Molecular cloning and characteristics of DnaJa1 and DnaJb1 in Coilia nasus: possible function involved in oogenesis during spawning migration

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Abstract

Background: Coilia nasus oogenesis/spawning migration is a well-defined synchronous arrangement process. DnaJs are indispensable molecular chaperones for oogenesis process. However, how DnaJs involved the anadromous spawning migration mechanism is outstanding and plausible.

Results: In this regard, two DnaJs (Cn-DnaJa1 and Cn-DnaJb1) are cloned from the Coilia nasus’s ovary. Their structure both contains J domain, G/F domain and ZF domain. Their mRNA transcripts were found extensively expressed in all the sampled tissues and significantly highly in gonads, which probably mean that DnaJs involved in C. nasus’s gonad development basal metabolic processes. In the process of spawning migration, Cn-DnaJa1 and Cn-DnaJb1 mRNA transcripts were also expressed with significant differences during oogenesis with highest levels in the development phase, and maintaining high levels during the multiplication, mature and spawning phase. Further study showed that the DnaJa1 and DnaJb1 protein have high distribution in the onset phase and mainly distributed in the oocyte cytoplasm especially during the migration development phase’s.

Conclusions: This experiment study demonstrated that DnaJs participate in reproductive regulation during the spawning migration process in C. nasus and possibly play a vital role in the ovary development process. These findings also provided a base knowledge for further molecular mechanism study of spawning migration.

Keywords: Coilia nasus, DnaJa1 and DnaJb1, Oogenesis, Spawning migration

Background

Coilia nasus is widely distributed in the fresh water rivers such as the Changjiang River and coastal waters, which is a kind of anadromous fishery species [1, 2]. Fish anadromous migration is a process preparation for spawning and it is also a highly complex biological event [3]. It is well known that the C. nasus spawning migration process was season-depended and temporally based reproductive activity [4]. Sex matured fish usually upstream migrate for a long distance for spawning. During the upstream migration, the fish also undergoes the oogenesis process. When oocytes develop to maturation, fish spawn in the Changjiang River spawning ground. After fertilized, the oosperms float down and hatch along the river side [2, 4]. When up streaming, lots of water stress such as water flow and temperature change will induce numerous adaptive genes such as heat shock proteins (HSPs) up or down regulation to meet the migration behavior need [5–7].

It has been proven that fish oogenesis is a well-defined synchronous arrangement biological process, which can also provide a good biologic model for finding out the
regulated mechanism of related genes on oocyte differentiation [8]. During _C. nasus_ spawning migration process, there were distinctive phases can be distinguished [8]. It is well known that oogenesis process is regulated by different kinds of HSPs [9]. Various types of HSPs are functioned in a stage-specific and developmentally regulated manner during oogenesis in mouse, rats and humans [10–12].

HSP40 family is an important molecular chaperone in HSPs super family. HSP40 homolog usually contained a highly conserved J domain, which is also named DnaJs [13]. It has a typical N-terminal consensus sequence that facilitates interactions with HSP70 [14]. Although they are moderately conserved, the DnaJ homolog do vary in structures and can be divided into different subtypes [15]. Different types of DnaJs contain different conserved domains, further study found that DnaJ proteins are cofactors for HSP70 [14, 16]. Through its interaction with HSP70, DnaJs are involved in DNA transcription, cell proliferation, signal transduction and other biological processes [1, 9].

Previous studies on ovary transcriptomic analysis in _C. nasus_ have shown that DnaJs were up-regulated significantly during spawning migration [17]. Based on this bias, the full length cDNA of _Cn-DnaJa1_ and _Cn-DnaJb1_ was cloned, tissues depended and temporal mRNA expression patterns in ovaries during different spawning migration phases were investigated; and finally _Cn-DnaJa1_ and _Cn-DnaJb1_ protein during different development phases were clarified. Combination with data from other literatures on fish anadromous migration biology, our findings can support more evidence for the anadromous spawning migration mechanism.

**Results**

**Characteristics of DnaJa1 and DnaJb1 cDNA**

Using RACE PCR method, the complete _Cn-DnaJa1_ cDNA was 1471 bp in length, which has a 1227 bp open reading frame (ORF), a 79 bp 5′-untranslated region (UTR), and a 165 bp 3′-UTR. The predicted ORF encoded a protein containing 408 amino acids with a calculated molecular weight (Mw) of 45.9 kDa and a theoretical predicted isoelectric point (pl) of 6.68. Conserved domain analysis identified that DnaJa1 have an N-terminal, a J domain (residues 6–65 aa) containing the conserved histidine proline aspartic acid (HPD) motif (residues 34–36 aa) (Fig. 1a), a Glycine-rich region profile (67–101 aa), a central Zinc Finger domain (126–210 aa) formed by four Zinc Finger CR-type profile (repeats of a CXXCGXG motif, ZF domain), and a DnaJ C-terminal region (residues 225–335 aa). The _Cn-DnaJa1_ cDNA and predicted protein sequence had been submitted to GenBank and the accession number is MH748547. The full-length cDNA of _Cn-DnaJb1_ was 1376 bp, which containing a predicted ORF of 1095 bp, beginning with a methionine codon at position 124 and ending with a TGA termination codon at position 1204 (Fig. 1b). Its GenBank accession number is MH748548. The encoded 364 amino acid polypeptide had an Mw of 39.4 and a pl of 8.94. As expected, the predicted protein of _Cn-DnaJa1_ and _Cn-DnaJb1_ had several similar DnaJ family domains, including J domain, HPD domain, G/F domain (glycine- and phenylalanine-rich domain), ZF domain (a cysteine-rich zinc finger domain) and the C domain (a less well-conserved C-terminal) (Fig. 1a & b).

**Homology and phylogenetic analyses**

Phylogenetic trees were constructed by analyzing the amino acid sequences of _C. nasus_ DnaJa1 and DnaJb1 with those from other species, respectively. BLAST analysis suggested that _Cn-DnaJa1_ had high sequence homology similarity with _Clupea harengus_ (90%), _Salvelinus alpinus_ (81%), _Salmo salar_ (81%), _Sinocyclocheilus grahamii_ (80%), _Oncorhynchus mykiss_ (80%), _Oryzias latipes_ (76%), _Pelodiscus sinensis_ (70%), _Otolemur garnettii_ (69%), respectively. _Cn-DnaJb1_ shared high similarity with that from _Clupea harengus_ (81%), _Oncorhynchus mykiss_ (72%), _Danio rerio_ (72%), _Scleropages formosus_ (72%), _Cynoglossus semilaevis_ (71%), _Oreochromis niloticus_ (69%), _Otolemur garnettii_ (69%), _Bos taurus_ (64%), respectively. DnaJa1 and DnaJb1 protein sequences were obtained from NCBI data base. The produced Neighbor-Joining (NJ) phylogenetic tree indicated that the evolution of _DnaJa1_ and _DnaJb1_ was almost in accordance with the evolution of species (Fig. 2a & b).

**DnaJa1 and DnaJb1 mRNA expression patterns**

_DnaJa1_ and _DnaJb1_ mRNA transcripts were extensively expressed and testified. Expression levels were found highest in ovary, and higher level in testis, liver and blood than in brain, stomach and intestine, and the gill has the lowest expression level (Fig. 3a). During the spawning migration cycle, _Cn-DnaJa1_ and _Cn-DnaJb1_ mRNA transcripts expression pattern showed a similar way (Fig. 3b). They were sharply up-regulated to peak level in the development phase, and maintained high levels during the multiplication, mature and spawning phase. Then expression was down-regulated in the resting phase significantly.

**Western blotting (WB) results**

For WB analysis, the crude protein was extracted from _C. nasus_ ovaries in different spawning phase. Anti-DnaJa1, anti-DnaJb1 and anti-serum were also used to recognize the reaction. When the crude protein extracts were transferred to the nitrocellulose membrane and immunoblotted with anti-DnaJa1 and anti-DnaJb1,
Fig. 1 Nucleotide and deduced amino acid sequences of Cn-DnaJa1 and Cn-DnaJb1. The deduced amino acid sequence is shown under the nucleotide sequence. a is for Cn-DnaJa1 and b is for Cn-DnaJb1. The termination codon is marked by an asterisk. J domain is showed underlined, HPD motif is boxed, Glycine-rich region profile is signed on the dotted line, Zinc finger CR-type profile (DnaJ CXXCXGXG central domain, 4 repeats) are in bold.
similar bands were observed on the immunoblot membrane (Fig. 4a). Control serum for the pre-immunized rabbit did not recognize any bands when encountered C. nasus ovaries extracts. DnaJa1 protein immunoblotted bands were considerately heavier than the DnaJb1’s. These two proteins showed different expression patterns when immunoblotting. DnaJa1 protein increased firstly and then declined, while DnaJb1 protein declined consistently when fish spawning. Between different spawning phases, DnaJa1 and DnaJb1 protein levels both reached the peak in the onset phase and presented declining tendency after that (Fig. 4b).

Localization of DnaJa1 and DnaJb1

Immunohistochemistry (IHC) method was used to identify the distribution of the DnaJa1 and DnaJb1 protein in different types of oocyte. Whole ovary sections were shown in Fig. 5, which were stained with hematoxylin-eosin (H&E, Fig. 5, O1) and immunolabeled with anti-DnaJa1 and DnaJb1 (Fig. 5, O3 & O4, counterstained with H&E), respectively. In the normal mature ovary, immunoblotting positive signals were shown in brown both for the DnaJa1 and DnaJb1 protein (Fig. 5, O3). Interestingly, the DnaJa1 protein was more widely distributed than the DnaJb1’s distribution, and they mainly located in the cytoplasm of different oocyte especially obvious in the development phase (Fig. 5, O3 & O4). There are no positive signals in the negative control, which was incubated with pre-immune rabbit serum (Fig. 5, O2).

**Fig. 2** Phylogenetic trees of DnaJa1 and DnaJb1 family members. Phylogenetic tree constructed by the MEGA 4.0 program by the neighbor-joining (NJ) distance method. 2a is for DnaJa1’s and 2b for DnaJb1’s. The statistical robustness of the tree was estimated by bootstrapping with 1000 replicates. Bootstrap values were indicated by genetic distance.
Discussion
In the present study, the complete cDNA sequence and characteristics of DnaJa1 and DnaJb1 gene in C. nasus were reported. Four conserved domains were also proven to be existed in Cn-DnaJs, which are thought to be involved in substrate binding [13, 16]. By stimulating ATPase activity via this J domain, which can change the conformation of HSP70, and then lead to the folding of substrate polypeptides and the binding to HSP70 [14]. Motif studies on Cn-DnaJa1 and DnaJb1 indicated that the highly conserved HPD motif may play a vital role in the activation of HSP70 via the J domain [18]. It is proposed that the HPD domain make special interactions with certain motifs in the ATPase domain of HSP70 through the formation of hydrogen bonds [19]. It was found that
these mentioned domains also functioned in Cn-DnaJa1 and DnaJb1 structures. These results suggest that Cn-DnaJa1 and DnaJb1 probably act as a partner in the HSP70 chaperone complex via a mechanism similar to that observed in E. coli DnaJ, which employs the HPD motif as the contact site for HSP70 [20]. In the silkworm, it was detected that there existed two DnaJ proteins with different posttranslational modifications in particular silkworm tissues [21]. In C. nasus, two kinds of DnaJ shared high similarity with other fish species sequences and have different post translation modification functions. The results suggested that these structure domains may be essential components in physiology and development biological behavior such as fish spawning migration [19, 22].

As tissue dependent expression patterns analysis showed (Fig. 3a), Cn-DnaJa1 and Cn-DnaJb1 mRNA transcripts were higher expressed in gonads than in other examined tissues. These results indicated that Cn-DnaJa1 and Cn-DnaJb1 were potentially synthesized constitutively to support the basic metabolism and development in gonads [12]. Moreover, the expression of Cn-DnaJa1 and Cn-DnaJb1 mRNA transcripts was significantly up-regulated in the onset migration phase (Fig. 3b). In our previous transcriptome analysis, DnaJ homologs were the most abundant chaperone at the mRNA level in the normal ovary [17]. During C. nasus spawning migration process, fish will encounter lots of stress especially such as water temperature change and rising tide. When C. nasus begins migration process, the water temperature is often rising. During fish spawning migration, water temperature is the main inducible factor which induces C. nasus oogenesis and spawning [3]. HSPs almost are thermal inducible and ubiquitously existed [11]. It was seen that not only in the development phase but also till the mature phase, both DnaJs temporal expressions are maintained high levels (Fig. 3b). Cn-DnaJa1 and Cn-DnaJb1 mRNA transcripts were up-regulated sharply in the onset
migration phase and the highest level was observed in the development phase, which is in keeping with the fish oogenesis/migration process. These results may imply that DnaJa1 and DnaJb1 may be functioned and essential in mediating anadromous migration initiate [23]. These findings also truly supported that DnaJa1 and DnaJb1 are both constitutive and inducible in the migration process.

When directly exposed to temperature and oxidative stress in the water environment, it has been proven that greater abundance of the DnaJ/HSP70 protein complex could be related to better protection of oocytes in C. gigas [24]. Furthermore studies in R. decussatus indicated that higher expressions of DnaJ homologs found in good quality oocytes, which might implicate their protective roles in oocytes development [25]. In the present study, WB and IHC results revealed that there were higher DnaJa1and DnaJb1 protein signals in the primary and secondary oocytes (the onset and development phase, Figs. 4 & 5). That is to say, these two proteins are abundant existing in oocyte cells from the beginning of the meiotic stage, and indicated that they

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Fig. 5 Localization of DnaJa1 and DnaJb1 in the mature ovary. Immunohistochemical positive signals of DnaJa1 and DnaJb1 immunolabeling are shown in brown. a: the whole ovary section stained with H&E; b: negative control (NC); c: different part and developmental phase of ovary for IHC with anti-DnaJa1; d: different part and developmental phase of ovary for IHC with anti-DnaJb1. PO: Primary Oocyte, SO: Secondary Oocyte, and MO: Mature Oocyte. Scale bar = 200μm
are essential for oocyte proliferation and differentiation [11, 26, 27]. These findings are also in close agreement with data from mammals and suggested that Cn-DnaJa1 and Cn-DnaJb1 proteins are primarily needed on the initial step of migration and gametogenesis in C. nasus [11, 27]. Moreover, during the oocyte development and proliferative phase, the very active cytoplasmic protein resembling happened, so that the high level of DnaJa1 and DnaJb1 proteins are needed to produce for cell division [28, 29]. Taken together, these findings make biological sense in protein assembling or proceeding, and therefore these proliferating oocytes would require more DnaJa1 / DnaJb1 protein to guarantee for mature and daughter oocytes for C. nasus [8, 11]. Rapid morphological changes are undergoing during oogenesis. Therefore as a result, an increasing high level of DnaJa1 and DnaJb1 protein in the ovary might lead to high reproductive efficiency [8, 12]. These results indicated that DnaJa1 and DnaJb1 are not only very sensitive to the oogenesis onset but also necessary to the ovary maturation [29].

DnaJa1 and DnaJb1 reflect highly synthetic activity and protein transport when cytoplasmic accumulation in the development phase [30]. In Bombyx mori, BmDnaJ1 protein was mainly found in the cytoplasm of blood cells, which suggested that DnaJ1 is a kind of cytoplasm protein [21]. It is likely that rearrangements of DnaJa1 and DnaJb1 in the oocyte may be required for transport of molecular particles between the germ cytoplasm and cell nucleus [8]. DnaJa1 and DnaJb1 are involved in stress-denatured protein folding [16]. When fish up-streaming, they encounter water temperature, tide and water flow stress. To avoid the stress disadvantage, a particular set of HSPs such as DnaJa1 and DnaJb1 will accumulate to protect germ cells against protein denature under these stress during their passage into new development pathways [29, 30].

In conclusion, we characterized DnaJa1 and DnaJb1 molecular structure and their mRNA temporal expression patterns; found that DnaJa1 and DnaJb1 were functional, inducible and essential in the C. nasus ovary development and migration process, suggested their compulsory roles in this process. The findings also indicated that DnaJa1 and DnaJb1 are highly expressed in the maintenance of developing oocytes; which implicated greater abundance of the DnaJs protein could better oocyte quality when fish anadromous migration. Our results expanded and supported earlier reports on DnaJs indispensable function to oocyte development.

**Conclusions**

In the present study, Cn-DnaJa1 and Cn-DnaJb1 complete cDNAs were cloned from the C. nasus ovary. Their typical structures and mRNA transcripts were found extensively expressed with significantly higher in gonads. The results proved that Cn-DnaJs involved in C. nasus’s gonad development basal metabolic processes. Further temporal expression and their protein distribution analysis indicated that DnaJs participate in reproductive regulation during the spawning migration process in C. nasus and possibly play a vital role in the ovary development process. These findings would provide a basic knowledge for further molecular mechanism study of spawning migration.

**Methods**

**Fish sampling and tissue collection**

Using flow drift net, required fishes were sampled by local fishermen. In short, six reach sections in the Yangtze River were selected for sampling fishes during the anadromous migration period (April to July, 2017) [31]. After fish catching, the sampled fish is dead immediately out of water. All fishes were embedded in ice and dissected to classify the reproductive period and then immediately sampled surgically with the needed tissues (including the blood, brain, gill, liver, testis, ovary, intestine and stomach). After washing by fluidic phosphate-buffered saline (PBS), all tissues were immediately throw into the liquid nitrogen and then transferred to the lab in dry ice boxes storing at −80°C for later use. All fish experiment procedures were authorized and approved by the Yangtze River Fish Committee in China.

**Nucleic acid preparation**

RNA Extraction kit (Invitrogen, CA, USA) was used to extract total RNA from different sampled tissues. The acquired RNA quality and concentration were identified by agarose gel electrophoresis and the spectrophotometry, respectively. First-strand cDNA was synthesized with reverse transcriptase (Takara Bio Inc., Shiga, Japan) according to the protocol. Target fragments of Cn-DnaJa1 and Cn-DnaJb1 were acquired from our constructed transcriptome library. After using BLAST program [32], the target and cloning sequence were verified and selected to obtain the full-length cDNA. After verifying the obtained sequence, the Real-time PCR Kit (Takara, Dalian, China) were used for real-time quantitative RT-PCR (qPCR) analysis. All used primers were designed by Primer Premier 5.0 and synthesized by Wuxi Tianlin Biotech Co. Ltd. (Table 1).

**Gene cloning of DnaJa1 and DnaJb1**

Rapid amplification of cDNA ends (RACE) technology was used to acquire the full length sequence of DnaJa1 and DnaJb1 cDNA. Two pairs of gene-specific primers (Gsp) for DnaJa1 and DnaJb1
Table 1 Sequences of primers used in the present study

| Gene-specific Primer pairs for RACE (Gsp) | DNA-Sequence 5’-3’ | Annealing Temperature (°C) | Fragment Size (bp) |
|-----------------------------------------|--------------------|---------------------------|-------------------|
| Gspdnaja1-5’                           | 5’-TTACCGTGCTTGTAGGGAAGTCT-3’ | 61.0                     | -                 |
| Gspdnaj1-3’                            | 5’-GCAAATCCCATAGCATAACCCAG-3’ | 59.2                     | -                 |
| Gspdnajb1-5’                           | 5’-GCCAGCAGTGGCCACAAACA-3’ | 68.6                     | -                 |
| Gspdnajb1-3’                           | 5’-GGGATGTTGGGTGGCCTCGTCT-3’ | 69.4                     | -                 |

**Primers for RT-qPCR**

| Primer Name | DNA-Sequence 5’-3’ | Annealing Temperature (°C) | Fragment Size (bp) |
|-------------|--------------------|---------------------------|-------------------|
| Dnaja1-F    | 5’-AAAAACCATCAGAGAGGACAA-3’ | 58.9                     | 259               |
| Dnaja1-R    | 5’-AGTCTCAACAGACACGGTAAG-3’ | 59.6                     | -                 |
| Dnajb1-F    | 5’-GCCAGCAGCAGCACAACCAACCA-3’ | 68.6                     | 151               |
| Dnajb1-R    | 5’-GGGACGCTCAGTTCAACACCA-3’ | 68.7                     | -                 |

**18sRNA primers**

| Primer Name | DNA-Sequence 5’-3’ | Annealing Temperature (°C) | Fragment Size (bp) |
|-------------|--------------------|---------------------------|-------------------|
| 18sRNA-R    | 5’-TGATTGGGACTGGGGAT-3’ | 59.2                     | 232               |
| 18sRNA-F    | 5’-TACGCGACCCGGGGTGCTG-3’ | 62.4                     | -                 |

(GspDnJa1-5’, GspDnJa1-3’; GspDnJa1-5’ and GspDnJa1-3’; Table 1) were used for their cDNA full-length sequence cloning. Their RACE PCR reactions were performed according to the SMARTer™ RACE cDNA amplification kit protocol (Clontech, Madison, USA). The positive PCR products were sequenced by Wuxi Tianlin Biotech Co. Ltd. and ultimately mixtures were injected into a New Zealand white rabbit at intervals of 2 weeks. After fourth injections, the rabbit was bled, and serum samples were sampled. The antibody titers were verified by enzyme-linked immunosorbent assay (ELISA) method. Under a dissecting microscope, ovaries were dissected from different phases (n = 3) in cold PBS. Ovaries were washed in cold 10 mM Tris–HCl (4°C, pH 7.4) and grinded, the homogenates were dissolved in electrophoresis sample buffer and run in the polyacrylamide gels. Gels were transferred to nitrocellulose membrane for immunoblotting. The primary antibodies diluted at 1:200 for incubation, and then followed by treatment with goat anti-rabbit IgG (1:80) and determine with the DAB method. PBS contained 0.5% Bull serum albumin (BSA) was used for a blocking solution. Embedded membranes were observed by gel imaging and analysis system (Bio-Rad Laboratories-Segrate, Italy) and gray value was counted by the software Image J2x 2.1. The average gray value was used to analysis the protein difference expression.

**Immunohistochemistry (IHC)**

Ovaries were dissected out from fish in the mature stage and fixed in 0.05 M PBS containing 4% paraformaldehyde at 4°C for 24 h (n = 3). Frozen sections were conducted for IHC analysis. Briefly, after washing with fluidic PBS three times, samples were dehydrated in different gradient saccharose-PBS solutions (from 30 to 10%) at room temperature overnight, and then embedded in organ optimal cutting temperature (OCT) compound (Sakure, CA, USA). Standard synthetic C-terminal peptide (DFYQGGRGQGCRQTS for anti-Dnaja1; SARDDTTAQLPAS for anti-Dnajb1) conjugated with keyhole limpet hemocyanin (KLH), and then emulsified with complete (for first immunization) and incomplete (for second to fourth) Freund adjuvant;
frozen sections (8 μm in thickness) were performed using a Leica microtome (Leica CM1900, Germany). IHC procedure steps were clarified briefly as follows. After washing 10 min in 0.02 M PBS for three times, sections were immersed in 0.01 M citric acid buffer (pH 6.0, containing 0.1% Tween 20), and then autoclaved for 8 min. Then sections were incubated with anti-DnajA1 (1:200) and anti-DnajB1 (1:200) at 4°C overnight. After that, sections were rinsed 10 min with 0.02 M fluidic PBS three times for each washes. Subsequently, sections were incubated with horseradish peroxidase conjugated IgG (goat anti-rabbit) for 30 min, ultimately sections were rinsed with 0.02 M fluidic PBS three times for 15 min. Sections were stained with H & E for good visualization. Immunoreactive signals were identified using dianamobenzidine (DAB) as the substrate. For the negative control, sections were incubated with pre-immune rabbit serum instead of the antibody as the above methods.

Statistical analysis
Data is shown as mean ± one standard error (SE). Statistical significance was determined by one-way ANOVA and post-hoc Duncan multiple range tests [34]. Significance was set at P = 0.05.

Abbreviations
DAB: Diaminobenzidine; ELISA: Enzyme-linked immunosorbent assay; GS: Gene-specific Primer; H&E: Hematoxylin-eosin; HSPs: Heat shock proteins; IHC: Immunohistochemistry; KLH: Keyhole limpet hemocyanin; Mw: Molecular weight; OCT: Optimal cutting temperature; ORF: Open reading frame; PBS: Phosphate-buffered saline; PMSF: Phenylmethylsulfonylfluoride; qPCR: Real-time quantitative RT-PCR; RACE: Rapid amplification of cDNA ends; RT-PCR: Reverse transcriptase; UTR: Untranslated region; WB: Western blotting; ZF: domain; Zinc finger domain

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Availability of data and materials
The datasets used and analyzed in this current study are available from the corresponding author on reasonable request.

Authors’ contributions
FDA conceived and designed the whole manuscript, and FXT write the manuscript, YXJ and WQW were responsible for experiments, data scoring and analyses. ZYF, XDP and RJU helped selecting the fish sample, RNA extraction, doing WB and IHC experiments during manuscript preparation. All authors have read and approved the final manuscript.

Ethics approval
All fish experimental procedures were authorized and approved by the Yangtze River Fish Committee in China.

Consent for publication
All authors have read the final manuscript and approved for publication.

Competing interests
The authors declare that they have no competing interests.

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