Isolation of Differentially Expressed Transcripts by cDNA-AFLP and Expression Analysis of DEAD Box ATP-Dependent RNA Helicase 48 in Ovaries of the Giant Tiger Shrimp *P. monodon*

Sirawut Klinbunga¹, Sirikan Prasertlux¹, Sirithorn Janpoom¹, Puttawan Romgmung¹, Bavornlak Khamnamtong¹,*

¹Aquatic Molecular Genetic and Biotechnology Team, National Center for Genetic Engineering and Biotechnology (BIOTEC), National Science and Technology Development Agency (NSTDA), 113 Paholyothin Road, Khlong Nueng, Khlong Luang, Pathum Thani 12120, Thailand

Abstract

Genes involved in ovarian development of the giant tiger shrimp (*P. monodon*) were isolated by cDNA-AFLP analysis. In total, 75 primer combinations were screened against cDNA of different stages of ovaries (premature ovaries of juveniles and previtellogenic, vitellogenic and mature ovaries of wild intact and eyestalk-ablated broodstock; \( N = 6 \) for each group). Eighteen cDNA-AFLP fragments were cloned and sequenced. Nucleotide sequences of ten fragments significantly matched EF-1\(\alpha\) (7e-63), DEAD box ATP-dependent RNA helicase 48 (*PmDdx48*; 8e-61), tyrosine phosphatase n9 (1e-42), autophagy related protein Atg4-like protein (3e-36), nuclear pore complex protein nup154 (6e-20), GK14382 gene product (1e-20), L-3-hydroxyacyl coenzyme A dehydrogenase short chain (8e-13) and three different hypothetical proteins. The remaining fragments did not match any previously deposited sequence (\( E\)-value > 10\(^{-4}\)). The full-length cDNA of *PmDdx48* was successfully characterized. It was 1646 bp containing an open reading frame (ORF) of 1209 bp corresponding to 402 amino acids.

Introduction

Breeding of pond-reared giant tiger shrimp (*P. monodon*) is rather difficult due to reduced degrees of reproductive maturation of females in captivity (Withyachumnarnkul et al., 1998; Coman et al., 2006). This crucially prohibits the development of effective selective breeding programs of this species. Eyestalk ablation was practically applied for stimulation of ovarian maturation in female penaeid shrimp (Benzie, 1998; Brody, 1998). Nevertheless, an issue on the animal welfare on this approach leads to the requirement of an alternative technique to the control of reproductive maturation without eyestalk ablation (Brody, 1998).

Genetic improvement of *P. monodon* cannot be achieved without knowledge on the control of reproduction. The basic knowledge on molecular mechanisms controlling ovarian development and maturation of *P. monodon* is important and can be directly applied to the shrimp industry (Quackenbush, 2001; Preechaphol et al., 2010). Therefore, identification and characterization of differentially expressed genes during ovarian development of this species is necessary.

Previously, differentially expressed transcripts of reproduction-related genes in ovaries of *P. monodon* were examined using typical EST analysis of vitellogenic ovaries (Tassanakajon et al., 2006; Prechaphol et al., 2007), subtraction subtractive hybridization (SSH) between cDNAs in stages I (previtellogenic) and III (late vitellogenic) (Leelatanawit et al., 2004) and 454 pyrosequencing of those in stages I and IV (mature)
ovaries (Uengwetwanit et al., 2018). Many reproduction-related transcripts were identified. However, isolation of differentially expressed genes involving ovarian (and oocyte) development of *P. monodon* using a simple method like complementary DNA-amplified fragment length polymorphism (cDNA-AFLP) has not been reported.

The cDNA-AFLP was derived from a typical AFLP analysis (Vos et al., 1995). It is a rapid technique for screening of differentially expressed transcripts which does not require any prior knowledge of gene sequences. In this study, we used cDNA-AFLP to isolate transcripts differentially expressed during ovarian development of *P. monodon*. The full-length cDNA of DEAD box ATP-dependent RNA helicase 48 (*PmDdx48*) was further characterized and reported for the first time in this commercially important species. The expression profile of *PmDdx48* was examined in different ovarian developmental stages using quantitative real-time PCR.

**Materials and Methods**

**Experimental Animals and Design**

For cDNA-AFLP analysis, 5-month-old juveniles were collected from Shrimp Genetic Improvement Center (SGIC), Surat Thani located in Southern Thailand (*N* = 6). In addition, wild broodstock were collected from the Andaman Sea and acclimated in the laboratory conditions for 7 days. For the eyestalk ablation group, wild shrimp were unilateral eyestalk-ablated. Ovaries were dissected out from cultured juveniles (*N* = 6) and wild intact and eyestalk-ablated broodstock and weighed. The ovarian developmental stage of juveniles was regarded as premature ovaries while those of broodstock were classified into different stages according to gonadosomatic indices (GSI, ovarian weight/body weight×100): <2, 2-4 and >6% for previtellogenic (stage I), vitellogenic (stage II) and mature (stage IV) ovaries (*N* = 6 for each stage), respectively.

For determination of the expression level of *PmDdx48* during ovarian development, female broodstock of *P. monodon* were live-caught from the Andaman Sea and acclimated under the farm conditions for 7 days. Intact and eyestalk-ablated broodstock were prepared as described above. Ovarian developmental stages were classified by GSI and further confirm by conventional histology and divided to previtellogenic (stage I, *N* = 10 and 4 for normal and eyestalk-ablated broodstock, respectively), vitellogenic (stage II, *N* = 7 and 7), late vitellogenic (stage III, *N* = 7 and 10) and mature (stage IV, *N* = 10 and 11) ovaries, respectively. In addition, juvenile (approximately 5-month-old) shrimp were purchased from a commercial farm and included in the gene expression analysis.

For tissue distribution analysis, various tissues of wild intact females and testes of intact males (*N* = 5 for each sex) were collected. Shrimp tissues were immediately placed in liquid N₂ and kept at -80 °C until needed.

**Total RNA and First-Strand cDNA Synthesis**

Total RNA was extracted from ovaries and other tissues of *P. monodon* using TRI-Reagent (Molecular Research Center). The concentration of extracted total RNA was spectrophotometrically measured. One microgram of DNase I-treated total RNA (0.5 U/μg of total RNA at 37 °C for 30 min) was reverse-transcribed using an Improm-II™ Reverse Transcriptase System (Promega).

**Preparation of cDNA-AFLP Template and Analysis**

Messenger (m) RNA was further purified from ovarian total RNA of juvenile and wild broodstock (stages I, II and IV of intact and eyestalk-ablated shrimp; *N* = 3 for each stage in duplicate) using an Illustra™ QuickPrep Micro mRNA Purification Kit (GE Healthcare). One microgram of mRNA was reverse-transcribed to the first-strand cDNA by combined with 3’ SMART™ CDS Primer II A (5’-AAGCAGTGGTATCAGACAGTACT-3’, 15 μM final concentration), SMART II™ Oligonucleotide (5’-AAGCAGTGGTATACACCGAGTACCGGG-3’, 15 μM final concentration). The reaction was incubated at 72°C for 2 min and cooled on ice for 5 min and 5X First-Strand Buffer, dithiothreitol (DTT), dNTPs and BD PowerScript™ Reverse Transcriptase (Clontech) were added and gently mixed to the final concentrations of 1X, 2 mM and 1 mM, respectively. The reaction mixture was incubated at 42°C for 1.5 h. Two microliters of the reverse-transcribed product were subjected to the second-stranded cDNA synthesis by PCR. The reaction components (50 µl) containing 1X Advantage 2 PCR Buffer, 0.2 mM of each dNTP, 0.6 µM 5’ PCR Primer II A (5’-AAGCAGTGGTATCAGACAGTACT-3’) and 1X BD Advantage 2 Polymerase Mix (Clontech), respectively. The reaction mixture was incubated at 95°C for 1 min followed by 95°C for 15 s, 65°C for 30 s and 68°C for 6 min for 3, 5 and 7 cycles. The synthesized cDNA (3 cycles) was purified by a phenol-chloroform extraction method. The cDNA was ethanol-precipitated and resuspended in 10 μl of TE buffer (10 mM Tris-HCl and 0.1 mM EDTA, pH 8.0). One microgram of the synthesized cDNA was digested with 6 units of Eco RI and Mse I (Vos et al., 1995). The digested cDNA was ligated with restriction site-specific adaptors. Preamplification was carried out utilizing adaptor specific primers with no selective base; E (5’-GACTGCGTACCAATT-3’) and M (5’-GATGAGTCCTGAGTA-3’) or a single selective base on the 3’ terminus of each primer; E_A (5’-GACTGCGTACCAATT-3’) and M_C (5’-GATGAGTCCTGAGTA-3’). The preamplification product was diluted 25-fold, and selectively amplified with primers with additional 2 selective bases at the 3’ terminus of each primer (E_A and M_A and E_C and M_C). The cDNA-AFLP fragments were size-
fractionated through 5% denaturing polyacrylamide sequencing gels and visualized by silver staining (Sambrook & Russell, 2001).

Cloning of Differentially Expressed cDNA-AFLP Fragment

A cDNA-AFLP band was excised from polyacrylamide gel, placed a sterile microcentrifuge tube containing 100 µl sterile deionized water and incubated at 60°C for 1 h. The eluted DNA fragment was re-amplified by PCR. The resulting product was ligated to the pGEM®-T Easy Vector (Promega) in a 10 µl reaction volume containing 5 µl of 2x Rapid Ligation Buffer (60 mM Tris-HCl, pH 7.8, 20 mM MgCl2, 20 mM DTT, 2 mM ATP and 10% PEG8000), 25 ng of the pGEM®-T Easy vector, 3 unit of T4 DNA ligase and 1-3 µl DNA insert. The ligation mixture was incubated overnight at 16°C before transformation into E. coli JM 109 (Sambrook & Russell, 2001). Plasmid DNA was extracted and sequenced for both directions using an automated DNA sequencer.

Rapid Amplification of cDNA End-Polymerase chain reaction (RACE-PCR) of PmDdx48

Gene specific primers for 5’RACE-PCR (R: 5’-GTCAGGCCAGTCCACCTT-3’) and 3’RACE-PCR (F: 5’-TACCGTTACCTCCACGACG-3’) of PmDdx48 were designed. RACE-PCR was carried out using a SMART RACE cDNA Amplification Kit following the protocol recommended by the manufacturer (BD Bioscience Clontech). The amplified fragment was electropheretically analyzed, eluted from the gel, cloned into pGEM-T Easy vector and sequenced. Nucleotide sequences of EST and RACE fragments were assembled and searched against previously deposited sequences in GenBank using BlastX (Altschul et al., 1990). The p/I value and molecular mass of the deduced P. monodon Ddx48 (PmDdx48) protein were examined using ProtParam (http://www.expasy.org/tools/protparam.html). The protein domain and signal peptide in the deduced PmDdx48 protein were predicted using SMART (http://smart.embl-heidelberg.de).

Quantitative Real-Time PCR

Expression of PmDdx48 in different tissues of wild intact females and different ovarian developmental stages of wild intact and eyestalk-ablated P. monodon was examined using quantitative real-time PCR. Standard curves representing 10⁷-10⁸ copies of recombinant plasmids of the target genes (PmDdx48) and the reference gene (EF-1α) were constructed. The expression level of PmDdx48 (F: 5’-TACCGTTACCTCCACGACG-3’ and R: 5’-GTCAGGCCAGTCCACCTTACG-3’) and EF-1α (F: 5’-TACCGTTACCTCCACGACG-3’ and R: 5’-CTTACAGACAGTTCCTCCGTG-3’) in ovaries of each shrimp were amplified in a 10 µl reaction volume contained 5 µl of 2x LightCycler 480 SYBR Green I Master (Roche), 50 (PmDdx48) or 5 ng (EF-1α) of the first-strand cDNA template, 0.3 µM each of gene-specific primers. The thermal profile for quantitative real-time PCR was 95°C for 10 min followed by 40 cycles of 95°C for 30 s, 58°C for 30 s and 72°C for 30 s. The specificity of PCR products was confirmed by melting curve analysis performed at 95°C for 15 s, 65°C for 1 min and 98°C for a continuous fluorescent reading (ramp time = 0.11 °C/s). The cooling process was carried out at 40°C for 30 s. Quantitative real-time PCR of each specimen was carried out in duplicate. The relative expression level (copy number of PmDdx48 and that of EF-1α) between shrimp possessing different stages of ovarian development were statistically tested using one way analysis of variance (ANOVA) followed by Duncan’s new multiple range test. Significant differences between groups of samples were considered when P<0.05.

Results and Discussion

Isolation of Differentially Expressed Transcripts in Ovaries of P. monodon

Analysis of differentially expressed transcripts provides the basic information required for further analysis of molecular reproduction in P. monodon. In this study, cDNA-AFLP analysis was applied for identification of differentially expressed transcripts in ovaries of P. monodon. This method is highly versatile in that both up-regulated and down-regulated genes can be identified by visualization of side-by-side comparisons using various primer combinations.

Initially, the first-strand cDNA was reverse-transcribed from mRNA of P. monodon having different ovarian developmental stages. The second-strand cDNA was synthesized by long distance PCR for 3, 5 and 7 cycles. Smear amplification products were observed in all samples and products from the lowest amplification cycle was chosen for further study (data not shown).

A total of 75 primer combinations were screened. Thirty-five primer combinations yielded positive amplification products were further analyzed by denaturing polyacrylamide gel electrophoresis. Ninety-two cDNA-AFLP fragments showed differential expression between groups of samples (Table 1). The expression patterns of cDNA-AFLP fragments were classified to 5 different groups; those only expressed in juveniles, those preferentially expressed in juvenile than broodstock ovaries and vice versa and those preferentially expressed in ovaries of intact- than eyestalk-ablated broodstock and vice versa, respectively (Table 1).

In the Pacific oyster (Crassostrea gigas), cDNA-AFLP was applied for identifying genes differentially expressed in response to Vibrio tubiashii exposure or between different families. In total, 92 fragments were found from 32 primer combinations. Fourteen cDNA-
AFLP fragments were cloned and sequenced. Genes involved in immune response and metabolic processes and their expression levels were characterized. Most of the differences in transcription patterns between stress-tolerant and stress-sensitive families were found following bacterial exposure (Taris et al., 2009).

Likewise, a large number of differentially expressed cDNA-AFLP fragments were found in the present study. Eighteen cDNA-AFLP fragments including one cDNA-AFLP fragments found only in juveniles, five fragments which were more preferentially expressed in juveniles than broodstock, four fragments displaying a greater expression in wild intact than eyestalk-ablated broodstock and eight fragments showing a greater expression in wild eyestalk-ablated than intact broodstock were cloned and sequenced (Table 1).

Nucleotide sequences of ten fragments were significantly matched elongation factor-1 alpha of P. monodon (E-value = 7e-63), DEAD box ATP-dependent RNA helicase 48 of Aedes aegypti (Ddx48; E-value = 8e-61), tyrosine phosphatase n9 of Culex quinquefasciatus (E-value = 1e-42), autophagy related protein Atg4-like protein of Bombyx mori (E-value = 3e-36), hypothetical protein of Branchiostoma floridae (E-value = 3e-27), hypothetical protein CBG10789 of Caenorhabditis brigasae AF16 (E-value = 5e-24), hypothetical protein LOC100125170 of Xenopus tropacalis (E-value = 2e-24), nuclear pore complex protein nup154 of Nasonia vitripennis (E-value = 6e-20), GK14382 gene product from transcript GK14382-RA of Drosophila willistoni (E-value = 1e-20), L-3-hydroxyacyl coenzyme A dehydrogenase short chain of Ciona intestinalis (E-value = 8e-13) (Table 2). In contrast, those of eight cDNA-AFLP fragments did not match previously deposited sequences in GenBank (E-value > 10e-4) (Table 2).

DEAD box proteins are putative ATP-dependent RNA unwinding proteins whose primary biochemical function is the alteration of RNA secondary structure. They have been implicated in translation initiation, ribosome assembly, RNA splicing, and RNA stability
(Cordin et al., 2006; Cesar Lopez et al., 2008). DEAD-box RNA helicases are required for germ cell functions (Cordin et al., 2006). Some members of this family are believed to be involved in embryogenesis, spermatogenesis, and cellular growth and division. In *Drosophila melanogaster*, a null Ddx1 mutation flies are smaller than the normal and heterozygous flies. Female Ddx1 null flies have reduced fertility with egg chambers undergoing autophagy, whereas males are sterile due to disrupted spermatogenesis (Germain et al., 2015).

The *vasa* gene, which encodes an ATP-dependent RNA helicase belonging to the DEAD-box protein family, was identified in the Pacific white shrimp, *Litopenaeus vannamei* (Alfaro et al., 2007), Chinese shrimp, *Fenneropenaeus chinensis* (Zhou et al., 2010) and the giant freshwater prawn, *Macrobrachium rosenbergii* (Nakkrasae & Damrongphol, 2007). *Vasa* was only expressed in gonads of adults of these species and was proposed to be a potential biomarker for germ cell development. In *Drosophila*, mutations in *vasa* may cause sterility in females with severe defects in oogenesis. Less damaging mutations were also found where oogenesis allows to proceed but the eggs develop into embryos lacking primordial germ cells (PGCs) (Hay et al., 1988; Lasko et al., 1988).

The partial cDNA sequence of *PmDdx48* has been identified from EST analysis of *P. monodon* from both conventional and SSH libraries (Preechaphol et al., 2007 and 2010). However, data on isolation and characterization of this DEAD-box protein gene and its expression profiles during ovarian development of *P. monodon* has not been reported.

### Isolation and Characterization of the Full-Length cDNA of *PmDdx48*

In this study, RACE-PCR was carried out and discrete bands of 900 bp and 1000 bp fragments were obtained from 5' and 3' RACE-PCR of *PmDdx48*. Nucleotide sequences of RACE-PCR fragments were assembled with that of the cDNA-AFLP clone. The full-length cDNA of *PmDdx48* were successfully characterized and reported for the first time in *P. monodon*. It was 1646 bp in length containing an ORF of 1209 bp corresponding to a polypeptide of 402 amino acids with 5' and 3' UTRs of 53 and 418 bp long (excluding the poly A tail), respectively (GenBank accession no. KY982623, Figure 1). Its closest similar sequence was Ddx48 (also called *eukaryotic initiation factor 4A-III*) of *Zootermopsis nevadensis* (*E*-value = 0.0 and identity = 93%). The calculated *pl* and molecular mass of *PmDdx48* was 5.91 and 46.11 kDa, respectively. The predicted signal peptide was not found in the deduced PmDdx48 protein. DEXDc and HELICc domains were found at positions 48-245 (*E*-value = 3.84e-55) and 282-363 (*E*-value = 1.07e-35) of the deduced PmDdx48 protein, respectively. The conserved motif, (V/I)-L-D-E-AD- X-(M/L)-L-X-X-G-F, was observed in all members of the DEAD box protein family (Linder et al., 1989; Tanner and Linder, 2001) was also found in PmDdx48.

### Tissue Expression Analysis of *PmDdx48*

Gene expression and tissue distribution analysis are important and provide the basic information for further analysis of functionally important genes (Klinbunga et al., 2009). Based on the fact that a particular gene may express in several tissues and possesses different functions in different tissues. *PmDdx48* was abundantly expressed in subcuticular epithelium and thoracic ganglion followed by eyestalk. It was comparably expressed in ovaries, testes, and other tissues (hemocytes, heart, pleopod, gill, lymphoid organ, intestine, hepatopancreas and stomach; Figure 2). This suggested that *PmDdx48* should be a

### Table 2. Similarity analysis of candidate differentially expressed cDNA-AFLP fragments in ovaries of *P. monodon*

<table>
<thead>
<tr>
<th>Fragment</th>
<th>Closest similarity</th>
<th>Species</th>
<th><em>E</em>-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>EACT/MCAC-366</td>
<td>L-3-hydroxyacryl coenzyme A dehydrogenase short chain</td>
<td>Ciona intestinalis</td>
<td>8e-13</td>
</tr>
<tr>
<td>ET/A/M210-381</td>
<td>Elongation factor-1 alpha</td>
<td><em>Penaeus monodon</em></td>
<td>7e-63</td>
</tr>
<tr>
<td>ET/A/M10-404</td>
<td>DEAD box ATP-dependent RNA helicase 48</td>
<td><em>Aedes aegypti</em></td>
<td>8e-61</td>
</tr>
<tr>
<td>ET/T/M1-374</td>
<td>Tyrosine phosphatase n9</td>
<td><em>Culex quinquefasciatus</em></td>
<td>1e-42</td>
</tr>
<tr>
<td>ET/T/M2-456</td>
<td>Nuclear pore complex protein nup154</td>
<td><em>Nasonia vitripennis</em></td>
<td>6e-20</td>
</tr>
<tr>
<td>ET/G/M1-462</td>
<td>GK14382 gene product</td>
<td><em>Drosophila willistoni</em></td>
<td>1e-20</td>
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<tr>
<td>ET/G/M1-413</td>
<td>Autophagy related protein Atg4-like protein</td>
<td><em>Bombyx mori</em></td>
<td>3e-36</td>
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<tr>
<td>ET/G/M1-292</td>
<td>Hypothetical protein CBG10789 AF16</td>
<td><em>Caenorhabditis briggsae</em></td>
<td>5e-24</td>
</tr>
<tr>
<td>ET/G/M1-539</td>
<td>Hypothetical protein LOC100125170</td>
<td><em>Xenopus tropicalis</em></td>
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<tr>
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<td>-</td>
<td>&gt;1e-04</td>
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</table>
multifunctional protein playing different functions in various tissues of \textit{P. monodon}.

**Expression Levels of \textit{PmDdx48} During Ovarian Development of Wild \textit{P. monodon} Broodstock**

Transcripts preferentially expressed in different stages of ovarian development can be used as the responsive indicators for reproductive maturation of \textit{P. monodon}. The expression level of \textit{PmDdx48} in ovaries of cultured juveniles was not significantly different from stage I (previtellogenic) ovaries of wild broodstock (\textit{P} > 0.05) but significantly lower than that of subsequent ovarian stages of wild intact adults (\textit{P} < 0.05). This critically implied that \textit{PmDdx48} is functionally involved in ovarian development of \textit{P. monodon}.

In wild intact broodstock, the \textit{PmDdx48} transcript was up-regulated in stages III (late vitellogenic) and IV (mature) ovaries (\textit{P} < 0.05). Its expression level was then decreased at the post-spawning stage (\textit{P} < 0.05). In eyestalk-ablated broodstock, \textit{PmDdx48} was up-regulated in stages IV ovaries (\textit{P} < 0.05). The expression...
profiles of *PmDdx48* in ovaries of wild intact and eyestalk-ablated broodstock suggested that it plays a functionally role during late vitellogenic and mature stages of ovarian development. Eyestalk-ablation resulted in significant reduction of *PmDdx48* in stages III and IV ovaries compared to that of the same stages in intact broodstock (Figure 3). This opens the possible use of RNA interference (RNAi) for induction of ovarian development of *P. monodon*.

In the present study, several reproduction-related transcripts in ovaries of *P. monodon* were identified by cDNA-AFLP analysis. The full-length cDNA and expression profiles of *PmDdx48* were examined. The functional involvement of additional genes in ovarian development of *P. monodon* found in the present study should be further studies for better understanding the reproductive maturation of *P. monodon* in captivity.

**References**


