Identification of candidate genes of male sexual development from androgenic gland in *Macrobrachium nipponense* through performing long-reads and next generation transcriptome sequencing after eyestalk ablation

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**Abstract**

The eyestalk of crustacean species contained many neurosecretory structures, affected the process of reproduction, molting, metabolism of glucose and other function in crustaceans. In this study, we aimed to selected important metabolic pathways and candidate genes involved in the male sexual development through performing the long-reads and next generation transcriptome sequencing of androgenic gland after the ablation of eyestalk from *Macrobrachium nipponense*. qPCR analysis revealed that the mRNA expression of Mn-IAG was significantly increased after ablation both of the single-side (SS) and double-side (DS)
eyestalk, compared with that of control group (CG). The long-reads transcriptome generated 49,480 non-redundant transcripts. A total of 1,319, 2,092 and 4,351 differentially expressed genes (DEGs) were identified between CG vs SS, SS vs DS and CG vs DS, respectively, indicating the ablation of double-side eyestalk has more important regulatory roles than that of single-side ablation on male sexual development, which was consistent with that of qPCR analysis. Cell cycle, Cellular senescence, Oxidative phosphorylation, Glycolysis/Gluconeogenesis and Steroid hormone biosynthesis were the main enriched metabolic pathways in all of these three comparisons, and the important genes from these metabolic pathways were also selected. The qPCR verifications of 10 GEDs were as the same as that of RNA-seq. The qPCR and RNAi analysis of Hydroxysteroid dehydrogenase like 1 (HSDL1) revealed that HSDL1 has the positive regulatory effect on testis development. This study provided valuable evidences on male sexual development in *M. nipponense*, promoting the studies on male sexual development in other crustacean as well.

**Key words:** Macrobrachium nipponense; Long-reads transcriptome; Eyestalk ablation; Androgenic gland; Male sexual development

**Introduction**

The oriental river prawn, *Macrobrachium nipponense* (Crustacea; Decapoda; Palaemonidae), is widely distributed in China and other Asian countries [1-3], which is an important commercial species with the annual aquaculture production reached of 205,010 tons in 2016 [4]. As the same as other *Macrobrachium* species, male prawns grow faster and reach larger size at the harvest time [2]. Thus, male prawns are preferred in the *M. nipponense* aquaculture. In addition, the rapid development of testis in the reproductive season is another main problem, restricted the sustainable development of *M. nipponense*. The previous studies revealed that the testis of a new born *M. nipponense* can reach sexual maturity within 40 days after hatching [5]. Thus, inbreeding will be happened between the new born prawns. Inbreeding will lead to
the decrease of the ability of resistance to adversity in their offspring, the small scale of market
prawn, and the degradation of germplasm resources. Therefore, it is urgently needed to fully
understand the male sexual differentiation and development mechanism, with the aims of
establishment of the technique to produce all male progeny on a commercial scale, and to
regulate the process of testis development in *M. nipponense*.

Androgenic gland is a special tissue in crustacean species. It has been proven to play
essential roles in male sexual differentiation and development in crustacean species. Many
studies reported that the androgenic gland and its secreted hormones promote the driving of
male sexual differentiation, the establishment of male sexual characteristics, and the
development of the testes in crustacean species [6-7]. The ablation of androgenic gland from
male *Macrobrachium rosenbergii* resulted in the sex reversal to “neo-female”. Insulin-like
androgenic gland hormone (IAG) is an important hormone, secreted by androgenic gland. IAG
was proven to promote the male sexual differentiation and development in many crustacean
species [8-10]. The knockdown of IAG expression by RNAi in male *M. rosenbergii* can also
result in the sex reversal [11]. Based on the importance of androgenic gland in male sexual
differentiation and development in crustacean species, the studies on androgenic gland have
been became the hot topic in recent years. A series of transcriptomes of androgenic gland have
been constructed in *M. nipponense* [12-14], and a series of important genes from androgenic
gland have been showed to play essential roles in male sexual development [15-18]. In addition,
the histological observations during different post-larval developmental stages indicated that
the development of androgenic gland has regulatory roles on the development of testis [5].

The eyestalk of crustacean species has many neurosecretory structures. The X-organ–SG
complex (XO–SG) was identified as a principal neuroendocrine gland located in the eyestalk
in crustaceans [19]. It stores and releases the crustacean hyperglycemic hormone (CHH)
superfamily neurohormones, including CHH, iontransport peptides (ITP), gonad-inhibiting
hormone (GIH), molt inhibiting hormone (MIH), and mandibular organ-inhibiting hormone (MOIH), playing essential roles in reproduction [20-22], molting [23-25], metabolism of glucose [26-27] and other function [28-30]. Knockdown the expression of GIH by RNAi promote the ovarian development in *M. nipponense* [31]. Knockdown the expression of MIH by RNAi promote the molting in *M. nipponense* [32]. CHH has been proven to promote testis development in *M. nipponense* [33].

In this study, we aimed to select the vital metabolic pathways and genes involved in the male sexual differentiation and development in *M. nipponense* through performing the long-reads and next generation transcriptome profiling analysis of androgenic gland after the ablation of single-side and double-side eyestalk. The functions of Hydroxysteroid dehydrogenase like 1 (HSDL1) were further analyzed in depth by using qPCR analysis and RNAi. This study provided valuable evidences on the studies of male sexual differentiation and development in *M. nipponense*, as well as other crustacean species.

**Results**

**The expression analysis of Mn-IAG after eyestalk ablation**

The mRNA expression of Mn-IAG were measured in three groups, including CP, SS and DS (Figure 1). The mRNA expression of Mn-IAG were increased with time of eyestalk ablation in SS group and DS group. The mRNA expressions of Mn-IAG were about 5-folder higher at day 4 and day 7 than that of day 1 in both SS and DS group (P < 0.01). However, the Mn-IAG expression was only slightly higher at day 7 than that of day 4 in both SS and DS group, and showed no significant difference (P > 0.05). The Mn-IAG expression in the DS group was almost 2-folder higher than SS group at the same day, and showed significant difference (P < 0.05).

**Long-read transcriptome**
A total of 22.83 GB clean data was generated in the long-reads transcriptome. A total of 160,496 high-quality transcripts were obtained with a mean length of 2,230 bp. Finally, 49,480 non-redundant transcripts were identified in the long-reads transcriptome. All of the non-redundant transcripts were compared with the non-redundant protein database and nucleotide sequences in NCBI, in order to identify their putative functions. A total of 37,355 (74.94%) unigenes were annotated in Nr database. The other unannotated transcripts represent novel genes whose functions need further investigation.

GO and COG analysis aimed to provide a structured vocabulary to describe gene products. A total of 19,673 (39.76%) unigenes were assigned to GO database, comprised of 52 functional groups (Figure 2). The number of unigenes in each functional group ranged from 1 to 10,057. A total of 13,395 (27.07%) unigenes were highly matched with the known proteins in COG database, classified into 25 functional groups (Figure 3). The number of unigenes in each functional group ranged from 1 to 6,793. KEGG analysis aimed to revealed the regulatory relationship between the unigenes in the long-reads transcriptome. A total of 18,618 (36.72%) unigenes were highly matched the known genes in KEGG database, mapped onto 264 metabolic pathways.

**Identification of differentially expressed genes**

The differentially expressed genes were identified, using the criterion of > 2.0 as up-regulatory genes and < 0.5 as down-regulatory genes, and P-value < 0.05. A total of 1,319 differentially expressed genes (DEGs) were identified between CG and SS, including 713 up-regulated genes and 606 down-regulated genes. A total of 2,092 DEGs were identified between SS and DS, including 1,036 up-regulated genes and 1,056 down-regulated genes. A total of 4,351 DEGs were found between CG and DS, including 2163 up-regulatory genes and 2188 down-regulatory genes. KEGG analysis revealed that Cell cycle, Cellular senescence, Oxidative...
phosphorylation, Glycolysis/Gluconeogenesis and Steroid hormone biosynthesis were the main enriched metabolic pathways in all of these three comparisons.

A total of 15 DEGs were selected from these enriched metabolic pathways, which were listed in Table 3. These genes were differentially expressed in at least two of these three comparisons. Cyclin B3, MAD2A, Polo-like kinase 1, Cyclin A, Cdc2 kinase, and Cyclin B were mainly found in the metabolic pathways of Cell cycle and Cellular senescence, which were differentially expressed in all of these three comparisons. SDHB, Cytochrome c oxidase assembly protein COX11 and Cytochrome c oxidase subunit 7A1 were mainly selected from the metabolic pathways of oxidative phosphorylation. Acetyl-coenzyme A synthetase 2-like, Fructose-bisphosphate aldolase, and Alcohol dehydrogenase class-3 were mainly differentially expressed in the metabolic pathways of Glycolysis/Gluconeogenesis. Estrogen sulfotransferase, 3 beta-hydroxysteroid dehydrogenase and HSDL1 were identified from the metabolic pathways of steroid hormone biosynthesis.

**qPCR verification**

qPCR analysis was used to verify the expressions of important DEGs in the androgenic gland from the CG, SS and DS prawns. The qPCR analysis showed the same expression pattern with that of RNA-seq (Figure 4). Six DEGs from the metabolic pathways of Cell cycle and Cellular senescence showed the lowest expressions in the CG prawns, and highest expressions in the DS prawns, including Cyclin B3, MAD2A, Polo-like kinase 1, Cyclin A, Cdc2 kinase, and Cyclin B. The mRNA expressions of these DEGs in the DS prawns showed significant difference with that of CG prawns and SS prawns (p < 0.01). The mRNA expressions of estrogen sulfotransferase, and alcohol dehydrogenase class-3 showed no significant difference between the CG prawns and the SS prawns (p > 0.05), whereas showed significant difference with that of DS prawns (p < 0.05). The mRNA expression of SDHB and HSDL1 between the
prawns of SS and DS showed no significant difference \((p > 0.05)\), whereas showed significant
difference with that of CG prawns \((p < 0.05)\).

**Expression analysis of Mn-HSDL1**

The previous study has been reported that the Mn-HSDL1 mRNA showed the highest
expression level in the hepatopancreas, followed by the testis, which showed significant
difference with other tested tissues \((p < 0.05)\). The expression in the hepatopancreas and testis
was 187 and 90-folder higher than that in the brain [48]. The mRNA expressions of Mn-HSDL1
in different developmental stages were measured by using qPCR (Figure 5). The Mn-HSDL1
expression in the larval developmental stages were generally higher than that of post-larval-
developmental. The highest expression level was observed in larval day 5 (L5), while it showed
no significant difference with other tested stage \((p > 0.05)\). During the post-larval
developmental stages, the lowest expression level was observed in post-larval day 5 (PL5), and
then gradually increased. The highest expression level was observed in PL25♂, which was 3.72
and 1.94-folder higher than that of PL5 and PL25♀, respectively.

**RNAi analysis of Mn-HSDL1**

RNAi was used to analyze the functions of Mn-HSDL1 on male sexual development in *M. nipponense*. qPCR analysis revealed that the expression of Mn-HSDL1 remained stable in the
control group after the injection of Mn-HSDL1 dsRNA \((P > 0.05)\). However, the expression of
Mn-HSDL1 significantly deceased at day 7 and 14 after the injection of Mn-HSDL1 dsRNA.
The decrease reached to 96% and 90% at day 7 and 14, respectively, compared with that in
control group (Figure 6-A).

The expressions of Mn-IAG were also measured in the cDNA template from the same
prawns (Figure 6-B). According to the qPCR analysis, the expression of Mn-IAG at day 1 in
control group was slightly higher than that of day 7 and day 14, while it generally remained
stable. In RNAi group, the expressions of Mn-IAG were significantly decreased at day 7 and
day 14 after the injection of Mn-HSDL1 dsRNA. The expression decreased about 61% and 54% at day 7 and 14, respectively, compared with that in control group.

Histological observations of testis after RNAi

According to the histological observations, sperms were the dominant cell type in the testis from control group, and only a limited number of spermatogonias and spermatocytes were observed. In RNAi group, the number of sperms were gradually deceased with the time of Mn-HSDL1 dsRNA treatment, and sperms were rarely found at day 14 after Mn-HSDL1 dsRNA treatment. However, the number of spermatogonias were increased (Figure 7).

Discussion

The eyestalk of crustaceans secreted many neurosecretory structures, mediated the reproduction, molting and metabolism of glucose in crustaceans [19-30]. The important neurosecretory structures include CHH, ITP, GIH, MIH and MOIH. Previous study reported that RNAi was used to knockdown the expression of GIH in female M. nipponense, promoting the ovarian developmental process [31]. This study aimed to analyze the effects of eyestalk on male sexual development. qPCR analysis revealed that the mRNA expression of Mn-IAG significantly increased at day 4 and day 7 after eyestalk ablation in both single-side and double-side ablation, compared with that of day 1, and the expression in double-side ablation was significantly higher than that of single-side ablation and normal prawn at the same day, which was consistent with the previous studies [34-36]. However, the expression between day 4 and day 7 showed no significant difference in both single-side and double-side ablation. IAG has been reported to promote the male sexual differentiation and development in crustaceans [8-10]. Thus, the increase of Mn-IAG expression after the ablation of eyestalk indicated that eyestalk has negative effects on male sexual differentiation and development in M. nipponense, which has the similar mediated functions on ovarian development in M. nipponense.

To the best of our knowledge, this is the first long-reads transcriptome in M. nipponense.
The combination of long-reads and next generation transcriptome sequencing can obtain transcripts with better integrity and quality for further gene structure and function analysis. The accuracy and length of the transcripts of the long-reads transcriptome are further improved and optimized through correcting by the next generation transcripts. Thus, it is a suitable strategy for the species without reference genome. The genes related to the male sexual development were predicted to be mainly found in the functional groups of Cell, Cell part, Cellular process and Binding in the GO assignment, and in the functional groups of General function prediction only, Signal transduction mechanisms and Posttranslational modification, protein turnover, chaperones in the COG classification, which were consistent with the previous studies [37-38]. The gene sequences from this long-reads transcriptome provide valuable information for the analysis of gene structure and gene function.

The number of DEGs between CG vs DS were 4,351, which were significantly more than the number of DEGs between CG vs SS and SS vs DS, indicating the ablation of double-side eyestalk has more regulatory roles on male sexual development in *M. nipponense*, which was consistent with the qPCR analysis. KEGG analysis revealed that Cell cycle, Cellular senescence, Oxidative phosphorylation, Glycolysis/Gluconeogenesis and Steroid hormone biosynthesis were the main enriched metabolic pathways in all of these three comparisons. Previous studies have been predicted the important roles of Oxidative phosphorylation, Glycolysis/Gluconeogenesis and Steroid hormone biosynthesis in the mechanism of male sexual development in *M. nipponense* [37-38], predicting the DEGs from these metabolic pathways in this study may play essential roles in male sexual development in *M. nipponense*.

The male sexual development will be vigorous after ablation the eyestalks. The transcriptome profiling analysis revealed that cell cycle and cell senescence were the most enriched metabolic pathway in all of these three comparisons. The cell cycle is a ubiquitous and complex process to ensure the correct during cell proliferation, in order to prevent the
copies of DNA damage, genetic derangement and other errors. Cyclins and cyclin-dependent kinase play essential roles in this process [39-40]. Cellular senescence is defined as irreversible cell cycle arrest caused by different forms of stress. These stresses include telomere shortening, other forms of genotoxic stress, or mitogens or inflammatory cytokines, catalyzing the activation of the p53 tumor suppressor and/or the cyclin-dependent kinase inhibitor p16 [41-42]. The dramatic enrichment of DEGs in these two metabolic pathways indicated that cell cycle and cell senescence play essential roles in the proofreading process when cells make copies of themselves. Four DEGs were enriched in both of the cell cycle and cell senescence, including cyclin A, cyclin B, cyclinB3 and cyclin-dependent kinase 2 (cdk2). Cyclin A is a vital component of the cell-cycle machinery, which can activate two different cyclin-dependent kinases (Cdk1 and Cdk2), functioning in both S-phase and mitosis [43-45]. Cdk1/cyclin B, which is also known as maturation promoting factor (MPF), is one of the main protein kinases. It activates and serves as master regulator for the M-phase transition, phosphorylating and activating other downstream protein kinases, and directly phosphorylating several structural proteins involved in cellular reorganization [46-48]. Cdk family includes 8 cdk genes which can combine with different types of cyclins to form complexes, regulating the process of cell transition from G1 phase to S phase or G2 phase to M phase and exit from M phase. Cdk2 is a member of a highly conserved family of protein kinases, regulating the eukaryotic cell cycle [49-51].

ATP is an unstable high-energy compound that is the most direct energy source in organisms. It is widely acknowledged that ATP is essential for the activities in an organism, including male differentiation and development. In the present study, oxidative phosphorylation and glycolysis/gluconeogenesis were the main enriched metabolic pathways in all of the three comparisons. Oxidative phosphorylation occurs in the inner membrane of mitochondria of eukaryotic cells or in the cytoplasm of prokaryotes. The energy released from
the oxidation of substances \textit{in vivo} promote the coupling reaction between ADP and inorganic phosphate to synthesize ATP through the respiratory chain \cite{52}. Glycolysis/gluconeogenesis promote the conversion of glucose (C$_6$H$_{12}$O$_6$) into pyruvate (CH$_3$COCOO$^- +$ H$^+$), releasing free energy to form the high energy molecules ATP and reduced nicotinamide adenine dinucleotide \cite{53}. Three DEGs were respectively selected from both of the metabolic pathways of oxidative phosphorylation and glycolysis/gluconeogenesis. SDHB is a DEG, which was down-regulated between CG vs SS and CG vs DS. SDHB was also predicted to be involved in the mechanism of male sexual development in \textit{M. nipponense}. SDHB is one of four protein subunits that form succinate dehydrogenase, which catalyzes the oxidation of succinate \cite{54-55}. Two subunits of cytochrome c oxidase were also differentially expressed in oxidative phosphorylation. Cytochrome c oxidase is located at the end of cytochrome c system in cell respiration. This enzyme directly transfers the electron of respiratory substrate to molecular oxygen through cytochrome system \cite{56-57}.

It is widely acknowledged that steroid hormones play essential roles in sexual development. It is generally divided into five main classes, including glucocorticoids, mineralocorticoids, androgens, estrogens, and progestogens. Natural steroid hormones, which are lipids, are generally synthesized from cholesterol in the gonads and adrenal glands \cite{58-59}. Hydroxysteroid dehydrogenase like 1 (HSDL1) was differentially expressed between CG vs SS and CG vs DS, indicating the expressions of HSDL1 were significantly regulated by the ablation of both single-side eyestalk and double-side eyestalk. HSDL1 was also reported to be involved in the mechanism of male sexual development in the previous study \cite{38}. The short-chain dehydrogenase/reductases family (SDR) is a very large enzyme family, which can affect the mammalian reproduction, hypertension, neoplasia and digestion \cite{60-61}. Hydroxysteroid dehydrogenase (HSD) is a subfamily of SDR, playing essential roles in sex-determination, the emergency and maintenance of the secondary sexual characters, and the regulation of endocrine
through catalyzing the metabolism of steroid hormone. Hydroxysteroid dehydrogenase like 1 (HSDL1) was an important gene in the metabolic pathway of steroid hormone [62]. qPCR verification revealed that the expression pattern of important DEGs from these metabolic pathways were as the same as that of RNA-seq, indicating the accuracy of the transcriptome profiling analysis.

Both of this study and previous study predicted the potentially vital roles of HSDL1 in the mechanism of male sexual development in *M. nipponense*. Thus, the potential roles of HSDL1 in the male sexual development were also analyzed by using qPCR and RNAi, combined with the histological observations in this study. Previous studies revealed that HSDL1 was proven to be highly expressed in reproductive tissues (i.e., testis and ovary) in human, as revealed by northern blot analysis [62]. *In situ* hybridization indicated that the expression of HSDL1 is higher in the prostate cancer than that in the normal prostate. In addition, this gene is involved in the development of the sheep fetus in the late gestational stage [63]. The qPCR analysis in different mature tissues revealed that the highest expression level of Mn-HSDL1 was observed in hepatopancreas, followed by testis, while the Mn-HSDL1 RNA were rarely measured in other detected tissues [38]. Vitellogenin was reported to play essential roles in embryonic growth and gonadal development, which was only expressed in the female hepatopancreas, hemolymph, and ovary of *M. nipponense* [64]. The similar expression pattern of HSDL1 in the male prawns predicted that HSDL1 may play similar roles in the male sexual development of *M. nipponense* as that of vitellogenin in female sexual development. In the different developmental stages, the expressions in the larval developmental stages were generally higher than that of post-larval developmental stages, indicating HSDL1 was involved in the metamorphosis process of *M. nipponense* [65-66]. The Mn-HSDL1 mRNA expression was gradually increased from PL5 to PL25. The period from PL5 to PL25 was proven to be the sex-differentiation sensitive period [5]. Thus, the increase from PL5 to PL25 indicated that HSDL1
plays essential roles in gonad differentiation and development. In addition, the gender can be
distinguished for the first time at PL25, and the expression in PL25♂ showed 2 times higher
than that of PL25♀, which also indicated that HSDL1 played more essential roles in male
sexual development. The mRNA expressions of Mn-HSDL1 were significantly decreased at
day 7 and day 14 after Mn-HSDL1 dsRNA injection, indicating the RNAi is efficient in this
study. The mRNA expression of Mn-IAG was also measured in androgenic gland from the
same prawn. The qPCR analysis revealed that the Mn-IAG expression was decreased with the
decrease of Mn-HSDL1, indicating HSDL1 has positive regulatory effects on IAG in M. nipponense. IAG is a hormone, secreted by androgenic gland, promoting male sexual
differentiation and development in many crustacean species [8-10]. Thus, HSDL1 was
predicted to be involved in the male sexual development in M. nipponense. According to the
histological observations, the number of sperms was decreased with the time of Mn-HSDL1
dsRNA injection. Compared with the control group, the sperms were rarely found at day 14
after Mn-HSDL1 dsRNA injection. This indicated that HSDL1 has positive regulatory effects
on testis development in M. nipponense.

In conclusion, the potentially candidate genes involved in the male sexual development
were selected through performing the long-reads and next generation transcriptome sequencing
of androgenic gland after eyestalk ablation in M. nipponense. qPCR analysis revealed Mn-IAG
was significantly increased after the ablation of both single-side and double-side eyestalk,
indicating the ablation of eyestalk has dramatically regulatory roles on male sexual
development in M. nipponense. The long-reads transcriptome generated 49,480 non-redundant
transcripts. A total of 1,319, 2,092, 4,351 DEGs were identified between CG vs SS, SS vs DS,
and CG vs DS, respectively, indicating the ablation of double-side eyestalk has more regulatory
roles on male sexual development in M. nipponense. Cell cycle, Cellular senescence, Oxidative
phosphorylation, Glycolysis/Gluconeogenesis and Steroid hormone biosynthesis were the
main enriched metabolic pathways in all of these three comparisons, and the important DEGs from these metabolic pathways were identified. qPCR analysis and RNAi analysis of Mn-HSDL1 indicated that HSDL1 has positive regulatory effects on testis development. Overall, this study provided valuable resources for the researches of the mechanisms underlying male sexual development in *M. nipponense* and other crustacean species.

**Materials and Methods**

**Ethics statement**

The permission was obtained from the Tai Lake Fishery Management Council and the committee of Freshwater Fisheries Research Center during the experimental programs. MS222 anesthesia was used to sedate the prawns and shear the tissues.

**Sample collection**

A total of 600 healthy male prawns of *M. nipponense* were collected from a wild population in Tai Lake, Wuxi, China (120°13′44″E, 31°28′ 22″N) with the body weights of 3.63–4.94 g. All the samples were randomly divided and transferred to three 500 L tanks and maintained in aerated freshwater for three days. The three groups were normal prawns (CG), single-side eyestalk ablation prawns (SS), double-side eyestalk ablation prawns (DS). The androgenic glands were respectively collected from these three groups after 7 days of eyestalk ablation, and immediately preserved in liquid nitrogen until used for long-reads and next-generation transcriptomic analysis. Different mature tissues included testis, ovary, hepatopancreas, muscle, eyestalk, gill, heart and brain. Specimens for the different stages of larval and post-larval developmental stages were from the full-sibs population, collected with their maturation process.

**Long-reads transcriptome analysis**

In order to provide sufficient RNA with an aim to establish a reference transcriptome for further analysis, equal amount of androgenic gland of CG, SS and DS (N ≥ 60) were pooled together
to perform the long-reads sequencing. According to the manufacturer’s instructions, UNIQ-10 Column Trizol Total RNA Isolation Kit (Sangon, Shanghai, China) was used to extract total RNA, and an Agilent RNA 6000 Nano kit and chips on a Bioanalyzer 2100 (Agilent Technologies, Santa Clara, CA, USA) were used to measure the RNA integrity. A PacBio RSII platform (Pacific Bioscience Inc., Menlo Park, CA, USA) was employed to construct the long-reads transcriptome. The detailed procedures for the construction of long-reads transcriptome and the analysis of raw sequence data have been well described in our previous study [67].

In the subsequent step, the contaminant sequences were removed by stepwise CLC [68], and the LRS isoforms were annotated [69]. Using Blastp, the transcriptome factors were aligned to the PlnTFDB database (http://plntfdb.bio.uni-potsdam.de/v3.0/), the AnimalTFDB database (http://bioinfo.life.hust.edu.cn/AnimalTFDB/), and the CARD database (https://card.mcmaster.ca/) for selection of genes, involved in the mechanism of male sexual development in *M. nipponense*, using the threshold of E-value $\geq 1 \times 10^{-10}$. Finally, all Blastp results were processed with BLAST2GO [70] for functional annotation.

**Transcriptomic profiling analysis**

The comparative transcriptome analysis of androgenic gland between the CG, SS, and DS were performed. In order to ensure the sufficient amount of RNA samples, androgenic gland from at least 30 prawns were pooled to form one biological replicate, and three biological replicates were sequenced for all of these three groups. The previously published studies have been well described the experimental process [12, 36].

Clean reads were assembled into non-redundant transcripts by using Trinity program (version: trinityrnaseq_r20131110) [71]. The NR protein, the Gene Ontology (GO), the Cluster of Orthologous Groups (COG), and the Kyoto Encyclopedia of Genes and Genomes (KEGG) database were then used to perform the gene annotation, using an E-value cut-off of $10^{-5}$ [12]. Blast2go software was used for functional annotation by GO terms [70]. Blast software was
employed to perform the functional annotation against the COG [72] and KEGG [73] database.

EB-seq algorithm was used to filter the differentially expressed genes, under the criteria of FDR (False discovery rate) < 0.05 [74].

**qPCR analysis**

qPCR was used to measure the relative mRNA expressions of Mn-HSDL1 in different developmental stages, and qPCR verification of important DEGs. The Bio-Rad iCycler iQ5 Real-Time PCR System (Bio-Rad) was used to carry out the SYBR Green RT-qPCR assay. The procedure has been well described in details in previous studies [17-18]. The primers used for qPCR analysis were listed in Table 1. The primers used for qPCR verification of important DEGs were listed in Table 2. EIF was used as reference gene in this study [75].

**RNA interference (RNAi) analysis**

RNAi was performed to analysis the potentially regulatory roles on Mn-HSDL1 in male sexual development in *M. nipponense*. Snap Dragon tools was used to design the specific RNAi primer with T7 promoter site (http://www.flyrnai.org/cgibin/RNAifind_primers.pl), and shown in Table 1. The Transcript Aid™ T7 High Yield Transcription kit (Fermentas, Inc, USA) was used to synthesize the Mn-HSDL1 dsRNA, followed by the procedures of the manufacturer. A total of 300 health mature male *M. nipponense* were collected with body weight of 3.21-4.78g, and divided into two groups. As described in previous study [76-77], the prawns from experimental group were injected with 4 μg/g Mn-HSDL1 dsRNA, while the prawns from control group were injected with equal volume of GFP. The HSDL1 Mrna expression were investigated in the androgenic gland by qPCR after the injection of 1, 7 and 14 days, in order to detect the interference efficiency (N ≥ 5). The Mrna expressions of Mn-IAG were also measured in the same Cdna templates, in order to analysis the regulatory relationship between Mn-HSDL1 and Mn-IAG.

**Histological observation**
The morphological changes of the testis between different days after RNAi treatment was observed by Hematoxylin and eosin (HE) staining. Five testicular samples were respectively collected after 1, 7, and 14 days of RNAi treatment for HE staining. The procedures have been well described in previous studies [78-79]. Olympus SZX16 microscope was used to observe the slides (Olympus Corporation, Tokyo, Japan). The various cell types were labelled based on morphological analysis [5].

Statistical Analysis

Quantitative data were expressed as mean ± SD. Statistical differences were estimated by one-way ANOVA followed by LSD and Duncan’s multiple range test. All statistics were measured using SPSS Statistics 23.0. A probability level of 0.05 was used to indicate significance ($p < 0.05$).

Additional files

Table S1: Summary of BLASTx results for unigenes of androgenic gland long-reads *M. nipponense* transcriptome.

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Available data and materials

The reads of *M. nipponense* transcriptome were submitted to NCBI with the accession number of PRJNA533885.

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Figure 1: Measurement of the expression of Mn-IAG after the ablation of eyestalk. The amount of Mn-IAG mRNA was normalized to the EIF transcript level. Data are shown as mean +SD (standard deviation) of tissues from three separate individuals. Capital letters indicated expression difference between different days in the same group. * ($p < 0.05$) and ** ($p < 0.01$) indicates significant expression difference between different groups at the sample day.

Figure 2: Gene ontology classification of non-redundant transcripts. By alignment to GO terms, 19,673 unigenes were mainly divided into three categories with 52 functional groups: biological process (19 functional groups), cellular component (16 functional groups), and molecular function (17 functional groups). The left y-axis indicates the percentage of a specific category of genes existed in the main category, whereas the right y-axis indicates the number of a specific category of genes existed in main category.

Figure 3: Cluster of orthologous groups (COG) classification of putative proteins.

Figure 4: Verification of the expressions of 10 differentially expressed genes (DEGs) between the androgenic gland of normal prawns, prawns of single-side ablation and prawns of double-side ablation by qPCR. The amounts of DEGs expression were normalized to the EIF transcript level. Data are shown as mean ±SD (standard deviation) of tissues in three separate individuals. Capital letter indicates expression.

Figure 5: Expression characterization of Mn-HSDL1 in different developmental stages. The amount of Mn-HSDL1 mRNA was normalized to the EIF transcript level. Data are shown as mean +SD (standard deviation) of tissues from three separate individuals. Capital letters indicate expression difference between different samples.

Figure 6: Expression characterization of Mn-HSDL1 and Mn-IAG at different days after Mn-HSDL1 dsRNA injection. The amount of Mn-HSDL1 and Mn-IAG mRNA was normalized to the EIF transcript level. Data are shown as mean +SD (standard deviation) of tissues from three separate individuals. Capital letters indicated expression difference between different days after
GFP injection in control group. Lowercase indicated expression difference between different days after Mn-HSDL1 dsRNA injection in RNAi group. * ($p < 0.05$) and ** ($p < 0.01$) indicates significant expression difference between the RNAi group and control group at the sample day.

A: Expression characterization of Mn-HSDL1 at different days after Mn-HSDL1 dsRNA injection. B: Expression characterization of Mn-IAG at different days after Mn-HSDL1 dsRNA injection.

Figure 7: The histological observations of testis between RNAi and control group. SG: Spermatogonia; SC: spermatocyte; S: sperm; CT: collecting tissue. Scale bars = 20 μm.
Table 1. Primers used for qPCR verification

| Primer   | Sequence                        |
|----------|---------------------------------|
| Cyclin B3-F | TGATGAAAGAAGACTCCGCGGT         |
| Cyclin B3-R | AGCGCACCTGGCATATCTTC          |
| MAD2A-F   | ACCCTCCTGAGTCCTTTCACTT        |
| MAD2A-R   | TGCACATGTCTGCCTCAAG          |
| Polo-F     | CGAACTACATCGCCCGAGAA          |
| Polo-R     | AGCGGTCCAATTCCTCGAAGG         |
| Cyclin A-F | CTGCCCTCATCAGTTGCCTTG        |
| Cyclin A-R | AGCTGTGATACCGAATGCCA         |
| Cdc2-F    | ATCAGCGCAGAGTTTTCACA        |
| Cdc2-R    | GAAGAACCTTCAGGTGCACGG       |
| Cyclin B-F | TGGGAGATGTGGGAAATCGG         |
| Cyclin B-R | CCTCAACCTTCGCTTTCTTG        |
| Estrogen-F | CTGCAAAAACCTGCGGGTCAAA    |
| Estrogen-R | CGAGACCTGGGACGTCATTC     |
| Alcohol-F  | CCTTCCTCCAGGGACTCGTA       |
| Alcohol-R  | CCTCATACGACTGACGACCG      |
| SDHB-F     | ACCGCAAGAAGTTGGATGGT         |
| SDHB-R     | TCGATGATCCAACGGGTAGGC       |
| PDHE1-F    | AGCCTAAGCGTTCCAACCTCC       |
| PDHE1-R    | TATTCAGCAGACCTCGTGCC       |
### Table 2. Primers used for HSDL1 analysis

| Primer name | Nucleotide Sequence (5′→3′) | Purpose |
|-------------|-----------------------------|---------|
| HSDL1-RTF   | AGCCTAAGCGTTCCAACCTCC       | FWD primer for GEM expression |
| HSDL1-RTR   | TATTCAGCAGACCTCGTGGG        | RVS primer for GEM expression |
| EIF-F       | CATGGATGTACCTGTGGTGAAC      | FWD primer for β-actin expression |
| EIF-R       | CTGTCAGCAGAAGGTCTCATT       | RVS primer for β-actin expression |
| HSDL1 RNAi-F| TAATACGACTCACTATAGGGGTCTTCAACCGGAAG | FWD primer for RNAi analysis |
| HSDL1 RNAi-R| TAATACGACTCACTATAGGGGTCTTAACCGGAAGG | RVS primer for RNAi analysis |
| Name                                      | Accession number | P-value    | CG vs SS | CG vs DS | SS vs DS | Fold change | Metabolic pathways                                                                 |
|-------------------------------------------|------------------|------------|----------|----------|----------|-------------|-----------------------------------------------------------------------------------|
| SDHB                                      | AIC55101.1       | 3.07E-08   | 0.48     | 0.43     |          |             | Oxidative phosphorylation; Citrate cycle                                          |
| cytochrome c oxidase assembly protein COX11 | XP_004522467.1   | 0.029      | 2.18     | 2.98     |          |             | Oxidative phosphorylation; Thermogenesis                                          |
| cytochrome c oxidase subunit 7A1          | XP_023170779.1   | 1.22E-16   | 2.73     | 2.68     |          |             | Oxidative phosphorylation; Thermogenesis; Parkinson disease                       |
| Acetyl-coenzyme A synthetase 2-like       | XP_018428753.1   | 3.72E-08   | 2.93     | 2.44     |          |             | Glycolysis/Gluconeogenesis; Pyruvate metabolism; Glyoxylate and dicarboxylate metabolism |
| Fructose-bisphosphate aldolase            | XP_018019177.1   | 1.41E-18   | 2.78     | 2.24     |          |             | Glycolysis/Gluconeogenesis; Glycerolipid metabolism;                               |
| alcohol dehydrogenase class -3           | ASW35082.1       | 4.40E-29   | 3.12     | 2.75     |          |             | Glycolysis/Gluconeogenesis; Tyrosine metabolism; Chemical carcinogenesis           |
| estrogen sulfotransferase                 | AJC52502.1       | 5.38E-07   | 4.43     | 3.09     |          |             | Steroid hormone                                                                     |
| 3 beta-hydroxysteroid dehydrogenase      | XP_008216462.1   | 0.001      | 3.07     | 3.12     |          |             | Steroid hormone; Cortisol synthesis and secretion; Aldosterone synthesis and secretion |
| HSDL1                                     | ADB44902.1       | 1.27E-48   | 2.71     | 2.91     |          |             | Steroid hormone                                                                    |
| cyclin-B3                                 | XP_018006504.1   | 1.61E-07   | 0.48     | 0.19     | 0.39     |             | Cell cycle; FoxO signaling pathway; Cellular senescence                           |
| Gene          | Accession     | p-value  | FDR   | Fold Change | Biological Processes                                                                 |
|--------------|---------------|----------|-------|-------------|-------------------------------------------------------------------------------------|
| MAD2A-like   | XP_023320668.1| 1.09E-13 | 0.45  | 0.17        | Cell cycle; Progesterone-mediated oocyte maturation; Oocyte meiosis                  |
| polo-like kinase 1 | AMO03195.1    | 5.47E-18 | 0.33  | 0.08        | Cell cycle; FoxO signaling pathway; Progesterone-mediated oocyte maturation; Oocyte meiosis |
| cyclin A     | AGG40744.1    | 1.21E-15 | 0.49  | 0.15        | Cell cycle; Human papillomavirus infection; Epstein-Barr virus infection; Progesterone-mediated oocyte maturation; Cellular senescence |
| Cdc2 kinase  | ADB44904.1    | 1.87E-27 | 0.45  | 0.13        | Cell cycle; Gap junction; Oocyte meiosis; p53 signaling pathway; Cellular senescence |
| cyclin B     | ADB44902.1    | 8.92E-32 | 0.37  | 0.10        | Cell cycle; Progesterone-mediated oocyte maturation; Oocyte meiosis; FoxO signaling pathway; Cellular senescence; p53 signaling pathway |