

1 **Title**

2 Expression of long-chain polyunsaturated fatty acid (LC-PUFA) biosynthesis genes
3 during zebrafish *Danio rerio* early embryogenesis

4

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21 **Keywords**

22 Development; Elovl2-like elongase; Elovl5-like elongase; fatty acyl desaturase; LC-
23 PUFA biosynthesis; zebrafish

24

25 **Summary**

26 Long-chain polyunsaturated fatty acids (LC-PUFAs) are essential in important
27 physiological processes, many of which are particularly vital during embryonic
28 development. This study investigated the expression of genes encoding enzymes
29 involved in LC-PUFA biosynthesis, namely fatty acyl desaturase (*Fad*) and *Elovl5*- and
30 *Elovl2*-like elongases, during early embryonic development of zebrafish. Firstly,
31 zebrafish *elovl2* cDNA was isolated and functionally characterised in yeast, showing
32 high specificity towards C20 and C22 PUFAs, compared to C18 substrates. Secondly,
33 spatial-temporal expression for *elovl2* and the previously cloned *fad* and *elovl5* were
34 studied during zebrafish early embryonic development. Temporal expression shows that
35 all three genes are expressed from the beginning of embryogenesis (zygote), suggesting
36 maternal mRNA transfer to the embryo. However, a complete activation of the
37 biosynthetic pathway seems to be delayed until 12 hpf, when noticeable increases of *fad*
38 and *elovl2* transcripts were observed, in parallel with high docosahexaenoic acid levels
39 in the embryo. Spatial expression was studied by whole-mount *in situ* hybridization in
40 24 hpf embryos, showing that *fad* and *elovl2* are highly expressed in the head area
41 where neuronal tissues are developing. Interestingly, *elovl5* shows specific expression
42 in the pronephric ducts, suggesting an as yet unknown role in fatty acid metabolism
43 during zebrafish early embryonic development. The yolk syncytial layer also expressed
44 all three genes, suggesting an important role in remodelling of yolk fatty acids during
45 zebrafish early embryogenesis. Tissue distribution in zebrafish adults demonstrates that
46 the target genes are expressed in all tissues analysed, with liver, intestine and brain
47 showing the highest expression.

48 **Introduction**

49 Long-chain polyunsaturated fatty acids (LC-PUFAs) are essential compounds that play
50 key roles in numerous metabolic and physiological processes ensuring normal cellular
51 function. Some LC-PUFAs, including arachidonic (20:4n-6, ARA) and
52 eicosapentaenoic (20:5n-3, EPA) acids, are precursors of eicosanoids, biologically
53 active compounds that modulate physiological processes including inflammation,
54 reproduction and hemostasis [1]. Increased dietary levels of n-3 LC-PUFAs including
55 EPA and docosahexaenoic acid (DHA, 22:6n-3) have being described as health
56 promoters related to cardiovascular, immune, and inflammatory conditions [2,3].
57 Additionally LC-PUFAs are constituents of cell membrane phospholipids, determining
58 in part fluidity, and activity of membrane proteins and enzymes involved in transport
59 and signal transduction [4]. This is critical in neuronal tissues where a unique degree of
60 fluidity and compressibility of cell membranes is provided by DHA-rich phospholipids
61 that enable rapid conformational changes required for neurotransmission and
62 photoreception [5].

63 The biosynthesis of LC-PUFAs in vertebrates involves consecutive desaturation and
64 elongation reactions that convert the essential fatty acids (EFAs) 18:3n-3 (α -linolenic
65 acid) and 18:2n-6 (linoleic acid) to longer-chain, more unsaturated fatty acids (FAs) of
66 the same series, including EPA, DHA and ARA (Fig. 1, [6,7]). Two types of enzymes
67 are responsible for these conversions, namely fatty acyl desaturases (Fad) and elongases
68 of very long fatty acids (Elovl). The former introduce a double bond in the fatty acyl
69 chain at C6 (Δ 6 Fad) or C5 (Δ 5 Fad) from the carboxyl group. On the other hand, Elovl
70 account for the condensation of activated FAs with malonyl-CoA in the FA elongation
71 pathway. Several members of the Elovl family are involved in PUFA biosynthesis in
72 mammals, those being Elovl5 with substrate specificity for C18 FAs and Elovl2 for C20

73 and C22 [8,9]. Additionally, Elovl4 has been speculated to participate in the elongation
74 steps required for synthesis of DHA in mammalian retina [9].

75 The importance of LC-PUFA in developing organisms is illustrated by their accretion
76 in neuronal tissues during embryogenesis [10-15]. Additionally, deficient production of
77 LC-PUFAs during development can cause neuromuscular defects, cuticle abnormalities,
78 reduced brood size, and altered biological rhythms in *Caenorhabditis elegans* mutants
79 that lack *fat-3*, the gene for $\Delta 6$ desaturase [16]. In mammals, it has been suggested that
80 LC-PUFAs are preferentially delivered from the mother to the fetus by transfer across
81 the placenta since fetal LC-PUFA biosynthetic capacity appears to be limited [12,17]. In
82 oviparous organisms such as avians, FAs present in yolk in the form of triacylglycerol
83 or phospholipid molecules are absorbed into the yolk sac membrane for delivery into
84 the embryonic circulation and utilisation for energy, membrane biogenesis, and fat
85 deposition [18]. Amounts of LC-PUFAs deposited by the hen are insufficient to fulfil
86 the requirements of the embryo, and therefore biosynthesis of LC-PUFA by the chicken
87 embryo is, contrary to human fetus, very active in order to compensate such a
88 deficiency [19,20].

89 In fish, studies have demonstrated that supply of LC-PUFAs to embryos is greatly
90 influenced by the diet of broodstock [21,22], and that suboptimal levels of LC-PUFA
91 delivered to larvae may compromise ability to capture prey in herring (*Clupea*
92 *harengus*) [23], delay response to visual stimuli in gilthead sea bream (*Sparus aurata*)
93 [24], and impair schooling behaviour in yellowtail (*Seriola quiqueradiata*) [25,26] and
94 Pacific threadfin (*Polydactylus sexfilis*) [27]. Despite the known importance of LC-
95 PUFA supply during embryonic development and their proven selective accumulation
96 in certain lipid classes [28], little is known about the capability of fish embryos for
97 endogenous biosynthesis to supplement preformed LC-PUFA present in the yolk.

98 Significant progress has been made in characterising the desaturases and elongases
99 involved in LC-PUFA synthesis in fish including freshwater [29-33] and marine species
100 [34-38]. Zebrafish (*Danio rerio*), a popular model organism in vertebrate developmental
101 biology, has recently been used to study aspects of lipid metabolism [39-42]. Two
102 enzymes involved in LC-PUFA biosynthesis have been characterised in zebrafish, a Fad
103 with dual $\Delta 5/\Delta 6$ activity unique among vertebrates [43], and an elongase with high
104 specificity towards C18 and, to a lesser extent, C20 PUFA [30], similar to elongases
105 found in several other fish species [31-32]. Recently, a cDNA for a second elongase
106 was isolated from salmon and shown to have high specificity towards C20 and C22
107 PUFA [33].

108 The present study aimed to investigate the expression of Fad and Elovl enzymes
109 involved in LC-PUFA biosynthesis during early development of zebrafish. Firstly, we
110 isolated and functionally characterised a second zebrafish elongase cDNA important in
111 the biosynthesis of DHA. Secondly, the spatial-temporal expression pattern of the newly
112 cloned elongase, together with the previously isolated Fad [43] and elongase [30], was
113 investigated during zebrafish embryogenesis. Expression of these three enzymes enable
114 zebrafish to synthesise all LC-PUFA from C18 EFA, and therefore zebrafish are an
115 excellent model to study early developmental regulation of LC-PUFA synthesis in
116 vertebrates.

117

118 **Materials and methods**

119

120 *Fish maintenance*

121 Adult AB wild-type zebrafish strain were maintained at the facilities of the Instituto de
122 Investigaciones Marinas (IIM-CSIC) as described previously [44]. Zebrafish embryos

123 collected from mating of single broodstock couples were isolated and raised at 28.5°C
124 and staged according to the number of hours post-fertilization (hpf) [43]. For whole-
125 mount *in situ* hybridization analyses, dechorionated embryos were fixed overnight at 4
126 °C in 4 % paraformaldehyde in 1xPBS, washed in PBS, and dehydrated through a
127 methanol series, and stored at -20 °C in 100 % methanol. To inhibit embryo
128 pigmentation, embryo medium was supplemented with 0.003 % 1-phenyl-2-thiourea
129 (PTU, Sigma, Alcobendas, Spain) [44].

130

131 *Zebrafish Elovl2: cloning and functional characterization by heterologous expression in*
132 *Saccharomyces cerevisiae*

133 PCR fragments corresponding to the ORF of the putative Elovl2 elongase
134 (gb|NP_001035452|) were amplified from zebrafish liver cDNA using specific primers
135 containing restriction sites (underlined) – Elovl2VF
136 (CCCAAGCTTAGGATGGAATCATATGAAAAATTGATAAG; *Hind*III) and
137 Elovl2VR (CCGCTCGAGTCACTGTAGCTTCTGTTTGGAG; *Xho*I). PCR was
138 performed using the high fidelity PfuTurbo[®] DNA polymerase (Stratagene, Agilent
139 Technologies, Cheshire, UK), with an initial denaturing step at 95 °C for 2 min,
140 followed by 35 cycles of denaturation at 95°C for 30 s, annealing at 57 °C for 30 s,
141 extension at 72 °C for 1 min 10 s, followed by a final extension at 72 °C for 5 min. The
142 DNA fragments were then digested with the corresponding restriction endonucleases
143 (New England BioLabs, Herts, UK) and ligated into a similarly restricted pYES2 yeast
144 expression vector (Invitrogen, Paisley, UK). The purified plasmids (GenElute[™]
145 Plasmid Miniprep Kit, Sigma) containing the putative Elovl2 ORF were then used to
146 transform *S. cerevisiae* competent cells (S.c. EasyComp Transformation Kit,
147 Invitrogen). Transformation and selection of yeast with recombinant pYES2-*elovl2*

148 plasmids, yeast culture and FA analysis was performed as described in detail previously
149 [28,41,44]. Briefly, cultures of recombinant yeast were grown in *S. cerevisiae* minimal
150 medium^{-uracil} supplemented with one of the following FA substrates: stearidonic acid
151 (18:4n-3), γ -linolenic acid (18:3n-6), EPA (20:5n-3), ARA (20:4n-6), docosapentaenoic
152 acid (22:5n-3) or docosatetraenoic acid (22:4n-6). Docosapentaenoic and
153 docosatetraenoic acids (>98-99% pure) were purchased from Cayman Chemical Co.
154 (Ann Arbor, USA) and the remaining FA substrates (>99% pure) and chemicals used to
155 prepare the *S. cerevisiae* minimal medium^{-uracil} were from Sigma Chemical Co. Ltd.
156 (Dorset, UK). FAs were added to the yeast cultures at final concentrations of 0.5 (C18),
157 0.75 (C20) and 1.0 (C22) mM. After 2-days, yeast were harvested and washed, and lipid
158 extracted by homogenization in chloroform/methanol (2:1, v/v) containing 0.01% BHT
159 as antioxidant. FA methyl esters were prepared, extracted, purified, and analysed by GC
160 in order to calculate the proportion of substrate FA converted to elongated FA product
161 as [product area/(product area +substrate area)] x 100. Identities of FA peaks were
162 based on GC retention times and confirmed by GC-MS as described previously [30,43].

163

164 *Sequence and phylogenetic analysis of Elovl2*

165 The amino acid (AA) sequence deduced from the zebrafish Elovl2 cDNA
166 (gb|NP_001035452|) was compared with human (gb|NP_060240|), mouse
167 (gb|NP_062296|) and rat (gb|NP_001102588|) ELOVL2s, amphibian *Xenopus laevis*
168 (gb|NP_001087564|) and *X. tropicalis* (gb|NP_001016159|) Elovl2s, bird *Taenopygia*
169 *guttata* (gb|XP_002186815.1|) and *Gallus gallus* (gb|XP_418947|) predicted Elovl2-like
170 proteins, and salmon Elovl2 (gb|FJ237532|) using the EMBOSS Pairwise Alignment
171 Algorithms tool (<http://www.ebi.ac.uk/Tools/emboss/align/>). A phylogenetic tree was
172 constructed on the basis of the AA sequence alignments between the putative zebrafish

173 Elov12, Elov12 orthologs and Elvol5 proteins, and using the Neighbour Joining method
174 [47]. Confidence in the resulting phylogenetic tree branch topology was measured by
175 bootstrapping through 1000 iterations.

176

177 *Temporal expression of fad, elov15, elov12 during zebrafish ontogeny*

178 To study the expression of the target genes during the embryonic development of
179 zebrafish, total RNA was extracted from pools of 20-30 embryos collected at 0, 3, 6, 9,
180 12, 14, 24, 48, and 72 hpf using Tri Reagent (Sigma) according to manufacturer's
181 protocol. Five μ g of total RNA was reverse transcribed into cDNA using M-MLV
182 reverse transcriptase first strand cDNA synthesis kit (Promega, Madison, USA).
183 Qualitative expression of *fad*, *elov15* and *elov12* transcripts during embryonic
184 development was determined by reverse transcriptase PCR (RT-PCR) on cDNA
185 samples, with an initial denaturing step at 95 °C for 2 min, followed by 35 cycles of
186 denaturation at 95 °C for 30 s, annealing at 60 °C for 30 s, and extension at 72 °C for 1
187 min 40 s, followed by a final extension at 72 °C for 5 min. Expression of β -actin was
188 also determined as reference gene [48]. Primers used for RT-PCR on embryos cDNA
189 samples are shown in Table 1.

190

191 *Spatial expression of fad, elov15, elov12, whole-mount in situ hybridization*

192 To examine the spatial expression of zebrafish *fad*, *elov15* and *elov12*, whole-mount *in*
193 *situ* hybridization (WISH) was performed on 24 hpf zebrafish embryos using
194 digoxigenin (DIG)-labelled antisense riboprobes as previously described [49].
195 Antisense riboprobes were made from linearised full length *Danio rerio fad*, *elov15* and
196 *elov12* cDNAs.

197

198 *Tissue distribution of fad, elovl5 and elovl2 mRNA transcripts in zebrafish adults*

199 Expression of the target genes was also measured in adult tissues by quantitative real-
200 time PCR (qPCR). Total RNA from eye, gill, liver, brain, ovary, testis, kidney, muscle,
201 intestine and adipose tissue was extracted as described above, and 1 µg of total RNA
202 reverse transcribed into cDNA (M-MLV reverse transcriptase, Promega). The qPCR
203 was performed using primers shown in Table 1. Copy numbers of target genes were
204 normalised with copy number of the reference gene *18s* rRNA [48]. PCR amplicons of
205 each gene were cloned into pBluescript II KS (Stratagene) that was then linearised,
206 quantified spectrophotometrically (NanoDrop ND-1000, Thermo Scientific,
207 Wilmington, USA), and serial-diluted to generate a standard curve of known copy
208 numbers. The qPCR amplifications were carried out in triplicate using a Quantica
209 machine (Techne, Cambridge, UK) in a final volume of 20 µl containing 5 µl diluted
210 (1/10) cDNA, 0.5 µM of each primer and 10 µl Absolute™ QPCR SYBR® Green mix
211 (ABgene, Epsom, UK). Amplifications were carried out with a systematic negative
212 control (NTC - no template control, containing no cDNA). The qPCR profiles contained
213 an initial activation step at 95 °C for 15 min, followed by 40 cycles: 15 s at 95 °C, 15 s
214 at the specific primer pair annealing T_m (Table 1) and 10-15 s at 72 °C. After the
215 amplification phase, a dissociation curve of 0.5 °C increments from 75 °C to 90 °C was
216 performed, enabling confirmation of the amplification of a single product in each
217 reaction. The qPCR product sizes were checked by agarose gel electrophoresis and their
218 identity confirmed by sequencing. No primer-dimer formation occurred in the NTC. All
219 reactions were carried out in triplicate and a linear standard curve was drawn, and
220 absolute copy number of the targeted gene in each sample was calculated.

221

222 *Fatty acid analyses of zebrafish embryos*

223 In order to monitor the FA changes during embryogenesis, pools of 150-200 embryos
224 were sampled at different stages (0, 9, 24, 48 and 72 hpf) and total lipid extracted, FA
225 methyl esters prepared and analysed as described above.

226

227 *Statistics*

228 For tissue expression profiles, results expressed as mean normalised values (\pm SE)
229 corresponding to the ratio between the copy numbers of *fad*, *elovl5* and *elovl2*
230 transcripts and the copy numbers of the reference gene, *18s* rRNA. A one-way analysis
231 of variance (ANOVA) followed by Tukey HSD test ($P < 0.05$) was performed to
232 compare the expression level among tissue samples (SPSS, Chicago, USA).

233

234 **Results**

235

236 *Zebrafish elongase (Elov12) sequence and phylogenetics*

237 The new zebrafish elongase ORF encodes a protein of 295 AA, sharing 73.6 % identity
238 in AA sequence to the salmon Elov12, 65.8 - 68.1 % AA identity to mammalian
239 homologues, and 66.9 - 68.4 % identity with predicted Elov12 sequences from
240 amphibians and birds. The phylogenetic tree (Fig. 2) shows that zebrafish Elov12
241 elongase clusters most closely with salmon Elov12, the only Elov12 elongase cloned and
242 characterised in fish so far. The fish Elov12 elongases cluster with the mammalian,
243 amphibian and bird Elov12-like elongases, and more distantly from Elov15-like
244 elongases from mammals and fish.

245

246 *Functional characterisation*

247 The zebrafish putative Elovl2 elongase was functionally characterised by determining
248 the FA profiles of *S. cerevisiae* transformed with pYES2 containing *elovl2* cDNA ORF
249 insert and grown in the presence of potential FA substrates. The FA composition of the
250 wild yeast consists essentially of 16:0, 16:1n-7, 18:0 and 18:1n-9 [43]. Control
251 treatments consisting of yeast transformed with pYES2 vector without elongase insert
252 contained these FA together with whichever exogenous FA was added as substrate (data
253 not shown), this result being consistent with the well established lack of PUFA elongase
254 activity in *S. cerevisiae* [30,32]. Zebrafish Elovl2 shows activity towards FA substrates
255 from 18 to 22 carbons, with the highest specificity on C20 and C22 substrates (Table 2).
256 The traces show the major endogenous FA (16:0, 16:1n-7, 18:0 and 18:1n-9) and
257 additional peaks corresponding to the substrate and elongation products (Fig. 3). Thus
258 exogenously added 18:4n-3 (Fig. 3A) and 18:3n-6 (Fig. 3B) were elongated to their
259 corresponding C20, C22 and C24 elongation products 20:4n-3, 22:4n-3 and 24:4n-3
260 (from 18:4n-3) and 20:3n-6, 22:3n-6 and 24:3n-6 (from 18:3n-6). Total conversion of
261 C18 substrates ranged from 20.1 - 23.0 % (Table 2). Higher elongation rates were
262 observed for C20 substrates 20:5n-3 (78.4 %) and 20:4n-6 (65.3 %), being elongated to
263 C22, C24 and C26 products (Fig. 3C-D). Elovl2 also elongated C22 FA substrates to
264 C24 and C26 elongation products. Thus, yeast transformed with pYES2-*elovl2*
265 converted 22:5n-3 to 24:5n-3 and 26:5n-3 (Fig. 3E), and 22:4n-6 was elongated to
266 24:4n-6 and 26:4n-6 (Fig. 3F). Comparison of peak areas of the endogenous fatty acids
267 in yeast indicates Elovl2 shows some capability to elongate monounsaturated fatty acids
268 such as 16:1n-7 to 18:1n-7 (5.2 - 7.0 %) and 18:1n-9 to 20:1n-9 (1.5 - 3.1 %). No
269 evidence for elongation of saturated FAs was observed with the zebrafish Elovl2.

270

271 *Spatial-temporal expression of fad, elovl5 and elovl2 in zebrafish*

272 Temporal expression of *fad*, *elovl5* and *elovl2* was studied by RT-PCR on cDNA
273 samples obtained from embryos at different developmental stages from 0 to 72 hpf (Fig.
274 4). Results reveal that all three genes are expressed from the zygote stage (0 hpf), with
275 transcripts detected throughout embryonic development. Although comparisons of
276 transcript levels from RT-PCR analyses have to be made cautiously, some temporal
277 patterns can be observed in the expression of *fad*, with a noticeable increasing
278 expression from 12 hpf onwards. Also obvious was the pattern shown by *elovl2*, which
279 showed low expression until 9 hpf, with evident increased expression during 12 to 72
280 hpf. Changes in expression of *elovl5* with development were less obvious, and β -*actin*
281 reference gene expression was constant during development of zebrafish embryos.

282 To examine the spatial expression of zebrafish *fad*, *elovl5* and *elovl2*, WISH was
283 performed on 24 hpf zebrafish embryos (Fig. 5). Zebrafish *fad* (Fig. 5B) and *elovl2*
284 transcripts (Fig. 5F) were widely distributed in the head region and specifically in the
285 yolk syncytial layer (YSL) (Fig. 5B, F insets). Similar to the expression patterns of
286 zebrafish *fad* and *elovl2*, zebrafish *elovl5* was also uniformly expressed in the YSL (Fig.
287 5D inset). However, unlike *fad* and *elovl2*, *elovl5* was specifically expressed in the
288 pronephric ducts of 24 hpf embryos (Fig. 6D). Embryos treated with control sense
289 probes did not show any signal (Fig. 5A, C, E).

290 Adult tissue distribution of *fad*, *elovl5* and *elovl2* mRNA transcripts was analysed by
291 qPCR (Fig. 6). Results indicate that these genes are expressed in all tissues analysed,
292 with significantly higher levels of these transcripts found in liver than any other tissue.
293 Although no significant differences were found, intestine and brain also showed high
294 levels of transcripts, especially *fad* and *elovl2*. Muscle and gill appear to be tissues with
295 very low expression of the three genes. Generally speaking, expression of zebrafish *fad*
296 gene was higher than those of elongase genes.

297

298 *Fatty acid composition of zebrafish embryos*

299 Activity of the enzymes involved in LC-PUFA biosynthesis during zebrafish
300 embryogenesis was estimated by comparing levels of C18 substrates (18:3n-3 and
301 18:2n-6) with levels of all potential desaturation/elongation products (Fig. 7). Total
302 amount of C18 precursors decreased by around 50% over the time-course of
303 embryogenesis, and the levels of products of the biosynthetic pathway showed a steady
304 increase as development proceeded (Fig. 7). Contents of DHA, the most abundant
305 PUFA in zebrafish embryos, initially decreased until 9 hpf, and then increased until the
306 end of embryonic development. The fatty acid profiles (μg of fatty acid per mg of total
307 lipid) of zebrafish embryos at different stages of development are shown in Table 3.

308

309 **Discussion**

310 Our overall objective is to elucidate the molecular mechanisms controlling LC-PUFA
311 synthesis in vertebrates. Using zebrafish as a model species, the specific aim of the
312 present study was to determine the ontogenic changes in expression of genes of the LC-
313 PUFA synthesis pathway during development. In order to do this, we examined all the
314 key genes of LC-PUFA synthesis pathway. Previously, we cloned a Fad cDNA from
315 zebrafish that was unique among vertebrate Fads in showing dual $\Delta 6/\Delta 5$ activity [43].
316 The enzyme product displayed all the fatty acyl desaturation activities required for the
317 synthesis of EPA and DHA [50]. Subsequently, a PUFA elongase cDNA was also
318 isolated from zebrafish [30]. In mammals, *ELOVL2* and *ELOVL5* have been shown to
319 participate in LC-PUFA biosynthesis [8,9,51,52]. Mammalian *ELOVL5* is
320 predominantly involved in the elongation of C18 and C20 PUFA, whereas *ELOVL2* has
321 greatest activity in the elongation of C20 and C22 PUFA and, therefore, appears to be a

322 critical enzyme for the synthesis of C22 and C24 LC-PUFAs [6,8, 51,52]. Functional
323 characterisation showed the first cloned zebrafish PUFA elongase [28] to be similar to
324 elongases found in several other fish species [31,32,38], now all designated as Elovl5
325 [33]. In contrast to mammalian Elovl5s, fish Elovl5s displayed C22 elongation activity,
326 albeit low, and so it was speculated that $\Delta 6/\Delta 5$ Fad and Elovl5 were the only desaturase
327 and elongase necessary for LC-PUFA synthesis in zebrafish [50]. However, whereas
328 sequence similarity searches against the zebrafish draft genome assembly (Zv7)
329 revealed no further Fad genes, a further elongase-like gene was present in chromosome
330 24 that, if expressed, could potentially participate in LC-PUFA production. We now
331 report the cDNA cloning and functional characterisation of this second zebrafish
332 elongase (gb|NP_001035452).

333 The AA sequence of the newly cloned zebrafish elongase shows high identity to the
334 recently cloned salmon elongase cDNA, which has been shown to be an Elovl2
335 orthologue [33], and relatively high identity to mammalian ELOVL2s. Phylogenetic
336 analysis groups the zebrafish elongase into a cluster with greatest similarity to salmon
337 Elovl2 and other Elovl2-like genes from mammals, amphibians and birds, and more
338 distantly from Elovl5 elongases. Functional characterisation of the zebrafish cDNA
339 confirms that the encoded protein elongated C20 and C22 PUFA and so the elongase is
340 designated as an Elovl2. Recombinant yeast containing zebrafish Elovl2 cDNA also
341 produced C26 PUFA from their corresponding C20 and C22 substrates, although these
342 conversions are unlikely to occur *in vivo* because of competition with $\Delta 6$ Fad for
343 intermediate C24 PUFAs [6]. As described for mouse and salmon, zebrafish *elovl2*
344 cDNA encodes an enzyme that also has C18-20 elongase activity [8,33]. This is in
345 contrast to human ELOVL2, which is only active towards C20 and C22 substrates [8].
346 Importantly, the major difference in comparison to zebrafish Elovl5 [30] and other fish

347 Elov15s, is the high activity towards C22 PUFA shown by zebrafish Elov12. Therefore,
348 Elov12 is a key component in the biosynthesis of DHA, where two consecutive
349 elongation steps from 20:5n-3 to 22:5n-3 and 22:5n-3 to 24:5n-3 are required, followed
350 by $\Delta 6$ desaturation and chain-shortening [6,53]. These results prove that zebrafish
351 possess all the enzymatic activities required for LC-PUFA synthesis [6], with $\Delta 6$ and $\Delta 5$
352 desaturation performed by a single protein [43], and elongation of PUFAs ranging from
353 C18 to 22 catalysed by Elov15 [30] and the herein characterised Elov12. The capability
354 of zebrafish for LC-PUFA biosynthesis was previously assessed in isotopic studies with
355 primary hepatocytes showing that the pathway for EPA and DHA synthesis was fully
356 functional [54]. This conclusion is supported by the molecular cloning of the $\Delta 6/\Delta 5$ Fad
357 [43], Elov15 [30], and the newly characterised Elov12.

358 Expression of all Fad and Elov1 activities required for LC-PUFA biosynthesis,
359 presents zebrafish as an excellent model to study relationships between expression of
360 these genes and important developmental events where high demands for LC-PUFA are
361 required, especially the formation of neuronal tissues critical for the viability of the
362 embryo [10,16]. In humans, such high requirements for LC-PUFAs are mostly delivered
363 to the fetus by transfer across the placenta, since fetus LC-PUFA biosynthesis capability
364 has been suggested to be insufficient [17]. Similar to avians, where embryos have been
365 demonstrated to biosynthesise LC-PUFA [55], our results suggest that LC-PUFA
366 biosynthesis occurs in zebrafish embryos, as supported by the presence of *fad*, *elov15*
367 and *elov12* transcripts during embryogenesis, and the dynamic FA composition of
368 embryos denoting endogenous production of LC-PUFA.

369 Temporal expression patterns show that genes of LC-PUFA biosynthesis enzymes in
370 zebrafish are detected at the zygote stage (0 hpf). The only explanation for this is that
371 maternal transfer of the target gene mRNA takes place in zebrafish, since zygotic gene

372 activation is delayed until midblastula transition, which begins at the 512 cell stage at
373 2.75 hpf [45]. This highlights that the maternal role in LC-PUFA supply to fish embryos
374 is not only transfer of preformed LC-PUFA [21,22], but also transfer of mRNA
375 transcripts that can potentially be translated to active proteins. Expression of *fad*, *elovl5*
376 and *elovl2* genes continues to the end of embryogenesis (72 hpf), and so the pathway
377 could be active throughout to assure the high demands of forming tissues such as brain
378 and retina for LC-PUFAs.

379 Beyond maternal mRNA transfer and its potential role in LC-PUFA biosynthesis in
380 early stage embryos, the results raise the question of when the embryo itself begins to
381 activate the pathway. Despite the steady increase in total LC-PUFA content during
382 embryogenesis, DHA initially decreases from 0 to 9 hpf. This could indicate that,
383 although mRNA transcripts of *fad*, *elovl5* and *elovl2* were detected during the early
384 developmental stages (0-9 hpf), the biosynthesis pathway is not fully active, at least for
385 producing C22 PUFAs. Supporting this idea is the fact that *elovl2* mRNA transcripts are
386 very low until 9 hpf, possibly limiting biosynthesis of specifically DHA during early
387 embryogenesis [8]. From 9 hpf onwards *de novo* transcription of embryonic genes likely
388 occurs as indicated by increased levels of *fad* and *elovl2* transcripts from 12 hpf. We
389 may speculate that the increase in expression of *fad* and *elovl2* is due to the
390 development of the central nervous system and retina, occurring in zebrafish at
391 gastrula:bud (10.0 - 10.33 hpf) and 5-9 somites (11.66 - 14.0 hpf), respectively [45].
392 The spatial expression of *fad* and *elovl2* in zebrafish embryos supports this hypothesis.

393 Spatial expression patterns of FA metabolism enzymes in zebrafish was first studied
394 by Hsieh and co-workers [56], who determined that stearoyl-CoA desaturase, the
395 enzyme responsible for the synthesis of 18:1n-9 from 18:0, is evenly expressed in all
396 embryo tissues. A more specific expression has now been observed for genes encoding

397 enzymes of the LC-PUFA biosynthesis pathway, with *fad* and *elovl2* genes highly
398 expressed in the head area of zebrafish embryos, probably related to the requirement for
399 ARA and DHA in developing neuronal tissues [10-17]. Interestingly, the Elov15
400 elongase was specifically expressed in the pronephric ducts of 24 hpf embryos.
401 Although Elov15 elongase has been reported to be expressed in kidney of adult fish
402 [33,36,46], there is no obvious explanation for such a specific expression in the
403 pronephric ducts of the embryonic kidney, and further investigations are required to
404 elucidate these findings.

405 The spatial gene expression data also reveals that the yolk syncytial layer (YSL) may
406 also be an important tissue for embryonic LC-PUFA biosynthesis in zebrafish. The
407 YSL, a structure unique to teleosts, forms a boundary layer between the embryo and the
408 yolk mass. Consequently, all nutrients contained in the yolk must pass through the YSL
409 before being utilised by the developing tissues in the embryo [57]. Indeed the presence
410 of proteolytic enzyme activities in teleost YSL has been reported previously, in
411 agreement with an active role in resorption of yolk lipoproteins [58,59]. Our results
412 show that YSL is likely also to be active in remodelling PUFA during zebrafish
413 embryogenesis. Thus, in addition to hydrolysis of the abundant lipids contained in the
414 yolk [60], the YSL may also influence the composition of the hydrolysed and absorbed
415 FA in a number of ways including conversion of C18 FA and alteration of EPA/DHA
416 ratio prior to transfer to the developing embryonic tissues. As aforementioned, retinal
417 membranes are composed by DHA-rich phospholipids [61,62], and therefore LC-PUFA
418 biosynthetic activity could be expected in developing eye. However, no clear expression
419 of *fad*, *elovl5* and *elovl2* genes in retina was detected in the present study. Previously,
420 zebrafish embryo retina/eye tissue was found to express Elov14 elongase [63],
421 speculated to be a photoreceptor-specific component of the LC-PUFA biosynthesis

422 pathway [9]. Recently it was shown that Elov14 was required for the production of C28-
423 C38 very long chain PUFA in retina, brain and sperm [64], and is implicated in the
424 synthesis of very long chain omega-hydroxylated fatty acids present in ceramides of the
425 epidermal permeability barrier in mammals [65].

426 The present study also demonstrates that adult zebrafish expressed $\Delta 6/\Delta 5$ *fad*, *elov15*
427 and *elov12* genes in all tissues analysed. In agreement with previous studies on
428 freshwater fish, our results show that the genes in zebrafish are predominantly
429 expressed in liver, intestine and brain implicating these tissues as the most active in LC-
430 PUFA biosynthesis [33,46]. This is consistent with liver and intestine being the major
431 sites of lipid synthesis and distribution. Furthermore, liver and intestine have been
432 described to be the primary tissues for LC-PUFA synthesis in salmonids [66,67].
433 Comparison of transcript levels indicates that *fad* expression is consistently higher than
434 that of both elongases. This could be related to the fact that zebrafish Fad, having dual
435 $\Delta 6/\Delta 5$ activity, is required for all desaturation steps necessary in LC-PUFA biosynthesis
436 [43].

437 In conclusion the present study demonstrates that zebrafish Elov12 shows substrate
438 specificity towards C20- and C22-PUFA, indicating its important role in synthesis of
439 LC-PUFA, particularly DHA. All three genes, *fad*, *elov15* and *elov12*, are ubiquitously
440 expressed in adult zebrafish tissues with highest expression levels in liver, intestine and
441 brain. Our results demonstrate the presence of *fad*, *elov15* and *elov12* transcripts from the
442 zygote stage indicating that maternal transfer of mRNA occurs in zebrafish. Subsequent
443 increases of *fad* and *elov12* transcript levels however, suggest endogenous embryonic
444 expression is activated at later stages when required for neuronal tissues development.
445 DHA levels during zebrafish embryogenesis and spatial expression of *fad* and *elov12*
446 support this hypothesis. The WISH data also indicated that other tissues such as YSL

447 and the pronephric ducts have roles in LC-PUFA metabolism in early embryogenesis in
448 *D. rerio*. Whereas the role of YSL appears obvious in remodelling of yolk FA, the role
449 of the pronephric ducts is both intriguing and obscure and requires further investigation.

450

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459

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655 **Legends to Figures**

656 Fig. 1. Biosynthesis pathways of long-chain polyunsaturated fatty acids from C18
657 precursors, 18:3n-3 and 18:2n-6 [6].

658

659 Fig. 2. Phylogenetic tree comparing the putative zebrafish Elovl2, Elovl2 orthologs and
660 Elvol5 proteins. The tree was constructed using the Neighbour Joining method [47]
661 using MEGA4. The horizontal branch length is proportional to amino acid substitution
662 rate per site. The numbers represent the frequencies (%) with which the tree topology
663 presented was replicated after 1000 iterations.

664 *Predicted proteins (GenBank).

665

666 Fig. 3. Functional characterisation of the zebrafish putative elongase Elovl2 in
667 transgenic yeast (*Saccharomyces cerevisiae*) grown in the presence of fatty acid
668 substrates 18:4n-3 (A), 18:3n-6 (B), 20:5n-3 (C), 20:4n-6 (D), 22:5n-3 (E) and 22:4n-6
669 (F). Fatty acids were extracted from yeast transformed with pYES2 vector containing
670 the ORF of the putative elongase cDNA as an insert. Peaks 1-4 represent the main
671 endogenous FAs of *S. cerevisiae*, namely 16:0 (1), 16:1n-7 (2), 18:0 (3) and 18:1n-9 (4).
672 Substrates (“*”) and their corresponding elongated products are indicated accordingly in
673 panels A-F. Vertical axis, FID response; horizontal axis, retention time.

674

675 Fig. 4. RT-PCR analyses of the temporal expression patterns of *fad*, *elvol5*, and *elovl2*
676 during zebrafish *Danio rerio* embryogenesis (0 to 72 hpf at 28.5 °C). Expression of the
677 housekeeping gene β -actin is also shown. hpf, hours post-fertilization; NTC, no
678 template control.

679

680 Fig. 5. Whole mount *in situ* hybridization showing the expression of *fad* (A, B), *elovl5*
681 (C, D), and *elovl2* (E, F) in 24 hpf embryos. Embryos were hybridised with either sense
682 (A, C, D) or antisense probes (B, D, F). Strong signal was observed in the head region
683 and yolk syncytial layer (B, F inset) of 24-hpf embryos when antisense *fad* and *elovl2*
684 probes were used (A), but no signal was observed for sense probe (E). Similar results
685 were observed for *elovl5* (C, D), however, its expression was specifically localised in
686 the pronephric ducts (D) and the yolk syncytial layer (D inset). Lateral views, dorsal
687 upward, anterior to the left (A-F). YSL, yolk syncytial layer; PD, pronephric ducts; H,
688 head; e, eye. Scale bars: 100 μ m.

689

690 Fig. 6. Tissue distribution of the *fad*, *elovl5* and *elovl2* transcripts (mRNA) in zebrafish
691 adults. Absolute copy numbers were quantified for each transcript and were normalised
692 by absolute levels of 18s RNA. Results are means \pm S.E. (n = 3). L, liver; I, intestine; B,
693 brain; E, eye; K, kidney; A, adipose; M, muscle; O, ovary; T, testis; G, gill. * P < 0.05
694 as determined by one-way ANOVA and Tukey's test.

695

696 Fig. 7. Fatty acid contents during zebrafish embryogenesis. Contents (μ g of fatty acid
697 per mg of total lipid) of substrates (sum of 18:3n-3 and 18:2n-6) and potential products
698 (sum of 18:4n-3, 18:3n-6, 20:3n-3, 20:4n-3, 20:2n-6, 20:3n-6, 20:5n-3, 20:4n-6, 22:4n-
699 3, 22:5n-3, 22:6n-3, 22:4n-6, 24:5n-3, 24:4n-6, 24:6n-3 and 24:5n-6) of long-chain
700 polyunsaturated fatty acid biosynthesis enzymes Fad, Elov15 and Elov12. Levels of
701 docosahexaenoic acid (DHA; 22:6n-3) are also shown.

Table1

Table 1. Sequence and annealing temperature (T_m) of the primer pairs used, size of the fragment produced and accession number of the sequence used as reference for primer design, for Elov12 ORF cloning, reverse transcriptase PCR (RT-PCR) performed in embryo samples, and quantitative real time PCR (qPCR) determinations of transcripts in adult tissues.

Aim	Transcript	Primer	Primer sequence	Fragment	T _m	Accession No ¹ .
<i>ORF cloning</i>	<i>elov12</i>	Elov12VF Elov12VR	5'-CCC <u>AAGCTT</u> AGGATGGAATCATATGAAAAAATTGATAAG-3' 5'-CCGCTCGAGTCACTGTAGCTTCTGTTTGGAG-3'	184 bp	60°C	NM_001040362
<i>RT-PCR</i>	<i>fad</i>	FadF1	5'-AGGAGGTGCAGAAACACACC-3'	1264 bp	60°C	AF309556
		FadR1	5'-CTCGCCAGATTTCTCCAAAG -3'			
	<i>elov15</i>	Elov15F1	5'-CTCAGGGTCACAGGATGGTT-3'	768 bp	60°C	NM_200453
		Elov15R1	5'-CTCCATTAGTGTGGCCGTTT-3'			
	<i>elov12</i>	Elov12F1 Elov12R1	5'-AAAGAGATACCCGCGTGAGA-3' 5'-TTGGAGTTGGCTCCGTTTAG-3'	810 bp	60°C	NM_001040362
<i>β-actin</i>	β-ActinF1 β-ActinR1	5'-CTCTTCCAGCCTTCCTTCCT-3' 5'-CACCGATCCAGACGGAGTAT-3'	246 bp	60°C	NM_131031	
	<i>qPCR</i>	<i>fad</i>	FadF2	5'-CATCACGCTAAACCCAACA-3'	158 bp	60°C
FadR2			5'-GGGAGGACCAATGAAGAAGA-3'			
<i>elov15</i>		Elov15F2	5'-TGGATGGGACCGAAATACAT-3'	173 bp	60°C	NM_200453
		Elov15R2	5'-GTCTCCTCCACTGTGGGTGT-3'			
<i>elov12</i>		Elov12F2 Elov12R2	5'-CACTGGACGAAGTTGGTGAA-3' 5'-GTTGAGGACACACCACCAGA-3'	184 bp	60°C	NM_001040362
<i>18s</i>	18sF1 18sR1	5'-CCGCTATTAAGGGTGTGGA-3' 5'-GGCGAGGGTTCTGCATAATA-3'	134 bp	62°C	NM_173234	

¹ GenBank (<http://www.ncbi.nlm.nih.gov/>)

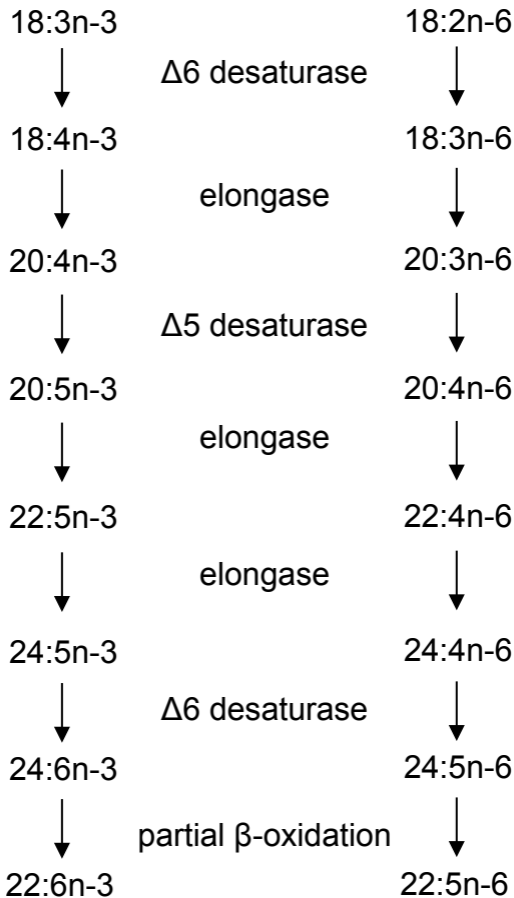
Table 2. Functional characterisation of the newly characterised Elovl2 elongase. Results are expressed as a percentage of total fatty acid (FA) substrate converted to elongated product. Percentage of stepwise conversion into intermediary products of the elongation pathway is also shown.

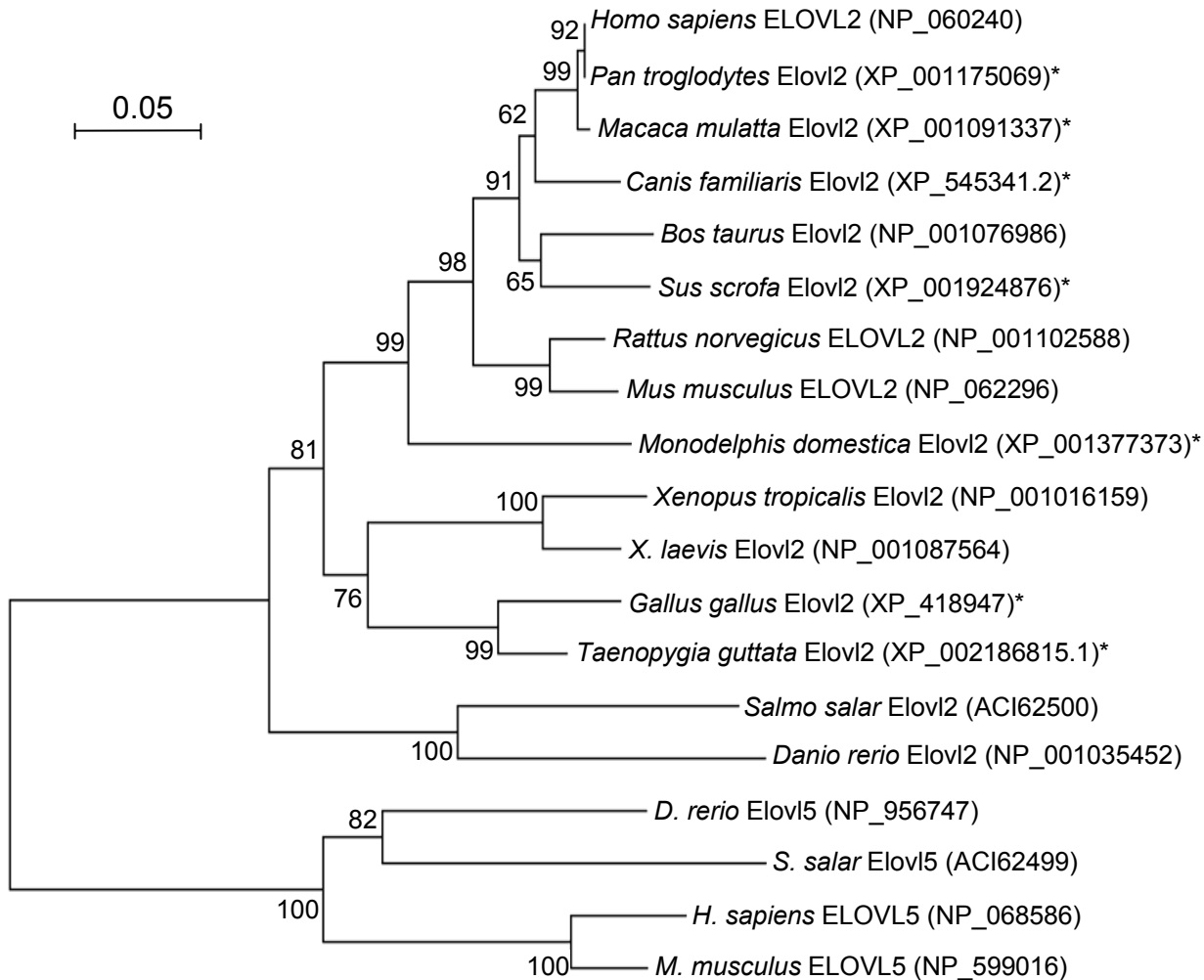
FA Substrate	Product	% Conversion	Activity
18:4n-3	20:4n-3	6.0	C18→20
	22:4n-3	7.0	C20→22
	24:4n-3	10.0	C22→24
	26:4n-3	0.0	C24→26
		Total: 23.0	
18:3n-6	20:3n-6	7.1	C18→20
	22:3n-6	4.2	C20→22
	24:3n-6	8.8	C22→24
	26:3n-6	0.0	C24→26
		Total: 20.1	
20:5n-3	22:5n-3	7.7	C20→22
	24:5n-3	63.1	C22→24
	26:5n-3	7.6	C24→26
		Total: 78.4	
20:4n-6	22:4n-6	3.9	C20→22
	24:4n-6	52.2	C22→24
	26:4n-6	9.2	C24→26
		Total: 65.3	
22:5n-3	24:5n-3	43.2	C22→24
	26:5n-3	11.0	C24→26
		Total: 54.2	
22:4n-6	24:4n-6	34.1	C22→24
	26:4n-6	9.3	C24→26
		Total: 43.4	

Table 3. Fatty acid composition of zebrafish embryos at different stages of development. Results are expressed in μg of fatty acid per mg of total lipid.

<i>Fatty acid</i>	<i>0 hpf</i>	<i>9 hpf</i>	<i>24 hpf</i>	<i>48 hpf</i>	<i>72 hpf</i>
14:0	2.3	5.1	5.1	4.4	3.2
15:0	1.2	2.0	2.1	1.4	1.7
16:0	141.2	130.8	120.7	121.4	122.8
18:0	53.6	47.3	42.5	41.4	42.2
20:0	0.0	0.5	0.6	1.0	1.2
Total saturated	198.4	185.6	171.1	169.6	171.0
16:1n-9	3.5	3.4	3.4	3.3	3.7
16:1n-7	8.2	18.6	18.0	14.1	12.2
18:1n-9	87.6	93.1	85.5	80.1	81.8
18:1n-7	18.9	24.8	24.3	21.6	19.6
20:1 ¹	3.0	5.3	6.8	4.3	3.2
22:1 ²	0.0	3.1	3.8	0.0	0.0
24:1n-9	0.0	0.4	0.5	0.3	0.3
Total monounsaturated	121.3	148.7	142.4	123.8	120.8
18:2n-6	41.5	21.8	22.7	23.5	17.6
18:3n-6	0.0	0.7	0.8	0.8	0.6
20:2n-6	2.5	1.5	1.8	1.7	1.9
20:3n-6	4.9	3.5	3.9	4.1	4.8
20:4n-6	11.7	14.2	15.6	16.3	16.3
22:4n-6	1.5	0.9	0.9	1.0	1.3
22:5n-6	1.0	4.1	4.2	5.1	5.1
Total n-6 PUFA	63.0	46.7	50.0	52.4	47.6
18:3n-3	3.2	4.5	3.4	3.1	2.6
18:4n-3	0.0	0.8	1.0	0.0	0.6
20:3n-3	0.0	1.0	0.9	0.8	0.9
20:4n-3	1.1	3.1	2.5	2.5	2.1
20:5n-3	19.3	44.4	43.4	41.3	42.9
22:5n-3	5.9	13.0	15.9	11.9	13.3
22:6n-3	91.5	63.1	74.4	86.1	89.7
Total n-3 PUFA	121.0	129.9	141.4	145.7	152.1
C16 PUFA	0.0	4.2	3.0	3.8	3.6
Total PUFA	184.1	180.8	194.4	201.9	203.3

¹ predominantly n-9 isomer; ² predominantly n-11 isomer;
PUFA, polyunsaturated fatty acid; hpf, hours post-fertilization





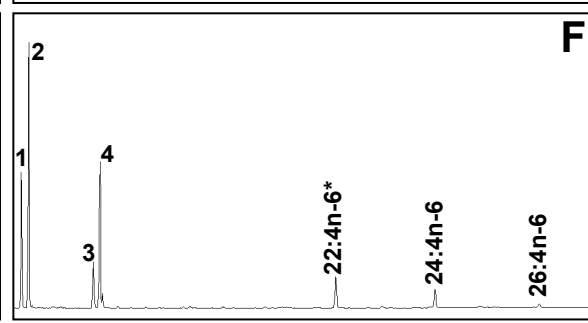
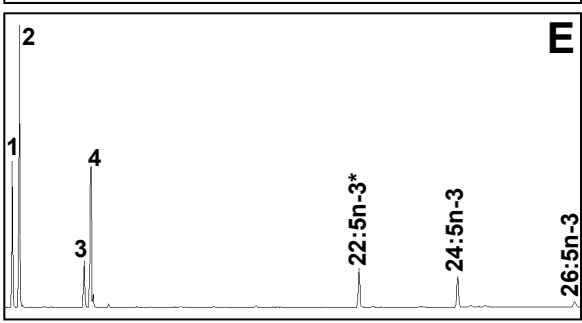
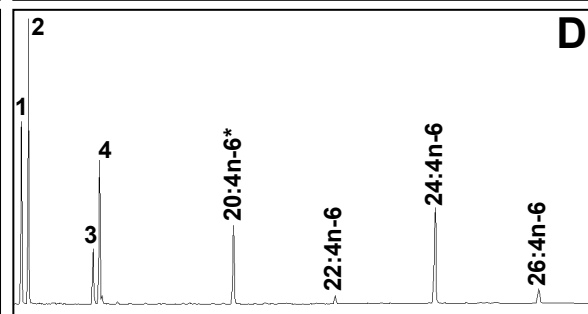
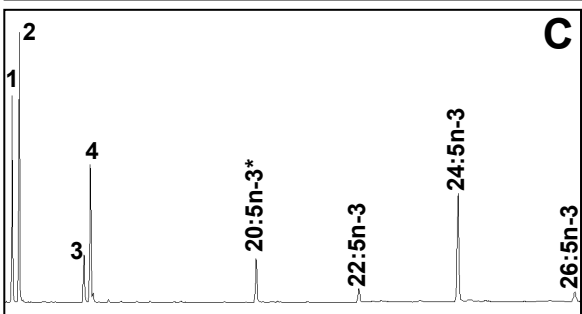
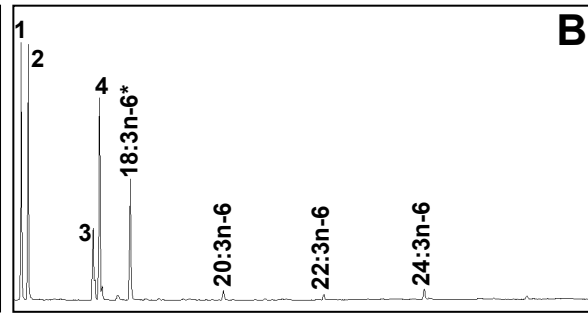
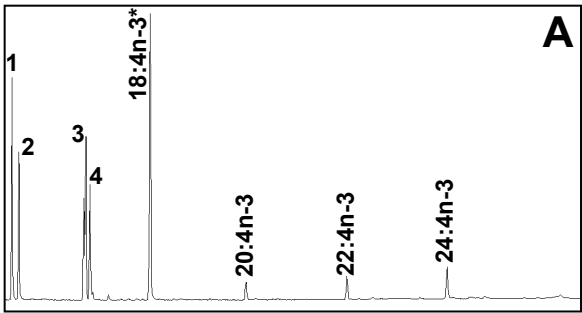


Figure 5 (revised)
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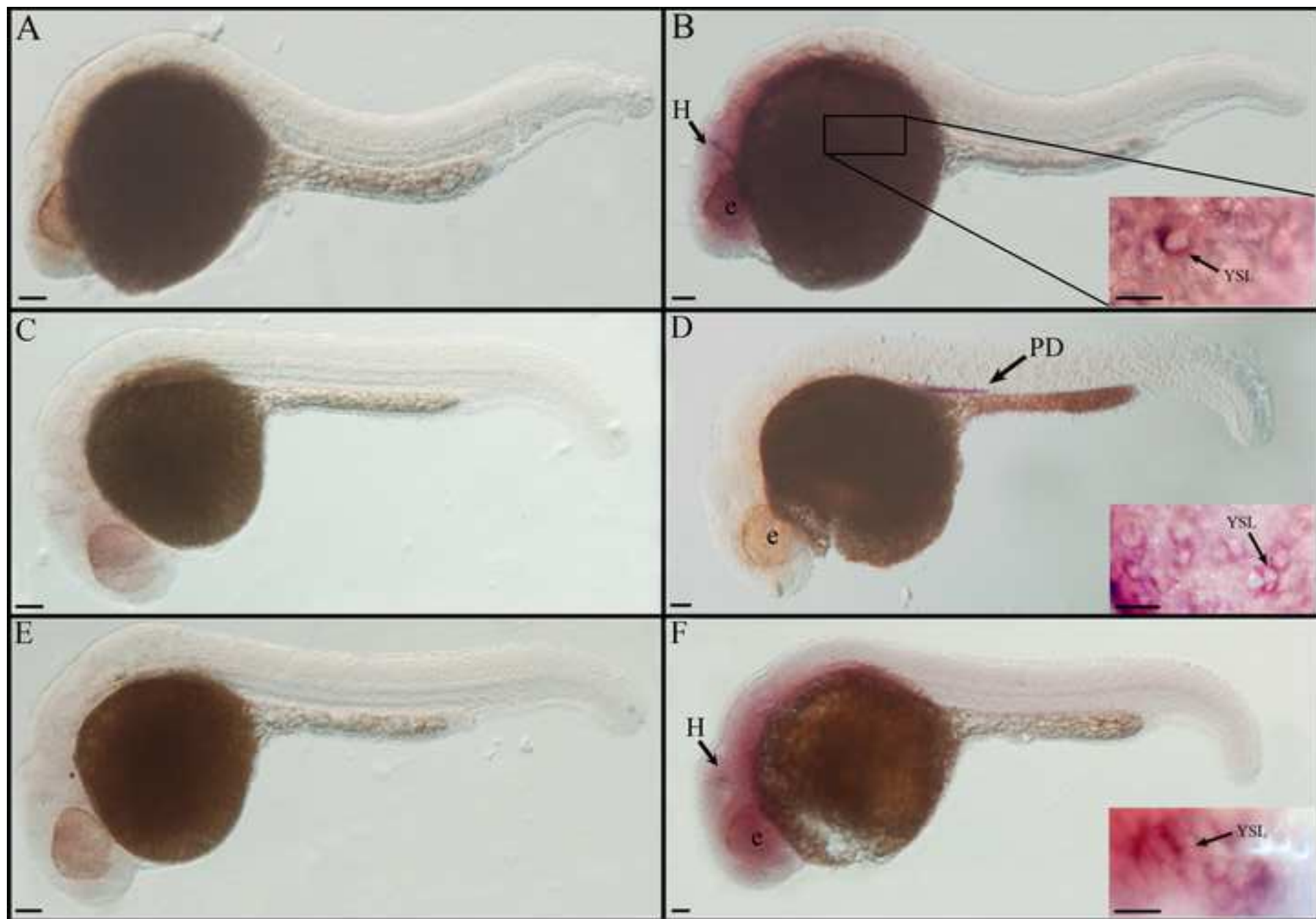


Figure6

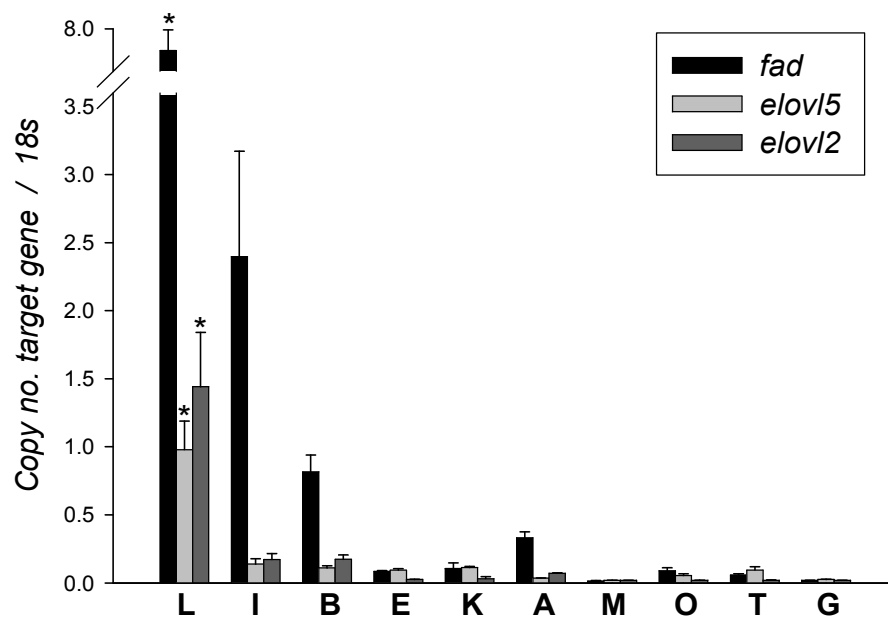


Figure7

