Data in Brief

Identification of myogenic regulatory genes in the muscle transcriptome of beltfish (Trichiurus lepturus): A major commercial marine fish species with robust swimming ability

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The beltfish (Trichiurus lepturus) is considered as one of the most economically important marine fish in East Asia. It is a top predator with a robust swimming ability that is a good model to study muscle physiology in fish. In the present study, we used Illumina sequencing technology (NextSeq500) to sequence, assemble and annotate the muscle transcriptome of juvenile beltfish. A total of 57,509,280 clean reads (deposited in NCBI SRA database with accession number of SRX1674471) were obtained from RNA sequencing and 26,811 unigenes (with N50 of 1033 bp) were obtained after de novo assembling with Trinity software. BLASTX against NR, GO, KEGG and eggNOG databases show 100%, 49%, 31% and 96% annotation rate, respectively. By mining beltfish muscle transcriptome, several key genes which play essential role on regulating myogenesis, including pax3, pax7, myf5, myoD, mrf4/myf6, myogenin and myostatin were identified with a low expression level. The muscle transcriptome of beltfish can provide some insight into the understanding of genome-wide transcriptome profile of teleost muscle tissue and give useful information to study myogenesis in juvenile/adult fish.

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1. Direct link to deposited data

http://www.ncbi.nlm.nih.gov/sra/SRX1674471 for muscle transcriptome.

2. Introduction

The muscle development process is spatially and temporally orchestrated by multiple myogenic regulatory factors. At cell fate determination stage of embryo, the external signals trigger mesodermal muscle progenitors transforming into myoblast. The interplay of pax3/pax7, myoD, myf5 and mrf4/myf6 promotes myoblast differentiating into multinucleate myotube (by a cell fusion process) which highly expressed myogenin and mrf4/myf6. The myostatin (also known as GDF-8), on the contrary, plays an inhibitory role on muscle differentiation. Later, the multinucleate myotube further assembles into muscle fiber and expresses variety of muscle specific proteins (MSPs) to build up the major architecture of muscle fiber [1,10,12,14,17]. Basically, the muscle development process is highly conserved between fish and high vertebrates. However, unlike their high vertebrate counterparts, fish continue their hypertrophic and hyperplastic muscle growth through adult stage [9,13]. Study on how muscle can have continuous cell growth...
and proliferation in fish can benefit better understanding on boosting the growth/repair of muscle tissue at juvenile/adult stage in high vertebrates.

Beltfish (Trichiurus lepturus) is a member of the cutlassfish family (Trichiuridae) and is a major commercial marine fish species with robust swimming ability. It is a long, slender fish found throughout the tropical and temperate oceans of the world. Beltfish is a major commercial species in Northwestern Pacific, especially in China, South Korea and Japan. Juvenile beltfish participate in the diet vertical migration, rising to feed on krill and small fish during the night and returning to the sea bed in the day. The movement pattern is reversed by large adults, which mainly feed on fish, squid and shrimp. Adults are highly carnivorous and will cannibalize younger individuals regularly [2]. Since Beltfish play as a top predator in the marine ecosystem, they provide a good model to monitor the bioaccumulation of heavy metals (like mercury and selenium) in the muscle tissues for a long time [4,15,16]. In this study, we took Beltfish as a marine fish model to explore muscle gene regulation at a molecular level by RNAseq approach. The establishment of muscle transcriptome provides not only useful information for evaluating the biological impact of heavy metal bioaccumulation on muscle, but also provides fundamental information for myogenic regulatory gene expression at juvenile/adult stages.

3. Experimental design, materials and methods

3.1. RNA extraction

The muscle tissue was dissected from one wild captured juvenile beltfish (body length around 15 cm) and immediately stored in RNAlater (Qiagen, Hilden, Germany) and then stored at –80 °C prior to RNA extraction. Total RNAs were extracted by using the TRIZOL Kit (Invitrogen, Carlsbad, CA, USA) according to the manufacturer’s instructions. Total RNA samples were then digested by DNase I to remove potential genomic DNA contamination. Integrity and size distribution were checked with Bioanalyzer 2100 (Agilent Technologies, Santa Clara, CA, USA).

3.2. RNA isolation, library construction and Illumina sequencing

Initially, about 2.5 μg of starting total RNAs were used to synthesize cDNA libraries by following the standard protocols of the Illumina TruSeq RNA Sample Preparation Kit (Illumina). The final library had an insert size about 200–300 bp. After qPCR quantitation and dilution, the library was sequenced on an Illumina NextSeq500 with 150 bp paired-end reads. A total of 57,851,358 raw paired-end reads were generated. Adapter sequences were trimmed; and reads with low quality were removed by cutadapt software [8]. After the removal of ambiguous nucleotides, duplicates and low-quality sequences (Phred quality scores < 20), a total of 57,509,280 cleaned reads (99.4%) were obtained. The raw transcriptome sequences in the present study were deposited in the NCBI SRA database (SRX1674471).

Fig. 1. Length distribution of the assembled unigene of beltfish muscle transcriptome.

![Fig. 1.](image1.png)

Fig. 2. Comparison of the gene annotation rate of unigene against NR, GO, KEGG, eggNOR and Swissprot databases.

![Fig. 2.](image2.png)
3.3. De novo transcriptome assembly and functional annotation of muscle expressed genes in belfast fish

Cleaned reads were de novo assembled into unigenes by Trinity software [5] with default parameters settings (k mer = 25 bp). The transcriptome was assembled into 26,811 unigenes, ranging from 201 to 11,019 bp in length and that the N50 length was 1033 bp. The unigene length distribution is shown in Fig. 1. The assembled transcriptomic unigenes were subjected to similarity search against non-redundant (NR) protein, Gene ontology (GO), KEGG, eggNOG [11] and Swissprot databases using BLASTX with an e-value cut-off of 1e−5. Gene names and descriptions were assigned to each unigene based on the BLASTX results. Gene ontology (GO) analysis was then conducted on the assembled unigenes by using Blast2GO [3]. KEGG pathways were assigned to assembled unigenes using online KEGG Automatic Annotation Server (KAAS) (http://www.genome.jp/tools/kaas/). The Bi-directional Best Hit (BBH) method was used to obtain KEGG Orthology (KO) assignment. BLASTX against NR, GO, KEGG and eggNOG databases showed 100%, 49%, 31% and 96% annotation rate, respectively. The systematic comparison of gene annotation rates was summarized in Fig. 2.

3.4. Identification of myogenic regulatory genes

The high-quality cleaned reads of RNA-seq library were mapped to the assembled unigenes by Bowtie2 program [6]. The counting of alignments was done using RSEM [7]. Unigenes with RPKM (reads per kilobase of exon per million reads mapped) ≥ 100 were defined as abundant expressed genes. In juvenile muscle transcriptome of belfast fish, we totally identified 15 unigenes with high similarities to either pax3, pax7, myoD, myf5, myogenin, mrf4/mrf6 or myostatin. To validate the identity of those myogenic regulatory gene homologs, we downloaded the complete protein dataset of 27 fish species with complete genome deposited in NCBI ftp sites (ftp://ftp.ncbi.nlm.nih.gov/genomes/), and combined into a single fish protein database to perform in-house BLAST (summarized in Table S1). Later, we constructed gene-specific phylogenetic tree by using Geneious software (http://www.geneious.com/) with 1000 bootstrap Neighbor-Joining calculation (Fig. S1). In total, we identified one pax3 unigene (c20928_g1_i2), five pax7 unigenes (c31666_g1_i1, c8716_g1_i1, c37452_g1_i1, c35122_g1_i1, c45670_g1_i1), two mrf4/myf5 unigenes (c14456_g1_i1 and c18379_g1_i2), two myoD unigenes (c23568_g1_i2 and c4591_g1_i1), two myf5 unigenes (c24505_g1_i1 and c20162_g1_i1), one myogenin unigene (c25933_g1_i3) and two myostatin unigenes (c18442_g1_i1 and c42164_g1_i1) (Fig. 3 and Fig. S2). The relative expression level of those myogenic regulatory genes in the juvenile belfast fish muscle is relatively low (RPKM < 20, summarized in Table S2) when compared to those highly expressed muscle specific proteins like parvalbumin 3 (c47515_g1_i1, RPKM = 65.269) and myosin heavy chain (c28260_g1_i1, RPKM = 57.107) etc. Based on monophyletic tree topology, we provided strong evidences to support the molecular identity of those myogenic regulatory genes identified in belfast muscle transcriptome. We believe this belfast fish muscle transcriptome and those identified myogenic regulatory genes will provide insights into the muscle development in belfast fish.

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.gdata.2016.04.005.

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