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2 Biosynthesis of long-chain polyunsaturated fatty acids in the African catfish Clarias gariepinus: Molecular cloning and functional characterisation of fatty acyl desaturase (fads2) 3 and elongase (elovl2) cDNAs 4 5 6 Authors Angela Oboh^{1,2}, Mónica B. Betancor¹, Douglas R. Tocher¹, Oscar Monroig¹*, 7 8 Addresses 9 ¹ Institute of Aquaculture, School of Natural Sciences, University of Stirling, Stirling FK9 10 4LA, Scotland, UK 11 ² Department of Biological Sciences, University of Abuja, P.M.B. 117, Nigeria 12 13 *Corresponding author 14 Oscar Monroig 15 Institute of Aquaculture, School of Natural Sciences, University of Stirling, Stirling FK9 16 4LA, Scotland, UK 17 Tel: +44 1786 467892; Fax: +44 1786 472133; E-mail: oscar.monroig@stir.ac.uk 18 19 20

21 Abstract

Fish differ in their capacity for endogenous synthesis of long-chain (C_{20-24}) polyunsaturated 22 fatty acids (LC-PUFA) from dietary C₁₈ precursors (α-linolenic and linoleic acids). 23 Understanding this capacity is of benefit to fish feed formulation. This, together with the 24 importance of fish as the primary source of omega-3 LC-PUFA in the human diet has 25 necessitated the rigorous study of the biochemical and molecular mechanisms involved in the 26 LC-PUFA biosynthesis pathway in fish species. Studies have shown the potential of a species 27 for LC-PUFA biosynthesis is associated with the complement and function of fatty acyl 28 29 desaturase (fads) and elongase of very long chain fatty acid (elovl) gene it possesses. The present study therefore aimed to investigate these genes in the African catfish (Clarias 30 gariepinus), the most commercially important farmed fish species in Sub-Saharan Africa. A 31 32 fads2 and an elovl2 cDNA were cloned containing open reading frames (ORF) of 1338 base pair (bp) and 864 bp specifying proteins of 445 and 287 amino acids, respectively. Functional 33 characterisation by heterologous expression in yeast showed that the Fads2 was bifunctional 34 with $\Delta 5\Delta 6$ activities catalysing the desaturation of both 18:3n-3 and 20:4n-3 and their 35 corresponding n-6 fatty acids, 18:2n-6 and 20:3n-6. The Elovl2 showed activity towards C₁₈, 36 C20 and C22 PUFA with highest activity towards C20 and C22 PUFA. Tissue expression 37 analysis showed a typical freshwater species expression pattern; higher expression in the liver 38 compared to brain and all other tissues with the exception of *elov15* which showed highest 39 40 expression in the intestine. Consistent with feeding studies of typical freshwater fish species that show their essential fatty acid requirement can be satisfied by dietary C₁₈ PUFA, the 41 present study confirms that the LC-PUFA biosynthesis pathway is active in the African 42 43 catfish C. gariepinus.

44

46 Keywords

- 47 Biosynthesis; Clarias gariepinus; elongase; essential fatty acids; desaturase; long-chain
- 48 polyunsaturated fatty acids

50 Introduction

Fish, like all vertebrates, are dependent on dietary sources of polyunsaturated fatty acids 51 (PUFA) such as α-linolenic (ALA, 18:3n-3) and linoleic (LA, 18:2n-6) acids as they lack the 52 $\Delta 12$ and $\Delta 15$ desaturases required for the synthesis of LA and ALA from oleic acid (18:1n-9) 53 (Tocher, 2010, 2015; Tocher and Glencross, 2015). However, whereas the C₁₈ PUFA, ALA 54 and LA, can satisfy essential fatty acid (EFA) requirements of some fish species, long-chain 55 (C₂₀₋₂₄) polyunsaturated fatty acids (LC-PUFA) including eicosapentaenoic acid (EPA, 20:5n-56 3), docosahexaenoic acid (DHA, 22:6n-3) and arachidonic acid (ARA, 20:4n-6), which play 57 58 physiologically important roles, are required in the diet to meet the EFA requirements of other species. This reflects the differing ability of fish species to endogenously synthesise 59 LC-PUFA from C_{18} precursors, associated with the complement of fatty acyl desaturases 60 61 (Fads) and elongation of very long-chain fatty acids (Elovl) enzymes they possess (Bell and Tocher, 2009; Castro et al., 2016; Tocher, 2010). This has important implications with 62 regards to feed formulation for fish farming. Species with active and complete biosynthetic 63 pathways can convert C₁₈ PUFA contained in vegetable oils (VO) that are now common 64 ingredients in aquafeeds, to LC-PUFA, and thus are less dependent on the inclusion of fish 65 oil (FO) to supply LC-PUFA in their diets. 66

67 The LC-PUFA biosynthesis pathways involves successive desaturation and elongation of the C₁₈ precursors catalysed by Fads and Elovl elongases (Castro et al., 2016; Monroig et al., 68 2011a; Tocher, 2003; Vagner and Santigosa, 2011). Fads enzymes introduce double bonds 69 (unsaturations) at specific positions of the fatty acyl chain (Nakamura and Nara, 2004). It has 70 been shown that all fads so far studied in teleost genomes are paralogues of fads2, a gene 71 encoding an enzyme that typically acts as $\Delta 6$ Fads in vertebrates, while *fads1*, another 72 vertebrate *fads* encoding an enzyme with $\Delta 5$ activity, appears to be absent in teleosts (Castro 73 et al., 2012, 2016). While most fish Fads2 enzymes functionally characterised are typically 74

 $\Delta 6$, others have been characterised as bifunctional $\Delta 6\Delta 5$ Fads2 (Fonseca-Madrigal et al., 2014; Hastings et al., 2001; Li et al., 2010; Tanomman et al., 2013) or monofunctional $\Delta 5$ Fads (Abdul Hamid et al., 2016; Hastings et al., 2005). In recent years, Fads2 with $\Delta 4$ activities have been found in a variety of teleost species (Fonseca-Madrigal et al., 2014; Kuah et al., 2015; Li et al., 2010; Morais et al., 2012). Furthermore, fish Fads2, as described in mammals (Park et al., 2009), also display $\Delta 8$ activity, an activity that appeared to be relatively higher in marine fish compared to freshwater fish species (Monroig et al., 2011a).

Elovl enzymes catalyse the condensation step in the elongation pathway resulting in the 82 addition of a two-carbon unit to the pre-existing fatty acid (Guillou et al., 2010). Functional 83 84 characterisation of fish Elovl2, Elovl4 and Elovl5, elongase enzymes that function in the LC-85 PUFA biosynthesis pathway, show that they display somewhat overlapping activities (Castro et al., 2016). Thus Elov15 generally elongate C18 and C20 PUFA, whereas Elov12 and Elov14 86 are more efficient towards C₂₀ and C₂₂ PUFA (Gregory and James, 2014; Monroig et al., 87 2011a, 2011b, 2009; Morais et al., 2009). While both *elov15* and *elov14* genes are present in 88 teleost genomes (Monroig et al., 2016), elovl2 appears to be lost in Acanthopterygii, a 89 phylogenic group that, with the exception of salmonids, includes the vast majority of the most 90 important farmed fish species (Castro et al., 2016). To the best of our knowledge, Elovl2 91 92 cDNAs have been characterised only in Atlantic salmon (Salmo salar) (Morais et al., 2009), zebrafish (Danio rerio) (Monroig et al., 2009) and rainbow trout (Oncorhynchus mykiss) 93 (Gregory and James, 2014). 94

Evidence indicates that the complement and functionalities of *fads* and *elovl* genes existing in any teleost species has been shaped by evolutionary drivers leading to the retention, subfunctionalisation or loss of these genes (Castro et al., 2016). Moreover, the habitat (marine *vs* freshwater), and specifically the availability of LC-PUFA in food webs, has also been implicated as influencing the LC-PUFA biosynthetic capability of fish (Bell and Tocher,

2009; Castro et al., 2016; Monroig et al., 2011b). Freshwater fish, having evolved on diets 100 low in LC-PUFA, are believed to have all the genes and/or enzymatic functionalities required 101 for endogenous LC-PUFA production (NRC, 2011; Tocher, 2015). Whereas, many marine 102 species have not retained all the genes and/or enzymatic functionalities required for 103 endogenous LC-PUFA production as a consequence of LC-PUFA being readily available in 104 their natural diets (NRC, 2011; Tocher, 2015). However, such dichotomy has been recently 105 seen as too simplistic and other factors including trophic level (Li et al., 2010) and trophic 106 ecology (Morais et al., 2015, 2012) also appear to influence LC-PUFA biosynthesis capacity 107 108 of fish species. Within an aquaculture nutrition context, investigations of the *fads* and *elovl* gene repertoire involved in LC-PUFA biosynthesis, as well as the functions of the enzyme 109 they encode, are necessary to ascertain whether the EFA requirements of a species can be 110 111 satisfied by C_{18} PUFA or whether dietary LC-PUFA are required.

The African catfish Clarias gariepinus, a freshwater species belonging to the family 112 Clariidae and order Siluriformes, is the most important aquaculture species in Sub-Saharan 113 Africa (FAO, 2012). Yet, neither its LC-PUFA biosynthetic pathway nor EFA requirement is 114 fully understood. Studies on C. gariepinus and other African catfishes suggest they can 115 effectively utilise C₁₈ PUFA contained in VO to cover their physiologically important LC-116 PUFA requirements (Baker and Davies, 1996; Sotolu, 2010; Szabo et al., 2009). Intriguingly, 117 lower growth performance has been reported for C. gariepinus fed diets with FO compared to 118 those fed diets containing VO (Hoffman and Prinsloo, 1995; Ng et al., 2003) in contrast to 119 most fish species including those with full LC-PUFA biosynthetic capability like salmonids 120 (Sargent et al., 2002; Tocher and Glencross, 2015). 121

The aim of this study was to investigate the functions of the genes encoding putative Fads and Elovl enzymes that account for the LC-PUFA biosynthetic capability of *C. gariepinus* and thus understand the potential of this species to utilise diets containing VO and low contents of LC-PUFA. Here, we report the cloning and functional characterisation of *fads2*and *elovl2* genes from *C. gariepinus*. We further investigated the mRNA tissue distribution of
the newly cloned genes, as well as that of the previously cloned *elovl5* (Agaba et al., 2005).

128

129 Materials and Methods

130 Sample collection and RNA preparation

Tissue samples were obtained from adult C. gariepinus (~1.8 kg) raised in the tropical 131 aquarium of the Institute of Aquaculture, University of Stirling, UK, on standard salmonid 132 diets. Eight C. gariepinus individuals were sacrificed with an overdose of benzocaine before 133 the collection of tissue samples including liver, intestine, pituitary, testis, ovary, skin, muscle, 134 gills, kidney, and brain. The samples were immediately preserved in an RNA stabilisation 135 136 buffer (3.6 M ammonium sulphate, 18 mM sodium citrate, 15 mM EDTA, pH 5.2) and stored at -80 °C prior to extraction of total RNA following homogenisation in TRI Reagent® 137 (Sigma-Aldrich, USA). Purity and concentration of total RNA was assessed using the 138 NanoDrop[®] (Labtech International ND-1000 spectrophotometer) and integrity was assessed 139 on an agarose gel. First strand complementary DNA (cDNA) was synthesised using High 140 Capacity cDNA Reverse Transcription Kit (Applied BiosystemsTM, USA) following the 141 manufacturer's instructions. 142

143 Molecular cloning of fads2 and elovl2 cDNAs

Amplification of partial fragments of the genes was achieved by polymerase chain reaction (PCR) using a mixture of cDNA from eye, liver, intestine and brain as template and primers FadCGF2F1 and FadCGF2R1 for *fads2*, and EloCGE2F1 and EloCGE2R1 for *elovl2* (Table 1). For clarity, it should be noted that the standard gene/protein nomenclature has been used in this study (Castro et al., 2016). Following conventions accepted for zebrafish (*Danio*

rerio), proteins are termed with regular fonts (e.g. Fads2) whereas genes are italicised (e.g. 149 fads2). Primers used for amplification of the first fragment of target genes were designed on 150 conserved regions of fish orthologues of *fads2* and *elovl2* according to the following strategy. 151 For *fads2*, sequences from the broadhead catfish (*Clarias microcephalus*) (gb|KF006248.1|), 152 spot pangasius (Pangasius larnaudii) (gb|KC994461.1|), striped catfish (Pangasianodon 153 *hypophthalamus*) (gb|JX035811.1|) and *Clarias* hybrid (*C. macrocephalus* and *C. gariepinus*) 154 (gb|KC994463.1|) were aligned with the ClustalW tool (BioEdit v7.0.9, Tom Hall, 155 Department of Microbiology, North Carolina State University, USA) for degenerate primer 156 157 design. For *elovl2*, homologous sequences from *D. rerio* (gb|NM 001040362.1|), *S. salar* (gb|NM 001136553.1)) and Mexican tetra (Astyanax mexicanus) (gb|XM 007260074.2)) 158 were retrieved from NCBI (http://ncbi.nlm.nih.gov), aligned (BioEdit) and conserved regions 159 160 used to retrieve expressed sequence tags (ESTs) from catfish species. Six Channel catfish (Ictalurus punctatus) ESTs (GenBank accession numbers GH651976.1, GH651977.1, 161 FD328544.1, FD284236.1, FD284235.1 and BM438219.1) were obtained and aligned with 162 BioEdit. Subsequently, the consensus *elovl2*-like sequences from *I. punctatus*, and those from 163 D. rerio, S. salar and A. mexicanus, were aligned for degenerate primer design. PCR 164 conditions consisted of an initial denaturation step at 95 °C for 2 min, followed by 33 cycles 165 of denaturation at 95 °C for 30 s, annealing at 57 °C for 30 s, extension at 72 °C for 1 min 30 166 s, followed by a final extension at 72 °C for 7 min. The PCR fragments were purified using 167 168 the Illustra GFX PCR DNA/gel band purification kit (GE Healthcare, Little Chalfont, UK), and sequenced (GATC Biotech Ltd., Konstanz, Germany). The primers used in this study and 169 their sequences are presented in Table 1. 170

In order to obtain full-length cDNA sequences, Rapid Amplification of cDNA Ends
(RACE) was performed with the FirstChoice[®] RLM-RACE RNA ligase mediated RACE kit
(Ambion[®], Life TechnologiesTM, USA). The 5' RACE outer primer and gene-specific primer

FadCGRF2R3 were used in a PCR using the 5' RACE cDNA as template (first round PCR) 174 for *fads2*. The resulting PCR product was then used as template for the second round PCR 175 with the 5' RACE inner primer and the gene-specific primer FadCGRF2R2. A similar 176 approach was followed to perform 3' RACE PCR, with primers FadCGRF2F1 and 177 FadCGRF2F2 used for first and second round PCR, respectively. For elovl2, the primers 178 CGRE2R3 and CGRE2R2 were designed and used for first and second round PCR, 179 respectively, for the 5' RACE PCR, while CGRE2F1 and CGRE2F2 were used for first and 180 second round PCR, respectively, for the 3' RACE PCR. The first fragments, 5' and 3' RACE 181 PCR fragments were then cloned into PCR 2.1 vector (TA cloning[®] kit, Invitrogen, Life 182 TechnologiesTM, USA) and sequenced as above. The full nucleotide sequences of the *fads2* 183 and *elovl2* cDNAs were obtained by aligning sequences of the first fragments, together with 184 185 those of the 5' and 3' RACE PCR positive products (BioEdit).

186

187 Sequence and phylogenetic analysis

The deduced amino acid (aa) sequences of the C. gariepinus fads2 and elovl2 cDNAs were 188 compared to corresponding orthologues from other vertebrate species and sequence identity 189 scores were calculated using the EMBOSS Needle Pairwise Sequence Alignment tool 190 (http://www.ebi.ac.uk/Tools/psa/emboss needle/). Phylogenetic analysis of the deduced aa 191 sequences of *fads2* and *elovl2* cDNAs from *C. gariepinus* and those from a variety of species 192 across vertebrate lineages were carried out by constructing trees using the neighbour-joining 193 method Nei, 1987), with the MEGA 4.0 (Saitou and software 194 (www.megasoftware.net/mega4/mega.html). Confidence in the resulting tree branch topology 195 was measured by bootstrapping through 1,000 iterations. 196

197

Functional characterisation of C. gariepinus fads2 and elovl2 by heterologous expression in
Saccharomyces cerevisiae

PCR fragments corresponding to the open reading frame (ORF) of C. gariepinus fads2 and 200 elovl2 were amplified from a mixture of cDNA synthesised from liver, intestine, eye and 201 brain total RNA, using the high fidelity Pfu DNA polymerase (Promega, USA) with primers 202 containing BamHI (forward) and XhoI (reverse) restriction sites (Table 1). PCR conditions 203 consisted of an initial denaturation step at 95 °C for 2 min, followed by 35 cycles of 204 denaturation at 95 °C for 30 s, annealing at 57 °C for 30 s, extension at 72 °C for 3 min 30 s 205 followed by a final extension at 72 °C for 7 min. The DNA fragments obtained were purified 206 as above, digested with the appropriate restriction enzymes, and ligated into similarly 207 digested pYES2 yeast expression vector (Invitrogen). 208

209 Yeast competent cells InvSc1 (Invitrogen) were transformed with the plasmid constructs pYES2-fads2 (desaturase) or pYES-elovl2 (elongase) or with empty vector (control) using 210 the S.c. EasyCompTM Transformation Kit (Invitrogen). Selection of yeast containing the 211 pYES2 constructs was performed on S. cerevisiae minimal medium minus uracil (SCMM^{-ura}) 212 plates. One single yeast colony was grown in SCMM^{-ura} broth for 2 days at 30 °C, and 213 subsequently subcultured in individual Erlenmeyer flasks until an optical density measured at 214 a wavelength of 600nm (OD₆₀₀) reached 1, after which galactose (2 %, w/v) and a PUFA 215 substrate were added. For both genes, final concentration of substrates were 0.5 mM (C_{18}), 216 0.75 mM (C₂₀) and 1.0 mM (C₂₂) to compensate for differential uptake related to fatty acyl 217 chain (Zheng et al., 2009). For the fads2, $\Delta 6$ (18:3n-3 and 18:2n-6), $\Delta 8$ (20:3n-3 and 20:2n-218 6), $\Delta 5$ (20:4n-3 and 20:3n-6), and $\Delta 4$ (22:5n-3 and 22:4n-6) Fads substrates were used. For 219 220 elovl2, substrates included C18, (18:3n-3, 18:2n-6, 18:4n-3 and 18:3n-6), C20 (20:5n-3 and 20:4n-6) and C₂₂ (22:5n-3 and 22:4n-6) PUFA. After 2 days, the yeasts were harvested, 221 washed and homogenised in chloroform/methanol (2:1, v/v) containing 0.01 % butylated 222

hydroxytoluene (BHT) and stored at -20 °C until further use. All FA substrates (> 98-99 %
pure) used for the functional characterisation assays, except for stearidonic acid (18:4n-3) and
eicosatetraenoic acid (20:4n-3), were obtained from Nu-Chek Prep, Inc. (Elysian, MN, USA).
Stearidonic acid (> 99 % pure) and yeast culture reagents including galactose, nitrogen base,
raffinose, tergitol NP-40 and uracil dropout medium were obtained from Sigma-Aldrich
(UK). Eicosatetraenoic acid was purchased from Cayman Chemical Co. (Ann Arbor, USA).

229

230 *Fatty acid analysis of yeast*

Total lipids extracted according to Folch et al. (1957) from yeast samples were used to prepare fatty acid methyl esters (FAME). FAME extraction, purification and analysis were performed as described by Li et al. (2010). Substrate FA conversion was calculated as the proportion of exogenously added FA substrate desaturated or elongated [all product peak areas / (all product peak areas + substrate peak area)] x 100 (Monroig et al., 2016). GC-MS was used to confirm double bond positions when necessary (Li et al., 2010).

237

238 *Gene expression analysis*

Expression of the newly cloned *fads2* and *elovl2* genes, as well as that of the previously 239 characterised elongase *elov15* (Agaba et al., 2005), were determined by quantitative real-time 240 PCR (qPCR). Extraction of RNA from tissues and cDNA synthesis were carried out as 241 described above. qPCR amplifications were carried out in duplicate using Biometra 242 Thermocycler (Analytik Jena company, Germany) and Luminaris Color Higreen qPCR 243 master mix (Thermo Scientific, CA, USA) following the manufacturer's instruction. The 244 qPCR was performed in a final volume of 20 μl containing 5 μl diluted (1/20) cDNA, 1 μl (10 245 µM) of each primer, 3 µl nuclease free water and 10 µl Luminaris Color Higreen qPCR 246 master mix. The qPCR conditions were 50 °C for 2 min, 95 °C for 10 min followed by 35 247

cycles of denaturation at 95 °C for 15 s, annealing at 59 °C for 30 s and extension at 72 °C for 248 30 s. After the amplifications, a dissociation curve of 0.5 °C increments from 60 to 90 °C was 249 performed, enabling confirmation of a single product in each reaction. Negative controls (no 250 251 template control, NTC) containing no cDNA were systematically run. Absolute copy number of the target and reference gene in each sample was calculated from the linear standard curve 252 constructed. Normalisation of each target gene was carried out by dividing the absolute copy 253 254 number of the candidate gene by the absolute copy number of the reference gene 28S rRNA (gb|AF323692.1|). In order to prepare solutions of known copy numbers, DNA concentration 255 256 linearised PCR 2.1 vectors containing a fragment of either candidate or reference genes was determined, and their molecular weights were estimated as 660 g bp x length (bp) of the 257 plasmid constructs. Primers used for qPCR analysis are also presented in Table 1. 258

259

260 *Statistical analysis*

Tissue expression (qPCR) results were expressed as mean normalised ratios (\pm SE) corresponding to the ratio between the copy numbers of the target genes (*fads2*, *elovl2* and *elovl5*) and the copy numbers of the reference gene, 28S rRNA. Differences in gene expression among tissues were analysed by one-way analysis of variance (ANOVA) followed by Tukey's HSD test at a significance level of P≤0.05 (IBM SPSS Statistics 21).

266

267 **Results**

268 Sequence and phylogenetic analysis

C. gariepinus Fads2 sequence was deposited in the GenBank database with the accession number KU925904. The full length of the *C. gariepinus* Fads2 was 1812 bp, comprising of a 5' untranslated region (UTR) of 162 bp, an ORF of 1338 bp encoding a putative protein of 445 aa, and a 3' UTR of 312 bp. The deduced *C. gariepinus* Fads2 enzyme showed distinctive structural features of fatty acyl desaturases including the three histidine boxes

HDFGH, HFQHH, and QIEHH (aa 181-185, 218-222 and 383-387, respectively) and 274 cytochrome b5-domain (aa 26-77) containing the heme binding motif HPGG (aa 54-57). 275 Pairwise aa sequence comparisons of C. gariepinus Fads2 with other Fads2-like proteins 276 showed highest identities with Fads from members of the catfish family such as C. 277 macrocephalus (97 %) and P. hypophthalamus (91.5 %). Comparisons with bifunctional 278 $\Delta 6\Delta 5$ Fads2 of *D. rerio* (gb|AF309556.1|) and *C. estor* (gb|AHX39207.1|), bifunctional $\Delta 5\Delta 4$ 279 Fads2 of C. striata (gb|ACD70298.1|) and S. canaliculatus (gb|ADJ29913.1|) and Δ4 Fads2 280 of S. senegalensis (gb|AEQ92868.1|) and C. estor (gb|AHX39206.1|) showed identities 281 282 ranging from 65.2-70.2 %. Lowest identities were observed when C. gariepinus Fads was compared to Fads1-like sequences from different vertebrate lineages. Phylogenesis of C. 283 gariepinus Fads with Fads from a variety of vertebrate species showed it clustered with all 284 285 other Fads2 in one group that was separate from the Fads1 group confirming that the newly cloned fads was a fads2 (Fig. 1). The C. gariepinus Fads2 clustered most closely with Fads2 286 from bony fish species (with the exception of the sarcopterygian, L. chalumnae which formed 287 a separate cluster with Fads2 from chondrichthyes (C. milli and S. canicula), mammalian (H. 288 sapiens, M. musculus and B. taurus) and avian species (G. gallus) (Fig. 1). 289

290

C. gariepinus Elovl2 sequence was deposited in the GenBank database with the accession 291 number KU902414. The full-length cDNA sequence of C. gariepinus elovl2 was 1432 bp (5' 292 293 UTR 91 bp, ORF 864 bp, 3' UTR 477 bp) encoding a protein of 287 aa. Analysis of the deduced as sequence of C. gariepinus Elovl2 revealed characteristic features of fatty acyl 294 elongases such as the highly conserved histidine box (HVYHH, aa 151-155) and the 295 carboxyl-terminal region, but the aa residues at the carboxyl terminus were KHKLQ, more 296 similar to the KXRXX found in Elov15 than to the KKXX in *H. sapiens* and *S. salar* Elov12 297 (Morais et al., 2009). Comparisons of C. gariepinus Elovl2 with homologues from A. 298

(gb|XP 007260136.1|), S. salar (gb|ACI62500.1|), D. 299 mexicanus rerio (gb|XP 005162628.1|), Clupea harengus (gb|XP 012671565.1|), and H. 300 sapiens (gb|NP 060240.3) showed identities of 81.7, 72.9, 72.7, 69.1 and 64.8 %, respectively. C. 301 gariepinus Elov12 shared 52 % identity with C. gariepinus Elov15. Phylogenetic analysis of 302 the Elovl2 with members of the Elovl family confirmed that the newly cloned elongase was 303 indeed an Elovl2 elongase. Thus, the C. gariepinus Elovl2 clustered together with all the 304 Elovl2 and more distantly from Elovl5 sequences including that from C. gariepinus (Agaba et 305 al., 2005) and even more distantly to Elovl4 enzymes (Fig. 2). 306

307

308 Functional characterisation of C. gariepinus Fads2 and Elovl2 in S. cerevisiae

Consistent with previous studies (Hastings et al., 2001), control yeast transformed with the 309 empty pYES2 vector did not show any activity towards any of the PUFA substrates assayed 310 (data not shown). Functional characterisation by heterologous expression in yeast revealed 311 that the C. gariepinus Fads2 had the ability to introduce double bonds at $\Delta 5$, $\Delta 6$ and $\Delta 8$ 312 positions in the appropriate PUFA substrates (Fig. 3; Table 2). The FA composition of the 313 yeast transformed with pYES2-fads2 showed peaks corresponding to the four main yeast 314 endogenous FA, namely 16:0, 16:1n-7, 18:0 and 18:1n-9, the exogenously added PUFA and 315 the corresponding PUFA product(s) (Fig. 3; Table 2). Thus, the C₁₈ PUFA substrates 18:3n-3 316 and 18:2n-6 were desaturated to 18:4n-3 (42 % conversion) and 18:3n-6 (23 %), respectively, 317 indicating the encoded protein had $\Delta 6$ Fads activity (Fig. 3A; Table 2). Moreover, the 318 transgenic yeast was able to desaturate 20:4n-3 and 20:3n-6 to 20:5n-3 (19 %) and 20:4n-6 319 (14 %), respectively, indicating the C. gariepinus Fads2 also had $\Delta 5$ activity (Fig. 3C; Table 320 2), and thus these results confirm that this Fads2 from C. gariepinus is a bifunctional $\Delta 6\Delta 5$ 321 Fads. Additionally, the C. gariepinus Fads2 showed $\Delta 8$ Fads activity as the yeast transformed 322 with pYES2-fads2 were able to desaturate 20:3n-3 and 20:2n-6 to 20:4n-3 and 20:3n-6, 323

respectively (Fig. 3B and Table 2). No additional peaks were observed when yeast expressing the *C. gariepinus fads2* were grown in the presence of 22:5n-3 and 22:4n-6 (Fig. 3D; Table 2).

The *C. gariepinus* Elovl2 showed the ability to elongate C_{18-22} PUFA substrates (Fig. 4; Table 3), with highest conversions towards the C_{20} substrates 20:5n-3 (73.4 %) (Fig. 4B) and 20:4n-6 (56 %). Conversion of the C_{22} substrate was 36.7 % for 22:5n-3 (Fig. 4C) and 9.7 % for 22:4n-6 (Table 3). Elongations of C_{18} PUFAs were generally lower compared to those for C_{20} and C_{22} substrates. Stepwise elongations derived from further activity of the *C. gariepinus* Elovl2 towards products of initial substrate elongation resulted in the production of several polyenes up to 24 carbons (Fig. 4; Table 3).

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335 *Tissue expression analysis of C. gariepinus fads2, elovl2 and elovl5*

Tissue distribution analysis of *C. gariepinus fads2*, *elovl2* and *elovl5* transcripts confirmed that these genes were expressed in all tissues analysed (Fig. 5). Liver and brain were found to contain the highest transcript levels of the *C. gariepinus fads2*, followed by pituitary, intestine and kidney. Liver, brain and pituitary were also found to contain the highest transcript levels of the *C. gariepinus elovl2*. Generally, gonads including testis and ovary showed the lowest transcript levels for both *fads2* and *elovl2* (Fig. 5). Intestine and liver exhibited the highest level of *elovl5*, while the lowest expression levels were found in muscle.

344 **Discussion**

Elucidating the LC-PUFA biosynthesis pathway in farmed fish is crucial for formulating diets that satisfy physiological requirements and thus ensure normal growth and development. These studies are particularly relevant in the current scenario whereby FO are being replaced by VO in aquafeed, the latter naturally devoid of essential LC-PUFA and thus potentially compromising both health of the fish and nutritional value for human consumers (Monroig et
al., 2011b; Tocher and Glencross, 2015). Relevant to the present study, identification and
production of fish that can efficiently utilise VO-based diets due to their high capacity for
LC-PUFA biosynthesis is a valid strategy to expand aquaculture considering that marine
ingredients (FO and fish meal) will be increasingly limited in the future (Tocher, 2015). *C. gariepinus* feed and grow well on a variety of feed ingredients and are, therefore, a good
model for studying the endogenous capacity for LC-PUFA synthesis of freshwater fish.

Phylogenetic analysis of the *fads*-like desaturase cDNA isolated from C. gariepinus, 356 together with the possession of all the main structural features common to the Fads2 protein 357 358 family confirmed it to be a Fads2. Sequence and phylogenetic analyses also showed that the C. gariepinus Fads2 shared highest aa sequence similarities with other catfish species, with 359 relatively low scores when compared with Fads from more distantly related fish lineages 360 (Betancur-R et al., 2013). Nevertheless, recent advances in functional analyses of fish Fads 361 have concluded that some Fads2 have acquired novel functions (subfunctionalisation) during 362 evolution and thus phylogeny of fish Fads2 does not necessarily correlate with their 363 functionalities (Castro et al., 2016). The herein reported functions of the C. gariepinus Fads2 364 further confirm such a conclusion. 365

Functional characterisation demonstrated that the C. gariepinus Fads2 is a bifunctional 366 $\Delta 6\Delta 5$ desaturase able to operate towards a range of substrates including n-3 (18:3n-3 and 367 20:4n-3) and n-6 (18:2n-6 and 20:3n-6) PUFA. Similar substrate specificities were previously 368 described in *D. rerio*, which represented the first ever report of dual $\Delta 6\Delta 5$ functionality in a 369 vertebrate Fads (Hastings et al., 2001). More recent studies have now shown that 370 bifunctionality appear to be a more common feature of fish Fads2 than originally thought. 371 Thus dual $\Delta 6\Delta 5$ Fads have been described in S. canaliculatus (Li et al., 2010), Nile tilapia 372 (Oreochromis niloticus) (Tanomman et al., 2013) and C. estor (Fonseca-Madrigal et al., 373

2014). Interestingly, fish Fads2 with $\Delta 4$ capability reported in S. canaliculatus (Li et al., 374 2010), S. senegalensis (Morais et al., 2012) and C. striata (Kuah et al., 2015) showed as well 375 some minor $\Delta 5$ activity and can thus be regarded as dual $\Delta 5\Delta 4$ Fads (Castro et al., 2016). In 376 contrast, other teleost Fads2 are single function $\Delta 6$ desaturases (González-Rovira et al., 2009; 377 Mohd-Yusof et al., 2010; Monroig et al., 2010; 2013; Zheng et al., 2009), in agreement with 378 Fads activities reported in mammalian FADS2 (Castro et al., 2016). Such substrate plasticity 379 exhibited amongst fish Fads2 is believed to be the result of a combination of multiple 380 evolutionary drivers including habitat, trophic level and ecology underlying the specific 381 382 phylogenetic position of each fish species (Castro et al., 2016, 2012; Li et al., 2010; Monroig et al., 2011b). In contrast, Fads1, another "front-end" Fads encoding a $\Delta 5$ Fads in mammals 383 (Castro et al., 2012, 2016), appears to have been lost during evolution of teleost and is absent 384 385 in the vast majority of farmed fish species (Castro et al., 2016).

The C. gariepinus Fads2 also exhibited $\Delta 8$ desaturation capability, an intrinsic feature of 386 vertebrate Fads2 (Monroig et al., 2011a; Park et al., 2009). Although conversions in yeast 387 might quantitatively vary from those occurring in vivo, it appeared that the C. gariepinus 388 Fads2 had lower efficiency as $\Delta 8$ Fads than as $\Delta 6$ Fads, in agreement with the " $\Delta 8$ pathway" 389 being regarded as a minor pathway compared to the more prominent $\Delta 6$ desaturation pathway 390 391 (Monroig et al., 2011a; Park et al., 2009). Interestingly, the $\Delta 8$ desaturation capabilities of C. gariepinus Fads2 towards 20:3n-3 (12.9 %) was relatively high leading to lower $\Delta 6\Delta 8$ ratio 392 (3.26), a parameter used to compare $\Delta 8$ desaturation capability among fish Fads2 enzymes 393 (Monroig et al., 2011a). Thus, the $\Delta 6/\Delta 8$ ratio of C. gariepinus Fads2 is more similar to that 394 of marine species like gilthead seabream (Sparus aurata) (2.7) and turbot (Psetta maxima) 395 (4.2). Whereas it is notably lower than those of freshwater or salmonid Fads2 including D. 396 rerio (22.4) and S. salar (12 and 14.7 for Fad b and Fad c, respectively) (Monroig et al., 397 2011a). These results suggest that the $\Delta 8$ pathway, while possibly not to such an extent as the 398

399 $\Delta 6$ pathway, can still contribute to the initial steps of LC-PUFA biosynthesis in *C*. 400 *gariepinus*. Note that $\Delta 8$ activity introduces the same double bond as $\Delta 6$ activity, after 401 elongation rather than before, and so a Fads having $\Delta 6\Delta 8$ activity is not regarded as 402 "bifunctional".

The ability of the C. gariepinus Fads2 to desaturate a range of $\Delta 5$, $\Delta 6$ and $\Delta 8$ Fads 403 substrates from both n-3 and n-6 series clearly shows it is a multifunctional enzyme. This is 404 emphasised by the stepwise desaturation reactions that occurred when transgenic yeast 405 expressing the C. gariepinus Fads2 were grown in the presence of certain FA substrates such 406 as 20:3n-3 and 20:2n-6. C. gariepinus Fads2 enzyme activity toward 20:3n-3 led to the 407 408 production of either 20:4n-3 ($\Delta 8$ desaturation) that was subsequently desaturated to 20:5n-3 ($\Delta 5$ desaturation), or the non-methylene interrupted (NMI) FA products $^{\Delta 5,11,14,17}20:4$ or 409 $^{\Delta6,11,14,17}$ 20:4 resulting from direct $\Delta5$ or $\Delta6$ desaturation, respectively. While the biological 410 significance of these pathways is difficult to determine, particularly for NMI FA biosynthesis, 411 the results further confirm that all the Fads capabilities ($\Delta 5$, $\Delta 6$ and $\Delta 8$) are present in the 412 characterised Fads2. 413

Moreover, we can further confirm that all the elongase activities required in the LC-PUFA 414 biosynthesis pathways also exist in C. gariepinus. Agaba et al. (2005) characterised an Elov15 415 from C. gariepinus that, like the vast majority of fish Elov15 investigated to date, showed C_{18} 416 and C₂₀ PUFA as preferred substrates, with markedly lower affinity towards C₂₂ substrates 417 (Castro et al., 2016). In contrast, the C. gariepinus Elovl2 showed higher elongation 418 efficiencies towards C₂₀ and C₂₂ PUFAs compared to C₁₈ substrates. Generally, these results 419 are consistent with the activities shown by the only three fish Elovl2 enzymes characterised 420 to date, i.e. S. salar, D. rerio and O. mykiss (Gregory and James, 2014; Monroig et al., 2009; 421 Morais et al., 2009). Although, similar to the human orthologue, the latter did not show any 422 activity on C₁₈ FA substrates (Leonard et al., 2002). The presence of Elov12 and particularly 423

its ability to elongate C₂₂ PUFA to a greater extent compared to Elov15 elongases has been 424 acknowledged as evidence supporting LC-PUFA biosynthetic capability in freshwater species 425 and salmonids (Morais et al., 2009). The plethora of genomic and transcriptomic sequences 426 currently available from a varied range of fish species and lineages strongly suggests that, 427 rather than the habitat (freshwater versus marine) of fish, it is the phylogeny of each species 428 that actually correlates with the presence or absence of Elovl2 within their genomes. Here we 429 430 show that marine species such as the Atlantic herring Clupea harengus (Fig. 2) possess a putative Elovl2, whereas freshwater species including O. niloticus or medaka (Oryzias 431 432 *latipes*) appear to have lost Elov12 from their genomes.

The functions of the herein reported Fads2 and Elovl2, together with the previously 433 characterised Elov15 (Agaba et al., 2005), allow us to predict the biosynthetic pathways of 434 LC-PUFA in C. gariepinus (Fig. 6). Thus, the dual $\Delta 6\Delta 5$ Fads2 catalyses the initial 435 desaturation of 18:3n-3 and 18:2n-6 ($\Delta 6$ desaturation), as well as the desaturation of 20:4n-3 436 and 20:3n-6 (Δ 5 desaturation). Although we cannot confirm whether the *C. gariepinus* Fads2 437 can desaturate 24:5n-3 and 24:4n-6 ($\Delta 6$ desaturation) required to synthesise 22:6n-3 and 438 22:5n-6, respectively, through the so-called Sprecher pathway (Sprecher et al., 2000). Such 439 ability of vertebrate Fads2 has been demonstrated in O. mykiss, S. salar and D. rerio (Bell 440 441 and Tocher, 2009; Buzzi et al., 1996; Tocher et al., 2003). Further studies will aim to elucidate whether the newly cloned Fads2 or other Fads potentially co-existing in the C. 442 gariepinus genome, have the ability to desaturate C_{24} PUFA in position $\Delta 6$. The Elovl2 was 443 able to catalyse the elongation of C₁₈ (18:3n-3, 18:2n-6, 18:4n-3 and 18:3n-6), C₂₀ (20:5n-3 444 and 20:4n-6) and C₂₂ (22:5n-3 and 22:4n-6) PUFA. Its activity towards C₁₈ PUFA was 445 however very low compared to activity towards C₂₀ and C₂₂ PUFA. This, together with the 446 activity of Elov15, which is high towards C₁₈ and C₂₀ PUFA (Agaba et al., 2005), confirm that 447

the activities required to catalyse all the elongation steps required for LC-PUFA synthesis arepresent in *C. gariepinus*.

Expression analysis showed *fads2*, *elovl2* and *elovl5* were expressed in all tissues analysed. 450 Consistent with the vast majority of freshwater species studied, the tissue distribution patterns 451 of C. gariepinus fads2 and elovl2 mRNAs showed liver as a major metabolic site for LC-452 PUFA biosynthesis. In contrast, marine fish species typically have brain as the tissue with 453 highest expression levels of LC-PUFA biosynthesis genes, with production of DHA from 454 EPA in brain being hypothesised as driving the retention of at least part of the LC-PUFA 455 biosynthetic pathway in species with high inputs of dietary LC-PUFA (Monroig et al., 456 457 2011b). An exception to this pattern is represented by the Nile tilapia fads2, with highest expression in the brain (Tanomman et al., 2013). C. gariepinus fads2 expression in liver was 458 approximately four-fold greater than in intestine, in contrast to salmonid *fads2* that have been 459 reported to be most highly expressed in intestine (Zheng et al., 2005). The expression of 460 elov15 was also high in liver but was highest in the intestine. 461

In conclusion, we have successfully cloned and characterised *fads2* and *elovl2* genes that encode enzymes with a broad range of substrate specificities from *C. gariepinus*. These two enzymes, and the previously reported Elov15, enable the African catfish *C. gariepinus* to carry out all the desaturation and elongation reactions required for endogenous LC-PUFA synthesis from C_{18} precursors, namely ALA and LA. These results strongly suggest that *C. gariepinus* has the ability to effectively utilise VO rich in C_{18} PUFA to satisfy essential LC-PUFA requirements.

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630 Solea senegalensis Fads2 AEQ92868.1 84 61 Chirostoma estor Fads2 AHX39206.1 631 31 Oreochromis niloticus Fads2 XP 005470690.1 Siganus canaliculatus Fads2 ABR12315.2 41 632 Rachvcentron canadum Fads2 ACJ65149.1 72 Scophthalmus maximus Fads2 AAS49163.1 bP Sparus aurata Fads2 ADD50000.1 633 Nibea mitsukurii Fads2 AX54437.1 57 98 Larimichthys crocea Fads2 NP_001290292.1 100 634 82 Salmo salar Fads2 NP 001165752.1 0.1 Gadus morhua Fads2 AAY46796.1 60 Clupea harengus Fads2 XP 012687541.1 635 Danio rerio Fads2 AF309556.1 Astyanax mexicanus Fads2 XP 007235183.1 636 79 Pangasianodon hypophthalmus Fads2 AFN21428.1 100 100 Clarias macrocephalus Fads2 AGR45589.1 100 637 100 - Clarias gariepinus Fads2 KU925904 Callorhinchus milli Fads2 XP 007885636.1 92 Scyliorhinus canicula Fads2 AEY94455.1 98 638 Latimeria chalumnae Fads2 XP 005988034.2 Gallus gallus FADS2 NP 001153900.1 99 639 Mus musculus FADS2 NP_062673.1 00 Homo sapiens FADS2 AAD20018.1 100 640 - Bos taurus Fads2 NP 001076913 88 Scyliorhinus canicula Fads1 AEY94454.1 Gallus gallus FADS1 XP_421052.4 641 98 Homo sapiens FADS1 AAF29378.1 82 Rattus norvegicus FADS1 AAG35068.1 100 642 100 Mus musculus FADS1 NP 666206.1 Mortierella alpina Fad AF465281 643

Fig. 1. Phylogenetic tree comparing the deduced amino acid sequence of *Clarias gariepinus fads2* with Fads from a range of vertebrates. The tree was constructed using the neighbourjoining method (Saitou and Nei, 1987) with the MEGA 4.0 software. The numbers represent the frequency (%) with which the tree topology presented was replicated after 10,000 iterations.



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Fig. 2. Phylogenetic tree comparing the deduced amino acid (aa) sequence of *Clarias gariepinus elovl2* with Elovl2, Elovl4 and Elovl5 from a range of vertebrates. The tree was constructed using the neighbour-joining method (Saitou and Nei, 1987) with the MEGA 4.0 software. The numbers represent the frequencies (%) with which the tree topology presented was replicated after 10,000 iterations.



Fig. 3. Functional characterisation of the newly cloned Clarias gariepinus Fads2 in yeast 686 (Saccharomyces cerevisiae). The fatty acid (FA) profiles of yeast transformed with pYES2 687 containing the coding sequence of *fads2* were determined after the yeast were grown in the 688 presence of one of the exogenously added substrates 18:3n-3 (A), 20:3n-3 (B), 20:4n-3 (C) 689 and 22:5n-3 (D). Peaks 1-4 represent the S. cerevisiae endogenous FA, namely 16:0 (1), 16:1 690 isomers (2), 18:0 (3) and 18:1n-9 (4). Additionally, peaks derived from exogenously added 691 substrates (*) or desaturation products are indicated accordingly. The peak indicated as 692 "20:4"" is a non-methylene interrupted FA ($^{\Delta 6,11,14,17}$ 20:4 or $^{\Delta 5,11,14,17}$ 20:4) (panel B). 693



Fig. 4. Functional characterisation of the of the newly cloned *Clarias gariepinus* Elovl2 in yeast (*Saccharomyces cerevisiae*). The fatty acid (FA) profiles of yeast transformed with pYES2 containing the coding sequence of *elovl2* were determined after the yeast were grown in the presence of one of the exogenously added substrates 18:3n-3 (A), 18:4n-3 (B), 20:5n-3 (C) and 22:5n-3 (D). Peaks 1-4 represent *S. cerevisiae* endogenous FAs namely 16:0 (1), 16:1 (2), 18:0 (3) and 18:1n-9 (4). Additionally, peaks derived from exogenously added substrates (*) or elongation products are indicated accordingly.



Fig. 5. Tissue distribution of fads2, elovl2 and elovl5 transcripts in Clarias gariepinus. Expression levels quantified for each transcript were normalised expression levels of the reference gene (28s rRNA) of the same tissue. The data are reported as mean values with their standard errors (n = 4). Within each target gene, different letters indicate statistically significant differences between expression levels (ANOVA and Tukey's HSD post hoc tests).





Fig. 6. The biosynthesis pathway of long-chain polyunsaturated fatty acids ($\leq C_{24}$) from α-linolenic (18:3n-3) and linoleic (18:2n-6) acids in *Clarias gariepinus*. Enzymatic activities shown in the scheme are predicted from heterologous expression in yeast of the herein investigated $\Delta 6\Delta 5$ fatty acyl desaturase 2 ($\Delta 6\Delta 5$ Fads2) and Elov12 elongase, and the previously reported Elov15 (Abaga et al., 2005). β-ox, partial β-oxidation; Elov1, fatty acyl elongase; Fads, fatty acyl desaturase.

* Enzymatic activities not yet demonstrated in *C. gariepinus*.

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749 Tables

750 Table 1

751 Sequences of primers used for cDNA cloning and tissue expression analysis (qPCR) of

752 *Clarias gariepinus fads2* and *elovl2*. Restriction sites *Bam*HI and *Xho*I are underlined.

Name	Direction	Sequence						
Initial cDNA cloning								
FadCGF2F1	Forward	5'-ATGGGCGGCGGAGGACAC-3'						
FadCGF2R1	Reverse	5'-GCATCTAGCCACAGCTCACC-3'						
EloCGE2F1	Forward	5'-TACTTGGGACCAAAGTACATGA-3'						
EloCGE2R1	Reverse	5'-AGATAGCGTTTCCACCACAG-3'						
5' RACE PCR								
FadCGRF2R2	Reverse	5'-CGATCACAACCCACTGATCA-3'						
FadCGRF2R3	Reverse	5'-CGTCCTCCAGGATGTCTTTT-3'						
EloCGRE2R3	Reverse	5'-AGCTTGCTGAAATAAGCTCCACT-3'						
EloCGRE2R2	Reverse	5'-TGTAGAAGGACAGCATGGTGAC-3'						
3' RACE PCR								
FadCGRF2F1	Forward	5'-CAGTCGCCATTCAACGATT-3'						
FadCGRF2F2	Forward	5'-GAACACCATCTCTTTCCCATG-3'						
EloCGRE2F1	Forward	5'-TTGTCCACCATTCCTTCAATG-3'						
EloCGRE2F2	Forward	5'-ACTGAACAGCTTCATCCATGTG-3'						
ORF cloning								
FadCGF5UF1	Forward	5'-AGAGGAGCGCAGTGATGAG-3'						
FadCGF3UR1	Reverse	5'-GTGGGAATTACAGAATTGTTATGG-3'						
FadCGFVF	Forward	5'-CCC <u>GGATCC</u> AAGATGGGCGGCGGAGGAC-3'						
FadCGFVR2	Reverse	5'-CCG <u>CTCGAG</u> TTATTTGTGGAGGTATGCATC-3'						
EloCGE2VF	Forward	5'-CCC <u>GGATCC</u> AACATGGATTTTATTGTGAAGAA-3'						
EloCGE2VR	Reverse	5'-CCG <u>CTCGAG</u> TCACTGCAGCTTATGTTTGGC-3'						
EloCGE25UF	Forward	5'-CCAGTTACATTAAGAGGCACCG-3'						
EloCGE23UR	Reverse	5'-AGATTAGTCAACATGAAAGGTGAA-3'						
Quantitative PCR								
FadCGqF2F1	Forward	5'-TCCTATATGCTGGAACTAATGTGG-3'						
FadCGqF2R1	Reverse	5'-AGGATGTAACCAACAGCATGG-3'						
EloCGqE2F1	Forward	5'-GCAGTACTCTGGGCATTTGTC-3'						
EloCGqE2R1	Reverse	5'-GGGACATTGGCGAAAAAGTA-3'						
EloCGqE5F1	Forward	5'-ACTCACAGTGGAGGAGAGC-3'						
EloCGqE5R1	Reverse	5'-GGAATGGTGGTAAACGTGCA-3'						
28SrRNAF1	Forward	5'-GTCCTTCTGATGGAGGCTCA-3'						
28SrRNAR1	Reverse	5'-CGTGCCGGTATTTAGCCTTA-3'						

754 Table 2

Substrate conversions of *Saccharomyces cerevisiae* transformed with *Clarias gariepinus fads2* coding region and grown in the presence of exogenously added substrate (18:3n-3, 18:2n-6, 20:3n-3, 20:2n-6, 20:4n-3, 20:3n-6, 22:5n-3 or 22:4n-6). Conversions were calculated according to the formula [individual product peak area / (all products peak areas + substrate peak area)] \times 100.

FA substrate	FA Product	Conversion (%)	Activity
18:3n-3	18:4 n -3	42.0	$\Delta 6$
18:2n-6	18:3 n- 6	22.5	$\Delta 6$
20:3n-3	20:4n-3	12.9 ^a	$\Delta 8$
20:2n-6	20:3n-6	2.5 ^a	$\Delta 8$
20:4n-3	20:5n-3	18.7	$\Delta 5$
20:3n-6	20:4n-6	13.8	$\Delta 5$
22:5n-3	22:6n-3	Nd	$\Delta 4$
22:4n-6	22:5n-6	Nd	$\Delta 4$

^a Conversions of Δ8 substrates (20:3n-3 and 20:2n-6) by Fads2 include stepwise reactions
due to multifunctional desaturation abilities. Thus, the conversion rates of 20:3n-3 and 20:2n6 include the Δ8 desaturation toward 20:4n-3 and 20:3n-6, respectively, and their subsequent
Δ5 desaturations to 20:5n-3 and 20:4n-6, respectively.
FA, Fatty acid; Nd, not detected.

Table 3

Substrate conversions of *Saccharomyces cerevisiae* transformed with *Clarias gariepinus elovl2* and grown in the presence of PUFA substrate exogenously added (18:3n-3, 18:2n-6, 18:4n-3, 18:3n-6, 20:5n-3, 20:4n-6, 22:5n-3 or 22:4n-6). Conversions were calculated for each stepwise elongation according to the formula [peak areas of first products and longer chain products / (peak areas of all products with longer chain than substrate + substrate peak area)] x 100.

FA Product	Conversion (%)
20:3n-3	7.5
20:2n-6	3.0
20:4n-3	15.2
20:3n-6	20.5
22:5n-3	73.4
22:4n-6	56.0
24:5n-3	36.7
24:4n-6	9.7
	FA Product 20:3n-3 20:2n-6 20:4n-3 20:3n-6 22:5n-3 22:4n-6 24:5n-3 24:4n-6

FA, fatty acid