Molecular cloning and analysis of four cDNAs from the heads of *Apis cerana cerana* nurse honeybees coding for major royal jelly proteins

Songkun Su\(^a\), Stefan Albert\(^b\), Shenglu Chen\(^a\)*, Boxiong Zhong\(^a\)

\(^a\) Laboratory of Apicultural Research, Animal Science College, Zhejiang University Hangzhou 310029, China  
\(^b\) Institute of Medical Radiation and Cell Research, University of Würzburg, Versbacherstrasse 5, 97078 Würzburg, Germany

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Abstract – A cDNA library was constructed from 8-day-old worker heads of *Apis cerana cerana*. A DIG-labeled probe derived from part of an *Apis cerana* mrjp3 genomic segment was used to screen the library. One hundred and twenty positive clones were identified and characterized. Thirty one clones were homologous with major royal jelly proteins (MRJPs) of *Apis mellifera*. The most abundant MRJP homologue was MRJP1 (11 clones), followed by MRJP3 (10 clones), MRJP2 (7 clones) and MRJP5 (3 clones). Clones containing *A. cerana cerana* MRJP1, MRJP2, MRJP3 and MRJP5 cDNAs were identified, completely sequenced, and analyzed with bioinformatics software. Several lines of evidence suggested that the identified cDNAs code for major royal jelly proteins of *A. cerana*. In addition to polymorphic regions of MRJP3 and MRJP5, another polymorphic repetitive region was found in AccMRJP2. The polymorphism of AcMRJP2 and AcMRJP5 repeat regions were tested by PCR with genomic DNAs of individual honeybees. Different properties of the repetitive regions of MRJP2 genes in two closely related *Apis* species were discussed.

*Apis cerana cerana* / royal jelly protein / polymorphism / VNTR / midisatellite / *Apis mellifera*

1. INTRODUCTION

Royal jelly (RJ), a secretion of both the hypopharyngeal and mandibular glands of nurse workers, is believed to play a central role in honeybee queen development (Knecht and Kaatz, 1990; Lensky and Rakover, 1983; Moritz and Southwick, 1992). Proteins are an important component of RJ, forming about 50% of the dry mass (Rembold, 1987). Five major proteins of *Apis mellifera* RJ, called MRJP1-5, form a significant part (~90%) of RJ total protein (Schmitzova et al., 1998). The cDNA sequences of MRJP1-MRJP5 were isolated from a *A. mellifera* cDNA library of nurse honeybee heads (Albert et al., 1999a; Klaudiny et al., 1994; Schmitzova et al., 1998). Corresponding MRJP1-5 proteins were identified in RJ separated by 2-D electrophoresis (Sano et al., 2004). Recently, three cDNAs coding for new but less abundant members of MRJP protein family were identified from a *A. mellifera* brain EST library (Albert and Klaudiny, 2004)

Biological activities of *A. mellifera* major royal jelly proteins have been studied in various systems. The protein fractions of royal jelly were confirmed to possess a high antioxidative activity and scavenging ability against reactive oxygen species (Nagai and Inoue, 2004). MRJP1 enhanced cell proliferation of rat hepatocytes (Kamakura et al., 2001b), stimulated the growth of human lymphocytes in a serum-free medium...
MRJP3 exhibited potent immunoregulatory effects in vitro and in vivo (Okamoto et al., 2003).

In contrast to *A. mellifera*, information on RJ of other honeybee species (including *Apis cerana*) is scarce. Takenaka and Takenaka (1996) reported that chemical composition, i.e., proteins, 10-hydroxydecenoic acid, and glucose/fructose ratio, differed between *A. mellifera* and *A. cerana* royal jelly. Recently an EST library was prepared from the hypopharyngeal glands of *A. cerana indica* (Srisuparbh et al., 2003). From this library, the MRJP1 homologue and apisimin cDNAs of *A. cerana indica* were isolated and sequenced. The MRJP1 of *A. cerana indica* showed 93% and 90% homology with MRJP1 of *A. mellifera* at the nucleotide and amino acid levels, respectively.

There are about two million *A. cerana cerana* colonies in China (Chen et al., 2002). *A. cerana* is widely used for commercial beekeeping in mountain areas of South China, primarily due to its resistance against diseases, wasps and bee mites.

In this report, we constructed a cDNA library from 8-day-old nurse honeybee heads of *A. cerana cerana*, screened it by hybridization and identified cDNAs encoding MRJP1, MRJP2, MRJP3, and MRJP5 homologues. We provided full cDNA sequences of these genes and partial genomic sequences of *A. cerana cerana* MRJP1, 5, and 7. The sequences were compared with the MRJPs of *A. mellifera* and several lines of evidence showed that isolated cDNAs encode functional homologues of MRJPs in *A. cerana*. Moreover, we identified and characterized an unexpected polymorphism of MRJP2 in *A. cerana*.

### 2. MATERIALS AND METHODS

#### 2.1. Biological samples

Nurse honey bees (*Apis cerana cerana*) were obtained from the Laboratory of Apicultural Research (Huajiachi campus, Zhejiang University, Hangzhou, China) as follows: newly emerged workers (less than 1-day-old) were marked with paint mark pen and put back into the colonies, to enable trophallactic contacts with other bees and normal development. Eight-day-old nurse bees were collected and anesthetized on ice, the heads were removed, frozen in liquid nitrogen and stored at –80 °C.

European honeybees of two putative races were obtained from colonies separated by several hundred kilometers, *Apis mellifera mellifera* from Göttingen, Germany, *Apis mellifera carnica* from Bratislava, Slovakia and Würzburg, Germany.

#### 2.2. Construction of a cDNA library from 8-day-old nurse bee heads

Total RNA from the heads of nurse honeybees was prepared using the TRIZOL Kit (Promega) according to manufacturer’s instructions. mRNA was extracted using PolyATtract TM (Promega) mRNA method. 5 μg of purified mRNA were reverse-transcribed into cDNA and Lambda-ZAP phage library was constructed using ZAP-cDNA Synthesis Kit and ZAP-cDNA Gigapack III Gold Cloning Kit (Stratagene) following the respective manuals (www.stratagene.com).

#### 2.3. PCR amplification of MRJP genomic fragments and cDNA library screening

Two primers termed P212 and P218 designed to prime within regions conserved among five MRJPs of *A. mellifera* (P212: AAA GT(G/A) T(T/G)G GAA G(T/A)C AAT CGA TG; P218: TGC CT(T/C) GG(C/T) ATA G(C/T)T TGT C) were used for PCR amplification of genomic DNAs of *A. cerana*. Reaction mixtures included commercial PCR buffer, containing 1 μM primers, 1 μg *A. cerana* genomic DNA and polymerase mixture (Taq:DeepVent = 5:1; AmpliTaq, Perkin Elmer and NEB, respectively). After initial heating at 94 °C and polymerases addition at 80 °C, 32 cycles of 30 s at 94 °C, 60 s at 52 °C and 120 s at 72 °C were run. Amplified products were cloned into PCR2.1 vector (Invitrogen).

Studies of the polymorphic MRJP2 and MRJP5 alleles were done as above with the following primers: P28 (TTA ATG AGA AAT ACT CAT TGC G) and P29 (AAC GAC GAA CTT GAT TAT CAT TC) for MRJP2. The primers P126 (AGA CTC TTC AAA CGG TCG TTG) and P127 (CTG TAA TTT CAT ACT TAA AGC CAT C) were designed to amplify the DRM repetitive region of MRJP5.

The amplified cDNA library was screened by hybridization using standard protocols. We used an mrjp3 gene fragment amplified from *A. cerana* genome as a hybridization probe employing DIG High Prime DNA Labeling and Detection Starter Kit I (Roche, Germany).
2.4. Primary characterization of positive clones

The phage extracts from the positive clones were transferred from plate into test tubes with 200 µL SM buffer. To determine the length of cDNA fragments inserted and check the purity of isolated clones, we amplified the cDNA inserts with T3 (5’-AAT TAA CCC TCA CTA AAG GG) and T7 (5’-TAA TAC GAC TCA CTA TAG GG) primers. Polymerization reactions were done as above in 50 µL volume with 5 µL positive clone extract (Lambda-ZAP, Stratagene).

2.5. DNA sequencing and bioinformatics analyzes

Plasmid DNAs were sequenced by the cycle sequencing method using the Prism Ready Reaction Dyedeoxy Terminator kit on an ABI PRISM 377 DNA Sequencer according to the manufacturer’s instructions. Obtained sequences were compared with GenBank using the BLASTN and BLASTX programs (http://www.ncbi.nlm.nih.gov).

DNA sequences were assembled with the help of DNATOOLs and DNASTAR program packages. To distinguish between proteins of different origin, the MRJPs of *Apis mellifera*, *Apis cerana indica*, *Apis cerana cerana* were termed AcMRJPs, AciMRJPs and AccMRJPs, respectively.

The cDNAs identified in this work were deposited in GenBank under following accession numbers:
- **AccMRJP1** (1 ×), **AccMRJP7** (1 ×), **AccMRJP3** (2 ×), and **AccMRJP5** (1 ×). We termed them AcMRJP1, AcMRJP7, AcMRJP3 and AcMRJP5, respectively.

We cloned the PCR products and sequenced five individual clones. Each of the clones contained an *mrjp*-like insert; two of them were identical. Cloned inserts showed homology to *mrjp1* (1 ×), *mrjp7* (1 ×), *mrjp3* (2 ×), and *mrjp5* (1 ×). We termed them AcMRJP1, AcMRJP7, AcMRJP3 and AcMRJP5, respectively.

The significantly larger size of the amplified genomic fragments (1350–1500 bp versus expected size of cDNA fragments ~ 680 bp) indicated the presence of introns, which was confirmed by DNA sequencing. Three introns located at the same positions were found in all amplified genomic fragments (see Figs. 2, 3).

3. RESULTS

3.1. Generation of MRJP genomic fragments of *Apis cerana*

To obtain initial information about *mrjp* genes of *A. cerana* and to prepare hybridization probes for screening a cDNA library, we looked for regions that were conserved among known *mrjp* cDNAs of *A. mellifera*. The rationale was that the regions conserved among *mrjps* of *A. mellifera* would also be conserved in other species. The cDNAs encoding *mrjp1–5* of *A. mellifera* were aligned to identify the regions of high conservancy. The primers P212 and P218 (see Materials and Methods), each with two degenerated positions were designed to anneal to these conserved regions. The PCR reaction was run at a relatively low annealing temperature (52 °C) using a mixture of proof-reading polymerase and Taq polymerase. A broad band of 1350–1500 bp was amplified, which might be a mixture of several products of similar size.

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3.2. Screening of the cDNA library of *Apis cerana cerana*

A cDNA library containing about 90% recombinant clones was prepared from the heads of 8-day-old nurse bees. More than 200 positive clones were found from the cDNA library with a DIG-labeled Ac*mrjp3* genomic fragment as a probe. One hundred and twenty of these clones were extracted into SM buffer and subjected to PCR with T3/T7 primers. Fifty five of them showed single cDNA inserts that were 1200–2500 bp in size. The PCR products amplified from these clones were sequenced. Thirty one clones were homologous to *mrjps* of *A. mellifera*. The most abundant *mrjp* homologues were *mrjp1* (11 clones) followed by *mrjp3* (10 clones), *mrjp2* (7 clones) and *mrjp5* (3 clones). The complete cDNA sequences of AcMRJP1, 2, 3, and 5 were obtained by sequencing of the clones #103, #46, #50 and #91 respectively.

3.3. Characterization of sequences encoding MRJPs of *Apis cerana cerana*

AccMRJP1 cDNA contained an open reading frame (ORF) of 1445 nucleotides (poly[A])
tail not included) encoding a protein of 433 amino acids, which was highly similar (99.5% and 90.5% identity) to its homologues from *A. cerana indica* (Srisuparbh et al., 2003) and *A. mellifera* (Schmitzova et al., 1998) at the protein level. According to N-terminal sequencing of the *A. cerana* RJ protein (Srisuparbh et al., 2003), the cleavage site for signal peptidase was localized between Ser30 and Ser31. Three potential N-linked glycosylation sites were found at amino acids 29, 145, and 178. The polyadenylation signal AATAAA was located 14 bp upstream of the poly(A) tail.

The sequence of AccMRJP2 cDNA was 1590 bp long and included an open reading frame (ORF) of 1404 nucleotides encoding a protein of 468 amino acids (see Fig. 1). The sequence of the encoded protein contained all three internal peptide sequences (Fig. 1, underlined) determined by sequencing of the protein from RJ of *A. cerana indica* (Srisuparbh et al., 2003). The putative signal peptidase cleavage site was between Gly17 and Ala18. Two potential N-linked glycosylation sites were found at asparagines 145 and 178. Interestingly, the repetitive region similar to that of MRJP3 (see below), consisting of 9 copies of NQKNN pentapeptide, was found at the C-terminal part of the deduced AccMRJP2 protein.

AccMRJP3 cDNA was 1977 bp long. The cDNA and inferred amino acid sequences were shown in Figure 2. The cDNA sequence contained an ORF (nucleotides 46–1824), which encoded a polypeptide of 593 amino acid residues. The deduced amino acid sequence of mature peptide began with AAVNHQRKSA (Schmitzova et al., 1998). Prompted by this observation, we set out to study the polymorphism of the MRJP2 repeat in detail. Two primers (P28 and P29; see Fig. 1, arrows) were designed to flank the repetitive region of MRJP2 and used in PCR with the genomic DNAs of *A. cerana* and *A. mellifera* individuals collected from geographically distant colonies. Figure 4A shows that there was a high variability of the repeat size in *A. cerana* even within individuals from the same colony, whereas the analogous region of *A. mellifera* showed the same size among all individuals originating from distant locations. For studying the polymorphism of the DRM repeat of MRJP5, PCR primers P126 and P127 (Materials and Methods) were designed to amplify this region. Similar to MRJP2, a clear size polymorphism with allele sizes ranging from ~560–650 bp was observed in *A. cerana* (Fig. 4B). MRJP5 of *A. mellifera* was also polymorphic (our unpublished data and Fig. 4B), but observed size differences among individual alleles were smaller. Additional bands of the intermediate size presumably representing

### 3.4. Studies of the polymorphic repeats of AccMRJP2 and AccMRJP5

The sequences of MRJP2 and MRJP5 cDNAs isolated from another *A. cerana* race, *A. cerana indica*, were deposited in GenBank by the group of S. Sittipraneed, Chulalongkorn University, Thailand (accession numbers [AF525777](https://www.ncbi.nlm.nih.gov/nuccore/AF525777) and [AY532369](https://www.ncbi.nlm.nih.gov/nuccore/AY532369) respectively). They were nearly identical with our cDNAs, except for their repetitive regions. For example, AccMRJP2 contained 9 copies of the reiterated pentapeptide unit, whereas Thailand isolation contained only 8 of them. This was surprising because no polymorphism has been reported for MRJP2 of *A. mellifera* (Schmitzova et al., 1998).
Figure 1. The sequence of AccMRJP2 cDNA and inferred protein. The vertical arrow points to deduced signal peptidase cleavage site. Consensus N-glycosylation sites are enclosed in boxes. The underlined are the peptide sequences determined by sequencing A2 protein of *A. cerana* royal jelly (Srisuparbh et al., 2003). Shorter regularly spaced lines highlight the repetitive units of the pentapeptide repeat region. Horizontal arrows show the positions of the PCR primers used for amplification of the polymorphic pentapeptide repeat region (see Fig. 4A). Triple dots indicate the continuation of the underlined feature in the following row.
heteroduplex DNAs (Kaiser et al., 2002) were observed in both PCR reactions. This phenomenon was more pronounced in MRJP5 products, possibly due to the different nature of the repetitive region or for other reasons that have not been investigated.

Figure 2. The sequence of AccMRJP3 cDNA and inferred protein. The vertical arrow points to signal peptidase cleavage site. N-glycosylation sites are boxed. The underlined is the N-terminal peptide sequence of A1 protein of A. cerana royal jelly (Srisuparbh et al., 2003). Shorter regularly spaced lines highlight the repetitive units of the pentapeptide repeat region. Vertical triangles show the positions of introns found in the genomic sequence.
Major royal jelly proteins of *Apis cerana*

At the beginning of this project, no proteins of RJ or genes encoding them of honeybees other than *A. mellifera* were known. Therefore we sought cDNAs encoding MRJPs in another economically important honeybee species, *Apis cerana*.

3.5. Differences among MRJPs of *Apis cerana* and *Apis mellifera*

Only 6 nucleotides and 2 amino acid residues differed between MRJP1 cDNAs of *A. cerana cerana* and *A. cerana indica* published in the meantime (Srisupabr et al., 2003). AccMRJP1 showed high homology to AmMRJP1 at both nucleotide (93.8%) and protein (90.5%) level. Taken together, MRJP1, the most abundant protein of RJ, was highly conserved among honeybee species and subspecies.

Figure 3. The sequence of AccMRJP5 cDNA and inferred protein. The vertical arrow points to putative signal peptidase cleavage site. Consensus N-glycosylation sites are enclosed in squares.
MRJP2 cDNAs were also nearly identical between A. cerana cerana and A. cerana indica (GenBank entry AF525777) except for the repetitive region (see above). The AccMRJP2 was also highly homologous with AmMRJP2 at both the nucleotide (89.2%) and protein (84.5%) levels.

The nearest homologue of AccMRJP3 was MRJP3 of A. mellifera. AmMRJP3 was shorter than AccMRJP3 due to the different length of..
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the repetitive region. As described for AmMRJP3 (Albert et al., 1999b), the repeat of AcMRJP3 was also highly polymorphic among individuals of *A. cerana* and other honeybees. (Albertova et al., unpublished data).

AcMRJP4 (Acc. Number **AY532368**) showed homology to its *A. mellifera* counterpart (90.0% and 80.0%). The most obvious difference between AcMRJP4 and AmMRJP4 was the length of encoded proteins. AcMRJP4 comprised 485 residues while AmMRJP4 only 464 residues. Again, the different length of the pentapeptide repetitive region between species caused the size difference.

The sequence identity of nucleotides and amino acid residues of MRJP5 was 99.3% and 97.2% between *A. cerana cerana* (Acc. Number **AY392757**) and *A. cerana indica* (Acc. Number **AY532369**). AccMRJP5 was homologous to MRJP5 of *A. mellifera* at nucleotide (90.7%) and protein level (83.6%), respectively. The DRM repetitive region (amino acids: 420–530) was highly variable between honeybee species and alleles of different size were also found among individuals (see Fig. 4B).

A phylogenetic tree was calculated from the aligned MRJP protein sequences of *A. cerana* (Fig. 5) and *A. mellifera* by neighbor-joining method using exhaustive maximum parsimony search with distantly related Yellow-f protein of *A. mellifera* (Albert and Klaudiny, 2004) as an outgroup (not shown). Each of the AcMRJPs formed a monophyletic group with its *A. mellifera* homologue. High bootstrap values supported the groups formed by *A. mellifera*-*A. cerana* pairs, further supporting the notion that the MRJPs of *A. cerana* isolated here were orthologs of the corresponding *A. mellifera* proteins.

4. DISCUSSION

4.1. The newly isolated cDNAs code for proteins of *A. cerana* royal jelly

Using PCR approach with primers priming to conserved regions of *A. mellifera* MRJPs, we have cloned and characterized fragments of four MRJP genes of *A. cerana*. Using one of the four genomic fragments (MRJP3) as a hybridization probe, we isolated complete cDNAs of AccMRJP1, 2, 3 and 5. On the other hand, screening of 120 positive clones was not sufficient for isolation of less abundant MRJP cDNAs, such as MRJP4, 6–8 (Albert and Klaudiny, 2004) from the cDNA library. Several lines of evidence, such as features of encoded proteins, homology with AmMRJPs, relative abundance of the isolated cDNAs in the cDNA library, and finally the presence of the peptide sequences proteins of *A. cerana* RJ (Srisuparbh et al., 2003) supported the view that the cDNAs presented here code for protein components of *A. cerana* RJ.

4.2. Are MRJP genes a heaven for polymorphic repetitive segments?

Phylogenetic analysis suggested that MRJP proteins evolved by subsequent but nearly simultaneous duplications (Albert et al., 1999a). Several MRJPs contained a pentapeptide repetitive region or its leftovers, which is extremely extended and highly polymorphic in MRJP3 (Albert et al., 1999b; Albert and Schmitz, 2002). Here we show that the same repetitive region is highly polymorphic in AcMRJP2. It was argued that in other MRJPs, the repeat units accumulated mutations which disabled their further rearrangements by a slipped strand mechanism. Three of five MRJP genes/proteins of *A. cerana* characterized so far (AcMRJP2, AcMRJP3, AcMRJP5) exhibit a repeat length polymorphism (Fig. 4 and unpublished data). The distribution of polymorphic alleles seems to be phenotypically neutral as there was no repeat length bias found in the population. Thus, the family of MRJP genes alone provides three midisatellite polymorphic loci of VNTR (variable number of tandem repeat) type, which are suitable for genotyping *A. cerana* individuals (Beye et al., 1998).

Of the remaining two known MRJPs of *A. cerana*, AcMRJP1 does not harbor any repetitive region. In line with this observation, the two AcMRJP1 cDNAs originating from two different *A. cerana* races (this work and Srisuparbh et al., 2003) are nearly identical and highly homologous with AmMRJP1. Another member of the family, AcMRJP4, although not tested yet, might also be polymorphic. At least in comparison to AmMRJP4, there is an apparent segment duplication seen in the repetitive region of AcMRJP4.
Figure 5. Alignment of AccMRJP proteins. The sequences were aligned using CLUSTAL W and visualized using BOXSHADE. Black-shaded residues are those identical between at least two proteins, grey-shaded indicate the conservative substitutions.
The advantage of using MRJP polymorphism for genotyping lies in easy detection of the polymorphic alleles by PCR and subsequent electrophoresis in standard agarose gels (Fig. 4). In addition, the polymorphism of MRJP genes affects the molecular mass of encoded proteins, although the resolution at the protein level is lower (Albert et al., 1999b). Therefore a precise analysis of A. cerana and towards length fixation and accumulation of mutations in A. mellifera. In line with the above statement, the basic units of the MRJP2 pentapeptide repeat region differ from each other in A. mellifera but are identical in A. cerana (Fig. 4A, alignments beneath the gel picture, black-shaded regions).

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Résumé – Clonage moléculaire et analyse de quatre ADNc issus de têtes de nourrices d’Apis cerana cerana codant pour les principales protéines de la gelée royale. La gelée royale (GR) est un composant crucial de la nutrition de l’abeille (Apis sp.). Elle est synthétisée par les glandes hypopharyngiennes des nourrices et cette sécrétion sert à nourrir la reine et les larves. Les principales protéines de la GR sont très proches les unes des autres et éloignées des protéines trouvées chez les autres insectes qui sont impliquées dans la pigmentation de la cuticule et dans d’autres processus physiologiques. Bien que les principales protéines de la gelée royale (MRJP) de l’abeille domestique (Apis mellifera) soient bien caractérisées sur le plan génétique et de leurs séquences protéiques, on sait peu de choses concernant les MRJP des autres espèces du genre Apis. Nous avons caractérisé les ADNc complémentaires (ADNC) et des portions des séquences génomiques qui codent pour les protéines de la GR d’Apis cerana cerana par le clonage, le séquençage et la PCR. Une bibliothèque d’ADNC a été construite à partir de nourrices d’A. c. cerana âgées de 8 j. Elle a été testée par hybridation à l’aide de protocoles standard, ce qui a permis d’identifier quatre ADNC codant pour les protéines d’A. c. cerana. Ces protéines ont été nommées AccMRJP1, AccMRJP2 (Fig. 1), AccMRJP3 (Fig. 2) et AccMRJP5 (Fig. 3), en fonction de leur ressemblance avec les protéines de la GR d’A. mellifera. Les protéines de la GR d’A. cerana caractérisées ici présentent une forte ressemblance avec les protéines respectives d’A. mellifera, y compris la présence de séquences répétitives chez certaines d’entre elles (Fig. 5). La longueur des séquences répétitives diffère selon l’espèce, mais aussi entre individus d’A. cerana en raison du nombre d’unités de répétition (Fig. 4). Nous avons pu montrer par la PCR et le séquençage de l’ADN que la région répétitive de MRJP2 avait évolué différemment après la scission des espèces mellifera et...
Apis cerana cerana / Apis mellifera / gelée royale / protéine / polymorphisme / midisatellite / VNTR


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