

Molecular characterization, tissue distribution, and immune reaction expression of karyopherins in the domestic silkworm (*Bombyx mori*)

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ABSTRACT. Karyopherins, including alpha and beta types, are transport proteins in the eukaryotic cell that carry cargoes across nuclear pore complexes into or out of the nucleus. In this study, full open reading frames of one beta and three alpha types of karyopherin were cloned from cDNA of the domestic silkworm (Bombyx mori). The one beta and three alpha types' open reading frames were 2661, 1563, 1515, and 1551 base pairs long, respectively, and coded 886, 520, 504, and 516 amino acids, respectively. The alphas all had one importin-beta-binding (IBB) domain, and eight, four, or seven armadillo/beta-catenin-like repeats. The beta had 19 HEAT repeat domains, which constructed one importin-beta-N-terminal domain and one IBB domain. The recombinant proteins were expressed in Escherichia coli cells. The molecular weight of the beta type was approximately 100 kDa, and the alphas weighed approximately 60 kDa. Phylogenic tree construction revealed that the alphas could be classified into three known karyopherin-alpha subfamilies. We detected mRNA of the four karyopherins in normal 3rd day of 5th instar larvae, and in larvae injected with Gram-positive bacteria, Gramnegative bacteria, viruses, and fungi using real-

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time fluorescence quantitative reverse transcriptase-polymerase chain reaction, and found that the four karyopherins were widely distributed, but their expression levels were related to tissues type, the microbe injected, and the time point.

Key words: *Bombyx mori*; Karyopherin; Gene clone; Protein expression; Real-time fluorescence quantitative RT-PCR; Immune reaction

INTRODUCTION

The nuclear envelope (NE) of eukaryotic cells has evolved in order to separate the cytoplasm and the nucleoplasm into two relatively independent regions. Genetic materials are restricted in the nucleus, so protein translation takes place in the cytoplasm. The proteins needed for DNA replication, mRNA transcription, gene regulation, nuclear skeleton repair, or for integrating into the inner nuclear membrane must be transported from the cytoplasm to the nucleoplasm through nuclear-pore complexes (NPCs), which penetrate the NE and connect the cytoplasm with the nucleoplasm. Proteins less than about 40 kDa in weight can go through NPCs directly, but most large proteins need carriers to take them into the nucleus (Pante and Kann, 2002; Terry et al., 2007). Karyopherins, also called importins (IMPs), are important carriers in eukaryotes. There are two types of karyopherins: karyopherin-beta (KPNB), also named importin-beta (IMPβ), which can carry cargos through NPCs and interact with FG nucleoporins or Ran-ATP to release their cargoes; and karyopherin-alpha (KPNA), also named importin-alpha (IMPa), is a cargo receptor. KPNA recognizes classic nuclear location sequences (cNLSs) on cargoes and KPNB recognizes importin-beta-binding (IBB) domains on KPNA. Cargoes, KPNA, and KPNB form a ternary complex. The mechanism behind the ability of the complex to go through NPCs is unclear, but probably relates to an interaction between KPNB and nucleoporins (Kapinos et al., 2014). In the nucleus, Ran-ATP binds to KPNB, and the ternary complex disassembles and the cargo is released.

Yeast has one type of KNPA, *Drosophila* has four, the mouse has six, and other mammals have seven. Based on their molecular characteristics and phylogenetics, KNPAs are divided into three subfamilies, and the cargoes they carry differ (Huenniger et al., 2010; Sachan et al., 2013; Zhou et al., 2013). Yeast, *Drosophila*, mice, and other mammals all have at least one KPNB. KPNAs and KPNBs play important roles in animal physiology, such as development, immunity, and stress responses (Gorjanacz et al., 2006; Ratan et al., 2008; Wirthmueller et al., 2013; Young et al., 2013; Angus et al., 2014; Schaller et al., 2014). Viruses also use karyopherins to carry their structures, or non-structural proteins, in and out of the nuclei of host cells in order to replicate themselves, e.g., the influenza virus nucleoprotein (Wang et al., 1997; Sasaki et al., 2013). The VP24 proteins coded by the Ebola virus can bind karyopherins to inhibit the immunological responses of an infected cell (Mateo et al., 2010).

The domestic silkworm (*Bombyx mori*) is one of the most important economic insects in China, Japan, and Southeast Asia, and has been used as a model Lepidopteran (Xia et al., 2004, 2009). We BLAST-searched the domestic silkworm genome database (SilkDB; http://silkworm. genomics.org.cn/) and found three predicted KPNAs (BGIBMGA003758-TA, BGIBMGA011351-TA, and BGIBMGA012556-TA) and one predicted KPNB (BGIBMGA002972-TA). We designed primers to clone the four genes, expressed them in *Escherichia coli* cells, and compared their molecular characteristics, tissue distributions, and expression patterns during immune reactions in the silkworm strain p50.

Genetics and Molecular Research 14 (4): 13049-13065 (2015)

MATERIAL AND METHODS

Experimental animals and treatments

The domestic silkworm used for the experiments was the p50 strain, which was maintained in the Key Laboratory of Sericultural Biotechnology, Anhui Agriculture University, China. The 5th 3-day-old larvae were divided into five groups, and each group contained at least 16 individuals. The larvae in Group 1 were injected with 4 μ L phosphate buffer solution (PBS; pH, 7.4). The larvae in Groups 2 to 5 were injected with 4 x 10₆ inactivated *E. coli*, 4 x 10₆ *Beauveria bassiana*, 4 x 10₆ nuclear polyhedrosis virus (NPV), and 2 μ g *Micrococcus luteus*, respectively. At the time points of 1, 4, 8, and 12 h after injection, the larvae of Groups 1 to 5 were dissected in order to collect blood and the fat body. Untreated 5th 3-day-old larvae were also dissected, not only to collect blood and the fat body but also the testes, ovaries, Malpighian tubules, midguts, silk glands, and integuments. At each time point, at least four larvae were dissected and the same tissues were mixed together. The experiment was repeated three times.

RNA extraction and cDNA preparation

Total RNA from the above tissues was extracted using the TRIzol method (Takara, Dalian, China). RNA quantity and quality were checked by detecting the values of OD260 and OD280, and also by RNA electrophoresis. The cDNA used for the cloned genes was produced using a TransScript_® First-Strand cDNA Synthesis SuperMix (Transgen, Beijing, China), and the cDNA used for the real-time fluorescence quantitative reverse transcriptase-polymerase chain reaction (RT-PCR) was produced using a PrimeScript_M RT Master Mix (Takara, Dalian, China).

Cloning and real-time fluorescence quantitative RT-PCR primer design

Primers were designed using Primer Premier 5.0 software, and the PCRs were verified by DNA sequencing. Real-time fluorescence quantitative RT-PCR primers of *B. mori* 18s rRNA have been reported previously (Zhang et al., 2012). The primers and their annealing temperatures are listed in Table 1.

Plasmid construction and recombinant prokaryotic protein expression

Open reading frames (ORFs) of the Bm-KPNAs and Bm-KPNB1 were cloned in a pGEM_®-T Easy Vector (Transgen) and amplified in Trans-T1 cells (Transgen). Plasmids were extracted using an AxyPrep_{TM} Plasmid Miniprep Kit (Axygen, Suzhou, China). Bm-KPNA1, Bm-KPNA3, and Bm-KPNB1 were double-digested by *Bam*Hl/ *Xho*l (Takara), and KPNA2 was digested by *Bam*Hl/ *Sal*l (Takara) at 37°C for 4 h. The prokaryote expression plasmid pET-28a+ was also double-digested by the above enzyme pairs, under similar conditions. Target fragments were collected by agarose electrophoresis and purified using an AxyPrep_{TM} DNA Gel Extraction Kit (Axygen). The targeted gene fragments and pET-28a+ were linked by T4 ligase (Takara) at 25°C for 1 h. The recombinant plasmids were verified by DNA sequencing and transferred into Transetta (DE3) cells (Transgen). Recombinant protein expression was induced by different concentrations of isopropyl β -d-1-thiogalactopyranoside (IPTG) (Sangon, Shanghai, China) at 30°C for 4 h.

Genetics and Molecular Research 14 (4): 13049-13065 (2015)

J. Li et al.

Table 1. Primers used for cloning and real-time fluorescence quantitative reverse transcriptase-polymerase chain reactions (RT-PCRs).

Gene name	Primer sequence (5' to 3')	Annealing temperature	PCR product size (bp
Open reading frame primers			
Bm-KPNA1	F: CGC <u>GGATCC</u> ATGTCTGGAGGTCAC	58°-0.2°C/cycle	1581
		35 cycles	
	R: CCG <u>CTCGAG</u> GTTAAAACTGAAAGC		
Bm-KPNA2	F: CGC <u>GGATCC</u> ATGGCTGAACGAACCTCG	60°-0.2°C/cycle	1533
		40 cycles	
	R: GACGTCGACCTAGAAGTGTATGGCTGC		
Bm-KPNA3	F: CGC <u>GGATCC</u> ATGGCGACTGATCAAGCGAAAAAC	60°-0.2°C /cycle	1569
		40 cycles	
	R: CCG <u>CTCGAG</u> CTAGAAGCGGA		
Bm-KPNB1	F: CGC <u>GGATCC</u> ATGCACGCAGAAAC	58°C	2679
		30 cycles	
	R: CCG <u>CTCGAG</u> TTACCAGCTGGCAAGAGG		
Real-time fluorescence			
quantitative RT-PCR primers			
Bm-KPNA1	F: GTTCTCCCACCCTTGCTT	60°C	231
	R: TTTCTCGTTCGGACCATCT		
Bm-KPNA2	F: CAGGCTGCGGCGAAATAC	60°C	201
	R: GAACTGATACTGCTCGTGGTC		
Bm-KPNA3	F: TGATTCCGTTGTTGTCGC	60°C	210
	R: GTTTATTACCTGCCGTGATGT		
Bm-KPNB1	F: TGCTTTGAGTATTGGTAGCGATTT	60°C	194
	R: GTCGTACCTGAGAACCTACACCT		
Bm-18S	F: CGATCCGCCGACGTTACTACA	60°C	180
	R: GTCCGGGCCTGGTGAGATTT		

Underlined text refers to enzyme digestion sites.

Protein electrophoresis and western blot analysis

The cells containing recombinant proteins were centrifuged at 12,000 rpm for 1 min. After adding 24 μ L PBS and 6 μ L 5X protein loading buffer, the precipitates were placed in boiling water for 10 min before being stored in ice. Subsequently, 10 μ L of the protein solution was used for 12% polyacrylamide gel electrophoresis (PAGE) and dyed with Coomassie brilliant blue R250 (Sangon), or used for western blot analysis.

Western blot analysis was initiated by transferring the proteins from the PAGE gels to polyvinylidene difluoride films (Millipore, USA). The films were washed with PBS with Tween® 20 (PBST) three times, each time for 10 min. A non-specific antibody recognized sites on the films blocked with 5% fat-free milk powder, and the films were then incubated with mouse anti-His monoclonal antibody (Transgen) for 1 h at room temperature. The films were washed with PBST another three times, and then incubated with HRP-labeled goat anti-mouse polyclonal antibody (Transgen). After again being washed three times with PBST, the films were colored using a DAB kit (Sangon).

Bioinformatics analysis

The proteins' molecular weights and isoelectric points were analyzed using the ExPASY Compute pl/Mw tool (http://web.expasy.org/compute_pi/). Domains and repeats were analyzed using Pfam (http://pfam.xfam.org/) (Punta et al., 2012; Finn et al., 2014). KPNA and KPNB protein sequences from different organisms were collected from online protein databases such as GenBank (http://www.ncbi.nlm.nih.gov/genbank/), MGI (http://www.informatics.jax.org/), FlyBase

Genetics and Molecular Research 14 (4): 13049-13065 (2015)

(http://flybase.org/), and BeeBase (http://hymenopteragenome.org/beebase/) for phylogenetic analysis. Multiple sequence alignments were conducted using ClustalX 2.1, and phylogenic trees were constructed based on amino acid (AA) sequences by MEGA 6.06, using the neighbor-joining method with a bootstrap test of 1000 replications (Tamura et al., 2011).

Statistical analysis

Statistical analysis was performed using PASW Statistics 18.0. Means were compared by one-way analyses of variance (ANOVA), and post-hoc multiple comparisons were conducted using the least significant difference (LSD) method. The significance level was set at 0.05, and each experiment was repeated at least three times.

RESULTS

Clones and sequence analysis

The cloned ORF sequences of Bm-KPNA1, Bm-KPNA2, Bm-KPNA3, and Bm-KPNB1 were 1563, 1515, 1551, and 2661 bp long, respectively, and coded 520, 504, 516, and 886 AAs, respectively. The Bm-KPNA ORF sequences were identical to BGIBMGA011351-TA, BGIBMGA011351-TA, and BGIBMGA012556-TA. The Bm-KPNB1 ORF was 96.06% similar to BGIBMGA002972-TA, and its AA sequence similarity was 95.49%. BGIBMGA002972-TA was short of two intron-like regions when aligned with BM-KPNB1. One was 40 bp long, and followed a "T" inserted after the 25th nucleotide (Figure 1A); the other was 63 bp long (Figure 1B). The nucleotides of the two regions were integer multiples of three, so there were only 19 and 21 AAs difference (Figure 1C, D).



Figure 1. Nucleotide and amino acid sequences aligned between Bm-KPNB1 and BGIBMGA002972 on SilkDB; A and B show the nucleotide differences, C and D show the amino acid differences.

Genetics and Molecular Research 14 (4): 13049-13065 (2015)

The aligned ORFs and protein sequences of the Bm-KPNAs revealed that the similarities between Bm-KPNA1 and BM-KPNA3 (52.16 and 45.71%, respectively) and BM-KPNA2 (49.21 and 40.65%, respectively) were higher than those between Bm-KPNA2 and Bm-KPNA3 (47.76 and 38.34%, respectively). These results support the hypothesis that KPNA2 and KPNA3 developed from KPNA1 (Mason et al., 2009). Bm-KPNB1 had the lowest similarity with the KPNAs (Table 2).

Table 2. Similarity (%) of open reading frames and protein sequences between Bm-KPNAs and Bm-KPNB1.					
	Bm-KPNA1	Bm-KPNA2	Bm-KPNA3	Bm-KPNB1	
Bm-KPNA1	100/100		-	-	
Bm-KPNA2	49.21/40.65	100/100	-	-	
Bm-KPNA3	52.16/45.71	47.76/38.34	100/100	-	
Bm-KPNB1	24.46/8.24	23.32/8.13	23.90/10.15	100/100	

Fifty-two KPNAs, belonging to 20 animal species from nine orders of three phyla, were collected to construct the phylogenic tree. The insects had three KPNAs, except for *Drosophila melanogaster* and *Pararge aegeria*, which had four and two, respectively. These KPNAs were classified into three clades that represented different alpha subfamilies, and each Bm-KPNA was classified into one of the subfamilies. Therefore, we named Bm-KPNAs according to the subfamily they belonged to (Figure 2). BM-KPNA1 and BM-KPNA2 were most similar to *P. aegeria* (81.6 and 68.2%, respectively), while KPNA3 was most similar to *N. vitripennis* (77.5%). Twenty-one KPNBs, from seven orders of three phyla that were also aligned, were similar. BM-KPNB1 was most similar to *Danaus plexippus* (86.3%) (Figure 3).



Figure 2. Phylogenic tree of KPNAs constructed using the neighbor-joining algorithm method.

Genetics and Molecular Research 14 (4): 13049-13065 (2015)

Characterization of karyopherins in B. mori



Figure 3. Phylogenic tree of KPNBs constructed using the neighbor-joining algorithm method.

KPNA1 had 201 identical AAs (Figure 4), KPNA2 had 68 (Figure 5), KPNA3 had 95 (Figure 6), and KPNB had 316 (Figure 7), which accounted for 38.65, 13.49, 18.41, and 35.66%, respectively. These results suggest that KPNA1 and KPNB were under greater evolutionary pressure than were KPNA2 or KPNA3. These AAs must play important roles for proteins to perform their normal functions.



Figure 4. Multiple alignments between 17 KPNA1. 1, *Homo sapiens* KPNA1; 2, *H. sapiens* KPNA6; 3, *H. sapiens* KPNA5; 4, *Mus musculus* KPNA1; 5, *M. musculus* KPNA6; 6, *Pararge aegeria* KPNA1; 7, *Tribolium castaneum* gi|91077058; 8, *Nasonia vitripennis* gi|156551639; 9, *Megachile rotundata* gi|383855034; 10, *Bombus impatiens* gi|350420265; 11, *Bombus terrestris* gi|340713271; 12, *Apis florea* gi|380020809; 13, *Ceratitis capitata* gi|498967720; 14, *Culex quinquefasciatus* gi|170065607; 15, *Aedes aegypti* gi|157127698; 16, *Drosophila melanogaster* KPNA1; 17, *Bombyx mori* KPNA1.

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13055

Genetics and Molecular Research 14 (4): 13049-13065 (2015)

J. Li et al.

13056



Figure 5. Multiple alignments between 18 KPNA2. 1, *Mus musculus* Kpna2; 2, *M. musculus* Kpna7; 3, *Homo sapiens* KPNA2; 4, *H. sapiens* KPNA7; 5, *Apis florea* gi|380029460; 6, *Ceratitis capitata* gi|498971975; 7, *Aedes aegypti* gi|157134097; 8, *Drosophila melanogaster* KPNA2/pendulin; 9, *Caenorhabditis elegans* Ima2; 10, *Pararge aegeria* KPNA3; 11, *Acyrthosiphon pisum* gi|193610929; 12, *Pediculus humanus corporis* gi|242023280; 13, *Tribolium castaneum* gi|91077394; 14, *Nasonia vitripennis* gi|345488160/; 15, *Megachile rotundata* gi|383858926; 16, *Bombus impatiens* gi|350396236; 17, *Bombus terrestris* gi|340722259; 18, *Bombyx mori* KPNA2.

Genetics and Molecular Research 14 (4): 13049-13065 (2015)



Figure 6. Multiple alignments between 17 KPNA3. 1, *Mus musculus* KPNA3; 2, *M. musculus* KPNA4; 3, *Homo sapiens* KPNA4; 4, *H. sapiens* KPNA3; 5, *Caenorhabditis elegans* Ima-3; 6, *Tribolium castaneum* gi|189237585; 7, *Nasonia vitripennis* gi|156553340; 8, *Megachile rotundata* gi|383854227; 9, *Bombus impatiens* gi|350420887; 10, *Bombus terrestris* gi|340718064; 11, *Apis florea* gi|380013155; 12, *Ceratitis capitata* gi|498980272; 13, *Culex quinquefasciatus* gi|170029939; 14, *Aedes aegypti* gi|157129937; 15, *Drosophila melanogaster* KPNA3; 16, *D. melanogaster* KPNA4; 17, *Bombyx mori* KPNA3.

13057

Genetics and Molecular Research 14 (4): 13049-13065 (2015)

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Figure 7. Multiple alignments between 21 KPNB. 1, Danaus plexippus gi|357625986; 2, Dendroctonus ponderosae gi|546683722; 3, Harpegnathos saltator gi|307202414; 4, Solenopsis invicta gi|322796716; 5, Bombus impatiens gi|350427743; 6, Apis dorsata gi|572262812; 7, Apis mellifera gi|571567903; 8, Bombus terrestris gi|340723427; 9, Culex quinquefasciatus gi|170036594; 10, Aedes aegypti gi|157135220; 11, Megachile rotundata gi|383857795; 12, Cryptoramorphus floridanus gi|307188424; 13, Ceratitis capitata gi|498932442; 14, Anopheles darlingi gi|568255141; 15, Drosophila grimshawi gi|195030120; 16, Drosophila willistoni gi|195434026; 17, Daphnia pulex gi|321465952; 18, Mus musculus gi|30931411; 19, Xenopus laevis gi|168693593; 20, Homo sapiens KPNB; 21, Bombyx mori KPNB.

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13058

Genetics and Molecular Research 14 (4): 13049-13065 (2015)

By analyzing the protein sequences using Pfam, we found that the Bm-KPNAs all had one IBB domain, and Bm-KPNA1, 2, and 3 had 8, 4, and 7 armadillo/beta-catenin-like repeats (ARMs), respectively (Figures 4, 5, and 6). Bm-KPNB1 had 19 HEAT repeats; Heat_2, part of HEAT_1 and HEAT_3, constructed an importin-beta-N-terminal domain (IBB_N), and HEAT_7 to HEAT_19 constructed an IBB-binding domain (Figure 7).

Prokaryotic protein expression

Cells with recombinant BM-KPNAs or BM-KPNB1 plasmids induced by IPTG at 30°C could correctly express recombinant proteins. The molecular weight of recombinant Bm-KPNAs was about 60 kDa, and that of BM-KPNB1 was about 100 kDa, which is in accord with the molecular weights predicted by ExPASY (Figure 8).



Figure 8. Recombinant protein expression of BM-KPNAs and BM-KPNB1. **A** to **D** represent BM-KPNA1, 2, 3, and Bm-KPNB1 recombination proteins dyed with Coomassie brilliant blue R-250; **E** to **H** represent western blot results of BM-KPNA1, 2, 3, and Bm-KPNB1 recombinant proteins. *Lanes M, 1, 2, 3, 4, 5,* and *6* = protein markers, cells with blank expression vectors, or cells with gene targeting induced by 0, 0.3, 0.6, 0.9, or 1.2 mM isopropyl β -d-1-thiogalactopyranoside, respectively.

Genetics and Molecular Research 14 (4): 13049-13065 (2015)

Tissue expression analysis

We checked Bm-KPNA and Bm-KPNB1 mRNA levels in the fat body, silk glands, Malpighian tubules, hemocytes, midguts, testes, ovaries, and integuments of 5th 3-day-old larvae, and found that the four kinds of protein were expressed in all eight tissue types, but the expression levels varied (Figure 9). The highest expression of all the proteins occurred in the testes. BM-KPNA2 was expressed approximately twice as much in the testes as were the other proteins, and its expression was 846.2-fold higher than that in the silk gland.



Figure 9. Relative mRNA expression levels of Bm-KPNAs and Bm-KPNB1. The 18s rRNA levels were defined as 1. Different letters represent significant differences (P < 0.05).

Relative mRNA levels in response to microbial injections

The relative mRNA levels of the Bm-KPNAs and Bm-KPNB1 in response to the microbes differed. In the fat body, Bm-KPNA mRNA levels were generally low at 1 and 8 h, but increased at 4 and 12 h. Bm-KPNA1 was most sensitive to *E. coli*, *B. bassiana*, and NPV, Bm-KPNA3 to *M. luteus*, *E. coli*, and *B. bassiana*, and Bm-KPNA2 was not as sensitive as the other two Bm-KPNAs to the four microbes. Bm-KPNB1 mRNA levels in response to *E.coli* and *M. luteus* exhibited no significant variation at 1, 4 or 8 h, but at 12 h they increased by 2.97-fold and 4.93-fold, respectively. When responding to *B. bassiana* and NPV, Bm-KPNB1 mRNA levels slightly fluctuated. In hemocytes, Bm-KPNA and Bm-KPNB1 mRNA levels were first downregulated and then upregulated, before decreasing again at 4 h to basal levels. They were most sensitive to *B. bassiana*, and were upregulated to 15.54, 19.32, 8.09, and 3.93-fold compared with control groups, respectively, at 8 h (Figure 10).

DISCUSSION

SilkDB predicted three Bm-KPNA ORFs and one Bm-KPNB1 ORF. We designed primers to amplify the four protein ORFs, and the Bm-KPNA sequencing results matched the sequences on SilkDB. The Bm-KPNB1 ORF was shorter (102 bp) than the sequence on SilkDB. Bm-KPNB1 probably has variable expression modes that code for a long and a short peptide. We designed PCR primers to amplify the two different regions, and tested them on 15% polyacrylamide gel; after testing 30 silkworms, we did not find the short Bm-KPNB1 (Figure 11).

Genetics and Molecular Research 14 (4): 13049-13065 (2015)



Figure 10. Relative mRNA expression levels of Bm-KPNAs and Bm-KPNB1 in response to four kinds of microbe. **A** to **D** represent responses in the fat body, **E** to **H** represent responses in the hemocyte. At each time point, the corresponding mRNA expression levels of larvae injected with PBS were defined as 1. Different letters represent significant differences (P < 0.05).

Continued on next page

Genetics and Molecular Research 14 (4): 13049-13065 (2015)





Genetics and Molecular Research 14 (4): 13049-13065 (2015)

Characterization of karyopherins in *B. mori*



Figure 11. DNA native polyacrylamide gel electrophoresis (Native-PAGE) of Bm-KPNB1. M represents a DNA marker. 1 to 9 represent cDNA templates from different larvae; each cDNA template was a cDNA pool from the sex glands, Malpighian tubules, midguts, silk glands, fat bodies, hemocytes, and integuments from 5th 3-day-old larvae. **A.** amplified from 470 to 893 bp; **B.** amplified from 1879 to 2550 bp of the Bm-KPNB1 open reading frame. **A.** forward primer 5'-TGGAGGCTATTGGCTACATT-3' and reverse primer 5'-TCTATTTCCTCATCGCTCACA-3'; **B.** forward primer 5'-GCAGGCAGTGTTCAGGAGGA-3' and reverse primer 5'-GTCGGTGAGCGGCTTGGTGT-3'. The annealing temperature of the two pairs of primers was 60°C.

Animal KPNAs are divided into three conserved subfamilies, named α 1, α 2, and α 3 (Köhler et al., 1997; Mason et al., 2009). The KPNAs of the domestic silkworm are also classed into three conserved subfamilies, and we named them karyopherin α 1 (BM-KPNA1), karyopherin α 2 (BM-KPNA2), and karyopherin α 3 (BM-KPNA3). Humans contain seven KPNAs, and KPNA1, 5, and 6 are in the α 1 subfamily, KPNA2 and 7 are in the α 2 subfamily, and KPNA3 and 4 are in the α 3 subfamily (Köhler et al., 1997; Mason et al., 2009; Kelley et al., 2010; Yang et al., 2010). The phylogenic tree we constructed confirmed this; however, *P. aegeria* KPNA3 was a special case, for it is named KPNA3 but is in the α 2 clade. To date, only three KPNAs have been found in insects, but *Drosophila* has an additional Dmel-KPNA4 (FlyBase, FBpp0076777). We found that Dmel-KPNA4 has not an obvious IBB domain, but other Dmel-KPNAs all have one IBB domain.

KPNAs and KPNBs in different organisms have conserved domains, which can identify them from other proteins. KPNAs have IBB and ARM repeats, and KPNBs have IBB_N, IBB-binding domain, and HEAT repeats. ARMs and HEATs are approximately 50 AA-long tandem repeated sequence motifs that probably have a common phylogenetic origin (Andrade et al., 2001). Several ARMs and HEATs construct protein-protein interaction domains, but no common features among the target proteins could be found. The domains constructed by ARMs in KPNAs recognize NLSs, and those constructed by HEATs in KPNBs recognize the IBB of KPNA and other proteins to be transported into the nucleus (Cingolani et al., 1999; Lee et al., 2003; Mynott et al., 2011). Many nuclear pore proteins contain FG sequence repeats that can bind to HEAT repeats within KNPBs,

Genetics and Molecular Research 14 (4): 13049-13065 (2015)

which is important for KNPB -mediated transport (Vetter et al., 1999; Bayliss et al., 2002; Isgro and Schulten, 2007; Lange et al., 2007). The IBB_N of KPNBs overlaps with HEATS, and interacts with Ran-GTP and Ran-GDP, which is important for cargo-KPNA-KPNB complex disassembly and KPNB relocation to the cytoplasm (Vetter et al., 1999). Similar to other KPNAs, Bm-KPNAs all have one IBB and several ARMs. Bm-KPNB1 has one IBB_N and 19 HEATs.

Silkworms lack adaptive immune systems, and possess innate immunity that only responds to common bacterial, fungal, and viral pathogens. Their responses involve cellular mechanisms, such as programmed cell death or autophagy, and the expression of a large set of genes. Our results show that Bm-KPNA and Bm-KPNB1 mRNA expression levels vary according to the pathogen, since *E. coli* is a Gram negative bacteria, *M. luteus* is a Gram positive bacteria, *B. bassiana* is a fungus, and NPV is a virus.

KNPAs and KNPBs are expressed a great deal in silkworm larvae, indicating that they play important roles in basic physiological processes. The fact that expression levels differed between tissues suggests that they probably play different roles in different tissues (Kimura and Imamoto, 2014). The probable reason that Bm-KPNA2 mRNA expression levels were high in the testes is that this organ is highly active, and KPNA2 interacts with proteins that are associated with mRNA processing, ribonucleoprotein complex biogenesis, chromatin modification, and transcription (Huang et al., 2013; Umegaki-Arao et al., 2013). Since the karyopherin family transports cargoes through the NE, and some cargoes are probably gene expression regulation factors, it is important to ascertain what cargoes they transport in different tissues.

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Genetics and Molecular Research 14 (4): 13049-13065 (2015)

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