Identification and expression analysis of the Broad-Complex core protein isoform 6 (BR-C Z6) gene in the giant tiger shrimp Penaeus monodon (Penaeidae: Decapoda)

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ABSTRACT. Broad-Complex (BR-C) is an early ecdysone-responsive gene encoding a family of zinc-finger transcription factors that function during metamorphosis in insects. We identified two full-length cDNAs of BR-C Z6 in the giant tiger shrimp (Penaeus monodon). They were 2422 and 2060 bp in length, containing open reading frames of 1440 and 1443 bp, corresponding to polypeptides of 479 and 480 amino acids, respectively. Tissue distribution analysis indicated that PmBR-C Z6 was abundantly expressed in hemocytes and ovaries in juveniles. In broodstock, PmBR-C Z6 was constitutively expressed in all tissues.
examined, and the highest expression was observed in ovaries. The expression of \( PmBR-C \text{ Z6} \) in ovaries was significantly greater than in testes in both juveniles and broodstock of \( P. \text{monodon} \). Quantitative real-time PCR indicated that the expression level of \( PmBR-C \text{ Z6} \) was significantly down-regulated in stages II and III of ovaries in intact wild broodstock and returned to the basal level in stage IV ovaries and after spawning. In eyestalk-ablated broodstock, \( PmBR-C \text{ Z6} \) was significantly up-regulated in stage IV (mature) ovaries. Moreover, the expression level of \( PmBR-C \text{ Z6} \) in vitellogenic, early cortical rod and mature (stages II-IV) ovaries of eyestalk-ablated broodstock was greater than that of the same ovarian stages in intact broodstock. \textit{In situ} hybridization revealed that \( PmBR-C \text{ Z6} \) transcripts were localized in oogonia and cytoplasm of previtellogenic and vitellogenic oocytes of both wild intact and eyestalk-abated broodstock. The effects of 20-hydroxyecdysone on expression of \( PmBR-C \text{ Z6} \) were examined. The expression level of \( PmBR-C \text{ Z6} \) in ovaries of juvenile \( P. \text{monodon} \) was significantly increased at 168 h post-injection. Taken together, these findings indicate that \( PmBR-C \text{ Z6} \) plays an important role in ovarian development of \( P. \text{monodon} \).

**Key words:** Broad-complex; Gene expression; \textit{In situ} hybridization; Real-time PCR; RACE-PCR

**INTRODUCTION**

Ecdysteroids primarily serve as molting hormones in arthropods (Huberman, 2000; Okumura and Sakiyama, 2004). In crustaceans, ecdysteroids are synthesized by the Y-organs, secreted into the hemolymph, and distributed to target tissues for conversion into the active form, 20-hydroxyecdysone (also called crustecdysone, ecdysterone; Subramoniam, 2000). There is evidence that ecdysteroids are also synthesized in the ovaries and testes of crustaceans (Styrishave et al., 2008; Brown et al., 2009). Their production is negatively regulated by the molt-inhibiting hormone, secreted from the X-organ, and positively regulated by methyl farnesoate.

In \( P. \text{monodon} \), the predominant form of ecdysteroids in circulation is 20-hydroxyecdysone (Kuo and Lin, 1996). The peak concentration of ecdysteroids during the \( P. \text{monodon} \) molting cycle coincides with stages D1 and D2 (proecdysis) followed by a rapid decline afterwards (Kuo and Lin, 1996). A similar pattern of hemolymph ecdysteroid concentration was observed in \textit{Litopenaeus vannamei} (Chan, 1995). Ecdysteroid concentrations were shown to be related to both vitellogenesis and molting. Gunamalai et al. (2004) monitored the concentration of 20-hydroxyecdysone in the hemolymph and ovaries through the molting cycle of the mole crab \textit{Emerita asiatica}. In \textit{Drosophila}, the balance between the concentration of 20-hydroxyecdysone and juvenile hormones (JH) seems to play a significant role in the development of the oocytes (Gilbert et al., 1998; Soller et al., 1999; Gruntenko and Rauschenbach, 2008).

\textit{Broad-Complex (BR-C)} is one of the ecdysyteroid-responsive genes, which is a key member of the 20-hydroxyecdysone-regulatory hierarchy that coordinates changes in gene expression during \textit{Drosophila} metamorphosis (Bayer et al., 1996). Typically, the \textit{BR-C} gene family can be divided into 4 isoforms, namely \textit{BR-C Z1}, \textit{Z2}, \textit{Z3}, and \textit{Z4}, which share an amino-
terminal core domain (a BTB domain), which is fused by alternative splicing of different C$_2$H$_2$
znf domains (DiBello et al., 1991; Zollman et al., 1994; Bayer et al., 1996).

Chen et al. (2004) found that the BR-C gene is involved in the 20-hydroxyecdysone-
regulatory hierarchy controlling vitellogenesis in the mosquito Aedes aegypti. Unlike ecdys-
steroid receptors (E74 and E75), the early gene expression of BR-C is activated in previtello-
genic females during a JH-dependent period.

In the cockroach (Blatella germanica), the role of BR-C in embryogenesis was studied
(Piulachs et al., 2010) and 6 isoforms of BR-C (Z1-Z6) were found. The temporal expression
patterns indicate that BgBR-C isoforms are present throughout the embryogenesis of B. germanica, although with weak fluctuations. Silencing all BgBR-C isoforms in the embryo through
parental RNAi elicits a diversity of phenotypes. These phenotypes suggest roles for BgBR-C in
the embryogenesis processes of B. germanica (Piulachs et al., 2010).

To examine the molecular involvement of the BR-C gene in ovarian development of
P. monodon, the full-length cDNA of PmBR-C Z6 was isolated and characterized and reported
for the first time in penaeid shrimp. The effects of eyestalk ablation and 20-hydroxyecdysone
administration on expression levels of ovarian PmBR-C Z6 in broodstock and juveniles were
examined, respectively. Localization of PmBR-C Z6 mRNA and protein in different stages of
oocytes was examined by in situ hybridization (ISH).

**MATERIAL AND METHODS**

**Experimental animals and design**

Female broodstock were wild-caught from the Andaman Sea and acclimated under
farm conditions for 2-3 days. The post-spawning group was immediately collected after shrimp
had ovulated (N = 6). Ovaries were dissected out from each broodstock and weighed. For the
eyestalk ablation group, shrimp were acclimated for 7 days prior to unilateral eyestalk abla-
tion. Ovaries of eyestalk-ablated shrimp were collected at 2-7 days after ablation. In addition,
cultured juveniles (4 months old, N = 6) and domesticated juveniles (6 months old, N = 5) and
broodstock of P. monodon 14 months old (N = 14) and 18 months old (N = 5) were collected
from the Broodstock Management Center, Burapha University (Chanthaburi, Thailand). The
gonadosomatic index (ovarian weight/body weight X100) of each shrimp was calculated. Ovarian
developmental stages were classified by conventional histology (Tan-Fermin and Pudadera, 1989;
Qiu et al., 2005) and divided into previtellogenic (I, N = 10 and 4 for intact and eyestalk-ablated
broodstock, respectively), vitellogenic (II, N = 7 and 6), early cortical rod (III, N = 7 and 9), and
mature (IV, N = 9 and 11) stages, respectively.

In addition, commercially cultured female juveniles of P. monodon (average body
weight of 17.56 ± 3.46 g) were acclimated in the fish tanks for 2 weeks. Five groups of juvenile
shrimp were single-injected intramuscularly with 20-hydroxyecdysone (1 µg/g body weight)
into the first abdominal segment of each shrimp and specimens were collected at 0, 6, 12, 24, 48
72, 96, and 168 h post-injection (hpi). Shrimp injected with 10% ethanol were also included as a
control (at 0 h). Ovaries of each sample were dissected out and immediately placed in liquid N$_2$.
The samples were stored at -80°C prior to RNA extraction and first-strand cDNA synthesis.

For tissue distribution analysis, various tissues of a female and testes of a male juvenile
and broodstock were collected, immediately placed in liquid N$_2$ and kept at -70°C until needed.
Total RNA extraction and first-strand cDNA synthesis

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

Rapid amplification of cDNA end-polymerase chain reaction (RACE-PCR)

Gene specific primers (5′PmBR-C Z6 and 3′PmBR-C Z6) were designed. 5′- and 3′RACE-PCR were carried out using a SMART RACE cDNA Amplification Kit following the protocol recommended by the manufacturer (BD Bioscience Clontech). The amplified fragments were electrophoretically analyzed, eluted from the gel, cloned into pGEM-T Easy vector, and sequenced. Nucleotide sequences of expressed sequence tag (EST) and 5′- and 3′RACE-PCR fragments were assembled and searched against previously deposited sequences in GenBank using BlastN and BlastX (Altschul et al., 1990; available at http://ncbi.nlm.nih.gov). The pI value and molecular weight of the deduced PmBR-C Z6 protein were examined using ProtParam (http://www.expasy.org/tools/protparam.html). The protein domain and signal peptide in the deduced PmBR-C Z6 protein were predicted using SMART (http://smart.embl-heidelberg.de).

RT-PCR and tissue distribution analysis

Expression of *PmBR-C Z6* (primers PmBR-C Z6-RT-F/R; 230 bp) in various tissues of female broodstock and testes of male broodstock was determined by RT-PCR. *EF-1α*500 (primers EF-1α500-F/R) was included as the positive control. The thermal profiles were 94°C for 3 min followed by 25 cycles (*PmBR-C Z6*) and 23 cycles (*EF-1α*500) of denaturation at 94°C for 30 s, annealing at 55°C for 45 s and extension at 72°C for 1 min. The final extension was carried out at 72°C for 7 min. The amplicon was electrophoretically analyzed on 1.5% agarose gels and visualized with a UV transilluminator after ethidium bromide staining (Sambrook and Russell, 2001).

Quantitative real-time PCR

Standard curves representing 10^3^-10^8 copies of *PmBR-C Z6* in triplicate (primers PmBR-C Z6-qRT-F/R; 244 bp) and the internal control, *EF-1α*214 (primers EF-1α214-F/R; Table 1), were constructed. *PmBR-C Z6* and the internal control, *EF-1α*214, in ovaries of each shrimp were amplified in a 10-μL reaction volume containing 5 μL 2X LightCycler 480 SYBR Green I Master (Roche), 100 (*PmBR-C Z6*) or 1 (*EF-1α*214) ng first-strand cDNA template, and 0.2 or 0.3 μM each of respective 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, 53°C (*PmBR-C Z6*) or 58°C (*EF-1α*214) for 30 s and 72°C for 30 s. Real-time PCR of each specimen was carried out in duplicate. The relative expression level (copy number of *PmBR-C Z6* and that of *EF-1α*) between shrimp with different stages of ovarian development was statistically tested using ANOVA (P < 0.05).
In situ hybridization

Ovaries of intact and eyestalk-ablated P. monodon broodstock were fixed in 4\% paraformaldehyde prepared in 0.1\% phosphate-buffered saline (PBS, pH 7.2) overnight at 4°C. The fixed ovarian tissue was washed four times with PBS at room temperature and stored in 70\% ethanol at -20°C until used. Conventional paraffin sections (5 \(\mu\)M) were prepared. The sense and antisense cRNA probes were synthesized from a PCR product using primers PmBR-C Z6-ISH-F/R (Table 1). PCR was carried out in a 25-\(\mu\)L reaction volume containing 10 ng recombinant plasmid containing the complete open reading frame (ORF) of PmBR-C Z6 as the template. PCR was initially performed by predenaturation at 94°C for 2 min followed by 30 cycles of denaturation at 94°C for 30 s, annealing at 60°C for 30 s and at 72°C for 1 min. The PCR product was purified using a MinElute PCR purification kit (QIAGEN). The cRNA probe was synthesized with a DIG RNA Labeling kit (Roche) using the protocol recommended by the manufacturer. The synthesized probes were purified using an RNase-free MinElute\textsuperscript{®} Cleanup kit (QiaGen). The amount of cRNA probes was roughly estimated by dot blot analysis. Tissue sections were dewaxed with xylene and dehydrated in absolute ethanol. The sections were prehybridized with 2X SSC containing 50\% deionized formamide, 1 \(\mu\)g/\(\mu\)L yeast tRNA, 1 \(\mu\)g/\(\mu\)L salmon sperm DNA, 1 \(\mu\)g/\(\mu\)L BSA, and 10\% (w/v) dextran sulfate at 50°C for 30 min and hybridized with either the sense or antisense probe in the prehybridization solution overnight at 50°C. After hybridization, the tissue sections were washed twice with 4X SSC for 5 min each and once with 2X SSC containing 50\% (v/v) formamide for 20 min at 50°C. The sections were immersed in prewarmed RNase A buffer (0.5 M NaCl, 10 mM Tris-HCl, pH 8.0, 1 mM EDTA) at 37°C for 30 min and treated with RNase A (10 \(\mu\)g/mL) at 37°C for 30 min. Tissue sections were washed four times with RNase A buffer (37°C, 10 min each) and twice with 2X SSC (50°C, 15 min each). High-stringent washing was carried out twice in 0.2X SSC at 50°C for 20 min each. The positive hybridization signals were detected with the DIG Wash and Block Buffer kit (Roche) (Qiu and Yamano, 2005).

Table 1. Nucleotide sequences of primers used for characterization and expression analysis of PmBR-C Z6.

<table>
<thead>
<tr>
<th>Primer</th>
<th>Primer sequence</th>
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<tbody>
<tr>
<td>RACE-PCR</td>
<td></td>
</tr>
<tr>
<td>5' PmBR-C Z6</td>
<td>R: 5'-TGATCGGACCACGTGCGAACCAG-3'</td>
</tr>
<tr>
<td>3' PmBR-C Z6</td>
<td>F: 5'-GCCACCAACCACGTGCGAACCAG-3'</td>
</tr>
<tr>
<td>RT-PCR</td>
<td></td>
</tr>
<tr>
<td>PmBR-C Z6-RT-F</td>
<td>F: 5'-ACGCCTACCTCCGCCCCAGGC-3'</td>
</tr>
<tr>
<td>PmBR-C Z6-RT-R</td>
<td>R: 5'-ACGCCTACCTCCGCCCCAGGC-3'</td>
</tr>
<tr>
<td>EF-1(_{\alpha})-F</td>
<td>F: 5'-ATGGTTGTTCAAATTTGCCACC-3'</td>
</tr>
<tr>
<td>EF-1(_{\alpha})-R</td>
<td>R: 5'-TTGACCTCTTGATACAGCAAC-3'</td>
</tr>
<tr>
<td>Quantitative real-time PCR</td>
<td></td>
</tr>
<tr>
<td>PmBR-C Z6-qRT-F</td>
<td>F: 5'-CGCAGAGGGGCCACCAACATCG-3'</td>
</tr>
<tr>
<td>PmBR-C Z6-qRT-R</td>
<td>R: 5'-CTCTGCCCTTCCCCCTCCAGACGTC-3'</td>
</tr>
<tr>
<td>EF-1(_{\alpha})-F</td>
<td>F: 5'-TCCGCTTCCCCCTCCAGACGTC-3'</td>
</tr>
<tr>
<td>EF-1(_{\alpha})-R</td>
<td>R: 5'-CTTTACAGACAGTTTCCTACGTTG-3'</td>
</tr>
<tr>
<td>In situ hybridization</td>
<td></td>
</tr>
<tr>
<td>PmBR-C Z6-ISH-F*</td>
<td>F: 5'-TATAACGACGTCACTATAGGGGGAGGAGATTATATCTCTGCAACAC-3'</td>
</tr>
<tr>
<td>PmBR-C Z6-ISH-R*</td>
<td>R: 5'-ATTTAGGTGACACATAGAACTGTTGACGTCTGCTGTTG-3'</td>
</tr>
</tbody>
</table>

*T7 and Sp6 promoter sequences are boldfaced. RACE-PCR = rapid amplification of cDNA end-polymerase chain reaction.
RESULTS

Isolation and sequence analysis of the full-length cDNA of *PmBR-C Z6*

The partial nucleotide sequence of *PmBR-C Z6* was initially obtained from EST analysis of the ovarian cDNA library of *P. monodon* (Preechaphol et al., 2007). This EST significantly matches the *BR-C Z6* gene of the European honey bee, *Apis mellifera* (*E*-value = 1e-110). Both 5' and 3'RACE-PCR of this gene homologue was further carried out. The positive amplification products of 158 and 1800 (3'RACE-PCR fragment 1) and 1437 (3'RACE-PCR fragment 2) bp in size were obtained. Nucleotide sequences of RACE-PCR fragments and the original EST were assembled. Two types of the full-length cDNAs of *PmBR-C Z6* were found (called *PmBR-C Z6-1* and *PmBR-C Z6-2*; GenBank accession Nos. JN638739 and JN638740, respectively). They were 2422 and 2060 bp in length, containing ORFs of 1440 and 1443 bp, corresponding to the polypeptides of 479 and 480 amino acids, respectively (Figure 1). The calculated pI and molecular mass of the deduced proteins were 5.54 and 53.64 kDa and 5.46 and 53.88 kDa, respectively. Both deduced proteins contained a BTB domain at positions 31-126 (*E*-value = 4.14e-22) and 4 zinc finger (ZnF) C_4H_2 domains (Figure 2). *PmBR-C Z6-1* and *PmBR-C Z6-2* significantly matched the Broad-Complex core protein isoform 6 of *Harpegnathos saltator* (*E*-value = 3e-31 and 4e-31, GenBank accession No. EFN83955). Sequence polymorphism was found including an indel at the amino acid position 248 owing to a tri-nucleotide repeat and 5 amino acid substitutions (positions 190, 203, 204, 440, and 470 where S, H, S, G, and L in *PmBR-C Z6-1* were replaced by P, R, N, E, and P in *PmBR-C Z6-2*, respectively) within the deduced proteins.

RT-PCR and tissue expression analysis of *PmBR-C Z6*

Tissue distribution analysis indicated that *PmBR-C Z6* was abundantly expressed in hemocytes and ovaries of cultured juveniles. A lower level of expression was observed in the hepatopancreas, stomach, lymphoid organs, and pleopods of a female and testes of a male juvenile *P. monodon*. In intact wild broodstock, *PmBR-C Z6* was constitutively expressed in all tissues examined, and the most abundant expression was observed in ovaries (Figure 3). The expression of *PmBR-C Z6* in ovaries was significantly greater than that in testes of both juveniles and broodstock of *P. monodon* (N = 4 for each group; P < 0.05) (Figure 4). In addition, the expression of *PmBR-C Z6* in ovaries of domesticated shrimp was also examined. Its expression level in ovaries of 6-month-old juveniles and 14- and 18-month-old domesticated broodstock was not significantly different (P > 0.05) (Figure 6).

Expression levels of *PmBR-C Z6* mRNA during ovarian development of *P. monodon*

Quantitative real-time PCR revealed that the expression levels of *PmBR-C Z6* in ovaries of juveniles and stage I ovaries of broodstock were comparable. *PmBR-C Z6* was significantly down-regulated in stages II and III of ovaries in intact wild broodstock (P < 0.05) and returned to the basal level in stage IV ovaries and after spawning. In eyestalk-ablated broodstock, its expression level in stage IV (mature ovaries) was significantly greater than that in stage I (previtellogenic) and II (vitellogenic) ovaries (P < 0.05). The expression level of *PmBR-C Z6* in vitellogenic, early cortical rod and mature (II-IV) ovaries of eyestalk-ablated broodstock was greater than that of the same stages of ovaries in intact broodstock (P < 0.05) (Figure 5). In addition, the expression of *PmBR-C Z6* in ovaries of domesticated shrimp was also examined. Its expression level in ovaries of 6-month-old juveniles and 14- and 18-month-old domesticated broodstock was not significantly different (P > 0.05) (Figure 6).
Figure 1. The full-length cDNA and deduced amino acids of *PmBR-C Z6-1* (A) and *PmBR-C Z6-2* (B). The start and stop codons are boldfaced and underlined. The polyA additional signal (AATAAA) is boldfaced, italicized and underlined. A BTB domain (positions 31-126) in both types of the deduced *PmBR-C Z6* is highlighted. Pairwise sequence alignment of *PmBR-C Z6-1* and *PmBR-C Z6-2* (C) is illustrated.
Figure 2. Schematic diagrams representing a BTB and 4 ZnF C₂H₂ domains in the deduced PmBR-C Z6-1 (A) and PmBR-C Z6-2 (B) proteins.

Figure 3. Tissue expression analysis of PmBR-C Z6 (A and C) and EF-1α (B and D) in various tissues of female (HE = hemocytes, GL = gills, HT = heart, OJ = ovaries of a juvenile, HP = hepatopancreas, ST = stomach, IT = intestine, LO = lymphoid organs, TG = thoracic ganglion, ES = eyestalk, PL = pleopods, OB = ovaries of a broodstock) and testes of male (TJ and TB = testes of a juvenile and broodstock, respectively) broodstock (A and B) and juveniles (C and D) of Penaeus monodon. Lanes M represent a 100-bp DNA marker; Lanes N represent a negative control (without the cDNA template).
Figure 4. The relative expression levels of PmBR-C Z6 in testes of cultured juveniles (TT-J) and wild broodstock (TT-B) and ovaries of cultured juveniles (OV-J) and wild broodstock (OV-B) of Penaeus monodon analyzed by RT-PCR (N = 4 for each group). Each histogram corresponds to a particular ovarian stage. The same letters above histograms reveal non-significant differences between groups of samples (P > 0.05).

Localization of PmBR-C Z6 transcript

The cellular localization of PmBR-C Z6 transcripts in ovaries of P. monodon broodstock was determined by ISH. The antisense PmBR-C Z6 probe gave a positive signal in oogonia and cytoplasm of previtellogenic and vitellogenic oocytes in different stages of ovaries of both intact broodstock (Figure 7) and eyestalk-ablated broodstock (data not shown).

Effects of 20-hydroxyecdysone administration on transcription of PmBR-C Z6 in ovaries of P. monodon juveniles

The effects of 20-hydroxyecdysone administration (1 µg/g body weight) on expression of PmBR-C Z6 in ovaries of juvenile P. monodon females were examined at 6, 12, 24, 48, 72, 96, and 168 h after administration. A late response effect of 20-hydroxyecdysone on expression of PmBR-C Z6 was observed at 168 hpi (Figure 8).

DISCUSSION

Characterization and primary structure of PmBR-C Z6

Closing the life cycle of cultured P. monodon is crucial to the sustainability of the shrimp industry. However, poor reproductive maturation of captive females and low quality of spermatozoa of captive males have limited the potential of genetic improvement, which in turn, resulted in remarkably slow domestication and selective breeding programs of P. monodon (Withyachumnarnkul et al., 1998; Preechaphol et al., 2007). Understanding the role of various genes during ovarian and oocyte development of P. monodon may lead to the possible ways to effectively induce ovarian maturation in this economically important species.
Figure 5. Histograms showing relative expression levels of *PmBR-C Z6* during ovarian development of intact (A) and unilateral eyestalk-ablated (B) *Penaeus monodon* broodstock analyzed by quantitative real-time PCR. Data of both intact and eyestalk-ablated broodstock were also analyzed together (C). Each histogram corresponds to a particular ovarian stage. The same letters above histograms reveal non-significant differences between groups of samples (P > 0.05). JN = cultured juveniles, I = previtellogenic; II = vitellogenic; III = early cortical rod; IV = mature ovaries of broodstock; PS = ovaries of intact post-spawning broodstock.
Figure 6. Histograms showing the relative expression levels of *PmBR-C Z6* in ovaries of domesticated juveniles (6 months old) and broodstock (14, and 18 months old) *Penaeus monodon* analyzed by quantitative real-time PCR. Each histogram corresponds to a particular ovarian stage. The same letters indicate that the expression levels were not significantly different (P > 0.05).

Figure 7. Localization of *PmBR-C Z6* transcript during ovarian development of intact *Penaeus monodon* broodstock visualized by *in situ* hybridization using the sense (A) and antisense (B–E) cRNA probes. Conventional hematoxylin/eosin staining was carried out for identification of oocyte stages (F). EP = early previtellogenic oocytes; ECR = early cortical rod oocytes; LCR = late cortical rod oocytes; OG = oogonia; VG = vitellogenic oocytes.
In this study, two full-length cDNAs of *PmBR-C Z6* were successfully characterized and reported for the first time in penaeid species. The *PmBR-C Z6* proteins contain a common N-terminal Bric-a-brac-Tramtrack-Broad (BTB) domain, which is an evolutionarily conserved protein-protein interacting domain (Zollman et al., 1994) that plays critical roles in cell differentiation and development (Costoya and Pandolfi, 2001) and four zinc-finger DNA-binding domains, which function as the transcription regulator. The latter domains directly interact with cis-acting-regulatory elements in the target genes (Zhu et al., 2007). Results from sequence analysis indicated that the different types of *PmBR-C Z6* are allelic variants and should be transcribed from a single locus rather than generated from the alternative splicing process. High sequence similarity between *PmBR-C Z6* and that of other taxa (data not shown) further suggested its functional importance on the signal transduction of the 20-hydroxyecdysone-regulatory pathway in shrimp (Palli et al., 2005; Zhu et al., 2007).

**Expression analysis of *PmBR-C Z6* during ovarian development of *P. monodon***

*BR-C* plays key roles both in coordinating the ecdysone response among tissues and in selective activation or silencing of the downstream effector genes (Thummel, 2002). The role of *BR-C* is well understood in *Drosophila* and moths (Riddiford et al., 2003), but there is limited information about its roles in adult insects.

Tissue distribution analysis revealed abundant expression of *PmBR-C Z6* in hemocytes and ovaries of female juveniles implying its primarily functional involvement in molting during that stage. Interestingly, more abundant expression of *PmBR-C Z6* in ovaries than in other tissues was observed in *P. monodon* female broodstock. This suggested that *PmBR-C Z6* may play the alternative role in reproduction in shrimp broodstock. *PmBR-C Z6* was more abundantly expressed in ovaries than testes in both juveniles and...
broodstock. Therefore, *PmBR-C Z6* may play a more important role in oogenesis than in spermatogenesis of *P. monodon*.

Ovarian maturation of *P. monodon* results from rapid synthesis and accumulation of a major yolk protein (vitellin) (Meusy and Payen, 1988; Yano and Hoshino, 2006). Unilateral eyestalk ablation is used in practice to induce ovarian maturation in penaeid shrimp, but this technique leads to the eventual loss of egg quality and the death of spawners (Benzie, 1998; Okumura and Sakiyama, 2004). Therefore, predictable maturation and spawning of captive penaeid shrimp without the use of eyestalk ablation is a long-term goal for the industry (Quackenbush, 2001).

The expression level of *PmBR-C Z6* fluctuated slightly during ovarian development (i.e., down-regulation in stages II and III before returning to the normal level in stage IV and after spawning) of intact broodstock. Eyestalk ablation clearly promoted the expression of *PmBR-C Z6* during vitellogenesis and final maturation of ovaries compared to intact broodstock for approximately 3-6 times. This suggested that gonad-inhibiting hormone, a polypeptide hormone synthesized in the eyestalk (Treerattrakool et al., 2008), has a prominent effect on the transcription of *PmBR-C Z6*. Accordingly, the expression level of this gene may be used as a molecular indicator for following the progression of ovarian development and the final maturation of *P. monodon* broodstock as a consequence of maturation-inducing feed (Meunpol et al., 2005) and/or exogenous hormonal administration (e.g., progesterone and 17β-estradiol; Yano, 1987; Yano and Hoshino, 2006; Meunpol et al., 2007).

The high level of expression of *PmBR-C Z6* was also observed in domesticated shrimp (6, 14 and 18 months old). The expression levels of *PmBR-C Z6* in ovaries of domesticated 14-month-old broodstock seem to be increased compared to other ages of domesticated stocks, but results were not statistically significant owing to a large standard deviation within each group of shrimp samples (P > 0.05).

The development of oocytes consists of a series of complex cellular events, in which different genes are expressed to ensure the proper development of oocytes and to store transcripts and proteins as maternal factors for early embryogenesis (Qiu et al., 2005). Recently, the full-length cDNA of *progestin membrane receptor component 1* (*Pgmrc1*) in *P. monodon* was characterized. *PmPgmrc1* was 2015 bp in length, containing an ORF of 573 bp, corresponding to a polypeptide of 190 amino acids. Quantitative real-time PCR indicated that the expression level of *PmPgmrc1* mRNA in ovaries of both intact and eyestalk-ablated broodstock was greater than that of juveniles (P < 0.05). Like *PmBR-C Z6*, *PmPgmrc1* was up-regulated in mature (stage IV) ovaries of intact broodstock (P < 0.05). Unilateral eyestalk ablation resulted in an earlier up-regulation of *PmPgmrc1* as of the vitellogenic (II) ovarian stage. Moreover, the expression level of *PmPgmrc1* in vitellogenic, early cortical rod and mature (II-IV) ovaries of eyestalk-ablated broodstock was greater than that of the same ovarian stages in intact broodstock (P < 0.05). *PmPgmrc1* mRNA was clearly localized in the cytoplasm of follicular cells and previtellogenic and early vitellogenic oocytes. Immunohistochemistry revealed positive signals of the PmPgmrc1 protein in the follicular layers and cell membrane of follicular cells and various stages of oocytes (Preechaphol et al., 2010b).

In addition, the full-length cDNA of *progesterone receptor-related protein p23* (*p23*), an essential component of the heat shock protein 90 (Hsp90) molecular chaperone complex with the progesterone receptor (Johnson and Toft, 1994), was recently character-
Pm-p23 was 1943 bp, comprising an ORF of 495 bp corresponding to 164 amino acid residues and the 5' and 3' UTRs of 7 and 1441 bp, respectively. Quantitative real-time PCR analysis revealed that the expression levels of Pm-p23 in ovaries of both intact and eyestalk-ablated broodstock were significantly greater than that of juveniles (4-month-old shrimp) (P < 0.05). Pm-p23 was up-regulated starting in stage II ovaries of intact and stage III ovaries of eyestalk-ablated P. monodon broodstock (P < 0.05). The mRNA level of Pm-p23 after spawning was not significantly different from stage II-IV ovaries of intact broodstock (P < 0.05). In situ hybridization indicated that Pm-p23 was localized in ooplasm of previtellogenic oocytes. The recombinant Pm-p23 protein was successfully expressed in vitro, and its polyclonal antibody was successfully produced. Western blot analysis indicated that the level of ovarian Pm-p23 protein peaked at the vitellogenic stage and decreased as oogenesis progressed (Preechaphol et al., 2010a).

Considering the findings of this study together with those of our previous studies, progesterone- and ecdysteroid-related genes seem to play an important role in ovarian development and maturation of P. monodon oocytes.

Localization of PmBR-C Z6 transcripts in ovaries of P. monodon

In situ hybridization was used to determine the cellular localization of PmBR-CZ6 mRNAs in ovaries of wild P. monodon broodstock. PmBR-C Z6 was localized in oogonia and ooplasm of previtellogenic and vitellogenic oocytes in different stages of ovarian development of both intact and eyestalk-ablated broodstock. Generally, more intense signals were observed in ovaries of eyestalk-ablated broodstock than intact broodstock. This also implied the effects of eyestalk ablation on the transcription of this gene. In situ hybridization signals of PmBR-C Z6 were not observed in follicular cells and more mature (early cortical rod and mature) stage oocytes.

Contradictory results from quantitative real-time PCR and ISH on the disappearance of PmBR-C Z6-1 hybridization signals from the ooplasm in oocytes at later stages (early cortical rod and mature oocytes) may have been due to a significant increase in oocyte size as oogenesis proceeded. In addition, real-time PCR detects gene expression with much greater sensitivity than ISH. A similar circumstance was also observed in PmPgmrc1 (Preechaphol et al., 2010a) and Pm-p23 (Preechaphol et al., 2010b).

Effects of ecdysteroid administration on expression of PmBR-C Z6 in ovaries of P. monodon

Domestication of P. monodon is impeded by poor reproductive maturation of both male and female brooders in captivity (Withyachumnarnkul et al., 1998; Preechaphol et al., 2010a). Effects of hormones (e.g., progesterone and other sex steroids) and neurotransmitters (e.g., serotonin and dopamine; Fingerman, 1997) on expression of reproduction-related genes and, in turn, on ovarian development of P. monodon are of interest by aquaculturists and could be directly applied to the shrimp industry.

In the mosquito (A. aegypti), the 5'-regulatory region of vitellogenin contains the binding site for the ecdysone receptor (EcR)/homologue of the retinoid X receptor, ultraspiracle (USP), BR and ecdysteroid receptors; E74 and E75 (Kokoza et al., 2001; Chen et al., 2004; Zhu et al., 2007). The EcR/USP dimer binds to the vitellogenin promoter and acts synergistically with
E74 to activate vitellogenin expression (Zhu et al., 2003, 2007). We hypothesized similar effects of 20-hydroxyecdysone on the stimulation of the *BR-C* Z6 gene in *P. monodon*. This should be proved when a large number of domesticated broodstock are available for the experiments.

Molecular effects of 20-hydroxyecdysone on the expression of reproduction-related genes in penaeid shrimp have not been reported. Due to the high cost (~$200USD/brooder) of a gravid female of *P. monodon* (Preechaphol et al., 2007), approximately 4-month-old commercially cultured shrimp were used to examine the effects of 20-hydroxyecdysone on the expression of ovarian *PmBR-C* Z6.

Treated juvenile shrimp showed immediate response to exogenous 20-hydroxyecdysone administration by molting within 48 h following the treatment in most individuals of all treatments. The expression level of *PmBR-C* Z6 was up-regulated at 168 hpi. Differential expression of *PmBR-C* Z6 should have reflected long duration effects of the ecdysteroid on the expression of these genes. The molecular effects of 20-hydroxyecdysone on this expression should be further examined in both wild and domesticated broodstock to evaluate the possible use of ecdysteroids for enhancing ovarian/oocyte development in *P. monodon*.

In the present study, the ecdysteroid-responsive gene of *P. monodon* was identified. The expression profile of *PmBR-C* Z6 in ovaries of intact and eyestalk-ablated *P. monodon* broodstock implied that it should contribute to ovarian development of *P. monodon*. Functionally, analysis of genes and proteins involving ovarian development can be further carried out for better understanding of the reproductive maturation of female *P. monodon* in captivity.

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