Molecular Structure and Organization of Crustacean Hyperglycemic Hormone Genes of *Penaeus monodon*

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The Crustacean hyperglycemic hormone (CHH) has been shown to exist as multiple molecular forms in several crustacean species. In *Penaeus monodon*, a gene encoding CHH (so-called *Pem-CHH1*) was recently described. In this study, the molecular structures of two other CHH genes (*Pem-CHH2* and *Pem-CHH3*) are reported. Both the *Pem-CHH2* and *Pem-CHH3* genes contain three exons that are separated by two introns that are similar to the structure of other genes in the same family. An analysis of the upstream nucleotide sequences of each *Pem-CHH* gene has identified the putative promoter element (TATA box) and putative binding sites for several transcription factors. The binding sites for CREB, Pit-1, and AP-1 were found upstream of all three *Pem-CHH* genes. A Southern blot analysis showed that at least one copy of each *Pem-CHH* gene was located within the same 10 kb genomic DNA fragment. These results suggest that the CHH genes are arranged in a cluster in the genome of *P. monodon*, and that their expression may be modulated by similar mechanisms.

**Keywords:** CHH, Gene organization, Gene structure, *Penaeus monodon*, Regulatory elements

**Introduction**

The crustacean hyperglycemic hormone (CHH) is the most abundant neuropeptide that is produced by a group of peptidergic neurons in the medulla terminalis X-organ of decapod crustacean (Charmantier *et al.*, 1997). The primary function of CHH is to regulate the glucose level in the hemolymph. CHH belongs to a peptide hormone family that is mainly comprised of the CHH, molt-inhibiting (MIH), gonad-inhibiting, (GIH) and mandibular organ-inhibiting hormones (MOIH) (Keller, 1992). Peptides in this family share unique characteristics: all are composed of 72-78 amino acids and possess six cysteine residues in identical positions (Chang, 1997).

The amino acid sequences of CHH from several species have been determined (Kegel *et al.*, 1989; Kegel *et al.*, 1991; Tensen *et al.*, 1991b; Huberman *et al.*, 1993; Martin *et al.*, 1993; Yang *et al.*, 1995; Gu and Chang, 1998). The presence of multiple molecular forms of CHH in a single species was observed. For example, two structurally different CHHs that were 92% identical in their amino acid sequences (so-called CHH-A and CHH-B) were reported in the lobster *Homarus americanus* (Tensen *et al.*, 1989). The cDNAs encoding these two forms of CHH in *H. americanus* were subsequently obtained (Tensen *et al.*, 1991a). Two distinctive forms of CHH were also found in the crayfish *Procambarus clarkii* (Huberman and Aguilar, 1988) and shrimp *Metapenaeus ensis* (Gu *et al.*, 2000). Moreover, in the shrimp *Penaeus japonicus*, at least five neuropeptides possessed hyperglycemic activity (Yang *et al.*, 1997). In addition, two isoforms, resulting from the configurational difference of a single amino acid residue (L- or D-Phe at position 3), were observed for each CHH-A and CHH-B in *H. americanus*. These isoforms had the same molecular mass and showed similar hyperglycemic activity (Soyez *et al.*, 1994). Similar stereo-inversion was observed in CHH from crayfish *P. clarkii* (Yasuda *et al.*, 1994).

Recently, a gene encoding CHH from the black tiger shrimp *P. monodon* was characterized (Udomkit *et al.*, 2000). We report here the identification and characterization of the genes encoding another two CHH peptides from *P. monodon*. The structure and putative promoter region of these three *Pem-CHH* genes were analyzed and compared. Preliminary data on the organization of these CHH genes was also described.

**Materials and Methods**

Amplification of *Pem-CHH2* and *Pem-CHH3* genes Genomic
DNA was prepared from abdominal muscle tissue of *P. monodon* by using the Qiagen Genomic-tip and Genomic DNA buffer set (Qiagen, Chatsworth, USA). The PCR reaction contained 150 ng of the genomic DNA template, 20 mM Tris-HCl (pH 8.4), 50 mM KCl, 2 mM MgCl₂, 200 nM each primer, and 200 μM each of dATP, dCTP, dGTP, and dTTP in a total volume of 50 μl. The reaction was heated to 95°C for 5 min then 2.5 units of *Tth* DNA polymerase (Biotools, Canada) were added. Amplification was achieved by 35 successive cycles of denaturation at 94°C for 1 min, annealing at 50°C for 1 min and extension at 72°C for 2 min, followed by a 10 min-final extension at 72°C. The primers that were used to amplify the genes were designed from the 5’ and 3’ ends of the corresponding cDNA. The nucleotide sequence of these primers were CHH-F: 5’ CGGAGATTCTCAGTGCAAGGGAGAGGC 3’ and CHH2-R: 5’ GCGGTACCTGCTTTATGAGACACTG 3’ for the *Pem-CHH2* gene and CHH-F and CHH3-R: 5’ ATGCTTTATGAGACACTAC 3’ for the *Pem-CHH3* gene.

**Genomic walking into the 5’ upstream sequences of *Pem-CHH* genes** The unknown 5’ DNA sequences (upstream of the *Pem-CHH1, Pem-CHH2, and Pem-CHH3* genes) were accessed by the strategy of the Universal Genome Walker™ Kit (CLONTECH, Palo Alto, USA). Genomic DNA of *P. monodon* was digested with either *DraI* or *SnaBI* I to generate blunt-ended genomic fragments. The *DraI* and *SnaBI* I-digested genomic DNA was then ligated to the Genome Walker Adapter in separate reactions, which resulted in pools of the adapter-ligated genomic fragments, which were referred to as *DraI* and *SnaBI* I Genome Walker libraries, respectively. The unknown flanking sequences of the individual *Pem-CHH* genes were amplified from the genome walker libraries with outer adapter (AP1) and outer gene specific primers (GSP1) that were designed in order to extend the amplification from the 5’ terminus of each Pem-CHH gene into their upstream sequences. Secondary PCR was performed with nested adapter (AP2) and nested gene specific primers (GSP2) in order to obtain specifically amplified products. The PCR reactions were carried out according to the manufacturer’s instruction. The gene specific primers for the *Pem-CHH2* gene were as follows: CHH2-GSP1: 5’ TAAAGGAAAA AATCGGAGCTTTGCTGAA A 3’ and CHH2-GSP2: 5’ CAAATGAGATAATGCATAACGTCGAGA A 3’ and for *Pem-CHH3* gene were CHH3-GSP1: 5’ AGGGGATATACAGGGCTTTAATCCA T 3’ and CHH3-GSP2: 5’ CAGGTGAAGCTGCAATTACAGAGC 3’.

**DNA sequence analysis** The PCR products were cloned into the pGEM®-T Easy vector (Promega, Madison, USA). Double-stranded DNA sequencing was performed by the ABI PRISM™ BigDye Terminator Cycle Sequencing Kit (Applied Biosystems, Foster City, USA). Searches for the transcription factor-binding site were performed using Tisitescan software on the MIRAGE WWW server (http://www.motif.org/cgi-bin/fti/Tisitescan.pl) and Signal Scan software on the BIMAS WWW server (http://bimas.dccr.nih.gov/molbio/signal).

**Southern blot analysis** About 10 μg of *P. monodon*’s genomic DNA was digested with selected restriction enzymes. The genomic fragments were separated on 0.7% agarose gel at 30 V for 21 h. They were subsequently transferred to the GeneScreen Plus™ hybridization transfer membrane (DuPont, Boston, USA) with 0.4 N NaOH using a vacuum blotter (Model 785, Bio-RAD, Hercules, USA).

Three DNA probes that are specific to the *Pem-CHH1, Pem-CHH2*, and *Pem-CHH3* genes were synthesized from the intronic region of each gene. The fourth probe was synthesized from the coding region of the mature *Pem-CHH1*, which is almost identical to the same region of the other two *Pem-CHH* genes. The primers that were used for amplification of each probe were as follows: forward *CHH1*-IF (5’ AACCCCAACCACTTTGTTGTTGTTGTTG 3’) and reverse *CHH1*-IR (5’ CCACCTGATGTTCTCAGAGG 3’) for the *CHH1*-specific probe, forward *CHH2*-IF (5’ TCCGATAGGG GATCCGA 3’) and reverse *CHH2*-IR (5’ TCAGTTTTTGGCGAGGAGG 3’) for *CHH2*-specific probe, forward *CHH4*-IF (5’ TCAGTGCAGAGGGAGAGG 3’) and reverse *CHH2*-IR followed by *AccI* digestion for the *CHH3*-specific probe and forward *CHH*-mat (5’ GGAATCCATAGGCTATCCTCAG 3’) and reverse CD-R (5’ CGGATCCTACTTGGCAGGCTC TG 3’) for the mature CHH1 probe.

The DNA probes were synthesized by the incorporation of Fluorescein-11-dUTP with the Gene Image random prime-labeling module (Amersham Biosciences, Upsala, Sweden) following the manufacturer’s protocol. Hybridization was performed in a hybridization solution (5 × SSC, 0.1% (w/v) SDS, 5% (w/v), dextran sulphate, 20-fold dilution of liquid block, and 100 μg/ml denatured salmon sperm DNA) at 65°C overnight. After hybridization, the first wash was performed in 1 × SSC, 0.1% (w/v) SDS at 65°C for 15 min, followed by the second wash in 0.1 × SSC, 0.1% (w/v) SDS at the same temperature for 15 min. The detection step was conducted according to the protocol of the Gene Images CDP-Star detection module (Amersham Pharmacia Biotech).

**Results**

**Nucleotide sequence analysis of *Pem-CHH2* and *Pem-CHH3* genes** Specific DNA fragments of about 1.2 kb were amplified from *P. monodon*’s genomic DNA using primers that were designed from both ends of the 5’ and 3’ cDNA encoding *Pem-CHH2* and *Pem-CHH3* (unpublished data). Figures 1A and 1B show the nucleotide sequences of these *Pem-CHH2* and *Pem-CHH3* genes, as well as the primers that were used for amplification. A sequence analysis using the Gene Finder Program of Baylor College of Medicine (BCM) via the worldwide web revealed that both genes contained three exons and two introns. The donor and acceptor consensus sequences of the splice site (GT-AG) were detected at all exon-intron boundaries. The first and second introns of *Pem-CHH2* were 406 and 268 bp long, whereas those of *Pem-CHH3* comprised 370 and 282 bp, respectively. The first intron interrupted the sequence encoding the signal peptide, while the second intron separated the mature peptide coding sequence in both the *Pem-CHH2* and *Pem-CHH3* genes (Figs. 1A and B). Figure 2 shows the schematic diagram comparing the structure of *Pem-CHH2* and *Pem-CHH3* genes to that of the previously characterized *Pem-CHH1* gene.
Analysis of the putative promoter and probable regulatory elements  The 5’ upstream sequences (about 500 nucleotides from the transcription start site (+1) of each *Pem-CHH* genes) were analyzed. Sequence similarities were observed between the 200 nucleotides upstream from the +1 position of the *Pem-CHH2* and *Pem-CHH3* genes, where the same region of *Pem-CHH1* showed a higher degree of variation (Fig. 3). In addition, three types of dinucleotide repeats (i.e. AC, AT, and GT) were found in the further upstream nucleotide sequence of the *Pem-CHH1* gene. No similar repeats were detected in the 5’ upstream sequence of the *Pem-CHH2* and *Pem-CHH3* genes that were analyzed in this study.

**Fig. 1.** Nucleotide and deduced amino acid sequences of the *Pem-CHH2* (A) and *Pem-CHH3* (B) genes of *P. monodon*. Introns are indicated by dashed lines. Amino acids are shown in a one-letter symbol and the asterisk marks the stop codon. Arrows indicate the nucleotide sequences corresponding to the primers used for amplification of each gene.

**Fig. 2.** Schematic diagram comparing the structure of the *Pem-CHH1* (Udomkit et al., 2000), *Pem-CHH2*, and *Pem-CHH3* genes. The hatched boxes represent the 5’ and 3’ untranslated regions. Exons and introns are shown in white and black boxes, respectively.
element (TATAA) that was located at 23 nucleotides upstream of the transcription start site of each gene. In addition, several potential recognition sites for transcription factors were identified (Fig. 4). The putative binding sites for the cAMP responsive element binding protein (CREB), pituitary-specific transcription factor-1 (Pit-1), and activator protein-1 (AP-1) were found in the upstream region of all three CHH-like genes. Moreover, recognition sequences for other transcription factors (i.e. NF-1, CP1, and CP2) have also been identified.

Analysis of the copy numbers and organization of Pem-CHH genes in the genome of P. monodon  A Southern blot analysis was performed in order to gain preliminary data on the copy numbers and organization of the Pem-CHH1, Pem-CHH2, and Pem-CHH3 genes in P. monodon’s genome. The nucleotide sequences that are specific to each Pem-CHH gene were used as a probe to detect individual Pem-CHH genes in the genome. These probes were unable to hybridize to one another (Fig. 5A). P. monodon’s genomic DNA was separately digested with three restriction enzymes that do not cut within any of the three Pem-CHH genes (i.e. BglI, EcoR I, and Hind III). The result in Fig. 5B shows that there were at least 3 copies of the Pem-CHH1 gene, one copy of the Pem-CHH2 genes, and 2 copies of the Pem-CHH3 gene in the genome. Some common DNA fragments were detected by more than one probe. A fragment of about 7 kb in the EcoR I-digested DNA was detected by both of the CHH2 and CHH3 probes, and the 9 kb band in EcoR I-digested DNA was detected by both of the CHH1 and CHH3 probes. In addition, the band above 10 kb in the Hind III-digested DNA was detected by all of the probes. All of the DNA fragments that were detected by each gene-specific probe, together with several additional bands, were hybridized to the mature CHH 1 probe (Fig. 5C).

Discussion

Despite the similarity in their amino acid sequences, Pem-CHH2 and Pem-CHH3 were shown to be encoded from genes that are structurally different from the one encoding Pem-CHH1. While Pem-CHH1 contains information that is encoded from two exons (Udomkit et al., 2000), Pem-CHH2 and Pem-CHH3 are both the translated products from three exons. The gene structure of CHH, MIH, and MOIH in other species is similar to that of Pem-CHH2 and Pem-CHH3 (i.e. containing three exons and two introns) (Gu and Chang 1998; Gu et al., 2000; Lu et al., 2000). Compared to the genes for other CHHs, the Pem-CHH1 gene lacks the first intron that separates the coding region for the signal peptide in the others. The only intron of the Pem-CHH1 gene is located at the same position as intron 2 of Pem-CHH2 and Pem-CHH3 genes.

In this study, about 500 nucleotides of the upstream regions from the transcription start site (+1) of the Pem-CHH1, Pem-CHH2, and Pem-CHH3 genes were characterized. A similarity in the nucleotide sequence within about 200 nucleotides upstream from the +1 position among the three Pem-CHH genes suggests that they occurred through gene
duplication events. Likewise, a high degree of similarity was also found in the 400 nucleotides 5' upstream regions of three CHH-like genes in *Metapenaeus ensis* (Gu and Chan, 1998). Although the coding regions are highly conserved among the three *Pem*-CHH genes, the 5' upstream region and the introns of *Pem*-CHH2 and *Pem*-CHH3 are more similar to each other than to that of *Pem*-CHH1. A comparison of the downstream sequences of these three CHH genes also showed that the nucleotide sequence in this region of *Pem*-CHH1 was less conserved than that of *Pem*-CHH2 and *Pem*-CHH3 (data not shown). Since *Pem*-CHH1 contains only a single intron (both *Pem*-CHH2 and *Pem*-CHH3 contain two introns), *Pem*-CHH1 seems to have gone through an independent evolutionary path from its related copies, *Pem*-CHH2 and *Pem*-CHH3.

The conserved 200 nucleotides upstream from +1 of each *Pem*-CHH gene may reflect the presence of common regulatory elements among the three *Pem*-CHH genes. On the other hand, the divergence in the more distal region (upstream from the 200 position) suggests that the nucleotides in this region may not be required for the regulatory function. It is, nevertheless, possible that these sequences may have undergone a divergence in order to acquire a specific regulatory function to each gene. However, no possible recognition sequences for any distinctive regulatory factors have been found within these distal regions.

The presence of a putative site for CREB (cAMP responsive element binding protein), a ubiquitous transcription factor that modulates the gene transcription via the cAMP/PKA signaling pathway (Silva et al., 1998) in the upstream region of *Pem-CHH* genes, suggests the role of cAMP in mediating the expression of these genes. This is supported by previous studies that show that the secretion of CHH in the crayfish *Procambarus clarkii* is modulated by serotonin concentration (Escamilla-Chimal et al., 2002), and that the interaction between serotonin and its receptors can modulate the cAMP level (Tierney, 2001). Pit-1 is one of the major pituitary transcription factors that control the expression of growth hormone and prolactin genes in vertebrates (Bradford et al., 2000). The presence of the sequence with high similarity to the Pit-1 binding site therefore suggests...
involvement of a tissue specific transcription factor in the regulation of the expression of the CHH-like genes. Another significant match is the homologue of the binding site for the activator protein 1 transcription factor (AP1). AP-1 is a heterodimer complex of two basic-region leucine-zipper (B-ZIP) factors, c-Fos and c-Jun (Steinmüller et al., 2001). Ap-1 is involved in the control cell growth, differentiation, and development in *Drosophila* (Ciapponi and Bohmann, 2002). The presence of the binding site for AP-1 in the upstream regulatory region of CHH genes thus implies that the CHH expression may be involved in the growth and development of crustaceans via the regulation of the AP1-like factor. Binding sites for similar transcription factors were also found in other characterized CHH, MIH, and MOIH genes (Gu and Chang, 1998; Gu et al., 2000; Lu et al., 2000). This demonstrates that the genes in the CHH/MIH/GIH family might be under the control of similar mechanisms. Further investigation of these putative transcription factors in *P. monodon* and other crustaceans is thus promising for a better understanding of the transcriptional control of the hormones in this family.

A Southern blot analysis showed that the genome of *P. monodon* contains at least three, one, and two copies of the *Pem-CHH1*, *Pem-CHH2*, and *Pem-CHH3* genes, respectively. In the shrimp *Metapenaeus ensis*, at least eighteen genes encoding two major types of CHH peptides, sixteen encoding MeCHH-A, and two encoding MeCHH-B were described. MeCHH-A and MeCHH-B share more than 98% identity in their amino acid sequences (Gu et al., 2000). These genes were arranged into two different clusters, each cluster contained at least six MeCHH-A genes and one copy of the MeCHH-B gene (Gu et al., 2000). In addition to CHH, other hormones in this family, MIH and MO-IH, of *Cancer pagurus* were also encoded from genes that are clustered within a 6.5 kb region of the genome (Lu et al., 2000). Our results also give preliminary information on the clustering of Pem-CHH genes in the genome of *P. monodon*. A single genomic band above 10 kb that was obtained from the Hind III digestion hybridized to all of the Pem-CHH probes. This suggests that
at least one copy of each of the Pem-CHH1, Pem-CHH2, and Pem-CHH3 genes are clustered within this Hind III-digested fragment. The DNA probe that was generated from the mature peptide-coding region of Pem-CHH1 can detect several extra bands in addition to those detected by each Pem-CHH-specific probe. This probe is 60-80% homologous in its nucleotide sequence to those of the other five CHH-like genes (Pm-sgp-I to -V) of P. monodon (Davey et al., 2000). It is therefore possible that the additional hybridizing bands correspond to the presence of Pm-sgp-I to -V and probably other genes in the CHH family.

In summary, this study showed that the Pem-CHH2 and Pem-CHH3 genes of P. monodon have similar structures to each other as well as to other genes in the same family that have been characterized, but is different from the previously reported structure of the Pem-CHH1 gene. The putative binding sites for regulatory factors that were identified within the 5 upstream sequence of all the Pem-CHH genes suggested that these genes are regulated by similar mechanisms. Moreover, clustering of the three Pem-CHH genes within a distance of at least 10 kb was also demonstrated by a Southern blot analysis. Further studies on the genomic organization of CHH-like genes in P. monodon would provide an implication for their evolution as well as the control of their expression.

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