Antimicrobial Peptide Evolution in the Asiatic Honey Bee
Apis cerana

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Abstract
The Asiatic honeybee, Apis cerana Fabricius, is an important honeybee species in Asian countries. It is still found in the wild, but is also one of the few bee species that can be domesticated. It has acquired some genetic advantages and significantly different biological characteristics compared with other Apis species. However, it has been less studied, and over the past two decades, has become a threatened species in China. We designed primers for the sequences of the four antimicrobial peptide cDNA gene families (abaecin, defensin, apidaecin, and hymenoptaecin) of the Western honeybee, Apis mellifera L. and identified all the antimicrobial peptide cDNA genes in the Asiatic honeybee for the first time. All the sequences were amplified by reverse transcriptase-polymerase chain reaction (RT-PCR). In all, 29 different defensin cDNA genes coding 7 different defensin peptides, 11 different abaecin cDNA genes coding 2 different abaecin peptides, 13 different apidaecin cDNA genes coding 4 apidaecin peptides and 34 different hymenoptaecin cDNA genes coding 13 different hymenoptaecin peptides were cloned and identified from the Asiatic honeybee adult workers. Detailed comparison of these four antimicrobial peptide gene families with those of the Western honeybee revealed that there are many similarities in the quantity and amino acid components of peptides in the abaecin, defensin and apidaecin families, while many more hymenoptaecin peptides are found in the Asiatic honeybee than those in the Western honeybee (13 versus 1). The results indicated that the Asiatic honeybee adult generated more variable antimicrobial peptides, especially hymenoptaecin peptides than the Western honeybee when stimulated by pathogens or injury. This suggests that, compared to the Western honeybee that has a longer history of domestication, selection on the Asiatic honeybee has favored the generation of more variable antimicrobial peptides as protection against pathogens.

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Introduction
The honeybee is a eusocial insect whose role in plant pollination is essential to global ecology. Through pollination of flowering plants, honeybees help maintain biodiversity and supply commodities such as honey, royal jelly, propolis, pollens, and wax. Publication of the genome sequence of the Western honeybee, A. mellifera L. facilitates genomic research on this economically important insect [1]. The discovery of a 100-million-year-old amber fossil pushes back the date that bees began pollinating flowers by millions of years [2], although species of Apis are much younger than this. Honeybees defend their nest, brood and stored food through many complicated and efficient defensive behaviors [3], but populations have decreased rapidly in many countries in recent years. Explanations for this decline include global warming caused by the greenhouse effect, indiscriminate use of pesticides, reduction or extinction of plant species and other human-induced disturbances. In addition, infection by strong pathogenic microorganisms (such as viruses) or pests (such as mites) have been cited as important direct or indirect reasons for honeybee collapse [4–9]. Protection of the honeybee species is a worldwide concern.

The Asiatic honeybee, A. cerana Fabricius, plays an important role in maintaining the biodiversity of plants in China. A. cerana is still found in the wild, where it nests in tree holes, fallen logs, and crevices, but it is also one of the few bee species that can be domesticated. It has acquired some incomparable advantages in its long history of evolution compared with other introduced Apis species. It has adapted to adverse climatic conditions, and can survive extreme fluctuations in temperature and long periods of rainfall. For example, it can survive when the air temperature is as low as ~0.1°C, a temperature lethal for the Western honeybee [10]. It is a natural host for, and can tolerate the mite, Varroa destructor, and the microsporidian, Nosema ceranae, both of which are the serious pests of the Western honeybee [11,12]. Having coevolved with these parasites, the Asiatic honeybee exhibits more careful grooming behavior than the Western honeybee, and appears to have other more effective defenses against these parasites. The Asiatic honeybee can also effectively pollinate mountain plants and crops, especially the early flowering fruits and vegetables in high altitude regions, such as the Himalayan region of China where temperatures are too low for the exotic A. mellifera [10]. Nevertheless, over the last 20 years, it has become a threatened species in China for many reasons, most likely because of competition from the introduced Western honeybee, A. mellifera. More research is needed and urgent measures must be taken to protect the Asiatic honeybee. It is noteworthy that the Asiatic
honeybee does not produce propolis, a highly valued resinous material that is used to seal small open spaces in the hive, and is believed to be involved in the honeybee defense system against pathogens in the hive.

Honeybees defend against many pathogens by producing antibiotic substances such as propolis and royal jelly [13,14]. The innate immune system is the first line of defense against pathogens in plants and invertebrate animals, and it is also critical for vertebrate immunity before the acquired immune system generates a specific response [15–17]. Various antimicrobial peptides are the key elements of the insect immune system [18–20]. After the honeybees are infected by pathogens, four antimicrobial peptide families are synthesized, representing a broad spectrum of antimicrobial activity in the haemolymph. All of these are cationic peptides identified as: apidaecins [21], abaecin [22], hymenoptaecin [23] and defensin [24]. Recently, two structurally different defensin genes were cloned from A. mellifera [25]. Almost all of the honeybee antimicrobial peptides and the antimicrobial peptide genes are reported from the Western honeybee; however, little research has been conducted on the Asiatic honeybee. As an ancient and important native honeybee species in China, many aspects of the species need to be explored. Here we examine the antimicrobial peptide gene families of the Asiatic honeybee and compare them with those of the Western honeybee.

Results

Analysis of defensin cDNA genes

In total, 29 different defensin cDNA sequences (Defensin1~29, GenBank: EU727268~EU727296) coding 7 different defensin peptides (termed as AcDe1~AcDe7) were amplified and identified. All of these cDNA genes have high sequence identity with the defensin cDNA gene of the Western honeybee. Nearly two-thirds (20/29) of the cDNA genes coded the major peptide AcDe4, and one or two cDNA genes coded for defensin peptides AcDe1~AcDe3 and AcDe5~AcDe7. The major peptide AcDe4 was identical to peptide AcDe8 reported for the Asiatic honeybee defensin cDNA gene (GenBank: ABS10820, direct submission). All of the inferred mature peptide sequences are highly homologous with the Western honeybee two defensin peptides (AmDe1 coded by one defensin cDNA gene, GenBank: U15955, and AmDe2 coded by one defensin cDNA gene, GenBank: AAR01214) and the royalisin peptide (termed as AmRo1) [26] (Figure 1A). Compared with the Western honeybee defensin peptides, one to three amino acid substitutions exist in each defensin peptide of the Asiatic honeybee. Mutations occur between the two hydrophobic amino acids, Phe and Leu, in AcDe1, AcDe2, AcDe3 and AcDe5, which may not change the property and structure. One acidic amino acid (Glu) substitutes the basic amino acid (Lys) at position 9 in AcDe6, which decreases the isoelectric point and may change antimicrobial properties. Three amino acid substitutions exist in AcDe7 as follows: one apolar amino acid (Gly) substitutes the polar amino acid (Cys) at position 3, one apolar amino acid (Pro) substitutes the polar amino acid (Thr) at position 41 and one natural mutation occurs at position 20 (Thr→Asn) in AcDe7 (Figure 1A). The first substitution may change the secondary structure of AcDe7, the second one would decrease the hydrophilic ability and may change other antimicrobial properties of AcDe7 and the last one may have no effects. There is one other unique Western honeybee defensin peptide peptide (AmDe3, GenBank: NP_001011638), which is an inferred peptide sequence for a recognized paralog of the definitive Western honeybee defensin gene [25]. It seems entirely unique compared with the amino acid sequences of other all defensins and royalisin. It has only 43 amino acids (VTCDVLSWQSKWLSIN-HSACAIRCLAQRKGGSCRNGVCICRK), while all others have 51 amino acids, and more amino acid substitutions among them. Moreover, the ORF of the Asiatic honeybee defensin has one extra amino acid (Gly) at the C-terminus that may be amidated as in the mature antimicrobial peptide of A. mellifera defensin [24,27].

The Asiatic honeybee defensin precursor gene is composed of three parts: a pre-region or signal region (coding 18 or 19 amino acids), a pro-region (between signal and mature region, coding 24 amino acids) and a mature region (coding 32 amino acids) (Figure 2). In total 303 nonsynonymous substitutions (NS) and 451 synonymous substitutions (SS) exist in the 29 precursor genes, 55 NS and 116 SS in the pre-pro-region, and 136 NS and 97 SS in the mature region. Most of the NS occupy the mature region, and most of the SS are in the pre-pro-region. The defensin mRNA transcript continues for an additional 41–126 bases (excluding the PolyA) after the stop codon (Figure 2).

Analysis of abaecin cDNA genes

In all, 11 different abaecin cDNA sequences (Abaecin1~Abaecin11, GenBank: EU714043~EU714053) coding 2 different abaecin peptides (termed as AcAb1 and AcAb2) were amplified and identified. Seven different cDNA genes coded for the major peptide AcAb1 and 4 different cDNA genes coded for peptide AcAb2. AcAb1 consists of 34 amino acids, while AcAb2 consists of 33 amino acids, and lacks the Gln at position 9 of Ab1 (Figure 1B). One natural mutation (Ile→Val) at position 7 exists in AcAb1 and AcAb2 compared with the Western honeybee abaecin peptide (termed as AmAb1 coded for by one Western honeybee abaecin cDNA gene, GenBank: NM_001011617) (Figure 1B), which may not change their structures and properties; one polar amino acid Gln is lacking in AcAb2, which would decrease the hydrophilicity of AcAb2. One polar amino acid Ser substitutes the apolar amino acid Ile at position 8 in AcAp2, and one apolar acid Ile substitutes the polar amino acid Asn. Both of these exist in the variable region, which would change the general antimicrobial spectrum while the conserved (constant) region remains the same. This would maintain general antimicrobial capacity [29,33]. One other Western honeybee abaecin peptide (termed as AmAb2, direct submission GenBank: U15954) is a special peptide, consisting of 33 amino acids with 5 amino acids (CAAA) extending at the N-terminal and 6 amino acids (KWPQGY). It is absent at the C-terminal compared with other honeybee abaecin peptides.

The Asiatic honeybee abaecin precursor gene is made up of two parts: a pre-region (coding 19 amino acids) and a mature region (coding 33 or 34 amino acids). It lacks a pro-region (Figure 3). In total, 22 NS and 48 SS exist in the 11 precursor genes, 3 NS and 6 SS in the pre-region and 4 NS and 10 SS in the mature region. The abaecin mRNA transcript continues for an additional 140 bases after the stop codon (excluding the PolyA) (Figure 3).

Analysis of apidaecin cDNA genes

In all, 13 different apidaecin cDNA sequences (Apidaecin1~Apidaecin13, GenBank: EU727255~EU727267) coding 4 different apidaecin peptides (termed as AcAp1~AcAp4) were amplified and identified. The apidaecin peptides from the Western honeybee are termed here as AmAp1~AmAp5. Of these, AmAp1~AmAp3 were isolated from hemolymph of Western honeybees that had been infected with bacteria [21] (Figure 1C). AcAp1 was coded for by 10 different cDNA genes, and AcAp2, AcAp3 and AcAp4 were coded for by one cDNA gene. AcAp1 is identical to AcAp3, and AcAp4 is identical to AmAp3. Compared with other honeybee apidaecin peptides, one amino substitution (Ser→Ile) exists at position 8 in AcAp2, one substitution (Asn→Ile) at position 8 in AcAp3, and one special substitution (Ser→Pro) at position 9 in
Figure 1. Alignment of mature peptides of the four antimicrobial peptide families of *A. cerana* and *A. mellifera*. (A): Alignment of mature peptides of honeybee defensins. AcDe1 to AcDe8 from the Asiatic honeybee; AmDe1, AmDe2 and AmRo1 from the Western honeybee. AcDe8, AmDe1, AmDe2 and AmRo1 downloaded from the NCBI website. (B): Alignment of mature peptides of honeybee abaecins. AcAb1 and AcAb2 from the Asiatic honeybee; AmAb1 and AmAb2 from the Western honeybee and downloaded from the NCBI website. (C): Alignment of mature peptides of honeybee apidaecins. AcAp1 to AcAp4 from the Asiatic honeybee; AmAp1—AmAp5 from the Western honeybee and downloaded from the NCBI website. (D): Alignment of mature peptides of honeybee hymenoptaecins. AcHy1—AcHy13 from the Asiatic honeybee; AmHy1 from the Western honeybee and downloaded from the NCBI website. Multiple sequence alignment performed using Clustal_X. Identical amino acids are in black dots. Different amino acids are shown in the alignment. Small dashes indicate gaps inserted for optimal alignment. Amino acid residues are numbered on the right.

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AmAp4 [28] (Figure 1C). AmAp4 is another apidaecin peptide in the Western honeybee, but it has never been detected in nature [29]. AmAp5 (GenBank: ABY84338), an apidaecin peptide, is unique in the Western honeybee, in that it consists of 17 amino acids and has a significantly different amino acid composition compared with other apidaecin peptides. It was successfully produced in the eukaryotic expression vector *Lactococcus lactis* [30].

The Asiatic honeybee apidaecin precursor gene is composed of two parts: a pre-pro-region and a variable number of repeated, 84 nt long, almost identical units (Figure 4). The pre-pro-region consists of a pre-region (coding for 19 amino acids) and a pro-region (coding for 13 amino acids). Each unit consists of coding regions for a spacer sequence and a basic dipeptide (RR) sequence, followed by coding sequence for the mature peptide. All the

![Figure 2. Nucleotide sequence and predicted amino acid sequence of Defensin1 (GenBank Accession EU727268).](image1)

![Figure 3. Nucleotide sequence and predicted amino acid sequence of Abaecin1 (GenBank Accession EU714043).](image2)
repeated units in the precursor gene of the Asiatic honeybee contain the same basic dipeptide (RR), except Apidaecin3, the nucleotide sequence of the dipeptide (CGT CGT) is mutated to TGT TGT, which codes for another dipeptide CR. In all, 117 NS and 162 SS exist in all the precursor genes, 108 NS and 141 SS in the pro-region and the repeated unit region, and 12 NS and 6 SS in the mature region. Most of the substitutions are in the pro-region.

The apidaecin mRNA transcript continues for about an additional 140 bases after the stop codon (excluding the PolyA) (Figure 4).

**Analysis of hymenoptaecin cDNA genes**

In all, 34 different hymenoptaecin cDNA sequences (Hymenoptaecin1–Hymenoptaecin18, GenBank: EU727297–EU727314; Hymenoptaecin19–Hymenoptaecin34, GenBank: EU835168–EU835183) coding for 13 different hymenoptaecin peptides (termed as AcHy1–AcHy13) were amplified and identified. Compared with the Western honeybee hymenoptaecin peptide (termed as AmHy1, coded for by one hymenoptaecin cDNA gene, GenBank: NM 001011615), two amino acid substitutions (Asn-Ser at position 16 and Val-Phe at 83) exist in all the Asiatic honeybee hymenoptaecin peptides, one substitution (Lys-Gln at position 9) in AcHy1–AcHy13, one substitution (Asn-Gln at position 37) in AcHy9, one substitution (Arg-His at position 53) in AcHy3–AcHy5, one substitution (Ser-Asn at position 43) in AcHy9, one substitution (Arg-His at position 53) in AcHy3–AcHy5, one substitution (Ser-Asn at position 43) in AcHy9, one substitution (Ser-Asn at position 43) in AcHy9, one substitution (His-Tyr at position 24) in AcHy11, and one substitution (Leu-Gln at position 50) in AcHy12 (Figure 1D). The Asiatic honeybee hymenoptaecin precursor gene is made up of three parts: a pre-region (coding for 17 amino acids), a pro-region (coding for 16 amino acids) and a mature region (coding for 93 amino acids). In all, 733 NS and 608 SS exist in all the precursor genes, 54 NS and 30 SS in the pre-pro-region, 499 NS and 448 SS in the mature region. The pre-pro-region is highly conserved, and most of the substitutions are in the mature region. The hymenoptaecin mRNA transcript continues for about an additional 230 bases after the stop codon (Figure 5), but only 75 bases (excluding the PolyA) after the stop codon are in the Hymenoptaecin28 sequences.

**Phylogenetic analysis**

The different precursor gene sequences from each of the cDNA genes coding for different mature peptides in the Asiatic honeybee and the complete or partial cds sequences in the Western honeybee downloaded from the NCBI website were selected to build the phylogenetic tree. The results show that there are three main branches (hymenoptaecin, defensin and apidaecin families), and the
abacin cDNA genes have a relatively close relationship with the hymenoptaecin cDNA genes (Figure 6). AmDef3 is one special sequence, which has only 46% possibility to group together with other defensin cDNA genes. Because of high homogeneity (similarities ranging from 96.5% to 100%) existing in each of the four antimicrobial peptide gene families, there are only a few branches and nodes within each of the four families (Figure 6). Although a few nucleotide sites vary within each of the four families, even only one nonsynonymous substitution in the relatively variable mature region will induce to generate a new peptide.

**Discussion**

As eusocial insects, honeybees have evolved both communal and individual traits that reduce the impact of their numerous parasites and pathogens. Among the individual traits, innate-immune responses have the potential to reduce both individual mortality and the spread of pathogens among colony members [3,31]. Propolis is a resinous substance variously composed of chewed tree buds mixed with pollen collected by honeybees as well as enzymes secreted by the bees. It has many functions in the defense of the nest, brood and stored food, and plays an important role in protecting the honeybee larvae by preventing diseases and parasites from entering the hive [32]. It’s puzzling that the Asiatic honeybee does not produce propolis. How does the Asiatic honeybee protect the larvae? When the workers visit plant flowers to collect pollen and honey, they are inevitably acting as potential disease vectors of different pathogens, such as bacteria, fungus and virus, and even some parasitic mites and insects. Thus the innate immune systems of adult workers are likely to play an important role not only in self defense, but also in helping to insure that the larvae and the queen are shielded from pathogens. In the wild, the Asiatic honeybee may pollinate many more plant species than the Western honeybee, and may therefore be at higher risk of contacting pathogens. It is easy to infer that the Asiatic honeybee may be under stronger selection to defend against pathogens and parasites than the Western honeybee.

Four antimicrobial peptide gene families were isolated from the Western honeybee and identified. Each gene family is responsible for a different duty in the innate immune system of the honeybee. Apidaecin is the most prominent component of honeybee humoral defense. Its unique precursor structure and/or processing makes...
Apidaecin easily triggered and potentially overproduced due to a low threshold for transcriptional activation [24]. In contrast, sizable experimental infections are required for induction of the normally silent defensin, abaecin and hymenoptaecin genes. Analysis of the apidaecin cDNA sequence of the Western honeybee indicated that up to 12 apidaecin peptides can be generated by processing single precursor proteins [28]. The same results were obtained when analyzing the apidaecin cDNA sequence of the Asiatic honeybee. However, only active antimicrobial peptides are detectable in adult honeybee haemolymph [28,33]. Apidaecin has strong antimicrobial activity to a wide spectrum of common plant-associated gram-negative bacteria [21,24], and it is the first line of defense against invading pathogens. Abaecin is another major antimicrobial peptide in the haemolymph of the honeybee. It is relatively weaker but as active as apidaecin against both gram-negative and gram-positive bacteria. It also has antimicrobial activity to some bacteria in vitro that are resistant to apidaecin [22].

There is no pro-region in the abaecin precursor gene, which means that the active peptide have been produced once it is translocated into the endoplasmic reticulum and without other processing [24]. An additional puzzling feature of abaecin is that it loses most of its antibacterial activity in the presence of phosphate-buffered saline or diluted bee haemolymph, media that resemble its physiological environment [22]. The weakly antimicrobial abaecin seems likely to function as a backup for bacterial resistance against apidaecin [22]. In relation to the major immune component (apidaecin), hymenoptaecin is largely complementary to apidaecin because it inhibits growth of certain gram-negative bacteria that are resistant to apidaecin [23]. Under honeybee physiological conditions, it inhibits the viability of gram-negative and gram-positive bacteria, including several human pathogens [23]. By comparison, bee defensin production is rather minimal and considerably delayed, representing the only component of the honeybee humoral defense specifically targeting gram-positive bacteria [24,26]. Steady state levels of defensin are probably sufficient to clear persistent gram-positive cells after 24 h of infection [24]. Under minor infection of pathogens in the wild, only apidaecin and hymenoptaecin genes were detected in the bodies of Western honeybee adults (abaecin and defensin were absent) [24]. The wide range of gram-negative bacteria targeted by apidaecin may account for the majority of honeybee infections in the wild [24]. Apidaecin and hymenoptaecin peptides might be important factors in restricting or killing different pathogens in the innate immune system of the honeybee. The abaecin peptide seems likely to function as a backup for the potential emergence of bacterial resistance against apidaecin [23]. The defensin peptide may play an important role in maintaining the immunological abilities of honeybees in the later stages of infections.

We have amplified and identified four families of antimicrobial peptide transcript genes from Asiatic honeybee adult workers infected by E. coli. Blastn results suggest that all of them share high sequence identity with the Western honeybee. The results of phylogenetic analysis suggest that all the cDNA genes in each of the four gene families in the same honeybee species are paralogous, and the peptides they encode for are allelic variants. For example, the high sequence similarities of defensin peptides and the identity of the pre-pro-regions of all peptides suggest that the defensin peptides might be variants encoded for by a single polymorphic gene [25]. In all, at least 29 different defensin cDNA genes encoding 7 defensin peptides, although one of the defensin variants (AcDe7) has a substitution at a cysteine residue that is definitive and widely conserved in nature, suggesting that this sequence might be from a pseudogene or allele with null activity. 11 different abaecin cDNA genes encoding 2 abaecin peptides, 13 different apidaecin cDNA genes encoding 4 apidaecin peptides and 34 different hymenoptaecin cDNA encoding 13 hymenoptaecin peptides exist in the Asiatic honeybee. In terms of the Western honeybee, in total 5 defensin cDNA genes (containing complete cds sequences, GenBank: NM_001011616, NM_001011638, AY333923, AY588474 and U15955) encoding 3 defensin peptides (AmDe1~AmDe3), 2 abaecin cDNA genes (containing complete cds sequences, GenBank: NM_001011617 and U15954) encoding one 1 abaecin peptide (AmAb1) and 1 directly submitted abaecin peptide (AmAb2, GenBank: AAL35348), 7 apidaecin cDNA genes (containing complete mature nucleotide sequences, GenBank: EU382095, NM_001011642, NM_001011613, X72577, X72576, X72575 and AF442148) encoding 5 apidaecin peptides, and 2 hymenoptaecin cDNA genes (containing complete cds sequences, GenBank: NM_001011615 and U15956) encoding 1 hymenoptaecin peptide have hitherto been reported. It is likely that additional antimicrobial peptide and cDNA genes exist in the Western

Figure 6. Neighbor-Joining (NJ) tree of four antimicrobial peptide cDNA gene families of A. cerana and A. mellifera. All the cDNA sequences for A. cerana were edited to begin at the initiation codon and end at the stop codon (containing complete cds). All the cDNA sequences for A. mellifera are downloaded from the NCBI website and edited to end in the stop codon (containing complete or partial cds). Numbers at nodes are bootstrap values based on 1000 replications. The cDNA gene names, GenBank accession numbers and the encoding peptide names are listed in Table 1. doi:10.1371/journal.pone.0004239.g006
honeybee, which may encode for the same mature peptides as reported here. However, there is no doubt that more antimicrobial peptides and their cDNA genes are found in the Asiatic honeybee than in the Western honeybee, specifically the hymenoptaecin peptides (13 versus 1). These results suggest that the long history of domestication of *A. mellifera* may have led to a lower standing diversity of antimicrobial peptides in its sequences. Additionally and/or alternatively, Asiatic honeybee adults may be under stronger selection pressure from pathogens and parasites, resulting in greater positive and diversifying selection on antimicrobial peptide sequences than in the Western honeybee.

### Materials and Methods

#### Insects

Colonies of the Chinese honeybee, *Apis cerana cerana* Fabricius, have been maintained on the campus of Zhejiang University since 2003. Adult honeybee workers were injected with viable *E. coli* (TG1) as described previously [21]. The fat bodies of the wasp adults were harvested 24 h after injection.

#### RT-PCR

Samples of the fat body were collected and washed twice with phosphate-buffered saline (PBS: 140 mM NaCl, 27 mM KCl, 8 mM Na2HPO4, 1.5 mM KH2PO4, pH 7.4). Total RNA was extracted from 40 live adult workers using a RNaseasy™ Mini Kit (Qiagen GmbH, D-40724 Hilden). The cDNA libraries were constructed using a Clontech, SMART™ PCR cDNA Synthesis Kit. After double stranded cDNA was synthesized, it was diluted 100 fold as the template. All the primers were designed from the four types of antimicrobial peptides gene families of *A. mellifera*. Primers used were as follows: P1 (for abaecin): GAT CTT CAC ACT CGA GGT CTT GCT (designed from a short stretch of sequence just upstream of the start codon of the *A. mellifera abaecin* gene), PCR conditions were 94°C, 2 min, 40× (94°C, 20 s; 54°C, 30 s; 72°C, 30 s), 72°C, 10 min; P2 (for apidaecin): TAA TCA TGA AGA ATT TTG CCT (designed from a short stretch of sequence just upstream of the start codon of the *A. mellifera apidaecin* gene), PCR conditions were 94°C, 2 min, 40× (94°C, 20 s; 54°C, 30 s; 72°C, 40 s), 72°C, 10 min; P3 (for hymenoptaecin): GAA ATC GAT CTC TAC AG (designed from a short stretch of sequence after the start codon of the *A. mellifera hymenoptaecin* gene), PCR conditions were 94°C, 2 min, 40× (94°C, 20 s; 50°C, 30 s; 72°C, 40 s), 72°C, 10 min; P4, P5 and P6 for amplifying complete defensin CDS sequences, P4: TGT CGG CCT TCT CTT CAT GG (designed from a short stretch of sequence after the start codon of the *A. mellifera defensin* gene), PCR conditions were 94°C, 2 min, 40× (94°C, 20 s; 50°C, 30 s; 72°C, 40 s), 72°C, 10 min; P4, P5 and P6 for amplifying complete defensin CDS sequences, P4: TGT CGG CCT TCT CTT CAT GG (designed from a short stretch of sequence after the start codon of the *A. mellifera defensin* gene), PCR conditions were 94°C, 2 min, 40× (94°C, 20 s; 50°C, 30 s; 72°C, 40 s), 72°C, 10 min; P4, P5 and P6 for amplifying complete defensin CDS sequences, P4: TGT CGG CCT TCT CTT CAT GG (designed from a short stretch of sequence after the start codon of the *A. mellifera defensin* gene), PCR conditions were 94°C, 2 min, 40× (94°C, 20 s; 50°C, 30 s; 72°C, 40 s), 72°C, 10 min; P5 (for 5'RACE): GAA ATC GAT CTC TAC AG (designed from a short stretch near the start codon of the sequence amplified by P4), PCR conditions were 94°C, 2 min, 40× (94°C, 20 s; 50°C, 30 s; 72°C, 30 s), 72°C, 10 min; P6 (for 5'RACE): GCC AAT ATG AAG ATC TAT TTG (designed from a short stretch near the start codon of the sequence amplified by P5), PCR conditions were 94°C, 2 min, 40× (94°C, 20 s; 50°C, 30 s; 72°C, 30 s), 72°C, 10 min. The 5'RACE primer in the kit was used as the forward primer and P5 as the reverse primer for 5'RACE of defensin. The CDS primer (ATT CTA GAG GCC GAG GCG GCC GAC ATG) in the kit was used as the reverse primer in all other PCR reactions. The PCR products were gel purified and cloned with TaKaRa pMD19-T Vector (TaKaRa Biotechnology, Dalian, China). Positive clones were first screened in blue/white colony screening, then positive clones were tested again with M13 vector primers in order to avoid, as far as possible, false positive clones. We sent 5–10 tested positive clones for each antimicrobial peptide gene family each time to Invitrogen Biotechnology Company (Invitrogen, Shanghai, China) for sequencing until no new cDNA sequence (coding different mature peptide) was found. In all, 156 clones that tested positive were sent to sequence, 56 clones for defensin, 23 clones for abaecin, 25 clones for apidaecin and 32 clones for hymenoptaecin.

#### Signal sequence cleavage sites

Signal peptide cleavage sites were predicted using the SignalP 3.0 server program [34].

### Table 1. Four antimicrobial peptide cDNA gene families of *A. cerana* and *A. mellifera* selected to construct the phylogenetic tree.

| Honeybee species | cDNA genes | GenBank accession numbers | Encoded peptide |
|------------------|------------|---------------------------|-----------------|
| *A. mellifera*   | AmDef1     | U15955                    | AmDef1          |
|                  | AmDef3     | AY588474                  | AmDef3          |
|                  | AmApi1     | NM_001011642              | AmApi1          |
|                  | AmApi3     | X72577                    | AmApi1          |
|                  | AmHym1     | U15956                    | AmHym1          |
|                  | AmDen2     | AY333923                  | AmDe2           |
|                  | AmAb1      | NM_001011617              | AmAb1           |
|                  | AmAb2      | NM_001011613              | AmAb1           |
|                  | AmAb4      | AF442148                  | AmAb1           |
| *A. cerana*      | AcDef1     | EU727268                  | AcDe1           |
|                  | AcDef6     | EU727273                  | AcDe3           |
|                  | AcDef8     | EU727275                  | AcDe5           |
|                  | AcDef7     | EU727274                  | AcDe7           |
|                  | AcAb1      | EU714043                  | AcAb2           |
|                  | AcApi3     | EU727257                  | AcAp2           |
|                  | AcApi7     | EU727261                  | AcAp4           |
|                  | AcHym3     | EU727299                  | AcHy2           |
|                  | AcHym5     | EU727301                  | AcHy4           |
|                  | AcHym8     | EU727304                  | AcHy6           |
|                  | AcHym16    | EU727312                  | AcHy8           |
|                  | AcHym18    | EU727314                  | AcHy10          |
|                  | AcHym28    | EU835177                  | AcHy12          |
|                  | AcDef4     | EU727271                  | AcDe2           |
|                  | AcDef3     | EU727270                  | AcDe4           |
|                  | AcDef2     | EU727269                  | AcDe6           |
|                  | AcAb2      | EU714044                  | AcAb1           |
|                  | AcApi1     | EU727255                  | AcAp1           |
|                  | AcApi4     | EU727258                  | AcAp3           |
|                  | AcHym1     | EU727297                  | AcHy1           |
|                  | AcHym4     | EU727300                  | AcHy3           |
|                  | AcHym7     | EU727303                  | AcHy5           |
|                  | AcHym9     | EU727305                  | AcHy7           |
|                  | AcHym17    | EU727313                  | AcHy9           |
|                  | AcHym25    | EU835174                  | AcHy11          |
|                  | AcHym19    | EU835168                  | AcHy13          |

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Author Contributions

Conceived and designed the experiments: PX MS. Performed the experiments: PX. Analyzed the data: PX MS. Wrote the paper: PX. Intellectual contributions during the design and implementation of this study, and during the writing of the manuscript: X-XC. Provided funding in support of this study: X-XC.

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