Molecular cloning and characterization of an inhibitor of apoptosis protein (IAP) from the tiger shrimp, *Penaeus monodon*☆

Jiann-Horng Leu, Yu-Chen Kuo, Guang-Hsiung Kou*, Chu-Fang Lo*

**Institute of Zoology, National Taiwan University, Taipei 10617, Taiwan, ROC**

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Abstract

The inhibitor of apoptosis proteins (IAPs) play important roles in both apoptosis and innate immunity. Here, we report the first cloning and characterization of a novel IAP family member, PmIAP, from *Penaeus monodon*. The full-length *PmIAP* cDNA is 4769 bp, with an ORF encoding a protein of 698 amino acids. The PmIAP protein contains three BIR domains and a C-terminal RING domain, and its mRNA was expressed in all analyzed tissues. In insect cells, PmIAP, together with *Spodoptera frugiperda* IAP, AcMNPV P35, and WSSV449 (or ORF390, an anti-apoptosis protein encoded by white spot syndrome virus), could all block the apoptosis induced by *Drosophila* Reaper protein (Rpr), whereas only P35 and WSSV449 could block the apoptosis induced by actinomycin D. Co-immunoprecipitation showed that PmIAP physically interacted with Rpr, and in an immunofluorescent analysis the two proteins produced co-localized punctate signals in the cytoplasm. Deletion analysis revealed that both the BIR2 and BIR3 domains of PmIAP could independently bind to and inhibit Rpr, whereas the BIR1 domain could not. These results strongly suggest that PmIAP blocks Rpr’s pro-apoptotic activity through mechanisms that are evolutionarily conserved across crustaceans, insects, and mammals.

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1. Introduction

Apoptosis is a genetically programmed cellular suicide process that eliminates unwanted or diseased cells, and it plays important roles in embryogenesis, homeostasis, insect metamorphosis, and immunity [1–3]. Cells undergoing apoptosis display a series of morphological changes, including cell shrinkage, chromatin condensation, DNA fragmentation, membrane blebbing, and, finally, formation of...
apoptotic bodies [4]. The concept of apoptosis was first introduced in 1972 [5]. Since then, hundreds of genes that control apoptosis have been identified in different species, which is good evidence that the mechanisms and the molecules involved in this suicide program have been highly conserved throughout metazoan evolution [6].

Inhibitor of apoptosis proteins (IAPs) are a conserved group of proteins that regulate apoptosis in both vertebrates and invertebrates. Since the first discovery of an IAP gene in a baculovirus [7], numerous cellular IAP homologs have been identified [8,9]. The hallmark of an IAP protein is the presence of one to three copies of a zinc-binding baculoviral IAP repeat (BIR) domain. These BIR domains bind directly to the caspases to inhibit them, and they are essential for the anti-apoptotic properties of the IAPs [10,11]. The interaction between BIR domains and caspases is negatively regulated by proteins that contain an IAP-binding motif (IBM), including Drosophila Reaper (Rpr), Grim, and Hid, and mammalian Smac/DIABLO. The only sequence homology shared by these IBM proteins is a short consensus sequence at their amino termini. This conserved sequence is the IBM, which is responsible for binding to the BIR domain and interfering with IAP-mediated caspase inhibition [12]. The IAPs also contain another zinc-binding domain known as the RING domain, which is invariably located at their C-termini. The RING domain has an ubiquitin E3 ligase activity, which attaches ubiquitin to proteins that bind to the IAPs, including caspases, some IBM proteins, and IAP itself. The ubiquitinated IAP and IBM proteins are then subject to proteasomal degradation, while the ubiquitinated caspases, instead of being degraded, become inactive [12–14].

In addition to acting as important regulators in apoptosis, recent evidence suggests that some IAPs also play important roles in innate immunity in both mouse and Drosophila [15–19]. In Drosophila, microbe infection initiates innate immune responses through two major, distinct signaling pathways: the Toll pathway and the immune deficiency (Imd) pathway. The Toll pathway is predominantly activated by fungal and Gram-positive bacterial infections, whereas the Imd pathway responds to infection by Gram-negative bacteria. The activation of these two pathways causes members of the nuclear factor kB (NF-kB) family to translocate to the nucleus to activate the transcription of antimicrobial peptide genes [20]. The importance of Drosophila IAP2 (DIAP2) in Drosophila innate immunity was first revealed by two independent studies using large-scale RNAi screening in cultured Drosophila cells to identify novel components in the Imd pathway. Both studies showed that diap2 RNAi abrogated the Imd-signaling antimicrobial peptide response [16,17]. Moreover, diap2 null mutant Drosophila fail to induce the synthesis of antimicrobial peptides and are highly susceptible to infection by Gram-negative bacteria [18,19].

In the present paper, based on annotations of a 5′EST database generated from a Penaeus monodon post-larvae cDNA library, we identified a cDNA clone that appeared to encode an IAP protein. Since no crustacean IAP has been reported, we therefore decided to clone the full-length cDNA sequence of this P. monodon IAP (PmIAP) and to characterize this protein. In a series of functional assays, other anti-apoptosis proteins were also included for comparison. One of these is a novel, recently reported anti-apoptosis protein (WSSV449, or ORF390) encoded by a shrimp viral pathogen, white spot syndrome virus (WSSV).

2. Materials and methods

2.1. Identification of a cDNA clone of a P. monodon inhibitor of apoptosis protein (PmIAP)

A cDNA library constructed from WSSV-infected postlarvae of P. monodon was subjected to large-scale 5′ and 3′ end sequencing, and the sequencing results were used to generate a 5′ and 3′ EST database. Analysis of the 5′ EST database using BLASTX against the NCBI non-redundant (nr) database revealed that one cDNA clone, PmTwl09F1, contained a partial nucleotide sequence that was highly homologous to the RING domain of the Bombyx mori inhibitor of apoptosis protein. Complete sequencing of this clone by primer walking showed that it contained only part of the nucleotide sequence of a putative P. monodon IAP gene.

2.2. Isolating the 5′ end of PmIAP cDNA

Pleopod (swimming leg) mRNA was purified using a Quick-Prep Micro mRNA Purification Kit according to the manual supplied by the manufacturer (GE Healthcare). Briefly, pleopods from adult shrimp were powdered in the presence of liquid nitrogen, and further homogenized in the extraction buffer supplied with the kit. The homogenate was diluted with elution buffer, clarified by centrifugation at 12,000g for 5 min, and then mixed with oligo(dT)-cellulose. After extensive washing with high- and low-salt buffers, the mRNA was eluted with elution buffer, precipitated with ethanol, and then quantified at A260. A 5′ RACE kit was then used to isolate the 5′ end of PmIAP cDNA according to the instructions provided by the manufacturer (Roche Molecular Biochemicals). Three gene-specific primers, I09F1-SP1, I09F1-SP2, and I09F1-SP3 (Table 1), were designed according to the nucleotide sequence of the PmTwl09F1 cDNA clone. The first-strand cDNA was synthesized from 100 ng pleopod mRNA using I09F1-SP1 primer, tailed at the 3′ end with dATPs using the terminal transferase, and then subjected to first-round PCR using oligo(dT)-anchor primer and I09F1-SP2 primer. The PCR product was diluted and subjected to second-round PCR using anchor primer and I09F1-SP3 primer. The final PCR products of 5′ RACE were cloned into the pGEM-T easy vector (Promega) and used to obtain the complete cDNA sequence of PmIAP.

2.3. Sequence analysis of PmIAP

The deduced amino acid sequence of PmIAP was analyzed with BLASTP against the NCBI nr database and ScanProsite. For multiple sequence alignment, the BIR1, 2, and 3 domains and RING domain from PmIAP were aligned using GeneDoc (ver. 2.6.002) with the corresponding domains from SfIAP (AX213188), DIAP1 (Q24306), DIAP2 (Q24307), and XIAP (P98170).
2.4. Tissue distribution of PmIAP mRNA

Total RNAs were extracted from various tissues using TRIzol reagent (Invitrogen, USA) according to the procedure provided by the supplier. After quantification at A260, about 10 μg of total RNAs were treated with DNase I, extracted with phenol–chloroform, and then precipitated with ethanol. The DNase I-treated total RNAs were primed with oligo-dT-anchor primer (Roche) and reverse-transcribed with SuperScript II (Invitrogen, USA) at 42 °C for 50 min. Aliquots of this cDNA were then used for PCR analysis. The PmIAP primer pair used for RT-PCR was 109F01R1 and 109F01F1 (Table 1). For internal control, an actin gene primer pair, P1882-actin F1 and P1883-actin R1 (Table 1), was designed based on P. monodon actin 2 (AF100987).

2.5. Plasmid construction

The pilot study showed that the insect cell expression vector, pIZ/V5-His (Invitrogen), was apparently unable to express any of the desired proteins in sufficient quantities (data not shown). We followed Clem and Miller [22] and designed expression constructs based on the promoter of the Drosophila heat shock protein 70 (hsp70) gene. The designed expression constructs based on the promoter and then ligated with the BspHI and HindIII-restricted hsp70 DNA fragment. The resulting plasmid was constructed by digesting the hsp70 promoter region (from −525 to +236, according to Torok and Karch [23], and GenBank accession no. J01103) was PCR amplified from D. melanogaster genomic DNA using forward primer 5'-CCTCATGAGTTGACAACAACAGTCTTGACAACCT-3' and reverse primer 5'-GGGAAGCTTCCTCGGTAACGACTTGTTGAAAGT-3' (underlined nucleotides correspond to cutting sites for BspHI and HindIII, respectively), and the PCR product was cloned into pGEM-T easy vector. To construct the heat-inducible expression vector, the plIZ/V5-His plasmid (Invitrogen) was digested with BspHI and HindIII to release the OPIE2 promoter and then ligated with the BspHI and HindIII-digested hsp70 DNA fragment. The resulting plasmid was named pDHsp/V5-His. A second heat-inducible expression vector, pDHsp/FLAG-His, was constructed by digesting the pDHsp/V5-His with SacI and MluI to release the V5 tag and ligated with the FLAG tag fragment (5'-GGGACTACAAAGACTTCTGAGTAAGAGCGACTCACAAG-3' and 5'-CCGCCTGACGTAGTCTCTTTAGGTCAAGTGTAG-3').

The genes analyzed in this study include Drosophila rpr (NM_079414), PmlIAP (EF114675), SfiAP (AX213188), AcMNPV p35 (P08160), and wssv449 (NP_477912). These genes were inserted into pDHsp/V5-His or pDHsp/FLAG-His by PCR cloning. The primers used for PCR and the corresponding constructs are listed in Table 2.

Various PmIAP deletion constructs were constructed by PCR using the plasmid pDHsp-PmIAP/V5 as template. The amplified fragments were cloned into pDHsp-His/V5 to generate the corresponding expression constructs, all of which produced a V5-tagged fusion protein. The constructs and the used primers are listed in Table 3.

2.6. Cell culture

Spodoptera frugiperda (Lepidoptera: Noctuidae) SF-9 insect cells were maintained at 27 °C in SF-900 II SFM medium (Invitrogen) supplemented with 10% heat-inactivated fetal bovine serum (Hyclone Laboratories).

2.7. Actinomycin D assay

SF-9 cells seeded in six-well plates (1 × 10^6 cells/well) were transfected with 4 μg of the constructed plasmids by Cellfectin reagent (Invitrogen) according to the procedure described by the supplier. At 18–20 h post-transfection, the cells were heat shocked in a 42 °C water bath for 30 min to induce the expression of the transfected genes placed under the hsp70 promoter, and then returned to 27 °C. At 4 h after heat shock, the media were removed and replaced with fresh media containing 150 ng/ml of actinomycin D (ActD). At 14–16 h after addition of ActD, the media was removed and the cells were washed with phosphate-buffered saline (PBS; 137 mM NaCl, 10 mM phosphate, 2.7 mM KCl, pH 7.4) and then stained with 0.04% trypan blue in PBS. The viable, intact cells were counted using a bright-field microscope equipped with a digital camera and AutoCell software. The total number of viable cells was determined from five evenly distributed fields of view at 200× magnification. The viability of the cells used to express the indicated proteins was calculated by dividing the total number of viable cells remaining after drug treatment by the total number of viable cells in control transfections at the time of drug addition. Results were expressed as mean ± S.D.; n = 3.

2.8. Viability assays for Rpr-induced apoptosis

SF-9 cells seeded in six-well plates (5 × 10^5 cells/well) were transfected with the appropriate plasmids using Cellfectin reagent (Invitrogen). After transfection for 16–18 h, the cells were heat shocked in a 42 °C water bath for 30 min and

<table>
<thead>
<tr>
<th>Primer name</th>
<th>Primer sequence</th>
<th>Usage</th>
</tr>
</thead>
<tbody>
<tr>
<td>I09F1-SP1</td>
<td>5'-TCACACTTCAGGATAACATAGGT-3'</td>
<td>IAP 5’ RACE</td>
</tr>
<tr>
<td>I09F1-SP2</td>
<td>5'-CTGAGTAGGACGGACTCAAAG-3'</td>
<td>IAP 5’ RACE</td>
</tr>
<tr>
<td>I09F1-SP3</td>
<td>5'-ACGAGCTCAGTTCTGAGCATCA-3'</td>
<td>IAP 5’ RACE</td>
</tr>
<tr>
<td>109F01R1</td>
<td>5'-AGAAGAATACGGCAAGTAGTGCT-3'</td>
<td>IAP RT-PCR</td>
</tr>
<tr>
<td>109F01F1</td>
<td>5'-GCCCTCTTATGGAAGTAGT-3'</td>
<td>IAP RT-PCR</td>
</tr>
<tr>
<td>P1882-actin F1</td>
<td>5'-CGCTCATCGGGTGATGCGT-3'</td>
<td>Actin RT-PCR</td>
</tr>
<tr>
<td>P1883-actin R1</td>
<td>5'-CCACGCTACGTGATCTTCA-3'</td>
<td>Actin RT-PCR</td>
</tr>
</tbody>
</table>
Proteins were separated by Tricine-SDS/12.5% polyacrylamide gel electrophoresis (PAGE) and transferred to Immobilon-P membranes (Millipore). V5-tagged fusion proteins were detected with rabbit anti-V5 antibody (1:10,000; Sigma) and goat anti-rabbit IgG–horseradish peroxidase (HRP) conjugate (1:10,000; Sigma). Rpr-FLAG protein was detected with mouse anti-FLAG antibody (1:5,000; Sigma) and goat anti-mouse IgG–HRP conjugate (1:10,000; Sigma).

2.10. Co-immunoprecipitation

SF-9 cells were transfected and subjected to heat shock as described above. At 1 h post-heat shock, the cells were washed with PBS and lysed in 100 μl of NP-40 lysis buffer (50 mM Tris-Cl, pH 8.0, 150 mM NaCl, 1.0% Nonidet P-40) supplemented with a protease inhibitor tablet (Roche). The lysis procedure was carried out on ice for 10 min with occasional shaking. The lysate was centrifuged at 12,000 g for 5 min and an aliquot of the supernatant (10 μl) was reserved for immunoblot analysis of the total cell lysate to confirm the expression of the transfected genes. An adequate amount of supernatant was then incubated with 10 μl of anti-FLAG M2 affinity gel (Sigma) or incubated with 1 μl of rabbit anti-V5 antibody (Sigma) and 20 μl of protein A

Table 2 Primer sequence list for the construction of various expression plasmids

<table>
<thead>
<tr>
<th>Gene</th>
<th>Primer sequence</th>
<th>Cutting site</th>
<th>Vector</th>
<th>Tag</th>
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<td>Rpr</td>
<td>F: 5’-GGAACGGTACCATGGCAGTCTCTACATACCCGA-3’</td>
<td>HindIII</td>
<td>pDHsp/V5-His</td>
<td>No</td>
</tr>
<tr>
<td></td>
<td>R: 5’-TCGGATCTCATTGGGATGGTCTGCGA-3’</td>
<td>BamHI</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>R: 5’-GGAACGGTACCATGGCAGTCTCTACATACCCGA-3’</td>
<td>HindIII</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>R: 5’-CGAACCGCGGGGCTGGAGTCTGCGATATTG-3’</td>
<td>Sacl</td>
<td></td>
<td></td>
</tr>
<tr>
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<td>F: 5’-GAAGCATCAGGTGGTATAATGCCAGAACG-3’</td>
<td>HindIII</td>
<td>pDHsp/FLAG-His</td>
<td>Flag</td>
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<td>NotI</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SfiAP</td>
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<td>pDHsp/V5-His</td>
<td>V5</td>
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<tr>
<td></td>
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<td>AcMNPV p35</td>
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<td>HindIII</td>
<td>pDHsp/V5-His</td>
<td>V5</td>
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<tr>
<td>wssv449</td>
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<td>HindIII</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>R: 5’-GCCACCTACTACTGAGCAGTTGCGTGG-3’</td>
<td>BamHI</td>
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<td></td>
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</table>

Table 3 Primer sequence list for the preparation of various PmlAP deletion constructs

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<th>Primer sequence</th>
<th>Cutting site</th>
<th>Region (aa)</th>
</tr>
</thead>
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<td>BIR</td>
<td>F: 5’-CTGCAACTAGTGAATAACAAAG-3’</td>
<td>NotI</td>
<td>1–358</td>
</tr>
<tr>
<td></td>
<td>R: 5’-AGGGGCGCGCGCTAGCTGCTGATCTTGGG-3’</td>
<td>NotI</td>
<td></td>
</tr>
<tr>
<td>RING</td>
<td>F: 5’-GAAGCTTACATGAGTACAGCAGAATGCGAAG-3’</td>
<td>HindIII</td>
<td>351–698</td>
</tr>
<tr>
<td></td>
<td>R: 5’-AGGGGCGCGCGCTAGCTGCTGATCTTGGG-3’</td>
<td>NotI</td>
<td></td>
</tr>
<tr>
<td>BIR1</td>
<td>F: 5’-CTGCAACTAGTGAATAACAAAG-3’</td>
<td>NotI</td>
<td>1–92</td>
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<tr>
<td></td>
<td>R: 5’-GGTCCCGCGCGCTAGCTGCTGATCTTGGG-3’</td>
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</tr>
<tr>
<td>BIR2</td>
<td>F: 5’-GGTCCCGCGCGCTAGCTGCTGATCTTGGG-3’</td>
<td>HindIII</td>
<td>86–227</td>
</tr>
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<td></td>
<td>R: 5’-GGTCCCGCGCGCTAGCTGCTGATCTTGGG-3’</td>
<td>NotI</td>
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</tr>
<tr>
<td>BIR3</td>
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<td>HindIII</td>
<td>233–358</td>
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<td></td>
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<tr>
<td>BIR1+2</td>
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<td>NotI</td>
<td>1–193</td>
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<td></td>
<td>R: 5’-GGTCCCGCGCGCTAGCTGCTGATCTTGGG-3’</td>
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<td></td>
<td>R: 5’-GGTCCCGCGCGCTAGCTGCTGATCTTGGG-3’</td>
<td>NotI</td>
<td></td>
</tr>
</tbody>
</table>

then returned to 27 °C. At 5 h after heat shock, the viability of the SF-9 cells was determined according to Clem and Miller [22]. Briefly, the media were removed, and the cells were resuspended in 1 ml PBS containing 0.04% trypan blue with a rubber policeman. The viable intact cells were counted with a hemocytometer, and eight grids were counted for each sample. The mean and standard deviation were calculated from three replicate samples. Relative percent viability was calculated by comparing the number of viable cells transfected with various expression plasmids with the number of viable cells transfected with the control plasmid. The relative percent viability for cells transfected with the control plasmid was set at 100%.

2.9. Immunoblot analysis

SF-9 cells seeded in six-well plates (1 × 10⁶ cells/well) were transfected with the appropriate plasmids using Cellfectin reagent (Invitrogen). After transfection for 16–18 h, the cells were heat shocked in a 42 °C water bath for 30 min and then returned to 27 °C. At 1 or 3 h post-heat shock, the cells were washed with PBS and lysed with SDS sample buffer. Proteins were separated by Tricine-SDS/12.5% polyacrylamide gel electrophoresis (PAGE) and transferred to Immobilon-P membranes (Millipore). V5-tagged fusion proteins were detected with rabbit anti-V5 antibody (1:10,000; Sigma) and goat anti-rabbit IgG–horseradish peroxidase (HRP) conjugate (1:10,000; Sigma). Rpr-FLAG protein was detected with mouse anti-FLAG antibody (1:5,000; Sigma) and goat anti-mouse IgG–HRP conjugate (1:10,000; Sigma).
agarose (Sigma) at 4 °C overnight with rotation. The gel or agarose was then washed five times in 150 μl of NP-40 lysis buffer. Aliquots of the total cell lysates and immunoprecipitates were separated by Tricine-SDS/12.5% PAGE and transferred to Immobilon-P membranes (Millipore). V5-tagged fusion proteins were detected with rabbit anti-V5 antibody (Sigma) and goat anti-rabbit IgG–HRP conjugate (Sigma). Rpr-FLAG protein was detected with mouse M2 anti-FLAG monoclonal antibody (Sigma) and goat anti-mouse IgG–HRP conjugate (Sigma).

2.11. In vitro binding assay

To prepare the cell lysates containing Rpr-FLAG, Rpr-FLAG and P35 expression constructs were cotransfected into SF-9 cells. To prepare the lysates containing the SfIAP or various truncated PmIAP proteins, the corresponding expression constructs were transfected into SF-9 cells. At 1 h post-heat shock, the cell lysates were prepared as described above. The lysates containing Rpr-FLAG were mixed with the lysates containing the appropriate proteins, and incubated with anti-FLAG M2 affinity gel at 4 °C overnight with rotation. The gels were then washed five times with lysis buffer, and the immunoprecipitates were analyzed by immunoblotting as described above.

2.12. Immunofluorescence microscopy

SF-9 cells seeded on cover glasses (10^5 cells per well in 4-well plates) were transfected with appropriate plasmids using Cellfectin reagent (Invitrogen) and heat shocked as described above. At 1 h after heat shock, cells were washed twice with PBS, fixed with 4% paraformaldehyde in PBS for 10 min at 4 °C, treated with 0.1% Triton X-100 in 4% paraformaldehyde/PBS for 3 min at 4 °C, and then washed twice with PBS. After blocking with PBS containing 5% bovine serum albumin and 2% normal goat serum for 16 h at 4 °C, the cells were treated with a polyclonal rabbit anti-V5 antibody (1:3000 in PBS; Sigma) or with a monoclonal mouse anti-FLAG M2 antibody (1:1000 in PBS; Sigma) or with both (3 h at room temperature). The cells were then washed 3 times (10 min each time) with PBST (0.2% Tween 20 in PBS) and reacted either with carboxymethylindocyanine (Cy3) dye-conjugated goat anti-rabbit IgG antibody (1:1000 in PBS; Sigma) or with fluorescein isothiocyanate (FITC)-conjugated goat anti-mouse IgG antibody (1:300 in PBS; Sigma), or else reacted with both for 2 h at room temperature. Counterstaining of the nucleus was performed with 4′,6-diamidino-2-phenylindole dihydrochloride (DAPI) (Vector Laboratories). After washing three times (10 min each time) with PBST, the cover glasses were wet mounted with anti-fade mounting media (Fluka). During all of the above procedures, the cells were kept in darkness. Fluorescence signals were examined using a fluorescence microscope (Olympus IX70).

2.13. Statistical methods

All quantitative data are presented as mean ± S.D. (n = 3). Significant differences between pairs of groups were determined using Student’s two-tailed t-test. A P-value < 0.05 was considered statistically significant.

3. Results

3.1. Isolation and sequence analysis of the PmIAP cDNA

Complete sequencing of the PmTw109F1 cDNA clone revealed a cDNA fragment of 2222 bp that encoded the last 60 amino acids of PmIAP, while most (2039 bp) of the rest of the sequence corresponded to the 3′ untranslated region (UTR). After 5′ RACE, the full-length 4769 bp transcript of PmIAP was obtained and its complete sequence was deposited in GenBank under accession no. EF114675. The full-length PmIAP transcript contains a 5′ UTR of 633 bp, an open reading frame (ORF) of 2097 bp, and a 3′ UTR of 2039 bp. The 3′ UTR contains a consensus polyadenylation signal (AATAAA) 11 bp upstream from the poly (A) tail. The ORF encodes a protein consisting of 698 amino acid residues with a calculated molecular mass of 76.8 kDa. The AUG start codon of this ORF is located within a favorable context for translation [24] and it is preceded by upstream stop codons in all three reading frames. ScanProsite analysis revealed that the predicted PmIAP protein contains three BIR domains, followed by a RING domain at its C terminus. This is structurally similar to Drosophila IAP2 (DIAP2) as well as to human X-chromosome-linked IAP (XIAP), cIAP1 (cellular inhibitor of apoptosis-1), and cIAP2 (Fig. 1).

The deduced amino acid sequences of the PmIAP BIR and RING domains were aligned with the corresponding domains of other IAP family members, including Spodoptera frugiperda IAP (SfIAP), Drosophila IAP1 and IAP2 (DIAP1 and DIAP2), and human XIAP (Fig. 2). The BIR2 and BIR 3 domains of PmIAP both have the conserved BIR motif of CX2CXCX6WX5DX4HXC (8). However, in the BIR1 domain, the conserved W residue has been replaced with K59 (see the asterisk in Fig. 2) and two extra amino acids have been inserted in front of the conserved D65 residue. The alignments showed that BIR1 is the least conserved of the three BIR domains, while BIR3 and RING are the most conserved.

3.2. Tissue distribution of PmIAP mRNA

To investigate the tissue distribution of PmIAP transcripts, total RNAs were extracted from various P. monodon tissues and subjected to RT-PCR analysis. Fig. 3 shows that the PmIAP transcript was expressed in all the analyzed tissues, including epithelium, gill, heart, hepatopancreas, hematocyte, intestine, lymphoid organ, nerve, and stomach.

3.3. Neither PmIAP nor SfIAP was able to block actinomycin D-induced apoptosis in SF-9 cells

In this study, to characterize the anti-apoptotic activities of PmIAP, three other anti-apoptosis proteins, SfIAP, P35, and WSSV449, were included for comparison. ActD induces apoptosis in SF cells [7], but this apoptotic response can be blocked by P35, as well as by some baculoviral and insect
**Fig. 1** Schematic representation of the structural motifs of PmIAP and several other IAP proteins. The approximate positions of the BIR, RING, CARD, and UBC domains are shown, and the total amino acid length is indicated. The possession of a RING domain in an IAP confers E3 ubiquitin protein ligase activity, while the possession of a ubiquitin-conjugating domain (UBC) is predicted to confer E2 activity. CARD refers to caspase recruitment domain and its function in cIAP1 and cIAP2 remains unknown. This figure is modified from Hay [43].

**Fig. 2** Multiple sequence alignments of the BIR1, BIR2, BIR3, and RING domains of PmIAP with the corresponding domains of other IAP family members. Identical residues are indicated in black. The amino acid sequence identities and similarities in each aligned domain are also indicated. The asterisk in the PmIAP BIR1 domain indicates the K59 residue that replaces the invariant W. GenBank accession numbers for SfIAP, DIAP1, and XIAP are AX213188, Q24306, Q24307, and P98170, respectively.
IAPs [22,25,26]. Here, we determined whether PmIAP could also block ActD-induced apoptosis in SF-9 cells. First we confirmed that all constructs successfully expressed the indicated V5-tagged proteins (Fig. 4A). Expression levels were similar, except for SfIAP, which was slightly lower (Fig. 4A, lane 3). For the P35, in addition to the major protein band, a number of higher molecular weight products were also detected (Fig. 4A, lane 4). We interpret these bands to be protein complexes formed from P35 and some cellular proteins, because similar P35-associated high molecular weight complexes were also reported when P35 was transiently expressed in mammalian cells [27]. We also note that the apparent molecular mass of PmIAP is about 110 kDa (Fig. 4A, lane 2), which is larger than its calculated molecular mass of about 77 kDa. Fig. 4B shows that P35 and, to a lesser extent, WSSV449 could significantly prevent the cells from undergoing ActD-induced apoptosis (both $P < 0.05$), while neither PmIAP nor SfIAP demonstrated any anti-apoptotic effect.

### 3.4. PmIAP inhibits Rpr-induced apoptosis in SF-9 cells

In SF-9 cells, both baculoviral and Drosophila IAPs have been shown to block the apoptosis induced by Rpr [28,29]. Therefore, in the present study, we investigated the ability of PmIAP to inhibit Rpr-induced apoptosis in SF-9 cells. Fig. 4C shows that PmIAP, SfIAP, P35, and WSSV449 all significantly blocked Rpr-induced apoptosis (all $P < 0.05$). However, while the blocking ability of SfIAP and P35 was quite strong, PmIAP was the least effective of these four proteins.

### 3.5. PmIAP interacts with Rpr and delays its degradation in insect cells

The physical interactions of baculoviral and Drosophila IAPs with Rpr not only inhibited its apoptotic activity but also prevented its degradation [28]. To investigate whether PmIAP and SfIAP could also interact with Rpr and prevent its degradation, we generated a construct that expressed an Rpr protein fused with a C-terminal FLAG-His6 tag (Rpr-FLAG). A time course study was then conducted to investigate the influence of these anti-apoptosis proteins on the stability of Rpr-FLAG. The results showed that, at 1 h post-heat shock, Rpr-FLAG proteins were detected in cells
that coexpressed PmIAP, SfIAP, P35, or WSSV449 (Fig. 5A, upper panel, lanes 3–6), while in cells expressing Rpr-FLAG alone the Rpr-FLAG protein level was much lower (Fig. 5A, upper panel, lane 2). At 3 h post-heat shock, Rpr-FLAG was detected only in cells that coexpressed PmIAP and SfIAP (Fig. 5A, lower panel, lanes 3 and 4). Fig. 5B shows the expression levels of the four anti-apoptosis proteins at 1 and 3 h post-heat shock. The amounts of PmIAP, P35, and WSSV449 increased during this time interval (Fig. 5B, lanes 3–6), while SfIAP appeared to be quite unstable in Rpr-FLAG-coexpressing cells. In the absence of Rpr-FLAG, SfIAP protein levels increased over time (Fig. 5B, lane 7), whereas in the Rpr-FLAG-coexpressing cells, at 3 h post-heat shock, most of the expressed SfIAP protein was truncated (Fig. 5B). We hypothesize that the truncated SfIAP was generated through cleavage of the full-length SfIAP by the activated caspase in the Rpr-induced apoptotic cells. This explanation is based on the fact that Drosophila DIAP1, which is structurally similar to SfIAP (Fig. 1), is degraded by caspases in apoptotic cells in response to Rpr expression [30,31], and we note that the predicted caspase-cleavage sites in SfIAP correspond to those that have been identified in DIAP1.

Next, we determined whether antibody to the Rpr-FLAG protein could co-immunoprecipitate these anti-apoptosis proteins. The result (Fig. 6A) showed that only V5-tagged PmIAP could specifically co-precipitate the FLAG-tagged Rpr (data not shown). In both of these experiments, the apparent inability of SfIAP to co-precipitate with Rpr-FLAG was quite surprising. Possible explanations might be the relatively low expression levels of SfIAP (compare with PmIAP, P35, and WSSV449 in Figs. 4A and 5B) and the instability of SfIAP in the presence of Rpr (Fig. 5B, lanes 4 and 7), so that although the Rpr-FLAG protein was efficiently precipitated by the anti-FLAG antibody, the amount of co-precipitated SfIAP may have been too low to be detected. To resolve this, an in vitro binding assay was conducted. The lysates from cells coexpressing Rpr-FLAG and P35 were mixed with the lysates from cells that expressed SfIAP, and then incubated with anti-FLAG antibody resin to immunoprecipitate the Rpr-FLAG proteins. Immunoblot analysis of the immunoprecipitates with anti-V5 antibody showed that SfIAP co-precipitated with Rpr-FLAG (Fig. 6B). A negative control without the addition of Rpr-FLAG showed the specificity of the in vitro binding assay.

### 3.6. PmIAP and Rpr share the same subcellular locations

To investigate whether PmIAP and Rpr are co-localized in the overexpressing cells, PmIAP was expressed in SF-9 cells with or without Rpr-FLAG and then assayed with indirect immunofluorescence microscopy. SfIAP, P35, and WSSV449 were also included for comparison. Immunofluorescent signals for Rpr-FLAG were detected in cells that coexpressed
PmIAP, SfIAP, P35, or WSSV449 (Fig. 7, column 3), whereas in cells expressing only Rpr-FLAG, no immunofluorescent signal was detected (data not shown). This suggested that the amount of Rpr-FLAG protein in cells that coexpressed PmIAP, SfIAP, P35, or WSSV449 was higher than in cells expressing only Rpr-FLAG, which is consistent with our immunoblot result (Fig. 5A). Column 3 of Fig. 7 suggests that Rpr-FLAG was generally diffused in the cytoplasm, while some cells also showed punctate signals. Signal intensity, the number of cells showing punctate signals, and the number of punctates themselves were highest when Rpr-FLAG was coexpressed with PmIAP. Row 1 of Fig. 7 shows that the staining pattern for PmIAP was also changed in the presence of Rpr-FLAG (cf cols 1 and 2). The strong punctate staining of PmIAP (row 1, col 2) almost exactly matches that of Rpr-FLAG (row 1, col 3), which suggests that both PmIAP and Rpr-FLAG were co-localized to the same subcellular locations. Conversely, the staining patterns of SfIAP, P35, and WSSV449 were mostly unaffected by the presence of Rpr (Fig. 7, cf cols 1 and 2).

3.7. BIR2 and BIR3 are the minimal regions for PmIAP to interact with Rpr and prevent its pro-apoptotic activity

To determine which regions of PmIAP are important for blocking Rpr-induced apoptosis and for binding to Rpr, a series of PmIAP truncation constructs were prepared (Fig. 8A). Rpr and the truncated PmIAPs were coexpressed in SF-9 cells, and the relative viability of the transfected cells was determined. All the constructs except RING and BIR1 were able to block apoptosis with an efficiency that was seemingly related to the number of BIR domains, and this blocking effect was statistically significant for the PmIAP, BIR, BIR1+2, and BIR2+3 constructs (Fig. 8B). BIR, which was a truncated PmIAP construct with three BIR domains and no RING domain, showed the highest inhibitory activity. Since this was even higher than that of the full-length PmIAP, it may be that the RING domain negatively regulates PmIAP’s blocking of Rpr-induced apoptosis. Constructs with two BIR domains (BIR1+2 and BIR2+3) had a lower inhibitory activity, while constructs with a single BIR domain (BIR2 and BIR3) had the lowest inhibitory activity of all. Although both BIR2 and BIR3 showed no statistically significant effect on Rpr, the relative viability of the corresponding transfected cells was always higher than that of the control (vector) cells. Interestingly, both the BIR1 and RING constructs actually enhanced the apoptosis induced by Rpr. This effect was statistically significant only for BIR1, but the relative viability of the RING-transfected cells was also constantly lower than that of the control cells. We note, however, that when BIR1 or RING was expressed in SF-9 cells in the absence of Rpr, no cell death was observed (data not shown).

The PmIAP truncation constructs were further tested for their ability to delay the degradation of Rpr-FLAG (Fig. 8C). At 1 h post-heat shock, Rpr-FLAG was detected in all of the cotransfected cells. The cells that coexpressed PmIAP, BIR, BIR3, BIR1+2, or BIR2+3 had high amounts of Rpr-FLAG, while the low expression levels in the RING, BIR1, and BIR2-coexpressing cells were similar to the expression levels in the cells expressing only Rpr-FLAG. At 3 h post-heat shock, the BIR2-coexpressing cells still had a low level of Rpr-FLAG, but no Rpr-FLAG was detected in cells coexpressing Bir1 or RING, or in cells expressing only Rpr-FLAG. Meanwhile, high amounts of Rpr-FLAG were still being detected in the PmIAP, BIR, BIR3, BIR1+2, and BIR2+3-coexpressing cells. The expression of each PmIAP truncation construct was also confirmed using anti-V5 antibody (Fig. 8C, lower panel).

In vitro binding assays were then performed to determine which of these truncated PmIAP proteins interact with Rpr-FLAG. Immunoblot analysis of the immunoprecipitates with
anti-V5 antibody showed that, except for RING and BIR1, all the other truncated proteins could co-precipitate with Rpr-FLAG (Fig. 9). This suggests that the PmIAP BIR2 and BIR3 domains are the minimal regions that bind to Rpr (Fig. 9A). In Fig. 9, the amounts of the co-precipitated BIR2, BIR3, and BIR1+2 are significantly lower than those of PmIAP, BIR, and BIR2+3. This is probably because BIR2, BIR3, and BIR1+2 each have only one Rpr-binding site, whereas the other three proteins each have two Rpr-binding sites. The results also showed that, among the three truncated proteins with a single Rpr-binding site, the amount of precipitated BIR1+2 was higher than that of BIR2 or BIR3 (compare the co-immunoprecipitated bands with the respective IgG bands in Fig. 9A and B, noting that the photographic exposure time was ~6× greater for the I.P. panel in Fig. 9A). This suggests that although BIR1 does not bind to Rpr itself, it nevertheless enhances the binding of BIR1+2 to Rpr.

4. Discussion

In this paper, we report the first cloning and characterization of a novel IAP family member, PmIAP, from the tiger shrimp, *P. monodon*. PmIAP consists of 698 amino acids and has three BIR domains followed by a RING domain at the C terminus (see Fig. 1). We demonstrated that PmIAP was able to inhibit apoptosis in SF-9 cells induced by Rpr, but not the apoptosis induced by ActD (Fig. 4B,C). Further study revealed that PmIAP and Rpr physically interacted with each other (Fig. 6A and 9), and that they both shared the same subcellular locations (Fig. 7). Truncation analysis showed that the PmIAP BIR2 and BIR3 domains played important roles in inhibiting Rpr-induced apoptosis in SF cells, and that the BIR1 domain increased this inhibitory ability even though BIR1 did not bind to Rpr directly (Figs. 8 and 9). Since ectopic expression of Rpr induces apoptosis in vertebrate cells as well as in invertebrate cells [29,32,33], the results presented here imply that PmIAP exerts its anti-Rpr activity by means of mechanisms that are evolutionarily conserved across crustaceans, insects, and mammals.

In this study, three other anti-apoptosis proteins were included for comparison. SfIAP is an endogenous IAP of *S. frugiperda* cells, while P35 is a well-studied, baculoviral inhibitor of many caspases [34–36]. By contrast, there have been no further studies of WSSV449 (ORF390), since it was first identified [21]. Wang et al. [21] showed that when
apoptosis was induced in SF-9 cells by infection with a p35-deficient AcMNPV, the expression of WSSV449 blocked this apoptosis and allowed the virus to replicate. From this, the authors concluded that WSSV449 was functionally similar to P35. Here, we have shown that, in SF-9 cells, WSSV449 blocks apoptosis when induced either by ActD or by Rpr (Fig. 4B, C). This suggests that WSSV449 functions through a conserved mechanism to block apoptosis induced by different stimuli. We have also found that, in SF-9 cells, WSSV449 directly binds to the *P. monodon* effector caspase (accession no. EF114674) and inhibits its proapoptotic activity (data not shown), and we therefore confidently conclude that WSSV449 is a caspase inhibitor. Thus, the two viral caspase inhibitors (WSSV449 and P35) blocked Rpr- and ActD-induced apoptosis, whereas the two cellular IAPs (PmIAP and SfIAP) only prevented apoptosis that was induced by Rpr (Fig. 4B, C). The inability of the cellular IAPs to block the ActD-induced apoptosis suggests that the caspase-inhibiting activity of these two IAPs is less effective than that of the two viral caspase inhibitors. We also note that, under natural conditions, shrimp cells would be the normal functional environment for the PmIAP and WSSV449 anti-apoptosis proteins. Thus, ideally, a shrimp cell line would have been used for the present studies, but, unfortunately, since no crustacean cell line has yet been established, we used SF-9 insect cells instead. It is therefore reasonable that, compared with the two insect anti-apoptosis proteins, the two shrimp anti-apoptosis proteins would function less effectively in SF-9 cells.

Fig. 8 Mapping the region[s] important for PmIAP to block Rpr-induced apoptosis in SF-9 cells. (A) Schematic diagram showing the various PmIAP truncation constructs. (B) SF-9 cells were cotransfected either with 0.5 μg Rpr and 2.5 μg of empty plasmid (vector) or PmIAP constructs, or transfected with empty plasmid (vector only). At 5 h after heat shock, the number of viable cells was determined and the relative viability was calculated. The asterisk indicates a statistically significant difference between the vector and various IAP constructs (t-test; P < 0.05). (C) SF-9 cells were cotransfected with Rpr-FLAG and various PmIAP constructs. At 1 and 3 h post-heat shock, cell lysates were harvested and subjected to immunoblot analysis probing with antibodies against anti-FLAG (upper panels) or anti-V5 (lower panel; lysates from 1 h post-heat shock).
functionally and biochemically different. In mammalian XIAP, the BIR2 domain inhibits caspase-3 and -7, and the BIR3 domain inhibits caspase-9, whereas the BIR1 domain displays no caspase-inhibiting activity and is also the least conserved of the three BIR domains [12]. Furthermore, Smac relieves the inhibition of caspases by binding to either XIAP BIR2 or BIR3, but not to BIR1, and its interaction with BIR3 is considerably stronger [37,38]. In the present study, the relative importance of the BIR3 domain in PmIAP is suggested by the amino acid sequence alignment (Fig. 2), which shows that the PmIAP BIR3 is highly conserved. Conversely, the BIR1 domain is the least well conserved. The results shown in Figs. 8 and 9 revealed that, among the three BIR domains, both BIR2 and BIR3 bind to Rpr and block its apoptotic activity, whereas BIR1 does not. Furthermore, BIR3 shows a slightly greater ability to delay the degradation of Rpr protein and to block the apoptosis induced by Rpr. These interactions between PmIAP and Rpr are similar to those between XIAP and Smac, suggesting that the mechanism by which an IAP interacts with an IBM protein is evolutionarily conserved across mammals and crustaceans. The interactions between IAPs and IBM proteins are important for functionally regulating many IAPs, and we have shown here (Figs. 6 and 7) that PmIAP also interacts with an IBM protein (Rpr). Thus, although the function[s] of PmIAP have not yet been experimentally determined, it is likely that these functions are also regulated by shrimp IBM protein[s] through direct interaction. Meanwhile, even though the biological function of PmIAP remains unknown, its importance in shrimp is suggested by its gene expression profile: PmIAP is expressed in at least two different developmental stages (postlarval and adult), and in the adult stage it is expressed ubiquitously. Until recently, very few methods were available to study in vivo gene function in shrimp, but now that RNAi technology has been successfully applied in shrimp to silence gene expression [42], the next step will be to use this technology to investigate the biological function[s] of PmIAP in shrimp.

Acknowledgments

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Fig. 9  In vitro binding assay to map the region[s] important for PmIAP to interact with Rpr. The lysates containing Rpr-FLAG were mixed with the lysates containing various PmIAP truncated proteins and subjected to immunoprecipitation with anti-FLAG M2 affinity resin. The immunoprecipitates were analyzed with immunoblot probed with anti-V5 or anti-FLAG antibodies. The input lysates were also analyzed with appropriate antibodies. (A) BIR1, BIR2, and BIR3 truncated proteins were assayed. Asterisks indicate the precipitated BIR2 and BIR3 proteins. (B) PmIAP and the truncated proteins BIR, BIR1+2, BIR2+3, and RING were assayed. The IgG band is indicated.

Compared with Drosophila IAPs, PmIAP is structurally related to DIAP1 and DIAP2, and is most similar to DIAP2 (Fig. 1). When overexpressed, both DIAP1 and DIAP2 inhibit apoptosis [39], but, in a normal cellular context, they have quite different biological functions. DIAP1 is essential to the survival of Drosophila cells in vivo [39,40] and in vitro [41], whereas although DIAP2 is required for the Imd pathway to trigger the synthesis of antimicrobial peptide, it is dispensable for Drosophila’s survival [16–19]. Despite their different biological functions, both DIAP1 and DIAP2 are negatively regulated by IBM proteins through similar mechanisms. The interactions between IAPs and IBM proteins are important for functionally regulating many IAPs, and we have shown here (Figs. 6 and 7) that PmIAP also interacts with an IBM protein (Rpr). Thus, although the function[s] of PmIAP have not yet been experimentally determined, it is likely that these functions are also regulated by shrimp IBM protein[s] through direct interaction. Meanwhile, even though the biological function of PmIAP remains unknown, its importance in shrimp is suggested by its gene expression profile: PmIAP is expressed in at least two different developmental stages (postlarval and adult), and in the adult stage it is expressed ubiquitously. Until recently, very few methods were available to study in vivo gene function in shrimp, but now that RNAI technology has been successfully applied in shrimp to silence gene expression [42], the next step will be to use this technology to investigate the biological function[s] of PmIAP in shrimp.
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