Small RNA and Degradome Deep Sequencing Reveals the Roles of microRNAs in Seed Expansion in Peanut (Arachis hypogaea L.)

Xingli Ma, Xingguo Zhang, Kunkun Zhao, Fapeng Li, Ke Li, Longlong Ning, Jialin He, Zeyu Xin* and Dongmei Yin*

College of Agronomy, Henan Agricultural University, Zhengzhou, China

Seed expansion in peanut is a complex biological process involving many gene regulatory pathways. MicroRNAs (miRNAs) play important regulatory roles in plant growth and development, but little is known about their functions during seed expansion, or how they contribute to seed expansion in different peanut lines. We examined seed miRNA expression patterns at 15 and 35 days after flowering (DAF) in two peanut eighth-generation recombinant inbred lines (RILs); 8106, a medium-pod variety, and 8107, a super-pod variety. Using high-throughput sequencing, we identified 1,082 miRNAs in developing peanut seeds including 434 novel miRNAs. We identified 316 differentially expressed miRNAs by comparing expression levels between the two peanut lines. Interestingly, 24 miRNAs showed contrasting patterns of expression in the two RILs, and 149 miRNAs were expressed predominantly in only one RIL at 35 DAF. Also, potential target genes for some conserved and novel miRNAs were identified by degradome sequencing; target genes were predicted to be involved in auxin mediated signaling pathways and cell division. We validated the expression patterns of some representative miRNAs and 12 target genes by qPCR, and found negative correlations between the expression level of miRNAs and their targets. miR156e, miR159b, miR160a, miR164a, miR166b, miR168a, miR171n, miR172c-5p, and miR319d and their corresponding target genes may play key roles in seed expansion in peanut. The results of our study also provide novel insights into the dynamic changes in miRNAs that occur during peanut seed development, and increase our understanding of miRNA function in seed expansion.

Keywords: peanut, deep sequencing, miRNA, degradome, seed expansion

INTRODUCTION

As one of the most important oil crops in the world, peanut (Arachis hypogaea L.) is grown on six continents but mainly in Asia, Africa, and America (Zhang et al., 2012; Zhao et al., 2015). Yield of peanut is determined primarily by two vital factors; (1) seed expansion, and (2) subsequent seed filling (Stanciel et al., 2000; Ramakrishna et al., 2006). However, compared to other food crops, peanuts show a distinct pattern of seed development. After fertilization, a new organ called the peg differentiates from the ovary (Brennan, 1969). Little mitotic division occurs in the embryo or endosperm during active geotropic peg growth until the peg penetrates the soil.
Rapid embryonic cell division then begins a few days later. This occurs approximately 10–12 days following fertilization (Smith, 1956). Seed developmental changes in water content, nucleic acid levels, enzyme activities, and storage-protein deposition patterns were studied from the earliest stage when kernels can be removed (Aldana et al., 1972; Basha and Cherry, 1976; Yamada et al., 1980). It has been shown that developmentally expressed genes in seeds are regulated at the level of transcription initiation (Falvey and Schibler, 1991). In addition, some important genes that contribute highly to seed development have also been identified in peanut (Bi et al., 2010; Zhang et al., 2012). Although some seed development genes have been studied in peanut, the roles of the upstream regulators of these coding genes, such as miRNAs, have not been well studied.

MicroRNAs (miRNAs) are a class of small (19–24 nt), non-coding RNAs that can regulate gene expression by cleavage or translational repression of the target gene mRNAs (Van Ex et al., 2011). miRNAs have been shown to play important roles in plant growth, development, and the response to environmental stresses (Sunkar and Zhu, 2004; Ding et al., 2012; Feng et al., 2014; Zhao et al., 2016). For example, in Arabidopsis, 33 different miRNA families have been detected in mature pollen, and several (miR156, miR2939, miR158, and miR845) showed elevated expression levels in pollen compared with leaves (Grant-Downton et al., 2009). In rice, miR167 may play a role in rice grain filling through the auxin-miR167-ARF8-OsGH3.2 regulatory pathway (Xue et al., 2009); miR397 is highly expressed in young panicles and grains (Zhang et al., 2013). In peanut, studies on miRNAs also have been reported (Zhao et al., 2010; Chi et al., 2011), and several miRNAs (miR160, miR164, miR393, miR396, miR397, miR482, and miR2118) have possible roles in the response to pathogen infection (Zhao et al., 2015). Although some miRNAs have been identified in peanut, to date, the mechanism of seed expansion involving miRNAs is unclear.

To understand the role of miRNAs in peanut seed expansion, we used two representative peanut lines in this study; 8106 and 8107 are eighth-generation recombinant inbred lines (RILs) from a cross between cultivars of Huayou7 (female) and Huayou4 (male), both of which are erect Virginia-types with 8–10 branches. The main difference between the two RILs is the pod size: line 8106 has medium-sized pods (3.2 cm long × 1.3 cm wide), and a 100-seed weight of 100 g, while line 8107 has large pods (5.5 cm wide), and a 100-seed weight of 182 g. These two typical peanut lines were investigated from the earliest stage when kernels can be removed (15 DAF) and 35 DAF. Four samples were frozen immediately in liquid nitrogen and stored at −80°C until use. Three replicates of samples were collected in this study.

Small RNA Library Construction and DNA Sequencing

Total RNA was extracted using TRK-1001 following the manufacturer’s instructions. RNA quantity and purity were assessed using a Bioanalyzer 2100 and the RNA 6000 Nano LabChip Kit (Agilent Technologies, Santa Clara, CA, United States) with RIN number > 7.0. Total RNA was ligated to the RNA 3′ and RNA 5′ adapters and reverse transcription, followed by PCR, was performed to make cDNA constructs of the small RNAs. The small cDNA fractions that ranged from 22 to 30 nt in length were then isolated via 6% denaturing polyacrylamide gel electrophoresis. Finally, the cDNA constructs were purified, and the library was validated. We then performed single-end sequencing (50 bp) on an Illumina HiSeq2500 at the LC-BIO (Hangzhou, China) following the vendor’s recommended protocol.

Identification of Known and Potential Novel miRNAs

The raw sequencing reads were subjected to the Illumina pipeline filter (Solexa 0.3), and the dataset was then further processed using an in-house program, ACGT101-miR (LC Sciences, Houston, TX, United States) to remove adapter dimers, low complexity reads, common RNA families (rRNA, tRNA, snRNA, snoRNA), repeats in the NCBI GenBank1, Repbase2, and Rfam3 databases. Subsequently, the remaining non-annotated sequences were mapped to specific species precursors in miRBase 21.04 by BLAST searches to identify known miRNAs. On the basis of sequence similarity to peanut genome, these miRNA sequences were classified into subgroups 1a, 1b, 2a, 2b, and 3 with high-to-mid confidence in order (Ambros et al., 2003; Li et al., 2017). The classification criteria for each subgroup were as follows: in gp1a, reads were mapped to miRNAs/pre-miRNAs of specific species in miRBase and the pre-miRNAs were further mapped to genome (known miRNAs); in gp1b, reads were mapped to (except for specific species) miRNAs/pre-miRNAs of selected species in miRBase and the pre-miRNAs were further mapped to genome (conserved miRNAs); in gp2a, reads were mapped to miRNAs/pre-miRNAs of selected species in miRBase and the mapped pre-miRNAs were not further mapped to a

---

1. [http://www.ncbi.nlm.nih.gov/Genbank/](http://www.ncbi.nlm.nih.gov/Genbank/)
2. [http://www.girinst.org/repbase/update/index.html](http://www.girinst.org/repbase/update/index.html)
3. [http://www.sanger.ac.uk/software/Rfam](http://www.sanger.ac.uk/software/Rfam)
4. [http://www.mirbase.org](http://www.mirbase.org)
genome, but the reads (and of course the miRNAs of the pre-miRNAs) were mapped to a genome. The extended genome sequences from the genomic loci may form hairpins (conserved miRNAs); in gp2b, reads were mapped to miRNAs/pre-miRNAs of selected species in miRbase and the mapped pre-miRNAs were not further mapped to a genome, but the reads (and of course the miRNAs of the pre-miRNAs) were mapped to a genome. The extended genome sequences from the genomic loci may not form hairpins (conserved miRNAs); in gp3, reads were mapped to miRNAs/pre-miRNAs of selected species in miRbase and the mapped pre-miRNAs were not further mapped to a genome, and the reads were also not mapped to a genome (conserved miRNAs). All non-annotated reads with lengths of 18–25 nt were mapped to the reference genome sequences of two diploid ancestors of cultivated peanut, *A. ipaensis* and *A. duranensis* using the Bowtie package; only perfectly matched sRNAs were used in further analyses. Novel miRNAs were identified using the MIREAP software based on their precursors, and the hairpin RNA structures containing sequences were predicted from the flanking 120 nt sequences using RNAfold software. The key criteria for miRNA prediction were based on those that had been previously reported in the literature (Meyers et al., 2008).

### Analysis of Differentially Expressed miRNAs

The raw reads for each small RNA sequence were first normalized using global normalization procedures (Huber et al., 2002), and the differentially expressed sequence counts at 15 and 35 DAF were then analyzed using the online web service IDEG6. Because we included three biological replicates in the sequencing, each miRNA derived from 15 and 35 DAF seeds were compared using the T-Test. Only p-values <0.05 were considered to be differentially expressed miRNAs.

### Reverse Transcription Reactions

Reverse transcription reactions were performed using the One Step PrimeScript miRNA cDNA Synthesis Kit (TaKaRa Co., Tokyo, Japan) following the manufacturer’s instructions. First-strand cDNA was synthesized in 20-µl reaction volumes containing 1 µg total RNA, 10 µl 2X miRNA reaction buffer mix, 2 µl 0.1% BSA, 2 µl miRNA PrimeScript RT enzyme mix, and RNase-free dH2O (to 20 µl). The reactions were incubated at 37°C for 60 min, followed by 85°C for 5 min, and then stored at −20°C until use.

### Validation of Differentially Expressed miRNAs

The qPCR reactions were performed with a SYBR PrimeScript miRNA RT-PCR Kit on a Fluorescence detection system (TianGen Biotech, Beijing, China). Each 20 µl reaction contained 1 µl cDNA template (∼100 ng), 1 µl 10 µM PCR forward primer, 1 µl 10 µM Uni-miR qPCR primer, 10 µl 2× SYBR premix EX TaqII, and 7 µl ddH2O. The amplification reactions were performed by first incubating at 95°C for 5 min, followed by 45 cycles of 95°C for 15 s, 60°C for 30 s, and 72°C for 45 s. Following amplification, a threshold was set and the threshold cycle (CT) was recorded automatically. All reactions were performed in triplicate for each sample. The relative expression levels of the miRNAs were calculated using the 2−ΔΔCT method (Livak and Schmittgen, 2001) and the data were normalized to the CT values for the *Actin* gene. The primer sequences for 16 differentially expressed miRNAs are given in Supplementary Table S13-1.

### Degradome Library Construction and Target Identification

Total RNA was extracted using TRK1001 following the manufacturer’s instructions. The total RNA quantity and purity were determined using a Bioanalyzer 2100 and RNA 6000 Nano
LabChip Kit (Agilent, Santa Clara, CA, United States) with RIN number > 7.0. Approximately 20 µg of total RNA was used to prepare the Degradome library. The method differed considerably from previous efforts (Addo-Quaye et al., 2008, 2009b) and followed the method of (Ma et al., 2010) with some modifications: (1) the poly(A)+ RNA was bound to the mRNA Capture Beads, (2) the 5’ adapter was ligated to only those RNAs containing 5’-monophosphates, (3) reverse transcription was performed using 3’-random primers, (4) the library was PCR amplified, and (5) the libraries were sequenced using the 5’ adapter only.

The purified cDNA library was used for cluster generation on Illumina’s Cluster Station and then sequenced on the Illumina Hisseq 2500 instrument following the manufacturer’s instructions. The extracted sequencing reads were stored in file SampleA_RawData.txt and were then used in e standard data analysis, which is described in Mao et al. (2012). A publicly available software package, CleaveLand3.0, was used to analyze the sequencing data generated. The key functions performed by this software and the relevant analysis results are described by Addo-Quaye et al. (2009a).

Gene ontology (GO) and KEGG analysis of target genes

Gene ontology (GO) annotations of target genes corresponding to the differentially expressed miRNAs were downloaded from the Gene Ontology8, NCBI11, and UniProt12. The KEGG13 pathway was analyzed through the ClueGO plug-in14 and Cytoscape software V2.8.25 to identify the significant pathways of the differential genes. GO terms and KEGG pathways were regarded to be significantly enriched with the corrected P-value ≤ 0.05, which was calculated using hypergeometric test (Hou et al., 2017).

Validation of the target genes

Expression analysis of selected target genes was performed by qPCR. First-strand cDNA was synthesized from 1 µg of RNA using TransScript First-Strand cDNA Synthesis SuperMix (TransGen Co., Beijing, China) following the manufacturer’s instructions. Each 20 µl qPCR reaction contained 1 µl cDNA, 0.5 µl 10 µM forward primer, 0.5 µl 10 µM reverse primer, 10 µl 2× TransStart Top Green qPCR SuperMix, and 8 µl double-distilled water. The amplification conditions were an initial denaturation step of 95°C for 5 min, followed by 45 cycles of 95°C for 15 s, 56°C for 30 s, and 72°C for 45 s. All reactions were performed in triplicate for each sample. Relative expression levels of the targets were quantified using the 2−ΔΔCT method (Livak and Schmittgen, 2001) normalized to Act1 CT values. The sequences of the primer pairs for the target genes are given in Supplementary Table S13-2.

RESULTS

Sequencing and Annotation of Peanut miRNAs

Lines 8106 and 8107 showed obvious difference in seed size (Figure 1). To investigate the dynamic variation of miRNAs during peanut seed expansion, two seed developmental stages, 15 DAF and 35 DAF, were selected to sequence the small RNAs using Solexa high-throughput sequencing technology. From RIL 8106 at the 15 DAF (C1) and 35 DAF (T1), we obtained 17,065,477 and 15,320,325 unfiltered sequence reads, respectively; RIL 8107 at 15 DAF (C2) and 35 DAF (T2) yielded 13,887,551 and 13,804,333 unfiltered sequence reads, respectively, from the miRNA libraries. We obtained 7,145,081, 6,008,924, 5,339,568, and 8,578,303 unique sequences from the C1, T1, C2, and T2 libraries, respectively (Supplementary Table S1). After discarding low-quality reads and further filtering the Rfam (rRNA, tRNA, snoRNA, snRNA, and other Rfam RNAs) and Repbase sequences, a total of 6,466,666, 5,121,789, 4,548,767, and 7,786,168 unique valid small RNA sequences remained, respectively (Supplementary Table S1). The lengths of the unique valid reads ranged from 18 to 25 nucleotides (nt), and the 21–24 nt sequences were predominant in the C1, T1, C2, and T2 libraries, with the 24 (nt) sequences being the most common (Supplementary Figure S1).

Identification of Known, Conserved, and Novel miRNAs

To identify the known miRNAs at different stages of seed expansion, the valid reads from each dataset were mapped to the precursor sequences in the peanut miRNA database available in miRBase (release 21). Based on the screening criteria of the miRNAs, we identified a total of 72 known miRNAs and 576 conserved miRNAs from the four small RNA libraries (Supplementary Table S2). Among these, 35 known miRNAs and 320 conserved miRNAs were shared in common (Supplementary Figures S2A,B). A total of 89 known and conserved miRNAs were confirmed in miRBase. In addition, 337 miRNAs were also confirmed, but their sequences were different from those reported in miRBase, including miR156b-5p_R-1, miR167-3p_L-1, miR394_L+1_2, miR408-5p_L+1R-1, and miR3508_2ss20CT21AT, among others. Furthermore, 222 new miRNAs were identified for newly reported 5p or 3p sequences, such as miR3510-5_p_1ss22GT, miR3513-p5, miR3515-p3, mir3516-p5, and mir3517-p5 (Supplementary Table S2).

To identify the novel or predicted candidate miRNAs (PC miRNAs), the criteria for annotation of plant miRNAs (Meyers et al., 2008) were used in our study. The classification criteria for the PC miRNAs are as follows: reads were not mapped to pre-miRNAs of selected species in miRBase, but the reads were mapped to the genome and the extended genome sequences from the genome loci may form hairpins. We predicted 434 PC miRNAs from the four small libraries (332 miRNAs in C1, 332 miRNAs in T1, 241 miRNAs in C2, and 321 miRNAs in T2) (Supplementary Table S3). Among these, 172 PC miRNAs...
Ma et al. Roles of microRNAs in Seed Expansion in Peanut

were found to be common to the four miRNA databases (Supplementary Figure S2C).

**Comparison of Differentially Expressed miRNAs Between the Two Peanut RILs**

We compared the frequencies of occurrence of differentially expressed miRNAs at the 15 and 35 DAF stages between the two lines based on a Poisson distribution approach (Audic and Claverie, 1997). We identified 130, 177, 139, and 175 differentially expressed miRNAs in the T1 vs. C1, T2 vs. C2, C2 vs. C1, and T2 vs. T1 comparisons, respectively (Supplementary Tables S4-1–S4-4). Among these, 26 differentially expressed miRNAs were common to the four comparisons (Figure 2). In order to dissect the miRNAs at the pod enlargement stage, we mainly focused on differentially expressed miRNAs at the 15 DAF and 35 DAF stages between the two peanut RILs. We identified 86 known or conserved miRNAs from RIL 8106 and 119 known or conserved miRNAs from RIL 8107 that were differentially expressed at the 35 DAF stage compared to 15 DAF (Supplementary Tables S4-1.4-2). Of these, 48 miRNAs were common to the two peanut lines (Supplementary Figure S3A). Moreover, 38 miRNAs and 71 miRNAs were specifically expressed in RILs 8106 and 8107, respectively (Supplementary Figure S3A). We used the following criteria to identify differentially expressed miRNAs: the adjusted p-value was <0.05 in at least one of the two comparisons. Finally, 143 miRNAs were differentially expressed at 35 DAF in the two lines (Supplementary Table S5), and 11 miRNAs showed up-regulated expression (Table 1), while another 21 miRNAs were down-regulated (Table 2). There are 16 miRNAs showed contrasting expression patterns in the two lines (Table 3); among them, 10 miRNAs, such as miR159a-5p_2ss19TA20TA, miR167-3p_L-1, and miR172c-5p, were down-regulated in RIL 8106 but up-regulated in RIL 8107, while six miRNAs including miR166a_1ss5AN, miR166b_L-1_1ss22AC, and miR6300_R+4_1 were down-regulated in RIL 8107 but were up-regulated in RIL 8106 (Figure 3A). Moreover, 95 miRNAs were expressed predominantly in only one of the RILs at the 35 DAF stage (Table 4 and Supplementary Table S6).

In addition, 44 PC miRNAs from RIL 8106 and 58 PC miRNAs from RIL 8107 were differentially expressed between the 15 and 35 DAF stages (Supplementary Tables S4-1-4-2). Among them, 20 miRNAs were common to the two peanut RILs (Supplementary Figure S3B), 24 miRNAs and 38 miRNAs were specifically expressed in RILs 8106 or 8107, respectively (Supplementary Figure S3B). Based on the screening criteria of the differentially expressed miRNAs, 74 miRNAs showed differential expression between the two lines (Supplementary Table S7). Of these, 12 miRNAs, such as PC-3p-12707_579 and PC-3p-55057_146, showed down-regulation in both lines at 35 DAF (Supplementary Table S8-1). Eight miRNAs were down-regulated in RIL 8106 but up-regulated in RIL 8107 (Supplementary Table S8-2). In addition, 54 miRNAs were expressed predominantly in only one line at the 35 DAF stage (Supplementary Table S8-3 and Figure 3A).

![FIGURE 2](https://www.frontiersin.org/NewMediaVideo恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨恨## Figure 2 | Differentially expressed miRNAs in two peanut RILs. Venn diagrams showing the number of common and specific miRNAs in comparisons of the four libraries (A). Volcanic diagrams showing the number of differentially expressed miRNAs in each comparison (B). The red dots indicate miRNAs with significant differences, and the blue dots indicate that the difference was not significant for miRNA expression. C1, RIL 8106 at 15 DAF; T1, RIL 8106 at 35 DAF; C2, RIL 8107 at 15 DAF; T2, RIL 8107 at 35 DAF.
Validation of Differentially Expressed miRNAs by qPCR

To confirm the high-throughput sequencing data and further comparative analyses, we verified the expression patterns of 16 randomly chosen miRNAs by qPCR. The qPCR results coincided with those of the high-throughput sequencing (Figure 4). For example, we confirmed that the levels of miR156e and miR530_1ss21CA were up-regulated in both peanut lines, whereas miR164a_L+1, miR319d_L+1R-2, PC-3p-12707_579, and PC-3p-55057_146 were down-regulated in both lines. Similarly, expression of miR172c-5p was shown by both methods to be down-regulated in RIL 8106 but up-regulated in RIL 8107. Conversely, miR166b_L+1_1ss22AC was shown by both methods to be down-regulated in 8107 but up-regulated in 8106. Moreover, miR160a, miR164a, miR168a, and miR171n_1ss20TC were expressed predominantly in RIL 8107. These results indicate that the frequency of occurrence as determined by high-throughput sequencing gave a reliable prediction of miRNA expression patterns.

Identification of Target Genes of miRNAs by Degradome Analysis

Target gene validation is important to further understand the biological functions of miRNAs. In this study, we constructed

**TABLE 1** miRNAs showing up-regulated expression in two peanut recombinant inbred lines at 35 DAF.

| miRNA ID          | Normalized value mean | Fold-change | Log₂ Fold-change | Log₂ p-Value | p-Value |
|-------------------|-----------------------|-------------|-----------------|--------------|---------|
|                   | C1 T1 C2 T2           | (T1/C1)     | (T1/C1)         | (T2/C2)      | (T2/C2) |
| ahymiR156b-3p     | 110 312 113 344       | 2.84        | 1.51            | 3.04         | 1.60    |
| ahymiR156b-5p_R+1 | 16 32 10 28           | 2.08        | 1.06            | 2.74         | 1.45    |
| ahymiR156b-5p_R+2 | 0 18 0 52             | inf         | inf             | inf          | inf     |
| ahymiR156e        | 1,500 7,809 1,535 5,764 | 5.21        | 2.38            | 3.75         | 1.91    |
| ahymiR1609c-L-1_1ss22GT | 1,452 2,256 1,933 1,933 | 1.55        | 0.64            | 2.04         | 1.03    |
| ahymiR8422a-5p-L-1_2ss11TG20AT | 27 76 57 1,366    | 2.79        | 1.48            | 24.03        | 4.59    |
| ahymiR8350_1ss21CA | 394 736 395 826        | 1.87        | 0.90            | 2.09         | 1.06    |

**TABLE 2** Down-regulated miRNAs in two peanut recombinant inbred lines at 35 DAF.

| miRNA ID          | Normalized value mean | Fold-change | Log₂ Fold-change | Log₂ p-Value | p-Value |
|-------------------|-----------------------|-------------|-----------------|--------------|---------|
|                   | C1 T1 C2 T2           | (T1/C1)     | (T1/C1)         | (T2/C2)      | (T2/C2) |
| ahymiR1590c-p5_2ss11AG18GA | 1,433 687 1,237 822 | 0.48        | -1.06           | 0.66         | -0.59   |
| ahymiR1560c-p3_2ss21GA23TA | 44 0 137 50          | inf         | inf             | 0.36         | -1.47   |
| ahymiR1609c-p5_3_1ss5GA    | 443 131 464 198       | 0.30        | -1.76           | 0.43         | -1.23   |
| ahymiR164a_L+1   | 104 53 73 12          | 0.50        | -0.99           | 0.17         | -2.58   |
| ahymiR164a_R+1  | 94 38 52 6            | 0.41        | -1.30           | 0.11         | -3.19   |
| ahymiR164a_1ss21AG | 81 26 59 3            | 0.33        | -1.62           | 0.06         | -4.10   |
| ahymiR166c-5p   | 1,350 308 1,115 401   | 0.23        | -2.13           | 0.36         | -1.47   |
| ahymiR166b-5p_2ss8TC10GT | 1,095 200 1,197 275 | 0.18        | -2.46           | 0.23         | -2.12   |
| ahymiR171a      | 21 0 46 24            | -inf        | -inf            | 0.52         | -0.95   |
| ahymiR171c-p5   | 71 32 142 33          | 0.45        | -1.16           | 0.24         | -2.08   |
| ahymiR172g_R+1_1ss9AT | 7 1 9 0             | 0.09        | -3.41           | -inf         | -6.77   |
| ahymiR3909      | 485 77 840 589        | 0.16        | -2.66           | 0.70         | -0.51   |
| ahymiR3913-p5_3_1ss19TC | 3 0 20 6            | inf         | inf             | 0.27         | -1.87   |
| ahymiR396p-p5   | 17 9 24 11            | 0.54        | -0.88           | 0.47         | -1.08   |
| ahymiR399c-p3   | 196 96 520 347        | 0.49        | -1.03           | 0.67         | -0.58   |
| ahymiR399b      | 505 167 1,789 98      | 0.33        | -1.60           | 0.05         | -4.20   |
| ahymiR399d-p3   | 16 0 48 0             | inf         | inf             | inf          | inf     |
| ahymiR1511-p5   | 632 136 668 257       | 0.22        | -2.21           | 0.38         | -1.38   |
| ahymiR1515_L-1  | 58 24 37 18           | 0.42        | -1.25           | 0.47         | -1.09   |
| ahymiR3508      | 328 168 4,428 311     | 0.51        | -0.97           | 0.07         | -3.83   |
two degradome libraries, DS1 (including C1 and T1) and DS2 (including C2 and T2). In total, we obtained 15,768,282 and 17,187,562 raw reads from DS1 and DS2, respectively. After removing the reads without the CAGCAG adaptor, 4,860,730 and 4,780,726 unique raw reads remained from the DS1 and DS2 libraries. The unique reads were aligned to the peanut genome database, and 4,824,026 and 4,746,614 reads were mapped to the genome, respectively. The mapped reads represented 2,916,767

| miRNA ID | Normalized value mean | Fold-change | Log2 | Fold-change | Log2 | p-Value | p-Value | Changing pattern |
|----------|-----------------------|-------------|------|-------------|------|---------|---------|-----------------|
| ahy-miR159a-5p_2ss19TA20TA | 737 225 229 267 | 0.30 | −1.71 | 1.17 | 0.22 | 1.49E-04 | 6.21E-04 | a |
| ahy-miR166a-5p_2ss10GT11CT | 40 6 195 1,043 | 0.15 | −2.69 | 5.36 | 2.42 | 1.78E-02 | 2.07E-02 | a |
| ahy-miR167-3p_L-1 | 1,691 386 1,300 8,060 | 0.23 | −2.13 | 6.20 | 2.63 | 9.14E-05 | 4.48E-03 | a |
| ahy-miR171m-p5_1ss17CT | 964 494 669 963 | 0.51 | −0.97 | 1.44 | 0.53 | 1.70E-03 | 1.59E-02 | a |
| ahy-miR172c-5p | 90 46 61 344 | 0.51 | −0.96 | 5.62 | 2.49 | 1.40E-03 | 2.54E-03 | a |
| ahy-miR395a_L-1R+1 | 4 1 0 8 | 0.19 | −2.41 | inf | inf | 2.95E-02 | 3.35E-02 | a |
| ahy-miR395p-p5 | 22 5 11 26 | 0.25 | −2.03 | 2.39 | 1.26 | 3.06E-03 | 1.87E-02 | a |
| ahy-miR1507-5p_R-1_2ssSTA18AG | 155 57 94 597 | 0.37 | −1.44 | 6.36 | 2.67 | 7.61E-03 | 2.47E-02 | a |
| ahy-miR2199_1ss17AG | 9 0 6 226 | −inf | −inf | inf | inf | inf | inf | a |
| ahy-miR3516-p5 | 33 5 146 3,097 | 0.15 | −2.77 | 21.24 | 4.41 | 6.35E-03 | 1.00E-02 | a |
| ahy-miR166a_1ss5AN | 171 271 228 131 | 1.59 | 0.67 | 0.58 | −0.80 | 2.21E-02 | 1.25E-02 | b |
| ahy-miR166b_L-1_1ss22AC | 192 296 212 129 | 1.55 | 0.63 | 0.61 | −0.72 | 9.89E-03 | 1.20E-02 | b |
| ahy-miR2118a-p5_2ss2AG20GT | 549 1,213 185 113 | 2.21 | 1.14 | 0.61 | −0.72 | 1.01E-02 | 1.99E-03 | b |
| ahy-miR6300_R+1_4 | 20 52 34 15 | 2.66 | 1.41 | 0.44 | −1.20 | 1.32E-04 | 1.09E-02 | b |
| ahy-miR6300_R+4_2 | 20 52 34 15 | 2.66 | 1.41 | 0.44 | −1.20 | 1.32E-04 | 1.09E-02 | b |
| ahy-miR6300_R+4_3 | 20 52 34 15 | 2.66 | 1.41 | 0.44 | −1.20 | 1.32E-04 | 1.09E-02 | b |

| TABLE 3 | miRNAs showing opposite expression patterns in peanut RILs 8106 and 8107 at 35 DAF. |
|-----------|-----------------------------------|----------|------|----------|------|--------|--------|---------------|
| miRNA ID  | Normalized value mean             | Fold-change | Log2 | Fold-change | Log2 | p-Value | p-Value | Changing pattern |
|-----------|-----------------------------------|----------|------|----------|------|--------|--------|---------------|
| ahy-miR159a-5p_2ss19TA20TA | 737 225 229 267 | 0.30 | −1.71 | 1.17 | 0.22 | 1.49E-04 | 6.21E-04 | a |
| ahy-miR166a-5p_2ss10GT11CT | 40 6 195 1,043 | 0.15 | −2.69 | 5.36 | 2.42 | 1.78E-02 | 2.07E-02 | a |
| ahy-miR167-3p_L-1 | 1,691 386 1,300 8,060 | 0.23 | −2.13 | 6.20 | 2.63 | 9.14E-05 | 4.48E-03 | a |
| ahy-miR171m-p5_1ss17CT | 964 494 669 963 | 0.51 | −0.97 | 1.44 | 0.53 | 1.70E-03 | 1.59E-02 | a |
| ahy-miR172c-5p | 90 46 61 344 | 0.51 | −0.96 | 5.62 | 2.49 | 1.40E-03 | 2.54E-03 | a |
| ahy-miR395a_L-1R+1 | 4 1 0 8 | 0.19 | −2.41 | inf | inf | 2.95E-02 | 3.35E-02 | a |
| ahy-miR395p-p5 | 22 5 11 26 | 0.25 | −2.03 | 2.39 | 1.26 | 3.06E-03 | 1.87E-02 | a |
| ahy-miR1507-5p_R-1_2ssSTA18AG | 155 57 94 597 | 0.37 | −1.44 | 6.36 | 2.67 | 7.61E-03 | 2.47E-02 | a |
| ahy-miR2199_1ss17AG | 9 0 6 226 | −inf | −inf | inf | inf | inf | inf | a |
| ahy-miR3516-p5 | 33 5 146 3,097 | 0.15 | −2.77 | 21.24 | 4.41 | 6.35E-03 | 1.00E-02 | a |
| ahy-miR166a_1ss5AN | 171 271 228 131 | 1.59 | 0.67 | 0.58 | −0.80 | 2.21E-02 | 1.25E-02 | b |
| ahy-miR166b_L-1_1ss22AC | 192 296 212 129 | 1.55 | 0.63 | 0.61 | −0.72 | 9.89E-03 | 1.20E-02 | b |
| ahy-miR2118a-p5_2ss2AG20GT | 549 1,213 185 113 | 2.21 | 1.14 | 0.61 | −0.72 | 1.01E-02 | 1.99E-03 | b |
| ahy-miR6300_R+1_4 | 20 52 34 15 | 2.66 | 1.41 | 0.44 | −1.20 | 1.32E-04 | 1.09E-02 | b |
| ahy-miR6300_R+4_2 | 20 52 34 15 | 2.66 | 1.41 | 0.44 | −1.20 | 1.32E-04 | 1.09E-02 | b |
| ahy-miR6300_R+4_3 | 20 52 34 15 | 2.66 | 1.41 | 0.44 | −1.20 | 1.32E-04 | 1.09E-02 | b |
and 2,979,274 annotated peanut genes in the DS1 and DS2 libraries, respectively.

We identified cleaved targets for miRNAs based on a method in the CleaveLand pipeline (Addo-Quaye et al., 2009b), in which a host gene with an alignment score of 4 or less was considered to be a potential target. In total, 1,766 and 1,616 targets were identified from the DS1 and DS2 libraries, respectively (Supplementary Tables S9, S10). For these targets, 501, 5, 859, 46, and 355 belonged to categories 0, 1, 2, 3, or 4 in the DS1 library, and 495, 45, 646, 40, and 390 belonged to a category <4 in the DS2 library (Supplementary Tables S9, S10). As expected, most of the transcripts targeted by the highly conserved miRNAs were associated with conserved target genes. For example, miR156e targets the SPL6 and SPL18 genes; miR164a_L+1 targets the NAC021 gene; miR166b_L+1_1ss22AC targets ATHB-15 gene; miR319d_L+1R-2 targets the TCP3 and TCP4 genes; miR159b_R-1_1ss7GT targets the GAM1 gene, miR160a targets the ARF10, ARF16, and ARF18 genes; miR164a targets the NAC100 gene; miR168a targets the AGO1 gene; and miR171n_1ss20TC targets the SCL22 gene. Also, the expression analysis of the 12 representative target genes by qPCR showed negative correlations with the levels of their corresponding miRNAs (Figure 5). These results show that several miRNAs may be directly or indirectly involved in peanut seed expansion by regulating expression of their target gene(s).

There were 1,401 differentially expressed target genes between the DS1 and DS2 libraries (Supplementary Table S11). Examples of “target plots” (T-plots) for the targets for peanut miRNAs are shown in Figure 6. In this T-plot, NAC100 (XM_016332293.1) and SCL22 (XM_016328632.1) were cleaved at 613 nt and 1196 nt in the two peanut lines by miR164a and miR171n_1ss20TC, respectively (Figure 6). Interestingly, the read numbers of these miRNAs that cleaved the target genes in RIL 8107 showed a significant decrease compared to that in RIL 8106, which is consistent with the expression level of miRNAs identified by RNA-Seq and qPCR (Figure 4). Also, higher expression levels of these two targets, NAC100 and SCL22 were observed in RIL 8107 compared to the levels in RIL 8106 (Figure 5). The results showed that these two miRNAs may be involved in the seed expansion process in peanut by negatively regulating NAC100 and SCL22.

For the novel peanut miRNAs, 129 transcripts were identified as targets for 26 PC miRNAs in the DS1 library, and 119 transcripts were identified as targets for 23 PC miRNAs in the DS2 library (Supplementary Tables S9, S10). For these targets, 13, 2, 73, 5, and 36 transcripts belonged to categories 0, 1, 2, 3, and 4 in DS1, whereas 10, 3, 49, 3, and 54 transcripts belonged to categories 0, 1, 2, 3, and 4 in DS2. These data suggest that the majority of the identified PC miRNA target genes are in categories 0, 1, 2, and 3.

**Function of the Potential miRNA Targets**

Gene ontology categories were assigned to 488 target genes for the 217 differentially expressed miRNAs...
FIGURE 4 | Comparisons of the expression levels of 16 miRNAs in seeds of two peanut RILs. The expression levels were normalized to the expression of Actin in qPCR. Red indicates the fold-changes of miRNA expression levels determined by qPCR. Blue indicates the miRNA expression fold-changes generated from the deep sequencing. The experiments were repeated three times, and vertical bars indicate the standard errors.

(Supplementary Table S12). Twelve classes of biological processes were identified, with the two most frequent being "regulation of transcription" and "cell division and differentiation" (Figure 3B). Moreover, other growth and development-related genes were also identified as miRNA targets, including "auxin mediated signaling pathway" and "cell cycle" (Figure 3B). These results imply the possible function of miRNAs in the regulation of biological processes involved in peanut seed expansion. In addition, based on the KEGG analysis, 208 target genes were significantly enriched in 15 pathways including "plant hormone signal transduction," "spliceosome," and "basal transcription factors" (Supplementary Figure S4). Examples of the plant hormone signal transduction pathway and the corresponding miRNAs are shown in Figure 3C. In this pathway, miR160a targets the ARF gene to respond to the IAA signal. Another miRNA, miR171n, is involved in ubiquitin-mediated proteolysis, and controls the accumulation level of the DELLLA protein in the GA pathway. These findings indicate that miRNAs have a significant effect on the regulation of peanut seed expansion by effecting hormone signaling transduction pathways.

DISCUSSION

Functions of miRNAs During Peanut Seed Expansion

Peanut seed expansion, including embryogenesis, cell division, and differentiation, and seed enlargement, is a complex biological process regulated by the expression of many genes. Previous studies have indicated that miRNAs and their predicted target mRNAs play important regulatory roles in plant growth and development (Jiang et al., 2014). Nevertheless, the function of miRNAs and their targets during peanut seed expansion has not been reported. In our study, we identified 72 known miRNAs, 576 conserved miRNAs, and 434 PC miRNAs from the four libraries (Supplementary Tables S2, S3). Interestingly, 222 miRNAs (including miR3513-p5, miR3515-p3, and miR3516-p5) were
first identified in peanut, indicating that they are preferentially expressed and are specific to peanut seed development. In addition, several PC miRNAs accumulated only in the 15 DAF or 35 DAF stages, for example, PC-5p-94368_80, PC-5p-69905_114, PC-3p-63255_126, and PC-5p-118138_60, indicating that these miRNAs may play key roles in peanut seed expansion.

**Presumptive miRNA–mRNA Modules Involved in Peanut Seed Expansion**

In this study, we identified 11 up-regulated miRNAs (Table 1) at 35 DAF in the two peanut lines. For example, the target gene of miR156e encodes the Squamosa promoter binding protein-like (SPL) transcription factor. Previous studies have shown that SPL family members (SPL3, SPL4, and SPL5) make a secondary contribution to the regulation of flowering and appear to function mostly in the control of flowering time and developmental phase change (Wu and Poethig, 2006; Wang et al., 2009). Another study also showed that OsSPL16 encodes a protein that promotes cell division, also with positive consequences for grain width and yield in rice (Wang et al., 2012). In this work, miR156e was up-regulated at the 35 DAF stage, while the expression pattern of the SPL gene was significantly down-regulated. The reduced expression of SPL might prolong the developmental phase associated with seed expansion.

Expression of 33 miRNAs showed down-regulation in the two peanut RILs at 35 DAF (Table 2 and Supplementary Table S8-1). The target gene of miR164a_L+1 encodes the NAC transcription factor, which plays a role in the early auxin response (Guo et al., 2005; Zhang et al., 2006). Another down-regulated miR319d_L+1R-2 is a member of the miR319 family and targets TCP4-like (TCP4) genes, which encode plant-specific transcription factors involved in jasmonic acid (JA) biosynthesis (Nag et al., 2009; Schommer et al., 2014). Previous studies have demonstrated that TCP positively regulates the JA level in plants (Schommer et al., 2008; Hao et al., 2012). JA has also been reported to affect grain filling in rice (Kim et al., 2009). In addition, miR319 might influence sweet corn seed vigor (Gong et al., 2015). In our present work, we found that the level of TCP4-specific mRNA increased compared with the decrease in miR319d_L+1R-2 (Figures 4, 5). The increased expression of TCP4 may enhance seed vigor by increasing the level of JA, which could then contribute to seed expansion in peanut.

There were 24 miRNAs that showed contrasting patterns of differential expression in the two peanut lines at 35 DAF (Table 3 and Supplementary Table S8-2). For example, miR172c-5p was down-regulated in the RIL 8106, but was up-regulated in RIL 8107 (Table 3 and Figure 4). The target gene of miR172c-5p is the splicing factor U2AF large subunit B-like (U2AF), which was up-regulated in RIL 8106 and down-regulated in RIL 8107 at the 35 DAF stage (Figure 5), indicating that U2AF may also be involved in peanut seed expansion. However, miR166b_L+1_1ss22AC was up-regulated in RIL 8106, but down-regulated in RIL 8107 (Table 3 and Figure 4).
miR166b_L+1_1ss22AC is a member of the miR166 family and targets the homeobox-leucine zipper protein ATHB-15 gene. In this study, we observed that expression of ATHB-15 increased with the decrease in miR166b_L+1_1ss22AC expression in RIL 8107 at 35 DAF, indicating that ATHB-15 may also play a role in peanut seed expansion by regulating early embryo development.

We also found that 95 known or conserved miRNAs (Table 4 and Supplementary Table S6) and 54 PC miRNAs (Supplementary Table S8-3) were expressed predominantly in only one of the two peanut RILs. For example, miR159b_R-L_1ss7GT and PC-3p-79977_98 were down-regulated in RIL 8106 at 35 DAF (Table 4 and Supplementary Table S8-3). miR159 negatively regulates the expression of GAMYB genes at the posttranscriptional level; GAMYB was first identified as a downstream GA signaling target in aleurone cells of barley (Hordeum vulgare L.) (Gubler et al., 1995). MYB family members were also found to play an important role in response to the presence of abscisic acid (ABA) during seed development in rice (Aldridge and Probert, 1993; Reyes and Chua, 2007). Our results indicate that the absence of miR159b_R-1_1ss7GT in RIL 8106 lead to increased expression of the GAMYB gene, which would be consistent with its role in peanut seed development. The target gene of PC-3p-79977_98, a novel miRNA, is the ubiquitin-conjugating enzyme E2-C (UCE2) which contributes to APC-dependent protein ubiquitylation in vivo during the cell cycle (Criqui et al., 2002) and cell differentiation (Wen et al., 2006). In this study, we found that UCE2 was up-regulated in concert with the down-regulation of PC-3p-79977_98 in RIL 8106 (Figures 4, 5), indicating that PC-3p-79977_98 may also participate in peanut seed expansion by regulating the cell cycle. Our findings suggest that the strong expression of these miRNAs

FIGURE 6 | Examples of T-plots of miRNA targets in two peanut RILs confirmed by degradome sequencing. The T-plots show the distribution of the degradome tags along the full length of the target mRNA sequence. The vertical red line indicates the cleavage site of each transcript and is also shown by an arrow. (A,B) The cleavage features in NAC domain-containing protein 100-like (XM_016332293.1) mRNA by miR164a in the two peanut RILs, 8106 and 8107, respectively. (C,D) The cleavage features in scarecrow-like protein 22 (XM_016328632.1) mRNA by miR171n_1ss20TC in RILs 8106 and 8107, respectively.
We also found that the expression of a number of miRNAs was down-regulated only in RIL 8107, including miR160a, miR168a, and miR171n_1ss20TC, etc. (Table 4 and Figure 4). The target gene of miR160a is a member of the auxin response factor (ARF) gene family. miR160 negatively regulates ARF10 (Liu et al., 2007), ARF16 (Wang et al., 2005), and ARF17 (Mallory et al., 2005) of the repressor ARF family, and plays a critical role in maintaining the process of seed germination and the normal developmental programs for embryos, roots, leaves, and floral organs. Furthermore, ARFs regulate the expression of early auxin responsive genes, including the AUX/IAA genes (Ulmasov et al., 1997; Tiwari et al., 2003), and frees ARFs from repression by AUX/IAA proteins. The accumulation of ARF16 resulting from the down-regulation of miR160a might enhance the auxin response and thus enhance seed development in peanut. Transcription of miR168a also was significantly decreased in RIL 8107, and its target is Argonaute 1 (AGO1) which is the core component of the RNA-induced silencing complex (RISC) (Vaucheret et al., 2006) and plays crucial roles in controlling cotyledon expansion and hormone signaling (Vaucheret et al., 2004; Kidner and Martienssen, 2005). Additionally, scarecrow-like protein 22 (SCL22) was predicted to be the target gene of miR171n_1ss20TC, and it can control seed development (Kim and Ahn, 2014). We found that miR171n_1ss20TC was down-regulated in RIL 8107, indicating that SCL may also be involved in peanut seed expansion. These results suggest that genotype-specific regulation of miRNAs might explain why the seeds of RIL 8107 are larger than seeds of RIL 8106. Altogether, our results suggest that at least 10 regulatory modules of these miRNA-targets play an important role in seed expansion in peanut.

Possible miRNA-Dependent Regulatory Pathways That Participate in Peanut Seed Expansion

Phytohormones can act as signaling compounds that promote and influence plant development. For example, auxin has been shown to be essential for plant growth and development by controlling cell division and cell elongation (Sauer et al., 2013). Gibberellic acid (GA) is involved in plant flowering and embryo development through its regulation of the expression of related genes (Blázquez and Weigel, 2000; White and Rivin, 2000). miR159 has been reported to regulate Arabidopsis floral development in the GA pathways; in addition, miRNA167 is involved in the auxin pathways during grain filling in rice (Achard et al., 2004; Xue et al., 2009). In this study, we also found that several miRNAs, including miR160a, miR171n, and miR156e, participate in auxin signal transduction, GA signal transduction, and BR signal transduction by regulating their target genes (Figure 3C). These results suggest that these miRNAs may play key roles in peanut seed expansion.

A number of other conserved and novel miRNAs may also be involved in peanut seed expansion, and we further assessed the functions of their potential target genes in seed development. Examples are miR164a-mediated cleavage of NAC, miR319d-mediated cleavage of TCP, miR166b-mediated...
cleavage of ATHB, miR159b-mediated cleavage of GAMYB, PC-3p-79977_98-mediated cleavage of UCE2, miR168a-mediated cleavage of AGO1, and miR172c-5p-mediated cleavage of U2AF. These miRNA–mRNA modules might be involved in regulating seed expansion in peanut (Figure 7). In summary, miRNA-target gene models, hormone transduction, and transcriptional regulation comprise a complex network that regulates seed expansion in peanut.

CONCLUSION

In this study, we found that 143 conserved miRNAs and 74 PC miRNAs were differentially expressed in the two peanut RILs at the 35 DAF stage. Of these, expression of 11 miRNAs was found to be up-regulated, while 33 miRNAs were down-regulated. Twenty-four miRNAs displayed contrasting expression patterns in the two peanut lines. Moreover, 149 miRNAs were expressed predominantly in only one of the two RILs. We also validated the expression of a number of representative miRNAs by qPCR. The potential miRNA target genes were verified by sequencing the two degradome libraries. Our results demonstrate that the expression patterns of some miRNAs can be very different even between two closely related inbred lines from the same species. The differentially expressed miRNAs and their target genes identified in this study could be important to the regulatory networks that control seed expansion in peanut.

AUTHOR CONTRIBUTIONS

DY designed and conceived the research. XM and ZX wrote the manuscript. KZ, FL, KL, and LN performed the experiments. ZX, JH, and DY edited the manuscript. All authors read and approved the final manuscript.

FUNDING

This work was financially supported by grants from the National Natural Science Foundation of China (No. 31471525) and Key Scientific and Technological Project in Henan Province (Nos. 161100111000 and S2012-05-03).

REFERENCES

Achard, P., Herr, A., Baulcombe, D., and Harberd, N. (2004). Modulation of floral development by a gibberellin-regulated microRNA. Development 131, 3357–3365. doi: 10.1242/dev.01206
Addo-Quaye, C., Eshoo, T. W., Bartel, D. P., and Axtell, M. J. (2008). Endogenous siRNA and miRNA targets identified by sequencing of the Arabidopsis degradome. Curr. Biol. 18, 758–762. doi: 10.1016/j.cub.2008.04.042
Addo-Quaye, C., Miller, W., and Axtell, M. J. (2009a). CleaveLand: a pipeline for using degradome data to find cleaved small RNA targets. Bioinformatics 25, 130–131. doi: 10.1093/bioinformatics/btn604
Addo-Quaye, C., Snyder, J. A., Park, Y. B., Li, Y. F., Sunkar, R., and Axtell, M. J. (2009b). Sliced microRNA targets and precise loop-first processing of MIR319 hairpins revealed by analysis of the Physcomitrella patens degradome. RNA 15, 2112–2121. doi: 10.1261/rna.177409
Aldana, A. B., Fites, R. C., and Pattee, H. E. (1972). Changes in nucleic acids, protein and ribonuclease activity during maturation of peanut seeds. Plant Cell Physiol. 13, 515–521. doi: 10.1093/oxfordjournals.pcp.a074764
Aldridge, C. D., and Probert, R. J. (1993). Seed development, the accumulation of abscisic acid and desiccation tolerance in the aquatic grasses Porteresia coarctata (Roxb.) Tateoka and Oryza sativa L. Seed Sci. Res. 3, 97–103. doi: 10.1017/S0960258500001641
Ambros, V., Bartel, B., Bartel, D. P., Burge, C. B., Carrington, J. C., Chen, X. M., et al. (2003). A uniform system for microRNA annotation. RNA 9, 277–279. doi: 10.1017/S0960258500001641
Audic, S., and Claverie, J. M. (1997). The significance of digital gene expression profiles. Genome Res. 7, 986–995. doi: 10.1101/gr.7.10.986
Basha, S. M. M., and Cherry, J. P. (1976). Composition, solubility, and gel electrophoretic properties of protein isolated from Flourunner (Arachis hypogaea L.) peanut seeds. *J. Agric. Food Chem.* 24, 359–365. doi: 10.1021/jf00240a058

Bi, Y. P., Liu, W., Xia, H., Su, L., Zhao, C. Z., Wan, S. B., et al. (2010). EST sequencing and gene expression profiling of cultivated peanut (Arachis hypogaea L.) Genome 53, 832–839. doi: 10.1006/jgen.2006.1074

Blázquez, M. A., and Weigel, D. (2000). Identification of floral inductive signals in *Arabidopsis*. Nature 404, 889–892. doi: 10.1038/35009125

Brennan, J. R. (1969). The peanut gyphonore. *Biologist* 51, 71–82.

Chu, X., Yang, Q., Chen, X., Wang, J., Pan, L., Chen, M., et al. (2011). Identification and characterization of microRNAs from peanut (Arachis hypogaea L.) by high-throughput sequencing. *PLoS One* 6:e27530. doi: 10.1371/journal.pone.0027530

Criqui, M. C., de Almeida Engler, J., Camasses, A., Capron, A., Parmentier, Y., Inzé, D., et al. (2002). Molecular characterization of plant ubiquitin-conjugating enzymes belonging to the UbcP4/E2-UCBx/UbcH10 gene family. *Plant Physiol.* 130, 1230–1240. doi: 10.1104/pp.011353

Ding, D., Wang, Y., Han, M., Fu, Z., Li, W., Liu, Z., et al. (2012). MicroRNA trancriptional analysis of heterosis during maize seed germination. *PLoS One* 7:e39578. doi: 10.1371/journal.pone.0039578

Falvey, E., and Schibler, U. (1991). How are the regulators regulated? *FASEB J.* 5, 309–314.

Feng, J., Liu, S., Wang, M., Lang, Q., and Jin, C. (2014). Identification of microRNAs and their targets in tomato infected with Cucumber mosaic virus based on deep sequencing. *Planta* 240, 1335–1352. doi: 10.1007/s00425-014-2158-3

Gong, S., Ding, Y., Huang, S., and Zhu, C. (2015). Identification of miRNAs and their target genes associated with sweet corn seed vigor by combined small RNA and degradome sequencing. *J. Agric. Food Chem.* 63, 5485–5491. doi: 10.1021/acs.jafc.5b00522

Grant-Downton, R., Le Trionnaire, G., Schmid, R., Rodriguez-Enriquez, J., Haifídhi, S., Mehdí, S., et al. (2009). MicroRNA and tasiRNA diversity in mature pollen of *Arabidopsis thaliana*. * BMC Genomics* 10:64. doi: 10.1186/1471-2164-10-64

Gübler, F., Kalla, R., Roberts, J. K., and Jacobsen, J. V. (1995). Gibberellin-enzymes belonging to the UbcP4/E2-C/UBCx/UbcH10 gene family. *Proc. Natl. Acad. Sci. U.S.A.* 100, 22534–22539. doi: 10.1073/pnas.9908718100

Ramakrishna, A., Tam, H. M., Wani, S. P., and Long, T. D. (2006). Effect of mulch on soil temperature, moisture, weed infestation and yield of groundnut in northern Vietnam. *Field Crops Res.* 95, 115–125. doi: 10.1016/j.fcr.2005.01.030

Reyes, J., and Chua, N. H. (2007). ABA induction of miR319 controls transcript levels of two MYB factors during Arabidopsis seed germination. *Plant J.* 49, 592–606. doi: 10.1111/j.1365-313X.2006.02980.x

Sauer, M., Robert, S., and Kleiwe-Vehn, J. (2013). Auxin: simply complicated. *J. Exp. Bot.* 64, 2565–2577. doi: 10.1093/jxb/ert139

Schommer, C., Juan, M., Debernardi, J. M., Bresso, E. G., Rodriguez, R. E., and Palatinik, J. F. (2014). Repression of cell proliferation by miR319-regulated TCP4. *Mol. Plant* 7, 1533–1544. doi: 10.1093/mp/suot084

Smillie, B. W. (1956). *Arachis hypogaea*. Embryogeny and the effect of peg elongation upon embryo and endosperm growth. *Am. J. Bot.* 43, 233–240.

Stanciel, K., Mortley, D. G., Hileman, D. R., Loretan, P. A., Bond, C. K., and Hill, W. W. (2000). Growth, pod and seed yield, and gas-exchange of hydroponically grown peanut in response to CO2 enrichment. *HorticScience* 35, 49–52.

Sunkar, R., and Zhu, J. K. (2004). Novel and stress-regulated microRNAs and other small RNAs from Arabidopsis. *Plant Cell* 16, 2001–2019. doi: 10.1105/tpc.104.022830

Tiwarri, S. B., Hagen, G., and Guilfoyle, T. (2003). The roles of auxin response factor domains in auxin-responsive transcription. *Plant Cell* 15, 533–543. doi: 10.1105/tpc.105.038041

Ulmavov, T., Hagen, G., and Guilfoyle, T. J. (1997). ARFI, a transcription factor that binds to auxin response elements. *Science* 276, 1865–1868. doi: 10.1126/science.276.5320.1865

Van Ex, F., Jacob, Y., and Martienssen, R. A. (2011). Multiple roles for small RNAs during plant reproduction. *Curr. Opin. Plant Biol.* 14, 588–593. doi: 10.1016/j.yqpb.2011.07.003

Vlahov, O., Mallory, A. C., and Bartel, D. P. (2006). AGO1 homeostasis entails coexpression of MIIR168 and AGO1 and preferential stabilization of miR168 by AGO1. *Mol. Cell* 22, 129–136. doi: 10.1016/j.molcel.2006.03.011

Vlahov, O., Vazquez, F., Crépé, P., and Bartel, D. P. (2004). The action of ARGONAUTE1 in the miRNA pathway and its regulation by the miRNA pathway are crucial for plant development. *Genes. Dev.* 18, 1187–1197. doi: 10.1101/gad.1201404

Wang, J. W., Czech, B., and Weigel, D. (2009). miR156-regulated SPL transcription factors define an endogenous flowering pathway in *Arabidopsis thaliana*. *Cell* 138, 738–749. doi: 10.1016/j.cell.2009.06.014

Ma et al.
Wang, J. W., Wang, L. J., Mao, Y. B., Cai, W. J., Xue, H. W., and Chen, X. Y. (2005). Control of root cap formation by MicroRNA-targeted auxin response factors in Arabidopsis. *Plant Cell* 17, 2204–2216. doi: 10.1105/tpc.105.033076

Wang, S., Wu, K., Yuan, Q., Liu, X., Liu, Z., Lin, X., et al. (2012). Control of grain size, shape and quality by OsSPL16 in rice. *Nat. Genet.* 44, 950–955. doi: 10.1038/ng.2327

Wang, S., Wu, K., Yuan, Q., Liu, X., Liu, Z., Lin, X., et al. (2012). Control of grain size, shape and quality by OsSPL16 in rice. *Nat. Genet.* 44, 950–955. doi: 10.1038/ng.2327

Wen, R., Newton, L., Li, G., Wang, H., and Xiao, W. (2006). Arabidopsis thaliana UBC13: implication of error-free DNA damage tolerance and Lys63-linked polyubiquitylation in plants. *Plant Mol. Biol.* 61, 241–253. doi: 10.1007/s11103-006-0007-x

White, C., and Rivin, C. (2000). Gibberellins and seed development in maize. II. Gibberellin synthesis inhibition enhances abscisic acid signaling in cultured embryos. *Plant Physiol.* 122, 1089–1097. doi: 10.1104/pp.122.4.1089

Wu, G., and Poethig, R. S. (2006). Temporal regulation of shoot development in Arabidopsis thaliana by miR156 and its target SPL3. *Development* 133, 3539–3547. doi: 10.1242/dev.02521

Xue, L. J., Zhang, J. J., and Xue, H. W. (2009). Characterization and expression profiles of miRNAs in rice seeds. *Nucleic Acids Res.* 37, 916–930. doi: 10.1093/nar/gkn998

Yamada, T., Aibara, S., and Morita, Y. (1980). Accumulation pattern of arachin and its subunits in maturation of groundnut seeds. *Plant Cell Physiol.* 21, 1217–1226. doi: 10.1093/oxfordjournals.pcp.a0676120

Zhang, B., Pan, X., Cobb, G. P., and Anderson, T. A. (2006). Plant microRNA: a small regulatory molecule with big impact. *Dev. Biol.* 298, 3–16. doi: 10.1016/j.ydbio.2005.10.036

Zhao, C., Xia, H., Cao, T., Yang, Y., Zhao, S., Hou, L., et al. (2015). Small RNA and degradome deep sequencing reveals peanut microRNA roles in response to pathogen infection. *Plant Mol. Biol. Rep.* 33, 1013–1029. doi: 10.1007/s11105-014-0806-1

Zhao, C. Z., Xia, H., Frazier, T. P., Yao, Y. Y., Bi, Y. P., Li, A. Q., et al. (2010). Deep sequencing identifies novel and conserved microRNAs in peanuts (Arachis hypogaea L.). *BMC Plant Biol.* 10:3. doi: 10.1186/1471-2229-10-3

Zhao, Z., Xue, Y., Yang, H., Li, H., Sun, G., Zhao, X., et al. (2016). Genome-wide identification of miRNAs and their targets involved in the developing internodes under maize ears by responding to hormone signaling. *PLoS One* 11:e0164026. doi: 10.1371/journal.pone.0164026

Conflict of Interest Statement: 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.

Copyright © 2018 Ma, Zhang, Zhao, Li, Li, Ning, He, Xin and Yin. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.