NOTE

Upregulation of actin-like gene expression in giant freshwater prawns *Macrobrachium rosenbergii* infected with *Metschnikowia bicuspidata*

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ABSTRACT: The giant freshwater prawn *Macrobrachium rosenbergii* is commercially cultured throughout the world including Taiwan. From 1992 to 1995, Taiwanese production decreased by approximately 50% due to disease. The yeast *Metschnikowia bicuspidata* is considered to be one of the major causes of white muscle disease, but the molecular mechanism of its pathogenesis is not known. Using RNA differential display (DD) with muscle and hepatopancreatic tissue, we identified a 324 nucleotide (nt) message specifically expressed by *M. rosenbergii* infected with *M. bicuspidata* but not in the controls. A ribonuclease protection assay (RPA) confirmed expression in both tissues. RPA data also revealed an additional 230 bp mRNA message that was not identified by DD. Using RNA ligase-mediated rapid amplification of 5' cDNA ends (5'-RACE), we successfully isolated a 1357 bp full-length gene (c57) that showed 92 and 87% sequence identity to the actin gene of the Kuruma shrimp *Marsupenaeus japonicus* (also called *Penaeus japonicus*) (GenBank accession number AB055975) and the beta-actin gene of the white shrimp *Litopenaeus vannamei* (also called *Penaeus vannamei*) (GenBank accession number AF300705), respectively. The deduced amino acid sequence of c57 showed 83% sequence similarity to *M. japonicus* and *L. vannamei* actin proteins. Based on this high homology, we suggest that upregulation of actin expression in the muscle and hepatopancreas is part of the shrimp response to *M. bicuspidata* infection. Increased expression may be related to repair of tissues damaged by yeast infection.

KEY WORDS: *Macrobrachium rosenbergii* · *Metschnikowia bicuspidata* · Actin

INTRODUCTION

Commercial culture of the giant freshwater prawn *Macrobrachium rosenbergii* is undertaken throughout the world including Taiwan, where production decreased by 47 to 52% from 1992–1995 due to diseases (New 1995) caused by yeast infection and *Lactococcus* infection (Cheng & Chen 1998, Lu et al. 1998). Yeast infections are associated with gross signs including yellow exoskeletons, a swollen hepatopancreas, milky hemolymph and opaque, whitish muscles (Cheng & Chen 1998, Lu et al. 1998). Yeast disease outbreaks have occurred since 1988 in southern Taiwan, most frequently in the cool season (i.e. October to March) and predominantly (i.e. 64%) during the winter (i.e. December to February) since 1994 (Lu et al. 1998). The rate of infection is the highest in adult prawns (i.e. 73%) and lower in juveniles (25%) and postlarvae...
Yeasts reported to be pathogenic for *Macrobrachium rosenbergii* include *Candida sake* I and II, *Pichia anoma*, *Endomyces fibuliger* and *Candida famata* (Lu et al. 1998). As identification of pathogenic yeast species based on cellular morphology and biochemical tests are often strain-specific (Lu et al. 1998), molecular approaches such as those based on the 26S rDNA gene (Kurtzman & Robnett 1997) are more useful. We used both approaches to identify individual species from co-infections of *Enterococcus faecium* and *Metschnikowia bicuspidata* in *M. rosenbergii* (Chen et al. 2003). However, the host response to *M. bicuspidata* infection and the mechanism of yeast pathogenicity have not been studied.

The technique of ribonucleic acid (RNA) differential display (DD) (Liang & Pardee 1992) was first used to study differences in gene expression during various developmental stages of tumor cells. In the present study, we use the technique to identify a gene that is upregulated in muscle and hepatopancreatic tissues of *Macrobrachium rosenbergii* infected with *Metschnikowia bicuspidata*.

**MATERIALS AND METHODS**

**Diseased prawn samples.** Diseased prawn samples (body weight 5 to 8 g, body length 8 to 10 cm) were collected for molecular analyses from 4 prawn farms (10 prawns were randomly selected from each farm) during disease outbreaks in 2001 (Chen et al. 2003). Infection with *Metschnikowia bicuspidata* was confirmed by sequence analysis of the D1/D2 domain of 26S rDNA and histopathological examination in diseased prawn tissues (Chen et al. 2003).

**Yeast strains.** The purified isolate derived from infected prawns in this study was as described by Chen et al. (2003) and labeled *Metschnikowia bicuspidata* AOD081MB.

**Experimental infections.** Healthy *Macrobrachium rosenbergii* (body weight 5 to 7 g) obtained from 1 prawn farm in Pingtung, Taiwan, were held at a density of 120 prawns in 3 continuously aerated 400 l aquaria containing 300 l of fresh water at 25°C for 7 d, until the prawns were acclimated to laboratory conditions. These prawns were fed twice daily with a commercial diet (Prawn Diet, Gold Brand Feed) and the waste was removed daily. Subsequently, 20 prawns with similar body length ranges were evenly divided into 2 groups: the Treatment Group and the Control Group. Ten-fold serial dilutions of the purified pathogen, isolate AOD081MB (Chen et al. 2003) (optical density [OD] = 1 at 620 nm), were prepared in sterile normal saline (0.85 % NaCl). Each prawn in the Treatment Group was injected intramuscularly (IM) with 0.1 ml of the diluted pathogen (approximately 1 × 10⁷ AOD081MB), whilst the Control Group was injected IM with 0.1 ml of sterile normal saline. After injection, the groups were maintained separately in a 80 l aquaria under the same conditions as described above for acclimation.

**RNA isolation and mRNA differential display.** In this study, the technique of mRNA DD was used to investigate changes in gene expression in muscle and hepatopancreatic tissue 6 h after IM injection in the Treatment and Control Groups. Total RNA was isolated from muscle and hepatopancreatic tissue samples derived from injection control, injection challenge and naturally infected prawns (10 prawns in each group) using 3-Zol reagent (MBio). Pooled RNA from muscle or hepatopancreatic tissue from each group was used for DD analysis. This RNA was treated with DNase I and quantified spectrophotometrically by absorbance at 260 nm. RNA integrity was verified by 1% agarose gel electrophoresis followed by ethidium bromide staining (MBio). Synthesis of the first strand cDNA was accomplished by reverse transcription (RT) using 300 U of Moloney murine leukemia virus reverse transcriptase (MMLV-RT) (Life Technologies) and 2.5 µM of 3′-primer. After heat inactivation of the RT at 95°C for 5 min, 2 µl of the sample was added to 18 µl of polymerase chain reaction (PCR) mix containing a 5′-primer and AmpliTaq DNA polymerase (Perkin Elmer). The PCR mix preparation was based on a non-radioactive procedure described by Doss (1996) except that the amount of AmpliTaq DNA polymerase was doubled. As DNase I-treated RNAs are sometimes contaminated with residual chromosomal DNA (Lieng et al. 1993), a control RT-PCR without reverse transcriptase was included. The RT-PCR products from each sample were run on a 5% denaturing polyacrylamide gel stained with silver (Su et al. 2004). Four 3′-primers (5′-T12AT-3', 5′-T12CA-3', 5′-T12CC-3', 5′-T12CG-3') and four 5′-primers (5′-AGTAGCTGGA-3', 5′-CTC-GGTACAC-3', 5′-ATGTGGTGGT-3', 5′-TCTGCCGT-GA-3') (Su et al. 2004) synthesized by MBio were used to accomplish 16 RT-PCRs. Differentially expressed gene product sizes were determined based on the procedures described by Bassam et al. (1991) using silver-stained gels with pGEM DNA markers obtained from Promega. DNA fragments that were differentially present in the sample groups were recovered from the gel using the methods of Weaver et al. (1994). The recovered DNA was PCR-reamplified and cloned using pCRII vector (Invitrogen) in *Escherichia coli* INVαF competent cells (Invitrogen) and the transformed cells were recovered in SOC medium (Invitrogen). Plasmid DNA was purified from selected trans-
formants by a modified mini alkaline-lysis and polyethylene glycol (PEG) precipitation procedure (Perkin-Elmer) using Terrific Broth (Tartof & Hobbs 1987) for the growth of bacteria and including PEG8000 in the precipitation step. Six independent clones from each original DNA fragment were sequenced with M13 forward and reverse primers using an automated sequencer (MDBio). The complete nucleotide sequence data of differentially expressed cDNA inserts were determined by aligning both strands of sequence data using the DNAMAN Program (Version 2.5, Lynnon BioSoft). Sequences were tested for homology to known sequences at GenBank using the basic local alignment search tool (BLAST) (Altschul et al. 1990).

**Amplification of 5' cDNA ends.** Total RNAs were extracted from muscle and hepatopancreatic tissue of experimentally inoculated and naturally infected prawns using Trizol reagent (Life Technologies). Isolation of the full-length differentially expressed cDNA clone was accomplished by 5'-RACE using a GeneRacer Kit (Version J, Invitrogen) according to the manufacturer’s instructions, except that the sequences of a 3' reverse gene-specific primer (5'-GTG GGG CAA TGA TCT TGA TGT GGG GAG TAG GAC TGG TAA CTG CAG-3') were designed from the differentially expressed cDNA sequence. After performing 5'-RACE, the PCR product was cloned in pCRII vector (Invitrogen). Plasmids from positive colonies were purified and sequenced by MDBio. The resulting sequences were tested for homology to known sequences at GenBank using BLAST (Altschul et al. 1990).

**Ribonuclease protection assay.** A non-radioactive ribonuclease protection assay (RPA) was performed to confirm the expression of c57 cDNA in the muscle and hepatopancreatic tissue of infected prawns using an RPA III kit (Ambion) following the manufacturer’s recommended protocol and the method described by Wundrack & Dooley (1992). The antisense RNA probe was prepared by transcription of an Apal-linearized plasmid template of c57 with Sp6 RNA polymerase using the Sp6/T7 transcription kit (Boehringer Mannheim) in the presence of digoxigenin (DIG)-UTP (Roche Molecular Biochemicals). The 464 nucleotide (nt) full-length hybridization probe contained 324 nt of 3' coding sequence of cloned c57 RNA and 140 nt of vector sequence. This antisense riboprobe was subsequently purified using a 1 ml Sephadex G-50 (Sigma) spin column as previously described (Su et al. 2001). After hybridization with RNA from muscle and hepatopancreatic tissue, the unhybridized single-stranded RNA was digested by RNases A and T1 (provided with the RPA III kit). Protected fragments were separated on a 6% polyacrylamide gel containing urea, transferred to a nylon membrane by electroblotting, and detected using a DIG Nucleic Acid Detection kit (Roche Molecular Biochemicals).

**RESULTS AND DISCUSSION**

**Identification and isolation of c57 from diseased prawns**

Using the primers, 5'-T12AT-3' and 5'-ATGTG-GTGGT-3', 1 up-regulated gene (c57) was identified in muscle and hepatopancreatic tissue of experimentally (Fig. 1, lanes 2 and 3 and 4 and 5, respectively) and naturally infected prawns (Fig. 1, lanes 8 and 9 and lane 10, respectively), but not in the tissues of healthy prawns (Fig. 1, lanes 6 and 7, respectively). As repeated experiments showed a similar display pattern (data not shown), c57 was selected for further work. Using the other 15 primers, 3 additional fragments were found in Metschnikowia bicuspidata-infected prawns but not consistently in repeated experiments.
Thus, they were excluded from further investigation. RNA sample authenticity was confirmed by the absence of cDNA amplification using pooled hepatopancreatic RNA from experimentally infected prawns in the absence of reverse transcription (Fig. 1, lane 1). Thus, cDNA products were fingerprints of tissue-derived mRNAs.

Nucleotide and amino-acid sequences of c57

The c57 fragment from a display gel was eluted, PCR-reamplified, cloned and sequenced from both strands. Sequence alignment revealed 100% complementarity between the 2 strands (data not shown). The sequence (nt 1027 to 1348) is shown in Fig. 2A with the flanking primer set shown in bold italics. There is 1 nucleotide mismatch at the penultimate base (lowercase g, Fig. 2A). This mismatch between the primer and c57 suggested that the primer may exhibit degeneracy during annealing in the RT step (Liang et al. 1993). The full-length c57 sequence (Fig. 2A) obtained by 5'-RACE had 92% identity to the nucleotide sequence of the actin gene of the Kuruma shrimp Marsupenaeus japonicus (also called Penaeus japonicus) (GenBank accession number AB055975) and 87% identity to the beta-actin gene of white shrimp Litopenaeus vannamei (also called Penaeus vannamei) (GenBank accession number AF300705). The deduced amino acid sequence of c57 shared 83% sequence similarity with residues of 78–381 of the M. japonicus and L. vannamei actin proteins (Fig. 2B).

![Fig. 2. Nucleotide sequence (GenBank accession number AY651918) and deduced amino acid sequence of c57 from Macrobrachium rosenbergii infected with Metschnikowia bicuspidata. (A) Start and stop codons are shown in bold letters. Nucleotide sequences of flanking mRNA mapping primers are shown in bold italics. Nucleotides 320 to 1070 showed 92% identity to the Kuruma shrimp Marsupenaeus japonicus actin gene and 87% identity to the white shrimp Litopenaeus vannamei beta-actin gene. : stop codon. (B) Data from alignment of the deduced protein sequence of c57 showing 83% similarity (highlighted regions) to the M. japonicus actin and Penaeus vannamei beta-actin protein sequences. PRO: protein sequence to each corresponding DNA](image)
Confirmation of c57 mRNA expression in the hepatopancreas and muscle

Using nonradioactive RPA to verify c57 mRNA expression in the hepatopancreas and muscle of infected prawns (Fig. 3), it was found that hybridization of a DIG-labeled c57 antisense RNA probe to RNA from infected hepatopancreatic and muscle tissue resulted in the formation of a 324 bp RNA-RNA hybrid that was protected from subsequent RNase digestion (Fig. 3, lanes 2 to 5). Expression of c57 detected by RPA was similar to that for the DD (Fig. 1). Visible changes such as whitish muscles and massive lesions have been found in Macrobrachium rosenbergii after Metschnikowia bicuspidata infection (Lu et al. 1997, 1998). It is possible that c57 was upregulated in response to yeast infection as a result of involvement in repair of tissues damaged by the yeast. The RPA data revealed another 230 bp message (lanes 2 to 5) that was not identified by the DD. Lack of expression for this message in DD may be due to poor optimization of conditions for the separation of DNA within the size range of 230 bp (i.e. DNA smearing seen on lanes 2 to 5 and 7 to 10, Fig. 1). As this 230 bp fragment was expressed together with c57 in the RPA, it is possible that both genes respond in a coordinated manner to M. bicuspidata infection. Further characterization of this DD fragment using different conditions would be required. RPA of hepatopancreatic tissue from healthy prawns failed to reveal any signal (lane 1). Taken together, our data demonstrate that ontogenic expression of c57 in the muscle and hepatopancreas of giant freshwater prawns M. rosenbergii was a consequence of the M. bicuspidata infection.

Ubiquitous expression of actin has been found in mammals, Drosophila and crustaceans and the nucleotide sequences of their actin genes show high homology (Kang & Naya 1993). In crustaceans, actin is believed to play an important role in developing cysts during the embryonic stage. Evidence has shown that at nauplius larva stage of the brine shrimp Artemia salina embryos, the proportion of actin mRNA and protein is increased 3-fold above levels found in the dormant cyst (Grosfeld & Littauer 1976). It was suggested that at the nauplius stage, actin may participate with myosin and other proteins to form muscles and this correlates with the acquisition of motility during this free-swimming larval stage (Grosfeld & Littauer 1976).

In addition to tissue repair, it is possible that expression of c57 may have some functional role in protecting Macrobrachium rosenbergii against external agents like the yeast Metschnikowia bicuspidata. The issue might be examined by employing in situ hybridization analysis to identify the cellular location of c57 expression. There is also evidence that expression of some immune-related proteins can be stimulated by yeast infections in shrimp. For example, Sequeira et al. (1996) reported that proliferating haemocytes were significantly increased in Marsupenaeus japonicus after stimulation by p43, an immunosuppressive lymphocyte mitogenic protein produced by Candida albicans. A similar response resulted from infection with Fusarium spp. (Sequeira et al. 1996). It is also known that betaglucan from yeast can initiate the shrimp prophenoloxidase (ProPO) defense cascade (Sritunyalucksana & Soderhall 2000). However, no role for actin has yet been identified in the shrimp immune response.

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