Transcriptome profiling of laser-captured germ cells and functional characterization of zbtb40 during MT-induced spermatogenesis in orange-spotted grouper (Epinephelus coioides)

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Abstract

Background: Spermatogenesis is an intricate process regulated by a finely organized network. The orange-spotted grouper (Epinephelus coioides) is a protogynous hermaphroditic fish, but the process of its spermatogenesis is not well-understood. In the present study, transcriptome sequencing of the male germ cells from orange-spotted grouper was performed to explore the molecular mechanisms underlying spermatogenesis.

Results: In this study, the orange-spotted grouper was induced to change sex from female to male by 17alpha-methyltestosterone implantation. During the artificial spermatogenesis, different cell types from cysts containing spermatogonia, spermatocytes, spermatids, and spermatozoa were isolated by laser capture microdissection. Subsequently, transcriptomic analysis for the isolated cells were performed. A series of genes was used to verify and investigate the expression patterns in spermatogenesis. Furthermore, we also analyzed the expression of the same set of genes involved with steroid metabolism and sex throughout spermatogenesis (early-mid, late, and maturing stages) in the orange-spotted grouper. Several generally female-related genes took significantly changes in sex reversal hinted that the female-related genes in previously recognized may also play vital roles in spermatogenesis and sex reversal. In the transcriptomic data, we focused on zbtb family genes, which may be related to the process of spermatogenesis. Their expression patterns and cellular localization were examined, and the location of Ecztb40 in different gonadal stages was investigated. We found that Ecztb40 was expressed throughout spermatogenesis. These preliminary findings suggest that Ecztb40 is highly conserved during vertebrate evolution and plays roles in spermatogenesis. Besides, the expression of Ecztb40 and Eccyp17a1a overlapped in male germ cells, especially spermatogonium and spermatocyte, which suggested that Ecztb40 might interact with Eccyp17a1a participant in spermatogenesis and sex reversal.
Conclusions: The present study first depicted RNA sequencing of the male germ cells from orange-spotted grouper, and identified many important functional genes and pathways involved in spermatogenesis. The Eczbtb40 gene was subjected to molecular characterization and expression pattern analysis. These results will contribute to future studies of the molecular mechanism of spermatogenesis and sex reversal.

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**Conclusions:** The present study first depicted RNA sequencing of the male germ cells from orange-spotted grouper, and identified many important functional genes and pathways involved in spermatogenesis. The *Eczbtb40* gene was subjected to molecular characterization and expression pattern analysis. These results will contribute to future studies of the molecular mechanism of spermatogenesis and sex reversal.

**Keywords:** Spermatogenesis, transcriptome, laser capture microdissection, grouper, *zbtb40*

**Background**

Transcriptome sequencing is a technology to catalogue all species of transcript, determine the transcriptional structure of genes, and quantify the expression of every transcript under different conditions [1]. Comparison of transcriptomes can provide both quantitative and qualitative information on genetic activity [2]. When gross tissue extracts are used as an mRNA source, tissue heterogeneity confounds assigning expressed genes to different cell populations [3]. Although some *in situ* technologies, such as *in situ* hybridization or immunohistochemistry, could confirm the spatial expression of specific gene, when numerous messages need to be examined, these technologies might not always be possible and is laborious and time consuming [4]. Hence, many researchers have tried to develop microdissection protocols to yield mRNA with sufficient quality for the subsequent transcriptomic analysis. Because of the speed, precision, and versatility, laser capture
microdissection (LCM) was applied in omics, including transcriptome [5, 6] and proteome [7].

LCM is a technology first developed in the late 1990s for obtaining pure populations of targeted cells from specific microscopic regions of tissue sections for subsequent analysis. LCM is a microscope-based technology involving sampling, sample embedding, cryosection, visualization of the target cells, adjustment of the cutting parameters, selecting the target areas, and collection of dissected cells into a tube [8]. The technology was first applied to study gene expression in human cancer cells and gene expression during spermatogenesis in rodents [9-11]. Subsequently, many researcher used LCM to acquire different male germ cells during spermatogenesis for detecting the expression of important genes in many other species, including European sea bass [12], zebrafish [13-15], and African catfish [16].

We first used LCM to obtain male germ cells from different developmental stages induced by 17alpha-methyltestosterone (MT) to profile the natural process of spermatogenesis in orange-spotted grouper (*Epinephelus coioides*). Grouper is a protogynous hermaphroditic fish underlying sex change from female to male in the life history [17]. It has been considered as a good model for the study of sex differentiation, ovarian development, and sex reversal. Because of lack of male fish, few studies of spermatogenesis has been conducted out in orange-spotted grouper. To profile the process of spermatogenesis in grouper, the LCM protocol was optimized to obtain the four germ cell types from male grouper induced by MT implantation, including spermatogonia (SG), spermatocytes (SC), spermatids (ST), and spermatozoa (SZ).

Spermatogenesis is a developmental process in which diploid male germ cells transform into haploid functional male gametes in a tight spatial and temporal organization. Spermatogonia develop into primary spermatocytes, which becomes haploid spermatids
through two meiotic divisions without DNA replication, finally transforming into mature spermatozoa. These processes are achieved by a complex network involving many genes, the germ cells themselves, and several somatic cell types (i.e. Sertoli cells and Leydig cells) [18]. The structural and functional aspects of spermatogenesis are highly conserved among the vertebrates [19]. The release of gonadotropic hormones (follicle-stimulating hormone, luteinizing hormone, growth hormone) released from pituitary stimulate the testis to produce the androgens that initiate spermatogenesis [19]. Hormonal control is a complicated network affected by many stage-specific and cell-specific factors [13, 20].

Previously, quantifying specific genes is difficult because the complex testis structure contains various cell types. Some approaches can be used to quantify and localize unique genes, including in situ hybridization [21, 22], serial analysis of gene expression (SAGE) [23, 24], and in vitro culture of the specific cell population [25]. However, these methods exist some problems to overcome, like the time it takes, the difficulty of the approach itself, etc. Subsequently, researchers have focused on LCM and established an optimized method to obtain specific cell types for studying spermatogenesis. Nevertheless, it is still difficult to isolate the somatic cells around the germ cells to mirror the intricate relationship between somatic cells and different male germ cells during spermatogenesis.

So in our study, transcriptome sequencing was conducted combining LCM to obtain specific cell type to explore the spermatogenesis induced by 17alpha-methyltestosterone (MT) in order to reveal the natural spermatogenesis in a new aspect. Transcriptomic analysis was performed with the goal of revealing the differential gene expression and regulatory networks and zbtb genes came into our sight. Zinc finger and BTB (broad complex, Tramtrack, and Bric-à-brac) (ZBTB) proteins are an evolutionarily conserved family of transcription factors.

Approximately 60 ZBTB proteins has been identified involving in diverse functions
including development, differentiation, and oncogenesis [26-28]. In recent years, ZBTB16 was found to play an essential role in spermatogenesis by controlling the self-renew and differentiation of spermatogonium [29-32]. In the transcriptome data, we found that several zbtb family genes was differently expressed. Further, the zbtb family genes were identified and their expression were investigated during the process of sex reversal and spermatogenesis, and the expression pattern and potential function of Eczbtb40 were determined in spermatogenesis.

Results

Developmental stages of gonads during MT-induced sex reversal

As shown in the Fig. 1, fish oocytes in the sham group remained in the primary-growth stage throughout the experimental period. In contrast, the fish in MT-implanted group underwent sex reversion from female to male. In the first week of MT implantation, the gonads were characterized by degeneration of oocytes and simultaneous proliferation of spermatogenic cysts (Week 1; Fig. 1C). At two weeks after MT implantation, the gonads entered into the intermediate transitional stage filled with SG and SC and numerous oocytes (Week 2; Fig. 1E). At three weeks after MT implantation, the gonads transformed into functional testis, comprised mostly of ST, a small number of SZ, and a few oocytes (Week 3; Fig. 1G).

Capture of target cells

The morphology of SG, SC, and ST were characterized by hematoxylin and eosin (H&E) (Fig.2A) and cryosection staining (Fig.2B). SZ was obtained from the functional testis. During spermatogonia differentiation, the density of heterochromatin reached a the maximum in type B late spermatogonia [33]. When spermatid changed into sperm, the DNA was maximally compacted [18]. Thus, the sizes of the four types of male germ cells
decreased (SG > SC > ST > SZ) and staining of the cell nucleus became darker. Four different male germ cells were detected in tissue slices based on these characteristics.

Validation of sample specificity

Six genes were used to examine the purity of the LCM-derived RNA samples (Fig.3A). *ef1a* is a reference gene and commonly used as an internal control for gene expression analysis. Its expression was detected in all four types of male germ cells at similar levels. Vasa is a germ cell marker [34] whose expression was also found in the four cell types. *slbp2* was specifically expressed in the oocyte of orange-spotted grouper [35], while no expression was detected in any male germ cells except for in the positive control (the gonad with mainly primary-growth stage oocytes). *dmrt1* was specifically expressed in the spermatogonial cells of orange-spotted grouper [36]. Here, *dmrt1* was only expressed in SG and SC with no expression in the positive or negative control.

Quality of transcriptomic data

Four cDNA libraries (SG, SC, ST, and SZ) were constructed with the respective RNA extracted from different male germ cell types of orange-spotted grouper by LCM. The cDNA libraries were sequenced on an Illumina Hiseq 2000 platform (San Diego, CA, USA). A total of 25,304,890 raw reads were obtained, and a total of 244,984,338 clean reads were produced after removing low-quality reads and adapter sequences (Table 1). The average Q20 and Q30 values were 95.74% and 90.57% respectively, and the content of GC was 44.08–47.47%, reflecting the accuracy of the transcriptomic data.

Differentially expressed genes (DEGs) among the four cell types (SG, SC, ST, and SZ) Fragments per kilobase million (FPKM$s$) were used to quantify the gene expression levels. The FPKM values of each gene in the four cell types were compared respectively (Additional file 1). There were 16,406 up-regulated genes and 11,054 down-regulated
genes in SG compared to SC. A total of 15,845 up-regulated genes and 6,895 down-regulated genes were identified in SC to ST. The STs had 8,320 up-regulated genes and 15,797 down-regulated genes compared to SZ. And SG had 22,599 up-regulated genes and 7,399 down-regulated genes relative to STs. There were 14,624 up-regulated genes and 12,413 down-regulated genes in SC compared to SZ. Among the four cell types, 4,483 DEGs were detected and analyzed.

GO and KEGG enrichment of DEGs

In the biological process and molecular function categories of the 4483 DEGs, cellular process (GO: 0009987) and binding (GO: 0005488) were the most enriched GO terms (Fig. S1). And the top 20 pathways were listed from KEGG enrichment (Fig. 4A). Among them, differentiation pathways, signal pathways, and metabolism pathways, which may play crucial roles during spermatogenesis, were observed.

Expression of functional genes putatively associated with sex differentiation and steroid metabolism

From the transcriptomic data, the expression of 20 genes putatively related to sex differentiation and steroid metabolism were analyzed (Fig.5 and 6). Eight of the sex differentiation genes (sox11a, nr0b1, era, erb, wnt9, gdf9, and bmp15) and 13 of the short-chain dehydrogenases/reductases (SDR) genes (cyp17a1, 3hd, sdr12, sdr11, p5cdh, cyp3a40, and sdr13) which are involved in steroid metabolism were expressed in different cell types with FPKM values ranging from 2.62 to 604.7. Most of these genes showed peak expression in SG and SZ (Fig. 4 B).

RT-PCR validation of gene expression inferred from transcriptomic data

RT-PCR was conducted to validate the expression patterns of 13 genes predicted to be
involved with sex differentiation and
steroid metabolism (asterisk-marked genes in Fig. 4) and
deduced from the transcriptomic data. As shown in Figs. 5 and 6, all of these
genes were significantly changed in the process of sex reversal, which was mostly
consistent with the transcriptomic data. These results indicate that the expressional
analysis based on RNA-seq data was credible in the present study.

Molecular cloning and sequence analysis of $Ec$zb$t$b40

Among the 4438 DEGs in four cell types, several $zbtb$ genes including $zbtb1$, $zbtb22$,
$zbtb40$, and $zbtb44$ attracted our attention. ZBTB family proteins are
transcription factors that participate in various important functions. It’s reported that $zbtb16$
($p$lz$f$) played an important role in the
self-renewal and differentiation of the undifferentiated spermatogonia [29].

Therefore, whether the $zbtb$ genes regulate

spermatogenesis of orange-spotted grouper was examined. We studied the
expression of the $zbtb$ genes ($zbtb1$, $zbtb22$, $zbtb40$, and $zbtb44$) to determine their
functions in spermatogenesis (Fig. S3). After further verification, $Ec$zb$t$b40 was came into
our sight, and then deeply functions and expression patterns were exploited.

The open reading frame (ORF) of $zbtb40$ was cloned from the testis of orange-spotted
grouper, which were denominated as $Ec$zb$t$b40. As shown in Fig. S2, $Ec$zb$t$b40 consisted of
a 2400-base pair ORF encoding a peptide of 799 amid acids. Amino acid sequence
alignment and comparison analysis indicated that $zbtb40$ contains a conserved domain
(Fig. S4). A phylogenetic tree was constructed based on amino acid sequences of the
known $zbtb40$ genes (Fig. 7A). On the tree, $Epinephelus coioides$ $zbtb40$ was clustered
together with $Larimichys crocea$ $zbtb40$. The $zbtb40$s contains variously conserved sites in
all selected species, among them the top ten motif sites were showed (Fig. 7B). The DNA sequence of each motif site was displayed in Tab. S1.

Tissue distribution of Ecztb40

Expression of Ecztb40 was examined in eight tissues by semi-quantitative PCR. The results revealed very high Ecztb40 expression in the testis, heart, and pituitary and weak expression in whole brain, head kidney, liver, and ovary (Fig. 3B).

Expression profiles of Ecztb40 in gonads during MT-induced sex reversal

The expression pattern of the Ecztb40 during the MT-induced sex reversal process was investigated. Ecztb40 expression showed no significant difference from the control groups (Fig. 6H). The mRNA levels in MT-treated fish were significantly higher than those in the control group in the second weeks.

In situ localization of Ecztb40 in gonads during sex reversal

At the early stage of sex reversal, there were still a large number of primary-growth stage oocytes which wasn’t detected the expression of Ecztb40 mRNA (Fig. 8B). At the middle stage of sex reversion, many spermatogonia and spermatocytes emerged, showing Ecztb40 mRNA expression (Fig. 8C). At the late stage of sex reversal, there were few oocytes in the gonad and Ecztb40 was abundantly expressed in SG, SC, ST, and SC (Fig. 8D). In comparison, no signal was detected in the ovary from the control group and sham group (Fig. 8A).

Co-localization of Ecztb40 and Eccyp17a1 in gonads during MT-induced sex reversal

Using the JASPAR CORE 2018 database of transcription factor DNA-binding sites, a match was found between an Ecztb40 MEME-generated consensus sequence and the Eccyp17a1 consensus DNA binding site (Tab. S1). The subcellular localization and co-localization of Ecztb40 and Eccyp17a1 in gonads cells were examined during MT-induced sex reversal by fluorescence in situ hybridization under a confocal microscope (Fig. 9). In ovary, Ecztb40
showed little signal, while Eccyp17a1 signals were clearly present in the cytoplasm of primary growth stage oocyte (Fig. 9A-D). In testis, the signals of Ecztb40 in the cytoplasm of SG and SC were overlapped with the Eccyp17a1 signals (Fig. 9E-H). Both Ecztb40 and Eccyp17a1 showed weak signals in ST and SZ. The signals were observed nearly exclusively in the cytoplasm but were barely detectable in the nucleus.

Discussion

In present study, transcriptome analysis was used to profile the process of spermatogenesis based on the LCM technology. A large number of differently expressed genes and signal pathways related to spermatogenesis were identified in this study.

Quality and purity of the LCM-derived RNA

The whole process of LCM must produce a sufficient amount of RNA with high quality to ensure the reliability of the transcriptome results. Single-cell LCM requires a long microdissection period and the yield of RNA is limited. In this study, we modified a previous protocol [37] to handle slides and tissues, samples staining and capture the cells for preserving the RNA integrity throughout microdissection.

To verify the purity of the four target cells, several cell markers were detected. The expression of ef1a indicated the homogeneity of concentration in four cell types. Vasa, a germ cell marker [38], is required for proper germ cells development. Its expression levels were decreased gradually with the process spermatogenesis which is consistent with the expression pattern of vasa in the male germ cells of the gibel carp [38] and brown-marbled grouper [34]. As an oocyte-specific marker [35, 39], slbp2 was used to distinguish the female germ cell and male germ cell. The non-expression of slbp2 in four cell types indicated that target cells didn't contaminate by female germ cells. dmrt1 was used as a spermatogenic cell marker [40] and is only expressed in SG and SC but not in ST and SZ in orange-spotted grouper. Based on the expression of these cell markers, the four target
cells were considered to be pure. The quality and purity of the LCM-derived RNA met the criterion for constructing the single cell transcriptome library after a series of detection.

**Transcriptome overview of spermatogenesis in orange-spotted grouper**

In the present study, male germ cells during MT-induced spermatogenesis were used to explore the natural spermatogenesis and profile the expression patterns of genes related to the sex differentiation and steroid metabolism. Omics resources with orange-spotted grouper are extremely limited so far. The transcriptome sequencing of spermatogenesis can generate a wealth of data, which is useful for further understanding the basic biological mechanisms of spermatogenesis in orange-spotted grouper. KEGG pathway analysis revealed that some biological pathways putatively involved in gonadal development were obtained, including osteoclast differentiation, apoptosis, and retinol metabolism, which is similar to the case in sea bass [12] and Senegalese sole [41]. In addition, the present data showed that 35 DEGs were enriched to Wnt signaling pathway, 5 DEGs were enriched to steroid biosynthesis, 25 DEGs were enriched to Estrogen signaling pathway, 25 DEGs were enriched to cell cycle, 24 DEGs were enriched to oocyte meiosis and 7 DEGs were enriched to ovarian steroidogenesis. These pathways should be evaluated to explore their roles in sex reversal and spermatogenesis.

The orphan nuclear receptor 0 B1 (Nr0B1) is related to many important functions including sex determination, embryonic development, neural differentiation and gonadal steroidogenesis in mammals [42, 43]. Nr0B1 homologues have been identified in several types of teleost such as zebrafish [44], Nile tilapia [45], European sea bass [46], rice-field eel [47], and scallop [48]. The expression pattern of *nr0b1* in rice-field eel revealed that it may be involved in the maintenance of testis function [47]. And *nr0b1* was mainly located in spermatogonia and spermatocytes of testis in scallop, indicating a potential role in spermatogenesis [48]. The significant decrease in *nr0b1* at one week after MT-
implantation (Fig. 5A) also verified its potential effect in the spermatogenesis of orange-spotted grouper. Estrogen receptors (ERs) are composed of two related subtypes, ERα and ERβ, which play roles in sex development and reproduction [49-51]. It was reported that the transcriptional activity of liganded ERs is inhibited by Nr0B1 possibly affecting the recruitment of corepressors [52]. ERα and ERβ were significantly increased after MT-implantation, likely because nr0b1 was decreased in orange-spotted grouper (Fig. 5B, C). This suggested that estrogen and ERs are important in both male and female.

Wnt genes encode a large family of secreted factors with diverse roles in governing cell fate, proliferation, migration, polarity, and death [53]. Although few studies have evaluated the role of wnt9 in testis, the significant increase in wnt9 during sex reversal revealed its potential role in spermatogenesis (Fig. 5D). A previous study showed that the oocyte-secreted growth differentiation factor (GDF) 9 and bone morphogenetic protein 15 (BMP15) regulate the growth, differentiation, and function of granulosa and thecal cells during follicular development in oocyte development, ovulation, fertilization, and embryonic competence [54]. An increasing number of studies has focused on the function of GDF9 in the testis. GDF9 mRNA in the testis of mouse, rat, and human is specifically detected in two of the germ cell types: large spermatocytes, and round spermatids [55]. Recombinant GDF9 was reported to disrupt the inter-Sertoli tight junction permeability barrier in vitro [56], suggesting that gdf9 regulates spermatogenesis in vivo. Similarly, there are several reports of the transcript or protein of BMP15 in testis and its role in testicular functions, indicating that bmp15 functions in the testis as in the ovary [57, 58]. In our study, the significant changes in gdf9 and bmp15 adequately indicate their potential roles in the sex reversal or spermatogenesis of orange-spotted grouper. Overall, the significantly changes in several generally female-related genes indicate that female-related genes may also play vital roles in spermatogenesis and sex reversal. These genes
should be further explored in the context of spermatogenesis and sex reversal.

Possible function of Eczbtb40 in spermatogenesis

Few studies have examined the function of ZBTB40. In our data, Eczbtb40 is only expressed in male germ cells and is important in the process of sex reversal. Gender-specific expression of Eczbtb40 indicated that it may participate in the process of spermatogenesis, too. The predicted result by software suggests that transcription factor Eczbtb40 may regulate Eccyp17a1. Hence, fluorescence in situ hybridization for the newly identified Eczbtb40 was co-localized with Eccyp17a1 in SG and SC during spermatogenesis further. cyp17a1 (steroid 17-alpha-hydroxylase/17, 20 lyase) catalyzes both the 17-alpha-hydroxylation and the 17, 20-lyase reaction. In cyp17a1-deficient zebrafish, male-typical mating behaviors and secondary sex characters (SSCs) were affected [59]. This result indicates the role of Eczbtb40 in spermatogenesis on secondary side. However, the specific function of zbtb40 requires further analysis in the whole process of spermatogenesis.

Conclusions

In the present study, four cell types (SG, SC, ST, and SZ) were captured in the process of spermatogenesis in orange-spotted grouper induced by artificial MT implantation. An optimized protocol was developed to separate different male germ cells in orange-spotted grouper by LCM. Through the transcriptome analysis, the expressional patterns of related genes in spermatogenesis were explored and Eczbtb40 was investigated.

Methods

Animals

Orange-spotted groupers were obtained from Guangdong Daya Bay Fishery Development Center (Huizhou 516081, Guangdong, China). The fish were kept in indoor pools under controlled water temperatures of 22.7~27.8 °C. All fish were anesthetized with MS222
until death in 10-20 minutes, then the fish were sacrificed. All animal experiments conducted were in accordance with the guidelines and approval of the respective Animal Research and Ethics Committees of Sun Yat-Sen University.

MT-induced sex reversal

In this study, the sex reversal was artificially inducted by MT (Sigma, USA) treatment. The fabrication of the slow-release strips and steps of MT implantation were referred to our previous paper with minor modification [60]. Fish (body weight, 1.90 ± 0.65 kg; body length, 43.75 ± 9.25 cm) were divided into two groups, sham group (n = 15) and MT implantation group (n = 15). The dosage of MT was 10 mg/kg body weight. Before implantation (Week 0), gonadal tissues of five fish were collected randomly. After MT implantation, five fish were sampled randomly every week respectively from two groups until 3 weeks. For each fish, one piece of gonadal tissue was fixed in Bouin’s solution for histological examination of the development stage of the gonad, another piece of the gonad was immobilized by 4% paraformaldehyde for florescence in situ hybridization (FISH), the other piece of gonadal tissue was embedded with OCT (Sakura, USA) then frozen immediately in liquid nitrogen for LCM. All the other tissues were frozen immediately in liquid nitrogen, and then stored at -80 °C until further use.

Histology analysis

Gonadal tissues were embedded in paraffin after being fixed 24 h in Bouin’s solution. The embedded blocks were sectioned at 5–6 μm and stained with hematoxylin and eosin (H&E). The gonadal sections were classified by light microscopy.

Visualization of the target cells

There are several factors influencing the RNA quality in the process of visualization, including the samplecryostat sections (the quality of fresh and quick-frozen samples usually is more optimal than fixed tissues), the complex histopathology, and issues arising
from the staining, and so on.

Cryostat sections of gonad for LCM

The RNase-free Membrane Slides (MMI, Switzerland) as a slide to mount the cryosections. A series of procedures were produced before sectioning. The slides were incubated in super clean bench under ultraviolet (UV) radiation for 30 min. Then the slides disposed of with 0.1mg/ml polylysine (Sigma, USA) for 5 min, after that rinsed slide by 0.1 % DEPC ((Sigma, USA, Diethyl pyrophosphate). At last, the slides dried and stored at a sealed box for further use [5].

Before sectioning, the microtome (Leica, Germany) was wiped down with RNase inhibitor (Ambion®, USA) to avoid cross-contamination, and a new blade (Lecia, Germany) treated with RNase inhibitor was used to cut each sample. The gonad blocks were put into Leica Microtomes 30 minutes to adjust the sectioning temperature (-20 °C~ -25 °C). The ovary was cryosectioned at 7 μm, and the testis was cryosectioned at 6 μm.

Cryosections Staining

After being desiccated for one minute, the sections stained by H&E Staining Kit Plus (MMI, Switzerland). The procedures were referred to instructions of the Kit and taken some adjustments to make sure the optimization of RNA quality. However, the sections of the two groups were performed different staining protocols. The whole process was controlled in 30 minutes [61].

Laser Capture Microdissection

First of all, all facilities and tools were wiped by RNase inhibitor. General sterile glass slides were put under the Membrane Slides as supported slides. Once the stained slides exposed in air, the surface of the slide emerged massive water droplets which accelerate degradation of RNA. Thus the whole process of microdissection must control in one hour, and dry the sections quickly. Then load the slides and LCM caps (diffuser caps, MMI,
Switzerland) to laser micro-cutting instrument (MMI, CellCut Plus, Switzerland). Find the cells of interest through adjusting microscope, at the same time optimize three important parameters (cell velocity, laser focus, and laser power). After circling the interesting area, the laser starts to capture the cells as many as possible. At last, unload of caps containing the captured tissue, and add 50 μl TPK Lysis Buffer (Micro Elute® RNA Kit, Omega, USA) immediately. Extract RNA instantly or store diffuser caps in –80 °C (<2 days).

RNA extraction

The procedures of RNA extraction were referred to as instructions of Micro Elute® RNA Kit (Omega, USA) with some adjustment.

Library preparation for transcriptome sequencing

A total amount of 1.5 μg RNA per sample was used as input material for the RNA sample preparation. NEBNext® Ultra™ RNA Library Prep Kit were used to generate sequencing libraries for Illumina® (NEB, USA) following the manufacturer's recommendations. And index codes were added to attribute sequences to each sample. Briefly, mRNA was purified from total RNA using poly-T oligo-attached magnetic beads. Fragmentation was carried out by divalent cations under elevated temperature in NEB Next First-Strand Synthesis Reaction Buffer (5X). First strand cDNA was synthesized using random hexamer primer and M-MuLV Reverse Transcriptase (RNase H-). Second strand cDNA synthesis used DNA Polymerase I and RNase H subsequently. Remaining overhangs were converted into blunt ends via exonuclease/polymerase activities. After adenylation of 3’ ends of DNA fragments, NEB Next Adaptor with hairpin loop structure was ligated to prepare for hybridization. In order to select cDNA fragments of preferentially 150~200 bp in length, the library fragments were purified with AMPure XP system (Beckman Coulter, Beverly, USA). Then 3 μl USER Enzyme (NEB, USA) was used with size-selected, adaptor-ligated
cDNA at 37°C for 15 min followed by 5 min at 95 °C before PCR. Then PCR was performed with Phusion High-Fidelity DNA polymerase, Universal PCR primers, and Index (X) Primer. At last, PCR products were purified (AMPure XP system) and library quality was assessed on the Agilent Bioanalyzer 2100 system.

The clustering of the index-coded samples was performed on a cBot Cluster Generation System using TruSeq PE Cluster Kit v3-cBot-HS (Illumina) according to the manufacturer’s instructions. After cluster generation, the library preparations were sequenced on an Illumina Hiseq platform and 150 bp paired-end reads were generated.

Processing of raw reads and quantification of differential gene expression levels

Raw data (raw reads) of the fastq format were first processed through in-house Perl scripts. In this step, clean data (clean reads) were obtained by removing reads containing adapter, reads containing ploy-N and low quality reads from raw data. At the same time, Q20, Q30 and GC content the clean data were calculated. Q20 indicates that every 100 bp of sequencing reads will have an error, and Q30 indicates that every 1000 bp of sequencing reads will have an error. All the downstream analyses were based on clean data with high quality.

The clean reads were mapped to the orange-spotted grouper (E. coioides) genome (Zhang Y. et al., unpublished data; using SOAP aligner 2.0 [62]), which has been de novo assembled. Index of the reference genome was built using Bowtie v2.2.3 and paired-end clean reads were aligned to the reference genome using TopHat v2.0.12. We selected TopHat as the mapping tool for that TopHat can generate a database of splice junctions based on the gene model annotation file and thus a better mapping result than other non-splice mapping tools.

HTSeq v0.6.1 was used to count the reads numbers mapped to each gene. And then FPKM of each gene was calculated based on the length of the gene and reads count mapped to
this gene. FPKM, expected number of Fragments Per Kilobase of transcript sequence per
Millions of base pairs sequenced, considers the effect of sequencing depth and gene
length for the reads count at the same time, and is currently the most commonly used
method for estimating gene expression levels [63].

Differential expression genes analysis

Prior to differential gene expression analysis, for each sequenced library, the read counts
were adjusted by edgeR program package through one scaling normalized factor.

Differential expression analysis of two conditions was performed using the DEGSeq R
package (1.20.0). The P values were adjusted using the Benjamini & Hochberg method.
Corrected P-value of 0.005 and log 2 (Fold change) of 1 were set as the threshold for
significant differential expression.

GO and KEGG enrichment analysis of differentially expressed genes

Gene Ontology (GO) enrichment analysis of differentially expressed genes was
implemented by the GOseq R package, in which gene length bias was corrected. GO terms
with corrected P value less than 0.05 were considered significantly enriched by differential
expressed genes. The identified DEGs were conducted for enrichment analysis
subsequently by GO: Termfinder software using the hypergeometric test [64, 65], and P-
values were corrected using the Bonferroni method [57]. Being selected significantly
enriched GO terms with Q-value < 0.05.

KEGG is a database resource for understanding high-level functions and utilities of the
biological system, such as the cell, the organism and the ecosystem, from molecular-level
information, especially large-scale molecular datasets generated by genome sequencing
and other high-through put experimental technologies (http://www.genome.jp/kegg/). We
used KOBAS software to test the statistical enrichment of differential expression genes in
KEGG pathways.
Different expression genes analyzed by real-time PCR

To validate the accordance with those genes inferred from RNA-seq data, the relative mRNA levels of 13 differentially expressed genes (nr0b1, era, erb, wnt9, gdf9, bmp15, cyp17a1, 3hd, sdr12, sdr11, p5cdh, cyp3a40, and sdr13) were examined by quantitative real-time PCR (RT-PCR) in sex reversal of orange-spotted grouper. Total RNA of gonad was extracted by TRIzol (Invitrogen, USA) and then 1 µg RNA from each sample was reverse transcribed with random primers by using the First Strand cDNA Synthesis Kit (Roche, USA) according to the manufacturer’s instruction. The RT-PCR reaction was performed in a 10 µl reaction volume using the SYBR Green PCR master mix (Roche, USA). The amplification regime was 95 °C for 5 min, followed by 40 cycles of amplification at 95 °C for 10 s, 58 °C for 15 s and 72 °C for 20 s. The specificity of RT-PCR amplification was confirmed by melt-curve analysis, agarose gel electrophoresis, and sequencing of PCR products. All mRNA quantification data were normalized to ef1a and presented as a relative control group. The specific primers used in this study were listed in Table 3.

Open

Reading

Frames (ORF) cloning and sequence analysis of Eczbtb40 cDNAs

Total RNA of the gonad was extracted by TRIzol (Invitrogen, USA). RNA was reversed to cDNAs with First Strand cDNA Synthesis Kit (Roche, USA). The reverse transcription process was as follow 37 °C for 15 min, 98 °C for 5 min, 4 °C for 5 min. The amplification regime was 35 cycles of 94 °C for 20 s, 55 °C for 10 s, and 72 °C for 20 s, followed by further amplification at 72 °C for 5 min. Based on the cDNA fragments in RNA-seq data, specific upstream and downstream primers (Table 2) were designed. The primers were used to amplify the ORFs of Eczbtb40. The PCR amplification procedures were performed.
as follows: denaturation at 94 °C for 2 min, followed by 35 cycles at 94 °C for 30 s, 52~58 °C (depending on the melting temperature of primers) for 30 s, and 68 °C for 1 min. The reactions were completed with a final extension of 10 min at 68 °C. To separate the desired band in the amplification product, 2% agarose gel electrophoresis was used, and the band of the desired size was purified by the E.Z.N.A. Gel Extraction Kit (Omega, USA). The purified product was then subcloned into the pGEM-Easy vector (Fermentas, USA). According to the sequencing result, the ORFs of *Eczbtb40* were obtained.

The putative amino acid sequences were predicted by DNAMAN software and multiple sequence alignments of amino acids were performed in the ClustalX (1.81) software. Meanwhile, protein phylogenetic analysis was conducted with MEGAX using the method of neighbor-joining method and the top ten motif sites were predicted by motif-based sequence analysis tools (MEME).

**Tissue distribution of *Eczbtb40***

To detect the tissue distribution of *Eczbtb40*, eight tissues were dissected, including whole brain, heart, head kidney, liver, kidney, pituitary, ovary, and testis. Total RNA from eight tissues was extracted. RNA was reversed to cDNA with First Strand cDNA Synthesis Kit (Roche, USA). The reverse transcription process was as follow 37 °C for 15 min, 98 °C for 5 min, 4 °C for 5 min. The amplification regime was 35 cycles of 94 °C for 20 s, 55 °C for 10 s, and 72 °C for 20 s, followed by further amplification at 72 °C for 5 min. The specific primers used in this study were listed in Table 2.

**Expression profile of *Eczbtb40* in gonads during MT-induced sex reversal**

The expression profiles of *Eczbtb40* in the gonad were detected by RT-PCR during MT-induced sex reversal.

**In situ localization of *Eczbtb40* in gonads during MT-induced sex reversal**

Fluorescence in situ localization (FISH) referred to previous papers with minor
modifications [66]. The DIG label was tested with an alkaline phosphatase conjugated Flu-anti-DIG antibody (Roche Diagnostics; diluted 1:1000) and colored the signal with Fluorescence Systems (Roche, USA), and sections were counterstained by 4′6-diamidino-2-phenylindole (DAPI) for cell nuclear staining to confirm the number and status of germ cell. At last, sections were mounted with the Gold Anti-fade reagent (Invitrogen, USA) and imaged by laser scanning confocal microscope (Leica TCS-SP5, Germany).

Dual-label in situ hybridization of Ecztb40 and Eccyp17a1 in gonads

The protocol of dual-label in situ hybridization of Ecztb40 and Eccyp17a1 was referred to previous study [67]. Expression of Ecztb40 was performed using digoxigenin (DIG)-labeled mRNA probes in combination with biotin-labeled Eccyp17a1 mRNA probes.

Statistical analysis

All data were expressed as mean values ± SEM. Significant differences were checked by one-way analysis of variance (ANOVA) and student’s t-test was used, and a probability level less than 0.05 (P < 0.05) was used to indicate significance. All data were performed using GraphPad Prism5.0 (GraphPad Software, San Diego, CA) and analyzed by SPSS17.0 (SPSS, Chicago, IL, USA).

Abbreviations

KEGG: Kyoto encyclopedia of genes and genomes database
RNA: Ribonucleic Acid
cDNA: complementary deoxyribonucleic acid
PCR: Polymerase Chain Reaction kg: kilogram cm: centimeter
PBS: phosphate buffered solution DEPC: diethyl pyrocarbonate
μm: micrometer μg: microgram ml: milliliter μl: microliter
min: minute s: second h: hour
Tris-HCl: Tris (hydroxymethyl) aminomethane
NaCl: Sodium chloride  
PH: potential of hydrogen

NBT: Nitro-Blue-Tetrazolium  
BCIP: 5-bromo-4-chloro-3-indolyl-phosphate

Declarations

Ethics approval and consent to participate
All the procedures in this manuscript had been approved by the Committee for Animal Experiments in Sun Yat-Sen University, China. The methods used in this study were carried out in accordance with the Laboratory Animal Management Principles of China.

Consent for publication
Not applicable.

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

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

Authors’ contributions
XW and YY analyzed the sequencing data; XW, CYZ and YG collected and prepared the samples; XW wrote the manuscript; SSL and XCL supervised the study. All authors read
and approved the final manuscript for publication.

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**Figure Legends**

Fig. 1. Histological effects of MT treatment on gonads of the orange-spotted grouper. (A, B, D and F) Histology of gonads in control fish. (C, E and G) Histology of gonads after MT implantation. PO, primary-growth stage oocyte; PVO, the cortical-alveolus stage oocyte; SG, spermatogonia; SC, spermatocyte; ST, spermatid; and SZ, spermatozoa. Scale bars = 50 um.

Fig. 2. Gonadal structure by different staining methods. (A) Histological structure of gonad in MT treatment by H&E staining. (B) Histological structure of fresh gonad in MT treatment
by cry-sectioning. SG, spermatogonia; SC, spermatocyte; ST, spermatid; SZ, spermatozoa; and PO, primary-growth stage oocyte. Scale bars = 20 um.

Fig. 3. Expression of several sex-related genes and tissue distribution of Ecztbb40. (A) Expression of several sex-related genes in four cell types by laser capture. Ef1a was used as the reference gene. M, maker 2000; C+, the cDNA of ovaries (stayed at the stage with the portion of mostly PO) as a positive control; C-, template-free as a negative control; SG, spermatogonia; SC, spermatocyte; ST, spermatid; SZ, spermatozoa. (B) Tissue distribution of Ecztbb40 in orange-spotted grouper. The ovaries of the fish stayed at the stage with the portion of mostly primary-growth stage oocytes (PO) and the testis from the mature male. Ef1a was used as the reference gene. M, maker 2000; 1, the whole brain; 2, heart; 3, head kidney; 4, liver; 5, kidney; 6, pituitary; 7, ovary; 8, testis.

Fig. 4. Analysis data of transcriptome. (A) The top 20 pathways in KEGG analysis with the significant difference. (B) A heat map was constructed based on the selected genes. Red indicated high expression and green low expression.

Fig. 5. Validation of selected genes using RT-PCR. (A) nuclear receptor subfamily 0 group B member 1 (nr0b1); (B) estrogen receptor alpha (era); (C) estrogen receptor beta (erb); (D) protein Wnt-9 (wnt9); (E) growth and differentiation factor 9 (gdf9); (F) Bone morphogenetic protein 15 (bmp15).

Fig. 6. Validation of selected genes using RT-PCR. (A) Steroid 17-alpha-hydroxylase (cyp17a1); (B) 3-hydroxyisobutyrate dehydrogenase (3hd); (C) dehydrogenase/reductase (SDR) family member 12 (sdr12); (D) SDR family member 11 (sdr11); (E) delta-1-pyrroline-5-carboxylate dehydrogenase (p5cdh); (F) cytochrome P450 3A30-like (cyp3a30); (G) SDR family member 13 (sdr13); (H) zinc finger and BTB domain containing 16 (zbtb40).

Fig. 7. Phylogenetic relationships and conserved DNA motifs in zbtb40 genes from 14 species. (A) Phylogenetic tree of zbtb40swere conducted by MEGAX using the neighbor-
joining likelihood method with 500 bootstrap replicates. Numerals at nodes were bootstrap values. The species names are followed by sequence accession numbers. (B) The top ten motif composition of *zbtb40* s. The motifs, numbers 1-10, are displayed in different colored boxes. The sequence information for each motif is provided in Supplementary table 1.

Fig. 8. ISH analysis the *Eczbtb40* mRNA expression in gonads at the sex reversal process by artificial MT implantation. (A) Gonadal stage with the portion of mostly primary-growth stage oocytes, (B) gonadal stage after one weeks of MT implantation, (C) gonadal stage after two weeks of MT implantation, (D) gonadal stage after three weeks of MT implantation. Red signals indicate *Eczbtb40*, blue staining indicates nuclei. PO, primary-growth stage oocyte; ST, spermatids; SZ, spermatozoa. Scale bars = 50 μm.

Fig. 9. Co-localization of *Eczbtb40* and *Eccyp17a1* by FISH in the gonad of ovary (A-D) and testis (E-H). Gonad sections were stained red for *Eczbtb40* mRNA, green for *Eccyp17a1* mRNA and blue for DAPI. Red and green colors generated a yellow color. PO, primary-growth stage oocyte; SG, spermatogonia; SC, spermatocyte; ST, spermatid. Scale bars = 25 μm.

Figures
Histological effects of MT treatment on gonads of the orange-spotted grouper. (A, B, D and F) Histology of gonads in control fish. (C, E and G) Histology of gonads after MT implantation. PO, primary-growth stage oocyte; PVO, the cortical-alveolus stage oocyte; SG, spermatogonia; SC, spermatocyte; ST, spermatid; and SZ, spermatozoa. Scale bars = 50 um.
Gonadal structure by different staining methods. (A) Histological structure of gonad in MT treatment by H&E staining. (B) Histological structure of fresh gonad in MT treatment by cry-sectioning. SG, spermatogonia; SC, spermatocyte; ST, spermatid; SZ, spermatozoa; and PO, primary-growth stage oocyte. Scale bars = 20 um.
Expression of several sex-related genes and tissue distribution of Ecztb40. (A) Expression of several sex-related genes in four cell types by laser capture. Ef1a was used as the reference gene. M, maker 2000; C+, the cDNA of ovaries (stayed at the stage with the portion of mostly PO) as a positive control; C-, template-free as a negative control; SG, spermatogonia; SC, spermatocyte; ST, spermatid; SZ, spermatozoa. (B) Tissue distribution of Ecztb40 in orange-spotted grouper. The ovaries of the fish stayed at the stage with the portion of mostly primary-growth stage oocytes (PO) and the testis from the mature male. Ef1a was used as the reference gene. M, maker 2000; 1, the whole brain; 2, heart; 3, head kidney; 4, liver; 5, kidney; 6, pituitary; 7, ovary; 8, testis.
Analysis data of transcriptome. (A) The top 20 pathways in KEGG analysis with the significant difference. (B) A heat map was constructed based on the selected genes. Red indicated high expression and green low expression.
Validation of selected genes using RT-PCR. (A) nuclear receptor subfamily 0 group B member 1 (nr0b1); (B) estrogen receptor alpha (era); (C) estrogen receptor beta (erb); (D) protein Wnt-9 (wnt9); (E) growth and differentiation factor 9 (gdf9); (F) Bone morphogenetic protein 15 (bmp15).
Validation of selected genes using RT-PCR. (A) Steroid 17-alpha-hydroxylase (cyp17a1); (B) 3-hydroxyisobutyrate dehydrogenase (3hd); (C) dehydrogenase/reductase (SDR) family member 12 (sdr12); (D) SDR family member 11 (sdr11); (E) delta-1-pyrroline-5-carboxylate dehydrogenase (p5cdh); (F) cytochrome P450 3A30-like (cyp3a30); (G) SDR family member 13 (sdr13); (H) zinc finger and BTB domain containing 16 (zbtb40).
Phylogenetic relationships and conserved DNA motifs in zbtb40 genes from 14 species. (A) Phylogenetic tree of zbtb40s were conducted by MEGAX using the neighbor-joining likelihood method with 500 bootstrap replicates. Numerals at nodes were bootstrap values. The species names are followed by sequence accession numbers. (B) The top ten motif composition of zbtb40s. The motifs, numbers 1–10, are displayed in different colored boxes. The sequence information for each motif is provided in Supplementary table 1.
ISH analysis the Eczbtb40 mRNA expression in gonads at the sex reversal process by artificial MT implantation. (A) Gonadal stage with the portion of mostly primary-growth stage oocytes, (B) gonadal stage after one weeks of MT implantation, (C) gonadal stage after two weeks of MT implantation, (D) gonadal stage after three weeks of MT implantation. Red signals indicate Eczbtb40, blue staining indicates nuclei. PO, primary-growth stage oocyte; ST, spermatids; SZ, spermatozoa. Scale bars = 50 μm.
Co-localization of Ecztb40 and Eccyp17a1 by FISH in the gonad of ovary (A-D) and testis (E-H). Gonad sections were stained red for Ecztb40 mRNA, green for Eccyp17a1 mRNA and blue for DAPI. Red and green colors generated a yellow color. PO, primary-growth stage oocyte; SG, spermatogonia; SC, spermatocyte; ST, spermatid. Scale bars = 25 μm.

Supplementary Files

This is a list of supplementary files associated with the primary manuscript. Click to download.

Tables 1 - 3.pdf
Supplementary Table.xlsx
Supplementary Figures.pdf