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Molecular and functional characterization of a fads2 orthologue in the Amazonian teleost, *Arapaima gigas*

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Abstract

The Brazilian teleost *Arapaima gigas* is an iconic species of the Amazon. In recent years a significant effort has been put into the farming of arapaima to mitigate overfishing threats. However, little is known regarding the nutritional requirements of *A. gigas* in particular those for essential fatty acids including the long-chain polyunsaturated fatty acids (LC-PUFA) eicosapentaenoic acid (EPA) and docosahexaenoic acid (DHA). The ability to biosynthesize LC-PUFA is dependent upon the gene repertoire of fatty acyl desaturases (Fads) and elongases (Elov1), as well as their fatty acid specificities. In the present study we characterized both molecularly and functionally an orthologue of the desaturase fatty acid desaturase 2 (*fads2*) from *A. gigas*. The isolated sequence displayed the typical desaturase features, a cytochrome b$_{5}$-domain with the heme-binding motif, two transmembrane domains and three histidine-rich regions. Functional characterization of *A. gigas fads2* showed that, similar to other teleosts, the *A. gigas fads2* exhibited a predominant Δ6 activity complemented with some capacity for Δ8 desaturation. Given that *A. gigas* belongs to one of the oldest teleostei lineages, the Osteoglossomorpha, these findings offer a significant insight into the evolution LC-PUFA biosynthesis in teleosts.
Introduction

Long-chain polyunsaturated fatty acids (LC-PUFA) play vital roles in numerous biological processes. They participate in structural functions as major components of biomembranes and are also involved in processes such as the inflammatory response, reproduction (Wall et al., 2010; Robinson and Mazurak, 2013), and neural development (Perica and Delaš, 2011) and can have beneficial effects in pathological conditions such as cardiovascular disease (Psota et al., 2006; Jump et al., 2012). LC-PUFA are often defined as compounds with 20 to 24 carbon atoms and three or more double bonds (unsaturations) and can be classified into two main groups: the omega-6 (ω6 or n-6) and the omega-3 (ω3 or n-3) LC-PUFA, based upon the position of the first double bond in relation to the methyl end carbon (CH₃) (Monroig et al., 2011a). LC-PUFA of the n-6 and n-3 series can be of dietary origin or, alternatively, they can be biosynthesized from dietary essential fatty acids (EFA) such as linoleic acid (LA, 18:2n-6) and α-linolenic acid (ALA, 18:3n-3), respectively, through a series of sequential biochemical reactions, mediated by elongation of very long-chain fatty acid protein (Elovl) and fatty acyl desaturases (Fads).

The ability to endogenously synthesize LC-PUFA from dietary fatty acids (FA) differs markedly among vertebrate species (Rivers et al., 1975; Bauer, 1997; Tocher, 2003; Burdge and Calder, 2005; Fonseca-Madrigal et al., 2014; Castro et al., 2016; Monroig et al., 2016a; Monroig et al., 2016b). This variation may be primarily attributed to differences in the eloavl and fads gene repertoire, as well as their associated fatty acid substrate specificities. For instance, mammals have several FADS genes of which FADS1 encodes a Δ5 desaturase and FADS2 encodes a desaturase with Δ6 preference, in addition to Δ4 activity reported in some mammals (Park et al., 2009; Park et al., 2015). In contrast, teleost fish examined to date have been found to possess exclusively FADS2 orthologues (Castro et al., 2012; Castro et al., 2016). However, while mammalian FADS enzymes are essentially mono-functional, mechanisms of bifunctionalization (i.e., acquisition of additional/alternative substrate specificities) have been described in several teleost Fads2. Thus, Fads2 with dual Δ6Δ5
desaturase activities have been described in *Danio rerio* (Hastings et al., 2001), *Siganus canaliculatus* (Li et al., 2010), *Oreochromis niloticus* (Tanomman et al., 2013), *Chirostoma estor* (Fonseca-Madrigal et al., 2014) and *Clarias gariepinus* (Oboh et al., 2016). In addition, *S. canaliculatus* and *C. estor* possess a duplicated *Fads2* that exhibit Δ4 desaturase activity (Li et al., 2010; Fonseca-Madrigal et al., 2014), a type of enzyme also found in *Solea senegalensis* (Morais et al., 2012) and *Channa striata* (Kuah et al., 2015). Moreover, in agreement with the abilities reported in the baboon Δ6-desaturase (Park et al., 2009), the majority of teleost *Fads2* desaturases have been demonstrated to possess the capability for Δ8 desaturation (Monroig et al., 2011b). Overall the complement of LC-PUFA biosynthetic enzymes, namely FADS and ELOVL, as well as their functionalities, dictates the ability of a species for the conversion of C18 PUFA (LA and ALA) into physiologically important LC-PUFA including arachidonic acid (ARA, 20:4n-6), eicosapentaenoic acid (EPA, 20:5n-3) and docosahexaenoic acid (DHA, 22:6n-3) (Bell and Tocher, 2009; Castro et al., 2016).

Importantly, the investigation of *Fads* and *Elov1* in fish has primarily focused on farmed species since both *Fads* and *Elov1* capabilities underpin the efficiency of these fish species to utilize the C18 PUFA present in vegetable oils (VO) currently used as sustainable replacements for dietary fish oils (FO) in aquafeeds (Tocher, 2010). Therefore a clear understanding of LC-PUFA biosynthesis pathways is critical to understand the potential limitations of farmed fish species and for the implementation of dietary strategies to fulfil essential requirements and ensure normal growth and development in captivity.

An iconic species of the Amazon, so-called “pirarucú” (*Arapaima gigas*), is one of the largest freshwater and air-breathing fishes in the world, and has been extensively fished since the 18th century (Verissimo, 1895; Goulding, 1980). In the early 1970’s over-exploitation of *A. gigas* led to its near extinction (Goulding, 1980) and listing in CITES (Convention on International Trade in Endangered Species of Wild Fauna and Flora). To overcome this threat, considerable effort has been put into developing the sustainable farming of this species. However, despite some important advances, critical knowledge in key areas such as
physiology and nutrition is still scarce in this species. Much of the published research on *A. gigas* has focused on the understanding and evolution of the air-breathing capacity (Brauner et al., 2004; Gonzalez et al., 2010), general health and aquaculture practices (Ribeiro et al., 2011; Bezerra et al., 2014) and, more recently, the potential use of *A. gigas* scales as biomaterials (Torres et al., 2015). In contrast, few studies have addressed the dietary requirements of *A. gigas* (Ituassú et al., 2005; Andrade et al., 2007; Ribeiro et al., 2011), stressing the need for a broader understanding of the metabolism of this carnivorous species.

Here, we describe the isolation and functional characterization of a cDNA from *A. gigas* orthologous to *fads2* desaturases, key enzymes in LC-PUFA biosynthetic pathways and crucial elements in determining EFA requirements in this species. The phylogenetic position of *A. gigas* within one of the most ancient teleost lineages, the Osteoglossomorpha, brings new insights into the evolution of the LC-PUFA biosynthesis cascade in both fish and vertebrates in general.

**Materials and Methods**

**Molecular cloning of the *A. gigas* fads gene**

Total RNA was extracted from a range of *A. gigas* tissues using the Illustra RNAspin Mini kit (GE Healthcare, UK). The RNA extraction process included an on-column DNase I treatment (provided in the kit). RNA integrity was assessed on a 1 % agarose TAE gel stained with GelRed™ nucleic acid stain (Biotium, Hayward, CA, USA). The Quant-iT™ RiboGreen® RNA Assay Kit (Life Technologies, Carlsbad, CA, USA) was used to measure total RNA concentration. Reverse transcription reactions were performed with the iScript cDNA Synthesis Kit (Bio-Rad, Hercules, CA, USA).

*Arapaima gigas* FADS gene was isolated in three main steps. First, degenerate primers targeting the Fads gene were designed using CODEHOP (Rose et al., 2003) available at http://blocks.fhcrc.org/codehop.html. The initial polymerase chain reaction (PCR) was
performed with a degenerate primer set and Flash High-Fidelity PCR Master Mix (Thermo Fisher Scientific, Waltham, USA), set for a final volume of 20 µl, with 500 nM of sense and antisense primers, and 1 µl of A. gigas cDNA pool (see Table 1 for primers, PCR conditions). In the second step, the partial \( fads \) sequence was further extended by Rapid amplification of cDNA ends (RACE) PCR using as template 5’ and 3’ RACE ready cDNA prepared with SMARTer™ RACE cDNA Amplification Kit (Clontech, CA, USA). Gene specific primers for RACE were designed using the previously isolated fragment and RACE PCR was performed with Flash High-Fidelity PCR Master Mix (Thermo Fisher Scientific) using 1 µl of gene specific primer combined with 2 µl Universal primer mix (Clontech) (see table 1 for primers and PCR conditions). The resulting 5’and 3’sequences were assembled to produce the full open reading frame (ORF) \( fads \)-like cDNA. In the final step, the full ORF of \( A. \) gigas FADS was isolated using 1 µl of A. gigas cDNA pool, and Flash High-Fidelity PCR Master Mix (Thermo Fisher Scientific, Waltham, USA), set for a final volume of 20 µl, with 500 nM of sense and antisense primers (see table 1 for primers and PCR conditions). In each step resulting PCR products were analysed in 1 % agarose gel, purified with NZYGelpure (NZYTech, Lisbon, Portugal) and confirmed by sequencing (GATC Biotech Constance, Germany). The final, full ORF sequence was translated and submitted to pFAM and NCBI for blastp searches retrieving Fads-like profile (Accession number: KX809739).

**Sequence collection, phylogenetic and 2D structural analysis**

Fads amino acid (aa) sequences were retrieved from Genbank and Ensembl (for accession numbers see Table 2). Sequences were aligned with MAFFT using the L-INS-i method (Katoh and Toh, 2008). The sequence alignment was stripped from all columns containing gaps leaving 374 gap-free sites for phylogenetic analysis. Maximum likelihood phylogenetic analysis was performed in PhyML v3.0 server (Guindon et al., 2010) using smart model selection resulting in LG +G+I+F, and branch support was calculated using 1000 bootstraps.
Using the same alignment a second Bayesian phylogenetic analysis was performed using MrBayes v3.2.3 available in CIPRES Science Gateway V3.3 (Miller et al., 2015). MrBayes was run for 1 million generations with the following parameters: rate matrix for aa=mixed, nruns=2, nchains=4, temp=0.2, sampling set to 500 and burin to 0.25. The resulting trees were visualized in Fig Tree V1.3.1 available at http://tree.bio.ed.ac.uk/software/figtree/ and rooted at mid-point. *A. gigas* aa sequence was submitted to TOPCONS web server for prediction of 2D topology, with all parameters set to default (http://topcons.net/) (Tsirigos et al., 2015), and results visualized using Potter web application (http://wlab.ethz.ch/protter) (Omasits et al., 2014).

**Yeast expression assays and fatty acid analysis**

The *A. gigas fads* ORF was isolated with two sequential PCR with Flash High-Fidelity PCR Master Mix (Thermo Fisher Scientific, USA) as described above. The first PCR was performed with an *A. gigas* cDNA pool and primers (AgigasFADS_ORF_F and AgigasFADS_ORF_R, Table 1) targeting the full ORF. The PCR product was diluted (1:50) and used as template for the second PCR performed with primers containing restriction sites for *KpnI* (AgigasFADS_pYES_KpnI_F) and *XbaI* (AgigasFADS_pYES_XbaI_R) (Table 1). The final PCR product was purified and digested with the appropriate restriction enzymes and cloned into the yeast expression vector pYES2 (Invitrogen, CA, USA). Transformation and culture of yeast *Saccharomyces cerevisiae* were conducted as previously described (Hastings et al., 2001; Agaba et al., 2004; Oboh et al., 2016). Briefly, transgenic yeast expressing the *A. gigas fads* ORF were grown in the presence of PUFA including Δ6 (18:3n-3 and 18:2n-6), Δ8 (20:2n-6 and 20:3n-3), Δ5 (20:4n-3; 20:3n-6) and Δ4 (22:5n-3 and 22:4n-6) desaturase substrates. PUFA substrates, added as sodium salts, were supplemented in the yeast medium at final concentrations of 0.5 mM (C18), 0.75 mM (C20) and 1.0 mM (C22) as uptake efficiency decreases with increasing chain length (Zheng et al., 2009). After 48 h of incubation, yeast were harvested, washed and total lipid extracted by homogenization in chloroform/methanol (2:1, v/v) containing 0.01 % BHT (Monroig et al., 2013). Fatty acyl methyl esters (FAME)
were prepared from total lipids extracted from harvested cells and identified based on GC retention times and confirmed by GC-MS as described previously (Hastings et al., 2001; Li et al., 2010). FA desaturation efficiencies from exogenously added PUFA substrates were calculated by the proportion of substrate FA converted to a desaturated product as (product area/(product area + substrate area)) x 100.

Results

Sequence conservation and topology prediction

The isolated A. gigas sequence was translated and submitted to BLASTp and to PFam to validate the fads-like profile and identify the main protein domains. BLASTp searches showed that the A. gigas sequence had highest identity scores with fads2 desaturases from other teleost species (results not shown), while the PFam search identified two main domains typical of Fads enzymes: a cytochrome b$_5$-like heme/steroid binding domain (15 - 88 aa) and FA desaturase domain (150 - 412 aa). To further characterize, the A. gigas Fads-like protein was aligned with four known and fully characterized Fads aa sequences from D. rerio (NCBI Protein accession no NP_001117047.1), Salmo salar (NCBI Protein accession no NP_004256.1) and O. niloticus (NCBI Protein accession no AGV52807.1) and Homo sapiens (NCBI Protein accession no NR_033434.1) (Fig. 1A). The A. gigas sequence showed highest degree of pairwise identity with the S. salar Fads2 (86.1 %), followed by Fads2 from O. niloticus (83.9 %), D. rerio (82.8 %) and H. sapiens (79.3 %), revealing a high degree of cross-species conservation. Additionally, using H. sapiens FADS2 sequence as a reference, several sequence signature motifs of Fads enzymes were identified: the heme binding motif HPGG and three histidine boxes HXXXH, HXXHH and QXXHH, which are presumed to form the Fe-binding active center of the enzyme (Los and Murata, 1998; Pereira et al., 2003) (Fig. 1A). The heme binding motif was totally conserved in Fads from all species analyzed including A. gigas. In the first histidine box two distinct patterns were observed: HDYGH in H. sapiens and S. salar, while A. gigas, D. rerio and O. niloticus showed the signature
HDGFH with the replacement of a tyrosine (Y) by a phenylalanine (F) (Fig. 1A). In the
second histidine box, all analyzed species presented HFQHH with the exception of O. niloticus, whose Fads2 presents HFRRHH (Fig. 1A). Full conservation of the third histidine box was found across all the analyzed species.

Regarding the 2D topology prediction, all calculation methods were consistent in predicting that A. gigas Fads-like displayed four membrane spanning domains, and that the N- and the C-terminals, as well as the three histidine motifs, were oriented towards the cytosol (Supplementary Material 1). Interestingly, the residues involved in regioselectivity were localized at the base of the third membrane spanning domain (Fig. 1B). The topology predicted for the A. gigas Fads2 was thus consistent with the structural organization proposed in previous reports for other Fads-like desaturases (Los and Murata, 1998; Meesapyodsuk et al., 2007; Lim et al., 2014).

Phylogenetic analysis of Fads-like ORF from A. gigas

Two phylogenetic analyses were conducted using the same data set consisting of aa sequence alignment between the newly cloned A. gigas putative Fads with FADS1 and FADS2 desaturase sequences from eighteen vertebrate species (mammals - H. sapiens, M. domestica birds – G. gallus, reptiles - A. sinensis, coelacanth - L. chalumnae, teleosts - G. morhua, T. maccocyii, O. niloticus, S. salar, and D. rerio, chondrichthyans - S. canicula, C. milii and one invertebrate (B. floridae). In both cases the tree topology showed two well-supported clades, one corresponding to the FADS1 and the second corresponding to the FADS2, being both trees out grouped by invertebrate FADS from B. floridae. The A. gigas Fads-like sequence strongly grouped (930 bootstraps or 1 posterior probabilities) within the teleost group composed of all Fads2 sequences. Out grouping the teleost clade we find tetrapod and chondrichthyans Fads2 desaturases, indicating that the A. gigas putative Fads is a true fads2 orthologue. However, desaturases with different substrate preferences, for example D. rerio and O. niloticus Fads2 that are bifunctional Δ6Δ5 desaturases (Hastings et
al., 2001; Tanomman et al., 2013), and G. morhua and S. salar Fads2 that have been reported as unifunctional Δ6 desaturases (Zheng et al., 2005; Monroig et al., 2010) were found within the teleost clade.

Functional analysis of Fads2 in A. gigas

Functional characterization of the A. gigas desaturase was performed with using a well-established heterologous system consisting of yeast S. cerevisiae expressing the ORF of the A. gigas fads2 and grown in the presence of potential desaturase PUFA substrates (Hastings et al., 2001; Agaba et al., 2004; Fonseca-Madrigal et al., 2014). FA profile of yeast transformed with the empty pYES2 plasmid (control) consisted of the yeast endogenous FA including 16:0, 16:1 isomers (16:1n-9 and 16:1n-7), 18:0, and 18:1 isomers (18:1n-9 and 18:1n-7) and whichever exogenously PUFA substrate was added (data not shown). These results confirmed that the yeast endogenous enzymes were not active on the exogenously added PUFA substrates (Agaba et al., 2005). On the other hand, yeast transformed with the ORF of the A. gigas fads2 showed additional peaks when grown in the presence of 18:3n-3, 18:2n-6, 20:3n-3 and 20:2n-6 (Fig. 3). Thus, transgenic yeast expressing the fads2 had the ability to desaturate 18:3n-3 and 18:2n-6 to 18:4n-3 (Fig. 3A) and 18:3n-6 (Fig. 3B), respectively, showing this enzyme has Δ6 desaturase activity. Moreover, transgenic yeast supplemented with 20:3n-3 and 20:2n-6 produced additional peaks identified as 20:4n-3 (Fig. 3C) and 20:3n-6 (Fig. 3D), respectively, showing that the A. gigas fads2 had also Δ8 desaturase activity. Therefore, the data confirmed that the cloned A. gigas fads2 encoded an enzyme with Δ6 and Δ8 desaturase specificities. Conversions obtained in the yeast expression system suggested that the A. gigas Fads2 has Δ6 as the most prominent activity and a preference for n-3 fatty acid substrates compared with n-6 substrates for each homologous FA substrate pair (Δ6 or Δ8) considered (Table 3). Neither Δ5 nor Δ4 activities were detected in yeast (Fig. 3E-H).
Discussion

Fads are, together with Elovl, key enzymes in LC-PUFA biosynthetic pathways (Castro et al., 2016; Monroig et al., 2016b). The sequential and concerted action of both enzymes defines the ability of a given species to endogenously synthesize physiologically relevant LC-PUFA including ARA, EPA or DHA (Bell and Tocher, 2009). The investigation of the molecular components of LC-PUFA biosynthetic pathway in fish has been an active field of research over the last decade (Agaba et al., 2005; Zheng et al., 2009; Monroig et al., 2011b; Castro et al., 2012; Monroig et al., 2012; Carmona-Antonanzas et al., 2013; Castro et al., 2016). This is particularly true in farmed fish species where a full understanding of LC-PUFA biosynthesis capacities is crucial to successfully grow fish on diets that are necessarily being formulated with ever-increasing levels of VO (rich in C₁₈ PUFA but devoid of LC-PUFA) as primary lipid sources to replace FO (Turchini et al., 2009). Overall, these studies have highlighted a surprisingly diverse and interesting pattern among Fads substrate specificities (Fonseca-Madrigal et al., 2014).

The primary objective of the present study was the molecular cloning and functional characterization of a desaturase of the Amazonian teleost A. gigas. This freshwater species with aquaculture potential (Cavero et al., 2003) has been barely investigated in terms of nutritional requirements. In addition, A. gigas belongs to the Osteoglossiformes, a teleost order that has been considered to be the most basal of living teleosts (Nelson, 1994), therefore bringing a fresh perspective on the functional diversification of the desaturases in teleosts. The isolated Fads2 sequence of A. gigas showed all the typical features of fatty acyl (also known as “front-end”) desaturases when subjected to BLASTp and to PFam searches.

Furthermore, detailed sequence alignment analysis revealed that the unique structure of Fads-like enzymes was preserved in A. gigas Fads2 that contained three highly conserved histidine boxes, as well as the heme motif within the cytochrome b₅-like domain, which are considered to be involved in the formation of the desaturase catalytic centre (Shanklin et al., 1994; Los and Murata, 1998; Tocher et al., 1998). The 2D topology analysis of A. gigas Fads2 predicted
four transmembrane domains TM1: 124-145, TM2: 151-172, TM3: 258-279, TM4: 300-321, that oriented the three histidine boxes and the cytochrome b6-like domain to the cytosol, consistent with the structural organization proposed in previous reports (Los and Murata, 1998; Meesapyodsuk et al., 2007; Lim et al., 2014). Among the three histidine boxes, two distinct patterns were observed in the first histidine box in the Fads2, with *A. gigas*, *D. rerio* and *O. niloticus* having the signature HDEFGH, whereas a replacement of a phenylalanine (F) by tyrosine (Y) occurs for *H. sapiens* and *S. salar* Fads2. This replacement was predicted to not affect the mandatory/canonical histidine residues within each box. Additionally the abovementioned aa substitution was not expected to have any major functional impact, possibly due to the fact that these two aa residues share very similar biochemical properties (Betts and Russell, 2003). In contrast, differences were found within the residues previously proposed to participate in the regioselectivity of these enzymes (Hsa:279Phe - 282Gln; Dre:279Phe - 282Gln, Oni: 280Phe - His283, Ssa: 289Phe-292Gln; Agi: 273Phe - 276Gln) (Meesapyodsuk et al., 2007; Lim et al., 2014), possibly accounting for the different Fads activities observed in these species.

All *fads* characterized so far from teleosts are orthologous to *FADS2*, which performs primarily Δ6 desaturations in mammals (Guillou et al., 2010). This is further supported by the herein phylogenetic analysis of *A. gigas fads*, together with phylogenetic analyses reported previously (Zheng et al., 2004; Monroig et al., 2011b; Liu et al., 2014). However, the teleost Fads exhibit a wide range of PUFA specificities (Hastings et al., 2001; Hastings et al., 2004; Li et al., 2010; Monroig et al., 2012; Xie et al., 2014), underscoring a “*functional plasticity*” that has been previously attributed as a consequence of adaptation to availability of LC-PUFA in variable habitats and trophic levels (Tocher, 2010; Monroig et al., 2011b; Castro et al., 2012; Monroig et al., 2012; Fonseca-Madrigal et al., 2014). Thus, Fads2 with dual Δ6Δ5 activity have been cloned from *D. rerio* (Hastings et al., 2001), *S. canaliculatus* (Li et al., 2010), *O. niloticus* (Tanomman et al., 2013), *C. estor* (Fonseca-Madrigal et al., 2014), and *C. gariepinus* (Oboh et al., 2016). Moreover, teleost Fads2 with Δ4 desaturase activity have been
found in *S. canaliculatus* (Li et al., 2010), *S. senegalensis* (Morais et al., 2012) and *C. striata* (Kuah et al., 2015). Interestingly, the human *FADS2* gene product has been recently demonstrated to have the ability for direct Δ4 desaturation of 22:5n-3 to 22:6n-3 (Park et al., 2015). Nevertheless, the majority of functionally characterized teleost Fads2 are essentially Δ6 desaturase enzymes as reported in a variety of teleost fish species including gilthead seabream, rainbow trout, Atlantic salmon (three genes), turbot, cobia, European seabass, barramundi, black seabream, nibe croaker, Northern bluefin tuna, meagre, Japanese eel and orange spotted grouper (Castro et al., 2016). In agreement, the *A. gigas* Fads2 was demonstrated to be a Δ6 desaturase able to convert 18:3n-3 and 18:2n-6 to 18:4n-3 and 18:3n-6, respectively.

However, in addition, the *A. gigas* Fads2 showed capability for Δ8 desaturation, since it was capable of converting both 20:3n-3 and 20:2n-6 into 20:4n-3 and 20:3n-6, respectively. This activity was first reported in the baboon FADS2 (Park et al., 2009) and subsequently described in a range of fish Fads2 enzymes (Monroig et al., 2011b). The capability for Δ8 desaturation appears widespread in Fads2 characterized from fish (Monroig et al., 2011b; Monroig et al., 2013; Wang et al., 2014; Kabeya et al., 2015; Oboh et al., 2016), with few exceptions represented by the Atlantic salmon and rainbow trout Δ5 Fads2, as well as the striped snakehead Δ4 Fads2 (Monroig et al., 2011b; Kuah et al., 2015; Abdul Hamid et al., 2016). Interestingly, it appeared that, generally, Fads2 from marine teleosts had relatively high Δ8 desaturase ability compared to their freshwater and salmonid counterparts (Monroig et al., 2011b). Consequently, the Δ6:Δ8 desaturation ratio varies among teleost Fads2, with marine species having relatively low Δ6:Δ8 ratios, while freshwater and salmonid species having higher Δ6:Δ8 ratios. The *A. gigas* Fads2 had a Δ6:Δ8 ratio of 4.4 for n-3 PUFA substrates (25.8 : 5.8), and thus more within the range of marine teleosts such as turbot (4.2) or gilthead seabream (2.7) and far from freshwater species like rainbow trout (91.5) and zebrafish (22.4). While it is unclear what the evolutionary drivers are for the high capacity for Δ8 desaturation in *A. gigas* Fads2, having a Fads2 with the ability to operate as a Δ6
desaturase on ALA and LA, and as a Δ8 on 20:3n-3 and 20:2n-6, may confer an advantage to
this species enabling production of 20:4n-3 and 20:3n-6, respectively, through two different
pathways. Both 20:4n-3 and 20:3n-6 are substrates of Δ5 desaturase, an enzyme that, despite
being absent in the vast majority of teleosts, is likely to be retained in basal teleosts such as
Osteoglossidae, the family to which A. gigas belongs. In fact, a close relative to A. gigas, the
Asian arowana (Scleropages formosus) also a basal teleost belonging to the Osteoglossidae,
presents two predicted Fads-like sequences recently deposited in GenBank KPP61181.1 and
KPP71333.1 (not included in phylogenetic analysis due to their partial nature) annotated as
FADS2-like and delta 6 desaturase-like respectively. However, no functional characterization
these genes is yet available. Further studies are required to fully confirm the presence or
absence of Fads1 in basal teleost lineages.

In conclusion, we herein demonstrate that A. gigas possess a fads2 gene with all the typical
features of front-end desaturases. Moreover, the functional assays of the A. gigas Fads2 in
yeast confirmed that, like the majority of teleost Fads2, the A. gigas orthologue exhibited Δ6
and Δ8 desaturase activities. Along with the Fads2 from the Japanese eel (Wang et al., 2014),
the herein reported A. gigas represents the most ancient representative of the Fads gene family
being investigated within the teleost clade.

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Table 2. Accession number of sequences used phylogenetic analysis.

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<th>Species</th>
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<td>GGA- <em>Gallus gallus</em></td>
<td>NP_001153900.1 XP_421052.4</td>
</tr>
<tr>
<td>LCH- <em>Latimeria chalumnae</em></td>
<td>XP_005988034.1 XP_005988035.1</td>
</tr>
<tr>
<td>CMI - <em>Callorhinchus milii</em></td>
<td>XP_007885636.1 XP_007885635.1</td>
</tr>
<tr>
<td>SCA- <em>Scyliorhinus canicula</em></td>
<td>AEY94455.1 -</td>
</tr>
<tr>
<td>DRE- <em>Danio rerio</em></td>
<td>NP_571720.2    -</td>
</tr>
<tr>
<td>SSA- <em>Salmo salar</em></td>
<td>NP_001117047.1 -</td>
</tr>
<tr>
<td>ONI- <em>Oreochromis niloticus (a)</em></td>
<td>XP_005470661.1 -</td>
</tr>
<tr>
<td>ONI- <em>Oreochromis niloticus (b)</em></td>
<td>XP_003440520.1 -</td>
</tr>
<tr>
<td>TMA - <em>Thunnus maccoyii</em></td>
<td>ADG62353.1 -</td>
</tr>
<tr>
<td>GMO - <em>Gadus morhua</em></td>
<td>AAY46796 -</td>
</tr>
<tr>
<td>BFL - <em>Branchiostoma floridae</em></td>
<td>XP_002586930.1 -</td>
</tr>
</tbody>
</table>
### Table 1. Primer sets and corresponding PCR conditions.

<table>
<thead>
<tr>
<th>Primer set function</th>
<th>Primer name</th>
<th>Primer sequence</th>
<th>Initial denaturation</th>
<th>Cycles</th>
<th>Denaturation</th>
<th>TM</th>
<th>Extension (size bp)</th>
<th>Final extension</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Degenerate primers</strong></td>
<td>FADS2degen_F</td>
<td>GCGCCTCGCCAAYtggtaayc</td>
<td>98°C /10s</td>
<td>40</td>
<td>98°C /1s</td>
<td>54°C/5s</td>
<td>72°C/10s</td>
<td>72°C/1min</td>
</tr>
<tr>
<td></td>
<td>FADS2degen_R</td>
<td>TGGCCGGAGAAACcartcrra</td>
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<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td><strong>Gene specific Race primers</strong></td>
<td>3RC_AgigasFADS_F</td>
<td>ACCTAAAGGGTGCTTCAGCCAACT</td>
<td>98°C /10s</td>
<td>20</td>
<td>98°C /1s</td>
<td>62°C/5s</td>
<td>72°C/15s</td>
<td>72°C/1min</td>
</tr>
<tr>
<td></td>
<td>5RC_AgigasFADS_R</td>
<td>GTTCGGAACACGCCCTCTTTCCT</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Nested Gene specific Race Primers</strong></td>
<td>N3RC_AgigasFADS_F</td>
<td>GTTTCTGGAGAGCCACTGTTTGT</td>
<td>98°C /10s</td>
<td>35</td>
<td>98°C /1s</td>
<td>62°C/5s</td>
<td>72°C/8s</td>
<td>72°C/1min</td>
</tr>
<tr>
<td></td>
<td>N5RC_AgigasFADS_R</td>
<td>CTGCCTTTCTGCGGTCTAAG</td>
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</tr>
<tr>
<td><strong>Full ORF</strong></td>
<td>AgigasFADS_ORF_F</td>
<td>ATATGCGAGGATGGATG</td>
<td>98°C /10s</td>
<td>20</td>
<td>98°C /1s</td>
<td>56°C/5s</td>
<td>72°C/22s</td>
<td>72°C/1min</td>
</tr>
<tr>
<td></td>
<td>AgigasFADS_ORF_R</td>
<td>GGGCCTCATCACATTCAATAAA</td>
<td></td>
<td></td>
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<td></td>
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<td></td>
</tr>
<tr>
<td><strong>Restriction site primers for cloning</strong></td>
<td>AgigasFADS_pYES_KpnI_F</td>
<td>CCCGGTACCAAGATGGGCGGCCGGGGGCA</td>
<td>98°C /10s</td>
<td>35</td>
<td>98°C /1s</td>
<td>67°C/5s</td>
<td>72°C/20s</td>
<td>72°C/1min</td>
</tr>
<tr>
<td></td>
<td>AgigasFADS_pYES_Xbal_R</td>
<td>CCCCTTAGGGGTACTTGTGGAGATACGCATC</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Table 3: Functional characterization of the *Arapaima gigas* Fads2 in yeast. Conversions were calculated according to the formula (product area / (product area + substrate area)) × 100.

<table>
<thead>
<tr>
<th>FA substrate</th>
<th>FA product</th>
<th>% conversion</th>
</tr>
</thead>
<tbody>
<tr>
<td>18:3n-3</td>
<td>18:4n-3</td>
<td>25.8</td>
</tr>
<tr>
<td>18:2n-6</td>
<td>18:3n-6</td>
<td>16.1</td>
</tr>
<tr>
<td>20:3n-3</td>
<td>20:4n-3</td>
<td>5.8</td>
</tr>
<tr>
<td>20:2n-6</td>
<td>20:3n-6</td>
<td>3.8</td>
</tr>
<tr>
<td>20:4n-3</td>
<td>20:5n-3</td>
<td>nd</td>
</tr>
<tr>
<td>20:3n-6</td>
<td>20:4n-6</td>
<td>nd</td>
</tr>
<tr>
<td>22:5n-3</td>
<td>22:6n-3</td>
<td>nd</td>
</tr>
<tr>
<td>22:4n-6</td>
<td>22:5n-6</td>
<td>nd</td>
</tr>
</tbody>
</table>

nd, not detected
Figure 1. Sequence analysis of *Arapaima gigas* Fads2. A, FADS sequence alignment, white: Cytochrome b5-like domain, green: heme binding motif, orange: conserved histidine boxes, and yellow reported regioselectivity residues. B, Predicted 2D topology of *Arapaima gigas* Fads color code is maintained.
**Figure 2: Molecular phylogenetic analysis.** A - Maximum likelihood phylogenetic analysis, node values indicate bootstrap replicates; B - Bayesian phylogenetic analysis node values indicate posterior probabilities. HSA - Homo sapiens, MDO - Monodelphis domestica, GGA - Gallus gallus, ASI - Alligator sinensis, LCH - Latimeria chalumnae, DRE - Danio rerio; AGI - Arapaima gigas; ONI - Oreochromis niloticus; SSA - Salmo salar; GMO - Gadus morhua; TMA - Thunnus maccoyii; CMI - Callorhinus milii; SCA - Scyliorhinus canicula. BFL - B. floridæ.
Figure 3. Functional characterization of *Arapaima gigas* Fads2 in yeast (*Saccharomyces cerevisiae*). Fatty acid (FA) profiles were determined after the yeast were grown in the presence of exogenously added substrates indicates in each case by (*). Peaks 1-4 in all panels correspond to yeast endogenous FA, namely 1 - (16:0), 2 - (16:1n-7), 3 – (18:0) and 4 – (18:1n-9). FA derived from the exogenously added substrates or elongation products are indicated accordingly in each panel above the corresponding product.

**Supplementary material 1**

2D topology prediction results
**Method**

**TM-helix position starting from 1**

*TOPCONS*:
- TM1: 124-145
- TM2: 151-172
- TM3: 258-279
- TM4: 300-321

*OCTOPUS*:
- TM1: 124-145
- TM2: 146-167
- TM3: 257-278
- TM4: 290-321

*Philius*:
- TM1: 126-147
- TM2: 152-173
- TM3: 259-281
- TM4: 300-324

*PolyPhobius*:
- TM1: 125-148
- TM2: 152-172
- TM3: 259-282
- TM4: 296-321

*SCAMPI*:
- TM1: 123-144
- TM2: 152-173
- TM3: 258-279
- TM4: 300-321

*SPOCTOPUS*:
- TM1: 124-145
- TM2: 146-167
- TM3: 257-278
- TM4: 290-321