Acyl-CoA Binding Protein (ACBP) Encoding Gene the Relationship Between Expression Analysis and Different Conditions in *Onychostoma Macrolepis*

Xiaofei Yang, Shaogang Xu, Wentong Li, and Guiqiang Yang

Abstract—One Acyl-CoA binding protein (ACBP) encoding gene was isolated from testis of *Onychostoma macrolepis* using homologous cloning and the RACE method. The full *Oma-ACBP* cDNA (GenBank accession no.: JN254628) was 503 bp long and comprised 37 bp of the 5'-Untranslated Regions (UTR), 267 bp of the coding sequence (CDS) encoding a 88 amino acid proteins, and 199 bp of the 3' - Untranslated Regions (UTR) with the polyA tail. A condensed phylogenetic tree show that *Oma-ACBP* had similarities with vasa of fish species and even mammal and amphibian species, The *Oma-ACBP* shared 92% sequenced identity with *Cyprinus carpio* and 91% with *Carassius auratus*, including a conserved FABP domain, the result enriches our understanding in the study of sequence classification in *Onychostoma macrolepis*. The Quantitative real-time PCR analysis demonstrated that *Oma-ACBP* was highly expressed in intestine, spleen and liver, but weakly in testis, ovary, heart, brain, cheek, muscle and eye. It was highly expressed in liver and spleen (42% of dietary protein), weakly expressed in intestine (52% of dietary protein). In starvation challenge, the expression was weakly expressed from 5 day to 8 day, and highly expressed from 3 day to 5 day, and the expression kept stabilizing until the end of the experiment.

Index Terms—ACBP, onychostoma macrolepis, cloning, qRT-PCR, expression.

I. INTRODUCTION

Acyl-CoA binding protein (ACBP) was known as diazepam binding inhibitor (DBI)/endozepine (EP), is a highly conserved, approximately 10 kDa cytosolic protein [1]. ACBP binds saturated and unsaturated C14-C22 acyl-CoA esters in a one-to-one binding mode with high specificity and affinity (Kd=2×10⁻⁶ nm) [2]. Acyl-CoA-binding protein (ACBP) is involved in lipid metabolism, ACBP plays a key role in multiple cellular tasks including modulation of fatty acid biosynthesis, enzyme regulation, and gene regulation [3]-[5]. ACBP is also apparently involved in fatty acid elongation, sphingolipid synthesis, protein sorting and vesicle traffic [6]-[8]. In mammals, ACBP was first characterized in liver (L-ACBP), homologous distinct ACBP genes have been identified in testes (T-ACBP) [9] and brain (B-ACBP) [10].

Now, the ACBP has been identified in many fishes, such as *emiatostella vectensis* [11], *Danio rerio* [12], *Ictalurus punctatus* [13], *Oncorhynchus mykiss*, *Salmo salar* and *Cyprinus carpio* were submitted to Genbank in the aquatic animals (http://www.ncbi.nlm.nih.gov). Sequence analysis comparisons suggest that ACBP is highly conserved across fish species in protein sequence. The largescale shoveljaw fish (*Onychostoma macrolepis*) is an omnivorous species belonging to the family *Cyprinidae*, it is a valuable wild species [14]. However, no data is available on the ACBP and its expression of *Onychostoma macrolepis*. In the present study, in order to clarify the role of the ACBP in energy metabolism, we isolated the cDNA and characterized the expression of ACBP in the tissues, starvation and dietary challenge.

II. MATERIALS AND METHODS

A. Materials Collection

Largescale shoveljaw fish (*Onychostoma macrolepis*) 21± 4 g in body weight, 12.3± 3.2 cm in body lengh, were obtained from National Engineering Research Center for Freshwater Fisheries in Fangshan District (Beijing, China), reared in 0.2 m³ class tanks and temperature (19–22°C) were maintained at the same levels. Fishes were held for a week to adjust to the temperature and environment conditions, and were fed four times daily with common fish feed.

B. Dietary and Starvation Challenge Experiment

For the study of ACBP in tissues, the testis, ovary, liver, heart, brain, cheek, intestine, muscle, eye and spleen were collected from the mature fish. In dietary challenge experiment, two levels of dietary protein (42% and 52%) were set, the fish was put in twelve sets of four class tanks that were controlled at the same temperature and environment conditions, 20 fish are in each tanks. Following the 12-week feeding, the liver, intestine and spleen were maintained in a concrete condition until 5-hour post-feed, the diet of Tubifex (*Limnodrilus hoffmeisteri*) as the control group. In the starvation challenge experiments, three fishes from each group, and the liver were collected at 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13 and 14 day post-starvation from the fishes. All of samples were snap frozen in liquid nitrogen, and stored at −80°C until use.

C. RNA Extraction

Tissue species were homogenated by IKA-T10...
Homogenizer (IKA, Staufen, Germany). Total RNAs from different tissues were extracted by the RNeasy Prep Kit (Tiangen, Beijing, China) technique following the manufacturer's instructions and preserved for quantitative real-time PCR. Purity and integrity were assessed by the A260/280 nm ratio using a spectrophotometer (BioRad, Hercules, USA) and integrity on a 1.2% agarose gel.

D. Isolation of a cDNA Encoding Oma-ACBP by PCR

First-strand cDNA synthesis was performed by Reverse Transcriptase kit (TaKaRa, Dalian, China). First strand cDNA was generated in a 25 μl reaction volume containing, 500 ng total RNA, 5×PrimeScript Buffer, 50 μM oligoT Primer, 100 μM Random 6 mers, PrimeScript RT Enzyme Mix I, the reaction was conducted at 37°C for 15 min and 85°C for 5sec. Specific primer pairs were designed based on the sequence available in NCBI to amplify a cDNA fragment of approximately 155 bp from testis (Fig. 1 and Fig. 2). PCR was performed under the following conditions: denaturation at 94°C for 3min, followed by 40 cycles of 30sec at 94°C, 30sec at 55°C, 2min at 72°C, final extension of 7min at 72°C, using a Eppendorf PCR cycler (Eppendorf, Germany). The RACE method was used to obtain the full length Oma-ACBP sequence by using BD SMART RACE Kit (Clontech, Toronto, Canada). Dissociation melting curves analysis of amplification products were performed at the end reaction to confirm that primer-dimers and nonspecific amplification were not detected. The relative quantification between ACBP and β-actin were calculated using the comparative CT method.

Primers Sequence(5′→3′) Position Product(bp)
For the cloning of conserved sequence
PF1 AAGACGAGCAGAGGAGGAGGAGG 59-155 155
PR1 CTAGTCCCCCATTGGG 55-203 148
For the cloning of 3′end RACE and 5′end RACE
P5RACE- GCCGGGAGTGTGAGA 168-50 336
1 TTTCAC
P3RACE- TCAGAAAGCAGCAGAGGAAGGTCGCTGCTTGG 55-203 149
2 GAGCTC
PSRACE- GCCGGGAGTGTGAGAATACTCC 1-178 198
1 TG
2 TAAC
Primers used for quantitative real-time PCR
P-QF TGAGGACGAGCAGGAGG 55-198 144
P-PQR GCCGTCCAGTGAATC 121
actinF TCCCTCCAATGCCAAG 121
actinR GGAGTCCATCACGATA 121

Fig. 1. Oma-ACBP and β-actin genes, Corresponding PCR primers used for gene fragment cloning and quantitative real time PCR.

E. Phylogenetic Analysis of Oma-ACBP

The PCR, RACE and quantitative real-time PCR primers were designed using Primer Premier 5(PREMIER Biosoft, Palo Alto, USA). Similarity analysis was performed with Blastn(http://www.ncbi.nlm.nih.gov/). Gene translation and prediction of the deduced protein were performed with Open Reading Frame Finder (ORF Finder, http://www.ncbi.nlm.nih.gov/gorf/). Sequence assembly was performed using DNAMan software (Lynnon Biosoft). The Molecular Evolution Genetics Analysis (MEGA) version 4.0 and Clustal X were used to construct a phylogenetic tree using the neighbor-joining method. Quantitative data were expressed as means±SD (standard deviation).

F. Quantitative Real-Time PCR Assay

Total RNAs were isolated from the different tissues of fish. First strand cDNA was synthesized and stored at −20°C until used for real-time PCR, using PrimeScript® RT Enzyme (Takara, Dalian, China). The first strand cDNA was synthesized as described above and stored at −20°C until used for real-time-PCR. The mRNA expression of Oma-ACBP in different tissues was measured by real-time PCR, the real-time primers were designed based on the cloned Oma-ACBP (Genbank accession: JN254628).

A pair primers (P-QF and P-QR) were used to amplify a PCR product, The β-actin was used as a reference gene for internal standardization with the primers actinF and actinR (Fig.1). The qRT-PCR was performed in a volume of 20 μl containing 10 μl 2×SYBR Premix Ex Taq, 0.4 μl ROX, 6 μl DNase-RNase free water (Takara Dalian China), 2 μl cDNA, 0.8 μl each of 10 μM forward and reverse primer, Quantitative real-time PCR(qRT-PCR) was performed in a Funglyn FTC-3000 real-time cycler (Funglyn Biotech, Toronto, Canada). Dissociation melting curves analysis of amplification products were performed at the end reaction to confirm that primer-dimers and nonspecific amplification was not detected. The relative quantification between ACBP and β-actin were calculated using the comparative CT method.

cDNA of each sample from the experiment fishes were used for analysis in real-time PCR, the β-actin transcript levels were used to normalize samples. The real-time standard curve of each gene was prepared using the Oma-ACBP cDNA as a template. The relative quantification between ACBP and β-actin transcript in all cDNAs were calculated from comparative CT method using a FTC-3000 real-time cycler System Software (Funglyn Biotech, Toronto, Canada), to maintain repeatability, the baseline was set autoscale by the software.

G. Tissue Expression Analysis

Expressions of Oma-ACBP mRNA in tissues were demonstrated by quantitative real-time PCR. The samples of untreated from the gonad, ovary, liver, heart, brain, cheek,
intestine, muscle, eye and spleen were collected. The samples of dietary challenge from liver, intestine and muscle were collected, starvation challenge from liver, intestine and muscle were collected. Total RNA at 500 ng was reverse transcribed to the first strand cDNA, a pair of primers (P-QF and P-QR) was used to amplify a PCR product, and the Primers (actinF and actinR) were used to amplify the β-actin fragment. The real-time PCR reaction was the same as that described above.

Fig. 3. Analysis of Oma-ACBP expression in different tissues by quantitative real-time RT-PCR. Each bar represents the mean±SD (standard deviation), \( n=3 \).

Fig. 4. Analysis of Oma-ACBP expression in dietary challenge by quantitative real-time RT-PCR at the protein level of 42%, 52%.

Fig. 5. Analysis of Oma-ACBP expression post starvation challenge from 0day to 14day. Each bar represents the mean±SD (standard deviation), \( n=3 \).

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A. Phylogenetic Analysis of Oma-ACBP with Other Species

To examine the relationship of fish sequences to the various members of the ACBP family, a phylogenetic tree was constructed using the neighbor-joining method (NJ) with the amino acid sequences of fish and other known ACBP.

Phylogenetic analysis revealed that the Oma-ACBP formed an exclusive group. One group consisted of fish ACBP (Carassius auratus, Cyprinus carpio, Danio rerio, Ictalurus punctatus, Epinephelus coioides, Oncorhynchus mykiss, Salmo salar), group another consisted of mammal and amphibian ACBP (Homo sapiens, Xenopus laevis; Fig. 9).
in rainbow trout, ACBP could be used as markers to prevent excess muscle fat accumulation and protein content, ACBP expression changed in muscle and liver, the genes of domesticated fish mainly for the regulation of structural genes [19]. In the present study, there were obvious difference of Oma-ACBP transcript post dietary challenge between 42% and control group in liver and spleen, 52% and control group in intestine; There was no significant difference between 52% and control group in liver, 42% and 52% in intestine and spleen. The metabolism was active during the dietary protein at the content of 42% in liver and spleen. This may indicate that through changes of dietary protein content, ACBP expressions of tissue were regulated, thereby changing the metabolic intensity, the level expression of ACBP may serve as an indicator of feed selection, the expression patterns in the Onychostoma macrolepis remain to be further studied. On the other hand, the ACBP over expression affects metabolic responses to diets with distinct difference in their fatty acid chain lengths, the ACBP-induced tissue-specific regulation of expression of PPARs and SREBP [20]. In mice, the ACBP expression level decreased significantly after 24 hours of hunger, and then fedded, the ACBP expression level reach the minimum point after 12 hours, the ACBP expression returned to normal levels after 24 hours [16]. Another study, cute fasting dramatically reduces ACBP/DBI mRNA levels in the hypothalamus and the ependyma bordering the third and lateral ventricles [21]. In Strongylocentrotus internedius, ACBP(HM208169) was weekly expressed in gonads of female sea urchins after 2-day starvation, the expression of ACBP(HM208170) in gonads male urchins was weak after 2-day starvation, turned to high after 4-day starvation, and kept high from 13th to 30th starvation [22]. In the present study, the expression of Oma-ACBP was impacted by the time of hunger extension, this may indicate that fish adopt to the stress response of starvation by altering gene expression.

In summary, this study reports the first cloning the ACBP gene(JN254628), from Onychostoma macrolepis. We obtained 503 bp long sequences (excluding the poly-A tail), 267 bp of the coding sequence(CDS) encoding a 88 amino acid proteins. By quantitative real-time PCR, we showed a significant changes in mRNA expression of two different conditions showing similarity in organs of Onychostoma macrolepis. The expression analysis of Oma-ACBP indicated that it play a key role in metabolism system. So, this study provides another insight on the gene of Onychostoma macrolepis, it has a role in the study of molecular biology and organizations, recently, the Oma-ACBP mRNA is highly expressed in intestine, this may indicate that mRNA expression of ACBP is up regulated to adjust to intensity of fat metabolism in intestine.

In insect, the Rhodnius prolixus ACBP gene expression increased in the first day after blood meal and then decreased to unfed levels in the seventh day after meal, and ACBP gene was expressed in all analyzed tissues and quantitative PCR showed that expression was highest in posterior midgut [13]. In rats fasting for 24 hours significantly decreased tissue levels of ACBP in the liver, whereas feeding of a high-fat diet for 48 hours caused ACBP levels to increase [18]. In rainbow trout, the growth rate of domesticated species were higher than wild species, ACBP expression changed in muscle and liver, the genes of domesticated fish mainly for the regulation and control, and a strong ability to control cell division and regulation of structural genes [19]. In the present study, there were obvious difference of Oma-ACBP transcript post dietary challenge between 42% and control group in liver and spleen, 52% and control group in intestine; There was no significant difference between 52% and control group in liver, 42% and 52% in intestine and spleen. The metabolism was active during the dietary protein at the content of 42% in liver and spleen. This may indicate that through changes of dietary protein content, ACBP expressions of tissue were regulated, thereby changing the metabolic intensity, the level expression of ACBP may serve as an indicator of feed selection, the expression patterns in the Onychostoma macrolepis remain to be further studied. On the other hand, the ACBP over expression affects metabolic responses to diets with distinct difference in their fatty acid chain lengths, the ACBP-induced tissue-specific regulation of expression of PPARs and SREBP [20]. In mice, the ACBP expression level decreased significantly after 24 hours of hunger, and then fedded, the ACBP expression level reach the minimum point after 12 hours, the ACBP expression returned to normal levels after 24 hours [16]. Another study, cute fasting dramatically reduces ACBP/DBI mRNA levels in the hypothalamus and the ependyma bordering the third and lateral ventricles [21]. In Strongylocentrotus internedius, ACBP(HM208169) was weekly expressed in gonads of female sea urchins after 2-day starvation, the expression of ACBP(HM208170) in gonads male urchins was weak after 2-day starvation, turned to high after 4-day starvation, and kept high from 13th to 30th starvation [22]. In the present study, the expression of Oma-ACBP was impacted by the time of hunger extension, this may indicate that fish adopt to the stress response of starvation by altering gene expression.

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metabolism of ACBP. However, further studies are needed to define mechanisms for Lipids metabolism and the relevant impact of FABP transcription conditions in *Onychostoma macrolepis*.

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**REFERENCES**


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