction buffer using a bead tissue disruptor (Bio Spec, Bartlesville,
307 Oklahoma, USA). Total RNA was isolated following manufacturer’s instructions and quantity and
308 quality determined by spectrophotometry using a Nanodrop ND-1000 (Labtech Int., East Sussex,
309 UK) and electrophoresis using 200 ng of total RNA in a 1 % agarose gel. cDNA was synthesized
310 using 2 μg of total RNA and random primers in 20 μL reactions and the high capacity reverse
311 transcription kit without RNase inhibitor according to the manufacturer’s protocol (Applied
312 Biosystems, Warrington, UK).

313

314 2.10. qPCR analysis

315 Several genes related to lipid and fatty acid metabolism, antioxidant and digestive enzymes,
316 as well as growth markers were evaluated in the present study. Quantitative real-time PCR (qPCR)
317 was carried out on transcription factors peroxisome proliferator-activated receptor alpha (*ppara*),
318 peroxisome proliferator-activated receptor gamma (*ppary*), liver X receptor (*lxr*), retinoid X receptor
319 (*rxr*), sterol regulatory element-binding protein 1 (*srebp1*) and sterol regulatory element-binding
320 protein 2 (*srebp2*); LC-PUFA biosynthesis genes delta-6 fatty acyl desaturase (*fads2d6*) and fatty
321 acyl elongase 5 (*elovl5*), and fatty acid metabolism genes fatty acid synthase (*fas*), carnitine palmitoyl
322 transferase I (*cpt1*), acyl coA oxidase (*aco*), fatty acid binding protein 2 (intestinal; *fabp2*), fatty acid
323 binding protein 4 (adipocyte; *fabp4*), fatty acid binding protein 7 (brain; *fabp7*), lipoprotein lipase
324 (*lpl*) and 3-hydroxy-3-methylglutaryl-CoA lyase (*hmgcl*); the antioxidant enzymes superoxide
325 dismutase (*sod*), catalase (*cat*), glutathione peroxidase 1 (*gpx1*) and glutathione peroxidase 4 (*gpx4*);
326 myogenesis growth indicators myosin heavy chain (*myhc*) and tropomyosin (*tropo*); and digestive
327 genes trypsin (*tryp*), amino peptidase (*anpep*), alkaline phosphatase (*alp*), amylase (*amy*), pancreatic
328 lipase (*pl*), phospholipase A2 (*pla2*), bile salt activated lipase 1 (*ball1*) and bile salt activated lipase 2
329 (*bal2*) (see Supplementary Table 2).

330 Expression of genes of interest was determined by qPCR of all the RNA samples with
331 *Elongation factor-1 α* (*elf1 α*) and *β -actin* used as reference genes. The cDNA was diluted 20-fold with
332 milliQ water. The efficiency of the primers for each gene was previously evaluated by serial dilutions
333 of cDNA pooled from the samples to guarantee it was > 85 % for all primer pairs. qPCR was
334 performed using a Biometra TOptical Thermocycler (Analytik Jena, Goettingen, Germany) in 96-
335 well plates in duplicate 20 μ L reaction volumes containing 10 μ L of Luminaris Color HiGreen qPCR
336 Master Mix (Thermo Scientific, Hemel Hempstead, UK), 1 μ L of the primer corresponding to the
337 analyzed gene (10 pmol), 3 μ L of molecular biology grade water and 5 μ L of cDNA (1/20 diluted).
338 In addition, amplifications were carried out with a systematic negative control (NTC, no template
339 control) containing no cDNA. Standard amplification parameters contained an UDG pre-treatment at

340 50 °C for 2 min, an initial denaturation step at 95 °C for 10 min, followed by 35 cycles: 15 s at 95
341 °C, 30 s at the annealing T_m and 30 s at 72 °C.

342

343 *2.11 Statistical analysis*

344 Results for biometry, lipid class and fatty acid compositions are presented as means \pm SD (n
345 = 20 for biometry and n = 3 for survival, lipid class and fatty acid compositions). The data were
346 checked for homogeneity of the variances by the Bartlett test and, where necessary, arc-sin
347 transformed before further statistical analysis. Relations between dietary components and the
348 different variables measured were surveyed by correlation and linear regression analysis.
349 Differences between mean values were analyzed by t-test and one-way ANOVA followed by
350 Tukey's multiple comparisons test performed using GraphPad Prism version 7.00 for Mac OSX
351 (GraphPad Software, La Jolla California USA, www.graphpad.com). Differences were reported as
352 statistically significant when $P < 0.05$ (Zar, 1999). Gene expression results were analyzed using the
353 relative expression software tool (REST 2009), which employs a pairwise fixed reallocation
354 randomization test (10,000 randomizations) with efficiency correction (Pfaffl et al., 2002) to
355 determine the statistical significance of expression ratios (gene expression fold changes) between
356 two treatments.

357

358 **3. Results**

359 *3.1 Analyzed composition of the different dietary treatments (rotifer, Artemia, copepods and gilthead 360 sea bream yolk sac larvae)*

361 Gross composition (% dry mass), energy content (kcal.g⁻¹ dry mass), taurine (mg.g⁻¹ dry
362 mass), vitamin E and selenium contents ($\mu\text{g.g}^{-1}$ dry mass) and lipid class composition (total lipid %)
363 of the different live preys used in the present study are shown in Table 1. Total protein content was
364 significantly highest in *Acartia* copepod (63.2 %) followed by sea bream yolk sac larvae (58.7 %)

365 and then enriched rotifer and *Artemia* (about 52 %). In contrast, total lipid content was highest in sea
366 bream yolk sac larvae and enriched *Artemia* (about 18 %), followed by enriched rotifer (12 %) and
367 copepod (8.8 %). Carbohydrates (generally mostly chitin) was highest in enriched rotifers, followed
368 by *Acartia* and *Artemia* and, as expected, lowest in sea bream yolk sac larvae. Total caloric content
369 was significantly highest in sea bream yolk sac larvae and enriched *Artemia* (about 56 kcal.g⁻¹), due
370 to the highest lipid caloric contribution, followed by enriched rotifers and *Acartia* (about 52 kcal.g⁻¹).
371 The significantly highest total polar lipid contents and polar/neutral lipid ratios were found in
372 *Acartia* nauplii and sea bream yolk sac larvae, mainly due to presenting highest levels of PC and PE.
373 Enriched *Artemia* and rotifer had the significantly lowest polar/neutral lipid ratios, due to their high
374 triacylglycerol (TAG) contents. Free cholesterol was highest sea bream yolk sac larvae >*Acartia*
375 nauplii >enriched *Artemia* metanauplii > rotifers. Vitamin E content was highest in sea bream yolk sac
376 larvae, followed by enriched rotifers, *Acartia* nauplii and lastly enriched *Artemia*. Se content was
377 highest in enriched rotifers followed by enriched *Artemia* and yolk sac sea bream larvae and then
378 *Acartia* copepods. Highest taurine content was found in gilthead sea bream yolk sac larvae (11 mg.g⁻¹
379 dry mass), followed by enriched *Artemia* (6.1 mg.g⁻¹ dry mass), *Acartia* nauplii (4.1 mg.g⁻¹ dry
380 mass) and finally enriched rotifer (2.5 mg.g⁻¹ dry mass).

381

382 3.2 Amino acid contents in live preys and ABT larvae

383 Highest total amino acid content was presented by *Acartia* nauplii (571.5 mg.g⁻¹ dry mass) >
384 gilthead sea bream yolk sac larvae (528.3 mg.g⁻¹ dry mass) > enriched *Artemia* metanauplii (411.6
385 mg.g⁻¹ dry mass) > enriched rotifer (334.3 mg.g⁻¹ dry mass) (Table 1 and Supplementary Table 3).
386 However, the highest total essential amino acid content was shown by sea bream yolk sac larvae
387 (280.4 mg.g⁻¹ dry mass) > *Acartia* nauplii (267.1 mg.g⁻¹ dry mass) > enriched *Artemia* metanauplii
388 (214.6 mg.g⁻¹ dry mass) > enriched rotifer (157.5 mg.g⁻¹ dry mass). The most deficient content of
389 essential amino acids, according to the sufficiency indices, was presented by enriched rotifers (Table
390 2). Enriched *Artemia* showed some insufficiencies with regards to valine, leucine, phenylalanine,

391 threonine and methionine. *Acartia* nauplii showed shortfall of histidine, and methionine, whereas sea
392 bream yolk sac larvae only showed a slight shortage of valine.

393

394 3.3 Total lipid fatty acid compositions and contents of the live preys used to feed ABT larvae in Trials
395 1 and 2

396 Total lipid fatty acid composition (percentage of weight) of rotifer *B. rotundiformis* enriched
397 with Algamac 3050[®], nauplii of the copepod *A. tonsa* fed with the microalgae *R. baltica*, *Artemia*
398 metanauplii enriched with Algamac 3050[®] and gilthead sea bream (*S. aurata* L.) yolk sac larvae used
399 to feed ABT larvae are presented in Table 3. The highest values of total saturated fatty acids were
400 shown by *Acartia* nauplii and gilthead sea bream yolk sac larvae (27.8 % and 26.0 %, respectively),
401 followed by enriched rotifer (23.9 %) and *Artemia* (22.4 %). The highest values for total monoenes
402 were presented by enriched *Artemia* and yolk sac larvae (27.5 % and 26.3 %, respectively), mainly
403 due to the large proportions of 18:1n-9 (14.9 % and 15.9 %, respectively). Copepods and enriched
404 rotifers showed monoene values significantly lower (11.5 % and 12.6 %, respectively). Total n-6
405 PUFA were highest in enriched rotifers, followed by *Acartia* nauplii, enriched *Artemia* and yolk sac
406 larvae, largely due to their contents of linoleic acid (18:2n-6, LA). Yolk-sac larvae showed a very
407 low level of arachidonic acid (20:4n-3, ARA) compared to the other live preys. Total n-3 PUFA were
408 highest in copepod nauplii (43.9 %) > enriched *Artemia* and yolk sac larvae (39.3 % and 37.2 %,
409 respectively) > enriched rotifers (32.9 %). Enriched *Artemia* presented the significantly highest value
410 (23.0 %) of linolenic acid (18:3n-3; LNA) characteristic of EG grade *Artemia*. The proportion of
411 eicosapentaenoic acid (20:5n-3, EPA) was significantly highest (7.2 %) and that of docosahexaenoic
412 acid (22:6n-3, DHA) second highest (25.1 %) in yolk sac larvae, whereas the proportion of
413 docosahexaenoic acid was highest in *Acartia* (27.0 %). The DHA/EPA ratio was highest in *Acartia*
414 (6.1), followed by yolk sac larvae (3.5), enriched rotifers (2.8) and finally enriched *Artemia* (1.9),
415 respectively (Table 3). Quantitative results as total lipid fatty acid content (μg fatty acid.mg dry mass⁻¹)
416 ¹) of the live preys used to feed ABT larvae in Trials 1 and 2 are shown in Supplementary Table 4.

417

418 *3.4 Growth performance and survival rates of ABT larvae in Trial 1 (13 dah) and Trial 2 (18 dah).*

419 Growth performance and survival of ABT larvae in Trial 1 (13 dah) and Trial 2 (18 dah) are
420 presented in Table 4. In Trial 1, 13 dah ABT larvae fed on *Acartia* copepods showed significantly
421 higher values for total length, total weight and flexion index than larvae fed on enriched rotifers.
422 However, no significant differences were observed in survival rates. In Trial 2, 18 dah ABT larvae,
423 previously fed on copepods in Trial 1 and then on sea bream yolk sac larvae (treatment CY), showed
424 the significantly highest values for total length, total weight and survival (32 %), and second lowest
425 for dry mass %. The next best set of values for growth and survival were obtained for ABT larvae fed
426 on treatment CA, followed by those of larvae fed on treatment CC, with poorest performance obtained
427 with ABT larvae fed treatment RA.

428

429 *3.5 Total lipid content and lipid class composition of 13 dah and 18 dah ABT larvae under different*
430 *live prey feeding protocols in Trials 1 and 2.*

431 Total lipid content (% of live and dry mass) and lipid class composition (total lipid %) of ABT
432 larvae 13 dah fed on treatments R and C in Trial 1, and ABT larvae 18 dah that had been fed on
433 treatments RA, CA, CC and CY in Trial 2, are presented in Table 5. In Trial 1, ABT 13 dah larvae
434 fed either on treatment R or C did not show significant differences in total lipid content. Neither
435 showed significant differences in total polar or total neutral lipids. Nevertheless, total polar lipids
436 predominated (56.5 and 58.5 %, respectively) over total neutral lipids (43.5 and 41.5 %, respectively)
437 in ABT larvae fed on treatments R or C, respectively. Moreover, ABT larvae fed on treatment R
438 showed significant higher level of TAG than larvae fed on treatment C (14.5 % vs 10.7 %). In Trial
439 2, ABT 18 dah fed on treatments CY and RA showed significant highest lipid contents followed by
440 those ABT larvae that were fed on treatments CC and CA. Total polar lipid showed highest (59.4 %)
441 in ABT larvae from treatment CC, followed by ABT larvae from treatments CA and RA (56.4 % and

442 54.6 %, respectively) and lowest value for larvae from CY treatment, due to their highest level of
443 total neutral lipid (51.8 %) primarily TAG (22.4 %) (Table 5).

444

445 *3.6 Total lipid fatty acid composition of 13 dah and 18 dah ABT larvae under different live prey*
446 *feeding protocols in Trials 1 and 2.*

447 In Trial 1, 13 dah ABT larvae fed treatments R or C showed no significant differences between
448 total saturated fatty acids, although larvae from treatment R showed significantly higher 16:0 contents
449 than larvae from treatment C (Table 6). Total monoenes were significantly higher in larvae from
450 treatment R, mainly due to higher levels of 16:1n-7 and 18:1n-7 in total lipids compared to larvae fed
451 treatment C. The same trend was observed for total n-6 PUFA, which were significantly higher in
452 ABT larvae fed treatment R, due to higher values of LA, ARA and n-6 docosapentaenoic acid (22:5n-
453 6, DPA) compared to larvae fed treatment C. In contrast, total n-3 PUFA content was highest in ABT
454 larvae fed treatment C (34.2 % vs 19.5 %), mainly due to contain higher levels of LNA, SDA, EPA
455 and DHA. The DHA/EPA ratio was almost 14-fold higher in ABT larvae fed treatment C.

456 In Trial 2, larvae fed treatments CC and CY showed significantly higher total saturated fatty
457 acids, mainly due to higher levels of 16:0, followed by ABT larvae of the RA and CA treatments.
458 Similarly, high levels of total monoenoic fatty acids (about 20 % of total fatty acids, with half
459 represented by 18:1n-9) were found in larvae from treatments RA, CA and CY, with larvae from
460 treatment CC showing lower levels. Total n-6 PUFA were highest in larvae from treatment RA (14.9
461 %) > larvae fed treatments CC and CA (12.5 % and 12.1 %, respectively), and were lowest in larvae
462 fed treatment CY (9.8 %). These values mainly reflected the level of 18:2n-6 that followed the same
463 order per treatment. ARA was significantly highest in larvae fed treatment CA (1.9 %), with no
464 significant differences among the other treatments. Total n-3 PUFA was significantly higher in ABT
465 larvae fed treatment CC (33.2 %) > CA, CY and RA (30.9 %, 30.7 % and 28.2 %, respectively).
466 These values reflected the major contribution of DHA, which was highest in larvae fed treatments

467 CC and CY (24.1 % and 22.2 %, respectively), followed by larvae fed treatments CA and RA (15.8
468 % and 10 %). Levels of EPA in ABT larvae showed, in decreasing order of abundance, treatments
469 RA and CY (4.7 % and 4.4 %, respectively, and not significantly different) followed by treatments
470 CA and CC (3.4 % and 2.1 %, respectively). The DHA/EPA ratio was significantly highest in 18 dah
471 ABT larvae fed treatment CC (11.3), followed by larvae fed treatments CY (5.1), CA (4.6) and RA
472 (2.1).

473

474 *3.7 Relative abundance (δ) of stable isotopes ^{15}N and ^{13}C and C:N ratios of enriched rotifer *B.*
475 *rotundiformis*, *A. tonsa* nauplii and 13 dah ABT larvae*

476 The relative abundance (δ) of ^{15}N and ^{13}C (‰) and C:N ratio of rotifer *B. rotundiformis*
477 enriched with Algamac 3050[®], nauplii of the copepod *A. tonsa* fed on the microalgae *R. baltica* and
478 ABT larvae 13 dah fed on *B. rotundiformis* and *A. tonsa* nauplii, as well as isotopic enrichment (Δ)
479 of ABT larvae in relation to its preys are presented in Table 7. The relative abundance of ^{15}N was
480 significantly higher in *A. tonsa* nauplii than in enriched rotifers, as was the case for 13 dah ABT
481 larvae fed *Acartia* compared to larvae fed enriched rotifer. In contrast, the relative abundance of ^{13}C
482 showed significantly higher values in enriched rotifers than in *Acartia*. The same was observed for
483 ABT larvae, with ABT larvae fed enriched rotifers presenting significantly higher values than larvae
484 fed copepod nauplii. On the other hand, the C:N ratio was significantly higher in enriched rotifers
485 and ABT larvae fed rotifers than in copepod nauplii and ABT larvae fed copepods. The isotopic
486 enrichments $\Delta^{15}\text{N}$ and $\Delta^{13}\text{C}$ of 13 dah ABT larvae in relation to their preys were also significantly
487 higher in larvae fed copepods than in larvae fed rotifers.

488

489 *3.8. Gene expression*

490 *3.8.1. Myogenic genes*

491 The expression of both *myhc* and *tropo* was significantly higher in ABT larvae fed *A. tonsa*
492 nauplii in Trial 1 (Fig. 1). On the other hand, in Trial 2, the expression of *myhc* in 18 dah ABT larvae

493 was significantly up-regulated in larvae from treatment CY whereas no significant differences were
494 observed among the expression of this gene in larvae from the others treatments (RA, CA and CC).
495 With regard to the expression of *tropo* in 18 dah ABT larvae, no differences were observed among
496 larvae fed the different treatments.

497 3.8.2. Fatty acid synthesis genes.

498 No significant nutritional regulation was detected at 13 dah between larvae fed on treatments
499 R and C with regard to the relative expression of the 3 fatty acid synthesis genes *fas*, *fad2d6* and
500 *elovl5* (Fig. 2). The expression values for the same 3 genes followed a similar pattern at 18 dah, with
501 ABT larvae fed CA displaying the highest expression levels, albeit not significantly different to
502 expression in larvae fed RA and CC in the case of *fad2d6*. The lowest transcript copy numbers of *fas*
503 and *elovl5* were found in larvae fed CY, whereas the expression of *fad2d6* did not differ between this
504 group and larvae fed RA or CC.

505

506 3.8.3. Fatty acid catabolism genes

507 In relation to fatty acid catabolism, the expression of *cpt1* did not show dietary regulation at
508 13 dah among larvae fed R or C in Trial 1 (Fig. 3), whereas at 18 dah the highest mRNA copy numbers
509 were observed in larvae fed CY > RA > CA = CC. In contrast, relative expression of *aco* was
510 significantly lower in 13 dah ABT larvae fed treatment C and, in Trial 2, larvae fed CY.

511

512 3.8.4. Fatty acid and lipid transport genes

513 In Trial 1, the relative transcript abundance of *fabp2*, *fabp7* and *lpl* was not significantly
514 different between larvae fed treatments R and C (Fig. 4). In addition, the expression of *fabp4* and
515 *hmgcl* were significantly lower in larvae fed treatment C. In Trial 2, the lowest copy numbers for
516 *fabp2*, *lpl* and *hmgcl* were observed in larvae fed CY, whereas these larvae showed the highest
517 expression levels of *fabp4* and *fabp7*, albeit not different to those of larvae fed CA in the case of
518 *fabp7* (Fig. 4).

519

520 3.8.5. *Transcription factors genes regulating lipid metabolism*

521 In Trial 1, the expression levels of *ppara*, *ppary*, *srebp1* and *srebp2* were significantly lower
522 in larvae fed treatment C (Fig. 5). In contrast, the expression of *rxr* was significantly higher in larvae
523 fed treatment C, whereas no significant differences were observed for expression of *lxr*. In Trial 2,
524 lower expression of *ppara*, *srebp1*, *ppary* and *lxr* was observed in CY-fed larvae. The relative
525 expression value of *srebp2* was also lowest for larvae fed treatment CY, albeit not different to the
526 level in larvae fed CC. No regulation was observed in the expression levels of *rxr* at 18 dah (Fig. 5).

527

528 3.8.6. *Antioxidant defence enzyme genes*

529 In Trial 1, the expressions of *sod*, *gpx1* and *gpx4* were significantly lower in larvae fed *Acartia*
530 nauplii (C), whereas the expression of *cat* was not regulated between larvae fed treatments R or C
531 (Fig. 6). In Trial 2, *sod* expression was significantly higher in ABT larvae fed treatment CY, whereas
532 abundance of *cat* and *gpx4* was lower in the same larvae. The highest *gpx1* mRNA copy number was
533 found in larvae fed RA, albeit not different to that of larvae fed CY (Fig. 6).

534

535 3.8.7. *Digestive enzyme genes*

536 In Trial 1, the expression values of *tryp*, *alp*, and *ball1* showed no regulation between 13 dah
537 ABT larvae fed treatments R or C (Fig.7). In contrast, the expression levels of *anpep*, *amy*, *pl*, *pla2*
538 and *bal2* were significantly lower in ABT larvae fed treatment C. In contrast, in Trial 2, larvae fed
539 CY displayed the lowest expression levels for all the digestive genes, excepting for *alp*, where no
540 differences were found among larvae fed the different dietary treatments. ABT larvae fed treatment
541 CC displayed the highest expression levels of *tryp*, *pla2* *ball1* and *bal2* (Fig. 7).

542

543 **4. Discussion**

544 Previous studies had shown that copepods supported better growth performance than other

545 live prey for larvae of ABT at first feeding (Yufera et al., 2014; Betancor et al., 2017a,b). The present
546 study was consistent with these data, as larvae fed this live prey displayed larger biometric data,
547 advanced stage of development (highest flexion index), and highest expression of the myogenic
548 genes, *myhc* and *tropo* than ABT larvae fed on enriched rotifer at 13 dah. Additionally, in the present
549 study we evaluated the effect that different live prey can have on later stages (18 dah) of ABT larvae
550 performance in combination with the earlier use of either rotifers or copepods. A higher C:N ratio in
551 the rotifer coincides with a significantly higher total lipid content in this prey than in the copepod, as
552 was the case for rotifer-fed 13 dah ABT larvae, whose average total lipid content was higher than
553 ABT larvae fed copepods, albeit not statistically significant. The $\delta^{15}\text{N}$ has been used to examine the
554 trophic position of organisms in food webs (Owens 1987; Hobson and Welch, 1992; Michener and
555 Kaufman, 2007), with ^{15}N values of consumers generally increasing with trophic level, with an
556 average 3.2 ‰ of enrichment per trophic level (Michener and Kaufman, 2007). In the present study,
557 ABT larvae fed *A. tonsa* nauplii showed higher values of enrichment of N ($\Delta^{15}\text{N}$) and C ($\Delta^{13}\text{C}$) than
558 ABT larvae fed on enriched *B. rotundiformis* (2.79 vs. 1.46 for $\Delta^{15}\text{N}$, and 3.06 vs. 0.48 for $\Delta^{13}\text{C}$,
559 respectively). This is also consistent with *Acartia* nauplii having a significantly higher content of total
560 protein (63.2 % vs. 51.3 % on a dry matter basis) and total amino acids (571.5 vs. 334.3 mg.g⁻¹ dry
561 mass) than rotifers. Gilthead sea bream yolk sac larvae, as a live prey showed the highest content of
562 essential amino acids with indication of a slight deficiency only for valine. Thus, the stable isotope
563 data support growth performance data, where ABT larvae fed on copepods grew and developed faster
564 than ABT larvae fed on enriched rotifers as they were prey of a supposedly “higher trophic position”,
565 and had better nutritional value, in consequence fixing more N and C.

566 Fast growth in ABT larvae during early life stages, under standard rearing conditions, occurs
567 after notochord flexion, when piscivory is attained (Uriarte et al., 2016). The results of the present
568 study showed that ABT larvae fed *Acartia* nauplii attained 54.5 % of flexioned larvae *versus* only
569 35.8 % for ABT larvae fed enriched rotifers at 13 dah. This implies an advantage to reach the faster
570 growth phase after flexion with larvae that had been fed copepods attaining higher growth rates than

571 those fed rotifers at 18 dah. Indeed, highest growth was obtained in ABT larvae fed copepods first
572 (to 13 dah) and then on gilthead sea bream yolk sac larvae (to 18 dah), reflecting the contribution of
573 piscivory in the diet. Once ABT larvae attained piscivory, a very high growth performance was
574 obtained, in agreement with previous reports with Pacific Bluefin tuna (PBT; *Thunnus orientalis*)
575 larvae (Tanaka et al., 2007, 2010, 2014, 2015). Moreover, results on PBT larvae suggested that fast-
576 growing larvae at the onset of piscivory could survive in the mass culture tank and were characterized
577 by growth-selective mortality based on direct evidence by comparing feeding (nitrogen stable isotope
578 ratios) and growth between live and dead fish (Tanaka et al., 2017).

579 In the present trial, survival was not different among treatments at 13 dah, since the variability
580 of average survival rate among replicates was relatively large. Such a difference in survival is
581 currently the norm in the tuna hatchery phase at the present time and limited the number of fish
582 available for the second phase of the trial, 13-18 dph, which was consequently carried out in single
583 replicate tanks to provide fish density to promote adequate feeding response. Great differences in
584 survival/growth have been apparent among different cohorts of fish produced since the first eggs were
585 hatched in 2009. These differences appear related to broodstock, as presently eggs are collected from
586 floating cages and therefore there is no accurate control of each broodstock contribution. However,
587 the mortality of ABT larvae at first feeding should not be quantitatively related to the nutritional
588 quality of the live prey. Survival of tuna larvae in artificial rearing has been reported to be relatively
589 low compared to other fish species (Miyashita, 2002). Multiple factors are thought to contribute to
590 mortality of larval tuna in rearing systems during the early larval stages, such as suboptimal physical
591 condition, including increased bacterial loads, surface adhesion and, of course, malnutrition as an
592 added negative effect (Margulies et al., 2016; Honryo et al., 2017). At these early stages, the vitality
593 and strength of the larvae also depends to a great extent on broodstock nutrition and the quality of the
594 eggs and yolk sac larvae produced, and this may vary with and within every spawned batch. Some
595 other neglected zootechnical factors dealing with rearing conditions in individual replicates/tanks
596 may affect swim bladder inflation and larval survival. Besides, mortality and poor development

597 during larval stages has also been attributed to antioxidant status (Mourente et al., 1999; Fernandez-
598 Díaz et al., 2006; Mazurais et al., 2015; Penglase et al., 2015), as discussed below. In any case, the
599 provision of a high-quality nutritional enhancement of planktonic prey during ABT larvae first
600 feeding, a period of high metabolic demand required to support exponential growth, is of paramount
601 importance.

602 Dietary fatty acid composition is known to have a modulatory effect on enzyme activities
603 controlling fatty acid biosynthesis and bioconversion pathways in cultured teleost species (Zheng et
604 al., 2004; Thanuthong et al., 2011; Ayisi et al., 2018). In fish, fatty acids can arise from two sources;
605 either synthesized *de novo* from non-lipid carbon sources, or directly from dietary lipid. Fatty acid
606 synthase (Fas) plays a key role in the process of *de novo* lipogenesis, and the rate of fatty acid *de*
607 *novo* synthesis has been shown to be negatively correlated to the level of dietary lipids (Henderson,
608 1996). In trial 2 in the present study, there was a strong negative correlation between dietary total
609 lipid content and *fas* expression ($r = -0.63$; $P = 0.05$), indicating an inhibition of *fas* expression by
610 increased dietary lipid levels, as stated above. In contrast, in Trial 1, *fas* expression showed no
611 differences in larvae fed on either rotifers or copepods, possibly due to sufficient lipid content in these
612 preys. On the other hand, *fas* is a direct target gene of the transcription factor *srebp1*, and both genes
613 followed a similar pattern of expression ($r = 1.0$; $P = 0.08$). These data were in contrast to the results
614 of our previous studies with ABT larvae (Betancor et al., 2017a,b).

615 As a general consideration, increased expression of *fads2d6* has been observed previously in
616 fish fed low levels of dietary n-3 LC-PUFA, whereas high dietary levels of these fatty acids has been
617 associated with reduced relative transcript abundance (Morais et al., 2012; Betancor et al., 2015). In
618 the present study, in Trial 1, no dietary regulation was detected at 13 dah among ABT larvae fed
619 treatments R or C with regard to the relative expression of the 3 fatty acid synthesis genes *fas*, *fas2d6*
620 and *elovl5*. These results could be interpreted in the following manner: i) both prey (enriched rotifers
621 and copepod nauplii) supplied sufficient lipid to maintain *fas* expression at a baseline level and ii)
622 both prey supplied sufficient n-3 and n-6 PUFA to keep *fads2d6* and *elovl5* expression at a baseline

623 level. This may also mean that prey lipid and essential fatty acid contents in Trial 1 satisfied ABT
624 larvae requirements for these nutrients at first feeding and did not significantly affect fatty acid or
625 LC-PUFA synthesis gene expression. In contrast, in Trial 2, the expression of *fad2d6* in 18 dah ABT
626 larvae was higher in larvae fed enriched *Artemia* for 5 days after feeding on copepod nauplii diet
627 (treatment CA), and those fed treatments RA and CC were also higher than larvae fed CY. This seems
628 logical taking into account that the *Artemia* diet presented the lowest level of total n-3 PUFA and
629 DHA. In contrast, ABT larvae fed treatment CY showed lower expression of *fads2d6*, possibly due
630 to higher levels of DHA and total n-3 PUFA in both copepod nauplii and sea bream yolk sac larvae
631 than other live prey. A similar pattern of expression to dietary treatments was shown by *elov5* in 18
632 dah ABT larvae, which further supports up-regulation of the LC-PUFA pathway in response to the
633 low dietary LC-PUFA in treatment CA. This suggests that n-3 LC-PUFA requirement of 18 dah ABT
634 larvae could be above the level supplied by treatment CA (mainly *Artemia*) and below or similar to
635 that of CY. In agreement, a previous study showed that an increased DHA/EPA ratio in ABT larvae
636 in parallel with increased expression of *fad2d6* and *elov5* during development of unfed ABT larvae
637 was associated with elongation and desaturation of EPA in order to maintain adequate DHA levels
638 (Morais et al., 2011).

639 High dietary levels of n-3 LC-PUFA, particularly DHA, can act as ligands for transcription
640 factors such as *ppara* and *srebp1*, down-regulating the biosynthesis of LC-PUFA (Worgall et al.,
641 1998; Hihi et al., 2002; Cunha et al., 2013; Peng et al., 2014), and regulating the expression of their
642 target genes such as *fas*, *cpt1*, *aco* or *lpl*. In the present study, *srebp1* was strongly positively correlated
643 with the expression of *fas* ($r = 1.0$; $p = 0.08$) and *elov5* ($r = 1.0$; $p = 0.08$) in Trial 2. However, no
644 correlation was observed in Trial 2 between dietary DHA levels and the expression values of *ppara*
645 and *srebp1*. Previous studies in teleosts have described an inhibition in hepatic *fas* expression when
646 fish were fed in a restricted manner (Tian et al., 2013; He et al., 2015; Gong et al., 2017) or with
647 increased dietary lipid (Leng et al., 2012), including our previous results with ABT larvae at first
648 feeding (Betancor et al., 2017b). Although the transcription factor genes *ppara* and *srebp1* did not

649 show nutritional regulation in Trials 1 and 2, *fas*, a direct target of *srebp1*, was strongly regulated by
650 the different dietary treatments in Trial 2 and followed a similar pattern of expression as *srebp1* in
651 response to diet.

652 *Ppar γ* , a transcription factor involved in adipocyte function and differentiation and lipid
653 storage by adipocytes (Nedergaard et al., 2005; Ji et al., 2011; Agawa et al., 2012; Ayisi et al., 2018),
654 showed lower expression in ABT larvae fed copepods in Trial 1 and 2, with its expression negatively
655 related to lipid content. Thus, taking into account that copepods are among the natural live prey of
656 ABT larvae in the wild (Uotani et al., 1990; Catalán et al., 2011; Tilley et al., 2016; Kodama et al.,
657 2017), it is feasible to suggest that the high lipid content of enriched rotifers (treatment R in Trial 1
658 and treatment RA in Trial 2) triggered a response in ABT larvae that modulated lipogenetic/lipolytic
659 mechanisms in order to adapt to an energy-dense ration. In this sense, lower *ppar γ* expression was
660 also observed in copepod-fed ABT larvae in previous trials (Betancor et al., 2017a,b). In mammals,
661 targets directly regulated by *ppar γ* include genes that favour uptake of circulating fatty acids by
662 adipocytes (Schoonjans et al., 1996; Frohnert et al., 1999; Chui et al., 2005) and others that promote
663 recycling rather than export of intracellular fatty acids (Guan et al., 2002; Hibuse et al., 2005). These
664 paradoxical effects on adipocyte biology means that, apart from enhancing fatty acid deposition
665 similar to *ppara*, *ppar γ* can lead to increased fatty acid oxidation (Lehrke and Lazar, 2005). The β -
666 oxidation of fatty acids takes place in both mitochondria and peroxisomes, but mitochondrial β -
667 oxidation is quantitatively more important and can utilize a wider range of fatty acids as substrate
668 (Henderson, 1996). In this respect, *ppar γ* expression was down-regulated by dietary treatments with
669 higher n-3 LC-PUFA (CA, CC and CY) and the same trend was followed by the expression of *aco* in
670 Trial 2. Nevertheless, the *cpt1* expression pattern for larvae fed this treatment was the opposite, being
671 higher in ABT larvae fed yolk sac larvae. CPT-1 is considered to be the main regulatory enzyme in
672 long-chain fatty acid β -oxidation and the higher expression observed in larvae fed yolk sac larvae
673 might be explained by the superior content of LC-PUFA in this live prey.

674 SREBP1 preferentially regulates fatty acid and LC-PUFA synthesis, whereas SREBP2
675 regulates the expression of genes involved in cholesterol synthesis (Jeon and Osborne, 2012;
676 Carmona-Antoñanzas et al., 2014) and is up-regulated in response to reduced cholesterol (Minghetti
677 et al., 2011; Carmona-Antoñanzas et al., 2014). Consistent with this, in the present study, there was
678 an inverse correlation between cholesterol levels in enriched rotifers and copepods and *srebp2*
679 expression in ABT larvae in Trial 1. Moreover, in Trial 2, the expression of *srebp2* was also
680 negatively correlated ($r = -0.95$; $P = 0.17$) to dietary cholesterol levels.

681 LXR is important in controlling intermediary metabolism mediating cross-regulation between
682 sterol and fatty acid metabolism (Ayisi et al., 2018), and its ligands can be antagonized by fatty acids
683 (Cruz-Garcia et al., 2012). Thus, high levels of unsaturated fatty acids activate *lxr*, whereas activated
684 *lxr* induces cholesterol catabolism and *de novo* fatty acid biosynthesis in liver, which has led to the
685 suggestion that *lxrs* are sensors of the balance between cholesterol and fatty acid metabolism (Ayisi
686 et al., 2018). In Trial 2, a weak negative correlation ($r = -0.32$; $P = 0.67$) was observed between dietary
687 cholesterol levels and *lxr* expression. Although there was no significant correlation between dietary
688 cholesterol levels and *lxr* expression in Trial 2, *lxr* expression was positively correlated with the
689 expression of the bile acid activated lipases *ball* and *bal2*, ($r = 0.80$; $p = 0.33$). The lack of a clear
690 relation between dietary cholesterol and *lxr* expression may be due to the fact that LXR is activated
691 by several sterols, including intermediates in the synthesis of cholesterol (Carmona-Antoñanzas et
692 al., 2014) and, although the level of cholesterol differed among treatments, the levels of other sterols
693 that may activate *lxr* might be similar. In addition to the role of *lxr* in regulating cholesterol
694 catabolism, storage, absorption and transport through the transcriptional regulation of key target
695 genes involved in these processes, it may also act in fatty acid metabolism by increasing the
696 expression of the transcription factor *srebp1* or genes such as *fas* or *lpl*. In this sense, the mRNA
697 levels of *lxr* and *lpl* showed a strong positive correlation ($r = 1.0$; $p = 0.08$) and the same pattern of
698 expression in 18 dah ABT larvae. Lpl is a lipase highly expressed in muscle and liver of ABT
699 (Betancor et al., 2017a) that hydrolyzes TAG in plasma lipoproteins and supplies free fatty acids for

700 deposition in adipose tissue or for oxidation in other tissues (Nilsson-Ehle et al., 1980; Kersten, 2014;
701 Ayisi et al., 2018). In agreement, high levels of *lpl* expression and activity have been associated with
702 increased lipid utilization in darkbarbel catfish (*Pelteobagrus vachelli*) larvae fed high-lipid diets
703 (Zheng et al., 2010).

704 Fatty acid binding proteins (Fabp) in general are noted for the intracellular transport of fatty
705 acids and play an intermediary role in orchestrating gene transcription involved in lipid homeostasis
706 (Tocher 2003; Ayisi et al., 2018). In agreement with our previous studies (Betancor et al., 2017a,b),
707 *fabp4*, a carrier protein involved in fatty acid uptake, transport and metabolism in adipocytes (Glatz
708 and van der Vusse, 1996), was down-regulated in copepod-fed ABT larvae in Trial 1, possibly
709 reflecting increased uptake and accumulation of lipid into larval tissues. On the other hand, *fabp2*
710 (intestinal) and *fabp7* (brain) showed no differences in their expression between 13 dah ABT larvae
711 fed treatments R or C. In agreement, a recent study in Senegalese sole (*Solea senegalensis*) larvae
712 showed no regulation of *fabp2* expression when larvae were fed enriched *Artemia*, whereas up-
713 regulation of *fabp1* and *fabp3* was observed in larvae fed high levels of n-3 LC-PUFA (Bonacic et
714 al., 2016), which may indicate differential regulation of *fabp* at different developmental stages (André
715 et al., 2000). In Trial 2 in the present study, different patterns of regulation were presented by different
716 *fabp* genes in response to dietary treatments. However, the positive correlation ($r = 0.80$; $p = 0.33$)
717 between *fas* expression and *fabp2* (intestinal) expression in ABT larvae fed the different dietary
718 treatments in Trial 2 was noteworthy.

719 Rotifers and *Artemia* have low levels of several nutrients, including antioxidants, compared
720 to copepods (Hamre et al., 2013). Nutrient intake plays a role in regulating the redox system in fish
721 larvae, suggesting that nutrient-induced changes in the redox system may contribute to differences in
722 larval fish growth and development (Mourente et al., 2007; Izquierdo and Betancor, 2015; Penglase
723 et al., 2015). In Trial 1, C-fed larvae showed lower expression of *sod*, *gpx1* and *gpx4*, possibly in
724 response to lower contents of antioxidant nutrients such as Se or vitamin E. It was more complicated
725 in Trial 2 as the pattern of expression was variable among the different dietary treatments. Differences

726 could not only be attributed to the feed, but also to the stage of development of the larvae as it is
727 known that lipid peroxidation levels in larvae show diet and age/growth dependence in their responses
728 (Fernandez-Diaz et al., 2006). Therefore, the differences in growth among the larvae could influence
729 lipid oxidation levels. It is perhaps noteworthy that the Se content of live prey, either enriched rotifers
730 or *Acartia* nauplii, showed a significant negative correlation with final total length ($r = -0.9$; $P =$
731 0.0417), total weight ($r = -0.9$; $P = 0.008$) and flexion index ($r = -0.8$; $P = 0.053$). Although Se is an
732 essential micronutrient, it has the narrowest window of any element between requirement and toxicity
733 (Polatajko et al., 2006), with reduced growth being one of the first symptoms of toxicity (Jaramillo et
734 al., 2009). Thus, it would need to be established whether the Se levels used in enriched rotifers in the
735 present study were within safe limits for ABT larvae.

736 Molecular methodologies could contribute to the understanding of the real digestive capacities
737 of developing larvae under different dietary protocols. In the present study, expression of trypsin
738 (*tryp*), pancreatic alkaline protease (*alp*) and bile acid activated lipase 1 (*ball*) was not regulated by
739 dietary treatments in Trial 1. In contrast, in Trial 2, all digestive enzyme genes studied were
740 significantly up-regulated by treatment CC and down-regulated by treatment CY, which were the
741 largest larvae, which could indeed indicate that the expression levels are related to the developmental
742 stage. It is commonly accepted that the major digestive lipase in teleosts, including larvae, appears to
743 be bile salt-dependent lipases (*bal*) (Rønnestad and Morais, 2008). It was also reported that *Bal* were
744 the main enzymes involved in lipid digestion in the larval stage of PBT (Murashita et al., 2014). In
745 the present study, the expression patterns of both isoforms (*ball* and *bal2*) were nutritionally
746 regulated, showing a similar pattern.

747 In conclusion, and in agreement with our previous trials, the present study showed that
748 copepods (*Acartia*) were a superior live prey for first feeding ABT larvae compared to enriched
749 rotifers, as indicated by higher growth performance. This rapid growth would enable rapid attainment
750 of the piscivory feeding stage, which would also enable earlier weaning to inert formulated diet. This
751 may reflect the higher protein and essential amino acids, polar lipid and n-3 LC PUFA and other

752 nutrient contents of copepods as first prey. When the start of piscivory is delayed (larvae fed Artemia
753 or copepods) larvae previously fed copepods do not show any advantage in growth compared to larvae
754 fed with rotifers. The highest DHA contents were found in ABT larvae fed CC and CY, whereas the
755 lowest contents were found in RA-fed larvae. Although, RA-fed larvae showed the highest level of
756 the intermediate product n-3 DPA, this was not supported by gene expression data. Different
757 expression patterns of digestive enzymes between ABT larvae fed copepods and enriched rotifers
758 could be due to different lipid class/fatty acid compositions of the live prey or to differences in the
759 size/development of the larvae.

760

761 **Acknowledgements**

762 We wish to thank the technical staff at Laboratory of Marine Aquaculture (IEO), Puerto de Mazarrón
763 (Murcia, Spain) and Nutritional Analytical Services (NAS; Institute of Aquaculture, University of
764 Stirling, UK) that contributed to this work. This work was supported by the Consejería de Innovación,
765 Ciencia y Empresa de la Junta de Andalucía, Proyecto de Excelencia de Promoción General del
766 Conocimiento [Ref. RNM 733, 2012), and Programa Estatal de Investigación del Ministerio de
767 Economía y Competitividad [Ref. AGL2014-52003-C2-1-R, 2014 and Ref. RTC-2016-5835-2].

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769 **References**

- 770 Ackman, R.G., 1980. Fish lipids. In: Connell, J. J. (Ed.), *Advances in Fish Science and Technology:*
771 *Fishing News Books, Farnham, pp. 83-103.*
- 772 Agawa, Y., Honryo, T., Ishii, A., Kobayashi, T., Oku, H., Sawada, Y., 2012. Molecular identification
773 and tissue distribution of peroxisome proliferators activated receptor gamma transcript in cultured
774 *Thunnus orientalis*. *Aquacult. Res.* 43, 1145-1158.

- 775 André, M., Ando, S., Ballagny, C., Durliat, M., Poupard, G., Briançon, C., Babin, P.J., 2000.
776 Intestinal fatty acid binding protein gene expression reveals the cephalocaudal patterning during
777 zebrafish gut morphogenesis. *Int. J. Dev. Biol.* 44, 249–252.
- 778 AOAC (2000) Official methods of analysis. Association of Official Analytical Chemists,
779 Washington, DC.
- 780 Ayisi, C. L., Yamei, C., Zhao, J-L., 2018. Genes, transcription factors and enzymes involved in lipid
781 metabolism in fin fish. *Agri Gene* 7, 7–14.
- 782 Betancor, M.B., Caballero, M.J., Terova, G., Saleh, R., Atalah, E., Benítez-Santana, T., Bell, J.G.,
783 Izquierdo, M., 2012. Selenium inclusion decreases oxidative stress indicators and muscle injuries
784 in sea bass larvae fed high-DHA microdiets. *Br. J. Nutr.* 13, 1–14.
- 785 Betancor, M.B., Sprague, M., Sayanova, O., Usher, S., Campbell, P.J., Napier, J.A., Caballero, M.J.,
786 Tocher, D.R., 2015. Evaluation of a high-EPA oil from transgenic *Camelina sativa* in feeds for
787 Atlantic salmon (*Salmo salar* L.): Effects on tissue fatty acid composition, histology and gene
788 expression. *Aquaculture* 444, 1-12.
- 789 Betancor, M.B., Ortega, A., de la Gándara, F., Tocher, D.R., Mourente, G., 2017a. Lipid metabolism-
790 related gene expression pattern of Atlantic bluefin tuna (*Thunnus thynnus* L.) larvae fed on live
791 prey. *Fish Physiol. Biochem.* 43, 493-516.
- 792 Betancor, M., Ortega, A., de la Gándara, F., Tocher, D. R., Mourente, G., 2017b. Molecular aspects
793 of lipid metabolism, digestibility and antioxidant status of Atlantic bluefin tuna (*T. thynnus* L.)
794 larvae during first feeding. *Aquaculture* 479, 357-369.
- 795 Bonacic, K., Campoverde, C., Sastre, M., Hachero-Cruzado, I., Ponce, M., Manchado, M., Estevez,
796 A., Gisbert, E., Morais, S., 2016. Mechanisms of lipid metabolism and transport underlying

797 superior performance of Senegalese sole (*Solea senegalensis*, Kaup 1858) larvae fed diets
798 containing n-3 polyunsaturated fatty acids. *Aquaculture* 450, 383-396.

799 Carmona-Antoñanzas, G., Tocher, D.R., Martinez-Rubio, L., Leaver, M.J., 2014. Conservation of
800 lipid metabolic gene transcriptional regulatory networks in fish and mammals. *Gene* 534, 1-9.

801 Catalán, I.A., Tejedor, A., Alemany, F., Reglero, P., 2011. Trophic ecology of Atlantic bluefin tuna
802 *Thunnus thynnus* larvae. *Journal of Fish Biology*, 78(5), 1545-1560.

803 Christie, W.W., 1993, Preparation of derivatives of fatty acids for chromatographic analysis. In:
804 Christie, W.W. (Ed.), *Advances in Lipid Methodology-Two*. The Oily Press, Dundee. pp. 69-111.

805 Chui, P.C., Guan, H.P., Lehrke, M., Lazar, M.A., 2005. PPAR gamma regulates adipocyte cholesterol
806 metabolism via oxidized LDL receptor 1. *J. Clin. Invest.* 115, 2244-2256.

807 Cowey, C.B., Adron, J.W., Walton, M.J., Murray, J., Youngson, A., Knox, D., 1981. Tissue
808 distribution, uptake and requirement for a-tocopherol of rainbow trout (*Salmo gairdneri*) fed diets
809 with a minimal content of unsaturated fatty acids. *J. Nutr.* 111, 1556–1567.

810 Cruz-Garcia, L., Sánchez-Gurmaches, J., Gutiérrez, J., Navarro, I., 2012. Role of LXR in trout
811 adipocytes: target genes, hormonal regulation, adipocyte differentiation and relation to lipolysis.
812 *Comparative Biochemistry and Physiology, Part A* 163, 120–126.

813 Cunha, I., Galante-Oliveira, S., Rocha, E., Planas, M., Urbatzka, R., Castro, L.F.C., 2013. Dynamics
814 of PPARs, fatty acid metabolism genes and lipid classes in eggs and early larvae of a teleost. *Comp.*
815 *Biochem. Physiol.* 164B, 247-258.

816 De La Gandara F., Mylonas C., Coves D., Ortega A., Bridges C.R., Belmonte R.A., Vassallo-Agius
817 R., Papandroulakis N., Rosenfeld H., Tandler A., Medina A., Demetrio A., Corriero A., Fauvel C.,
818 Falcon J., Sveinsvoll K., Ghysen A., Deguara S., Gordin H., 2010. Seedling production of Atlantic
819 bluefin tuna (ABFT) *Thunnus thynnus*. The selfdott project. In: S Miyashita, S., Sakamoto, W.,
820 Biswas, A. (Eds.), *Joint International Symposium of Kinki University and Setouchi Town on the*
821 *40th Anniversary of Pacific Bluefin Tuna Aquaculture, Towards the Sustainable Aquaculture of*
822 *Bluefin Tuna*. University Press, Amami, Japan, pp. 45–52.

- 823 De La Gándara F., Ortega, A., Buentello, A., 2016. Tuna aquaculture in Europe. In: D. D. Benetti,
824 D.D., Partridge, G.J., Buentello, A. (Eds.), *Advances in Tuna Aquaculture: From Hatchery to*
825 *Market*. Elsevier AP, New York, pp. 273-321.
- 826 Fernandez-Diaz, C., Kopecka, I., Canavate, J.P., Sarasquete, C., Sole, M., 2006. Variations on
827 development and stress defences in *Solea senegalensis* larvae fed on live and microencapsulated
828 diets. *Aquaculture* 251, 573–584
- 829 Folch, J., Lees, M., Sloane-Stanley, G.H., 1957. A simple method for the isolation and purification
830 of total lipids from animal tissues. *J. Biol. Chem.* 226, 497-509.
- 831 Frohnert, B.I., Hui, T.Y., Bernlohr, D.A., 1999. Identification of a functional peroxisome proliferator-
832 responsive element in the murine fatty acid transport protein gene. *J. Biol. Chem.* 274, 3970-3977.
- 833 Glatz, J.F., van der Vusse, G.J., 1996. Cellular fatty acid-binding proteins: their function and
834 physiological significance. *Prog. Lipid Res.* 35, 243-282.
- 835 Gong, Y., Chen, W., Han, D., Zhu, X., Yang, Y., Jin, J., Liu, H., Xie, S., 2017. Effects of food
836 restriction on growth, body composition and gene expression related in regulation of lipid
837 metabolism and food intake in grass carp. *Aquaculture* 469, 28-35.
- 838 Guan, H.P., Li, Y., Jensen, M.V., Newgard, C.B., Stepan, C.M., Lazar, M.A., 2002. A futile
839 metabolic cycle activated in adipocytes by antidiabetic agents. *Nat. Med.* 8, 1122-1128.
- 840 Hamre, K., Yúfera, M., Rønnestad, I., Boglione, C., Conceição, L.E.C., Izquierdo, M., 2013. Fish
841 larval nutrition and feed formulation: Knowledge gaps and bottlenecks for advances in larval
842 rearing. *Rev. Aquac.* 5, S26-S58.
- 843 He, A.Y., Ning, L.J., Chen, L.Q., Chen, Y.L., Xing, Q., Li, J.M., Qiao, F., Li, D.L., Zhang, M.L., Du,
844 Z.Y., 2015. Systemic adaptation of lipid metabolism in response to low- and high-fat diet in Nile
845 tilapia (*Oreochromis niloticus*). *Physiol. Rep.* 3, e12485.

- 846 Henderson, R.J., 1996. Fatty acid metabolism in freshwater fish with particular reference to
847 polyunsaturated fatty acids. *Archives of Animal Nutrition/Archiv fur Tierernahrung* 49(1), 5-
848 22.
- 849 Henderson, R.J., Tocher, D.R., 1992. Thin layer chromatography. In: Hamilton, R.J., Hamilton, S.,
850 (Eds.), *Lipid analysis: a practical approach*: IRL Press, Oxford, pp. 65–111.
- 851 Henken, A.M., Lucas, H., Tijssen, P.A.T. and Machiels, M.A.M., 1986. A comparison between
852 methods used to determine the energy content of feed, fish and faeces samples. *Aquaculture* 5,
853 195-201.
- 854 Hibuse, T., Maeda, N., Funahashi, T., Yamamoto, K., Nagasawa, A., Mizunoya, W., Kishida, K.,
855 Inoue, K., Kuriyama, H., Nakamura, T., Fushiki, T., Kihara, S., Shimomura, I., 2005. Aquaporin
856 7 deficiency is associated with development of obesity through activation of adipose glycerol
857 kinase. *Proc. Natl. Acad. Sci. U.S.A.* 102, 10993-10998.
- 858 Hihi, A.K., Michalik, L., Wahli, W., 2002. PPARs: transcriptional effectors of fatty acids and their
859 derivatives. *Cell. Mol. Life Sci.* 59, 790-798.
- 860 Hobson, K.A., Welch, H.E., 1992. Determination of trophic relationships within a high Arctic marine
861 food web using $d^{13}C$ and $d^{15}N$ analysis. *Marine Ecology Progress Series* 84, 9–18.
- 862 Honryo, T., Kurata, M., Guillen, A., Tamura, Y., Cano, A., Stein, M., Margulies, D., V. Scholey, V.,
863 Sawada, Y., 2017. Optimal period for the effective promotion of initial swim bladder inflation
864 in yellowfin tuna, *Thunnus albacares* (Temminck and Schlegel), larvae. *Aquaculture Research*
865 48, 5443–5446.
- 866 Izquierdo, M.S., Betancor, M., 2015. Vitamin E. In: Lee, C.S., Lim, C., Gatlin, D., Webster, C.D.
867 (Eds.), *Dietary Nutrients, Additives and Fish Health*. Wiley-Blackwell, New Jersey, pp. 173-194.
- 868 Jaramillo, F., Peng, L., Gatlin, D., 2009. Selenium nutrition of hybrid striped bass (*Morone chrysops*
869 x *M. saxatilis*) bioavailability, toxicity and interaction with vitamin E. *Aquacult. Nutr.* 15, 160-
870 165.
- 871 Jeon, T.I., Osborne, T.F., 2012. SREBPs: metabolic integrators in physiology and metabolism.

872 Trends Endocrinol. Metab. 23, 65–72.

873 Ji, H., Liu, Y., Zhao, X., Zhang, M., 2011. N-acetyl-L-cysteine enhances the osteogenic
874 differentiations and inhibits the adipogenic differentiation through up regulation of Wnt 5a and
875 down regulation of PPAR γ in bone marrow stromal cells. Biomed. Pharmacother. 65, 369-374.

876 Kersten, S., 2014. Physiological regulation of lipoprotein lipase. BBA- Mol. Cell Biol. Lipids. 1841,
877 919–933.

878 Kodama, T., Hirai, J., Tamura, S., Takahashi, T., Tanaka, Y., Ishihara, T., Tawa, A., Morimoto, H.,
879 Ohshimo, S., 2017. Diet composition and feeding habits of larval Pacific bluefin tuna *Thunnus*
880 *orientalis* in the Sea of Japan: Integrated morphological and metagenetic analysis. Mar. Ecol. Prog.
881 Ser. 583, 211–226.

882 Lehrke, M., Lazar, M.A., 2005. The many faces of PPAR γ . Cell 123, 993-999.

883 Leng, X.J., Wu, X.F., Tian, J., Li, X.Q., Guan, K., Weng, D.C., 2012. Molecular cloning of fatty acid
884 synthase from grass carp (*Ctenopharyngodon idella*) and the regulation of its expression by dietary
885 fat level. Aquacult. Nutr. 18, 551-558.

886 Margulies, D., Scholey, V.P., Wexler, J.B., Stein, M.S., 2016. Research on the reproductive biology
887 and early life history of yellowfin tuna *Thunnus albacares* in Panama. Pages 77–114 in D. D.
888 Benetti, G. J. Partridge, and A. Buentello, editors. Advances in tuna aquaculture from hatchery
889 to market. Academic Press, Waltham, Massachusetts, USA.

890 Mazurais, D., Covès, D., Papandroulakis, N., Ortega, A., Desbruyeres, E., Huelvan, C., Le Gall,
891 M.M., De la Gándara, F., Cahu, C.L., 2015. Gene expression pattern of digestive and antioxidant
892 enzymes during the larval development of reared Atlantic bluefin tuna (ABFT), *Thunnus thynnus*
893 L. Aquacult. Res., 46, 2323-2331.

894 McMurray, C.H., Blanchflower, W.J., Rice, D.A., 1980. Influence of extraction techniques on
895 determination of α -tocopherol in animal feedstuffs. J AOAC 63, 1258–1261.

896 Michener, R. H. and Kaufman, L., 2007. Stable isotope ratios as tracers in marine food webs: an

897 update, *Stable Isotopes in Ecology and Environmental Science*, 2, 238–282.

898 Minghetti, M., Leaver, M.J., Tocher, D.R., 2011. Transcriptional control mechanisms of genes of
899 lipid and fatty acid metabolism in the Atlantic salmon (*Salmo salar* L.) established cell line, SHK-
900 1. *BBA- Mol. Cell Biol. Lipids*. 1811, 194-202.

901 Miyashita, S., 2002. Studies on the seedling production of the Pacific bluefin tuna, *Thunnus thynnus*
902 *orientalis*. *Bulletin of the Fisheries Laboratory of Kinki University*, 8:1–171.

903 Morais, S., Mourente, G., Ortega, A., Tocher, J.A., Tocher, D.R., 2011. Expression of fatty acyl
904 desaturase and elongase genes, and evolution of DHA/EPA ratio during development of unfed
905 larvae of Atlantic bluefin tuna (*Thunnus thynnus* L.). *Aquaculture* 313, 129-139.

906 Morais, S., Castanheira, F., Martinez-Rubio, L., Conceição, L.E.C., Tocher, D.R., 2012. Long chain
907 polyunsaturated fatty acid synthesis in a marine vertebrate: Ontogenetic and nutritional regulation
908 of a fatty acyl desaturase with $\Delta 4$ activity. *Biochim. Biophys. Acta - Mol. Cell Biol. Lipids* 1821,
909 660–671.

910 Mourente, G., Tocher, D.R., Díaz, E., Grau, A., Pastor, E., 1999. Study of the (n-3) HUFA
911 requirement and antioxidant status of *Dentex dentex* larvae at the *Artemia* feeding stage.
912 *Aquaculture*, 179, 291-307.

913 Mourente, G., Bell, J.G., Tocher, D.R., 2007. Does dietary tocopherol affect fatty acid metabolism in
914 fish? *Fish Physiol. Biochem.* 33, 269-280.

915 Murashita, K., Matsunari, H., Kumon, K., Tanaka, Y., Shiozawa, S., Furuita, H., Oku, H., Yamamoto,
916 T., 2014. Characterization and ontogenetic development of digestive enzymes in Pacific bluefin
917 tuna *Thunnus orientalis* larvae. *Fish Physiol. Biochem.* 40, 1741–1755.

918 Nedergaard, J., Ricquier, D., Kozak, L.P., 2005. Uncoupling proteins: current status and therapeutic
919 prospects. *EMBO Rep.*, 6, 917–921.

- 920 Nilsson-Ehle, P., Grafinkel, A.S., Schotz, M.C., 1980. Lipolytic enzymes and plasma lipoprotein
921 metabolism. *Annu. Rev. Biochem.* 49, 667-693.
- 922 Ortega, A., 2015. Cultivo Integral de dos especies de escómbridos: Atún rojo del Atlántico (*Thunnus*
923 *thynnus*, L. 1758) y Bonito Atlántico (*Sarda sarda*, Bloch 1793). PhD Thesis, Universidad de
924 Murcia, Murcia (Spain).
- 925 Oser, B.L. (1959) An Integrated Essential Amino Acid Index for Predicting the Biological Value of
926 Proteins. In: Albanese, A.A., Ed., *Amino Acid Nutrition*, Academic Press, New York, 295-311.
- 927 Owens, N.J.P., 1987. Natural variation in ^{15}N in the marine environment. *Advances in Marine*
928 *Biology* 24, 389–451.
- 929 Penglase, S., Edvardsen, R. B., Furmanek, T., Rønnestad, I., Karlsen, Ø. Van der Meeren T., Hamre,
930 K., 2015. Diet affects the red-ox system in developing Atlantic cod (*Gadus morhua*) larvae. *Redox*
931 *Biology*, 5, 308–318.
- 932 Peng, M., Xu, W., Mai, K., Zhou, H., Zhang, Y., Liufu, Z., Zhang, K., Ai, Q., 2014. Growth
933 performance, lipid deposition and hepatic lipid metabolism related gene expression in juvenile
934 turbot (*Scophthalmus maximus* L.) fed diets with various fish oil substitution levels by soybean
935 oil. *Aquaculture* 433, 442-449.
- 936 Peterson, B.J., Fry, B., 1987. Stable isotopes in ecosystem studies. *Annu. Rev. Ecol. Syst.* 18, 293–
937 320.
- 938 Pfaffl, M.W., Morgan, G.W., Dempfle, L., 2002. Relative expression software tool (REST) for group-
939 wise comparison and statistical analysis of relative expression results in real-time PCR. *Nucleic*
940 *Acids Res.* 30, e36.
- 941 Polatajko, A., Jakubowski, N., Szpunar, J., 2006. State of the art report of selenium speciation in
942 biological samples. *J. Anal At. Spectrom.* 21, 639-654.
- 943 Post, D.M., Layman, C.A., Arrington, D.A., Takimoto, G., Quattrochi, J., Montaña, C.G., 2007.
944 Getting to the fat of the matter: models, methods and assumptions for dealing with lipids in stable
945 isotope analyses. *Oecologia* 152, 179–189.

- 946 Rønnestad, I., Morais, S., 2008. Digestion. In: Finn, R.N., Kapoor, B.G. (Eds.), Fish Larval
947 Physiology, Science Publishers, Enfield, NH, USA, pp. 201-262.
- 948 Schoonjans, K., Peinado-Onsurbe, J., Lefebvre, A.M., Heyman, R.A., Briggs, M., Deeb, S., Staels,
949 B., Auwerx, J., 1996. PPAR alpha and PPAR gamma activators direct a distinct tissue-specific
950 transcriptional response via a PPRE in the lipoprotein lipase gene. EMBO J. 15, 5336-5348.
- 951 Tanaka, Y., Gwak, W.S., Tanaka, M., Sawada, Y., Okada, T., Miyashita, S., Kumai, H., 2007.
952 Ontogenetic changes in RNA, DNA and protein contents of laboratory reared Pacific bluefin
953 tuna *Thunnus orientalis*. Fish. Sci., 73, 378–384.
- 954 Tanaka, Y.T., Inami, H.M., Shihi, Y.I., Umon, K.K., Ba, T.E., Ishi, A.N., Ikaido, H.N., Hiozawa,
955 S.S., 2010. Prey utilization by hatchery-reared Pacific bluefin tuna larvae in mass culture tank
956 estimated using stable isotope analysis, with special reference to their growth variation.
957 Aquaculture Sci., 58, 501–508.
- 958 Tanaka, Y., Minami, H., Ishihi, Y., Kumon, K., Higuchi, K., Eba, T., Nishi, A., Nikaido, H.,
959 Shiozawa, S., 2014. Relationship between prey utilization and growth variation in hatchery-
960 reared Pacific bluefin tuna, *Thunnus orientalis* (Temminck et Schlegel), larvae estimated using
961 nitrogen stable isotope analysis. Aquac. Res. 45, 537–545.
- 962 Tanaka, Y., Kumon, K., Higuchi, K., Eba, T., Nishi, A., Nikaido, H., & Shiozawa, S. (2015).
963 Influence of the prey items switched from rotifers to yolk-sac larvae on growth of laboratory-
964 reared Pacific bluefin tuna. Aquaculture Science, 63, 445–457.
- 965 Tanaka, Y., Kumon, K., Ishihi, Y., Eba, T., Nishi, A., Nikaido, H., Shiozawa, S., 2017. Mortality
966 processes of hatchery-reared Pacific bluefin tuna *Thunnus orientalis* (Temminck et Schlegel)
967 larvae in relation to their piscivory. Aquac. Res. 49, 11–18.
- 968 Thanuthong, T., Francis, D.S., Manickam, E., Senadheera, S.D., Cameron-Smith, D., Turchini, G.M.,
969 2011. Fish oil replacement in rainbow trout diets and total dietary PUFA content: II effects on

- 970 fatty acid metabolism and in vivo fatty acid bioconversion. *Aquaculture* 322–323, 99–108.
- 971 Tian, J., Wen, H., Zeng, L.B., Jiang, M., Wu, F., Liu, W., Yang, G.C., 2013. Changes in the activities
972 and mRNA expression levels of lipoprotein lipase (LPL), hormone-sensitive lipase (HSL) and
973 fatty acid synthetase (FAS) of Nile tilapia (*Oreochromis niloticus*) during fasting and re-feeding.
974 *Aquaculture* 400-401, 29-35.
- 975 Tilley, J.D., Butler, C.M., Suárez-Morales, E., Franks, J.S., Hoffmayer, E.R., Gibson, D.P., Comyns,
976 B.H., Ingram, G.W., Jr., Blake, E.M., 2016. Feeding ecology of larval Atlantic bluefin tuna,
977 *Thunnus thynnus*, from the central Gulf of Mexico. *Bulletin of Marine Science*, 92(3), 321-334.
- 978 Tocher, D.R., 2003. Metabolism and functions of lipids and fatty acids in teleost fish. *Rev. Fisheries*
979 *Sci.* 11, 107-184.
- 980 Tocher, D.R., Harvie, D.G., 1988. Fatty acid composition of the major phosphoglycerides from fish
981 neutral tissues: (n-3) and (n-6) polyunsaturated fatty acids in rainbow trout (*Salmo gairdneri* L.)
982 and cod (*Gadus morhua* L.) brains and retinas. *Fish Physiol. Biochem.* 5, 229-239.
- 983 Uotani, I., Saito, T., Hiranuma, K., Nishikawa, Y., 1990. Feeding habit of bluefin tuna *Thunnus*
984 *thynnus* larvae in the western North Pacific Ocean. *Nippon Suisan Gakkaishi* 56, 713-717.
- 985 Uriarte A., García A., Ortega A., de la Gándara F., Quintanilla J., Laiz-Carrión R., 2016. Isotopic
986 discrimination factors and nitrogen turnover rates in reared Atlantic bluefin tuna larvae (*Thunnus*
987 *thynnus*): effects of maternal transmission. *Sci. Mar.* 80(4), 447-456.
- 988 Van Beijnen (2017). The Closed Cycle Aquaculture of Atlantic Bluefin Tuna in Europe: current
989 status, market perceptions and future potential. 95p.
- 990 Varela, J. L., de la Gándara, F., Ortega, A., Medina, A., 2012. ¹³C and ¹⁵N analysis in muscle and liver
991 of wild and reared young-of-the-year (YOY) Atlantic bluefin tuna. *Aquaculture* 354–355, 17–21.
- 992 Varela, J.L., Rodríguez-Marín, E., Medina, A., 2013. Estimating diets of pre-spawning Atlantic
993 bluefin tuna from stomach content and stable isotope analyses. *J. Sea Res.* 76, 187–192.

994 Worgall, T.S., Sturley, S.L., Seo, T., Osborne, T.F., Deckelbaum, R.J., 1998. Polyunsaturated fatty
995 acids decrease expression of promoters with sterol regulatory elements by decreasing levels of
996 mature sterol regulatory element-binding protein. *J. Biol. Chem.* 273, 25537–25540.

997 Yufera, M., Ortiz-Delgado, J., Hoffman, T., Sigüero, I., Urup, B., Sarasquete, C., 2014.
998 Organogenesis of digestive system, visual system and other structures in Atlantic bluefin tuna
999 (*Thunnus thynnus*) larvae reared with copepods in mesocosm system. *Aquaculture* 426, 126-137.

1000 Zar, J.H., 1999. *Biostatistical Analysis* 4th Edition Prentice-Hall, New Jersey.

1001 Zhao, C., Dahlman-Wright, K., 2010. Liver X receptor in cholesterol metabolism (Review). *Journal*
1002 *of Endocrinology*, 204, 233–240.

1003 Zheng, X., Seiliez, I., Hastings, N., Tocher, D.R., Panserat, S.C., Dickson, A., Bergot, P., Teale, A.J.,
1004 2004. Characterization and comparison of fatty acyl D6 desaturase cDNAs from freshwater and
1005 marine teleost fish species. *Comp. Biochem. Physiol. B* 139, 269–279.

1006 Zheng, K., Zhu, X., Han, D., Yang, Y., Lei, W., Xie, S., 2010. Effects of dietary lipid levels on
1007 growth, survival and lipid metabolism during early ontogeny of *Pelteobagrus vachelli* larvae.
1008 *Aquaculture* 299, 121-127.

1009

1010 **Figure legends**

1011 **Fig. 1.** Nutritional regulation of myosin heavy chain 2 (*myhc*) and tropomyosin (*tropo*) gene
1012 transcription in Atlantic bluefin tuna (ABT) larvae 13 days after hatch (Trial 1) fed with rotifer
1013 *B. rotundiformis* enriched with Algamac 3050[®] (R), *A. tonsa* copepod nauplii (C), and ABT
1014 larvae 18 days after hatch (Trial 2) fed with rotifer *B. rotundiformis* and then *Artemia* metanauplii
1015 enriched with Algamac 3050[®] (RA), *A. tonsa* copepod nauplii and then *Artemia* enriched with
1016 Algamac 3050[®] (CA), *A. tonsa* copepod nauplii and then followed with *A. tonsa* nauplii and
1017 copepodites (CC), and *A. tonsa* nauplii and then gilthead sea bream (*S. aurata*) yolk sac larvae
1018 (CY). Values are normalized expression ratios, corresponding to an average of 8 pooled samples
1019 in Trial 1 (n = 8) and 6 pooled samples in Trial 2 (n = 6) with standard errors (SEM). Different
1020 superscript letters denote differences among the dietary treatments.

1021
1022 **Fig. 2.** Nutritional regulation of fatty acid synthase (*fas*), delta-6 fatty acyl desaturase (*fads2d6*) and
1023 fatty acyl elongase 5 (*elovl5*) gene transcription in Atlantic bluefin tuna (ABT) larvae 13 days
1024 after hatch (Trial 1) fed with rotifer *B. rotundiformis* enriched with Algamac 3050[®] (R), *A. tonsa*
1025 copepod nauplii (C), and ABT larvae 18 days after hatch (Trial 2) fed with rotifer *B.*
1026 *rotundiformis* and then *Artemia* metanauplii enriched with Algamac 3050[®] (RA), *A. tonsa*
1027 copepod nauplii and then *Artemia* enriched with Algamac 3050[®] (CA), *A. tonsa* copepod nauplii
1028 and then followed with *A. tonsa* nauplii and copepodites (CC), and *A. tonsa* nauplii and then
1029 gilthead sea bream (*S. aurata*) yolk sac larvae (CY). Values are normalized expression ratios,
1030 corresponding to an average of 8 pooled samples in Trial 1 (n = 8) and 6 pooled samples in Trial
1031 2 (n = 6) with standard errors (SEM). Different superscript letters denote differences among the
1032 dietary treatments.

1033
1034 **Fig.3.** Nutritional regulation of carnitine palmitoyl transferase I (*cptI*) and acyl coA oxidase (*aco*)
1035 gene transcription in Atlantic bluefin tuna ((ABT) larvae 13 days after hatch (Trial 1) fed with

1036 rotifer *B. rotundiformis* enriched with Algamac 3050[®] (R), *A. tonsa* copepod nauplii (C), and
1037 ABT larvae 18 days after hatch (Trial 2) fed with rotifer *B. rotundiformis* and then *Artemia*
1038 metanauplii enriched with Algamac 3050[®] (RA), *A. tonsa* copepod nauplii and then *Artemia*
1039 enriched with Algamac 3050[®] (CA), *A. tonsa* copepod nauplii and then followed with *A. tonsa*
1040 nauplii and copepodites (CC), and *A. tonsa* nauplii and then gilthead sea bream (*S. aurata*) yolk
1041 sac larvae (CY). Values are normalized expression ratios, corresponding to an average of 8
1042 pooled samples in Trial 1 (n = 8) and 6 pooled samples in Trial 2 (n = 6) with standard errors
1043 (SEM). Different superscript letters denote differences among the dietary treatments.

1044
1045 **Fig.4.** Nutritional regulation of fatty acid binding protein 2, 4 and 6 (*fabp2*, *fabp4* and *fabp7*
1046 respectively), lipoprotein lipase (*lpl*) and 3-hydroxy-3-methylglutaryl-CoA lyase (*hmgcl*) gene
1047 transcription in Atlantic bluefin tuna (ABT) larvae 13 days after hatch (Trial 1) fed with rotifer
1048 *B. rotundiformis* enriched with Algamac 3050[®] (R), *A. tonsa* copepod nauplii (C), and ABT
1049 larvae 18 days after hatch (Trial 2) fed with rotifer *B. rotundiformis* and then *Artemia* metanauplii
1050 enriched with Algamac 3050[®] (RA), *A. tonsa* copepod nauplii and then *Artemia* enriched with
1051 Algamac 3050[®] (CA), *A. tonsa* copepod nauplii and then followed with *A. tonsa* nauplii and
1052 copepodites (CC), and *A. tonsa* nauplii and then gilthead sea bream (*S. aurata*) yolk sac larvae
1053 (CY). Values are normalized expression ratios, corresponding to an average of 8 pooled samples
1054 in Trial 1 (n = 8) and 6 pooled samples in Trial 2 (n = 6) with standard errors (SEM). Different
1055 superscript letters denote differences among the dietary treatments.

1056
1057 **Fig.5.** Nutritional regulation of peroxisome proliferator-activated receptor alpha (*ppara*), gamma
1058 (*ppar γ*), sterol regulatory element-binding protein 1 and 2 (*srebp1* and *srebp2* respectively),
1059 retinoid X receptor (*rxr*) and liver X receptor (*lxr*) gene transcription in Atlantic bluefin tuna
1060 (ABT) larvae 13 days after hatch (Trial 1) fed with rotifer *B. rotundiformis* enriched with
1061 Algamac 3050[®] (R), *A. tonsa* copepod nauplii (C), and ABT larvae 18 days after hatch (Trial 2)

1062 fed with rotifer *B. rotundiformis* and then *Artemia* metanauplii enriched with Algamac 3050[®]
1063 (RA), *A. tonsa* copepod nauplii and then *Artemia* enriched with Algamac 3050[®] (CA), *A. tonsa*
1064 copepod nauplii and then followed with *A. tonsa* nauplii and copepodites (CC), and *A. tonsa*
1065 nauplii and then gilthead sea bream (*S. aurata*) yolk sac larvae (CY). Values are normalized
1066 expression ratios, corresponding to an average of 8 pooled samples in Trial 1 (n = 8) and 6 pooled
1067 samples in Trial 2 (n = 6) with standard errors (SEM). Different superscript letters denote
1068 differences among the dietary treatments.

1069
1070 **Fig.6.** Nutritional regulation of superoxide dismutase (*sod*), catalase (*cat*) and glutathione peroxidase
1071 1 and 4 (*gpx1* and *gpx4* respectively) gene transcription in Atlantic bluefin tuna (ABT) larvae 13
1072 days after hatch (Trial 1) fed with rotifer *B. rotundiformis* enriched with Algamac 3050[®] (R), *A.*
1073 *tonsa* copepod nauplii (C), and ABT larvae 18 days after hatch (Trial 2) fed with rotifer *B.*
1074 *rotundiformis* and then *Artemia* metanauplii enriched with Algamac 3050[®] (RA), *A. tonsa*
1075 copepod nauplii and then *Artemia* enriched with Algamac 3050[®] (CA), *A. tonsa* copepod nauplii
1076 and then followed with *A. tonsa* nauplii and copepodites (CC), and *A. tonsa* nauplii and then
1077 gilthead sea bream (*S. aurata*) yolk sac larvae (CY). Values are normalized expression ratios,
1078 corresponding to an average of 8 pooled samples in Trial 1 (n = 8) and 6 pooled samples in Trial
1079 2 (n = 6) with standard errors (SEM). Different superscript letters denote differences among the
1080 dietary treatments.

1081
1082 **Fig.7.** Nutritional regulation of trypsin (*tryp*), amino peptidase (*anpep*), amylase (*amy*), pancreatic
1083 lipase (*pl*), phospholipase A₂ (*pla2*) and bile salt activated lipase 1 and 2 (*ball* and *bal2*
1084 respectively) gene transcription in Atlantic bluefin tuna (ABT) larvae 13 days after hatch (Trial
1085 1) fed with rotifer *B. rotundiformis* enriched with Algamac 3050[®] (R), *A. tonsa* copepod nauplii
1086 (C), and ABT larvae 18 days after hatch (Trial 2) fed with rotifer *B. rotundiformis* and then
1087 *Artemia* metanauplii enriched with Algamac 3050[®] (RA), *A. tonsa* copepod nauplii and then
1088 *Artemia* enriched with Algamac 3050[®] (CA), *A. tonsa* copepod nauplii and then followed with

1089 *A. tonsa* nauplii and copepodites (CC), and *A. tonsa* nauplii and then gilthead sea bream (*S.*
1090 *aurata*) yolk sac larvae (CY). Values are normalized expression ratios, corresponding to an
1091 average of 8 pooled samples in Trial 1 (n = 8) and 6 pooled samples in Trial 2 (n = 6) with
1092 standard errors (SEM). Different superscript letters denote differences among the dietary
1093 treatments.
1094

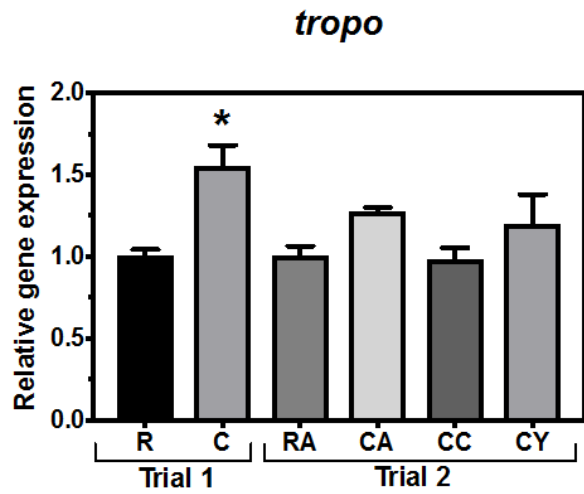
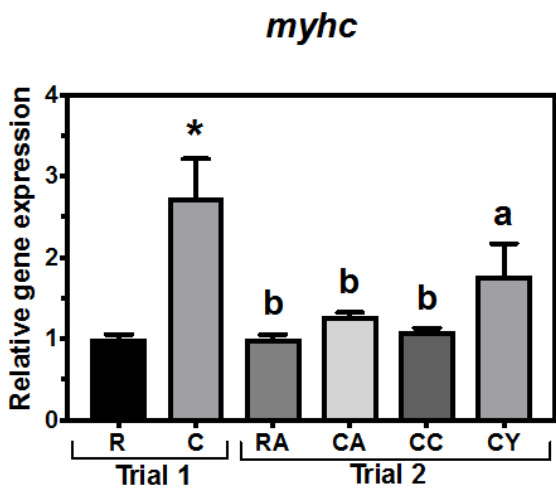


Fig.1

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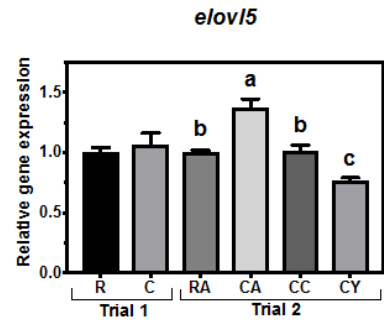
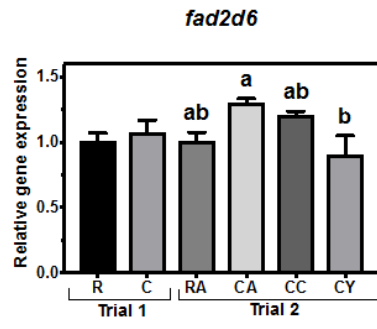
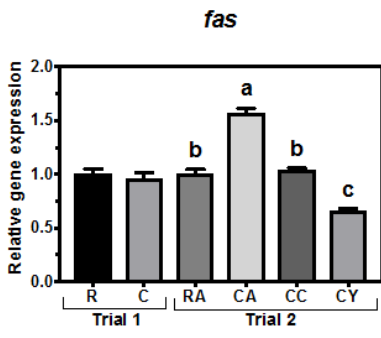


Fig.2

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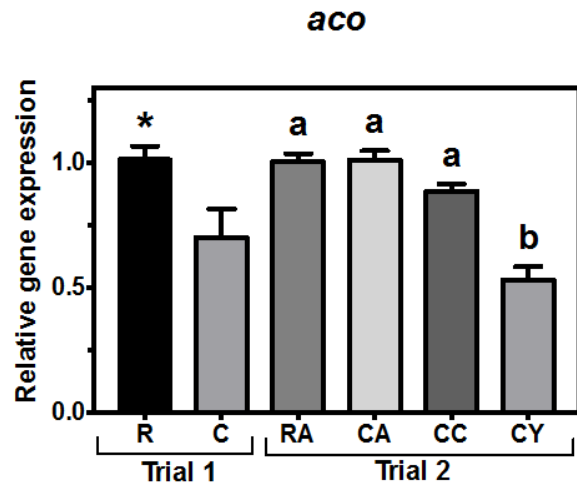
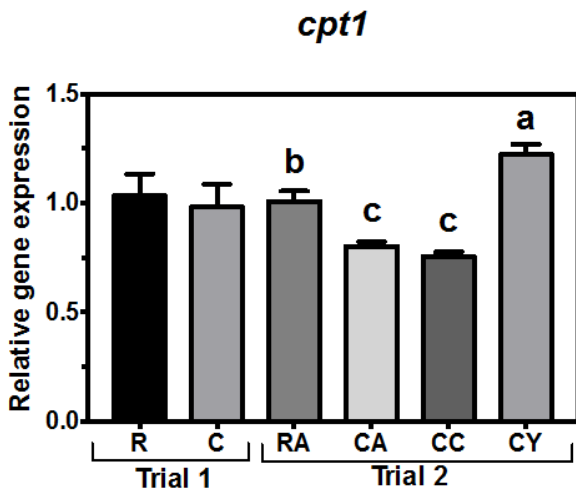


Fig.3

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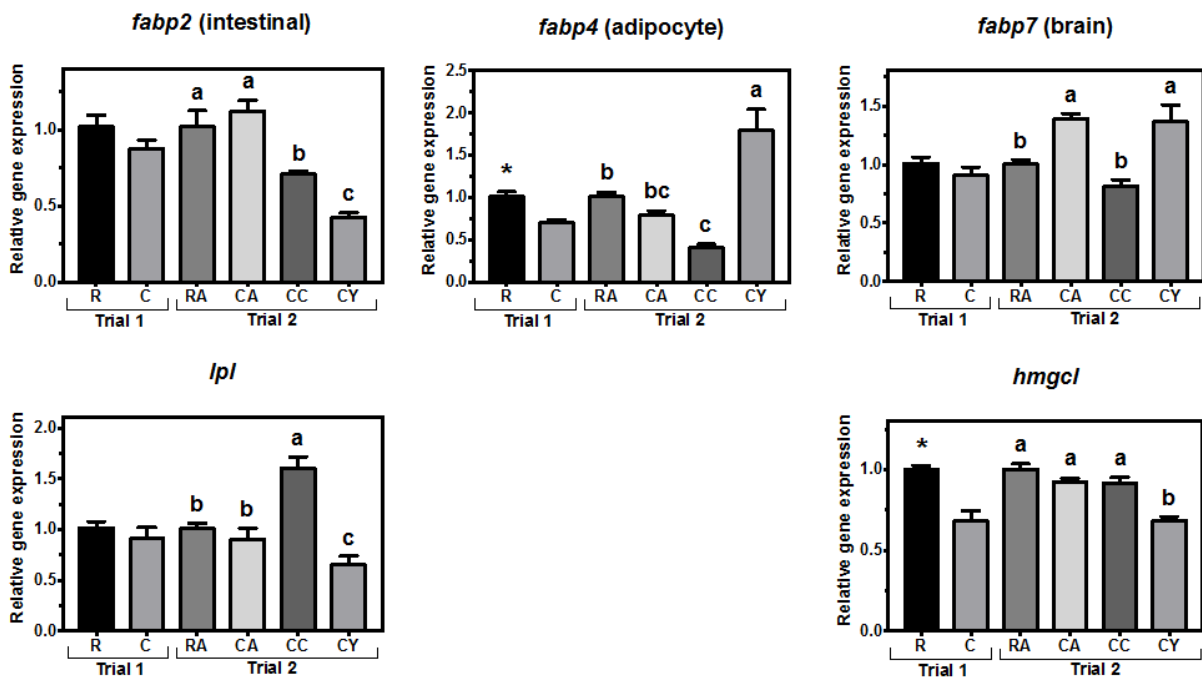


Fig.4

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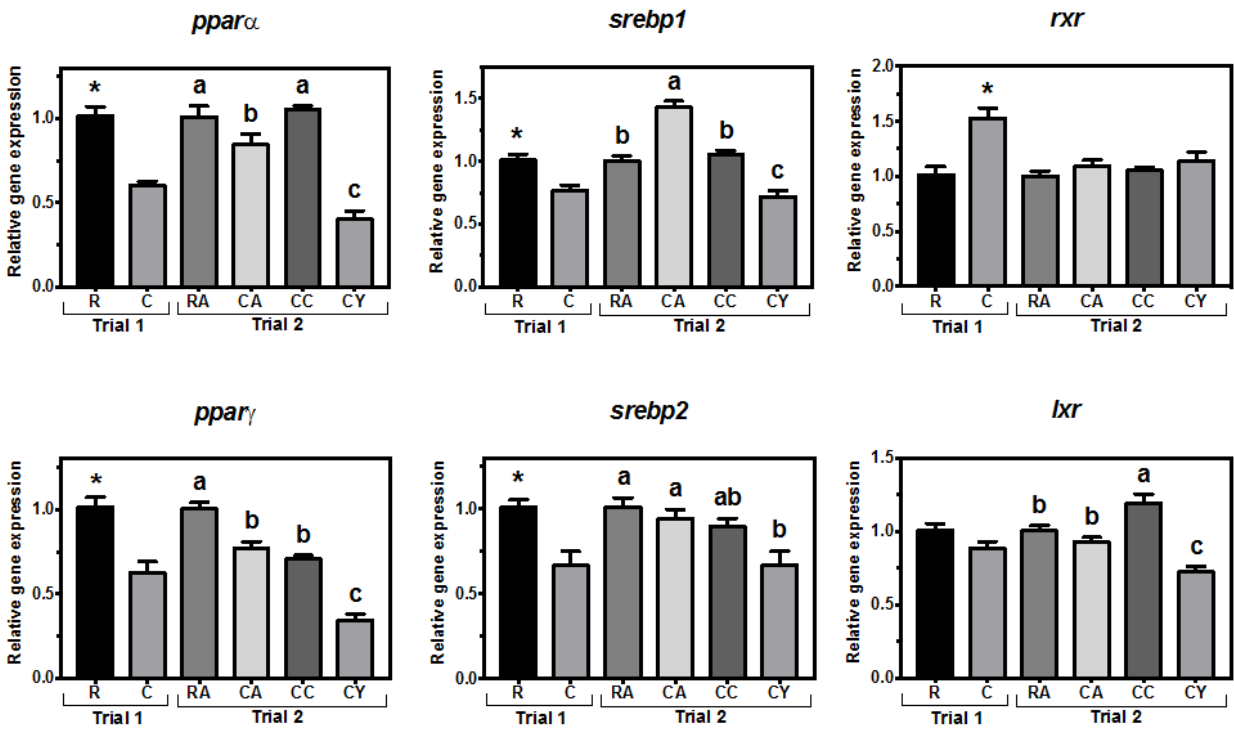


Fig.5

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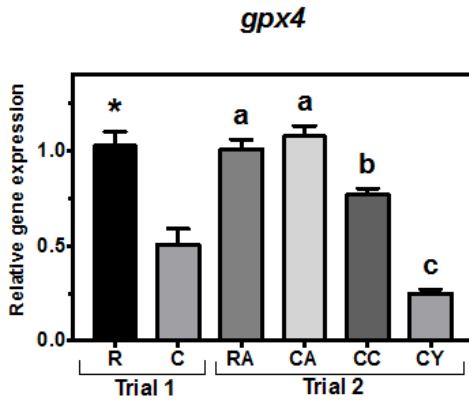
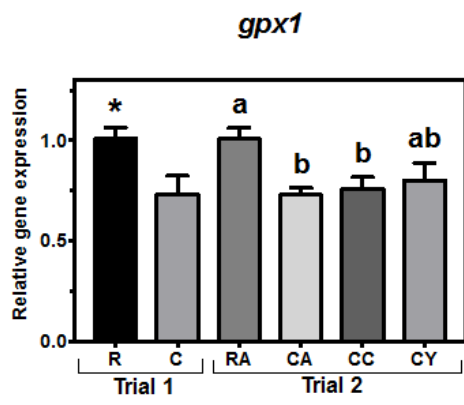
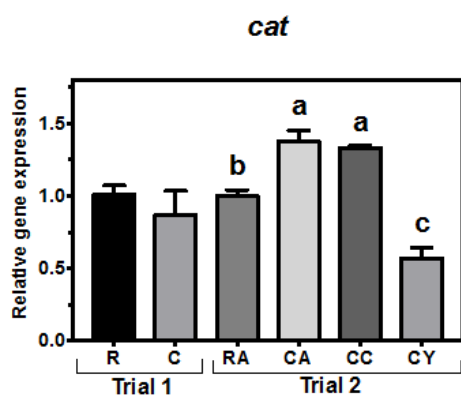
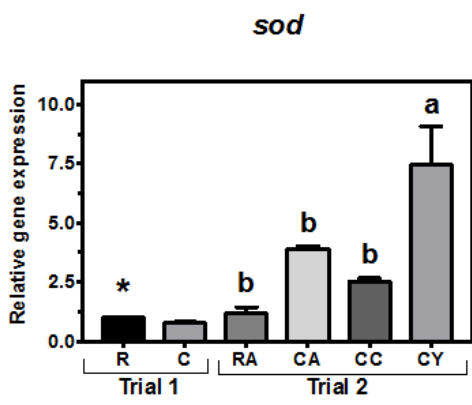


Fig.6

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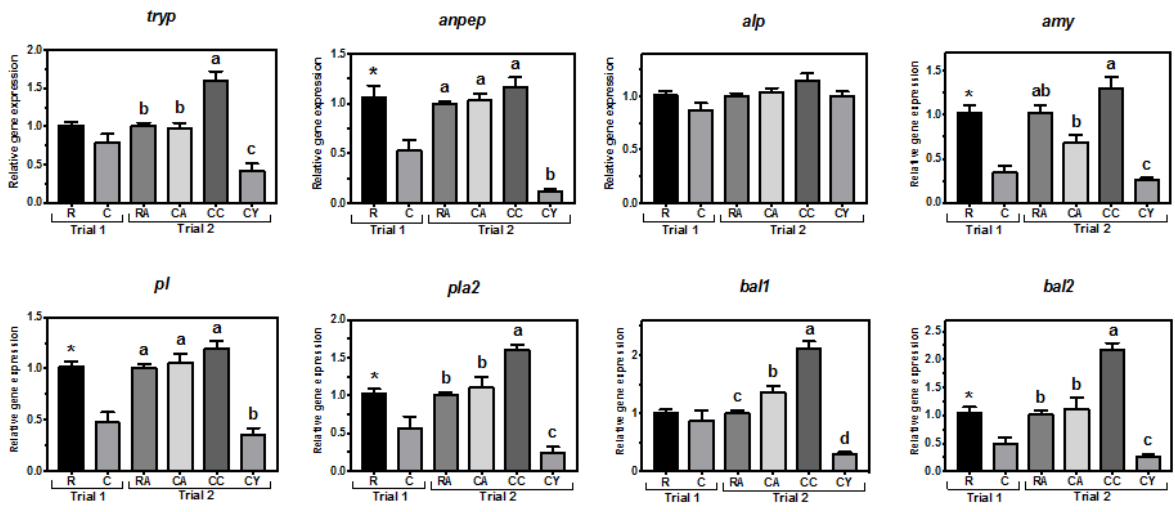


Fig.7

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1109 **Table 1.** Total gross composition (% dry mass), energy content (kcal.g⁻¹), taurine (mg.g⁻¹), vitamin
 1110 E (µg.g⁻¹dry mass), Se (µg.g⁻¹dry mass) and lipid class composition (total lipid %) of rotifer
 1111 *Brachionus rotundiformis* (ROT) enriched with Algamac 3050®, nauplii of the copepod *Acartia*
 1112 *tonsa* (COP) fed with the microalgae *Rhodomonas baltica*, *Artemia* metanauplii (ART) enriched with
 1113 Algamac 3050®, and gilthead sea bream (*Sparus aurata* L.) yolk sac larvae (YSL) used to feed
 1114 Atlantic bluefin tuna (*Thunnus thynnus* L.) larvae.
 1115

	ROT	COP	ART	
1116				
1117				
1118				YSL
1119				
1120	Composition (% dry mass)			
1121	51.3 ± 0.4 ^c	63.2 ± 2.1 ^a	53.8 ± 0.8 ^c	58.7 ± 0.4 ^b
1122	12.0 ± 0.5 ^b	8.8 ± 0.3 ^c	18.1 ± 0.9 ^a	18.9 ± 0.4 ^a
1123	30.1 ± 0.6 ^a	20.4 ± 1.1 ^b	20.0 ± 0.7 ^b	13.7 ± 0.3 ^c
1124	6.7 ± 0.3 ^c	7.6 ± 0.3 ^b	8.1 ± 0.1 ^a	8.7 ± 0.2 ^a
1125				
1126	Energy (kcal.g ⁻¹)			
1127	28.6 ± 0.7 ^c	35.3 ± 0.9 ^a	30.2 ± 0.6 ^c	33.0 ± 0.4 ^b
1128	11.3 ± 0.4 ^b	8.5 ± 0.3 ^c	16.9 ± 0.7 ^a	17.9 ± 0.6 ^a
1129	12.6 ± 0.3 ^a	8.4 ± 0.3 ^b	8.4 ± 0.5 ^b	5.9 ± 0.3 ^c
1130	52.5 ± 0.6 ^b	52.2 ± 0.7 ^b	55.5 ± 0.6 ^a	56.8 ± 0.5 ^a
1131				
1132	2.5 ± 0.2 ^d	4.1 ± 0.2 ^c	6.1 ± 0.2 ^b	11.0 ± 0.8 ^a
1133	232.3 ± 1.7 ^b	170.1 ± 6.5 ^c	78.9 ± 5.8 ^d	308.7 ± 4.6 ^a
1134	4.8 ± 0.2 ^a	0.4 ± 0.0 ^c	1.8 ± 0.5 ^b	1.4 ± 0.1 ^b
1135				
1136				
1137	Lipid classes (% total lipid)			
1138				
1139	12.3 ± 0.3 ^c	16.2 ± 0.3 ^b	9.2 ± 0.2 ^d	18.7 ± 1.0 ^a
1140	13.1 ± 1.1 ^b	11.8 ± 0.2 ^b	7.7 ± 0.2 ^c	17.1 ± 0.4 ^a
1141	41.0 ± 0.3 ^c	52.9 ± 0.7 ^a	28.6 ± 1.0 ^d	47.9 ± 0.5 ^b
1142				
1143	6.6 ± 0.3 ^c	11.9 ± 0.7 ^b	11.8 ± 0.4 ^b	21.7 ± 0.8 ^a
1144	40.2 ± 0.3 ^b	23.3 ± 0.3 ^c	46.3 ± 0.5 ^a	10.0 ± 0.7 ^d
1145	59.0 ± 0.3 ^b	47.1 ± 0.7 ^d	71.4 ± 1.0 ^a	52.1 ± 0.5 ^c
1146				
1147	0.7 ± 0.1 ^{ab}	1.1 ± 0.2 ^a	0.4 ± 0.1 ^b	0.9 ± 0.2 ^a
1148				

1149
 1150 Results are means ± SD (n = 3). An SD of 0.0 implies an SD of < 0.05. Values bearing different
 1151 superscript letter are significantly different (P < 0.05). TNL, total neutral lipids; PC,
 1152 phosphatidylcholine; PE, phosphatidylethanolamine; TPL, total polar lipids.
 1153
 1154

1155 **Table 2.** Sufficiency Index (SI) of essential amino acids (Oser, 1959) of rotifer *Brachionus*
 1156 *rotundiformis* (ROT) enriched with Algamac 3050®, nauplii of the copepod *Acartia tonsa* (COP) fed
 1157 with the microalgae *Rhodomonas baltica*, *Artemia metanauplii* (ART) enriched with Algamac 3050®
 1158 and gilthead sea bream (*Sparus aurata* L.) yolk sac larvae (YSL) used to feed Atlantic bluefin tuna
 1159 (*Thunnus thynnus* L.) larvae in Trials 1 and 2.
 1160

	ROT	COP	ART	YSL
1164 Taurine	60.5	99.3	147.7	266.3
1166 Valine	68.0	107.9	80.2	97.2
1168 Isoleucine	78.4	970.1	835.8	947.8
1169 Leucine	73.3	101.5	79.3	110.7
1170 Phenylalanine	87.4	116.1	91.4	114.3
1171 Histidine	54.1	93.7	74.6	121.9
1172 Lysine	74.6	113.8	100.2	124.4
1173 Arginine	79.4	127.6	110.4	127.6
1174 Threonine	69.3	121.6	79.6	115.0
1175 Methionine	48.5	95.9	63.5	101.1

1177
 1178 Results are means (n = 3). Values equal or above 100 indicate sufficient amount of that amino acid in
 1179 the live prey, whereas values below 100 show insufficiency for that amino acid.
 1180
 1181

1182 **Table 3.** Total lipid fatty acid composition (weight %) of rotifer *Brachionus rotundiformis* (ROT)
 1183 enriched with Algamac 3050[®], nauplii of the copepod *Acartia tonsa* (COP) fed with the microalgae
 1184 *Rhodomonas baltica*, *Artemia metanauplii* (ART) enriched with Algamac 3050 and gilthead sea
 1185 bream (*Sparus aurata* L.) yolk sac larvae (YSL) used to feed Atlantic bluefin tuna (*Thunnus thynnus*
 1186 L.) larvae.

	ROT	COP	ART	YSL
1190 Fatty acid				
1191				
1192 14:0	2.3 ± 0.1 ^b	9.8 ± 0.3 ^a	1.5 ± 0.1 ^c	2.1 ± 0.2 ^b
1193 16:0	15.1 ± 0.4 ^b	13.0 ± 0.3 ^c	14.9 ± 0.3 ^b	18.7 ± 0.3 ^a
1194 18:0	4.8 ± 0.2 ^a	3.0 ± 0.1 ^b	5.1 ± 0.1 ^a	4.6 ± 0.4 ^{ab}
1195 Total saturated ¹	23.9 ± 0.3 ^b	27.8 ± 0.8 ^a	22.4 ± 0.4 ^b	26.0 ± 0.4 ^a
1196				
1197 16:1n-7	1.9 ± 0.2 ^c	3.3 ± 0.2 ^b	2.3 ± 0.1 ^c	4.8 ± 0.2 ^a
1198 18:1n-9	2.2 ± 0.1 ^c	4.2 ± 0.1 ^b	17.2 ± 0.2 ^a	15.9 ± 0.4 ^a
1199 18:1n-7	1.6 ± 0.2 ^c	2.0 ± 0.1 ^{bc}	6.5 ± 0.1 ^a	2.8 ± 0.2 ^b
1200 20:1n-9	0.9 ± 0.2 ^a	0.3 ± 0.0 ^c	0.6 ± 0.1 ^a	0.6 ± 0.1 ^a
1201 Total monoenes ²	12.6 ± 0.4 ^b	11.5 ± 0.3 ^b	27.5 ± 0.6 ^a	26.3 ± 0.4 ^a
1202				
1203 C16 PUFA	6.3 ± 0.3 ^a	3.1 ± 0.1 ^b	0.8 ± 0.2 ^c	0.9 ± 0.2 ^c
1204				
1205 18:2n-6	16.3 ± 0.7 ^a	5.1 ± 0.3 ^c	4.3 ± 0.1 ^d	7.1 ± 0.3 ^b
1206 20:4n-6	1.1 ± 0.1 ^c	1.4 ± 0.1 ^b	1.8 ± 0.2 ^a	1.6 ± 0.2 ^{ab}
1207 22:5n-6	2.9 ± 0.4 ^a	3.3 ± 0.2 ^a	3.2 ± 0.2 ^a	0.3 ± 0.1 ^b
1208 Total n-6PUFA ³	24.8 ± 0.7 ^a	13.6 ± 0.3 ^b	10.0 ± 0.3 ^{bc}	9.7 ± 0.6 ^c
1209				
1210 18:3n-3	3.9 ± 0.2 ^b	4.7 ± 0.2 ^b	23.0 ± 0.2 ^a	0.9 ± 0.1 ^c
1211 18:4n-3	0.2 ± 0.0 ^c	4.8 ± 0.3 ^a	2.8 ± 0.1 ^b	0.6 ± 0.1 ^c
1212 20:4n-3	1.0 ± 0.1 ^a	0.6 ± 0.1 ^b	0.6 ± 0.1 ^b	0.6 ± 0.1 ^b
1213 20:5n-3	5.1 ± 0.3 ^b	4.4 ± 0.3 ^{bc}	4.1 ± 0.3 ^c	7.2 ± 0.2 ^a
1214 22:5n-3	3.2 ± 0.4 ^a	0.2 ± 0.0 ^b	0.2 ± 0.0 ^b	2.6 ± 0.2 ^a
1215 22:6n-3	14.1 ± 0.6 ^b	27.0 ± 0.9 ^a	8.0 ± 0.2 ^c	25.1 ± 0.9 ^a
1216 Total n-3PUFA ⁴	32.9 ± 0.9 ^c	43.9 ± 1.6 ^a	39.3 ± 0.3 ^b	37.2 ± 0.8 ^b
1217				
1218 Total PUFA	57.7 ± 1.6 ^a	57.6 ± 1.4 ^a	50.1 ± 0.5 ^b	47.7 ± 0.9 ^b
1219				
1220 n-3/n-6	1.3 ± 0.1 ^c	3.2 ± 0.2 ^b	3.9 ± 0.1 ^a	3.8 ± 0.2 ^a
1221 DHA/EPA	2.8 ± 0.3 ^c	6.1 ± 0.7 ^a	1.9 ± 0.1 ^d	3.5 ± 0.2 ^b

1222 Results are means ± SD (n = 3). An SD of 0.0 implies an SD of < 0.05. Values bearing different
 1223 superscript letters are significantly different (P<0.05). ¹, Totals include 15:0, 20:0, 22:0 and 24:0; ²,
 1224 Totals include 16:1n-9, 18:1n-11, 20:1n-7, 22:1 isomers and 24:1; ³, Totals include 18:3n-6, 20:2n-6
 1225 and 22:4n-6; ⁴, Totals include 20:3n-3 and 22:3n-3; DHA, docosahexaenoic acid; EPA,
 1226 eicosapentaenoic acid; PUFA, polyunsaturated fatty acid.
 1227
 1228
 1229

Table 4. Growth performance and survival of ABT larvae in Trial 1 (13 dah) and Trial 2 (18 dah). Total length (mm), total weight as live mass or dry mass per larvae (mg), dry mass (%) and survival rate (%) of ABT larvae 13 days after hatch fed on rotifer *B. rotundiformis* enriched with Algamac 3050® (treatment R) or *A. tonsa* copepod nauplii (treatment C), and ABT larvae 18 days after hatch that were fed rotifer *B. rotundiformis* and then *Artemia* metanauplii enriched with Algamac 3050® (RA), *A. tonsa* copepod nauplii and then *Artemia* enriched with Algamac 3050® (treatment CA), *A. tonsa* copepod nauplii and then followed with *A. tonsa* nauplii and copepodites (treatment CC), and *A. tonsa* nauplii and then gilthead sea bream (*S. aurata*) yolk sac larvae (treatment CY).

Dietary treatments	Trial 1(13 dah ABT larvae)		RA	Trial 2 (18 dah ABT larvae)		
	R	C		CA	CC	CY
Total length (mm)	6.8 ± 0.3	7.6 ± 0.4 *	8.2 ± 0.4 ^b	9.7 ± 0.5 ^{ab}	8.9 ± 0.6 ^b	10.3 ± 0.5 ^a
Total weight (mg/larvae live mass)	4.3 ± 0.2	8.2 ± 0.5 *	9.7 ± 0.5 ^c	11.6 ± 0.5 ^b	11.4 ± 0.6 ^b	24.3 ± 0.9 ^a
Total weight (mg/larvae dry mass)	0.6 ± 0.1	1.1 ± 0.2 *	1.2 ± 0.3 ^c	2.5 ± 0.4 ^b	1.9 ± 0.3 ^{bc}	3.6 ± 0.4 ^a
Dry mass (%)	14.4 ± 0.8	12.8 ± 0.9	12.2 ± 0.7 ^d	21.2 ± 1.1 ^a	16.9 ± 0.9 ^b	14.9 ± 0.8 ^c
Flexion Index (%)	35.8 ± 3.9	54.5 ± 5.0 *	100	100	100	100
Survival at 13 dah (%)	13.9 ± 11.8	8.2 ± 3.8				
Survival at 18 dah (%)			2.4	2.1	2.3	3.6
Survival (13 -18 dah, 5 days) (%)			17.6	19.5	21.7	32.0

Results are means ± SD (n =4 in Trial 1 and n = 20 in Trial 2 for total length, total weight, dry mass and flexion index; n = 4 per treatment for survival in Trial A). An SD of 0.0 implies an SD of < 0.05. Values bearing an asterisk (*) between samples of 13 dah ABT larvae are significantly different (P < 0.05). Values of 18 dah ABT larvae bearing different superscript letters are significantly different (P < 0.05).

Table 5. Total lipid content (% of live and dry mass) and lipid class composition (total lipid %) of Atlantic bluefin tuna (*Thunnus thynnus* L.) larvae (ABT) 13 days after hatch fed with rotifer *Brachionus rotundiformis* enriched with Algamac 3050® (R), *Acartia tonsa* copepod nauplii (C), and ABT larvae 18 days after hatch that had been fed with rotifer *B. rotundiformis* and then *Artemia* metanauplii enriched with Algamac 3050® (RA), *Acartia* copepod nauplii and then *Artemia* enriched with Algamac 3050® (CA), *Acartia* copepod nauplii and then followed with *Acartia* nauplii and copepodites (CC), and *Acartia* nauplii and then gilthead sea bream (*Sparus aurata* L.) yolk sac larvae (CY).

Dietary treatments	13 dah ABT larvae		18 dah ABT larvae			
	R	C	RA	CA	CC	CY
Total lipid (% live mass)	1.3 ± 0.1	1.1 ± 0.1	1.3 ± 0.1	1.7 ± 0.2	1.5 ± 0.2	1.8 ± 0.3
Total lipid (% dry mass)	9.1 ± 0.6	8.0 ± 0.7	10.5 ± 0.9 ^a	7.4 ± 0.6 ^b	8.3 ± 0.5 ^b	12.1 ± 0.9 ^a
Lipid classes (total lipid %)						
Phosphatidylcholine	21.2 ± 1.1	22.2 ± 1.2	21.6 ± 1.0	21.6 ± 0.4	23.2 ± 0.7	20.6 ± 0.9
Phosphatidylethanolamine	14.5 ± 1.2	14.6 ± 0.8	14.7 ± 0.8 ^a	14.0 ± 0.5 ^a	14.6 ± 0.3 ^a	11.4 ± 0.3 ^b
Phosphatidylserine	6.6 ± 0.2	7.2 ± 0.7	5.8 ± 0.4 ^b	6.5 ± 0.3 ^{ab}	7.8 ± 0.4 ^a	4.8 ± 0.2 ^c
Phosphatidylinositol	4.6 ± 0.3	3.7 ± 0.2 *	4.2 ± 0.6	4.5 ± 0.5	4.6 ± 0.2	4.0 ± 0.5
Sphingomyelin	2.2 ± 0.1	3.8 ± 0.4 *	2.1 ± 0.1 ^c	3.3 ± 0.1 ^b	4.0 ± 0.2 ^a	2.5 ± 0.2 ^{bc}
Phosphatidic acid/cardiolipin	3.3 ± 0.2	3.3 ± 0.1	3.2 ± 0.3 ^a	2.9 ± 0.7 ^{ab}	2.5 ± 0.2 ^b	2.5 ± 0.3 ^b
Lysophosphatidylcholine	0.4 ± 0.1	0.6 ± 0.2	0.4 ± 0.1	0.5 ± 0.1	0.5 ± 0.1	0.4 ± 0.0
Pigmented material	3.6 ± 0.8	3.1 ± 0.3	2.6 ± 0.2 ^{ab}	3.3 ± 0.5 ^a	2.5 ± 0.3 ^{ab}	2.0 ± 0.1 ^b
Total polar lipids	56.5 ± 1.7	58.5 ± 3.4	54.6 ± 2.1 ^b	56.4 ± 1.8 ^b	59.4 ± 0.6 ^a	48.2 ± 3.2 ^{bc}
Cholesterol/sterols	18.1 ± 0.6	17.5 ± 1.4	16.4 ± 1.1 ^{ab}	16.1 ± 1.8 ^{ab}	18.5 ± 0.6 ^a	14.6 ± 2.1 ^b
Free fatty acids	4.9 ± 1.1	6.2 ± 0.1	5.6 ± 0.7	5.1 ± 1.1	6.0 ± 0.6	5.7 ± 0.8
Diacylglycerol	2.1 ± 0.1	1.8 ± 0.2	1.4 ± 0.1 ^b	1.8 ± 0.5 ^{ab}	2.0 ± 0.3 ^a	1.9 ± 0.2 ^a
Triacylglycerol	14.5 ± 0.7	10.7 ± 1.0 *	19.8 ± 1.2 ^a	15.4 ± 0.5 ^b	11.5 ± 0.3 ^c	22.4 ± 1.2 ^a
Wax/Sterol esters	3.9 ± 1.0	5.3 ± 0.7 *	2.2 ± 0.4 ^c	5.4 ± 1.3 ^b	2.7 ± 0.4 ^c	7.2 ± 0.9 ^a
Total neutral lipids	43.5 ± 1.7	41.5 ± 2.4	45.4 ± 1.1 ^b	43.7 ± 1.7 ^{bc}	40.6 ± 0.8 ^c	51.8 ± 2.3 ^a

Results are means ± SD (n = 4 in Trial 1 and n = 3 in Trial 2). An SD of 0.0 implies an SD of < 0.05. Values bearing an asterisk (*) between samples of 13 dah ABT larvae (R and C) are significantly different (P < 0.05). Values of 18 dah ABT larvae (samples RA, CA, CC and CY) bearing different superscript letters are significantly different (P < 0.05).

Table 6. Total lipid fatty acid composition (weight %) of Atlantic bluefin tuna (*T. thynnus* L.) larvae 13 days after hatch fed with rotifer *B. rotundiformis* enriched with Algamac 3050® (R), *A. tonsa* copepod nauplii (C), and bluefin tuna larvae 18 days after hatch fed with rotifer *B. rotundiformis* and then *Artemia* metanauplii enriched with Algamac 3050® (RA), *A. tonsa* copepod nauplii and then *Artemia* enriched with Algamac 3050® (CA), *A. tonsa* copepod nauplii and then followed with *A. tonsa* nauplii and copepodites (CC), and *A. tonsa* nauplii and then gilthead sea bream (*S. aurata*) yolk sac larvae (CY).

Dietary treatments	13 dah ABT larvae		18 dah ABT larvae			
	R	C	RA	CA	CC	CY
Fatty acid (weight %)						
14:0	1.1 ± 0.2	1.4 ± 0.2	0.4 ± 0.1 ^c	0.8 ± 0.2 ^b	1.6 ± 0.2 ^a	0.4 ± 0.1 ^c
16:0	24.9 ± 0.9	21.2 ± 0.6 *	17.7 ± 0.7 ^b	16.7 ± 0.5 ^b	20.5 ± 0.2 ^a	21.8 ± 0.3 ^a
18:0	11.4 ± 0.8	12.4 ± 0.4	12.9 ± 0.5 ^a	13.0 ± 0.4 ^a	11.4 ± 0.1 ^b	12.0 ± 0.2 ^{ab}
Total SFA ¹	39.5 ± 1.5	37.5 ± 1.2	33.7 ± 1.3 ^b	33.1 ± 0.3 ^b	36.9 ± 0.9 ^a	36.3 ± 0.5 ^a
16:1n-7	5.5 ± 0.2	2.2 ± 0.2 *	1.8 ± 0.1 ^b	1.8 ± 0.1 ^b	2.4 ± 0.3 ^a	2.1 ± 0.2 ^{ab}
18:1n-9	6.8 ± 0.2	6.9 ± 0.2	10.9 ± 0.6 ^b	10.8 ± 0.2 ^b	7.5 ± 0.5 ^c	13.2 ± 0.8 ^a
18:1n-7	3.1 ± 0.2	2.1 ± 0.1 *	4.8 ± 0.4 ^a	4.6 ± 0.5 ^a	2.0 ± 0.3 ^c	2.9 ± 0.3 ^b
20:1n-9	0.7 ± 0.2	1.1 ± 0.3	0.7 ± 0.1	0.4 ± 0.2	0.5 ± 0.1	0.6 ± 0.1
Total MUFA ²	20.0 ± 0.6	14.0 ± 0.4 *	20.2 ± 0.8 ^a	19.3 ± 0.4 ^a	14.2 ± 0.5 ^b	20.7 ± 1.4 ^a
C16 PUFA	2.1 ± 0.1	3.5 ± 0.5 *	3.1 ± 0.2 ^a	3.0 ± 0.2 ^a	3.5 ± 0.3 ^a	2.1 ± 0.1 ^b
18:2n-6	4.5 ± 0.3	3.0 ± 0.1 *	5.6 ± 0.3 ^a	4.8 ± 0.3 ^{ab}	4.6 ± 0.7 ^{ab}	4.1 ± 0.5 ^b
20:4n-6	5.1 ± 0.2	1.0 ± 0.1 *	1.6 ± 0.1 ^b	1.9 ± 0.2 ^a	1.1 ± 0.0 ^b	1.2 ± 0.1 ^b
22:5n-6	4.3 ± 0.3	3.1 ± 0.2 *	2.7 ± 0.2 ^b	3.0 ± 0.2 ^{ab}	3.6 ± 0.4 ^a	1.1 ± 0.1 ^c
Total n-6 PUFA ³	17.5 ± 0.6	10.1 ± 0.8 *	14.9 ± 0.6 ^a	12.1 ± 0.5 ^b	12.5 ± 0.3 ^b	9.8 ± 0.2 ^c
18:3n-3	0.3 ± 0.1	0.6 ± 0.1 *	6.9 ± 0.3 ^a	5.7 ± 0.4 ^b	1.8 ± 0.3 ^c	0.4 ± 0.1 ^d
18:4n-3	0.5 ± 0.1	1.5 ± 0.3 *	1.0 ± 0.2 ^b	1.8 ± 0.3 ^a	1.4 ± 0.3 ^{ab}	0.2 ± 0.0 ^c
20:4n-3	0.6 ± 0.1	0.6 ± 0.0	0.8 ± 0.2 ^a	0.8 ± 0.1 ^a	0.6 ± 0.1 ^{ab}	0.5 ± 0.1 ^b
20:5n-3	10.2 ± 0.8	3.1 ± 0.3 *	4.7 ± 0.2 ^a	3.4 ± 0.3 ^b	2.1 ± 0.2 ^c	4.4 ± 0.2 ^a
22:5n-3	0.3 ± 0.0	0.4 ± 0.1	2.3 ± 0.4 ^a	0.3 ± 0.0 ^b	0.4 ± 0.1 ^b	1.8 ± 0.3 ^a
22:6n-3	5.9 ± 0.4	25.3 ± 0.9 *	10.0 ± 0.4 ^c	15.8 ± 0.4 ^b	24.1 ± 1.0 ^a	22.2 ± 1.5 ^a
Total n-3 PUFA ⁴	19.5 ± 0.6	34.2 ± 1.8 *	28.2 ± 1.1 ^b	30.9 ± 1.4 ^{ab}	33.2 ± 1.3 ^a	30.7 ± 1.6 ^{ab}
Total PUFA	39.1 ± 1.7	47.5 ± 1.6 *	43.1 ± 1.6 ^{ab}	43.0 ± 0.7 ^{ab}	45.7 ± 1.0 ^a	40.5 ± 1.6 ^b
DHA/EPA	0.6 ± 0.1	8.2 ± 0.4 *	2.1 ± 0.1 ^b	4.6 ± 0.3 ^a	11.3 ± 0.7 ^b	5.1 ± 0.2 ^a
n-3/n-6	1.1 ± 0.2	3.4 ± 0.4 *	1.9 ± 0.3 ^c	2.6 ± 0.2 ^b	2.6 ± 0.2 ^b	3.1 ± 0.2 ^a
Unknown	1.4 ± 0.2	1.0 ± 0.2	3.0 ± 0.3 ^b	4.6 ± 0.2 ^a	3.2 ± 0.4 ^b	2.5 ± 0.4 ^c

Results are means ± SD (n = 4 in Trial 1 and n = 3 in Trial 2). An SD of 0.0 implies an SD of < 0.05. Values bearing an asterisk (*) between samples of 13 dah ABT larvae (a and b) are significantly different (P < 0.05). Values of 18 dah ABT larvae (samples c to f) bearing different superscript letters are significantly different (P < 0.05). ¹, Totals include 15:0, 20:0, 22:0 and 24:0; ², Totals include 16:1n-9, 18:1n-11, 20:1n-7, 22:1 isomers and 24:1; ³, Totals include 18:3n-6, 20:2n-6 and 22:4n-6; ⁴, Totals include 20:3n-3 and 22:3n-3; DHA, docosahexaenoic acid; EPA, eicosapentaenoic acid; MUFA, monounsaturated fatty acid; PUFA, polyunsaturated fatty acid; SFA, saturated fatty acid.

Table 7. Relative abundance (δ) of $^{15}\text{N}/^{14}\text{N}$ and $^{13}\text{C}/^{12}\text{C}$ (‰) and C:N ratio of rotifer *Brachionus rotundiformis* enriched with Algamac 3050® (R), nauplii of the copepod *Acartia tonsa* fed on the microalgae *Rhodomonas baltica* (C) and 13 dah Atlantic bluefin tuna (*Thunnus thynnus* L.) larvae from Trial 1 fed on *B. rotundiformis* (ABT R) and *A. tonsa* nauplii (ABT C), and isotopic enrichment (Δ) of ^{15}N and ^{13}C (‰) in ABT larvae in relation to its prey, rotifers and copepods.

C	R (e)	C	ABT R	ABT
$\delta^{15}\text{N}$ (‰)	-0.81 ± 0.02	4.39 ± 0.04 *	0.65 ± 0.05	7.18 ± 0.01 *
$\delta^{13}\text{C}$ (‰)	-11.09 ± 0.02	-20.17 ± 0.06 *	-10.61 ± 0.17	-17.11 ± 0.01 *
C:N	4.64 ± 0.03	3.74 ± 0.02 *	3.92 ± 0.01	3.67 ± 0.03 *
$\Delta^{15}\text{N}$ (‰)			1.46 ± 0.03	2.79 ± 0.02 *
$\Delta^{13}\text{C}$ (‰)			0.48 ± 0.08	3.06 ± 0.04 *

Results are means \pm SD ((n = 4 in Trial 1 and n = 3 in Trial 2). An SD of 0.0 implies an SD of < 0.05 . Values between live prey, or ABT larvae fed on these live prey, bearing an asterisk (*), are significantly different ($P < 0.05$).

Supplementary Table 1. Rearing conditions for feeding trials of Atlantic bluefin tuna (*Thunnus thynnus* L.) larvae.

	Trial A (0-13dph)	Trial B (13-18dph)
Age of initial fish (days after hatching)	just hatched larvae	13 dph
Experimental period	13 days	5 days
Tank volume (m ³)	1.4	1.4
Number of tanks per treatment	4	1
Number of fish (ind./tank)	11,900 eggs	600
Initial ABT larval density (larvae·L ⁻¹)	8.5	0.4
Live preys density (individuals·mL ⁻¹)		
Rotifers (<i>Brachionus rotundiformis</i>)	10	-(5)
Copepods (<i>Acartia tonsa</i>)	10	2.5
Artemia	-	0.5
Yolk sac larvae (<i>Sparus aurata</i>)	-	0.1
Water temperature (°C)	24.8±0.4°C	22.8±0.6°C
Dissolved oxygen (mg/l)	6.41±0.06	6.56±0.19
Photoperiod (hL:hD)	14:10	14:10
Exchange rate of sea water (% tank volume/day)	100-200%	100-200%

Supplementary Table 2. Sequence, annealing temperature (T_m) and size of the fragment produced by the primer pairs used for quantitative PCR (qPCR).

Name	Sequence (5'-3')	Amplicon size (bp)	T _m °C
<i>elovl5</i>	F: CCACGCTAGCATGCTGAATA	236	60
	R: ATGGCCATATGACTGCACAC		
<i>fads2d6</i>	F: CCGTGCACTGTGTGAGAAAC	152	60
	R: CAGTGTAAGCGATAAAAATCAGCTG		
<i>ppara</i>	F: TGGTCATGGAGGTGGAAGAC	152	60
	R: ATGGATGACGAAAGGAGGGG		
<i>ppary</i>	F: ACCTGACCAACATGGACTAC	118	60
	R: GAGAAAACAGGACTGTCAGC		
<i>lxr</i>	F: CACACTGGATCCACAACAGC	192	58
	R: ATCTCCTGCACCGACATGAT		
<i>rxr</i>	F: TGAGGGAAAAGGTCTACGCA	212	59
	R: TGTGATCTGATGTGGTGCCT		
<i>srebp1</i>	F: CCAGCTACACATGACAGGGA	153	59
	R: GCTTTGACCCTTAGAGCTGC		
<i>srebp2</i>	F: AGATCCAGTGAGTCGTTGGT	212	60
	R: CTACAGCCCCTTCTCCCTTC		
<i>fabp2</i>	F: CGCAGCGAGAATTATGACAA	244	55
	R: AGCATGTACACCCTCCATCTC		
<i>fabp4</i>	F: ACTGCAATGACCGAAAGACC	175	55
	R: CCTCCTTTCCGTAGGTCCTC		
<i>fabp7</i>	F: CCTACACCTGATGACCGACA	212	55
	R: GCTGGGATGATTTGCTCATT		
<i>cpt1</i>	F: TGGAGGCTGTCCACCAGTCA	211	60
	R: TGCTGGAGATGTGGAAGTTG		
<i>lpl</i>	F: CCGAAAGAACCGCTGCAATA	212	59
	R: GATCCTCCTTCTCTCCGTGG		
<i>fas</i>	F: ATACCGTGGCAATGTAACGC	188	59
	R: GTGAGCTGTGGATGATGCTG		
<i>aco</i>	F: AGCGCTATGACCAGGCTATT	164	59
	R: GTACAGGGTTGGGAGGAACA		
<i>hmgcl</i>	F: CGTGCCAACAGAGACGAAAA	173	59
	R: GGGTGAGGACTGGGTAAGAC		
<i>gpx1</i>	F: TGGAGAAAGTGGATGTGAACGG	309	55
	R: GTGCTGTGGAAGCTGTATGATGG		
<i>gpx4</i>	F: TGGGGAATAGCATCAAGTGG	206	55
	R: CGAGAAAGGAGGGAAACAGG		
<i>cat</i>	F: ATGGTGTGGGACTTCTGGAG		60
	R: ATGAAACGGTAGCCATCAGG		
<i>sod</i>	F: TCCCAGATCACCTACATGCC	182	59
	R: CTGCGGAGAGTTGCTTGATC		
<i>myhc</i>	F: GATTCAGCTGAGCCATGCCA	190	60
	R: TCTCAGCTCCTCAATCTCAG		
<i>anpep</i>	F: CCTGAGGTGGTGGAAATGACT		60
	R: GGGTTCAGCTTTGTCTGCTC		
<i>amy</i>	F: TCATGTGGAAGCTGGTTCAG		60

	R: AATATTGCCACTGCCAGTCC		
<i>tryp</i>	F: CCCCAACTACAACCCCTACA		60
	R: CCAGCCAGAGACAAGACACA		
<i>alp</i>	F: ACTCTGACAACGAGATGCCA	189	60
	R: TTCCGTCTTTTCTTGTGCCG		
<i>pl</i>	F: TTCCAGGACACTCCTGTTTCTGTGC	107	59
	R: ATCCCCAGACCAAGTTTGGAGTTGA		
<i>bal1</i>	F: CATGGATGGACACCTCTTACTGGT	126	59
	R: AAACCAGCCTGGCCCTTCTCTTTAG		
<i>bal2</i>	F: GGATGGGCACCTCTTCACATCACAG	120	59
	R: CCAGCTTGGCCCTTCTCTTTGGTAT		
<i>pla2</i>	F: GGATGATCTGGACAGGTGCT	217	59
	R: TCTGGCAAAACACTCAACGG		
<i>tropo</i>	F: AGAATGCCTTGGACAGAGCT	227	60
	R: ACGTCTGTTAAGGGAAGCGA		
<i>efla</i>	F: CCCCTGGACACAGAGACTTC	119	60
	R: GCCGTTCTTGGAGATACCAG		
<i>bactin</i>	F: ACCCACACAGTGCCCATCTA	155	61
	R: TCACGCACGATTTCCCTCT		

elovl5, fatty acyl elongase 5; *fads2d6*, delta-6 fatty acyl desaturase; *ppara*, peroxisome proliferator-activated receptor alpha; *ppary*, peroxisome proliferator-activated receptor gamma; *lxr*, liver X receptor; *rxr*, retinoid X receptor; *srebp1*, sterol regulatory element-binding protein 1; *srebp2*, sterol regulatory element-binding protein 2; *fabp2*, fatty acid binding protein 2 (intestinal); *fabp4*, fatty acid binding protein 4 (adipocyte); *fabp7*, fatty acid binding protein 7 (brain-type); *cpt1*, carnitine palmitoyl transferase I; *lpl*, lipoprotein lipase; *fas*, fatty acid synthase; *aco*, acyl coA oxidase; *hmgcl*, 3-hydroxy-3-methylglutaryl-CoA lyase; *gpx1*, glutathione peroxidase 1; *gpx4*, glutathione peroxidase 4; *cat*, catalase; *sod*, superoxide dismutase; *myhc*, myosin heavy chain; *anpep*, amino peptidase; *amy*, amylase; *tryp*, trypsin; *alp*, alkaline phosphatase; *pl*, pancreatic lipase; *bal1*, bile salt activated lipase 1; *bal2*, bile salt activated lipase 2; *pla2*, phospholipase A2; *tropo*, tropomyosin; *efla*, elongation factor 1 alpha; *bactin*, beta actin.

Supplementary Table 3. Total amino acid content, including taurine (mg/g dry mass) of 1 day post hatch ABT yolk sac larvae (ABT), rotifers *B. rotundiformis* enriched with Algamac 3050 Bio Marine® (ROT), *Artemia* metanauplii enriched with Algamac 3050 Bio Marine® (ART), 3 days post hatch nauplii of the copepod *Acartia tonsa* fed *Rhodomonas baltica* (COP) and gilthead sea bream (*Sparus aurata* L.) yolk sac larvae (YSL).

	(ABT)	(ROT)	(ART)	(COP)	(YSL)
Taurine	4.1 ± 0.1 ^c	2.5 ± 0.2 ^d	6.1 ± 0.2 ^b	4.1 ± 0.2 ^c	11.0 ± 0.8 ^a
EAA					
Valine	33.5 ± 0.3 ^{ab}	22.8 ± 0.8 ^c	26.9 ± 0.6 ^{bc}	36.2 ± 5.6 ^a	32.6 ± 1.2 ^{ab}
Isoleucine	2.7 ± 0.2 ^c	2.1 ± 0.1 ^c	22.4 ± 2.3 ^b	26.0 ± 1.3 ^a	25.4 ± 0.6 ^{ab}
Leucine	41.4 ± 0.2 ^b	30.3 ± 0.6 ^c	32.8 ± 1.1 ^c	42.0 ± 2.5 ^b	45.8 ± 1.3 ^a
Phenylalanine	22.3 ± 0.7 ^{bc}	19.5 ± 0.5 ^c	20.4 ± 0.6 ^b	25.9 ± 1.0 ^a	25.5 ± 0.7 ^a
Histidine	13.1 ± 0.2 ^b	7.1 ± 0.2 ^d	9.8 ± 0.3 ^c	12.3 ± 1.2 ^b	16.1 ± 0.1 ^a
Lysine	37.5 ± 0.2 ^b	28.0 ± 0.6 ^c	37.6 ± 0.7 ^b	42.7 ± 5.3 ^{ab}	46.7 ± 0.8 ^a
Arginine	27.8 ± 0.2 ^b	22.1 ± 0.4 ^c	30.7 ± 1.2 ^b	35.5 ± 2.2 ^a	35.5 ± 1.3 ^a
Threonine	21.2 ± 0.4 ^b	14.7 ± 0.6 ^c	16.9 ± 0.4 ^c	25.8 ± 1.8 ^a	24.4 ± 0.8 ^a
Methionine	17.3 ± 0.3 ^a	8.4 ± 0.1 ^c	11.0 ± 0.6 ^b	16.6 ± 1.7 ^a	17.5 ± 0.9 ^a
Total EAA	221.1 ± 2.6 ^c	157.5 ± 4.1 ^d	214.6 ± 2.8 ^c	267.1 ± 2.4 ^b	280.4 ± 4.6 ^a
NEAA					
Aspartic acid	35.8 ± 0.6 ^c	38.1 ± 0.8 ^c	39.4 ± 0.7 ^{bc}	54.5 ± 3.6 ^a	44.1 ± 1.3 ^b
Glutamic acid	56.1 ± 0.9 ^c	49.0 ± 1.4 ^d	57.3 ± 2.8 ^c	76.5 ± 4.5 ^a	67.5 ± 1.7 ^b
Serine	22.4 ± 0.5 ^a	16.1 ± 0.4 ^b	15.3 ± 0.5 ^b	23.5 ± 0.6 ^a	22.3 ± 0.9 ^a
Proline	19.6 ± 0.5 ^b	19.7 ± 0.7 ^b	21.1 ± 0.8 ^b	34.2 ± 1.7 ^a	21.7 ± 0.8 ^b
Glycine	18.6 ± 0.4 ^c	17.2 ± 0.4 ^c	24.0 ± 0.9 ^b	38.6 ± 1.8 ^a	26.0 ± 0.8 ^b
Alanine	34.2 ± 0.5 ^b	17.1 ± 0.5 ^d	23.2 ± 0.8 ^c	36.9 ± 1.1 ^a	33.8 ± 0.9 ^b
Tyrosine	20.8 ± 0.8 ^c	15.6 ± 0.7 ^d	13.0 ± 0.6 ^d	30.6 ± 2.6 ^a	26.4 ± 0.9 ^b
Cysteine	2.7 ± 0.4 ^d	4.0 ± 0.1 ^c	3.7 ± 0.2 ^{cd}	9.6 ± 0.6 ^a	6.1 ± 0.7 ^b
Total NEAA	210.2 ± 2.3 ^c	176.8 ± 1.6 ^c	197.0 ± 2.4 ^d	304.4 ± 2.7 ^a	247.9 ± 2.9 ^b
Total AA	431.2 ± 6.8 ^c	334.3 ± 9.1 ^e	411.6 ± 4.5 ^d	571.5 ± 7.3 ^a	528.3 ± 5.7 ^b
EAA/NEAA	1.05	0.48	1.09	0.52	1.13

Results are means ± SD (n = 3). An SD of 0.0 implies an SD of < 0.05. Mean values bearing different superscript letter are significantly different (P<0.05). AA, amino acid; EAA, essential amino acid; NEAA, non essential amino acid.

Supplementary Table 4. Total lipid fatty acid content (μg fatty acid/mg dry mass) of rotifer *Brachionus rotundiformis* enriched with Algamac 3050® (ROT), nauplii of the copepod *Acartia tonsa* fed with the microalgae *Rhodomonas baltica* (COP), *Artemia* metanauplii enriched with Algamac 3050 (ART) and sea bream (*Sparus aurata* L.) yolk sac larvae (YSL) used to feed Atlantic bluefin tuna (*Thunnus thynnus* L.) larvae.

Live prey	ROT (e)	COP	ART	YSL
Fatty acid				
14:0	1.8 \pm 0.2 ^c	5.1 \pm 0.3 ^a	1.4 \pm 0.1 ^c	2.6 \pm 0.2 ^b
16:0	11.8 \pm 0.9 ^c	6.8 \pm 0.3 ^d	13.5 \pm 0.4 ^b	22.3 \pm 0.6 ^a
18:0	3.7 \pm 0.3 ^c	1.6 \pm 0.1 ^d	4.6 \pm 0.2 ^b	5.5 \pm 0.2 ^a
Total saturated ¹	18.6 \pm 0.6 ^b	14.4 \pm 0.8 ^c	20.6 \pm 0.5 ^b	31.0 \pm 1.1 ^a
16:1n-7	1.5 \pm 0.2 ^b	1.7 \pm 0.1 ^b	2.1 \pm 0.2 ^b	5.8 \pm 0.2 ^a
18:1n-9	1.7 \pm 0.3 ^c	2.2 \pm 0.2 ^c	15.5 \pm 0.5 ^b	19.0 \pm 0.7 ^a
18:1n-7	1.3 \pm 0.2 ^c	1.1 \pm 0.1 ^c	5.9 \pm 0.1 ^a	3.3 \pm 0.1 ^b
20:1n-9	0.7 \pm 0.1 ^a	0.2 \pm 0.0 ^c	0.5 \pm 0.1 ^b	0.7 \pm 0.1 ^a
Total monoenes ²	9.8 \pm 0.7 ^c	5.9 \pm 0.3 ^d	24.9 \pm 0.6 ^b	31.4 \pm 1.2 ^a
C16 PUFA	4.9 \pm 0.7 ^a	1.6 \pm 0.1 ^b	0.7 \pm 0.1 ^c	1.0 \pm 0.1 ^c
18:2n-6	12.7 \pm 1.0 ^a	2.6 \pm 0.3 ^d	3.9 \pm 0.2 ^c	8.5 \pm 0.3 ^b
20:4n-6	0.8 \pm 0.1 ^b	0.7 \pm 0.1 ^b	1.6 \pm 0.2 ^a	1.9 \pm 0.2 ^a
22:5n-6	2.3 \pm 0.4 ^a	1.7 \pm 0.2 ^b	2.9 \pm 0.2 ^a	0.3 \pm 0.1 ^c
Total n-6PUFA ³	19.4 \pm 0.9 ^a	7.1 \pm 0.3 ^d	9.0 \pm 0.3 ^c	11.6 \pm 0.5 ^b
18:3n-3	3.0 \pm 0.5 ^b	2.4 \pm 0.2 ^b	20.8 \pm 0.4 ^a	1.1 \pm 0.1 ^c
18:4n-3	0.2 \pm 0.0 ^c	2.5 \pm 0.3 ^a	2.6 \pm 0.1 ^a	0.7 \pm 0.1 ^b
20:4n-3	0.8 \pm 0.1 ^a	0.3 \pm 0.1 ^b	0.6 \pm 0.1 ^a	0.7 \pm 0.1 ^a
20:5n-3	3.9 \pm 0.6 ^b	2.3 \pm 0.2 ^c	3.7 \pm 0.3 ^b	8.6 \pm 0.4 ^a
22:5n-3	2.5 \pm 0.4 ^b	0.1 \pm 0.0 ^c	0.2 \pm 0.0 ^c	3.2 \pm 0.1 ^a
22:6n-3	10.9 \pm 0.8 ^b	14.0 \pm 0.4 ^b	7.2 \pm 0.4 ^c	30.0 \pm 1.3 ^a
Total n-3PUFA ⁴	25.7 \pm 1.9 ^c	22.8 \pm 0.5 ^d	35.6 \pm 0.6 ^b	44.4 \pm 2.0 ^a
Total PUFA	50.1 \pm 1.9 ^b	31.5 \pm 1.4 ^d	45.3 \pm 0.7 ^c	57.0 \pm 2.5 ^a
n-3/n-6	1.3 \pm 0.1 ^b	3.2 \pm 0.2 ^a	3.9 \pm 0.1 ^a	3.8 \pm 0.4 ^a
DHA/EPA	2.8 \pm 0.3 ^c	6.1 \pm 0.7 ^a	1.9 \pm 0.1 ^d	3.5 \pm 0.2 ^b

Results are means \pm SD (n = 3). An SD of 0.0 implies an SD of < 0.05. Values bearing different superscript letter are significantly different (P<0.05). ¹, Totals include 15:0, 20:0, 22:0 and 24:0; ², Totals include 16:1n-9, 18:1n-11, 20:1n-7, 22:1 isomers and 24:1; ³, Totals include 18:3n-6, 20:2n-6 and 22:4n-6; ⁴, Totals include 20:3n-3 and 22:3n-3; DHA, docosahexaenoic acid; EPA, eicosapentaenoic acid; PUFA, polyunsaturated fatty acid.