A silkworm hemolymph protein is a prophenoloxidase activation blocker

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ABSTRACT
Melanization in insect hemolymph is triggered by the recognition of pathogen-associated molecular patterns via pattern recognition receptors. The signal transduction leads to the activation of the prophenoloxidase and hence the generation of melanin. The proPO activation process must be tightly controlled to minimize the host damage caused by reactive intermediates during melanin synthesis. The full-length cDNA sequence of a 20 kDa hemolymph protein from Bombyx (Bmhp20) was determined. Bmhp20 gene was expressed in larval fat body, integument, trachea, and ovary and was induced by the challenge of B. bombyseptieus. Binding of recombinant Bmhp20 to microbial cell wall components as well as gram-positive bacteria and fungi was confirmed. Phenoloxidase activity assay indicated that recombinant Bmhp20 blocked the proPO activation in hemolymph that was triggered by peptidoglycan or beta-1, 3-glucan. Our data suggest that Bmhp20 plays bifunctional roles in silkworm humoral responses: to participate in pattern recognition and to block the activation of proPO.

Keywords: Bombyx mori; Humoral Responses; Pattern Recognition Receptors; Phenoloxidase Activity

1. INTRODUCTION
Humoral responses of immunity system in insect such as melanization and induced production of plasma proteins are essential for the host to fight against the pathogens [1,2]. For its large body and easily obtained hemolymph, silk-worm is one of the earliest insects used to study humoral responses [3,4].

Pathogen-associated molecular patterns (PAMPs) are conserved structures present on the surfaces of microbes including peptidoglycan (PGN), beta-1, 3-glucan, lipoteichoic acid and lipopolysaccharide [5]. The humoral responses of silkworm are triggered by the recognition of PAMPs via pattern recognition receptors (PRRs) [6]. Yoshida and Ashida (1986) purified a beta-1, 3-glucan recognition protein in silkworm plasma, which can stimulate the activation of the prophenoloxidase (proPO) cascade [3]. Characterization of silkworm peptidoglycan recognition protein (PGRP) indicated that in the absence of PGRP, the proPO cascade could not be triggered by peptidoglycan [7]. Activation of proPO in insects by a cascade of proteolytic cleavages results in the generation of a phenoloxidase (PO). PO oxidizes phenolic compounds to produce quinines that polymerize to form melanin [8,9]. Insects used melanin to encapsulate non-self organisms and seal wounds. Since reactive intermediates for melanin synthesis are thought to be harmful to the host insects, the proPO activation process must be tightly controlled to minimize the damage of the host by melanin. Previous studies showed that the activation of proPO system is regulated by serine protease inhibitors and PO inhibitors [10-13].

The availability of silkworm genome makes identification of large number of PRR genes. PGRP genes in silkworm are clustered into two groups located on chromosome 1 (PGRP1-5) and 16 (PGRP 8-11) [14]. Three C-type lectins were identified by searching silkworm genome database and their microorganism recognition specificities were analyzed [15]. The expressions of two PGRP short-type proteins (PGRP-S2 and PGRPS5) and four beta-glucan recognition proteins were induced by the oral infection of Bacillus bombyseptieus [16]. In silkworm, mRNAs of most of PRRs usually are identified in fat body and epidermis. They are synthesized and secreted into the hemolymph, where the recognition occurred. Larval hemolymph contains a large number of proteins involved in humoral responses including hemolin, prophenoloxidase, PGRP-SA, serine proteinase-like proteins, and protease inhibitors [17,18]. In our previous studies, when we tried to isolate the proteins interacting with a protease inhibitor, we obtained a hemolymph protein (Bmhp20) and its amino terminal sequence was determined [19].

*The authors declare that they have no competing interests.
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The present research is focused on the characterization of this silkworm hemolymph protein. Its full-length cDNA sequence was obtained. We found that the deduced amino acid sequences of Bmhp20 are rich in putative glycosaminoglycan attachment sites. The expression of Bmhp20 gene was enhanced by B. bombyseptieus. In vitro binding assays revealed that recombinant Bmhp20 bound to gram-positive bacteria and fungi. The result of phenoloxidase activity assay indicated that the recombinant Bmhp20 blocked the proPO activation in hemolymph that was activated by PGN or β-1, 3-glucan.

2. EXPERIMENTAL PROCEDURES

2.1. Silkworm Larvae and Regents
Silkworm strain p50 is maintained at the State Key Laboratory of Silkworm Genome Biology, Southwest University, China. The larvae are reared on mulberry leaves at 25°C. The curdlan (Sigma, USA), cellulose (Sigma, USA), and chitin (NEB, USA) were purchased and separately suspended in 100 µl PBS (pH 7.4, 200 mM NaCl).

2.2. Database Searching, Gene Cloning, and 5’ RACE
The amino acid sequence data from Edman degradation analysis were blasted against the protein database at SilkDB (http://silkworm.swu.edu.cn/silkdb/). The identified protein sequence and its corresponding nucleic acid sequences (predicted gene BGIBMGA002175) were used as references for primer design.

Total RNA was extracted from the fat body of the fifth instar day 3 larvae using Trizol reagent (Invitrogen, China). RNA samples were treated with DNase I (DNA-free, Ambion) at 37°C for 1 h to remove the DNA contamination. The concentration of RNA was measured in a spectrophotometer (Amersham Biosciences, Sweden). Three micrograms of total RNAs were used to synthesize first-strand cDNA in 25 µl of reverse transcription PCR (RT-PCR) system with M-MLV Reverse Transcriptase (Promega, USA).

Two primers (forward: 5’-ATGGCACTGAAAATTACTTT-3’; reverse: 5’-GGAGCTCTAAGCTGTAAA-3’) were synthesized (Genscript, China) for PCR amplification using the fat body cDNA. PCRs were performed as follows: 94°C for 3 min, 35 cycles at 94°C for 40 s, 55°C for 40 s, 72°C for 40 s, followed by incubation at 72°C for 10 min. PCR products were cloned into pMD19-T simple vector (TaKaRa, China). According to the manual of 5’ RACE system Version 2.0 (Invitrogen, China), the partial sequenced sequence were used to design the primers GSP1 (5’-GTGGTCCAAATCCGAGATCTGGT-3’), GSP2 (5’-CGAACCCAGATCGACCGCAA-3’), and nested primer (5’-AACTGCCGAAGCCGCTATGA-3’). The first-strand gene specific cDNA was reverse-transcribed from fat body RNA by GSP1 using SuperScript™ II (Invitrogen, China). The specific cDNA was tailed with poly C. Tailed cDNA was used as template to reamplify the target gene using Abridged Anchor Primer 1 (5’-GGCCACGCGTCGACTACGGGGGGG-3’) and GSP2. The product was used for the third round PCR amplification using Abridged Universal Amplification Primer 2 (5’-GGCCACGCGTCGACTAGTAC-3’) and nested primer. The amplified nucleic acid fragment was cloned into pMD19-T simple vector (TaKaRa, China) and sequenced.

2.3. Gene Structure and Sequence Analysis
Gene translation and the prediction of molecular weight and theoretical pI were performed with ExPASy (http://www.expasy.ch/). Gene structure was drawn by SIM4 (http://pbil.univ-lyon1.fr/members/duret/cours/inserm210604/exercise4/sim4.html). Sequence homologous was analyzed on NCBI. Signal peptide prediction was carried out using the program SignalP 3.0 (http://www.cbs.dtu.dk/services/SignalP/). Functional sites in protein were predicted using ELM database (http://elm.eu.org/). TESS (http://www.cbi.upenn.edu/cgi-bin/tess/tess) was used to predict the Upstream regulatory sequences.

2.4. RNA Extraction and RT-PCR
To investigate the expression of Bmhp20 mRNA in tissues, the RNAs from ten different tissues (including fat body, integument, ovary, trachea, head, testis, silk gland, midgut, hemocyte, and malpighian tubule) of the fifth instar day 3 larvae were extracted. For developmental expression profile of Bmhp20, RNAs were prepared using whole larval body from the 1st to the 4th instar larva day 1 to adult. RNAs isolation and digestion were carried out as mentioned above. The amplification of the BmAthin3 served as an internal control. The Bmhp20 primers (forward: 5’-TTTGGAAATTCGCAACCCCT-3’, reverse: 5’-GATAACCGCCAATGAAATG-3’) and BmAthin3 primers (forward: 5’-AACACCCGCTCCGTACTG-3’, reverse: 5’-GGGCCGACGTGATTTTCCCT-3’) were used for RT-PCR. PCRs were performed for one cycle at 94°C for 3 min, 25 cycles at 94°C for 40 s; 55°C for 40 s; 72°C for 40 s; followed by incubation at 72°C for 10 min.

2.5. Expression, Purification of Recombinant Bmhp20, and Production of Antiserum
The cDNA fragment encoding the mature Bmhp20 was...
amplified using the fat body cDNA by the following primers: forward: 5'-CTAgctagcTTCCCTCATGGCAGTAGCG-3', reverse: 5'-CgagctcGTGTGACATGCGCTAGATG-3', and then cloned into pMD19-T simple vector (TaKaRa, China) for sequencing. The recombinant plasmid of pET28a-Bmhp20 was constructed. One Rosetta colony containing pET28a-Bmhp20 was incubated at 37°C in 400 ml of LB medium supplemented with 25 µg/ml kanamycin until the OD600 reached 0.4 - 0.8. A final concentration of 1 mM IPTG was added into the culture and then induced for 3 h at 37°C. Cells were harvested by centrifugation. All the following purification procedures were conducted at 4°C unless otherwise noted. Cells were washed for 3 times with 20 mM PBS buffer (pH 7.4, 500 mM NaCl), and then suspended in 20 ml of same PBS buffer. Cells were disrupted by sonication on ice and separated by centrifugation for 30 min at 16,000× g. The supernatant was loaded onto a Ni-NTA affinity agarose column (Invitrogen, China). The fractions eluted with 20 mM PBS (pH 7.4, 500 mM NaCl) containing 150 mM imidazole were collected. A gel filtration chromatography was subsequently performed using a Superdex 75 column (AKTA purify). The column was washed with 50 mM PBS (pH 7.4, 200 mM NaCl) at a linear velocity of 1ml/min. The fractions containing His-Bmhp20 were combined and the proteins were digested by thrombin (Novagen, USA). After the cleavage reaction, biotinylated thrombin was removed with Streptavidin Agarose, and then concentrated by ultrafilter (Amicon Ultra-15 Millipore, USA). The protein concentration was determined with BCA kit (Beyotime, China). The cleaved Bmhp20 served as antigen for the production of polyclonal rabbit antiserum. A total of 500 µg of cleaved Bmhp20 was equally mixed with complete Freund’s adjuvant (Sigma, USA). The resultant emulsified mixture was injected into rabbit. Subsequent booster injections in the rabbit were carried out for three times (250 µg per time, once a week). Seven days after the last injection, the rabbit was bled and the blood sample was collected. The antiserum was purified by MAbTrap™ Kit (GE, USA).

### 2.7. Microbial Challenge and Real Time PCR

The fifth instar day 3 larvae were injected with the formaldehyde-fixed *E. coli* (10⁸ cells per larvae), *B. bombyseptieu* (10⁷ cells per larvae), *B. bassiana* (10⁵ cells per larvae), respectively. Injection of 10 µl of 0.85% NaCl was used as a negative control. Fat body from challenged larvae was obtained 1, 3, 6, 12, 24 h after injection. The RNAs were isolated and first-strand cDNAs were synthesized as mentioned above. For real-time PCR, primers of Bmhp20 (forward: 5'-AGTAGCACAAGCTTCCCTCATG-3'; reverse: 5'-ACTGCCGAAGCCGCTATGAC-3') and primers of sw22934 (forward: 5'-TCGTACTGGCTTCTTCGT-3'; reverse: 5'-CAAAGTTGATAGCAATTCCCT-3') were designed. The primer annealing temperatures was 60°C and the size of product was 139 bp. Primer specificity was visualized by 2% agarose gel. The PCR efficiency was within 95% - 110%. The real-time PCRs were performed using the SYBR Premix Ex Taq kit (TaKaRa, China). Each reaction in 25 µl reaction cocktail including 12.5 µl of Ex Taq, 70 ng cDNA, 10 mM Bmhp20 forward and reverse primers. The PCR reaction was performed on an ABI Prism 7000 Sequence Detection System (Applied Biosystems) using the following program: initial denaturation at 95°C for 10 s, and 40 cycles of 95°C for 5 s, 60°C for 31 s. The primers of sw22934 (transcription initiation factor 2 gene) was used as internal standardization.

### 2.8. In Vitro Binding Assay of Bmhp20 with Microbial Cell Wall Components and Microbes

In vitro binding assays were executed according to published protocol [20]. Six kinds of log phase microbes (3 ml), *S. aureus*, *B. bombyseptieus*, *S. marcescens*, *P. aeruginosa*, *B. bassiana*, and *P. pastoris* were collected and fixed with 2% formaldehyde for 2 h. The cells were washed by PBS buffer (pH 7.4, 200 mM NaCl) and suspended in 100 µl of the same PBS. In addition, 2 mg of curdlan, cellulose, and were suspended in 100 µl PBS (pH 7.4, 200 mM NaCl). Two micrograms of Bmhp20 was then added to the above cell/polysaccharide solution and incubated for 2 h at room temperature. After centrifugation, the pellets were washed for five times with PBS buffer. The remaining pellets were resuspended in 100 µl PBS buffer and boiled in SDS sample buffer for 5
min. The proteins detached from the pellets were analyzed by Western blotting.

2.9. In Vitro Phenoloxidase Activity Assay
Silkworm larva plasma (SLP) reagent (Wako, Japan) was used to measure the phenoloxidase (PO) activity. Two types of polysaccharide, PGN (53243 Sigma, USA) extracted from *M. luteus* and laminarin (L9634 Sigma, USA), a kind of β-1,3-glucan extracted from *L. digitata*, were used as inducers. The assays were performed as described in manufacturer’s instructions. In brief, 20 µl recombinant Bmhp20 (1.5 µg/µl), 20 µl PGN (0.2 ng/µl) or 20 µl laminarin (0.2 ng/µl), and 10 µl double distilled water (ddH₂O) were mixed together. After incubated at 27°C for 5 min, 50 µl of SLP reagent was added. Then the mixture was incubated for another 30 min at 27°C. PO activity was monitored at absorbance of 450 nm. The assay was repeated three times.

3. RESULTS AND DISCUSSION

3.1. Full-Length cDNA and Sequence Analysis of Bmhp20
The sequence of 20 residues “HGSSDVGDGGGEVEAVAGTLK” was used to search the silkDB (http://silkworm.swu.edu.cn/silkdb) with TBLASTN program. A protein (BGIBMGA002175) with the same deduced amino acid sequence was found. However, the 5’cDNA sequence of this gene is absent. We then performed 5’RACE to clone its full length cDNA. As a result, a 740-bp cDNA sequence was obtained, as shown in Figure 1(a). This gene has an open reading frame of 648 bp, encoding a 215-residue polypeptide with an 18-residue signal peptide predicted by SignalP 3.0. The mature protein has a calculated molecular weight of 21.6 kDa and a pI of 5.5. This protein was so named as *Bombyx mori* hemolymph protein 20 (Bmhp20, accession number: GU015849) based on its size and presence in hemolymph. Also shown in Figure 1(a), a predicted transcription start site A was found 13 nt upstream of the ATG codon. A polyadenylation sequence AATAAA was observed at the end of Bmhp20 mRNA.

Comparison of the cDNA and its genome sequences revealed that Bmhp20 had three exons separated by two introns (Figure 1(b)). The first exon consists of 13 bp 5’ untranslated region and partial coding sequence of the signal peptide. The second exon encodes residues 5 - 17 of the signal peptide. The last exon codes for the remaining residue of signal peptide and the mature protein of Bmhp20. It was found that Bmhp20 had only one copy in silkworm genome which was located on chromosome 15.

The deduced amino acid sequences of Bmhp20 were shown in Figure 1(c). The amino acid composition of Bmhp20 has a low complexity, which is rich in glycine (26.9%), serine (13.2%), phenylalanine (12.2%), aspartate acid (10.7%), and histidine (9.1%). It is worth to mention that Bmhp20 is rich of glycosaminoglycan attachment sites (DGSGE, DNSGF, NSGF, HSGF, DSGF, and DSAL) predicted by ELM [21], as shown in Figure 1(c).

3.2. The Temporal and Spatial Expression Profiles of Bmhp20 Gene
RT-PCR was carried out using RNAs extracted from fat body, integument, ovary, trachea, head, testis, silk gland, midgut, hemocyte, and malpighian tubule of the fifth instar day 3 larvae. As shown in Figure 2(a), the Bmhp20 transcripts were found to be expressed in larval fat body, integument, trachea, and ovary. Furthermore, the temporal expression profile of Bmhp20 was investigated by RT-PCR. The results in Figure 2(b) showed that Bmhp20 gene was expressed in all of the larval stages. When the larva entered the wandering stage, the expression of Bmhp20 was highly expressed at the first day and its expression was decreased at W2 and W3 (the second and third day of the wandering stage, respectively). At the pupal stages, the transcripts of Bmhp20 were undetectable from P1 to P3, up-regulated at P4, and then dramatically decreased from P5 to P7 (P1 - P7, day 1 - 7 of pupa).

3.3. The Transcriptional Regulation of Bmhp20 against Microbial Challenge
Real-time RT-PCR was carried out to measure the transcriptional regulation of Bmhp20 in the fat body challenged by *B. bombyseptieus*, *Beauveria bassiana*, and *Escherichia coli*. As illustrated in Figure 2(c), Bmhp20 was rapidly induced by all challenges after injections for 1 h. However, after injection for 3 to 6 h, the expressions of Bmhp20 were leveled off. Interestingly, in contrast to *B. Bassiana* and *E. coli*, *B. bombyseptieus* strongly induced the expression of Bmhp20 (up to 6-fold) after injection for 12 to 24 h. Our data indicated that Bmhp20 might involve in the immune responses of silkworm.

3.4. Production and Purification of Recombinant Bmhp20
His-Bmhp20 was expressed in prokaryotic expression system and purified by a two-step chromatography. His-tag at the N terminal of his-Bmhp20 was removed with biotinylated thrombin. As shown in Figure 3, on a SDS-PAGE, the band of purified recombinant Bmhp20 without his-tag migrated with a molecular size around 26 kDa which is different from the putative molecular weight of Bmhp20. We assumed that the delayed migration of Bmhp20 on SDS-PAGE may result from that the SDS-stable structures of Bmhp20.
Figure 1. Gene sequence of Bmhp20 and its deduced amino acid sequence. (a) The cDNA sequence of Bmhp20 protein. The deduced amino acid sequence is shown below the cDNA sequence. The transcriptional start site of A is indicated by an arrow and is designated +1. The polyadenylation sequence AATAAA is in grey frame. Predicted signal peptide is boxed and the N-terminal sequence determined by Edman degradation is double-underlined; (b) Genomic structure of Bmhp20 gene. Black boxes represent exons; (c) The deduced amino acid sequence of Bmhp20 mature protein. The predicted glycosaminoglycan attachment sites are shown in boldface italic.

3.5. Bmhp20 Is Secreted into Hemolymph

The distribution of Bmhp20 in different tissues was examined by Western blotting. Same amount of proteins were separated by SDS-PAGE and stained by silver as a loading control (Figure 4(b)). On the membrane, as shown in Figure 4(a), only one band in hemolymph was detected, indicating that the mature Bmhp20 was secreted into hemolymph. The Bmhp20 was previously annotated as a cuticular protein (CPRG16, accession number: NM_001167717) for its rich of glycine and owning a GGX/AAP repeat sequence [22]. The result of Western blotting assay obviously showed that Bmhp20 is a hemolymph protein, rather than a structural protein present in the silkworm cuticles.

3.6. Recombinant Bmhp20 Binds to Various Microorganisms in Vitro

As mentioned above, the analysis of the amino acid sequence of Bmhp20 revealed that it has abundant putative glycosaminoglycan attachment sites (Figure 1(c)). We assumed that Bmhp20 might be involved in the pattern recognition. Therefore we investigated the binding activity of Bmhp20 by three kinds of microbial cell wall components (Figure 5(a)). Bmhp20 was incubated with these polysaccharides separately. After washing for five times with PBS, the proteins eluted from polysaccharides were detected by Western blotting. The results showed that Bmhp20 was bound to chitin, cellulose, and curdlan. In vitro binding assays of recombinant Bmhp20 were thereafter performed with gram-positive bacteria (Staphylococcus aureus and B. bombesiepticus), gram-negative bacteria (Serratia marcescens, and Pseudomonas aeruginosa), and fungi (B. bassiana and Pichia pastoris). As shown in Figure 5(b), Bmhp20 bound to S. aureus, B. bombesiepticus, B. bassiana, and P. pastoris. Recombinant Bmhp20 also had weak binding activity to S. marcescens and there was no protein eluted from P. aeruginosa.
Figure 2. The expression of Bmhp20 gene. The spatial (a) and temporal (b) expression profiles of Bmhp20 were performed by RT-PCR as described in Material and Methods. (a) The tissue distribution of Bmhp20 in silkworm larvae at fifth instar day 3 including trachea (TR), head (HD), ovary (OV), testis (TE), fat body (FA), silk gland (SL), integument (IT), hemocyte (HC), malpighian tubule (MA), and midgut (MD); (b) The developmental expression profile of Bmhp20 gene from the first instar larvae, day 2 (I2) to adult day 1 (A1). I2 stands for day 2 of the 1st instar larvae, II2 day 2 of the 2nd, III2 and IV2 day 2 of the 3rd and 4th, respectively. V1 to V7, day 1 - 7 of the 5th instar larvae. W1-W3, day 1-3 of the wandering phase. P1 to P7, day 1 - 7 of the pupa. A1, day 1 of the adult; (c) Quantitative real-time PCR analysis of Bmhp20 mRNA in fat body after injection of *E. coli*, *B. bassiana* and *B. bombyseptieus* at the indicated time points. The error bars represent the standard error of the mean (n = 3).

suggesting little binding of Bmhp20 to this gram-negative bacterium.

3.7. Recombinant Bmhp20 Inhibits the PO Activity That Is Activated by Microbial Cell Wall in Vitro

PGN or β-glucan activates the PO cascade of silkworm larval plasma. The produced melanin can be easily detected with a substrate 3, 4-dihydroxyphenylalanine [23]. Silkworm larva plasma (SLP) reagent has been used for the detection of PGN in human plasma (Kobayashi et al., 2000). Since Bmhp20 bound to Gram-positive bacteria and fungi cell walls, we then used the SLP reagent to examine whether Bmhp20 was involved in the activation of PO cascade. The activated PO activity was monitored at absorbance of 450 nm after the SLP reagents incubated with the test samples at 27°C for 30 min. As shown in Figure 6, compared to SLP alone, when PGN or laminarin was added into the SLP, the PO activities dramatically increased. In the case of his-Bmhp20, the PO activity was not changed by recombinant protein itself. In this context, when PGN or laminarin was added into
Figure 3. The production of the recombinant Bmhp20 protein. Purified recombinant his-Bmhp20 and Bmhp20 without his tag were analyzed by 15% SDS-PAGE and the gel was stained with Coomassie Brilliant Blue.

the SLP reagent pre-incubated with his-Bmhp20, the PO activities were no longer increased, indicating that Bmhp20 acts as a proPO activation blocker. Previous studies of proPO-activating proteinases in silkworm revealed at least two enzymes, i.e. an esterase hydrolyzing N-α-benzoyl-L-arginine ethyl ester (BAEEase), a prophenoloxidase activating enzyme (PPAE) were involved in the activation pathway [25,26]. BAEEase activity was shown to be inactivated by silkworm antitrypsin, whereas the activation of PPAE was unaffected by protease inhibitors [27]. Phenoloxidase inhibitor (POI) was isolated from the housefly Musca domestica and Manduca sexta [12,13]. Bmhp20 has little sequence similarity with above protease inhibitors and low molecular weight peptides. Binding to microbial surfaces and blocking the proPO activation make Bmhp20 a distinct hemolymph protein involved in the silkworm humoral responses.

Figure 4. The expression of Bmhp20 protein. Western blotting analysis of the tissue distribution of Bmhp20 in the fifth instar day 3 silkworm larvae. The proteins were isolated from the trachea (TR), head (HD), ovary (OV), testis (TE), fat body (FA), silk gland (SL), integument (IT), heymocyte (HC), malpighian tubule (MA), hemolymph (HM), and midgut (MD).

Figure 5. In vitro binding assay of Bmhp20 protein with various polysaccharides and microbes. (a) Binding assay of Bmhp20 with cellulose, chitin, and curdlan; (b) Binding assay of Bmhp20 to gram-positive bacteria (S. aureus, B. bombyseptieus), gram-negative bacteria (S. marcescens, P. aeruginosa), and fungi (B. bassiana, P. pastoris). The bands were detected by Western blotting. S: supernatant; W5: fraction from the fifth wash; P: pellet.
To our knowledge, proteins with recognition activities have not yet been reported to inhibit the proPO-activating system of insect. It is concluded that for the first time, we identified a silkworm hemolymph protein participating in pattern recognition as well as blocking proPO activation.

4. AUTHORS’ CONTRIBUTIONS

NH conceived of the study and developed the study design. BL and XQ carried out the analysis and BL drafted the manuscript. QH did the PO activity assay. LJ performed the binding assay. ZX participated in the study design. NH contributed to the critical revision of the manuscript. All authors read and approved the final manuscript.

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LIST OF ABBREVIATIONS

PAMPs, Pathogen-associated molecular patterns; PRRs, pattern recognition receptors; proPO, prophenoloxidase; PGRP, peptidoglycan recognition protein; PO, phenoloxidase; PGN, peptidoglycan; IPTG, isopropyl β-D-thiogalactoside; SLP, silkworm larva plasma; BAEEase, Nα-benzoyl-L-arginine ethyl ester; PPAE, prophenoloxidase activating enzyme.