Transcriptional activation of zebrafish fads2 promoter and its transient transgene expression in yolk syncytial layer of zebrafish embryos

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The front-end desaturases (Fads) are rate-limiting enzymes responsible for production of long-chain polyunsaturated fatty acids (LC-PUFA). The full spectrum of the transcriptional regulation of fads is still incomplete, as cloning of fads promoter is limited to a few species. Here, we described the cloning and characterisation of the zebrafish fads2 promoter. Using 5′-deletion and mutation analysis on this promoter, we identified a specific region containing the sterol regulatory element (SRE) which is responsible for the activation of the fads2 promoter. In tandem, two conserved CCAAT boxes were also present adjacent to the SRE and mutation of either of these binding sites attenuates the transcriptional activation of the fads2 promoter. An in vivo analysis employing GFP reporter gene in transiently transfected zebrafish embryos showed that this 1754 bp upstream region of the fads2 gene specifically directs GFP expression in the yolk syncytial layer (YSL) region. This indicates a role for LC-PUFA in the transport of yolk lipids through this tissue layer. In conclusion, besides identifying novel core elements for transcriptional activation in zebrafish fads2 promoter, we also reveal a potential role for fads2 or LC-PUFA in YSL during development.

In eukaryotic cells, the front-end desaturases (Fads) catalyse the introduction of a double bond at fixed number of carbons from the carboxyl group. A typical Fads possesses characteristic features such as three histidine-rich boxes, transmembrane regions, and an N-terminal cytochrome b₅ domain containing the heme-binding motif HPGG. Long-chain polyunsaturated fatty acids (LC-PUFA) such as arachidonic acid (ARA, 20:4n-6), eicosapentaenoic acid (EPA, 20:5n-3) and docosahexaenoic acid (DHA, 22:6n-3) are generated from shorter polyunsaturated fatty acids (PUFA) through the actions of Fads and elongases (Elovl). LC-PUFA are crucial for maintenance of cellular membrane integrity, as precursors for eicosanoids, regulation of gene expression and signal transduction pathways. In aquaculture, the interest to decipher the activities of the LC-PUFA biosynthesis enzymes and transcription factors involved in LC-PUFA biosynthesis is driven by the pressing need to improve performance of vegetable oils in aquafeeds¹. These oils are lacking in LC-PUFA but rich in C₁₈ PUFA. Understandably, a greater understanding of distinctive bioconversion capacities of C₁₈ PUFA to LC-PUFA in different farmed species could theoretically improve the strategy of employing vegetable oil as feeds².

Among the vertebrates, numerous studies have reported the molecular cloning and characterisation of Fads from different fish species³. Fads from mammals are principally mono-functional, with Δ5 and Δ6 desaturations being carried out by two separate genes, Fads1 and Fads2, respectively⁴. In fish, fads1 have so far been isolated from only one species, a basal gnathostome, leading to the hypothesis that a complete loss of fads1 in teleosts have occurred following gnathostome radiation⁵. Isolation and characterisation of teleost fads2 to date have reported a broad range of substrate specificities, with Δ4, Δ5, Δ6 and Δ8 desaturation capacities reported. In addition, both mono-functional and bi-functional fads2 have been isolated from a myriad of teleost species⁶⁻⁷. As compared to freshwater species, marine teleosts seemed to display a lesser capacity for bioconversion of C₁₈ PUFA to LC-PUFA,
due to the loss of crucial fads from its genome or low activities of fads2 or elovl. This was postulated to be a result of richer LC-PUFA content in the marine environment, causing diminished capacity for LC-PUFA biosynthesis. Subsequent works however advocated the influence of feeding niche on LC-PUFA biosynthesis, when a catalogue of fads2 and elovl were isolated from marine fish species which possess a diet with limited DHA intake.

Despite the multiple interests in molecular characterisation of fads in teleost, there is a paucity of understanding on its regulation at the transcriptional level. It is conceivable that differences in regulatory activities by dissimilar transcription factors responding to different cues including nutritional status is responsible for the different LC-PUFA biosynthesis capacity in various fish species. These transcription factors themselves are subjected to regulation by the nutritional status of the animal. Therefore, efficient use of vegetable oils in aquafeeds will benefit from the ability to decipher the link between the overall lipid metabolic pathways modification caused by the diet and the subsequent influence on lipid homeostasis genes at the transcriptional level. To date, promoter sequences from teleost fads2 have been isolated from Atlantic salmon, Atlantic cod, rainbow trout, European sea bass, Japanese seabass, large yellow croaker and rabbit fish. Firstly, these works collectively showed the presence of binding sites for transcription factors known to play a role in mammalian cholesterol and lipogenic synthesis pathways such as sterol regulatory element-binding protein (Srebp), nuclear factor Y (NF-Y) and ubiquitous transcription factor Sp1 in the fads2 promoter. Secondly, these studies suggested an interplay between different transcription factors to mediate the regulation of fads2 transcription.

The zebrafish has gained reputation as a useful model in understanding the role of lipids and fatty acids during development. Largely driven by the existence of multiple homologous genes involved in conserved lipid and lipoprotein metabolism network, zebrafish have been used to gain insight on lipid adsorption, atherosclerosis, fatty liver disease and obesity. Zebrafish tissues are reported to contain significantly higher concentrations of PUFA than equivalent mammalian tissues. Molecular characterisation and expression profile of fads2 and several elovl genes during embryogenesis have been reported in zebrafish. However, our knowledge on the function of fads2/elovl and LC-PUFA during development is still fragmentary. Given how important transcriptional regulation in influencing the activities of fads2 may be and how little is known about the regulators and cellular conditions for expression, it is opportune to use zebrafish as a model to fill these gaps.

In this present study, we described the cloning and characterisation of the zebrafish fads2 promoter. This includes identification and functional characterisation of pivotal regulatory elements responsible for driving transcriptional activities using luciferase reporter gene in a zebrafish cell line. In addition, we also carried out an in vivo characterisation of the zebrafish fads2 promoter using a transient approach to track its expression in developing zebrafish embryos.

Results
Cloning and characterisation of the zebrafish fads2 promoter. We isolated a putative promoter region of 1754 bp, comprising 19 bp of the 5′-UTR of zebrafish ΔS/ΔΔfads2 and 1735 bp of the upstream region. For positioning purpose, +1 was given to the putative transcription start site (TSS) derived from comparison of the 5′-end of full-length cDNA (GenBank accession number: NM_131645.2) and DNA sequences of the fads2 gene. Within this fads2 promoter region, a classic transcription initiation element, the TATA box was identified at the location of −26 bp (Supplementary data 1).

Zebrafish Srebp upregulates fads2 promoter activity. Co-transfection of ZFL cells with the fads2-1735 promoter-luciferase reporter plasmid and expression plasmid containing either the nuclear form of zebrafish Srebpl (nSrebpl) or Srebp2 (nSrebp2) proteins, respectively, resulted in significantly higher reporter activities than cells transfected with empty pcDNA3.1 expression plasmid (Fig. 1). Among the two isoforms, the zebrafish nSrebp2 protein resulted in higher luciferase reading. This signified the importance of Srebp proteins in driving zebrafish fads2, and subsequent luciferase reporter transactivation assays were performed with the overexpression of zebrafish nSrebp2 protein.

Identification of cis-regulatory elements within the −214/−67 bp region of zebrafish fads2 promoter. To identify the presence of any potential cis-regulatory element, the nucleotide sequence within the −214/−67 bp region of zebrafish fads2 promoter was aligned and compared to corresponding sequences from rainbow trout (GenBank accession number: KT781408.1), Atlantic salmon (GenBank accession number: AK736067.2), European seabass (GenBank accession number: FP671139.1), large yellow croaker (GenBank accession number: KT781409.1), Japanese seabass (GenBank accession number: KT781410.1), Atlantic cod (GenBank accession number: FJ859898.1), human (GenBank accession number: AP002380.3) and mouse (GenBank accession number: AC135670.3). The alignment disclosed the presence of several conserved putative transcription factor binding sites, namely two inversed CCAAT boxes, an inverted ubiquitous transcription factor Sp1 site and a sterol regulatory element (SRE) within the cloned promoter region (Fig. 2). The result also showed that the CCAAT boxes were highly conserved across the mammalian and fish species while the SRE (TCGAACTGATC) found in the zebrafish fads2 promoter was identical to those in all the teleost species but differed from the mammalian SRE (CTCTGCTGATC) by three consecutive nucleotides. The Sp1 binding site was less conserved amongst the mammalian and fish species, while Sp1 binding site in the mammalian Fads2 promoters, while teleost fads2 promoters typically contain either a single Sp1 site or none.

−161/−67 bp region is critical for zebrafish fads2 promoter activation. We sought to determine a minimal fads2 promoter fragment capable of activating transcription by fusing a series of 5′-deleted promoter fragments to the luciferase reporter gene. Upon normalisation to values obtained with a promoter-less pGL3.1 plasmid, the intact fads2 promoter (−1735/+19 bp) drove a luciferase activity of approximately 200-fold (Fig. 3). Deletion of promoter up to 161 bp upstream of the TSS did not greatly diminish transcription activities, notwithstanding some reduction in...
luciferase readings as compared to the longer fragments. However, further deletion of the sequence within −161/−67 bp almost completely abolished the promoter activity, which indicated a mandatory site within the region of −161 to −67 bp for activation of the fads2 activities. This region contains both CCAAT boxes and the SRE (Supplementary data 2). This also means the removal of the SP1 was not detrimental to the activation of zebrafish fads2 promoter. CCAAT boxes and SRE mediates fads2 promoter activity. We introduced single or multiple site mutations for the SRE, proximal CCAAT box (CATp) and distal CCAAT box (CATd) to the fads2 promoter-luciferase reporter plasmid, followed by the dual luciferase reporter assay in ZFL (Fig. 4). Firstly, results showed that both CATp and CATd could drive expression of promoter, as seen in the fragment with mutated SRE, although the expression was lower than the fragment with all three binding sites intact. In comparison, disrupting both proximal and distal CCAAT bindings sites resulted in total abolishment of luciferase expression, an observation which was recapitulated when all three binding sites were mutated. This implies an obligation for both CCAAT and SRE to optimally drive zebrafish fads2, although the former was still able to activate a reduced measure of transcriptional activities. When SRE was present, CATp seemed to play a more prominent role as compared to CATd in driving promoter activation. A similar pattern was also observed when SRE was disrupted.

In vitro binding of ZFL nuclear proteins at the SRE and CCAAT regions in the zebrafish fads2 promoter. Based on the observations from the mutagenesis experiments, we ran EMSA to assess the binding capacities of transcription factors in ZFL nuclear extract to the fads2 promoter cis-regulatory elements. Results showed the formation of DNA-protein complex when ZFL cells nuclear extract was incubated with wild type oligonucleotide probe containing CATp binding site. There was no complex formed when the probe was incubated without the nuclear extract or when a probe with mutated CATp sequence was used with the nuclear extract. Similarly, the use of non-labelled wild type oligonucleotides in excess amount completely outnumbered the formation of labelled DNA-protein complex. A comparable pattern was shown with CATd sequence (Fig. 5). Unlike the CATp and CATd probes, DNA-protein complex was not observed when nuclear proteins were added into the sample containing oligonucleotides with only SRE site (result not shown). Taking this and the results from mutagenesis experiment into consideration, we used a longer oligonucleotide containing both SRE and the adjacent CATp. Results showed that binding of transcription factors and DNA occurred with the simultaneous presence of both the SRE and CATp binding sites (Fig. 6). Mutation on either site prevented the formation of any DNA-protein complex. When the wild type labelled probe was incubated with excessive non-labelled probe containing mutated SRE and intact CATp, a decrease in DNA-complex staining was observed due to competition for binding. This result encored the observation in cell-based luciferase assay which showed the requirement of CATp for activation of reporter gene expression. In contrast, when wild type labelled probe was incubated with excessive non-labelled probe with intact SRE and mutated CATp binding site, the formation of DNA-complex occurred. Therefore, in the absence of any conjugated protein binding to CATp, the binding of Srebp to the SRE was perturbed. Taken together, the results from mutagenesis and EMSA suggested a functional role for CCAAT box in the Srebp dependent transcriptional activation of zebrafish fads2 promoter.

Transient expression of fads2 promoter (−1735/+19 bp) in developing zebrafish embryos. We observed GFP signal in the yolk syncytial layer (YSL) of embryos beginning at 26 hpf for both the pZs−244 bp and pZs−1735 bp constructs, respectively. As embryos developed, higher percentage of embryos with YSL-GFP
expression were observed, along with increase in GFP signal intensity (Fig. 7). At 120 hpf, more than 95% of injected embryos showed GFP expression at the YSL (Table 1). Therefore, a fads2 promoter fragment containing response elements such as SRE, CATp and CATd proven to be essential for activation of expression is now shown to drive a spatio-temporal expression of fads2 in the zebrafish YSL.

Figure 2. Conserved cis-regulatory elements within the core promoter regions of the fads2 genes from zebrafish and other vertebrate counterparts. Sequence alignment was performed with web-based program, MAFFT (version 7). Numbers indicating the sequence positions are relative to TSS (+1), based on the cDNA sequence information deposited in the GenBank (Danio rerio: CU694371.13; Oncorhynchus mykiss: KT781408.1; Salmo salar:AY736067.2; Dicentrarchus labrax: FP671139.1; Larimichthys crocea: KT781410.1; Gadus morhua: FJ859898.1; Mus musculus: AC135670.3 and Homo sapiens: AP002380.3). Conserved regions for the SP1 binding site, CCAAT boxes and SRE consensus sequence are highlighted in grey, while identical nucleotides are indicated with asterisks.

Figure 3. 5′-deletion analysis of the zebrafish fads2 promoter. ZFL cells were transiently co-transfected with the fads2 5′-deletion promoter-luciferase reporter plasmids, nSrebp2 expression plasmid together with Renilla luciferase reference plasmid pRL-SV40. 5′-deletion plasmids were named according to respective position in relation to TSS (+1) and are represented by horizontal line on the left. Non-coding exon is indicated with open boxes. Luciferase activity of fads2 promoter in ZFL cells is expressed as normalised luciferase activity (to Renilla luciferase activity) relative to empty pGL3-Basic plasmid. Values are means ± S.D. (n = 3) (right panel). Groups indicated with different letters are significantly different (Tukey’s test; P < 0.05).
Discussion

The zebrafish *fads2* was first cloned and shown to have the capacity for *in vitro* Δ5/Δ6 desaturation. The transcripts were reported to express in several tissues known for biosynthesis of LC-PUFA during embryonic development. The mRNA transcript of zebrafish *fads2* is also seemingly regulated by dietary PUFA intake. Despite these studies, investigation on the regulation of *fads2* expression at transcription level is lacking. Here, we cloned...
a 1754 bp promoter region of the zebrafish *fads2* gene to determine response elements critical for control of *fads2* expression. We showed here that like all known teleost *fads2* promoters, zebrafish *fads2* promoter harbours SRE and CCAAT binding elements 10–13. Zheng et al. 10 also showed a high degree of sequence conservation between mammalian, amphibian and teleost species, which indicates conserved mechanism of *fads2* role. As with salmon 10 and trout 13, putative SP1 binding site was also identified in this region of zebrafish *fads2* promoter. In contrast, all sequenced promoters of *fads2* from marine teleost species so far do not contain SP1 element, which was postulated as a reason for lower *fads2* expression in marine fish species 11–13.

Results from the promoter deletion experiment demarcated regions responsible for transcriptional activation of the zebrafish *fads2*. One such region is the cis-element SRE, the binding site for Srebp transcription factors. Using mutagenesis approach, we further corroborate the obligation of the SRE region in activating transcription of zebrafish *fads2*. We also showed that this sequence forms a DNA-protein complex with nuclear extract of ZFL cells. In human, an E-box like SRE region, as opposed to the classical SRE region, was crucial for both activation and suppression of the FADS2 gene 24. Srebps are prominent regulators of lipid, fatty acid and cholesterol biosynthesis, and were first identified as nuclear proteins possessing binding abilities to SRE of low density lipoprotein receptor 26. Like mammals, fish have two distinct homologues of mammalian Srebp1 and Srebp2 27. In zebrafish, activation of hepatic Srebp induce the expression of both lipid and cholesterol biosynthesis genes 28. Our results also showed that in ZFL cells, Srebp2 activated a higher level of *fads2* transcription as compared to Srebp1. This is reminiscent of the findings in Atlantic salmon, where Srebp2 promoted higher *fads2* activities as compare to Srebp1 19. The importance of Srebp in mediating teleost *fads2* expression has been shown through the use of cell-based luciferase assays involving Srebp expression plasmids or mutation of the promoter’s SRE response element 13,14,29. In terms of tissue localisation, there was a parallel pattern between the distribution of *srebp* and *fads2* mRNA expression 13,29. The role of Srebp as a sensory mediator between dietary LC-PUFA intake and LC-PUFA biosynthesis was also observed in several fish species wherein limited dietary LC-PUFA intake led to increased Srebp level, subsequently resulting in higher *fads2* mRNA expression 13,29–32. In rat and human, dietary PUFA inhibits the rate of *Fads2* transcription through Srebp, probably by lessening the production of mature Srebp or accelerating the decay of *srebp* mRNA 24,33. Besides FADS2, Srebp have also been shown to regulate ELOVL 24,34.

**Figure 6.** Binding of ZFL nuclear proteins to the SRE on the zebrafish *fads2* promoter. Biotin-labelled probes containing SRE and proximal CCAAT box (CATp) were incubated with the nuclear extract of ZFL cells. Specific binding of DNA-protein complex was validated with site-mutated oligonucleotides (Lanes 3 and 4) and 100-fold molar excess of non-labelled mutated competition oligonucleotides (Lan 5 and 6).
Through a combination of deletion and mutagenesis, we demonstrate the importance of two CCAAT motifs in the core promoter region of zebrafish fads2 for transcriptional activation. The CCAAT box, named after the five nucleotides customarily found in the binding sites, is one of the most ubiquitous binding elements in eukaryotic promoters, with several corresponding binding proteins already isolated and characterised. The NF-Y is the major protein recognizing the CCAAT box, and is obligatory for the transcription of a considerable number of genes with ubiquitous or tissues-specific expression patterns. Functional CCAAT motifs recognised by NF-Y are among the most frequent boxes found in promoters. Mutation of putative NF-Y binding sites in fads2 promoter of two marine teleost species also resulted in weaker transcriptional activities.

Our present finding suggests that the putative NF-Y binding sites are required for formation of complex between Srebp and SRE, which then activates the transcription of reporter gene. This result resonated with earlier findings wherein Srebp was incapable of driving transcription of target genes if adjacent NF-Y binding sites were mutated. By themselves, Srebp proteins are inherently weak transcriptional activators, which underlines the need for Srebp proteins to interact with additional transcriptional factors to regulate the expression of lipogenic genes. Studies on promoters of cholesterol and fatty acid metabolism genes consistently highlight the role of NF-Y as a pivotal partner in Srebp-mediated regulation of transcriptional activities. Elsewhere, studies on murine stearoyl-CoA desaturase (Scd) and human FADS2 promoters both showed a requirement for putative NF-Y binding sites located adjacent to SRE for transcriptional activation. Promoters of teleost and amphibian fads genes also contain a conserved region bearing NF-Y and Srebp binding sites at similar positions relative to the transcriptional start site (TSS). It is speculated that direct interaction of Srebp and NF-Y, together with the

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**Figure 7.** Transient GFP expression of linearised fads2 promoter (−1735/+19 bp) localised in yolk syncytial later (YSL) of zebrafish embryos. Lateral view of 26, 48, 72 and 96 hpf of transient transgene embryo injected with linearised fads2 promoter (−1735/+19 bp) showing increased expression in YSL (A,C,E,G) and their corresponding control (B,D,F,H). Cross-sectional view of a 120hpf control embryo (K–N) and a 120hpf transient transgenic embryo (I,J) injected with linearised fads2 promoter (−1735/+19 bp) showing expression at YSL.

| Description          | Embryo number (%) |
|----------------------|-------------------|
| Injected             | 72                |
| Survival             | 63 (87.5%)        |
| Dead/Abnormal        | 9 (12.5%)         |
| GFP expressing embryos | 58 (92.1%)     |
| Localisation of GFP  |                   |
| YSL                  | 57 (98.3%)        |
| Non-specific (notochord) | 1 (1.7%)  |

**Table 1.** Transient expression of the linearised fads2 promoter (−1735/+19 bp) in zebrafish embryos. Figures are cumulative embryos scored from three independent microinjections at 120 hpf.
binding of both these transcription factors to neighbouring sites on the DNA lend stability to the DNA-protein complex\(^\text{25}\). In addition, our results showed that removal of the putative SP1 binding region did not supress the luciferase expression. The interaction between Srebp and SP1 in the regulation of \textit{Fads} have not been demonstrated this far\(^\text{13,33}\), although a synergistic type of activation between these two transcriptional factors have been elucidated in other lipid metabolism genes\(^\text{34}\). A role for SP1 in Srebp-mediated activation of \textit{Fads2} however cannot be ruled out as different isoforms of Srebp or nutritional conditions may utilise different co-regulatory factors to activate expression of \textit{Fads}, as shown in promoters of fatty acid synthase\(^\text{45}\).

Fatty acids from yolk storage are critical for energy and structural components for both fish embryo and larval stages\(^\text{46}\). Together with the head region, yolk sac has been reported to contain majority of the lipids during zebrafish embryonic stages\(^\text{18}\). While saturated and monounsaturated fatty acids (SFA, MUFA) are consumed for energy, LC-PUFA are essential for cellular membrane integrity, regulation of gene transcription, regulation of signalling pathways and production of eicosanoids. Previously, the yolk was regarded as a non-metabolic active site for nutrient reserve, where molecules such as lipids are stored within yolk cells and transported when needed\(^\text{46,47}\). However, emerging findings employing zebrafish as the model organism, are recognising the yolk as site for active lipid metabolism preceding transportation\(^\text{18}\). Using a transient transgenic approach in zebrafish embryos, we showed that the \(-1735/+19\) bp region of the \textit{fads2} promoter directed a specific expression in the YSL beginning at 26 hpf embryos. In support of this, \textit{fads2} and \textit{elovl} transcripts were reported to be expressed in YSL at the 24 hpf stage, alongside brain tissues\(^\text{18,21}\). During early embryonic development, some of the blastomeres situated at surface of the yolk collapse and fuse with the yolk cells, leading to the formation of YSL\(^\text{46,47}\). It then undergoes a series of changes in terms of thickness and nuclei distribution, before going into the process of degradation upon the exhaustion of yolk supplies\(^\text{46}\). YSL governs major roles during development, including yolk metabolism, epithelial specification of germ layers, organogenesis of yolk and innate immunity\(^\text{46,47}\). A major role of YSL in teleost development is on the transport of nutrients from yolk to embryonic cells and tissues. Transcripts of genes related to lipid metabolism and transportation\(^\text{33–38}\) are expressed in YSL during early development to facilitate the hydrolysis and transport of yolk lipids. When these genes are disrupted, embryos show reduced yolk consumption, delayed embryonic growth and failure in lipid absorption\(^\text{46,49}\).

The presence of GFP signal in YSL also suggested a role for LC-PUFA, the product of \textit{fads2} promoter activities, in transportation of yolk lipids. Detailed analyses of lipid classes in zebrafish yolk imply active \textit{de novo} synthesis of lipids, in which fatty acids are resynthesised into major lipid classes between 0–48 hpf stage, followed by packaging into very low-density lipoproteins for supply to whole embryo beyond 48hpf\(^\text{15,17}\). In \textit{Caenorhabditis elegans}, transportation of yolk lipoprotein to oocytes was disrupted in mutants with PUFA deficiency, a malaise restored by feeding worms with n-6 PUFA\(^\text{40}\). These findings, viewed alongside our results showing localisation of \textit{fads2} promoter activity at the YSL, provides the collective interpretation that LC-PUFA is required for transportation of yolk lipids during development.

In adult teleost, the liver is a major site for LC-PUFA biosynthesis, with both \textit{fads} and \textit{elovl} transcripts being reported in adult zebrafish liver and early larval stages\(^\text{18,21}\). Since the injected embryos only showed expression in the YSL, the \textit{fads2} promoter fragment lacks the regulatory region responsible for directing expression of \textit{fads2} in other organs such as liver, intestine or brain. It is plausible that the regulatory elements for the expression of these tissues are located in more distal regions or/and introns\(^\text{39,40}\). Multiple studies have shown that recapitulation of endogenous expression by a promoter in zebrafish embryos depends on the length of the promoter fragment\(^\text{40–44}\). In conclusion, we successfully isolated a 1754 bp upstream region of the zebrafish \textit{fads2} promoter and identified several response elements critical to drive the activation of transcription. We also showed the capacity of this fragment to direct GFP expression at the YSL region of 26 hpf and older stages embryos. Since zebrafish relies on yolk as sole lipid source prior to onset of exogenous feeding capacity, it potentially provides a platform to further elucidate the role of \textit{fads2} or LC-PUFA in transportation of yolk lipids across the YSL.

Materials and Methods

Cell line maintenance and subculture. The zebrafish liver cell line, ZFL (ATCC® CRL-2643™) was purchased from the American Type Culture Collection (ATCC, USA) and maintained in the complete growth medium at 28 °C following the protocol of ATCC. Routine subculture was performed once the ZFL cells have achieved 80 to 90% confluence in a 25 cm² flask.

Zebrafish maintenance and embryo collection. Wild type zebrafish (AB line) were purchased from the Institute of Molecular and Cell Biology (IMCB, Singapore), and maintained in the ZebTec Stand Alone System (Tecniplast, USA). The fish were kept on a 13:11 h light/dark cycle at 28.5 °C, and fed till visual satiation twice daily with a combination of commercial micro pellet (Aquadene, Malaysia), frozen bloodworms and \textit{Artemia} nauplii. Breeding and embryo collection were carried out according to Westerfield (2000) with adaptation\(^\text{65}\).

Animal ethics approval. Husbandry, handling and use of animals in this manuscript comply with the guidelines and requirements of the Animal Ethics Committee, Universiti Sains Malaysia and were approved by the same committee (PA/ASC/002/2011).

Zebrafish \textit{fads2} promoter cloning and promoter-luciferase reporter plasmid construction. A promoter region of 1754 bp, corresponding to \(-1735/+19\) bp relative to the transcription start site of the zebrafish \textit{fads2} gene was amplified with PCR utilising the \textit{i-Taq}\(^\text{TM}\) Plus DNA Polymerase (iNtRON, Korea). DNA extracted from the ZFL cells was used as the template for the amplification, together with the forward primer F1735 containing the restriction site for \textit{KpnI}, and a reverse primer R19 containing the restriction site for \textit{XhoI} (Supplementary data 3). The reaction mixtures were subjected to a thermal cycling programme consisted of an
Lipofectamine™ 2000 (Life Technologies, Germany) at a volume of 0.15 μl and incubated overnight for cell attachment. Transient transfection was carried out using transfection reagent was used and each binding reaction containing 1X binding buffer, 1 (v/v) NP-40, 50 mM KCl, 5 mM MgCl₂, 20 mM EDTA, 5 nM nucleotides in a total volume of 20 μl binding reaction. The DNA-protein binding complexes were then incubated at room temperature for 30 min before size-fractioning on a 5% (v/v) non-denaturing polyacrylamide gel at 4 °C in the Mini-PROTEAN II Slab Electrophoresis Cell (Bio-Rad, USA). Finally, biotin-labelled DNA was detected with the aid of Chemiluminescent Detection System (Promega, USA). Firefly luminescence reading for each sample was first normalised to Renilla luminescence measurements were performed in the GloMax® Multi + Detection System (Promega, USA). Prior to electrophoretic mobility shift assay (EMSA), 3′-end of each complementary strand of the oligonucleotides (Supplementary data 6) were labelled with biotin using the Biotin 3′ End DNA Labelling Kit (Thermo Scientific, USA) following descriptions of the manufacturer. Subsequently, biotin-labelled double-stranded oligonucleotides (10 μM) were used and each binding reaction containing 1X binding buffer, 1 μg Poly (dI·dC), 5% (v/v) glycerol, 0.05% (v/v) NP-40, 50 mM KCl, 5 mM MgCl₂, 20 mM EDTA, 5 μl nuclear extract and 100 fmol biotin-labelled oligonucleotides in a total volume of 20 μl ddH₂O was prepared. For the negative control binding reaction, nuclear extract was added into 1X annealing buffer containing 10 mM Tris (pH 7.5), 1 mM EDTA and 50 mM NaCl, and the mixture was heated in the Thermomix (Eppendorf, Germany) for 5 min at 95 °C before gradually cooling down to room temperature over a period of approximately 1 h. Nuclear extract was prepared from the transfected ZFL cells using the NE-PER® Nuclear and Cytoplasmic Extraction Reagents (Thermo Scientific, USA) according to the manufacturer's instructions. For EMSA, the LightShift® Chemiluminescent EMSA Kit (Thermo Scientific, USA) was used and each binding reaction containing 1X binding buffer, 1 μg Poly (dI·dC), 5% (v/v) glycerol, 0.05% (v/v) NP-40, 50 mM KCl, 5 mM MgCl₂, 20 mM EDTA, 5 μl nuclear extract and 100 fmol biotin-labelled oligonucleotides in a total volume of 20 μl ddH₂O was prepared. For the negative control binding reaction, nuclear extract was excluded, and 100-fold molar excess of non-labelled oligonucleotides was included to the competition binding reaction. The DNA-protein binding complexes were then incubated at room temperature for 30 min before size-fractioning on a 5% (v/v) non-denaturing polyacrylamide gel at 4 °C in the Mini-PROTEIN II Slab Electrophoresis Cell (Bio-Rad, USA). Finally, biotin-labelled DNA was detected with the aid of Chemiluminescent Nucleic Acid Detection Module (Thermo Scientific, USA) and luminescent signal was captured using the CCD camera (Bio-Rad, USA).
GFP promoter-reporter plasmid construction. DNA fragments from the fads2−244 and −1735 promoter-luciferase reporter plasmids were PCR amplified using both forward and reverse primers containing the restriction site for XhoI and BamHI (Supplementary data 3), respectively. The resulted PCR products were gel-purified and cloned into the pZsGreen1-1 GFP promoter-reporter plasmid.

Transient expression of GFP promoter-reporter plasmid in zebrafish embryo. In vivo promoter activity of both promoter fragments was determined by microinjection of GFP promoter-reporter plasmids into zebrafish embryos, respectively. Prior to microinjection, GFP promoter-reporter plasmids were linearized with the restriction enzymes XhoI and NotI. Each pZsGreen1-1 GFP promoter-reporter construct was adjusted to 10 ng/μl with dH2O containing 1× Danieau’s buffer and 0.2% phenol red. Approximately, 4.6 nl of the DNA solution was delivered into embryos at early one-cell stage by using the Nanoliter 2000 Microinjector (World Precision Instrument, USA).

The injected embryos were incubated in E3 medium at 28.5 °C. The embryos were periodically monitored under the MVX10 Fluorescence Macro Zoom Microscope equipped with ColorView III Soft Imaging System (Olympus, Japan) until 120 hpf. For cryosectioning, zebrafish young larvae were fixed with 4% paraformaldehyde, treated and sectioned at a thickness of 20 μm using the Leica CM1850 UV Cryostat (Leica Biosystems, Germany).

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Author Contributions
Shu-Chien A.C., M.K. Kuah and S.S. Tay designed the studies. S.S. Tay and M.K. Kuah conducted the research. All authors analysed the data, wrote, read and approved the final manuscript.

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