Identification and characterization of melon circular RNAs involved in powdery mildew responses through comparative transcriptome analysis

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ABSTRACT

Circular RNAs (circRNAs) are a class of newly discovered non-coding RNAs that are typically derived from a genome’s exonic, intronic, and intergenic regions. Recent studies of circRNAs in animals and plants have shown that circRNAs are vital in response to various abiotic and biotic stresses. Powdery mildew disease (PM) is a serious fungal disease threatening the melon industry. We performed whole transcriptome sequencing using the leaves of a PM-resistant (M1) and a PM-susceptible (B29) melon to identify circRNAs and determine their molecular functions. A total of 303 circRNAs were identified and >50% circRNAs were derived from exonic regions. Expression levels were significantly altered in 17 and 23 circRNAs after PM infections in B29 and M1, respectively. Melon circRNAs may participate in the response to biotic stimuli, oxidation reduction, metabolic processes, and the regulation of gene expression based on the functional annotation of circRNA parental genes. Furthermore, 27 circRNAs were predicted to be potential targets or ‘sponges’ for 18 microRNAs (miRNAs). Our results are the first to identify and characterize circRNA functions in melon and may contribute to a better understanding of the role and regulatory mechanisms of circRNAs in resisting PM.

INTRODUCTION

Non-coding RNAs (ncRNAs), including microRNA (miRNA), small interfering RNA (siRNA), circular RNA (circRNA), and long noncoding RNA (lncRNA), account for a large portion of the transcriptome sequences in eukaryotic organisms. They have been shown to play important roles in the regulation of protein coding gene expression at transcriptional and post-transcriptional levels (Ponting, Oliver & Reik, 2009; Djebali et al., 2012; Axtell, 2013). CircRNAs are a class of newly discovered endogenous non-coding RNAs characterized by the formation of a closed loop structure, which can be classified as exonic circRNA, intronic circRNA, or intergenic circRNA according to its origin and
position in the genome (Zhang et al., 2013a; Zhang et al., 2013b; Jeck & Sharpless, 2014; Suzuki & Tsukahara, 2014; Chen, 2016). Recent studies indicate that circRNAs are much more stable than linear RNAs and often present distinct expression patterns in specific cell, tissue, and at specific developmental stages (Salzman et al., 2013; Jeck & Sharpless, 2014).

CircRNAs have been studied in humans, mice, mosquitos, and sheep (Salzman et al., 2012; Gruner et al., 2016; Jiang et al., 2014; Li et al., 2017) with little attention given to plant circRNAs. The rapid development of deep-sequencing technology and bioinformatics has led to more circRNAs reported in plants including Arabidopsis, rice, wheat, barley, maize, soybean, cotton, and tomato (Dou et al., 2017; Lu et al., 2015; Wang et al., 2017a; Wang et al., 2017b; Darbani, Noeparvar & Borg, 2016; Chen et al., 2018; Zhao et al., 2017; Xiang et al., 2018; Yin et al., 2018). Many circRNAs have been functionally characterized in model species such as Arabidopsis, rice, and tomato, suggesting that circRNAs play vital roles in multiple biological processes by regulating the expression of their parental genes or acting as miRNA ‘sponges’ to affect the accumulation of target mRNAs. Liu et al. (2017) identified 6,012 circRNAs using publicly available RNA-seq data from Arabidopsis leaves and discovered the function of circRNA in the regulation of the development and senescence of leaves. Tomato is the model plant for studying fruit ripening. Over-expression of a ripening-related circRNA generated from Phytoene Synthase 1 (PSY1) in tomato leads to a significant decrease of PSY1 mRNA abundance and lycopene content (Tan et al., 2017).

CircRNAs have been shown to play important roles in response to various environmental stresses, including biotic and abiotic stresses. Many circRNAs in maize responded to viral infections and phosphate-starvation conditions (Ye et al., 2015; Ghorbani et al., 2018). In wheat, circRNAs can act as ‘sponges’ of corresponding miRNAs to increase resistance to dehydration in wheat seedlings (Wang et al., 2017a; Wang et al., 2017b). Furthermore, 1934 circRNAs in cucumber responded to salt stress. The functional annotation of parental genes revealed that circRNAs may respond to salt stress by mediating signal transcription, metabolism adaptation, and cell cycle and ion homeostasis-related pathways (Zhu et al., 2019). These studies revealed that circRNAs are essential in plant growth and development and aid in their resistance against various environmental stresses. However, information about circRNA has not been reported in melon and little is known about the function of circRNA in resisting powdery mildew disease (PM).

Melon (Cucumis melo L.) is an important fruit crop and is very vulnerable to PM in its later developmental stages. PM can decrease fruit yield and quality and has restricted the development of the melon industry worldwide (Zhang et al., 2013a; Zhang et al., 2013b). In the present study, we selected a PM-resistant and a PM-susceptible cultivated melon to study circRNAs. Comparative transcriptome analysis was performed to identify differentially expressed circRNAs. The parental genes of differentially expressed circRNAs were functionally characterized, which provided a foundation for further analysis of the functional and regulatory mechanisms of circRNAs in resisting PM.
MATERIALS AND METHODS

Plant materials and pathogenic fungus infection

Plant materials were collected from melon seedlings with a highly resistant genotype (M1) and a highly susceptible genotype (B29) to powdery mildew fungus. Melons were grown in a greenhouse on an experimental farm at a temperature of 28 °C/20 °C (day period/night period) and a photoperiod of 16/8 h (day/night). Powdery mildew fungus (*Podosphaera xanthii, race 1*) was collected from the leaves of cultivated melons and was inoculated onto melon seedlings with two or three true leaves at a concentration of $1 \times 10^6$ spores/mL, as previously described (*Cohen, 1993*). The control seedlings were treated with water. The leaves of M1 and B29 treated with water and powdery mildew fungus were harvested at 24 h and 48 h post-inoculation (identified as mock, M24, M48, B24, and B48, respectively). Three leaves were collected from independent seedlings and pooled as a single biological replicate. Three individual biological replicates were prepared for each treatment and were stored at −80 °C for the experiments.

Total RNA extraction and library construction

Trizol reagent was used to extract total RNA from all samples according to the manufacturer’s instructions (Invitrogen, CA, USA). The concentration and quality of RNA was measured using a NanoPhotometer spectrophotometer (IMPLEN, CA, USA) and Bioanalyzer 2100 system (Agilent Technologies, CA, USA). RNA integrity was checked using 1% agarose gel. Three µg RNA per sample was used as input material to construct the library. Ribosomal RNAs were removed using the rRNA Removal Kit (Epicentre, WI, USA) following the manufacturer’s instructions, and linear RNAs were digested using RNase R (Epicentre, WI, USA) at 37 °C for one hour. The remaining RNAs were used as templates for reverse transcription in accordance with the manufacturer’s instructions for the RNA-Seq sample preparation kit (Illumina, San Diego, USA). All libraries were sequenced on the Illumina Hiseq 4000 platform (Novogene, Tianjin, China) with a 150 bp paired-end strand-specific sequencing method.

Bioinformatics analysis and identification of circRNA

To identify circRNA, the raw reads were first filtered using the Fastx-toolkit pipeline (http://hannonlab.cshl.edu/fastx_toolkit/) to remove low-quality reads and to trim the adapter sequences. All clean reads were mapped to the melon reference genome (http://cucurbitgenomics.org/organism/18) using Bowtie2 (v2.2.8; http://bowtie-bio.sourceforge.net/bowtie2/index.shtml), and only uniquely mapped reads with no more than two mismatches were retained for further analysis. We used find_circ (v1.2, https://github.com/marvin-jens/find_circ) and CIRI2 (https://sourceforge.net/projects/ciri/) to analyze and identify the candidate circRNAs with the default parameters (*Gao, Zhang & Zhao, 2018*).

Experimental validation of circRNAs

Total RNAs were extracted using Trizol reagent and treated with DNase I at 37 °C for 30 min according to the manufacturer’s instructions (Invitrogen, CA, USA). Genomic DNA
(gDNA) was isolated using the Plant DNA Mini Kit (TRANSGEN BIOTECH, Beijing, China). A total of 50 µg DNase I-treated RNA was treated with RNase R for 40 min to remove the rRNA. The first-strand cDNA was synthesized from 2 µg of RNase R-treated RNA with random primers using the cDNA Synthesis SuperMix (TRANSGEN BIOTECH, Beijing, China). PCR amplification was conducted using divergent primers designed on the flanking sequences of head-to-tail splicing sites of circRNAs to validate the head-to-tail back-spliced site of circRNAs in melon. Sanger sequencing was performed to further confirm the presence of the back-spliced junction sites. The divergent primers used for circRNA validation are listed in Table S1.

**Quantification of circRNAs and differential expression analysis**

Transcript per million mapped reads (TPM) values were calculated to obtain the expression quantity of all circRNAs. TPM values were calculated based on the read counts mapped to the circRNA. DESeq2 R package (version 2.14, http://www.bioconductor.org/packages/release/bioc/html/DESeq.html) was used for differential expression analysis of circRNAs based on the negative binomial distribution model and the \( P \)-values were adjusted using Benjamini and Hochberg’s approach to control the false discovery rate. circRNAs with differential expression levels were identified using a criterion of adjusted \( P \)-values < 0.05 and \(|\log_2 \text{fold change}| \geq 1\).

**qRT-PCR validation of differentially expressed circRNAs**

Total RNAs were extracted from melon leaves using a Trizol reagent and were treated with DNase I to digest genomic DNA according to the manufacturer’s instructions (Invitrogen, CA, USA). A total of 50 µg DNase I-treated RNA was treated with RNase R for 40 min to remove rRNA. The first-strand cDNA was synthesized from 2 µg of RNase R-treated RNA with random primers using the cDNA Synthesis SuperMix (TRANSGEN BIOTECH, Beijing, China). qRT-PCR was performed to access the relative expression of circRNA using SYBR Green Master Mix (Bio-Rad, Hercules, California). The qRT-PCR procedure was as follows: 95 °C for 5 min, followed by 40 cycles of 95 °C for 5 s, 58 °C for 30 s, and 72 °C for 30 s on an ABI 7500 Real Time PCR system (Applied Biosystems, Waltham, Massachusetts). *CmActin* was used as an internal reference gene. The \( 2^{-\Delta\Delta C_t} \) method was used to calculate the relative expression quantity with three biological replicates and Duncan’s multiple range test was performed to assess whether the results were statistically different \((P < 0.01)\). Primer sequences used in qRT-PCR experiments are listed in Table S1.

**Functional annotation of circRNA parental genes**

The genes producing circRNAs were defined as parental genes. Gene Ontology (GO) enrichment analysis of parental genes of differentially expressed circRNAs was carried out using the GOseq R package (https://bioconductor.org/packages/release/bioc/html/goseq.html), and GO terms with adjusted \( P \)-value < 0.05 were considered to be significantly enriched.
RESULTS

Characterization of the phenotype and resistance against PM of M1 and B29

To explore the mechanism underlying PM resistance in melon at the transcriptome level, high-throughput sequencing and comparative transcriptome analysis were performed between genotypes that were highly resistant (M1) and a highly susceptible (B29) to PM fungus. M1 is a homozygous inbred line with a thick rind and high net density that has been self-pollinating for thirteen generations (Fig. 1A). B29 is a homozygous inbred line, distinct from a cultivated melon, with a smooth, thin rind (Fig. 1B). The genetic background of both lines is highly stable and homozygous. The PM resistance of these two inbred lines was confirmed when we observed the amount of bacterial plaque on the leaves after PM fungus inoculation in the greenhouse. Our results showed different symptoms on the leaves of M1 and B29 7 days after PM infection. No obvious bacterial plaque was observed on the leaves of M1 seven days after PM infection (Fig. 1C), suggesting that M1 is a highly resistant genotype. However, the B29 leaves showed an intense bacterial plaque (Fig. 1D), indicating that B29 is a highly susceptible genotype to PM fungus.

Identification and characterization of circRNAs in melon

To detect circRNAs and explore their functions in resisting against PM disease, 18 rRNA-depleted libraries were generated from the leaves of PM-resistant lines and PM-susceptible lines under both control and PM infected conditions, yielding approximately 100 million reads for each library (Table S2). After the low-quality reads were removed, clean reads were mapped to the reference genome and approximately 83% to 86% of clean reads were uniquely mapped to the melon reference genome in each sample (Table S2). A total of 303 circRNAs were identified using the circRNA identification tool find_circ and CIRI2 a. M1 and B29 shared 261 circRNAs, and only 23 and 19 unique circRNAs were specifically accumulated in M1 and B29, respectively (Fig. 2A). CircRNAs were classified into three groups based on their genomic location and orientation, namely exonic circRNAs, intergenic circRNAs, and intronic circRNAs. Interestingly, in both B29 and M1 melon, exonic circRNAs predominated (54.9% and 54.7%, respectively) compared to intergenic and intronic circRNAs (Figs. 2B and 2C). This was consistent with the conclusion that circRNAs are mainly generated from coding regions in both monocot and dicot plants (Lu et al., 2015; Dou et al., 2017). The length of the circRNAs in melon ranged from 150 to 83,568 nt, but most (86.8%) were <2,000 nt (Fig. S1). To validate the head-to-tail back-spliced site of circRNAs identified in melon, divergent primers were designed for three circRNAs to perform PCR amplification. The PCR products were further analyzed by agarose gel electrophoresis and Sanger sequencing. The results showed that all circRNAs had expected size and validated back-spliced junction sites (Fig. 3), indicating that these circRNA were credible.
Differential expression of circRNAs in response to PM infection and qRT-PCR validation

We quantified the expression of circRNAs in all samples to explore the biological function of melon circRNAs in response to PM infection and found that all melon circRNAs were expressed (TPM > 0) in at least one sample. Their differential expression profiles were evaluated between mock and PM infected samples, and a total of 13, 15, 13, and 13 circRNAs were found to be significantly up-regulated in M24, M48, B24, and B48, respectively. Only two, one, one, and two circRNAs were significantly down-regulated in B24, B48, M24, and M48, respectively (Figs. 4 and 5). Additionally, the expression levels of two and six circRNAs specifically changed in the PM-susceptible melon and PM-resistant melon, respectively (Fig. 4). Furthermore, qRT-PCR results confirmed that the expression changes of Cmcirc0260, Cmcirc0881, Cmcirc1283, Cmcirc2417, Cmcirc3971, and Cmcirc4093 in the PM-resistant melon were more noticeable than that in the PM-susceptible melon after PM infection, which was consistent with the transcriptome profiling results (Fig. 6).

Functional characterization of parental genes of differentially expressed circRNAs

The molecular function of most circRNAs is not well defined, although some circRNAs have been shown to participate in the regulation of their parental gene expression. To explore the putative function of melon circRNAs, we identified the parental genes of melon
circRNAs from the location where the circRNAs derived in the genome. One hundred and fifty-two parental genes were obtained for the 23 differentially expressed circRNAs and GO categories analyses were performed to explore their functions. The circRNA parental genes were mainly involved in the biological processes of gene expression, oxidation–reduction process, single-organism cellular process, response to biotic stimulus, and pathogenesis. Gene ontology of cellular components of parental genes showed that most of the gene products are cell parts, intracellular parts, membrane parts, and the macromolecular complex. The enriched GO terms for molecular function included catalytic activity, oxidoreductase activity, peroxidase activity, kinase activity, and protein binding (Table 1).

**Identification of CircRNAs act as miRNA targets or ‘sponges’**

circRNAs have been shown to interact with miRNAs to prevent them from targeting mRNAs and therefore may control target gene expression (Hansen et al., 2013). Binding between miRNA and circRNA is based on the complementarity between their nucleotide sequences. Many circRNAs can be cleaved by miRNA and were named as miRNA targets.
Figure 3  Experimentally validation of Cmcirc0260 (A), Cmcirc0881 (B) and Cmcirc2417 (C). Divergent primers were used for amplification of the cDNA and gDNA. Agarose gel electrophoresis and Sanger sequencing were performed to examine the size and sequence of PCR product.
However, many circRNAs can compete with mRNA to bind to specific miRNA without cleavage, thus blocking cleavage and alleviating the repression of its target mRNA. These circRNAs were called miRNA 'sponges'. To identify circRNAs that may act as targets or 'sponges' of miRNA, we searched for the circRNAs and their complementary sequences against the melon miRNAs using the BLAST algorithm and found that 27 circRNAs can be bound by 18 miRNAs (Table 2). We predicted that some highly conserved miRNAs, such as miR156, miR159, miR161 and miR172, that had been functionally characterized, could be targeted by specific circRNAs. Of these circRNAs, some had more than one miRNA binding sites. For instance, Cmcirc03865 can be targeted by CmmiR419, CmmiR5021, and CmmiR5658. In addition, specific miRNA in melon could bind to different circRNA. For example, CmmiR5021 could bind with 11 different circRNAs and CmmiR5998 could bind to four different circRNAs.

**DISCUSSION**

Recently, the identification and functional characterization of circRNAs has been widely reported in both animals and plants, demonstrating that circRNAs participate in multiple
Clustering and differential expression patterns of 23 melon circRNAs after PM infection. The expression values were measured as TPM and presented as log2 (TPM value +1). Biological processes (Memczak et al., 2013; Chu et al., 2018). However, the molecular functions and regulatory mechanisms of circRNAs underlying PM resistance in melon is largely unknown. High-throughput sequencing and comparative transcriptome analysis were performed to identify putative circRNAs and their target genes involved in PM resistance. A total of 303 circRNAs were detected from the melon leaf. The number of circRNAs in melon was lower than in Arabidopsis (6,012), rice (12,037), cucumber (2,787), and soybean (5,372), which may be attributed to our more restrictive filter conditions for circRNA identification and the use of only leaf tissue for sequencing. Previous studies revealed that circRNA was expressed in a highly tissue- or stage-specific manner in most organisms (Tong et al., 2018).

It has been reported that circRNA can be generated from exons, introns, and intergenic regions of the genome and are named exonic circRNA, intronic circRNA, or intergenic circRNA, respectively (Zhang et al., 2013a; Zhang et al., 2013b; Jeck & Sharpless, 2014). In
Figure 6  Experimental validation of six differential expressed circRNAs by qRT-PCR. (A) Cmcirc0260, (B) Cmcirc0881, (C) Cmcirc21283, (D) Cmcirc2417, (E) Cmcirc3971, (F) Cmcirc4093. CmActin was selected as internal reference. The relative expression quantity of circRNAs was normalized to that in mock. The red broken lines represent the RNA-seq values presented as log2 (TPM value + 1). Error bars indicate ± SD of three biological replicates. Asterisks indicated a significant change (*P < 0.05; **P < 0.01) between different samples.

Table 1  GO enrichment analysis and functional categories of circRNA parental genes in melon.

| GO:accession | Functional category | Term type       | P-value     | Gene count |
|--------------|---------------------|-----------------|-------------|------------|
| GO:0004096   | catalase activity   | molecular_function | 2.28E−03   | 34         |
| GO:0004601   | peroxidase activity | molecular_function | 5.30E−04   | 9          |
| GO:0016684   | Oxido-reductase activity | molecular_function | 5.30E−04   | 6          |
| GO:0005515   | protein binding     | molecular_function | 6.62E−05   | 26         |
| GO:0003824   | kinase activity     | molecular_function | 9.01E−04   | 15         |
| GO:0032991   | macromolecular complex | cellular_component | 6.99E−05   | 9          |
| GO:0044425   | membrane part       | cellular_component | 7.61E−04   | 11         |
| GO:0044424   | intracellular part  | cellular_component | 8.95E−05   | 5          |
| GO:0044464   | cell part           | cellular_component | 9.22E−04   | 17         |
| GO:0009607   | response to biotic stimulus | biological_process | 8.62E−05   | 12         |
| GO:0009405   | pathogenesis        | biological_process | 1.14E−05   | 5          |
| GO:0006952   | defense response    | biological_process | 1.15E−04   | 6          |
| GO:0055114   | oxidation–reduction process | biological_process | 2.18E−05   | 4          |
| GO:0044710   | single-organism metabolic process | biological_process | 4.61E−04   | 9          |
| GO:0010467   | gene expression     | biological_process | 4.95E−05   | 5          |
| GO:0044763   | single-organism cellular process | biological_process | 6.71E−04   | 18         |

In our study, exonic circRNAs were predominant (approximately 54.9%) compared to the intergenic (approximately 10%) and intronic (approximately 30%) circRNAs (Fig. 2). These proportions are similar to those in Arabidopsis, rice, cucumber, and tomato (Lu et al., 2015; Dou et al., 2017; Yin et al., 2018). In contrast, approximately 51% and 55% of the circRNAs were intergenic circRNAs in wheat and kiwifruit, which may be due to
Table 2  Predicted interaction between circRNAs and miRNA in melon.

| CircRNA ID  | miRNA ID  | E-value  | Sequence complementarity (%) |
|-------------|-----------|----------|------------------------------|
| Cmcirc00352 | CmmiR5021| 2.15E−04 | 100                          |
| Cmcirc00352 | CmmiR8170| 6.36E−03 | 98.45                        |
| Cmcirc00422 | CmmiR855 | 1.46E−04 | 90.28                        |
| Cmcirc00429 | CmmiR865 | 4.80E−04 | 94.42                        |
| Cmcirc00464 | CmmiR845 | 8.96E−04 | 100                          |
| Cmcirc00713 | CmmiR5021| 9.63E−05 | 93.24                        |
| Cmcirc00840 | CmmiR447 | 9.85E−04 | 100                          |
| Cmcirc00916 | CmmiR5021| 1.59E−04 | 91.02                        |
| Cmcirc01282 | CmmiR3434| 4.56E−04 | 100                          |
| Cmcirc01283 | CmmiR417 | 2.72E−04 | 88.95                        |
| Cmcirc01290 | CmmiR161 | 1.48E−05 | 94.26                        |
| Cmcirc01351 | CmmiR161 | 1.80E−04 | 89.95                        |
| Cmcirc01495 | CmmiR5021| 1.45E−05 | 96.66                        |
| Cmcirc01695 | CmmiR5658| 6.95E−04 | 89.69                        |
| Cmcirc01695 | CmmiR5998| 5.48E−04 | 87.58                        |
| Cmcirc01723 | CmmiR156 | 5.70E−05 | 100                          |
| Cmcirc01723 | CmmiR5658| 2.00E−05 | 93.28                        |
| Cmcirc01736 | CmmiR5021| 2.37E−04 | 95.78                        |
| Cmcirc01736 | CmmiR5658| 8.70E−06 | 87.94                        |
| Cmcirc02030 | CmmiR159 | 7.57E−06 | 100                          |
| Cmcirc02063 | CmmiR5021| 4.76E−04 | 95.21                        |
| Cmcirc02076 | CmmiR5021| 3.37E−06 | 90.14                        |
| Cmcirc02152 | CmmiR5021| 5.90E−04 | 90.55                        |
| Cmcirc02438 | CmmiR5021| 4.79E−04 | 88.56                        |
| Cmcirc02855 | CmmiR5021| 4.23E−05 | 97.84                        |
| Cmcirc02891 | CmmiR417 | 4.79E−05 | 93.45                        |
| Cmcirc02905 | CmmiR5633| 1.78E−04 | 86.58                        |
| Cmcirc03115 | CmmiR172 | 1.45E−03 | 100                          |
| Cmcirc03503 | CmmiR779 | 6.67E−04 | 93.89                        |
| Cmcirc03756 | CmmiR414 | 3.78E−04 | 97.25                        |
| Cmcirc03865 | CmmiR419 | 4.00E−05 | 87.56                        |
| Cmcirc03865 | CmmiR5021| 8.89E−04 | 90.26                        |
| Cmcirc03865 | CmmiR5658| 4.78E−03 | 96.55                        |

wheat’s large genome size with comparably fewer annotated genes (Wang et al., 2017a; Wang et al., 2017b). It has been deemed that the intergenic regions of a genome are more likely to contain missed genes that have not been annotated. Meanwhile, most of the total circRNAs were intronic circRNAs in soybean, which may be attributed to the genome duplication event responsible for generating multiple gene copies (Zhao et al., 2017). Our results indicated that the molecular basis of circular RNA biogenesis in plants varies among different species.
Differential expression analysis of circRNAs was conducted between mock and PM-inoculated samples of both PM-resistant and PM-susceptible genotypes. A total of 23 circRNAs were found to be responsive to PM infection, suggesting that these circRNAs may play important roles in PM resistance. RNA-seq data and qRT-PCR results confirmed that six circRNAs, including Cmcirc0260, Cmcirc0881, Cmcirc1283, Cmcirc2417, Cmcirc3971, and Cmcirc4093 in PM-resistant melon were more highly induced than that in PM-susceptible melon after PM infection, suggesting that a different disease response mechanism might exist between these melon genotypes. The functional annotation results showed that a large number of parental genes of circRNAs were involved in multiple redox processes such as the peroxidase gene and the glutathione reductase gene. Peroxidase and glutathione reductase play an essential role in the ROS scavenging pathway to prevent oxidative damage during the process of pathogen infection (Cui et al., 2017). The overexpression of a glutathione reductase gene (SIGRE21) prevented ROS accumulation and enhanced resistance against P. infestans in tomato (Cui et al., 2017). We also identified eight parental genes that encode pathogen-related proteins and the LRR receptor, implying that circRNAs could regulate the gene expression to resist against PM infection in melon.

Previous reports have shown that circRNAs can regulate gene expression in a miRNA-mediated manner. They may act as targets or ‘sponges’ of miRNAs to restrict the cleavage of target genes mediated by miRNA and thus promote target gene expression (Hansen et al., 2013). Similar to the results in Arabidopsis, cotton, and other plants, many melon circRNAs are thought to be miRNA ‘sponges’. Twenty-seven circRNAs were predicted to be potential targets of 18 family miRNAs including miR159, miR161, miR417, miR419, miR845, miR5021, and others. A few miRNAs including miR845, miR855, miR5021, and miR5658 were confirmed to be involved in the response to various biotic and abiotic stresses (Borges et al., 2018). Our results suggested that direct interactions between miRNAs and circRNAs may also exist in melon, providing new information for further investigation into the function and mechanisms of circRNAs in PM resistance.

**CONCLUSIONS**

Our study was the first to report the existence of abundant circRNAs in the melon leaf and to characterize their possible regulatory roles in response to PM disease. Nevertheless, it should be noted that only leaf tissue was included in our study, which may limit the amount and properties of circRNAs in melon that we were able to find. Future studies should be conducted to detect circRNAs from more tissues and various developmental stages and to determine the mechanism of melon circRNAs in resisting PM. Our findings improved the understanding of circRNAs in cucurbitaceae species and provided new insights into the regulatory roles of circRNAs in PM resistance.
ADDITIONAL INFORMATION AND DECLARATIONS

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Competing Interests
The authors declare there are no competing interests.

Author Contributions
• Jianlei Sun performed the experiments, prepared figures and/or tables, and approved the final draft.
• Yumei Dong performed the experiments, analyzed the data, prepared figures and/or tables, and approved the final draft.
• Chongqi Wang analyzed the data, prepared figures and/or tables, and approved the final draft.
• Shouhua Xiao analyzed the data, authored or reviewed drafts of the paper, contributed analysis tools, and approved the final draft.
• Zigao Jiao and Chao Gao conceived and designed the experiments, authored or reviewed drafts of the paper, and approved the final draft.

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The raw data are available in the NCBI Sequence Read Archive database: SRR9129105–SRR9129122.

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REFERENCES

Axtell MJ. 2013. Classification and comparison of small RNAs from plants. *Annual Review of Plant Biology* 64:137–159 DOI 10.1146/annurev-arplant-050312-120043.

Borges F, Parent J, Van Ex F, Wolff P, Martinez G, Köhler C, Martienssen R. 2018. Transposon-derived small RNAs triggered by miR845 mediate genome dosage response in Arabidopsis. *Nature* 50(2):186–192 DOI 10.1038/s41588-017-0032-5.

Chen LL. 2016. The biogenesis and emerging roles of circular RNAs. *Nature Reviews Molecular Cell Biology* 17:205–211 DOI 10.1038/nrm.2015.32.

Chen L, Zhang P, Fan Y, Lu Q, Li Q, Yan J, Muehlbauer GJ, Schnable PS, Dai M, Li L. 2018. Circular RNAs mediated by transposons are associated with transcriptomic and phenotypic variation in maize. *New Phytologist* 21(2):299–306 DOI 10.1111/nph.14901.

Chu Q, Shen E, Ye C, Fan L, Zhu Q. 2018. Emerging roles of plant circular RNAs. *The Plant Cell* 1(1):1–14.

Cohen R. 1993. A leaf disk assay for detection of resistance of melons to *Sphaerotheca fuliginea* race 1. *Plant Disease* 77:513–517 DOI 10.1094/PD-77-0513.

Cui J, Luan Y, Jiang N, Bao H, Meng J. 2017. Comparative transcriptome analysis between resistant and susceptible tomato allows the identification of IncRNA16397 conferring resistance to *Phytophthora infestans* by co-expressing glutaredoxin. *Plant Journal* 89(3):577–589 DOI 10.1111/tpj.13408.

Darbani B, Noeparvar S, Borg S. 2016. Identification of circular RNAs from the parental genes involved in multiple aspects of cellular metabolism in barley. *Frontiers in Plant Science* 7:776 DOI 10.3389/fpls.2016.00776.

Djebali S, Davis CA, Merkel A, Dobin A, Lassman T, Mortazavi A. 2012. Landscape of transcription in human cells. *Nature* 489(7414):101–108 DOI 10.1038/nature11233.

Dou Y, Li S, Yang W, Liu K, Du Q, Ren G, Yu B, Zhang C. 2017. Genome-wide discovery of circular RNAs in the leaf and seedling tissues of Arabidopsis thaliana. *Current Genomics* 18(4):360–365 DOI 10.2174/1389202918666170307161124.

Gao Y, Zhang J, Zhao F. 2018. Circular RNA identification based on multiple seed matching. *Briefings in Bioinformatics* 19(5):803–810 DOI 10.1093/bib/bby014.

Ghorbani A, Izadpanah K, Peters JR, Dietzgen RG, Mitter N. 2018. Detection and profiling of circular RNAs in uninfected and maize Iranian mosaic virus-infected maize. *Plant Science* 274:402–409 DOI 10.1016/j.plantsci.2018.06.016.

Gruner H, Cortés-López M, Cooper DA, Bauer M, Miura P. 2016. CircRNA accumulation in the aging mouse brain. *Scientific Reports* 6:38907 DOI 10.1038/srep38907.

Hansen T, Jensen T, Clausen B, Bramsen J, Finsen B, Damgaard C, Kjems J. 2013. Natural RNA circles function as efficient microRNA sponges. *Nature* 495(7441):384–388 DOI 10.1038/nature11993.
Jeck WR, Sharpless NE. 2014. Detecting and characterizing circular RNAs. *Nature Biotechnology* 32(5):453–461 DOI 10.1038/nbt.2890.

Jiang X, Peery A, Hall AB, Sharma A, Chen XG, Waterhouse RM, Komissarov A, Riehle MM, Shouche Y, Sharakhova MV, Lawson D, Pakpour N, Arensburger P, Davidson VL, Eiglmeier K, Emrich S, George P, Kennedy RC, Mane SP, Maslen G, Oringanje C, Qi Y, Settlage R, Tojo M, Tubio JM, Unger MF, Wang B, Vernick KD, Ribeiro JM, James AA, Michel K, Riehle MA, Luckhart S, Sharakhov IV, Tu Z. 2014. Genome analysis of a major urban malaria vector mosquito. *Anopheles stephensi*. *Genome Biology* 15(9):459 DOI 10.1186/s13059-014-0459-2.

Li C, Li X, Yao Y, Ma Q, Ni W, Zhang X, Cao Y, Hazi W, Wang D, Quan R, Hou X, Liu Z, Zhan Q, Liu L, Zhang M, Yu S, Hu S. 2017. Genome-wide analysis of circular RNAs in prenatal and postnatal muscle of sheep. *Oncotarget* 8(57):97165–97177 DOI 10.3389/fgene.2013.00307.

Liu T, Zhang L, Chen G, Shi T. 2017. Identifying and characterizing the circular RNAs during the lifespan of Arabidopsis leaves. *Frontiers in Plant Science* 8:1278 DOI 10.3389/fpls.2017.01278.

Lu T, Cui L, Zhou Y, Zhu C, Fan D, Gong H, Zhao Q, Zhou C, Zhao Y, Lu D, Luo J, Wang Y, Tian Q, Feng Q, Huang T, Han B. 2015. Transcriptome-wide investigation of circular RNAs in rice. *RNA* 21(12):2076–2087 DOI 10.1261/rna.052282.115.

Memczak S, Jens M, Elefsinioti A, Torti F, Krueger J, Rybak A, Maier L, Mackowiak S, Gregersen L, Munschauer M, Loewer A, Ziebold U, Landthaler M, Kocks C, Le Noble F, Rajewsky N. 2013. Circular RNAs are a large class of animal RNAs with regulatory potency. *Nature* 495(7441):333–338 DOI 10.1038/nature11928.

Ponting CP, Oliver PL, Reik W. 2009. Evolution and functions of long noncoding RNAs. *Cell* 136(4):629–641 DOI 10.1016/j.cell.2009.02.006.

Salzman J, Chen RE, Olsen MN, Wang PL, Brown PO. 2013. Cell-type specific features of circular RNA expression. *PLOS Genetics* 9(9):e1003777 DOI 10.1371/journal.pgen.1003777.

Salzman J, Gawad C, Wang PL, Lacayo N, Brown PO. 2012. Circular RNAs are the predominant transcript isoform from hundreds of human genes in diverse cell types. *PLOS ONE* 7(2):e30733 DOI 10.1371/journal.pone.0030733.

Suzuki H, Tsukahara T. 2014. A view of pre-mRNA splicing from RNase R resistant RNAs. *International Journal of Molecular Sciences* 15(6):9331–9342 DOI 10.3390/ijms15069331.

Tan J, Zhou Z, Niu Y, Sun X, Deng Z. 2017. Identification and functional characterization of tomato circRNAs derived from genes involved in fruit pigment accumulation. *Scientific Reports* 7(1):8594 DOI 10.1038/s41598-017-08806-0.

Tong W, Yu J, Hou Y, Li F, Zhou Q, Wei C, Bennetzen J. 2018. Circular RNA architecture and differentiation during leaf bud to young leaf development in tea (Camellia sinensis). *Planta* 248(6):1417–1429 DOI 10.1007/s00425-018-2983-x.

Wang Z, Liu Y, Li D, Li L, Zhang Q, Wang S, Huang H. 2017b. Identification of circular RNAs in kiwifruit and their species-specific response to bacterial canker pathogen invasion. *Frontiers in Plant Science* 8:413 DOI 10.3389/fpls.2017.00413.
Wang Y, Yang M, Wei S, Qin F, Zhao H, Suo B. 2017a. Identification of circular RNAs and their targets in leaves of Triticum aestivum L. under dehydration stress. Frontiers in Plant Science 7:2024 DOI 10.3389/fpls.2016.02024.

Xiang L, Cai C, Cheng J, Wang L, Wu C, Shi Y, Luo J, He L, Deng Y, Zhang X, Yuan Y, Cai Y. 2018. Identification of circular RNAs and their targets in Gossypium under Verticillium wilt stress based on RNA-seq. PeerJ 6:e4500 DOI 10.7717/peerj.4500.

Ye CY, Chen L, Liu C, Zhu QH, Fan L. 2015. Widespread noncoding circular RNAs in plants. New Phytologist 193:1049–1063 DOI 10.1111/j.1469-8137.2011.04006.x.

Yin J, Liu M, Ma D, Wu J, Li S, Zhu Y, Han B. 2018. Identification of circular RNAs and their targets during tomato fruit ripening. Postharvest Biology and Technology 136:90–98 DOI 10.1016/j.postharvbio.2017.10.013.

Zhang CQ, Ren Y, Guo SG, Zhang HY, Gong GY, Du YC, Xu Y. 2013a. Application of comparative genomics in developing markers tightly linked to the Pm-2F gene for powdery mildew resistance in melon (Cucumis melo L.). Euphytica 190:157–168 DOI 10.1007/s10681-012-0828-4.

Zhang Y, Zhang XO, Chen T, Xiang JF, Yin QF, Xing YH, Zhu SS, Yang L, Chen LL. 2013b. Circular intronic long noncoding RNAs. Molecular Cell 51(6):792–806 DOI 10.1016/j.molcel.2013.08.017.

Zhao W, Cheng YH, Zhang C, You QB, Shen XJ, Guo W, Jiao YQ. 2017. Genome-wide identification and characterization of circular RNAs by high throughput sequencing in soybean. Scientific Reports 7(1):5636 DOI 10.1038/s41598-017-05922-9.

Zhu Y, Jia J, Yang L, Xia Y, Zhang H, Jia J, Zhou R, Nie P, Yin J, Ma D, Liu L. 2019. Identification of cucumber circular RNAs responsive to salt stress. BMC Plant Biology 19(1):164 DOI 10.1186/s12870-019-1712-3.