Comparative Proteomics for Identification of Reproduction-Related Proteins in Testes of the Giant Tiger Shrimp *Penaeus monodon*

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## Introduction

The giant tiger shrimp *Penaeus monodon* is an economically important species (Klinbunga et al., 2001; Rosenberry, 2003; Clifford and Preston, 2006). Farming of *P. monodon* in Thailand relies almost entirely on wild-caught broodstock for supply of juveniles because of poor reproductive maturation of pond-reared *P. monodon* (Withyachumnarnkul et al., 1998; Preechaphol et al., 2007). This issue is one of crucial problems for sustainability of the shrimp farming industry (Leelatanawit et al., 2017). In addition, reduced reproductive maturation in cultured *P. monodon* have limited the ability to selectively improve or maintain important genetic traits in this species.

The production of *P. monodon* is largely constrained by the current dependency on wild-caught broodstock which varies in both quality and quantity (Klinbunga et al., 2012; Leelatanawit et al., 2009; Preechaphol et al., 2007). The domestication and selective breeding programs of this species would provide a more reliable supply of juveniles and improvement of the production efficiency (Makinouchi and Hirata, 1995; Clifford and Preston, 2006; Coman et
al., 2006). The use of selectively bred stocks having improved culture performances on commercially desired traits rather than the reliance on wild-caught stocks is a major mean of sustainability of the shrimp industry (Benzie, 1998; Clifford and Preston, 2006; Coman et al., 2006). Nevertheless, genetic improvement of P. monodon is slow owing to the lack of the basic information related with its gonad development and maturation.

The information related to testicular development and sperm quality in penaeid shrimp is rather limited. Accordingly, an initial step towards understanding molecular mechanisms of testicular development in P. monodon is identification and characterization of reproduction-related genes/proteins expressed in testes of this economically important species.

Proteomic analysis is a powerful and widely used method for identification of differential expressed proteins. Proteomics also provide the basic information on post-translational modification of interesting proteins which cannot be inferred from genomic and transcriptomic analyses. This information is important for understanding molecular mechanisms controlling testicular development of P. monodon. Here, protein profiles in testes of wild and domesticated P. monodon broodstock were identified by GeLC-MS/MS. Expression of serine/arginine-rich protein kinase 3 (PmSrpk3) which play a role in a signal transduction pathway, in testes of wild and different ages of domesticated shrimp was determined. Effects of serotonin (5-HT) and progesterone on its expression in domesticated 18-month-old broodstock were also examined. The information obtained allows better understanding of the reproductive maturation of male P. monodon in captivity.

Materials and Methods

Sampling

For proteomics based on one-dimensional gel electrophoresis (SDS-PAGE) and nano-electrospray ionization-LC-MS/MS (nanoESI-LC-MS/MS), wild male P. monodon broodstock (N=6, average body weight=122.11±7.15; gonadosomatic index, GSI=0.67±0.09%) were collected and further divided to two groups according to the SDS-PAGE protein patterns (N=3 for each group with the average body weight of 123.55±9.36 g, GSI=0.66±0.18% for group A and average body weight of 120.67±11.09 g, GSI=0.68±0.09% for group B, respectively). In addition, domesticated broodstock: 14-month-old (N=3, average body weight=69.84±2.76 g and GSI=0.37±0.03% for group C) and 18-month-old (N=3, average body weight=82.18±2.88 g and GSI=0.37±0.01% for group D) were also included in the experiment.

For expression analysis of PmSrpk3 mRNA, male domesticated P. monodon juveniles (N=3, average body weight=37.80±1.85 g, 6-month-old) and broodstock: 10-month-old (N=3, average body weight=51.24±3.27 g), 14-month-old (N=4, average body weight=62.40±3.87 g), and 18-month-old (N=5, average body weight=74.10±4.28 g) and wild broodstock (N=5, average body weight=126.16±10.75 g), were analyzed. Testes were dissected out from each shrimp, placed in liquid N₂ and kept at -80°C until needed.

To examine effects of serotonin (5-HT) on expression of PmSrpk3, domesticated 18-month-old P. monodon males (average body weight of 74.18±1.85 g, N=40) were sampled and acclimated for 7 days at the laboratory conditions (28-30°C and 30 ppt seawater under natural daylight) in 500-liter aquaria. Eight groups of female shrimp were injected intramuscularly into the first abdominal segment with 5-HT (50 μg/g body weight; the working solution=25 μg/μl in 0.85% NaCl). Specimens were collected at 0, 0.5, 1, 3, 6, 12, 24, 48 and 72 h post injection (hpi). Shrimp injected with the 0.85% saline solution (at 0 hpi) were included as the vehicle control (VC).

Moreover, four groups of acclimated domesticated male shrimp (18-month-old; average body weight of 67.37±2.14 g, N=20) were injected intramuscularly into the first abdominal segment with progesterone (0.1 μg/g body weight; the working solution=1 μg/μl in absolute ethanol). Specimens were collected at 12, 24, and 72 hpi. Uninjected shrimp (at 0 hpi) and those injected with absolute ethanol (at 12 hpi) were included as the negative (NC) and vehicle (VC) controls, respectively. Procedures for animal use in the present study were performed according to the regulation of the Institutional Animal Care and Use Committee (IACUC) of National Center for Genetic Engineering and Biotechnology (BIOTEC).

Total Protein Extraction, Size Fractionation and Mass Spectrometry

Approximately 50 mg of frozen testes of P. monodon was ground to the fine powder in the presence of liquid N₂, suspended in 500 μl of 10% (w/v) trichloroacetic acid in acetone (TCA) containing 0.1% dithiothreitol (DTT) and left at -20°C for 1 h. The homogenate was centrifuged at 10000 g for 10 min at 4°C. The supernatant was collected. An equal volume of 0.85% TCA was added and left at -20°C overnight. The mixture was centrifuged at 10000 g for 30 min at 4°C. The supernatant was discarded and the pellet was washed in acetone containing 0.1% DTT. The sample was centrifuged at 10000 g for 30 min at 4°C. The pellet was air-dried and dissolved in the lysis buffer (30 mM Tris-HCl, 2 M Thiourea, 7 M Urea, 4% CHAPS, w/v). The total amount of extracted proteins was measured using a modification of Lowry’s method (Peterson, 1977). The extracted proteins (10 μg) from wild and captive broodstock were size-fractionated by 12.5% SDS-PAGE (Laemmli, 1970) using a low molecular weight protein standard marker (Bio-Rad) to estimate protein sizes. At the end of each run, the protein gels were silver-stained.
The protein bands were excised according to marker proteins ranges (>225, 176–225, 150–176, 102–150, 76–102, 52–76, 38–52 and <38 kDa) and 4–5 pieces of approximately 1 mm² gel slices were subjected to in-gel digestion with 20 μl of a trypsin solution (10 ng/μl trypsin in 50% acetonitrile/10 mM ammonium bicarbonate). The protein digest was injected into an Ultimate 3000 LC System (Dionex, USA) coupled to an ESI-ion Trap MS (HCT Ultra PTM Discovery System, Bruker, Germany) with electrospray at a flow rate of 300 nl/min to a nanocolumn (Acclaim PepMap 100 C18, 3 μm, 100A, 75 μm inside diameter ×150 mm). A solvent gradient (solvent A: 0.1% formic acid in water; solvent B: 80% acetonitrile in 0.1% formic acid) was run in 40 min.

**Database Search and Quantitation of Characterized Proteins**

After data acquisition, MS/MS ions from nanoLC-MS/MS were analyzed using DeCyder MS Differential Analysis software (DeCyderMS, GE Healthcare) for quantitative protein analysis (Johansson et al., 2006; Thorsell et al., 2007). Acquired LC-MS raw data were converted and the PepDetect module was used for automated peptide detection, charge state assignments, and quantitation based on the peptide ions signal intensities in MS mode. The analyzed MS/MS data from DeCyderMS were subjected to database searches using Mascot software (Matrix Science, London, UK; Perkins et al., 1999) against data of the P. monodon database (http://pmonodon.biotech.or.th), the NCBI nonredundant protein sequence database (nr; http://www.ncbi.nih.gov) and the SWISSPROT database (http://expsys.org/) for protein identification. Database interrogation selections included taxonomy (all entries), enzyme (trypsin), variable modifications (carbamidomethyl, C or oxidation of methionine residues, M), mass values (monoisotopic), protein mass (unrestricted), peptide mass tolerance (1.2 Da), fragment mass tolerance (±0.6 Da), peptide charge state (1+, 2+ and 3+) and max missed cleavages (1). Proteins were considered matched if at least two peptides were found with an individual mascot score corresponding to P<0.05 and P<0.1, respectively.

**RT-PCR and Tissue Distribution Analysis**

Expression of *PmSrpk3* (F: 5′-ATGTTGGTGAAGTGCTGGTGC-3′ and R: 5′-TTTATAGGGAACCCAGTGGC-3′) and *EF-1α* (F: 5′-ATGTTGGTGAAGTGCTGGTGC-3′ and R: 5′-TTTATAGGGAACCCAGTGGC-3′) in various tissues of wild male broodstock and ovaries of female broodstock were analyzed by RT-PCR. The thermal profiles were predenaturation at 94°C for 3 min followed by 30 cycles of 94°C for 30 s, 53°C for 45 s and 72°C for 30 s. The final extension was carried out at 72°C for 7 min. Five microliters of the amplification product was analyzed by 1.5% agarose gel electrophoresis. **Quantitative Real-Time PCR**

Standard curves representing 10²–10⁸ copies of recombinant plasmids of *PmSrpk3* (Primers *PmSrpk3*-F/R) and the internal control, *EF-1α* (F: 5′-TCCGCTTCCCTTCCGACCCTGC-3′ and R: 5′-TTTACAGACAGTTTCTCAGTTG-3′), were constructed. Each concentration was carried out in triplicate. The expression of *PmSrpk3* and *EF-1α* in ovaries of each shrimp was amplified in a 10 μl reaction volume containing 5 μl of 2x LightCycler 480 SYBR Green I Master (Roche), 300 ng (target) or 10 (EF-1α) ng the first strand cDNA template, 0.2 μM each primer. The thermal profile for quantitative real-time PCR was 95°C for 10 min followed by 40 cycles of 95°C for 15 s, 55°C for 30 s and 72°C for 30 s. Real-time PCR of each specimen was carried out in duplicate. The relative expression level of *PmSrpk3* and *EF-1α* mRNA of each specimen was evaluated from their standard curves. The relative expression level (copy number of *PmSrpk3* and *EF-1α*) between different groups of domesticated male shrimp (N=4 each of 6, 10, 14- and 18-month-old), wild (N=4), domesticated 18-month-old shrimp in serotonin injection (vehicle control, VC and 0.5, 1, 3, 6, 12, 24 and 48 hpi; N=4 for each group), and domesticated 18-month-old shrimp in progesterone injection (negative control, NC and VC and 12, 24 and 72 hpi; N=4 for each group), were calculated.

**Statistical Analysis**

The relative expression levels of *PmSrpk3* between male shrimp possessing different stages of testicular development and those in different groups in serotonin and progesterone injection were statistically tested using one-way analysis of variance (ANOVA) and Duncan’s new multiple range test (P<0.05). Results were considered significant when P<0.05.

**Results**

**Size-fractionation of testicular proteins by SDS-PAGE and peptide sequencing by nanoESI-LC-MS/MS**

Total proteins from testes of each shrimp were analyzed by 12.5% SDS-PAGE. The electrophoresed bands were visualized by silver staining (Figure 1). Electrophoretic patterns of size-fractionated testicular proteins of wild shrimp were separated to two groups (A and B). Total soluble proteins of *P. monodon* testes were further analyzed using nanoESI-LC-MS/MS. The intensity of the protein spectrum from testes of wild broodstock group A was used to normalize that of other sample groups. A few thousands of different proteins were identified and approximately 50 proteins that showed differential (up-regulation and down-regulation) expression profiles among sample groups for each molecular weight range were annotated (Supplementary Table 1).
Protein Annotation and Functional Classification

In total, 344 differentially expressed proteins were annotated. Of these, 222 (64.53%) significantly matched known proteins in the database and 122 (35.47%) proteins did not match any protein in the NCBI database and were considered as unknown proteins (Supplementary Table 1). Interestingly, 11 proteins (e.g. p97/VCP-binding protein p135, lipoxigenase homology domains 1, dipeptidyl-peptidase, SEPArease family member, euchromatic histone-lysine N-methyltransferase and gravin; 4.95%) were found in both groups A and B of wild broodstock but not in domesticated broodstock while 152 (68.47%) known proteins were commonly found in all groups of samples (Figure 2 and Supplementary Table 1).

Several proteins seem to be more abundantly expressed in domesticated broodstock than wild broodstock. They were, for example, kinesin like protein 67a, RUN domain containing 2A, GPBP-interacting protein 130b and brain cyclic nucleotide gated 1. In contrast, proteins that the expression level seems to be decreased in domesticated broodstock were, for instance, zinc finger protein 184, glutathione S-transferase alpha 1, seven transmembrane helix receptor, cortactin-binding protein 2, nuclear receptor subfamily 3, group C, member 2 and syntaxin 5 (Supplementary Table 1).

Known proteins in this study were further categorized according to the biological functions of their homologues using the Gene Ontology Categorizer (GoCat software) and 222 differentially expressed

Figure 1. A 12.5% SDS-PAGE showing expression patterns of testes of wild broodstock pattern A (lanes 1-3, panel A) and domesticated 14-month-old (lanes 4-6, panel A), wild broodstock pattern B (lane 1-3, panel B) and domesticated 18-month-old (lanes 4-6, panel B). Lanes M is the protein marker.

Figure 2. Functional classifications of 345 proteins identified from testes of wild (groups A and B; WB-A and WB-B) and domesticated (14- and 18-month-old; DB-14 and DB-18) P. monodon broodstock.
proteins identified in testes of wild and domesticated broodstock of *P. monodon* were able to be classified to 11 functional categories (Figure 3).

These included transport and binding proteins (57 proteins accounting for 16.57%; e.g. arginine kinase 2, brain cyclic nucleotide gated 1, asparagine-rich antigen, deltex 2 and DNA methyltransferase), cell division/DNA synthesis, repair and replication (41 proteins accounting for 11.92%; e.g. zinc finger protein 184, zinc finger RNA binding protein, transcription factor 25, serine/threonine protein kinase and RUN domain containing 2A), signal transduction (36 proteins accounting for 10.47%; e.g. Ran GTPase activating protein 1, vomeronasal type-1 receptor 1, F-box A protein family member fbxa-218, protein tyrosine phosphatase 99A CG2005-PB isoform B and GTP-binding protein alpha subunit gna), structural protein (27 proteins accounting for 7.85%; e.g. calmodulin-regulated giantin, chromosome-associated kinesin KIF4A, dynamin and circadian clock protein PER3), metabolic process (22 proteins accounting for 6.40%; e.g. fatty-acid amid hydrolase, ATP synthase subunit alpha mitochondrial precursor, ATPase, H+ transporting, lysosomal V0 subunit A2, glutathione S-transferase alpha 1 and phosphate transporter), defense and homeostasis (11 proteins accounting for 3.20%; e.g. collagen type IV CG4145-PA isoform A, peptidoglycan recognition protein-ic, immunity-related GTPase M9, ectonucleoside triphosphate and intersecticin long isoform 1), oxidation-reduction (8 proteins accounting for 2.33% including 2,4-dichlorophenol hydroxylase, cytochrome P450 family 4 subfamily A polypeptide 11, NADPH oxidase 4, NADH dehydrogenase Fe-S protein 1 75 kDa precursor isoform 1, oxidoreductase, short chain dehydrogenase/reductase family protein, CG4009 and GH16376), RNA processing (8 proteins accounting for 2.03% including nuclear cap-binding protein subunit 1, nucleoporin 133, glutamyl-tRNA synthetase, glutamyl-tRNA cleavage and polyadenylation specificity factor 1, initiation factor 4B and sfrs8 protein), biosynthetic process (7 proteins accounting for 2.03% including 5'-nucleotidase, dedicator of cytokinesis family protein, GI13543, guanylate cyclase, phosphoribosylformylglycinamidine synthase isoform CRA_a, midasin homolog and regulatory solute carrier protein, family 1, member 1), catabolic process (4 proteins accounting for 1.16% including Uba1a protein, WW and C2 domain containing 2, inositol polyphosphate-4-phosphatase, type II and ubiquitin specific peptidase 38), chaperone (2 proteins accounting for 0.58% including heat shock 40 kDa protein 1 and GL21472), respectively (Figure 3 and Supplementary Table 1).

**Tissue distribution and expression of serine/arginine rich protein-specific kinase 3 (PmSrpk3) in ovaries and testes of *P. monodon***

Serine/threonine protein kinases were identified in our previous (based on 2-DE and nanoESI-LC-MS/MS; Klinbunga et al., 2012) and present (based on Gel-LC-MS/MS) studies. In addition, the partial cDNA sequence of *PmSrpk3* (also called *serine/threonine-protein kinase 23, Stpk23*) transcript was initially identified from EST.

![Figure 3. Categorized classifications of 345 proteins identified from testes of different groups of samples (wild broodstock groups A and B, and domesticated 14-month-old and 18-month-old broodstock) of *P. monodon.*](image-url)
analysis (clone no. TT-N-S01-0903-N, Leelatanawit et al., 2009). It significantly matched serine/threonine-protein kinase 23, Stpk3 of Apis mellifera (E-value=4e-90). Therefore, expression profiles of PmSrpk3 was chosen to examine for demonstration of the contribution of serine/threonine protein kinase genes on testicular development of P. monodon.

At the transcriptional level, PmSrpk3 was constitutively expressed in all examined tissues of P. monodon broodstock and its expression was more preferentially expressed in ovaries than testes of P. monodon juveniles and broodstock (Figure 4).

Expression Profiles of PmSrpk3 in Different Stages of Testicular Development of P. monodon

The expression level of PmSrpk3 in 6-month-old juveniles was not significantly different from that in 10-, 14- and 18-month-old shrimp (P>0.05) but its expression in testes of domesticated shrimp was significantly lower than from that of wild broodstock (P<0.05) (Figure 5).

Effects of Exogenous Progesterone and 5-HT Injection on Expression Levels of PmSrpk3

The expression levels of PmSrpk3 in testes of domesticated 18-month-old P. monodon was not significantly induced by progesterone injection. Although its expression levels in shrimp at 72 hpi seemed to be higher than that in the negative and vehicle controls, the results were not significant owing to large standard deviation between groups of samples (P>0.05) (Figure 6A).

In contrast, the expression level of PmSrpk3 in testes of domesticated P. monodon (18-month-old) upon injecting with serotonin (5-HT) for 1 h was significantly greater than that of the vehicle control (at 0 hpi). Its expression level at other time-intervals seemed to be greater but was not significantly different from the control (P>0.05) (Figure 6B).

Discussion

Comparative Proteomics for Isolation of Differentially Expressed Proteins in Testes of P. monodon

Due to difficulties in reproductive maturation of captive P. monodon, molecular mechanisms of this process have long been of interest by aquaculture industry (Preechaphol et al., 2007). Isolation and characterization of reproduction-related genes in testes of P. monodon have been reported based on express sequence tag (EST), suppression subtractive hybridization (SSH) (Leelatanawit et al., 2004; 2008 and 2009; Wongsurawat et al., 2010) and transcriptomic (Uengwetwanit et al., 2018) analyses. More direct information of molecules functioned on testicular development of P. monodon can be obtained based on cellular proteomics. However, data on reproduction-related proteins in testes of P. monodon is still limited.

In our previous study, typical 2-DE gel electrophoresis and nanoESI-LC-MS/MS was applied for isolation and characterization of protein profiles in testes of wild (GSI=1.08±0.18%, N=3) and domesticated (GSI=0.37±0.05%, N=3 and GSI=0.31±0.05%, N=3) broodstock of P. monodon. In total, 642 protein spots
were characterized and 287 spots (44.70%) significantly matched protein sequences in the databases ($P < 0.05$).

Several reproduction-related proteins, for example, progesterone receptor-related protein p23 (p23), farnesoic-O-methyltransferase (FAmT), cyclophilin A, NADP-dependent leukotriene B4 12-hydroxydehydrogenase (LTB4DH), receptor for activated protein kinase C (RACK), 14-3-3 like protein and several members of ubiquitin-proteosome pathways (e.g. proteosome alpha 3, proteasome beta 6) were identified (Klinbunga et al., 2012).

**Figure 5.** Histograms showing relative expression levels of $PmSrpk3$ during testes development of 6-month-old male juveniles (DJ-6), and 10-month-old (DB-10), 14-month-old (DB-14) and 18-month-old (DB-18) domesticated male broodstock and wild (WB) male broodstock of *P. monodon*.

**Figure 6.** (A) Mean relative expression levels of $PmSrpk3$ in testes of domesticated 18-month-old shrimp injected with progesterone and assessed at 12, 24, and 72 hpi (0.1 μg/g body weight; 18-month-old, $N = 4$ for each group). VC = shrimp injected with absolute ethanol at 12 hpi (vehicle control). Acclimated shrimp without any treatment were included as the negative control (NC). (B) Time-course relative expression levels of $PmSrpk3$ in ovaries of domesticated broodstock at 0, 0.5, 1, 3, 6, 12, 24, 48 and 72 hpi of 5-HT (50 μg/g body weight; 18-month-old, $N = 4$ for each stage). Shrimp injected with 0.85% saline solution at 0 hpi were included as the vehicle control (VC). Bars labeled with the same letters above the histograms reveal non-significant differences between groups of samples ($P < 0.05$).
The use of 2-DE for proteomic analysis is tedious and time consuming. In addition, it is difficult to identify proteins with very low and high molecular weight or those exhibiting very low or high pI simultaneously. In the present study, we use a GeLC-MS/MS (Talahkun et al., 2014) for testicular proteome of *P. monodon*. Extracted proteins are electrophoresed through SDS-PAGE and further characterized by LC-MS/MS. This speed up the process and more cost-effective than the use of a typical 2-DE for proteomic research. In addition, proteins showing differential expression during testicular development of *P. monodon* could be inferred from the protein spectra. This resolves problems from the inability to achieve quantitative results due to a non-stoichiometric binding of silver ions to proteins based on silver staining (Lopez et al., 2000; Choe and Lee, 2003).

Previously, expression patterns of proteasome alpha 3 subunit (*PmPsma3*) and proteasome beta 6 (*PmPsmb6*) mRNAs in domesticated (10-, 14- and 18-month-old) and wild broodstock were examined. Results suggested reduced degrees of reproductive maturation of 18-month-old captive males compared to other groups of male broodstock of *P. monodon* (Klinbunga et al., 2012). In the present study, 14- and 18-month-old males were included to identify reproduction-related proteins that showed differential expression in comparison with wild males.

In total, 344 differentially expressed proteins were annotated. Of these, 222 (64.53%) proteins significantly matched known proteins in the databases. Several reproroduction-related proteins were identified for example, vasa-like protein, p97/VCP-binding protein p135, Wee1-like protein kinase, serine/threonine protein kinase, mitogen activated protein kinase kinase 2, GTP-binding protein alpha subunit, and members of ubiquitin-proteasome pathways (e.g. ubiquitin-specific peptidase).

The *vasa-like protein* encodes an ATP-dependent RNA helicase belonging to the DEAD-box family that, in many organisms, is specifically expressed in germline cells throughout the life cycle. Afalio et al. (2007) characterized the partial cDNA of *vasa-like protein* in *L. vannamei* and it was only expressed in gonads. Valosin-containing protein (VCP also known as p97) belongs to the ATPase-associated with diverse cellular activity (AAA) family of ATPase (Bug and Meyer, 2012). In humans, p97/VCP/Cdc48 has been reported to be required for the mitotic M-phase (Wójcik et al., 2004). The p97/VCP protein is required not only for progression of meiotic metaphase I but also for chromosome condensation at the diakinesis phase in meiotic prophase I (Sasagawa et al., 2007). It functions as a binding protein of sperm-activating and sperm-attracting factor (SAAF) in eggs of the ascidian (*Ciona intestinalis*) (Kondoh et al., 2008).

The gonadosomatic index (GSI) values of wild broodstock were greater than those of domesticated broodstock implying a possible reduction of the maturation potential in domesticated shrimp. The expression profiles of proteins found only in wild broodstock may be used as biomarkers for reduced reproductive maturation of *P. monodon* in captivity. In addition, negative or positive effects of the key proteins on the progression of testicular development may also be inferred from up- or down-regulated proteins compared between wild and domesticated *P. monodon*. Importantly, the preliminary data on differential expression profiles of key testicular proteins of *P. monodon* should be further confirmed by Western blot analysis or enzyme-linked immunosorbent assay (ELISA).

### Expression Levels of Functionally Important Genes in Testes of Wild and Domesticated *P. monodon*

One difficulty in identifying compounds that stimulate crustacean reproduction is the lack of adequate biological markers for reproductive maturation particularly, in *P. monodon*. Gene expression and tissue distribution analysis are important and provide the basic information to set up the priority for further analysis of functional genes.

Expression levels of several reproduction-related genes in testes of male wild and domesticated broodstock of *P. monodon* were examined. *Prohibitin2* (*PmPHb2*) showed an increased expression in captive males (P < 0.05) while reduced expression levels of *meiotic recombination protein DMC1/LIM15 homolog isoform 1* (*PmDmc1*), *cyclophilin A* (*PmCYA*), *small ubiquitin modifier 1* (*PmSUMO1*), *spermatagonial stem-cell renewal factor*, proteasome alpha 3 (*PmPsma3*) and *proteasome beta 6* (*PmPsma6*) in captive *P. monodon* males were reported (P < 0.05; Klinbunga et al., 2012; Leelatanawit et al., 2009).

Protein phosphorylation plays a key role in the signal transduction pathway in most cellular activities including division, proliferation, apoptosis, and differentiation (Hanks et al., 1988; Manning et al., 2002). *Srpk3* is member of cell cycle-regulated protein kinases which phosphorylates serine/arginine (SR) domain-containing proteins in nuclear speckles and mediate the pre-mRNA splicing (Grosso et al., 2008; Jang et al., 2008; Kuroyanagi et al., 1998; Tang et al., 2000). In mammals, SRPK1 and SRPK2 proteins are co-predominantly expressed in testis but highly expressed in pancreas and brain, respectively, whereas SRPK3 is specifically expressed in the heart and skeletal muscle from embryogenesis to adulthood (Bassel-Duby and Olson, 2006; Mylonis and Giannakouros, 2003; Nakagawa et al., 2005; Wang et al., 1998).

A particular gene may express in several tissues and it may possess a different function in different tissues. Tissues distribution analysis of *PmSrpk3* were examined in various tissues of a male broodstock and ovaries of a female broodstock. *PmSrpk3* was not specifically expressed in gonads but widely expressed in various tissues. This suggested that *PmSrpk3* protein may play multifunctional properties in different tissues of *P. monodon*. 
To better understand molecular aspects of the PmSrpk3 gene in reproductive maturation of P. monodon, its expression profiles in different groups of male shrimp were examined. The transcriptional levels of differentially expressed genes in testes could be used as the responsive indicators for reproductive maturation of P. monodon. PmSrpk3 mRNA was not significantly different in different groups of domesticated P. monodon males (P>0.05). Nevertheless, its expression was significantly lower than that of wild broodstock (P<0.05). Expression profiles of testicular PmSrpk3 indicated that reproductive maturation of domesticated male shrimp possibly reduced disregarding the cultivation periods.

Effects of Progesterone and Serotonin (5-HT) Injection on Expression of PmSrpk3

Understanding the induction mechanisms of reproduction-related genes will be useful in developing effective methodologies for inducing reproductive maturation in P. monodon (Ibara et al., 2007). Progesterone is a sex steroid hormone that plays the important roles in gonad development and maturation (Fingerman, 1997; Miura et al., 2006; Yano, 1985). Progesterone and 17α-hydroxyprogesterone injection induced ovarian maturation and spawning in Metapenaeus ensis (Yano, 1985). However, effects of progesterone in reproduction of male shrimp are limited. In this study, effects of progesterone injection on expression of PmSrpk3 transcript were examined. The expression of this gene seemed to be slightly increased from the control in shrimp injected with progesterone for 12, 24 and 72 h. However, results were not statistically significant (P>0.05).

Serotonin (5-HT) has an indirect impact on reproductive maturation by influencing the release of relevant hormones or by modulating the responses of the target tissues to the hormones (Chang et al., 2007; Fingerman, 1992; Sathyanandam et al., 2008). Effects of exogenous serotonin (5-HT) injection on the reproductive performance and maturation of the fiddler crab, Uca pugilator and crayfish, P. clarkii were reported (Sarojini et al., 1993, 1994 and 1995).

The ability to induce reproduction-related genes by exogenous neurotransmitter or steroid hormone administration should reflect an increase in reproductive maturation of male P. monodon adults. In the previous study, positive effects of 5-HT injection on the expression of proteasome alpha 3 (PmPsma3) and proteasome beta 6 (PmPsma6) mRNAs in testes of domesticated 18-month-old P. monodon were reported. Their expression levels upon injecting with 5-HT for 24 h were significantly greater than those of the negative and vehicle controls at 0 hpi (P>0.05) (Klinbunga et al., 2012). In this study, a more rapid effect of inducing gene expression was observed for PmSrpk3 as its expression was significantly increased at 1 h post injection. Practically, biomarkers to indicate male maturation should be developed based on the non-lethal sampling method. This could be done by further analysis on the expression profiles of PmSrpk3, for example, in hemocytes and pleopods of male P. monodon.

In this study, a large number of proteins in testes of P. monodon were identified. The differential expression profiles of proteins identified in testes of P. monodon implied that these proteins may have contributed testicular development in P. monodon. Serotonin administration could induce levels of testicular PmSrpk3 and other reproduction-related transcripts in 18-month-old shrimp. The results suggested that serotonin may be used to promote reproductive maturation of male P. monodon. The information on proteomics of testicular proteins allows further studies to resolve problems on reduced reproductive maturation of P. monodon in captivity.

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