Identification of Immune Response to Sacbrood Virus Infection in Apis cerana Under Natural Condition

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Chinese sacbrood virus (CSBV) is a serious threat to eastern honeybees (Apis cerana), especially larvae. However, the pathological mechanism of this deadly disease remains unclear. Here, we employed mRNA and small RNA (sRNA) transcriptome approach to investigate the microRNAs (miRNAs) and small interfering RNAs (siRNAs) expression changes of A. cerana larvae infected with CSBV under natural condition. We found that serine proteases involved in immune response were down-regulated, while the expression of siRNAs targeted to serine proteases were up-regulated. In addition, CSBV infection also affected the expression of larvae cuticle proteins such as larval cuticle proteins A1A and A3A, resulting in increased susceptibility to CSBV infection. Together, our results provide insights into sRNAs that they are likely to be involved in regulating honeybee immune response.

Keywords: Chinese sacbrood virus, serine protease, Apis cerana, transcriptome, siRNA, immune response

INTRODUCTION

The Asian honeybee, Apis cerana, is an important pollinator to maintain plant biodiversity in Southeast Asian countries, especially to wild flowering plants and crops (Ai et al., 2012). However, honeybees have been suffering colony decline in recent years due to an increasing number of infections from multiple pathogens (Diao et al., 2018a). Among honeybee pathogens, Chinese sacbrood virus (CSBV), a Chinese strain of sacbrood virus (SBV), is the major factor threatening the survival of A. cerana colony (Sun et al., 2018). SBV is the single-strand positive RNA virus, which can infect honeybee larvae and lead to larvae death (Chen and Siede, 2007). SBV belongs to the genus Iflavirus, the family Iflaviridae within the order Picornavirales (Bailey et al., 1964). The full-length genome of SBV was sequenced in 1999 (Ghosh et al., 1999).

The clinical symptom of CSBV-infected A. cerana larvae was ecdisial fluid aggregated with typical “sac,” resulting in failure to pupate (Ma et al., 2013). CSBV was identified firstly from A. cerana larvae in Guangdong province of China in 1972 (Yang et al., 1988), and frequently caused extensive larvae death and colony decline and recently re-emerged as one of the causative agents of larvae disease outbreaks in Liaoning province in 2008 (Ma et al., 2011).
frequently detected in *A. cerana* and remains a major threat to *A. cerana* in China (Hu et al., 2016; Diao et al., 2018a). Usually, viral infection can induce host cell apoptosis and tissue damage as well as functional disorder of target genes. Then, all these alterations can be displayed through gene expression changes (Tan et al., 2007; Francis et al., 2012).

Like many other social insects, honeybees have no adaptive immune response known in vertebrates (Evans and Armstrong, 2006). To survive under the persistent viral infection, honeybees have evolved several defense mechanisms that are activated via different immune pathways (Evans et al., 2006; Zou et al., 2006). Honeybee antiviral defense mechanisms include RNA interference (RNAi), Janus kinase/Signal Transducer and Activator of Transcription (JAK/STAT) pathway, Toll pathway, Immune Deficiency (Imd) pathway, c-Jun N-terminal kinase (JNK) pathway, Mitogen-Activated Protein Kinases (MAPK) pathway and melanization, as well as reactive oxygen species generation (Brutzer et al., 2015). Serine protease (SP) and serine protease homolog (SPH) play an important role in innate immune response that includes coagulation, melanization, and the Toll signaling pathway in honeybee (Zou et al., 2006; Gao et al., 2017).

Experimental evidence has shown that CSBV infection induced routine immune responses by enhancing the expression of antimicrobial peptides (AMPs) (Liu et al., 2017). Generally, activation of the Toll, Imd and JAK/STAT pathways result in the production of AMPs and other effector proteins (Brutzer et al., 2015). Nevertheless, knowledge about the comprehensive immune responses to CSBV infection is limited. Omics technique is a useful tool to measure the dynamic changes of whole gene expression associated to viral infection (Sheyn et al., 2018). Using the proteomic technique, it was found 142 proteins and 12 phosphoproteins down-regulated in CSBV-infected larvae, which were significantly affected in carbohydrate, energy and fatty acid metabolism (Han et al., 2013). Lately, the full-genome sequence of *A. cerana* was determined, providing new insights into physiological resistance to Varroa mites, and was found to have six more immune genes than *Apis mellifera* (Diao et al., 2018b).

In addition, transcriptome technology was applied to several fields of honeybees, such as gland development (Liu et al., 2014), Varroa mite control (Campbell et al., 2015), and *Ascposphaera apis* pathogenesis (Cornman et al., 2012). Previous studies have confirmed that chemical stressors and pathogens can lead to up-regulate of AMPs (Siede et al., 2012; Li et al., 2016). Galbraith et al. (2015) identified the transcript and epigenetic responses of honeybees to Israeli acute paralysis virus (IAPV) infection and found that honeybees employed distinct immune pathways such as JAK-STAT to resistance viral infection besides universal immune responses. Recently, Rutter et al. (2019) confirmed that IAPV infection combined with diet quality can affect the immune gene expression of Notch and JAK-STAT signaling pathways. Nevertheless, artificial infection with acute bee paralysis virus (ABPV) did not induce a humoral immune response of larvae and workers (Azzami et al., 2012). In addition, artificial infection frequently caused the unexpected transcription changes of AMPs as well as non-target genes (Boncristiani et al., 2013).

Small RNAs (sRNAs) include microRNAs (miRNAs) and small interfering RNAs (siRNAs) that are involved in regulating gene expression in most organisms (Ding and Voinnet, 2007; Flenniken and Andino, 2013; Fletcher et al., 2016). miRNAs are a group of sRNAs with 22 nt in size and are important regulators of diverse biological processes, including development and interactions between host and virus (Ding and Voinnet, 2007). siRNA is produced by double-stranded RNA (dsRNA), which can be from either the viral genome itself or an intermediate dsRNA product generated during viral replication (Fung et al., 2018). Deep sequencing on honeybee samples found that vsiRNAs were matched to several honeybee viruses such as Deformed wing virus (DWV) and IAPV (Chejanovsky et al., 2014; Fung et al., 2018). Virus-derived small interfering RNAs (vsiRNA) produced during the viral infection are a group of siRNA ranged from 21 to 24 nt in size that may hijack the host RNAi pathway (Aliyari et al., 2008). In brief, vsiRNA guides the RNA induced silencing complex (RISC) to target viral genomes, which can potentially alter the host transcriptome responses (Ding, 2010; Yang et al., 2018). However, little research has been done on the sRNAs from CSBV. Thus, the mechanisms underlying the host responses to CSBV infection are unknown, especially the effects of sRNAs on CSBV infection under natural condition. Here, we examined the transcriptional responses and the abundance of siRNAs of *A. cerana* larvae to CSBV infection under natural conditions. We characterized (1) the transcriptomic variation of genes related to larval development and immune, (2) the host’s metabolism, and (3) determined whether there is a relationship between siRNA and CSBV infection. Our results provide insights into up- and down-regulated in cuticle protein and serine proteinase during CSBV infection. These findings significantly broaden our knowledge of virus-host interactions and provide novel targets for the control of CSBV.

### MATERIALS AND METHODS

#### Sample Collection

The samples were collected from Liaoning and Guangdong provinces in October 2016. The larvae of *A. cerana* were collected from three apiaries in different regions (Longmen, Meizhou, and Conghua) in Guangdong province in China. CSBV-infected larvae were taken from the colonies with obvious cystic symptoms and confirmed it using RT-PCR with a pair of specific primers, 5′-CCTGGGAAAGTTGGCTAGTTACG-3′ and 5′-CCTATCACCATCCATGGGTACG-3′ according to the described by Ma et al. (2013) and the healthy larvae were collected from three different apiaries (Huludao and Benxi in Liaoning Province, and Guangzhou in Guangdong province). For RNA sequencing, the larvae of *A. cerana* were mixed into three groups with the same genetic background. To avoid interferon by other viruses, six common honeybee viruses were tested before RNA sequencing (Supplementary Table 1).

The 2 or 3-day-old larvae were used for transcriptome analysis since they were most susceptible to CSBV (Sun et al., 2018). The age of larvae was identified by confining a queen to lay eggs within 24 h. Twenty 2-day-old larvae were considered as one treatment.
RNA Extraction, Library Construction, and RNA-seq

Total RNA of each sample were isolated from the larvae using the Trizol Reagent (Life technologies, California, CA, United States). RNA quality was measured using an Agilent 2100 Bioanalyzer (Agilent Technologies, Inc., Santa Clara, CA, United States). The mRNA was obtained by NEBNext Poly (A) mRNA Magnetic Isolation Module (NEB, Ipswich, MA, United States). Briefly, the enriched mRNA was fragmented into approximately 200-nt RNA inserts, which were used to synthesize the cDNA. The end-repair/dA-tail and adaptor ligation were performed to the double-stranded cDNA. The corresponding fragments were obtained by Agencourt AMPure XP beads (Beckman Coulter, Inc., United States) and PCR amplification. PCR was performed by using Phusion High-Fidelity DNA polymerase, Universal PCR primers and Index (X) Primer. PCR products were purified (AMPure XP system) and library quality was assessed on the Agilent Bioanalyzer 2100 system. Then, the clustering of the index-coded samples was performed on a cBot Cluster Generation System using TruSeq PE Cluster Kit v4-cBot-HS (Illumina, Inc., United States) according to the manufacturer's instructions and the library preparations were sequenced on an Illumina Hiseq 2500 platform and single-end reads were generated.

For sRNA library construction, about 5 μg total RNA we used to construct sRNA library using the Small RNA Sample Prep Kit (Illumina). Since the sRNA has a phosphate and hydroxyl group at the 5' and 3' end, respectively, the T4 RNA ligase 1 and ligase 2 (truncated) were respectively ligated to corresponding ends of the sRNA. After ligation, the ligated RNA fragments were reverse transcribed using M-MLV reverse transcriptase (Invitrogen, Inc., United States) and then the resultant cDNA products were amplified with two primers corresponding to the ends of the adapter sequences. Polyacrylamide gel was used to get sRNA libraries by electrophoresis and rubber cutting recycling. Finally, to perform clustering and sequencing of sRNA, cluster generation was performed by using the TruSeq PE Cluster Kit v4-cBot- HS (Illumina, Inc., United States) according to the manufacturer's protocols. After cluster generation, the library construction was loaded to an Illumina Hiseq 2500 platform and sequencing single-end reads were created.

Raw data with FASTQ format were analyzed through in-house Perl scripts. Clean reads were obtained by removing those reads containing adapter, ploy-N, and low-quality. Reads with smaller than 18 nt or longer than 30 nt were trimmed and cleaned. Meanwhile, several parameters of the clean data were calculated such as Q20, Q30, GC-content and sequence duplication level. Low-quality reads, such as only adaptor, unknown nucleotides >5%, or Q20 < 20% (percentage of sequences with sequencing error rates <1%), were removed by Perl script. All the downstream analyses were based on clean data with high quality. The clean reads filtered from the raw reads were mapped to A. cerana genome (Park et al., 2015; Diao et al., 2018b) and A. mellifera genome (Aml-4.5) (Elsik et al., 2014) using Tophat2 software. The aligned records from the aligners in BAM/SAM format were further examined to remove potential duplicate molecules. Gene abundance were calculated based on the value of the transcripts per million (TPM).

Identification of Differential Gene Expression and Sequence Annotation

DESeq (Anders and Huber, 2010) and Q-value were employed to evaluate differential gene expression between CSBV and control groups. Gene expression levels were estimated using FPKM values (fragments per kilobase of exon per million fragments mapped) using Cufflinks software (Trapnell et al., 2010). The false discovery rate (FDR) control method was used to identify the threshold of the P-value in multiple tests to compute the significance of the differences. Here, only genes with an absolute value of log2 fold change ≥1.5 and FDR significance score <0.05 were used for subsequent analysis. Genes were compared against various protein databases by BLASTX, including the National Center for Biotechnology Information (NCBI) non-redundant protein (Nr) database, and Swiss-Prot database with a cut-off E-value of 10-5. Furthermore, genes were searched against the NCBI non-redundant nucleotide sequence (Nt) database using BLASTn by a cut-off E-value of 10-5. Genes were retrieved based on the best BLAST hit (highest score) along with their protein functional annotation. Differentially expressed honeybee genes were analyzed against a background set of genes, which are all the Drosophila orthologs (Drosophila genome vs. Dmel r5.42) in the honeybee genome (A. cerana and A. mellifera genome) (Elsik et al., 2014; Park et al., 2015; Diao et al., 2018b). The hierarchical clustering heatmap analysis of expression level of all genes was performed by MeV.1

Identification of Small RNA Targeted Genes

Here, the sequences of ribosomal RNA (rRNA), transfer RNA (tRNA), small nuclear RNA (snRNA), small nucleolar RNA (snoRNA), and other kinds of non-coding RNAs were identified using a basic local alignment search tool (Bowtie tools soft, Version 1.1.2) against known non-coding RNAs deposited in the Silva database, GtRNAdb database, Rfam database and Repbase database, respectively, and unannotated reads of sRNA were obtained (Justin et al., 2014). MiRDeep2 software (Version 2.0.5) was mainly used for the prediction of animal miRNAs (Friedländer et al., 2012). Next, we mapped sRNA reads with lengths of 18–30 nt to the A. cerana genome (Park et al., 2015; Diao et al., 2018b) and CSBV genome (GenBank: KU574662.1) to get miRNA and CSBV-specific vsiRNA by using miRDeep2. Mapped reads were used to identify known miRNAs and novel

1https://sourceforge.net/projects/mev-tm4/
miRNAs using miRBase 20.0 (Griffiths-Jones, 2006) and miREvo (Wen et al., 2012). Usually, these known miRNA data have been deposited in miRBase. Based on the distribution information of reads on the precursor sequence (miRNA production characteristics, mature, star, loop) and energy model of precursor structure (RNAfold randfold), the Bayesian model was used to score and authenticate the miRNAs and vsiRNA (Liu et al., 2009). We then classified these sRNAs into several different categories according to their annotations.

Last, we used psRobot to identify potential target genes of vsiRNAs in A. cerana (Wu et al., 2012), and used a well-established miRNA-target database MiRanda (v3.3a) and TargetScan (Version:7.0) to predict the target genes of the identified miRNAs, and investigated the possible functions using the method of gene function annotation (Agarwal et al., 2015). The vsiRNA profile was used as a query in sRNA target prediction module psRobot (Wu et al., 2012; see text footnote 3). Target gene function was annotated based on the following databases: Nr (NCBI non-redundant protein sequences); Nt (NCBI non-redundant nucleotide sequences); Pfam (Protein family); KOG/COG (Clusters of Orthologous Groups of proteins); Swiss-Prot (A manually annotated and reviewed protein sequence database); KO (KEGG Ortholog database); GO (Gene Ontology). The analysis of miRNA-mediated mRNA was performed based on the predicted miRNA-mRNA relationships. We selected miRNA-mRNA pairs showing opposite expression changes. The absolute value of log2 fold change ≥1 was used as a cutoff score.

GO and KEGG Analysis

Genes were annotated in the NCBI non-redundant hits with GO (Conesa et al., 2005). Perl script was then used to plot GO functional classification for the genes with a GO term hit to view the distribution of gene functions. The obtained annotation was enriched and refined using TopGo (R package). The gene sequences were also aligned to the COG database to predict and classify functions (Tatusov et al., 2000), Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways were assigned to the assembled sequences by Perl script. The interaction networks of corresponding proteins of targeted genes were predicted using STRING.

Verification of Expression Difference Genes by qPCR

For the analysis of miRNA expression, cDNA synthesis was performed using the miRCURY LNA Universal cDNA Synthesis kit II (Exiqon, Woburn, MA, United States) according to the manufacturer’s instructions and using specific reverse transcription primer (5′-CTCAACTGCTGGGCTGAGTGC GCAATTCCGTTGACCCCGCAAA-3′) (Fu et al., 2012). Stem-loop reverse transcription quantitative PCR (qPCR) were performed as follows: 94° for 2 min followed by 30 cycles of 94°C for 30 s, 56°C for 35 s, and 72°C for 20 s. Specific sequence

\[\text{ Target miRNA 5′-ACAATCCAGCTGGGGCTGAGTGC GCAATTCCGTTGACCCCGCAAA-3′ }\]

\[\text{ Reference gene U6 (F: 5′-AACGCTTCAGGAATTTGCGT-3′) }\]

\[\text{ Primer set for qPCR (10 μM) }\]

\[\text{ Amplification reactions }\]

\[\text{Product size }\]

\[25 \mu L \text{ of 2× SYBR Premix Ex TaqTM II (Takara, Dalian, China), 0.5 μL of upstream and downstream primers (10 mM), respectively, 2 μL of template cDNA, and 9.5 μL of double-distilled H2O in a total volume of 25 μL. Real-time qPCR was performed using a cycling sequence of 95°C for 30 s, followed by 40 cycles of 95°C for 5 s, 54-56°C for 25 s, and 72°C for 25 s. β-actin was used as housekeeping gene and three technical replicates were performed. To verify the reliability of the RNA-seq result, twenty 2-day-old larvae were used to measure the expression of immune genes, while five larvae each of day-old (4th, 5th, 6th, and 7th) were used to quantify the expression of vsiRNA. Three replicates were run.}

Statistical Analysis

The limma algorithm was used to analyze the differentially expressed mRNAs and sRNAs according to the significant analysis with FDR analysis (Wettenhall et al., 2006). All data analysis meets the following two criteria: (1) log2 fold change ≥1.5 or P < 0.05; and (2) FDR < 0.05. The relative expression of the target genes in the infected and control groups were calculated with the 2^−ΔΔCt method. Statistical comparisons and Pearson correlation coefficient analysis were performed using GraphPad Prism 7 (GraphPad Software Inc., San Diego, CA, United States). Multiple t-tests were used to compare the significance of differential expression of genes between CSBV and control groups. The asterisks in the figures indicate significant differences (* P < 0.05; ** P < 0.01).

RESULTS

The Overview of Expressed mRNA and Small RNAs

A total of six RNA-seq libraries were obtained using Illumina sequencing platforms, but we subjected two pooled controls and two pooled CSBV infected samples to further analysis due to the two poor data with other virus contamination (data not show). For mRNA sample sequencing, a total of 18,270,587 (T01), 19,574,141 (T02), 17,646,189 (T03), and 20,618,929 (T04) clean reads were obtained from the healthy and CSBV-infected mRNA libraries (Supplementary Table 3) (T01 and T02, control groups; T03 and T04, e CSBV-infected group). Mapped reads to A. cerana genome were more than 70% among all samples
GO and KEGG Analysis on Differentially Expressed mRNA Involved in Response to CSBV Infection

A total of 10,262 mRNA transcripts were produced (FDR < 0.05) (Supplementary Table 6), and 203 significantly differently expressed genes (DEGs) (Supplementary Table 7). The cutoff of log2-fold change > 1.5 and P-value > 0.8 was used to filter genes that were differentially expressed between CSBV-infected and the control samples (Tarazona et al., 2011). Among the 203 DEGs, 83 genes were successfully matched to GO terms (Figure 2A), and the top 10 GO terms (P ≤ 0.01) of molecular function belong to the structural constituent of cuticle, serine-type endopeptidase, serine-type peptidase, and serine hydrolase activity (Figure 2A). Based on GO-directed acyclic graph the relationship among these 10 terms in molecular function showed that the structural constituent of cuticle and serine-type endopeptidase activity were significantly enriched (P < 0.01) (Figure 2B). Ten genes related to the structural constituent of cuticle were significantly up-regulated including larval cuticle protein A1A, larval cuticle protein A2B, flexible cuticle protein 12-like, and larval cuticle protein A3A (Supplementary Table 7). While seven SP genes were significantly down-regulated except easter was up-regulated.

Kyoto Encyclopedia of Genes and Genomes analysis on 203 DEGs found that fatty acid metabolism, biosynthesis of unsaturated fatty acids, AMP-activated protein kinase (AMPK) signaling pathway, and peroxisome proliferator-activated receptors (PPAR) signaling pathway were mainly enriched in 20 categories (Figure 2C). Two acyl-CoA Delta 11-desaturase-like genes (ACSNU05780 and ACSNU05780) involving in fatty acid metabolism and PPAR signaling pathway were significantly up-regulated to fourfold. Furthermore, we also found that two genes related to fatty acid biosynthesis were significantly down-regulated to 0.12 fold at least (Supplementary Table 7).

GO Analysis and KEGG Annotation on Target mRNA of Differentially Expressed miRNA

Using miRDeep2 and DESeq software, we mapped clean sRNA reads to the A. cerana genome against the miRanda database (Park et al., 2015; Diao et al., 2018b). The compositions of those reads of sRNA are shown in Supplementary Figure 3, and the most abundant class of sRNAs were 22-nt in size, including known miRNA and new identified miRNA (Supplementary Figures 3A,B). The 260 miRNAs were obtained included 23 differently expressed miRNAs (Supplementary Table 8, log2 fold change > 0.6 or <−0.6, P < 0.05). The 260 miRNAs were targeted 1035 mRNAs and 23 differently expressed miRNAs were predicated toward 227 target mRNAs that involved in biological process, cellular component, and molecular function (Figure 3A). KEGG annotation analysis showed that 227 target genes mainly participate in ECM-receptor interaction, ribosome biogenesis in eukaryotes and endocytosis (Figure 3B).

We identified a miRNA, ame-miR-3759, was significantly up-regulated to more than onefold (Table 1), while the expression of the two target genes, LOC410515 and putative uncharacterized
FIGURE 1 | Summary of the types and features of genes from different immune pathways. (A) The analysis of expression level of samples based on FPKM value. T01 and T02, T03 and T04 indicate control and CSBV-infected groups. (B) Correlation analysis of the expression of all mRNA between healthy groups and CSBV-infected groups. (C) The 144 differentially expressed immune genes were shared between our study and Evans et al. (2006) described. (D) The interaction network of 144 shared genes was predicted and found four major terms enriched ($P < 0.05$). Green, serine proteases and trypsin domain; red, Jak-STAT signaling pathway; blue, innate immune response; yellow, Toll receptor homology domain. (E) The analysis of the regulated genes related to serine proteases, melanization, and Toll pathway, Jak-STAT signaling pathway, innate immune response, and IMD pathway after CSBV infection. The values represent standardized expression levels based on FPKM mean values. Green and red indicate decreased and increased in expression levels, respectively. (F) Heat map of the regulated genes related to serine proteases, melanization, and Toll pathway, Jak-STAT signaling pathway, innate immune response, and IMD pathway. (H) The relative expression levels of serine protease and defensin genes from RNA-seq (T02, T03, and T04) and qPCR analyses. Actin was used as the internal reference gene. *$P < 0.05$, **$P < 0.01$. (I) The differentially expressed serine protease and defensin genes (*$P < 0.05$, FDR < 0.05) were compared with those of SBV-induced (Ryabov et al., 2016) (*$P < 0.05$, FDR < 0.05).
protein DDB_G0277255-like, were significantly down-regulated (P < 0.01). LOC410515 may have a similar function in immune with a homologous protein, Apis dorsata chorion peroxidase-like (XP_006611124.1) (NCBI blast sequence identify >75%), which played an important role in melanin synthesis (Feng et al., 2018).

Identification and Analysis on Target mRNA of CSBV-Specific siRNA

We aligned all the cleaned sRNA-seq reads to the CSBV genome (GenBank: KU574662.1) and we identified 319 out of 2,467 unique vsiRNAs targeted 290 genes. Most of targets genes were enriched in involved in DNA binding, transcription factor activity, apoptosis, G-protein coupled receptor activity, transporter activity, and SP activity (Figure 4A and Supplementary Table 10). KEGG analysis on target mRNA of CSBV-specific siRNA showed that purine metabolism, hippo signaling pathway and fatty acid biosynthesis were significantly enriched (P < 0.05) (Figure 4B).

Although the expression of Dicer-like and Argonaute-2 (Ago2) genes associated to the RNAi pathway were up-regulated (Supplementary Table 5 and Supplementary Figure 4), we focused on siRNAs related to serine/threonine kinase, serine-type endopeptidase and serine-type peptidase genes based on the analysis of mRNA and miRNA data. Generally, the complete expression profile of vsiRNAs was categorized into “low” (TPM value <50), “high” (TPM value ≥50), and “very high expression” (TPM value ≥1,000) (relative abundance) based on the TPM value of siRNAs (Ramesh et al., 2017). For example, the expression levels of vsiRNA_1441 (TPM value is 2,932) were extremely significant higher than other siRNAs (3–20 times higher, mean value is 405) (Figure 4C and Supplementary Table 11). Then, we found that the target mRNA of vsiRNA_1441 was extracellular serine/threonine
FIGURE 3 | GO and KEGG annotation on target genes of differentially expressed miRNA. (A) The top GO terms enriched \((P < 0.05)\) on target genes of differentially expressed miRNA in molecular function, biological process and cellular component. (B) The KEGG analysis of target genes of differentially expressed miRNA \((P < 0.05)\).

TABLE 1 | The opposite expression in miRNA-mRNA pairs.

| miRNA         | miRNA_log2 fold change | Target mRNA       | mRNA_log2 fold change | Annotation            | Homologous protein      |
|---------------|------------------------|-------------------|-----------------------|------------------------|-------------------------|
| ame-miR-3759  | 1.09                   | ACSNU05809        | −3.40                 | DDB_G0277255-like      | Peptidyl-prolyl cis-trans isomerase |
| ame-miR-3759  | 1.09                   | ACSNU05034        | −3.61                 | LOC410515              | Chorion peroxidase      |

proteins protein kinase FAM20C-like of Apis, which potentially regulate many processes including cell survival, growth and metabolism. Subsequently, qPCR analysis confirmed that the expression of extracellular serine/threonine protein kinase FAM20C was significantly down-regulated (Figure 4D). Next, the expressions of vsiRNA_1441 of CSBV-infected larvae at different day-old were identified by RT-PCR (Figure 4E). Similarly, the expression of the extracellular serine/threonine protein kinase FAM20C gene was significantly down-regulated at the 4-, 5-, 6-, and 7-day-old after CSBV infection (Figure 4F).

DISCUSSION

Healthy larvae are crucial for the development and growth of the *A. cerana* colony population. However, CSBV was considered as one of factors contributed to recent declines of the *A. cerana* colony (Diao et al., 2018b). A few studies have shown that vsiRNAs negatively regulate host mRNAs and effectively silence host genes to gain more proliferation (Yang et al., 2018). Here, to understand the immune response of the CSBV infection, we presented the first sRNA analysis to CSBV under natural condition. The RNA-seq results showed that 203 DEGs were significantly altered between the healthy and infected larvae. Of these, all cuticle protein genes related to structural constituent of cuticle were significantly up-regulated, while SP genes related to serine-type endopeptidase activity, serine-type peptidase activity and serine hydrolase activity were significantly down-regulated. Meanwhile, 23 differently expressed miRNAs and 319 effective vsiRNAs were identified, which some of vsiRNAs targeted serine/threonine kinase or serine-type endopeptidase-related genes. These findings provide a novel insight into how CSBV resists host immune responses and result in larvae unable to the pupae.

Cuticle proteins might play a key role in resistance to CSBV infection. Cuticle is a barrier against viral invasion in animals such as white spot syndrome virus infection in shrimp (Corteel et al., 2009). Cuticle proteins play a critical role in keeping the body from pathogens and serving as a barrier to resistance them (Zhao et al., 2017). It has been reported that natural resistance-associated macrophage protein (NRAMP) was a cellular receptor of Sindbis virus in insects (Rose et al., 2011). In aphidiae, cuticle protein 4 is essential to entry of cucumber mosaic virus (CMV) because it was considered as a viral putative receptor (Liang and Gao, 2017). Although NRAMP was not enriched in our study, our results showed that the expression levels of larvae cuticle protein genes were greatly elevated including larval cuticle protein A2B, larval cuticle protein A1A and larval cuticle protein A3A. While Ryabov et al. (2016) found that the genes involved in cuticle and muscle development were down-regulated at a later stage after...
SBV infection (9 dpi). This was similar with that deformed wing virus infection will induce a cuticular protein down-regulated, apidermin 3 (Tomas et al., 2019). Thus, we infer that cuticle protein may have an important role to resistance CSBV infection during the development of larvae.

Serine proteases were significantly inhibited by CSBV infection. SPs with 60–400 members in insects form a large family of enzymes that hydrolyze peptide bonds at different rates (Rawlings and Barrett, 1993; Cao et al., 2017). SPs and SPHs, especially extracellular SPs, have a great influence on insect development and innate immunity (Ahola et al., 2015; Dudzic et al., 2019). Experimental evidence showed that SPs of *A. mellifera* may be involved in the embryonic development, melanization, and immune responses (Zou et al., 2006; Rodriguez-Andres et al., 2012). Our results suggested that most immunity genes related to the SPs were overlapped with Evans’ study (Evans and Armstrong, 2006) and two SPs, serine-type endopeptidase and peptidase activity, were significantly down-regulated. RT-qPCR results confirmed also that CSBV infection caused down-regulation of SP genes, such as SP stubble and transmembrane protease serine 9. This was consistent with that of sacbrood virus infection (Ryabov et al., 2016).

Honeybees can initiate humoral immunity involving SPs that include coagulation, melanization, and the Toll signal pathway (Rodriguez-Andres et al., 2012). SPs exist in hemolymph in the form of zymogens and then activate downstream genes such as *ppo*. Although Gao et al. (2017) identified an SP gene, *Accsp10*, related to development and pathogens resistance of *A. cerana*, they did not confirm its function. However, our results showed that the *ppo* gene of melanization pathway were down-regulated. Additionally, we also found that the expression level of the *ppo* gene was significantly down-regulated at the 4-, 6-, and 7-day-old in CSBV-infected larvae and suggested that CSBV infection significantly suppressed the melanization response by inhibiting the expression of SP and PPO. This was similar to that of semliki forest virus inhibited on the phenoloxidase cascade of mosquito (Rodriguez-Andres et al., 2012). However, intermediate steps of CSBV infection inhibited the SPs need to be further investigated.

Small RNAs have been confirmed to be related to a diversity of biological processes including cell proliferation and apoptosis through the post-transcriptional regulation of gene expression (Charkhpour et al., 2011; Zhang et al., 2014; Yang et al., 2017). RNA profile analyses suggested that these sRNAs were involved in biological processes associated with development.
and immunity of honeybees (Chen et al., 2017). The key initiator of the RNAi pathway is dsRNA, which produced by proliferation process are processed into vsiRNA duplexes by dicer-2 (Xu and Zhou, 2017). Viral small interfering RNAs are processed into 21-nt through Dicer-2 on viral dsRNA and then are integrated into insect Ago2 and guide the Ago2 onto target RNAs to cause their degradation (Bartholomay and Michel, 2018). It has also been reported that vsiRNAs can regulate the gene expression related to the host RNAi pathway and enhance the pathogenesis. For example, Xu et al. (2017) found that siRNAs derived from Southern rice black-streaked dwarf virus targeted several types of genes related to host resistance such as receptor-like protein kinases. Chejanovsky et al. (2014) identified the viral siRNA population from honeybee colony with colony collapse disorder and indicated those siRNA responses were against viral infection. Recently, Chen et al. (2014) feed siRNA target the viral suppressor of RNAi and significantly suppressed IAPV replication. In our study, a total of 290 vsiRNAs targeted mRNA were related to DNA binding, transcription factor activity, apoptosis, G-protein coupled receptor activity, transporter activity, serine/threonine kinase activity and serine-type endopeptidase, and suggested various vsiRNA could affect the expression of host genes by RNAi pathway. For example, the expression of Dicer-like and Ago2 genes were up-regulated. Especially, vsiRNA_1441 significantly suppressed the expression of extracellular serine/threonine kinase protein FAM20C-like.

Previous studies investigated mainly host response to viral infection. However, metabolites are another important indicator that can reflect physiological response to pathogens (Marelli-Berg et al., 2012). Fatty acid metabolism and biosynthesis are known to play a vital role in viral infections and proliferation (Fritsch and Weichhart, 2016; Wang et al., 2019). It has reported that fatty acids can influence host immune by affecting the inflammatory repertoire of the host, substrates for biosynthesis of inflammatory mediators and activation of cell receptors (Huang et al., 2012). In our study, KEGG analysis showed that 203 DEGs were significantly enriched in fatty acid metabolism and biosynthesis, which was found in accordance with integrated KEGG analysis of DEGs and vsiRNA target mRNA. In addition, two acyl-CoA Delta 11-desaturase-like genes related to fatty acid metabolism and biosynthesis of unsaturated fatty acids were significantly up-regulated after CSBV infection, while two long chain fatty acids genes were significantly down-regulated. It was reported that fatty acids were used to control the American foulbrood and other bee diseases of honeybee (Kuzyšinová et al., 2016). These evidences suggested that fatty acid metabolism and biosynthesis in host immune response could be critical factors to resistance CSBV. However, further studies are needed to clarify the detail process.

According to result from this study combined with previous studies, we propose that CSBV deploys sRNAs to modulate honeybee immune by targeting genes associated with multiple biological processes. One, down-regulated serine/threonine protein kinase and serine-type endopeptidase are key genes of SP cascade and affect melanization response. The other is to down-regulate the expression of acyl-CoA Delta 11-desaturase and long chain fatty acids genes, progress to disruption of the fatty acid biosynthesis and metabolism. Our study offers important insights into understanding the mechanism of pathogenicity of CSBV and may lead to new molecular tools for both viral diagnosis and control. However, further studies will test the functions of SPs and cuticle proteins during the viral infection.

**DATA AVAILABILITY STATEMENT**

The datasets generated or analyzed during the current study are available from the public database, http://gsa.big.ac.cn/ (CRA002310).

**AUTHOR CONTRIBUTIONS**

CH conceived and revised the manuscript. YD, HZ, SS, SY, DY, and SD performed the experiments. YD analyzed the data and wrote the manuscript. All authors contributed to the article and approved the submitted version.

**FUNDING**

This research was funded by the National Natural Science Foundation of China (31572471), the Agricultural Science and Technology Innovation Program (CAAS-ASTIP-2020-IAR), the Fundamental Research Funds for CAAS (Y2020PT17), and Young Talent Program of CAAS, GDAS Special Project of Science, and Technology Development (2018GDASCX-0107).

**ACKNOWLEDGMENTS**

This manuscript has been released as a pre-print at bioRxiv (Deng et al., 2020).

**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fgene.2020.587509/full#supplementary-material

**Supplementary Figure 1** | The correlation analysis \( r = 0.8, P-value = 0.05 \) between qPCR validation and RNA-seq data to identify the reliability of the result.

**Supplementary Figure 2** | Expression level of ppo gene at different day-old after CSBV infection. **P < 0.01**.

**Supplementary Figure 3** | The length distribution of siRNA. (A) The length distribution of known mRNA (B), new identified miRNA (C) and CSBV-specific siRNA (vsiRNA).

**Supplementary Figure 4** | Expression level of genes related to RNAi pathway based on FPKM value.

**Supplementary Table 1** | Primers used for detection the presence of common honey bee viruses.

**Supplementary Table 2** | Primers used to run qPCR for target genes of siRNA.

**Supplementary Table 3** | The list of genes and their annotations from identified miRNA between control and CSBV infected larvae.
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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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