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25 Summary

26 Polyunsaturated fatty acids (PUFA) have been identified as key nutrients for the 27 common octopus (Octopus vulgaris), particularly for its early life-cycle stages 28 (paralarvae). Our overarching aim is to establish the essential fatty acid (FA) 29 requirements for octopus paralarvae through determination of the enzymes of 30 endogenous PUFA biosynthetic pathways. We here report on the molecular cloning and 31 functional characterisation of a cDNA encoding a putative elongase of very long-chain 32 fatty acids (Elovl), a critical enzyme that mediate the elongation of FA including PUFA. 33 Our results suggested that the octopus Elovl is phylogenetically related to Elov15 and 34 Elovl2, two elongases with demonstrated roles in PUFA biosynthesis in vertebrates. 35 Further evidence supporting a role of the octopus Elovl in PUFA biosynthesis was 36 provided through functional characterisation of its activity in yeast. It was confirmed 37 that expression of the octopus Elovl conferred on yeast the ability to elongate some C18 38 and C20 PUFA, while C22 PUFA substrates remained unmodified. The substrate 39 specificities exhibited by the octopus elongase were consistent with those of vertebrate 40 Elov15. Interestingly, the octopus Elov1 elongated n-6 PUFA substrates more efficiently 41 than their analogous n-3 substrates, suggesting that n-6 PUFA may have particular 42 biological significance in O. vulgaris. Finally, we investigated the potential role of the 43 newly cloned Elovl in the biosynthesis of non-methylene-interrupted FA, compounds 44 typically found in marine invertebrates and confirmed to be also present in the common 45 octopus.

46 Introduction

47 Cephalopods have emerged as prime candidates for diversifying aquaculture. Among the species studied, the common octopus (Octopus vulgaris, Cuvier, 1797) has received 48 49 special attention and relevant aspects for its culture such as husbandry (Iglesias et al., 50 2006; Estefanell et al., 2012), behaviour (Di Cristo et al., 2005; Valverde and García, 51 2005), reproduction (Otero et al., 2007; Wodinsky, 2008; Estefanell et al., 2010), 52 pathologies (Castellanos-Martínez and Gestal, 2011) and nutrition (Villanueva, 1994, 53 Navarro and Villanueva, 2000, 2003; Villanueva et al., 2004, 2009; Ouintana, 2006; 54 Villanueva and Bustamante, 2006; Seixas et al., 2010; Estefanell et al., 2011; Fuentes et 55 al., 2011; Viciano et al., 2011) have been studied. Despite considerable effort, the 56 production of the common octopus in captivity is limited to on-growing wild-captured 57 specimens in floating cages (Iglesias et al., 2007), as the octopus life cycle has not yet 58 been closed. While limited success in the production of juvenile octopuses has been 59 achieved (Villanueva, 1995; Iglesias et al., 2002, 2004; Lenzi et al., 2009), the massive 60 mortalities occurring during early life-cycle stages (paralarvae) have become an, as yet, 61 unresolved zootechnical issue that requires further investigation.

62 Polyunsaturated fatty acids (PUFA) have been previously suggested to be critical 63 dietary components for octopus paralarvae (Navarro and Villanueva, 2003). We have 64 recently initiated a series of studies to establish the essential fatty acid (FA) 65 requirements for octopus paralarvae, so that balanced diets matching their endogenous 66 biosynthetic capability can be formulated and thus promote paralarval survival and 67 development. Due to the obvious difficulties in conducting feeding trials with octopus 68 paralarvae, our approach aims to characterise the gene products encoding enzymes 69 involved in the PUFA biosynthetic pathway, which themselves dictate the ability of 70 species to endogenously produce long-chain PUFA (LC-PUFA) (Bell and Tocher,71 2009).

72 Previously, we reported the molecular cloning and functional characterisation of a 73 fatty acyl desaturase (Fad) from O. vulgaris (Monroig et al., 2012a). The substrate 74 specificity of the octopus Fad revealed that this enzyme was a Δ 5-like Fad and thus we 75 provided for first time molecular evidence of such an enzymatic activity in molluscs 76 (Monroig et al., 2012a). Interestingly, the $\Delta 5$ Fad enables the common octopus to 77 endogenously convert 20:4n-3 and 20:3n-6 into eicosapentaenoic acid (EPA, 20:5n-3) 78 and arachidonic acid (ARA, 20:4n-6), respectively, the latter regarded as critical LC-79 PUFA in a variety of physiological processes ensuring normal cellular function (Funk, 80 2001). Rather than a role in the biosynthesis of EPA, we hypothesised that $\Delta 5$ Fad 81 activity may actually be retained in the octopus for the endogenous biosynthesis of 82 ARA, as high contents of ARA encountered in adult octopus tissues were unlikely to be 83 exclusively from dietary origin. In addition to the potential participation of the octopus 84 $\Delta 5$ Fad in ARA biosynthesis, the common octopus $\Delta 5$ Fad might also have a role in the 85 biosynthesis of non-methylene interrupted (NMI) FA, compounds with unusual 86 unsaturation features that have been found in a variety of marine invertebrates 87 (Barnathan, 2009; Kornprobst and Barnathan, 2010).

The biosynthesis of PUFA including NMI FA in marine molluscs has been investigated previously (De Moreno et al., 1976; Waldock and Holland, 1984; Zhukova, 1986, 1991, 2007). Although the biosynthetic ability of molluscs was species-specific (Waldock and Holland, 1984), it has been shown that some molluscs have active PUFA biosynthetic pathways and, in addition to the above mentioned $\Delta 5$ desaturase, active FA elongation systems also appear to be present. Using radioactive FA, it was reported that the clam *Mesoderma mactroides* could elongate both 18:3n-3 and 18:2n-6 (De Moreno

95 et al., 1976). Later, Waldock and Holland (1984) demonstrated that the Pacific oyster 96 Crassostrea gigas had the ability to produce 20:5n-3 and 22:6n-3. Investigations in 97 other bivalves like Scapharca broughtoni, Callista brevisiphonata and Mytilus edulis 98 (Zhukova, 1986, 1991) demonstrated that the biosynthesis of the NMI dienes $\Delta 7,13$ 99 22:2 and $\Delta 7,15$ 22:2 was achieved by elongation from $\Delta 5,11$ 20:2 and $\Delta 5,13$ 20:2, 100 respectively. In addition to biochemical assays with radiotracers, indirect evidence of 101 FA elongase activity in molluscs was provided analytically (Joseph, 1982). For 102 instance, the unusual NMI FA $\Delta 5.9.15$ 24:3 and $\Delta 5.9.17$ 24:3 found in the limpets 103 Cellana grata and Collisella dorsuosa were suggested to derive from the typical NMI 104 dienes $\Delta 7,13$ 22:2 and $\Delta 7,15$ 22:2, respectively, by chain elongation and subsequent $\Delta 5$ 105 desaturation (Kawashima, 2005).

106 The elongases of very long-chain fatty acids (Elovl), a protein family with seven 107 distinct members (Elovl 1-7) in vertebrates, account for the condensation of 2C into 108 activated preexisting fatty acyl chains (Jakobsson et al., 2006). Investigation of FA 109 biosynthetic pathways has allowed the molecular and functional characterisation of a 110 number of genes encoding Elovl enzymes from vertebrates (see reviews by Jakobsson et 111 al., 2006; Guillou et al., 2010; Monroig et al., 2011a). In contrast, studies of elongase-112 encoding genes from non-vertebrate organisms are scarce, with only few examples such 113 as elongases from the nematode *Caenorhabditis elegans* (Beaudoin et al., 2000) and the 114 marine protist Thraustochytrium sp. (Heinz et al., 2001; Jiang et al., 2008), and no 115 elongases from molluscs have been reported.

In order to expand our knowledge of EFA requirements of common octopus, the present study reports the molecular cloning, functional characterisation and tissue distribution of transcripts (mRNA) of a cDNA encoding a putative elongase involved in PUFA biosynthesis. In order to understand a potential role of the newly cloned elongase 120 in the NMI FA biosynthesis in the common octopus, we also analysed the double bond

121 features of NMI FA found in specific tissues of octopus adult specimens.

122 Materials and methods

123 *Tissue samples*

Tissue samples from common octopus were obtained from the dissection of two (male and female) adult individuals (~1.5 kg) captured through artisanal fisheries along the Mediterranean East coast of Spain. The octopusses were cold anesthetised and sacrificed by direct brain puncture and tissues including nerve, nephridium, hepatopancreas, brain, caecum, gill, muscle, heart and gonad were sampled and immediately frozen at -80 °C until further analysis.

130 Elongase cDNA cloning

131 Total RNA was extracted from octopus tissues using TRIzol® (Gibco BRL, Grand 132 Island, NY, USA) reagent following manufacturer's instructions. Subsequently, first 133 strand cDNA was synthesised using a Verso[™] cDNA kit (ABgene, Rockford, IL, USA) 134 primed with random hexamers. In order to amplify the first fragment of the elongase 135 cDNA, the amino acid (aa) sequences of Elov15 proteins from Homo sapiens (NP 068586.1), Rattus norvegicus (NP 599209.1), Bos taurus (NP 001040062.1), 136 137 Danio rerio (NP 956747.1) and Pagrus major (ADQ27303.1) were aligned using 138 BioEdit v5.0.6 (Tom Hall, Department of Microbiology, North Carolina State 139 University, USA). Conserved regions were used for in silico searches of mollusc 140 expressed sequence tags (EST) using NCBI tblastn tool (http://www.ncbi.nlm.nih.gov/). 141 Several EST displaying high homology with Elovl encoding genes were identified from 142 the molluscs *Mytilus galloprovinciallis* (gb|FL495089.1| and gb|FL499406.1|), 143 Euprymna scolopes (gb|DW256301.1|), and Lymnaea stagnalis (gb|FC701557.1|, gb|FC773093.1|, gb|FC770692.1| and gb|FC696214.1|). Additionally, a search of the 144

145 owl limpet Lottia gigantea genome was performed using the zebrafish Elov15 146 (NP 956747.1) sequence with the tblastn tool at http://genome.jgi-147 psf.org/Lotgi1/Lotgi1.home.html. After processing, the mollusc Elovl-like sequences 148 aligned (Bioedit) for the design of the primers **UNIEloF** (5'were 149 TTGTGGTGGTGGTATTACTTCTC-3') and **UNIEloR** (5'-150 GTAATATACTTTTTCCACCA-3') that were used for polymerase chain reaction 151 (PCR) using GoTaq® Colorless Master Mix (Promega, Southampton, UK), and using a 152 mixture of cDNA from gonads, brain, nerve and caecum as template. The PCR 153 consisted of an initial denaturing step at 95 °C for 2 min, followed by 35 cycles of 154 denaturation at 95 °C for 30 s, annealing at 50 °C for 30 s, extension at 72 °C for 1 min, 155 followed by a final extension at 72 °C for 5 min. The PCR fragment was sequenced at 156 the DNA Sequencing Service of the IBMCP-UPV (Valencia, Spain) and gene-specific 157 primers were designed for 5' and 3' rapid amplification of cDNA ends (RACE) PCR 158 (FirstChoice® RLM-RACE kit, Ambion, Applied Biosystems, Warrington, UK) to 159 produce a full-length cDNA. Details of all primers used for RACE PCR are given in 160 Table 1.

161 For 5'RACE PCR, a positive fragment was obtained by two-round PCR. The first 162 round PCR was performed using the adapter-specific 5'RACE OUTER primer and the 163 gene-specific forward primer OVEloR1, with an initial denaturing step at 95 °C for 2 164 min, followed by 32 cycles of denaturation at 95 °C for 30 s, annealing at 58 °C for 30 s, 165 extension at 72 °C for 75 s, followed by a final extension at 72 °C for 5 min (GoTag[®]) 166 Colorless Master Mix, Promega). First round PCR products were used as template for 167 nested PCR with primers 5'RACE INNER and OVEloR2 in a 32-cycle reaction under 168 the same thermal conditions as above. For 3'RACE PCR, a similar two-round approach 169 was followed with first round PCR performed with primers OVEloF1 and 3'RACE OUTER, with an initial denaturating step at 95 °C for 1 min, followed by 32 cycles of denaturation at 95 °C for 30 s, annealing at 56 °C for 30 s, extension at 72 °C for 2 min, followed by a final extension at 72 °C for 5 min (GoTaq[®] Colorless Master Mix, Promega). First round PCR product was then used as template for nested PCR with primers 5'RACE INNER and OVEloF2, with thermal conditions as above. RACE PCR products were cloned into pGEM-T Easy Vector (Promega) and sequenced as above.

176 Sequence and phylogenetic analyses

177 Using ClustalW (Bioedit), the deduced as sequence of the newly cloned O. vulgaris 178 elongase cDNA was aligned with those of a predicted elongase found in the gastropod 179 owl limpet (termed 'L. gigantea Elovl transcript 1', jgi|Lotgi1|224291|), as well as those 180 of protein homologues including the human ELOVL5 (gb|NP 068586|) and ELOVL2 181 (gb|NP 060240|), and the zebrafish Elovl5 (gb|NP 956747|) and Elov₁₂ 182 (gb|NP 001035452|). The aa sequence identity between Elovl-like proteins was 183 compared using the EMBOSS Needle Pairwise Sequence Alignment tool 184 (http://www.ebi.ac.uk/Tools/psa/emboss needle/). Phylogenetic analysis of the aa 185 sequences deduced from the Elovl-like cDNA from common octopus and those from 186 other organisms including several marine invertebrates was performed by constructing a 187 tree using the Neighbour Joining method (Saitou and Nei 1987), with confidence in the 188 resulting tree branch topology measured by bootstrapping through 10000 iterations.

189 Functional characterisation of the octopus elongase by heterologous expression in
190 Saccharomyces cerevisiae

PCR fragments corresponding to the open reading frame (ORF) of the putative elongase were amplified from a mixture of cDNA synthesised from gonads, brain, nerve and caecum RNA extracts, and using the high fidelity Pfu DNA Polymerase (Promega). PCR conditions consisted of an initial denaturing step at 95 °C for 2 min, followed by

195 35 cycles of denaturation at 95°C for 30 s, annealing at 57 °C for 30 s, extension at 72 196 °C for 2 min 15 s, followed by a final extension at 72 °C for 5 min. The primers 197 containing restriction sites (underlined in Table 1) OVEloVF (HindIII) and OVEloVR 198 (SacI) were used for PCR, and the DNA fragments produced were subsequently 199 purified, digested with the corresponding restriction enzymes (Promega), and ligated 200 into a similarly restricted pYES2 yeast expression vector (Invitrogen, Paisley, UK). The 201 purified plasmids (GenElute[™] Plasmid Miniprep Kit, Sigma) containing the octopus 202 elongase ORF were then used to transform Saccharomyces cerevisiae competent cells 203 (S.c. EasyComp Transformation Kit, Invitrogen). Transformation and selection of yeast 204 with recombinant pYES2-OVElo plasmids, and yeast culture were performed as 205 described in detail previously (Agaba et al., 2004).

206 In order to test the ability of octopus Elovl cDNA ORF to elongate either saturated or 207 monounsaturated FA, yeast transformed with pYES2 vector containing the octopus 208 elongase as an insert (pYES2-OVElo) and no insert (control) were grown in S. *cerevisiae* minimal medium^{-uracil} with no exogenously added FA substrates. 209 210 Additionally, the ability of *O. vulgaris* Fad to desaturate PUFA substrates was tested by 211 growing pYES2-OVElo transgenic yeast in medium supplemented with one of the 212 following substrates: 18:3n-3, 18:2n-6, 18:4n-3, 18:3n-6, 20:5n-3, 20:4n-6, 22:5n-3 and 213 22:4n-6. The FA were added to the yeast cultures at final concentrations of 0.5 (C18), 214 0.75 (C20) and 1.0 (C22) mM as uptake efficiency decreases with increasing chain 215 length (Zheng et al., 2009). Yeast transformed with empty pYES2 were also grown in presence of PUFA substrates as control treatments. After 2-day culture at 30 °C, yeast 216 217 were harvested, washed, and lipid extracted by homogenisation in chloroform/methanol 218 (2:1, v/v) containing 0.01% butylated hydroxy toluene (BHT) as antioxidant. All fatty 219 acid substrates, except stearidonic acid (18:4n-3), were purchased from Nu-Chek Prep,

Inc (Elysian, MN, USA). Stearidonic acid and chemicals used to prepare the S.
 cerevisiae minimal medium-^{uracil} were from Sigma Chemical Co. Ltd. (Dorset, UK),

except for the bacteriological agar obtained from Oxoid Ltd. (Hants, UK).

223 *Tissue distribution of elongase transcripts*

224 Expression of the octopus elongase was examined in adult tissues by RT-PCR. Total 225 RNA from a series of tissues including nerve, nephridium, hepatopancreas, brain, 226 caecum, gill, muscle, heart, and female and male gonads was extracted as described 227 above, and 1 µg of total RNA was reverse transcribed into cDNA (M-MLV reverse 228 transcriptase, Promega). In order to determine the mRNA distribution of the octopus 229 elongase, the tissue cDNAs were used as templates in PCR consisting of a denaturating 230 step at 95 °C for 1 min, followed by 35 cycles of denaturation at 95 °C for 30 s, 231 annealing at 58 °C for 30 s, extension at 72 °C for 30 s, followed by a final extension at 72 °C for 5 min (GoTag[®] Green Master Mix, Promega). Additionally, the expression of 232 233 the housekeeping β-actin was determined to check the cDNA integrity. Primers used for 234 RT-PCR are shown in Table 1.

235 Fatty acid analysis by GC-MS

236 FA from the transgenic yeast were analysed by preparing methyl esters (FAME) as 237 previously described (Hastings et al. 2001). Briefly, FAME were identified and 238 quantified using an Agilent 6850 Gas Chromatograph system coupled to a 5975 series 239 MSD (Agilent Technologies, Santa Clara, CA, USA). The elongation efficiency from 240 potential substrates including the yeast endogenous FA and the exogenously added 241 PUFA substrates (18:3n-3, 18:2n-6, 18:4n-3, 18:3n-6, 20:5n-3, 20:4n-6, 22:5n-3 and 242 22:4n-6) were calculated by the proportion of substrate FA converted to elongated FA 243 product as [product area/(product area + substrate area)] x 100. When further 244 confirmation of double bond positions was required, picolinyl esters were prepared from FAME according to the methodology described by Destaillats and Angers (2002)and modified according to Li et al. (2010).

247 In order to investigate the potential participation of the octopus elongase in the 248 biosynthesis of NMI FA, the FA compositions of specific tissues in which we had 249 previously detected NMI FA (Monroig et al., 2012a) were determined through 250 preparation of both methyl and picolinyl ester derivatives from polar lipid (PL) fractions 251 prepared as follows. Lipid extracts (2 mg) from nephrydium, male gonad, eye and 252 caecum were applied to 20x20 silica gel plates (Merck, Darmstadt, Germany) and 253 eluted with a solvent mixture of n-hexane / diethyl ether / glacial acetic acid (85:15:1.5, 254 v/v/v). PL fractions, identified by comparison with known standards, were scraped from 255 the plates and FAME prepared (Monroig et al., 2012a) and analysed as described above. FAME samples were subsequently derivatised to FA picolinyl prepared for 256 257 identification of the double bond patterns in NMI FA.

258

259 Results

260 Octopus elongase sequence and phylogenetics

261 The ORF of the newly cloned Elovl from O. vulgaris consisted of 885 bp encoding a 262 putative protein of 294 aa. Its sequence was deposited in the GenBank database with 263 accession number JX020803. The deduced as sequence from the octopus elongase 264 showed identity scores ranging from 39.3 and 43.2 % with several Elovl proteins 265 (Elov12, Elov14 and Elov15) from vertebrates including *H. sapiens*, *X. tropicalis* and *D.* 266 rerio. When compared with the two full-length elongases found in the genome of the 267 gastropod L. gigantea, the octopus Elovl was 58.1 % identical to the so-called 'L. 268 gigantea Elovl transcript 1' and 39.5 % identical to the 'Elovl transcript 2'. When the 269 octopus Elovl aa sequence was compared with incomplete elongase sequences from E.

scolopes, L. stagnalis, M. galloprovincialis and A. californica the identity scores were
relatively low, ranging from 31.9 to 43.9 %.

Similar to vertebrate Elovl-like proteins, the deduced aa sequence of the octopus elongase contained the diagnostic histidine box (HXXHH) conserved in all members of the Elovl protein family (Fig. 1). It also possessed two lysine (K) residues at the carboxyl terminus (KKXX), regarded as putative ER retrieval signals. Additionally, five putative transmembrane-spanning regions containing hydrophobic aa streches were predicted in residues 32-50, 65-83, 117-137, 158-192 and 239-259 by InterProScan (version 4.2) (Fig. 1).

279 A phylogenetic tree was constructed on the basis of aa sequence comparisons of the 280 octopus Elovl and other predicted elongases from molluscs, as well as several Elovl 281 types (Elovl 1-7) from a variety of vertebrates (Fig. 2). Our results showed that the 282 octopus Elovl protein clustered with other Elovl-like from molluscs including the 283 cephalopod E. scolopes and the gastropods L. stagnalis and L. gigantea ('transcript 1'), 284 altogether forming a group close to Elov12 and Elov15 proteins from vertebrates. More 285 distantly, three main clusters could be distinguished including Elov13/Elov16, 286 Elov11/Elov17 and Elov14 representatives. Interestingly, the Elov14 cluster included the 287 well-studied proteins from vertebrates, but also other mollusc Elovl-like proteins from 288 L. gigantea ('transcript 2'), A. californica and M. galloprovincialis.

289 Functional characterisation in yeast

The octopus Elovl-like encoding cDNA was functionally characterised by expressing the ORF in yeast *S. cerevisiae*. The FA composition of wild yeast consists basically of the main endogenous FA of *S. cerevisiae*, namely 16:0, 16:1 isomers (16:1n-9 and 16:1n-7), 18:0, 18:1n-9 and 18:1n-7 (Monroig et al., 2010a). Total lipid of yeast transformed with the empty pYES2 vector (control) contained these FA together with

whichever exogenous FA (if any) was added as substrate (data not shown), indicating that no elongase activity towards any of the exogenously added PUFA substrates assayed. This result is in agreement with the well-know inability of *S. cerevisiae* elongases to operate towards PUFA substrates (Hastings et al., 2001; Agaba et al., 2004).

300 In order to test the ability of the octopus Elovl to elongate saturated and 301 monounsaturated FA, yeast transformed with pYES2-OVElovl were grown in absence 302 of exogenously added substrates. Our results showed that none of the yeast endogenous 303 FA, whether saturated or monounsaturated, were elongated. Conversely, yeast 304 transformed with pYES2-OVElovl showed activity towards PUFA substrates producing 305 the corresponding 2-carbon elongation product. As shown in Fig. 3, the exogenously 306 added C18 (18:3n-3, 18:2n-6, 18:4n-3 and 18:3n-6) and C20 (20:5n-3 and 20:4n-6) 307 substrates were elongated to C20 (20:3n-3, 20:2n-6, 20:4n-3 and 20:3n-6) and C22 308 (22:5n-3 and 22:4n-6) products, respectively. Conversion rates derived from the yeast 309 assays suggested that the octopus Elovl generally elongated n-6 PUFA substrates more 310 efficiently than n-3 substrates for each pair of homologous substrates considered. Thus, 311 the substrates 18:2n-6, 18:3n-6, 20:4n-6 were consistently elongated at higher rates than 312 the corresponding n-3 PUFA substrates 18:3n-3, 18:4n-3 and 20:5n-3, respectively. 313 Interestingly, no activity towards C22 (22:5n-3 and 22:4n-6) PUFA substrates was 314 detected.

315 *Tissue distribution of octopus elongase transcripts*

Tissue expression of the common octopus Elovl was studied by RT-PCR on cDNA samples obtained from a range of tissues (Fig. 4). Except for nephridium, transcripts of the octopus Elovl gene were detected in all tissues analysed. Although RT-PCR

analyses should not be regarded as strictly quantitative data, our results indicate thatboth the male and female gonads showed higher expression signals.

321 Fatty acid composition from polar lipids of adult octopus tissues

322 FA from PL were analysed in several tissues of adult octopus individuals (Table 3).

323 DHA appeared the most abundant FA for each tissue considered, with up to 27.0 % of

324 total FA in eye PL. Other PUFA relatively abundant in the tissues studied were ARA

325 (with up to 16.4 % in male gonad PL) and EPA (up to 13.7 % in caecum PL).

326 Interestingly, 20:3n-3 content in eye was 13.5 % of total FA in the PL fraction.

GC-MS analysis of picolinyl esters enabled us to identify four different NMI FA in the octopus tissues, namely $\Delta 5,11$ 20:2, $\Delta 7,13$ 20:2, $\Delta 5,11,14$ 20:3 and $\Delta 7,13$ 22:2 (Table 3). Although we specifically analysed the PL fractions, where NMI FA are believed to accumulate (Klingensmith, 1982; Pirini et al., 2007), the amounts of all the NMI FA identified were generally low, and only relatively higher contents were detected for $\Delta 5,11$ 20:2 in nephrydium (1.8 %) and its corresponding elongation product $\Delta 7,13$ 22:2 in male gonad (2.2 %).

334 Discussion

335 The FA biosynthesis pathways have been investigated in both terrestrial (van der 336 Horst 1973, 1974; Weinert et al. 1993; Zhu et al. 1994) and aquatic mollusc species 337 (Chu and Greaves 1991; de Moreno et al. 1976; Waldock and Holland 1984; Zhukova 338 1986, 1991, 2007; Delaporte et al., 2005). It was shown that some molluscs have active 339 FA elongation systems (Waldock and Holland, 1984; Zhukova, 1986; Delaporte et al., 340 2005). In the present study we provide compelling evidence of the existence of an Elovl 341 cDNA that encodes an enzyme potentially involved in the biosynthesis of PUFA in the 342 cephalopod O. vulgaris.

343 The deduced aa sequence of the Elovl-like cDNA from O. vulgaris contains all the 344 features of the vertebrate Elovl protein family members, including five membrane-345 spanning regions, an ER retrieval signal at the C terminus containing lysine residues 346 (KKXX) and a diagnostic histidine box (HXXHH) (Leonard et al., 2004; Jakobsson et 347 al., 2006). Moreover, the histidine (H) box and its N-terminal side (QVTFLHVFHH) 348 show a typical as pattern of the PUFA elongase subfamily of eukaryotic elongases, with 349 a glutamine (Q) at position -5 and a leucine (L) at position -1 from the first H 350 (Hashimoto et al., 2008). Further evidence supporting a potential role of this octopus 351 Elovl cDNA in the PUFA biosynthetic pathways was provided by phylogenetic 352 analysis. Thus, the octopus Elovl aa sequence, as well as those of other mollusc 353 elongases, obtained by *in silico* searches, including the cephalopod *Euprymna scolopes* 354 and the gastropods Lymnaea stagnalis and Lottia gigantea (transcript 1), showed great 355 similarity to the sequences of Elov12 and Elov15 proteins, critical enzymes participating 356 in the biosynthesis of LC-PUFA in vertebrates (Leonard et al., 2004; Jakobsson et al., 357 2006). More distantly, the other elongase identified in the L. gigantea genome 358 (transcript 2) and also other Elovl-like proteins from A. californica and M. 359 galloprovincialis grouped together with vertebrate Elov14 elongases, another type of 360 elongase involved in the biosynthesis of very long-chain FA (C>24) including both 361 saturates and polyenes (Agbaga et al., 2008; Monroig et al., 2010b, 2011b, 2012b). 362 While these results suggest that another elongase with similarity to Elovl4 might also be 363 present in the common octopus, the functional characterisation of the present Elovl 364 cDNA confirmed, not only its participation in the PUFA elongation pathway, but also 365 that it has substrate specificities more similar to Elov15 than Elov12.

Clearly, transgenic yeast expressing the octopus Elovl efficiently converted C18 and
 C20 PUFA substrates to their corresponding 2-carbon elongated products, but no

activity towards C22 PUFA was detected. Generally, this pattern of substrate specificity 368 369 of the octopus elongase is consistent with that of vertebrate Elov15 proteins (Jakobsson 370 et al., 2006). For instance, the human ELOVL5 (also termed HELO1) and the rat 371 ELOVL5 (also termed rELO1) were shown to efficiently elongate C18 and C20 PUFA, 372 whereas C22 PUFA did not appear to be substrates for these enzymes (Leonard et al., 373 2000; Inagaki et al., 2002). Similarly, fish Elov15 demonstrated high activity for the 374 elongation of C18 and C20 PUFA substrates, whereas C22 substrates were only 375 elongated to a lesser extend (Agaba et al., 2004; Morais et al., 2009; Mohd-Yusof et al., 376 2010; Monroig et al., 2012b). Importantly, elongation of C22 PUFA including 22:5n-3 377 and 22:4n-6 in vertebrates is basically mediated by Elov12, whose substrate chain-length 378 specificity also includes C20, but not C18, PUFA substrates, the latter being only 379 marginally or not elongated (Tvrdik et al., 2000; Leonard et al., 2002; Monroig et al., 380 2009; Morais et al., 2009). Overall it can be concluded that the O. vulgaris elongase 381 cloned here is phenotypically an ElovI5-like elongase, but its sequence similarity to 382 vertebrate Elovl2 suggests an interesting evolutionary scenario that is worth exploring 383 in future investigations.

384 The functional characterisation of the octopus Elovl revealed, however, that the gene 385 product might have conserved/acquired a different PUFA family specificity compared 386 to vertebrate Elov15 proteins during evolution. Unlike mammalian (Leonard et al., 2000; 387 Inagaki et al., 2002) and fish Elov15 (Agaba et al., 2005; Mohd-Yusof et al., 2010; 388 Morais et al., 2011; Monroig et al., 2012b), which are generally more efficient in 389 elongating n-3 rather than n-6 FA substrates, the octopus Elovl exhibited higher 390 elongation rates towards n-6 compared to n-3 substrates for each homologous pair 391 considered. Thus 18:2n-6, 18:3n-6, 20:4n-6 were all elongated at higher rates than the 392 corresponding n-3 FA, namely 18:3n-3, 18:4n-3 and 20:5n-3, respectively. These results

393 emphasise that n-6 FA in general, and especially ARA (20:4n-6), might play 394 particularily important physiological roles in the common octopus. Consitent with this, 395 several studies have reported unexpectedly high levels of ARA in tissues of common 396 that were unlikely to derive purely from dietary origin and, thus, an active biosynthesis 397 of ARA in the common octopus was postulated (Milou et al., 2006; García-Garrido et 398 al., 2010; Monroig et al., 2012a). In the present study, the efficiency shown by the 399 octopus Elovl to elongate certain PUFA substrates indicates that this enzyme could 400 contribute to the endogenous biosynthesis of ARA in this species.

401 In vertebrates, ARA is biosynthesised from the dietary essential C18 PUFA 18:2n-6 402 through two alternative pathways, the 'classical' $\Delta 6$ -pathway ($\Delta 6$ desaturation \rightarrow 403 elongation $\rightarrow \Delta 5$ desaturation), or alternatively through the so-called ' $\Delta 8$ -pathway' 404 (elongation $\rightarrow \Delta 8$ desaturation $\rightarrow \Delta 5$ desaturation) (Monroig et al., 2011c). In addition 405 to the ability of the formerly characterised Fad cDNA to mediate the Δ 5-desaturation 406 steps of these pathways (Monroig et al., 2012a), we here demonstrate that the newly 407 cloned octopus Elovl can efficiently catalyse the elongation reactions required for ARA 408 biosynthesis from the dietary essential 18:2n-6, namely 18:3n-6 \rightarrow 20:3n-6 for the Δ 6-409 pathway and 18:2n-6 \rightarrow 20:3n-6 for the Δ 8-pathway. Although genes responsible for 410 elongation and $\Delta 5$ desaturation steps of these pathways have now been identified in 411 octopus, no Fad cDNA with $\Delta 6$ or $\Delta 8$ -desaturase activity has yet been identified and, 412 consequently, it remains unclear whether the common octopus can biosynthesise ARA 413 from the dietary essential 18:2n-6. This appears to be the case for some abalone species 414 (Dunstan et al., 1996; Durazo-Beltrán et al., 2003) but other species like C. gigas 415 (Waldock and Holland, 1984) and Mytilus edulis (Zhukova, 1991) appear unable to 416 biosynthesise ARA from 18:2n-6.

In addition to the biosynthesis of conventional PUFA, the octopus Elovl can also 417 418 have a role in the production of non-methylene-interrupted (NMI) FA. Thus, the 419 biosynthesis of $\Delta 7,13$ 22:2 encountered in male gonad, eye and caecum may be 420 accounted for by the elongation of $\Delta 5,11$ 20:2, as described for other marine 421 invertebrates (Kornprobst and Barnathan, 2010). Although we cannot directly conclude 422 that the octopus ElovI has the ability to elongate $\Delta 5.11$ 20:2 as this substrate was not 423 available, some of our results suggest a role for the elongase in the production of $\Delta 7,13$ 424 22:2 from $\Delta 5.11$ 20:2. First, the increased expression signal of Elovl in the male gonad 425 is consistent with this tissue containing the highest amount of $\Delta 7,13$ 20:2. Second, it is 426 reasonable to assume that, similar to the elongation rates exhibited towards other C20 427 PUFA like 20:4n-3 and 20:3n-6, the octopus Elovl might also efficiently operate 428 towards another C20 PUFA like $\Delta 5,11$ 20:2. Whereas these circumstantial data suggest 429 that the octopus Elovl may contribute to the endogenous biosynthesis of NMI FA in this 430 cephalopod, the extent to which this biosynthetic pathway is operative in the common 431 octopus is difficult to predict. On one hand, the ability of the octopus $\Delta 5$ Fad to convert 432 20:3n-3 (Δ 11,14,17 20:3) and 20:2n-6 (Δ 11,14 20:2) to the NMI FA Δ 5,11,14,17 20:4 433 and $\Delta 5,11,14$ 20:3, respectively (Monroig et al., 2012a), supports the hypothesis of a 434 notable production of NMI FA by O. vulgaris itself. On the other, the endogenous 435 biosynthesis of NMI FA in the common octopus appears to be limited as, despite the likely intake of preformed NMI FA through the diet, they still present relative low 436 437 levels compared to those found in some bivalves (Klingensmith, 1982) or nudibranchs 438 (Zhukova, 2007).

In summary, the present study demonstrates that the common octopus possesses an
Elovl-like cDNA with high homology to vertebrate Elovl5 and Elovl2 enzymes. The
functions of the octopus Elovl, while generally consistent those of vertebrate Elovl5,

have some novel particularities. Thus, the octopus Elovl showed higher elongation
efficiency towards n-6 than n-3 PUFA suggesting that these compounds, and especially
ARA, might play particularily pivotal physiological roles in the common octopus.
Moreover, the Elovl might be involved in the biosynthesis of NMI FA, although the
quantitative significance of this biosynthetic pathways in *O. vulgaris* requires further
investigation.

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- 683

684 Tables

685 Table 1. Sequences of the primer pairs used and accession numbers of the sequences used as references for primer design in the cloning of the octopus elongase of very long-686 687 chain fatty acids (Elovl) ORF and for RT-PCR analysis of gene expression in octopus

688 tissues.

| Aim | Transcript | Primer | Primer sequence | Ac |
|-------------|------------|---------|---|----|
| | | | | |
| RACE PCR | Elovl | OVEloF1 | 5'-GACTTGGTTCGGTGCTTGTT-3' | • |
| | | OVEloF2 | 5'-ATGGCCTGTCTGCTATACCAT-3' | |
| | | OVEloR1 | 5'-ATGGTATAGCAGACAGGCCAT-3' | |
| | | OVEloR2 | 5'-ATGATGGAAGACATGCAGGAA-3' | |
| ORF cloning | Elovl | OVEloVF | 5'-CCCAAGCTTAAAATGGCGGACGTTGTG-3' | |
| | | OVEloVR | 5'-CCG <u>GAGCTC</u> CTATTGAGCTTTCTTCACC-3' | |
| RT-PCR | Elovl | OVEloF1 | 5'-GACTTGGTTCGGTGCTTGTT-3' | |
| | 21011 | OVEloR3 | 5'-GTCTGCCTTTGATGTAAGCCTG-3' | • |
| | ß-actin | OVACTF | 5'-CTTGACTCCGGAGATGGTGT-3' | ł |
| | | OVACTR | 5'-CGCATTTCATGATGGAGTTG-3' | 1 |

689 ^a GenBank (<u>http://www.ncbi.nlm.nih.gov/</u>)

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Table 2. Functional characterisation of the octopus elongase in Saccharomyces 691

692 cerevisiae. Results are expressed as a percentage of total fatty acid (FA) substrate

693 converted to elongated products.

| FA Substrate | Product | % Conversion | Activity |
|--------------|---------|--------------|----------|
| 18:3n-3 | 20:3n-3 | 13.4 | C18→20 |
| 18:2n-6 | 20:2n-6 | 40.8 | C18→20 |
| 18:4n-3 | 20:4n-3 | 36.9 | C18→20 |
| 18:3n-6 | 20:3n-6 | 52.3 | C18→20 |
| 20:5n-3 | 22:5n-3 | 2.4 | C20→22 |
| 20:4n-6 | 22:4n-6 | 15.9 | C20→22 |
| 22:5n-3 | 24:5n-3 | 0.0 | C22→24 |
| 22:4n-6 | 24:4n-6 | 0.0 | C22→24 |

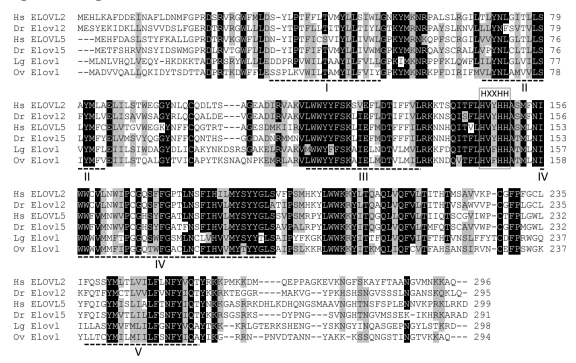
Table 3. Fatty acid (FA) composition (% of totals) from the polar lipids of tissues collected from *Octopus vulgaris* adult individuals. FA are designated using the 'n-' nomenclature, except for non-methylene-interrupted FA where the ' Δ ' nomenclature

698 was used.

| | Nephrydium | Male gonad | Eye | Caecum |
|------------------|------------|---------------|------|--------|
| 14:0 | 0.7 | 0.5 | 0.7 | 1.3 |
| 15:0 | 0.3 | 0.3 | 0.3 | 0.3 |
| 16:0 | 14.2 | 14.7 | 18.9 | 14.8 |
| 16:1n-9 | nd | 0.5 | 0.2 | 0.1 |
| 16:1 n- 7 | 0.5 | 0.4 | 0.4 | 1.9 |
| 16:0 iso | 0.2 | 0.1 | 0.2 | 0.2 |
| 16:0 anteiso | 0.2 | nd | 0.1 | nd |
| 17:0 | 2.7 | 1.4 | 1.2 | 1.9 |
| 17:1 | nd | 0.1 | 0.2 | nd |
| 17:0 iso | 0.3 | nd | 0.2 | 0.2 |
| 18:0 | 13.6 | 8.4 | 7.4 | 14.9 |
| 18:1n-13 | 0.6 | 0.9 | 1.4 | 0.3 |
| 18:1 n- 9 | 2.0 | 2.1 | 1.1 | 3.1 |
| 18:1n-7 | 2.0 | 1.3 | 1.4 | 2.6 |
| 18:1n-5 | 0.3 | 0.1 | nd | nd |
| 18:2n-6 | 0.1 | nd | 0.8 | 0.3 |
| 18:3n-3 | nd | nd | 0.1 | 0.2 |
| 18:4 n -3 | nd | nd | nd | 0.1 |
| 20:0 | 0.1 | 0.1 | 0.1 | 0.2 |
| 20:1n-11 | 0.5 | 0.5 | 0.5 | 1.0 |
| 20:1n-9 | 9.2 | 10.5 | 2.4 | 2.8 |
| 20:1n-7 | 0.2 | 0.2 | 0.1 | 0.2 |
| Δ5,11 20:2 | 1.8 | nd | nd | nd |
| Δ7,13 20:2 | nd | 0.2 | 0.1 | 0.1 |
| 20:2n-6 | 0.3 | 0.1 | 0.8 | 0.4 |
| Δ5,11,14 20:3 | 0.8 | nd | nd | nd |
| 20:3n-6 | 0.1 | nd | 0.2 | 0.1 |
| 20:4n-6 | 11.9 | 16.4 | 5.1 | 13.6 |
| 20:3n-3 | 0.1 | nd | 13.5 | 0.1 |
| 20:5n-3 | 10.0 | 7.3 | 11.4 | 13.7 |
| 22:0 | 0.2 | 0.1 | 0.2 | 0.3 |
| 22:1n-11 | 2.0 | 2.3 | 0.4 | 1.8 |
| 22:1n-9 | 0.1 | 0.1 | nd | 0.3 |
| Δ7,13 22:2 | nd | 2.2 | 0.4 | 0.8 |
| 21:5n-3 | 0.1 | nd | 0.1 | nd |
| 22:2n-6 | nd | nd | nd | 0.3 |
| 22:4n-6 | 1.1 | 7.7 | 0.3 | 1.1 |
| 22:5n-6 | 1.0 | 0.8 | 0.2 | 0.8 |
| 22:5n-3 | 1.3 | 2.0 | 1.0 | 1.5 |
| 22:6n-3 | 20.3 | 17.4 | 27.0 | 15.5 |
| 24:1n-9 | 0.1 | 0.1 | 0.2 | 0.4 |
| | | | | |

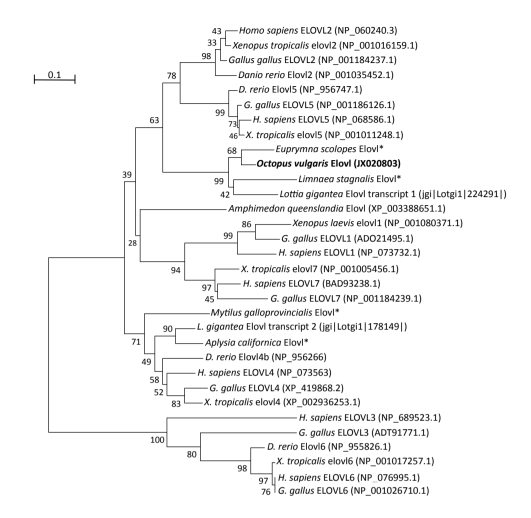
699 nd, no detected.

700 Legends to Figures

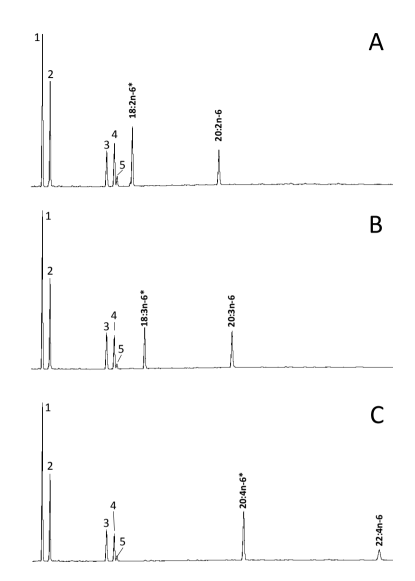


701

702 Fig. 1. Alignment of the deduced amino acid (aa) sequence of the elongase from 703 Octopus vulgaris (Ov). The aa sequence of the octopus Elovl-like protein was aligned 704 with the Homo sapiens (Hs) ELOVL2 (gb|NP 060240|) and ELOVL5 (gb|NP 068586|), 705 the Danio rerio (Ds) Elov12 (gb|NP 001035452) and Elov15 (gb|NP 956747) and the 706 so-called Elovl-like transcript 1 (jgi|Lotgi1|224291|) from Lottia gigantea (Lg) using 707 ClustalW (Bioedit). Identical residues are shaded black and similar residues are shaded 708 grey. Identity/similarity shading was based on the BLOSUM62 matrix, and the cut-off 709 for shading was 70%. The histidine box (HXXHH) conserved among Elovl family 710 members is highlighted with a grey square. Five (I-V) transmembrane-regions predicted 711 by InterProScan (http://www.ebi.ac.uk/Tools/pfa/iprscan/) are dot-underlined.



714 Fig. 2. Phylogenetic tree comparing the deduced amino acid (aa) sequence of the 715 Octopus vulgaris elongase of very long-chain fatty acids (Elovl)-like with a series of 716 protein sequences including representatives of the seven (1-7) Elovl subtypes and other 717 Elovl-like sequences from invertebrate organisms. All accession numbers are from 718 GenBank database, except for Lottia gigantea elongases where JGI protein ID are given 719 (http://genome.jgi-psf. org/Lotgi1/Lotgi1.home.html). Asterisks indicate the aa 720 sequences deduced from searches and subsequent assembly of expressed sequence tags 721 (EST) using NCBI tblastn tool (http://www.ncbi.nlm.nih.gov/) as described in Materials 722 and Methods. The tree was constructed using the Neighbour Joining method (Saitou and 723 Nei 1987) with MEGA4. The horizontal branch length is proportional to aa substitution 724 rate per site. The numbers represent the frequencies (%) with which the tree topology 725 presented was replicated after 10000 iterations.



727

728 Fig. 3. Functional characterisation of the Octopus vulgaris elongase of very long-chain 729 fatty acids (Elovl) in yeast (Saccharomyces cerevisiae). The fatty acid (FA) profiles of 730 yeast transformed with pYES2 containing the ORF of the putative Elovl cDNA as an 731 insert, were determined after the yeast was grown in the presence of one of the 732 exogenously added substrates 18:2n-6 (A), 18:3n-6 (B) and 20:3n-6 (C). Peaks 1-5 in all 733 panels are the main endogenous FA of S. cerevisiae, namely 16:0 (1), 16:1 isomers (2), 734 18:0 (3), 18:1n-9 (4) and 18:1n-7 (5). Additionally peaks derived from exogenously added substrates ("*") or elongation products are indicated accordingly in panels A-C. 735 736 Vertical axis, FID response; horizontal axis, retention time.

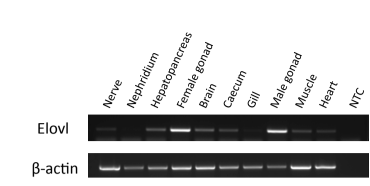


Fig. 4. RT-PCR analyses showing the tissue distribution of octopus elongase of very

739 long-chain fatty acids (Elovl) transcripts. Expression of the housekeeping gene β-actin

740 is also shown. NTC, no template control.