Genome-Wide Identification of miRNAs Responsive to Drought in Peach (*Prunus persica*) by High-Throughput Deep Sequencing

**Abstract**

Peach (*Prunus persica* L.) is one of the most important worldwide fresh fruits. Since fruit growth largely depends on adequate water supply, drought stress is considered as the most important abiotic stress limiting fleshy fruit production and quality in peach. Plant responses to drought stress are regulated both at transcriptional and post-transcriptional level. As post-transcriptional gene regulators, miRNAs (miRNAs) are small (19–25 nucleotides in length), endogenous, non-coding RNAs. Recent studies indicate that miRNAs are involved in plant responses to drought. Therefore, Illumina deep sequencing technology was used for genome-wide identification of miRNAs and their expression profile in response to drought in peach. In this study, four SRNA libraries were constructed from leaf control (LC), leaf stress (LS), root control (RC) and root stress (RS) samples. We identified a total of 531, 471, 535 and 487 known mature miRNAs in LC, LS, RC and RS libraries, respectively. The expression level of 262 (104 up-regulated, 158 down-regulated) of the 453 miRNAs changed significantly in leaf tissue, whereas 368 (221 up-regulated, 147 down-regulated) of the 465 miRNAs had expression levels that changed significantly in root tissue upon drought stress. Additionally, a total of 197, 221, 238 and 265 novel miRNA precursor candidates were identified from LC, LS, RC and RS libraries, respectively. Target transcripts (137 for LC, 133 for LS, 148 for RC and 153 for RS) generated significant Gene Ontology (GO) terms related to DNA binding and catalytic activities. Genome-wide miRNA expression analysis of peach by deep sequencing approach helped to expand our understanding of miRNA function in response to drought stress in peach and Rosaceae. A set of differentially expressed miRNAs could pave the way for developing new strategies to alleviate the adverse effects of drought stress on plant growth and development.

**Introduction**

Peach (*Prunus persica* L.) is considered to be one of the most widely grown and economically important stone fruit species in the Rosaceae, comprising more than 3,000 species in approximately 110 genera distributed worldwide [1]. In 2010, it was estaminated that world annual production of peaches and nectarines exceeded 19 million metric tons according to FAO statistics (FAOSTAT, http://faostat.fao.org). In addition to its ecological and high economic importance, peach is also emerging as a model tree species for both comparative genomic studies, evolutionary studies and plant development research owing to its small genome size of 300 Mb (just about twice comparing with *Arabidopsis thaliana*) and the relatively short reproductive time [2,3]. Peach has a haploid chromosome number of 8 [4] and the first draft of peach genome (peach v1.0, obtained from “Lovell” haploid) was unraveled by the International Peach Genome Initiative (IPGI), which are available from the Genome Database for Rosaceae (http://www.rosaceae.org/peach/genome). The genus *Prunus*, which includes peach, nectarine, apricot, weet and sour cherry, have stone fruits with fleshy mesocarp, but the growth and development of these fruits, especially large-fruited species like peach, are seriously affected by drought. For peach, drought stress is one of the major abiotic stresses limiting fruit production and quality during the 4–6 week period before harvesting [5,6].

As a major abiotic factor, drought can be described as basically the water deficiency or insufficient access to water and it has adverse effects on the growth of plants and crop production. However, plants growing in drought stress respond to dehydration and have to develop a variety of mechanisms at morphological and molecular level in order to remain alive. Deciphering the physiological processes and molecular genetic mechanisms involved in drought resistance has certainly made a significant progress in understanding of complex biological response of plants at the molecular and organism levels against the drought. The expression profile of protein-coding genes could highly fluctuate in response to drought at the transcriptional and post-transcriptional levels of miRNA [7,8,9]. The aberrant expression of genes may be regulated by an newly discovered small RNAs, termed micro-RNAs (miRNAs).

The miRNAs are an extensive class of small (19–25 nucleotides), single-strand, endogenous, noncoding RNAs which negatively
modulate gene expression at the post-transcriptional levels by
directing the cleavage of miRNAs or by inhibiting translation
dependent on the extent of complementarity between miRNA and
its target(s) [7,8,10–13]. In plants, biogenesis of miRNA necessitates
a multiple biological process to generate full-function mature
miRNAs by recruiting several evolutionary conserved protein
families. At first, plant miRNA genes are transcribed to primary
miRNAs (pri-miRNAs) by RNA polymerase II [14–16], then these
long pri-miRNAs are cleaved to hairpin-like miRNA precursor
(pre-miRNAs) and the loop-regions of the hairpin are excised by
RNase III enzyme DICER LIKE1 (DCL1) [17] with the aid of
HYL1 and SERRATE [18–21]. Eventually, released mature
miRNAs incorporate into ribonucleoprotein complex known as
the RNA-induced silencing complex (RISC), which inhibits
translation elongation or triggers the degradation of target mRNA
[12,22].

Considerable amount of plant miRNAs have been identified by
computational and/or experimental methods and these miRNAs
have been deposited in the latest release of miBase v18 (release
18.0 November 2011, only experimentally validated [23]) and
PMRD [Plant microRNA database, both experimental and
computational [24]; since the first miRNAs were discovered in
plants in 2002 [25,26]. At present, there are 4053 hairpin entries
pertaining to 52 plant species in the miBase v18.0. The
identification of any miRNAs has great importance for subsequent
research such as miRNA function, nature, target prediction and
biogenesis. In recent years, the innovative strategies and practical
methodologies have been developed for determining miRNA
eexpression. The main approaches of experimental methodologies
can be summarized as follows: direct-cloning [27–29], stem-loop
qRT-PCR [10,30–32], next-generation sequencing technology
[33–37] and hybridization-based detection, such as northern blotting [38], in situ detection [39,40] and microarray [31,41,42].
Although each of these methods has their own particular
advantages and disadvantages, the next-generation sequencing
technologies play an increasingly prominent role in discovering
novel miRNAs [36,43] and measuring quantitatively expression
levels of low-abundant [44] and species or tissue specific miRNAs
[35,45] comparing to other genome-wide transcriptome analysis
methods, such as miRNA-microarray [46,47]. In addition to
experimental approach, computational approach has also been a
preferred method because of its low cost, high efficiency and
speed, prior to experimental validation. The efficacy and power of
computational approach come from its major characteristic features: (i) high evolutionary conservation from mosses to
cudicots in plants (comparative genomics) [48,49], (ii) hairpin-shaped
stem-loop secondary structure [50,51], (iii) high minimal
folding free energy index [48].

Many recent studies have revealed that plant miRNAs have
pivotal roles in plant response to abiotic stresses, including drought
[9,52,53], salt [54,55,56], cold [33,57], oxidative stress [58–60]
and UV-B radiation [61]. The miRNAs whose expression level is
significantly altered in drought condition compared with normal
conditions have been well-reviewed in recent works [7,8]. It has
been indicated that a certain number of miRNAs involve in
response to drought stress by altering the gene expression
[52,62,63]. As for Prunus species, there is no comprehensive data
about the expression profiles of drought responsive miRNAs. The
aim of the present study is to determine the expression profile of
drought stress-responsive miRNAs in peach. Thus miRNA deep
sequencing by Illumina HiSeq 2000 were applied not only for
simultaneous evaluation of drought responsive miRNAs’
expressions, but also for providing comprehensive information about P. persica miRNA transcriptome on genome-wide scale. Stem-loop
real time qRT-PCR (ST-RT PCR) was also employed to further
validate the expression level of a set of miRNAs identified during
deep sequencing. Additionally, the identification and character-
ization of P. persica miRNAs and their target genes were established
by using computational methods combined with experimental validation.

Results
We used the Illumina Solexa sequencing platform to investigate the genome-wide identification and expression profiles of miRNAs
in peach, particularly for the drought-responsive miRNAs. Four
small RNA libraries were constructed by the use of total RNAs
isolated from control leaf (LC), drought-stressed leaf (LS), control
root (RC), and drought-stressed root (RS) tissues. Small-RNA
sequencing yielded a total of 53,878,885 high-quality raw sequence.
Total high-quality raw reads in each of LC, LS, RC and RS libraries
are 15,499,314, 12,473,137, 12,703,130 and 13,203,304,
respectively (Table 1). After removing low quality reads, adapters, poly-A
sequences and short RNA reads smaller than 18 nucleotides,
53,475,533 (99.23%) clean reads including 14,204,383 unique
sequences were obtained from the all libraries. Among the unique
sequences, 2,063,684 (49.01%), 1,599,019 (50.40%), 1,400,836
(51.96%) and 1,747,201 (42.36%) were mapped to the peach
genoome using SOAP2 for sequences generated from LC, LS, RC
and RS, respectively (Table 1 and Table S1). In order to get a
big-picture view of sequence distribution of all sRNA reads, all clean
reads were mapped against the peach genome database at Genbank
(http://www.ncbi.nlm.nih.gov/Genome/388), Rfam (http://rfam.
sanger.ac.uk/) and miRBase v18.0 (http://www.mirbase.org/), and
therefore, they are classified into seven annotation categories: non-
coding RNAs (tRNA, rRNA, snRNA and snoRNA), miRNA, exon-
sense, exon-antisense, intron-sense, intron-antisense, and unknown
sRNAs (Table 2, Figure S1). As shown in Table 2, the highest
abundance of unique conserved and potential non-conserved
miRNAs reads was found in root-drought stress library and leaf
control library, whereas most of the total miRNA reads were found
in leaf-stress library. The length distribution of unique sRNA reads
revealed that the majority of reads from each library were 18–25 nt
in length, of which the class of 24 nt was the most abundant group
accounted for average ~50% of total reads for each library and it
was followed by the group of 21 nt class (Figure 1). Although these
small RNAs unevenly distributed in four groups according to their
length, small RNAs in control and drought-exposed group for leaf
and root represent similar distribution within each of their own
group (Figure 1).

Identification of known miRNAs in peach
In order to identify known (both conserved and species-specific)
miRNAs from control and drought-exposed root and leaf tissues of
peach, small RNA sequences generated from each library were
independently aligned with currently known and experimentally
validated mature miRNAs deposited in miBase v18.0, including
4,014 viridiplantae miRNAs belonging to 52 plant species. After
homology search, a total of 531, 471, 535 and 487 miRNAs were
identified from LC, LS, RC and RS libraries, respectively (Table
S2). These miRNAs belong to 43 evolutionary conserved miRNA
families (Table 3), suggesting that miRNA-mediated biological
process are also present in peach as found in other plant species.
However, some miRNAs, such as miR116, miR347, miR441 and
miR529, were not detected in both leaf and root samples, suggesting that these miRNAs may be tissue-specific expression.
The expression levels varied from miRNAs to miRNAs from one
copy to more than one million of copies based on the deep

Identification of novel miRNAs in peach

A diversity of miRNAs were identified using the new sequencing approach (Table S2). A majority of miRNAs were detected with more than 50 copies; such as a total of 272 miRNAs for LC (51.22%), 229 miRNAs for LS (48.61%), 225 miRNAs for RC (46.20%) and 269 miRNAs for RS (50.28%) were sequenced more than 50 times. As previously reported, evolutionary conserved miRNAs have generally high expression abundances when compared with non-conserved miRNAs. Among the conserved miRNAs, total reads of miR335, miR157, miR166, miR156 and miR408 accounted for vast majority of total miRNAs; LC (51.22%), 229 miRNAs for LS (48.61%), 225 miRNAs for RC (75.54%) and RS (55.02%). Of these, miR335 was the most abundant miRNA in both control and drought-exposed libraries (Table S2).

Identification of novel miRNAs in peach

After obtaining known miRNAs in peach, the remaining sequences of four libraries, which are classified as “unannotated” (excluding known miRNAs and Rfam matching other non-coding RNAs), were taken into consideration to discover novel and potential peach-specific miRNA candidates. To accomplish this, these small RNA sequences were aligned with the *Prunus persica* genome to identify genomic regions potentially harbouring potential pre-miRNA sequences whose hairpin-like structures are widely used for distinguishing miRNAs from other small non-coding RNAs. The minimum of free energy (MFE) of the secondary structures was also considered to be another criteria for prediction of potential pre-miRNAs. After aligning these unannotated sequences to the genome, we obtained a total of 197, 221, 238 and 265 novel miRNA precursor candidates for LC, LS, RC and RS libraries, respectively (Table S4) and some of these novel miRNA candidates with characteristic features are listed in Table 4. All novel miRNA prediction were carried out according to the default parameters of MIREAP (MicroRNA Discovery By Deep Sequencing) software developed by BGI. In agreement with previously reported results, the uracil nucleotide is dominant in the first position of 5’ end for majority of these newly determined putative novel miRNAs. The first nucleotide bias analysis showed that uracil was the most frequently used first nucleotide in miRNAs of *P. persica*; with 10,528 uracil nucleotides (47%) for LC, 12,834 uracil nucleotides (43%) for LS, 21,571 uracil nucleotides (63%) and 13,014 uracil nucleotides (34%) for RS library (Table S3). Our sequence analysis for all libraries showed that the putative pre-miRNAs of each library greatly varied from 70 to 365 nucleotides in length. With the usage of software mFold, these pre-miRNA sequences were applied to predict the characteristic stem-loop secondary structures of pre-miRNA and their locations were also determined in the genomic loci (Tables S4 and S5). Some of the stem-loop secondary structures of predictive pre-miRNAs of *P. persica* determined via mFold can be found in Figure 2. We also calculated the minimum folding free energies of putative peach miRNA precursors for each libraries; ranging from −18.8 to −157.4 kcal/mol with an average of −53.1 kcal/mol for LC, from −18.3 to −171.2 kcal/mol with an average of −55.2 kcal/mol for LS, from −18.3 to −157.4 kcal/mol with an average of −50.9 kcal/mol for RC and from −18.11 to −181.01 kcal/mol with an average of −50.34 kcal/mol for RS (Table S4). In contrast with the common or evolutionarily conserved miRNAs, the predicted novel miRNAs are often expressed at a very low level as reported before. One possible explanation for this result was that many plant miRNAs are evolutionarily conserved and approximately one hundred miRNA

Table 1. Statistics of small RNA sequences for control and drought stress libraries from *Prunus persica* leaf and root.

| Library | Raw reads | High-quality reads | Clean reads | Unique sRNAs | Total sRNAs mapped to Genome | Unique sRNAs mapped to Genome |
|---------|-----------|--------------------|-------------|--------------|-----------------------------|-----------------------------|
| LC      | 15,521,503 | 15,499,314         | 15,470,689  | 4,210,911    | 10,264,244 (66.35%)         | 2,063,684 (49.01%)         |
| LS      | 12,492,645 | 12,473,137         | 12,428,654  | 3,172,346    | 8,673,228 (69.78%)          | 1,599,019 (50.40%)         |
| RC      | 12,726,680 | 12,703,130         | 12,539,747  | 2,696,057    | 9,324,699 (74.36%)          | 1,400,836 (51.96%)         |
| RS      | 13,233,471 | 13,203,304         | 13,036,443  | 4,125,069    | 8,157,867 (62.58%)          | 1,747,201 (42.36%)         |

Table 2. Classification of small RNA sequences from control and drought stress libraries.

| Category       | Unique (%) | Total (%) | Unique (%) | Total (%) | Unique (%) | Total (%) | Unique (%) | Total (%) |
|----------------|------------|-----------|------------|-----------|------------|-----------|------------|-----------|
| Exon antisense | 66583 (1.58)| 233813 (1.51)| 53440 (1.68)| 176740 (1.42)| 66331 (2.53)| 162078 (1.29)| 61648 (1.49)| 168170 (1.29)|
| Exon sense     | 104977 (2.49)| 534317 (3.45)| 85103 (2.68)| 403706 (3.24)| 102294 (3.79)| 321626 (2.56)| 115637 (2.80)| 379058 (2.90)|
| Intron antisense| 73179 (1.73)| 336082 (2.17)| 54652 (1.72)| 234564 (1.88)| 48174 (1.78)| 170265 (1.35)| 60037 (1.46)| 219032 (1.68)|
| Intron sense   | 99820 (2.37)| 736794 (4.76)| 76536 (1.40)| 508059 (4.08)| 68976 (2.55)| 384332 (3.06)| 82242 (1.99)| 419902 (3.22)|
| miRNA          | 28239 (0.67)| 2004049 (12.95)| 23627 (0.74)| 260206 (20.93)| 24359 (0.90)| 1971298 (15.72)| 31969 (0.77)| 963596 (7.39)|
| rRNA           | 52275 (1.24)| 2571204 (3.40)| 45662 (1.43)| 390134 (3.13)| 87516 (3.24)| 2223692 (17.3) | 20183 (4.90)| 2552133 (19.57)|
| snRNA          | 2030 (0.04)| 7435 (0.04)| 1443 (0.04)| 4365 (0.03)| 4080 (0.15)| 34663 (0.27)| 3827 (0.09)| 24744 (0.19)|
| snoRNA         | 732 (0.01)| 1619 (0.01)| 622 (0.02)| 1294 (0.01)| 837 (0.03)| 3522 (0.02)| 1700 (0.04)| 6349 (0.04)|
| tRNA           | 4841 (0.11)| 83704 (0.54)| 4822 (0.15)| 97317 (0.78)| 8763 (0.32)| 895226 (7.13)| 39282 (0.95)| 303823 (2.33)|
| Unannotated    | 3778235 (89.72)| 11006092 (71.14)| 2826619 (89.1)| 8010411 (64.4)| 2282727 (84.6)| 6373045 (50.82)| 3526154 (85.48)| 7999636 (61.36)|

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conserved miRNA families were found in the Arabidopsis [64]. These evolutionarily conserved miRNAs regulate target transcripts involved in many key metabolic processes which is commonly found in viridiplantea, thus their expression level may be higher than non-conserved miRNAs [65]. Of the 197 putative miRNAs for LC library, only nine miRNAs (LC-m0019, LC-m0049, LC-m0060, LC-m0131, LC-m0146, LC-m0156, LC-m0162, LC-m0164 and LC-m0225) were sequenced more than 500 times, whereas among 221 putative miRNAs belonging to LS library, eleven miRNAs (LS-m0019, LS-m0044, LS-m0051, LS-m0058, LS-m0145, LS-m0181, LS-m0185, LS-m0230, LS-m0234, LS-m0242 and LS-m0256) had more than 500 reads. As for root libraries, eight miRNAs (RC-m0028, RC-m0029, RC-m0077, RC-m0173, RC-m0177, RC-m0200, RC-m0231 and RC-m0253) were sequenced more than 500 reads in control library while eight putative miRNAs (RS-m0025, RS-m0056, RS-m0072, RS-m0153, RS-m0176, RS-m0207, RS-m0244 and RS-m0263) have more than 500 reads in stress library (Table S4).

Genome-wide expression patterns of drought-responsive miRNAs identified in peach

262 and 368 miRNAs were observed with more than 2 fold change response to drought treatment in peach leaf and root, respectively (Table S6; Figure 3). As reported in previous studies [9,42,56,66–72], a series of miRNAs, including miR156, miR159, miR160, miR163/miR166, miR167, miR168, miR169, miR170/miR171, miR390, miR393, miR395, miR396, miR397, miR398, and miR408 are considered to be drought-responsive miRNAs (Figure 4). Our analysis also revealed that miR165 and miR167 were commonly down-regulated in both leaf and root, whereas miR156 was slightly up-regulated in root and leaf after stress treatment. The expression level of miR159, miR169, miR393, miR397, miR398 and miR399 were only decreased in root under drought stress while the miR395 were only down-regulated in leaf in response to drought (Figure 4). The miR160 were solely up-regulated in root, whereas there were no changes in the expression level of miR168, miR390 in leaf and root tissues (Table 5). Since our results largely agree with previous studies [well reviewed in [7,8]], it has been said that peach has its own specific miRNA expression profile under drought-stress.

Target prediction and function analysis

The putative miRNA targets in peach were predicted using BlastN search (v2.2.22) against EST and cDNA sequences in P. persica genome annotation database (http://www.plantgdb.org/XGDB/phplib/download.php?GDB=Pc) based on the rules described in the section of methods. Based on this strategy, a total of 672, 1293, 1194 and 1719 putative miRNA targets were obtained for LC, LS, RC and RS libraries, respectively. Although the number of targets for each miRNA varied considerably among the libraries ranging from 65 to 1 for LC, from 457 to 1 for LS, from 433 to 1 for RC and from 563 to 1 for RS, most of miRNAs in each library have only one predicted target transcripts; LC (30%), LS (30%), RC (32%) and RS (37%) (Table S5). The sequence alignment between miRNA and its predicted target genes are also found in Tables S4 and S5. For comprehensive annotation, all putative target transcripts in each library were analyzed by Gene Ontology (GO) terms with the aid of Blast2GO program with default parameters. Among the peach miRNA targets identified, a total of 571 target transcripts (137 for LC, 133 for LS, 148 for RC and 153 for RS) generated significant GO terms for further analysis. Then, transcripts representing genes with a known function were categorized by biological process, cellular component and molecular function according to the ontological definitions of the GO terms. The putative target transcripts of miRNAs in the biological process category were
related to binding (65 terms for LC, 69 terms for LS, 74 terms for RC and 71 terms for RS), catalytic activity (58 terms for LC, 56 terms for LS, 69 terms for RC and 60 terms for RS), electron carrier activity (1 term for LC, 2 terms for LS, RC and RS), antioxidant activity (1 term for LC), molecular translacer activity (4 terms for LC, 7 terms for LS, 16 terms for RC and 6 terms for RS), transporter activity (3 terms for LC, 2 terms for LS, 4 for RC and 6 terms for RS), structural molecule activity (2 terms for RC and 1 term for RS), enzyme regulatory activity (2 terms for LC, 1 term for LS and RS and 3 terms for RC) and transcription regulator activity (1 term for RC and 2 terms for RS) (Figure 5A).

As shown in Figure 5, most of the miRNA target genes were assigned to the binding category whose present sequences appear to be involved in nucleic acid binding, protein binding and ion binding. Because these sequences encode transcription factors, this is in accord with previously reported notion explained as a large proportion of miRNA targets encode transcription factors.

Table 3. List of conserved miRNAs obtained from control and drought-stresses leaves and roots of *P. persica*.

| miRNA Family | miRNA sequence (5’-3’) | Length (nt) | Count | Fold Change (Log2) | miRNA Orthologs |
|--------------|------------------------|------------|-------|--------------------|-----------------|
|              |                        |            | LC vs LS | RC vs RS | Ath | Osa | Ptc | Mtr | Rco | Vvi |
| miR156       | UGACAGAAGAGAGAUGAGCAC  | 20         | 934112  | 1346965 | 83630 | 81033 | 0.84 | 0.18  | + | + | + | + |
| miR157       | UUGACAGAAGAGAGAGACAG   | 21         | 262981  | 645762  | 606233 | 87444 | 114.05* | -284.94* | + | + | + | - |
| miR159       | UUGUGAAUAGUGAGGCUCUA   | 21         | 3033    | 1889   | 2176  | 489   | -0.36 | -220.98* | + | + | + | + |
| miR160       | CGCUAGGAGGAGGACAGCAUA  | 21         | 2969    | 2235   | 2118  | 13322 | -0.09 | 259.80*  | + | + | + | + |
| miR162       | UGCUAUAACCCUUGCUACCAG | 21         | 321     | 310    | 261   | 92    | 0.26  | -156.03* | + | + | + | + |
| miR164       | UGGGAGAGGAGGAGGAGGUC   | 21         | 13187   | 13085  | 9571  | 3088  | 0.304 | -168.80* | + | + | + | + |
| miR166       | UGGCGAGCCUGAUACUCCCC   | 21         | 144793  | 176391 | 158327 | 109345 | 0.60  | -0.59 | + | + | + | + |
| miR167       | GAAGCGUCGACGAUGAUCUG   | 21         | 92812   | 71576  | 83715 | 5571  | 0.058 | -369.55* | + | + | + | + |
| miR168       | UGCUGUUGGCACUGUGGAGA   | 21         | 20085   | 21168  | 14751 | 16549 | 0.391 | 0.109  | + | + | + | + |
| miR169       | UAGGCGAAAGAUGACUGGU    | 21         | 907     | 804    | 1420  | 94    | 0.141 | -397.31* | + | + | + | + |
| miR171       | UUGGCGGGAGGAAUACUCC    | 21         | 262     | 128    | 90    | 393   | -0.707 | 207.08*  | + | + | + | + |
| miR172       | AGAUCUUGAUGAUGACUGUA   | 21         | 2874    | 1891   | 1719  | 1212  | -0.288 | -0.560  | + | + | + | + |
| miR173       | UGGACUGAAGGGACGCCUC   | 20         | 106     | 124    | 83    | 9     | 0.542  | -326.110 | + | + | + | + |
| miR180       | AGAUCUGAAGGGAGACGCC   | 21         | 4098    | 5471   | 1862  | 2659  | 0.732  | 0.457   | + | + | + | + |
| miR195       | CUGAAGUUGUUGGGGAACUC  | 21         | 120     | 42     | Not Detected | -119.869* | - | + | + | + | + |
| miR196       | GCUCAAGAAGAUCGUGGGGA  | 21         | 2993    | 3033   | 3388  | 539   | 0.355  | -270.811* | + | + | + | + |
| miR197       | UCAUUGAUGUCCAGUUGUA    | 21         | 10464   | 11428  | 36291 | 4151  | 0.443  | -318.412* | + | + | + | + |
| miR198       | UUGUCCUCAGGGUGCCCGUG   | 21         | 1741    | 1569   | 10259 | 586   | 0.165  | -418.588* | + | + | + | + |
| miR199       | GGCCAAAGAAGGUGUGCUCUA | 21         | 103     | 91     | 36    | 27    | 0.137  | -0.471  | + | + | + | + |
| miR203       | UUAUGACUAGCCAAACAGUC   | 21         | 140     | 144    | 109   | 54    | 0.356  | -106.935* | + | + | + | + |
| miR204       | AGCCCCAGAGGAGAAGCAU    | 21         | 77621   | 55268  | 71947 | 34715 | -0.174 | -110.741* | + | + | + | + |
| miR214       | GCAUCCCAUCUACAUCAUGU   | 19         | Not Detected | 58 | 10   | - | -259.206* | + | + | + | + |
| miR215       | AAAGAUCCAAGAAACAGCA    | 21         | 806     | 516    | 2     | 24    | -0.327 | 352.886* | + | + | + | + |
| miR218       | UUAUGUAGUAGAAGAGGAGC   | 22         | 163     | 138    | Not Detected | 0.075 | - | + | + | + | + |
| miR219       | UGAUGUUGCGUCAGAUGAC    | 21         | 52      | 15     | 54    | 66    | -147.767* | 0.233 | + | + | + | + |
| miR220       | AAACUAAACUGGGAAACUG    | 20         | Not Detected | 24 | 0   | - | -758.037* | + | + | + | + |
| miR244       | GGUGUUCUCAAGAUGUUCUC   | 21         | 14      | 149    | 176   | 0    | 372.773* | -1045.485* | + | + | + | + |
| miR272       | UCUUCCCAUCCUAUCACUGCC  | 22         | 833     | 676    | 1408  | 99    | 0.007  | -388.611* | + | + | + | + |
| miR279       | UGUAUUGUGUCGGUCCGUA    | 21         | 43      | 21     | 428   | 0    | -0.718  | 510.039*  | + | + | + | + |
| miR335       | UGACGACAGAGAAGAGACAG   | 21         | 989554  | 1438413 | 604358 | 277182 | 0.855 | -118.061* | + | + | + | + |
| miR327       | UUAUGAUCACUAACAAACA    | 21         | 156     | 51     | 273   | 14  | -129.711* | -434.146* | + | + | + | + |
| miR2118      | CUAAGCCUAGACUACCAUGCA  | 22         | 1358    | 1345   | 4343  | 1897  | 0.301  | -125.101* | + | + | + | + |

Abbreviations: LC; Leaf-control, LS; Leaf-stress, RC; Root-control, RS; Root-stress, Ath, *Arabidopsis thaliana*; Osa, *Oryza sativa*; Ptc, *Populus trichocarpa*; Mtr, *Medicago truncatula*; Rco, *Ricinus communis*; Vvi, *Vitis vinifera*. Note that the asterisk indicates a statistically significant difference between control and drought-stresses samples.

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### Table 4. Potential novel miRNAs found in *Prunus persica*.

| miRNA ID   | Sequence (5p)          | Sequence (3p)       | Length (nt) | Pre-miRNA Length (nt) | Count | Location * | Arm | MFE (kcal/mol) |
|------------|------------------------|---------------------|-------------|-----------------------|-------|------------|-----|----------------|
| LC-m0007   | -                      | UUGUUUAACCGGUGGCUCUA | 23          | 186                   | 7     | scaffold_1:895658038956765 | +   | −60.70         |
| LC-m0026   | GGATGTTTAGTGTGGCGATT   | TACGCGTGGCAACACATCCGG | 21          | 200                   | 248   | scaffold_1:2565112565710    | −   | −75.40         |
| LC-m0057   | -                      | CGAUAUUAUUCUGGCGGCAU | 22          | 197                   | 11    | scaffold_2:26713696718392   | +   | −98.80         |
| LC-m0066   | GGACAUCAUCAAGAUACACA   | -                   | 21          | 105                   | 7     | scaffold_2:22228576622228870 | +   | −51.40         |
| LC-m0085   | -                      | TCTGTGATAGAGCCTTGGAGA | 23          | 343                   | 15    | scaffold_3:304383043830457 | -   | −78.41         |
| LC-m0116   | -                      | UCUAGUUGGGAACAGGGGAG | 21          | 89                    | 18    | scaffold_3:315173951517481 | -   | −47.20         |
| LC-m0129   | -                      | GAUUCGUAGGCAUCAGGAUGG | 23          | 91                    | 9     | scaffold_4:41801175618001846 | +   | −25.90         |
| LC-m0207   | CAGTTGTTGCGTTGGAAATA   | -                   | 23          | 273                   | 8     | scaffold_5:79303778304050   | −   | −59.03         |
| LS-m0011   | GUGGGUGGUGGCAUGCCGGCGA | -                   | 23          | 73                    | 7     | scaffold_1:2302560223025674 | +   | −22.10         |
| LS-m0017   | GUGGUGAGGUGGUGGCGGGAAGA | UUUCCGAAACCUCCAAUCCAA | 22          | 120                   | 233   | scaffold_1:2964606229646181 | +   | −47.20         |
| LS-m0064   | UAAAAAGAAGAAGAUGCUAA   | -                   | 21          | 96                    | 12    | scaffold_1:222728272272832 | +   | −56.10         |
| LS-m0068   | UCAUUAAGGAGCUGCAGAGG   | -                   | 21          | 81                    | 18    | scaffold_2:35636453563725   | -   | −35.10         |
| LS-m0071   | UCAUUAAGGAGCUGCAGAGG   | -                   | 21          | 81                    | 18    | scaffold_2:35636453563725   | -   | −35.10         |
| LS-m0072   | UCAUUAAGGAGCUGCAGAGG   | -                   | 21          | 81                    | 18    | scaffold_2:35636453563725   | -   | −35.10         |
| LS-m0146   | -                      | ACCCGGGAACAGCACCAGAUC | 21          | 104                   | 32    | scaffold_4:42489662424898731 | +   | −76.80         |
| LS-m0229   | AUUUGGCAUUAUUGGGCGAUG | UCCGCGGAUUCUCAAAAGA | 21          | 112                   | 70    | scaffold_2:3198213818921494 | +   | −65.80         |
| RC-m0001   | GGCGCGGGGCCGUGGCGGUGGCG | GUGUUAUAGGUGGACGGCGGGAUG | 22          | 84                    | 9     | scaffold_2:1084263842706   | -   | −36.60         |
| RC-m0006   | UGGGGUGGGAAGGAGCGGCGGAG | -                   | 23          | 94                    | 10    | scaffold_1:3663482367575   | -   | −44.80         |
| RC-m0010   | -                      | ACGUUGCAUGGGAAGAAGUGU | 23          | 132                   | 12    | scaffold_1:421346542133956 | +   | −73.00         |
| RC-m0027   | GGUGAGGAGGAGGCGGGAAGA | UCCGGAACCUCCAUCCCAA | 22          | 120                   | 181   | scaffold_2:12964606229646181 | +   | −47.20         |
| RC-m0030   | -                      | GAAUUGGUGGGGAAGAAGGU | 21          | 84                    | 11    | scaffold_1:3044757130447654 | +   | −26.40         |
| RC-m0032   | UGAUUAUGAAGAACGAGACGGUG | GUCCGAGUCCUAAUAGACAGCG | 23          | 75                    | 11    | scaffold_2:3112339512334056 | +   | −34.30         |
| RC-m0063   | GCTTATGCTTCGTTAGGAGCG | TGAAGGAAAGTAGAAGCGCG | 21          | 127                   | 264   | scaffold_1:1401940241019528 | −   | −79.00         |
| RC-m0146   | TTATCATTATGATCTGTGCGT | -                   | 22          | 116                   | 66    | scaffold_4:418435218434467 | +   | −64.90         |
| RS-m0062   | -                      | UAGGGGAGGAAAGAAGUGA | 20          | 85                    | 31    | scaffold_2:22899244289328   | +   | −42.10         |
| RS-m0084   | AAAAGATTACAGTCCTGGTTACA | -                   | 23          | 344                   | 113   | scaffold_1:23555313555356 | −   | −75.30         |
| RS-m0110   | CGUGGUAUGAUGAAGUGUAUA | -                   | 21          | 100                   | 9     | scaffold_3:385754108575509 | +   | −42.00         |
| RS-m0136   | -                      | UCUAGUUGGAUGAAGGCGGAG | 21          | 89                    | 38    | scaffold_3:315173951517481 | −   | −42.00         |
| RS-m0154   | -                      | ACCCGGGAAGACACGCGACAG | 21          | 104                   | 147   | scaffold_4:324896284289731 | +   | −76.80         |
| RS-m0177   | -                      | UUAUGGUGGGAAGAGCGAAA | 21          | 104                   | 288   | scaffold_5:5190969190172   | +   | −56.80         |
| RS-m0189   | GGATGTTTAGTGTGGCGATT   | TGCAGCATGCTCAACACATCCGG | 21          | 206                   | 180   | scaffold_5:5316931317136 | −   | −84.10         |
| RS-m0263   | AUCAGUGACAGAAGACAGAAG | UGAUUCUGUGAGUACAGUGAUGU | 21          | 104                   | 2085  | scaffold_7:228871342887237 | −   | −69.40         |

*The location of the hairpin precursor(s) on reference of *Prunus persica* genomic scaffolds.

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Drought-Responsive Peach MicroRNAs

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process (9 terms) and response to abiotic stimulus (8 terms) for LC, oxidation-reduction process (12 terms), response to abiotic stimulus (7 terms), small molecule metabolic process (10 terms) and cellular catabolic process (6 terms) for LS, response to salt stress (7 terms), small molecule metabolic process (7 terms) and oxidation-reduction process (12 terms) for RC and oxidation-reduction process (6 terms), cellular response to stress (7 terms), response to salt stress (5 terms) and catabolic process (5 terms) for RS (Figure 5B).

Based on KEGG biochemical pathway analysis, a total of 351, 530, 563 and 597 target transcripts involved in different cellular pathways were determined for LC (124 pathways), LS (216 pathways), RC (211 pathways) and RS (226 pathways) libraries, respectively (Table S7). The pathway analysis for all four libraries show that target transcripts are commonly involved in some cellular pathways including: plant-pathogen interaction (18.75% for LC, 8.73% for LS, 9.95% for RC and 8.23% for RS), metabolic process (6.08% for LC, 8.40% for LS, 8.30% for RC and 7.70% for RS) biosynthesis of secondary metabolites (3.89% for LC, 4.41% for LS, 4.31% for RC and 4.19% for RS) and plant hormone signal transduction (4.22% for LC, 2.99% for LS, 3.60% for RC and 3.29% for RS). It is interesting to note that most of target genes were associated with plant hormone signal transduction because the environmental stress factors such as drought affect plant hormone balance and these biotic environmental stress factors are regulated by transcription factors, which are potential targets of most plant miRNAs.

**qRT-PCR validation of *P. persica* miRNAs and target transcripts**

We applied stem loop quantitative real-time RT-PCR (qRT-PCR) for further experimental verification of the presence of some conserved miRNAs and comparison of the expression pattern of these miRNAs with deep sequencing. Analysis of seven drought-responsive miRNAs by qRT-PCR show that the expression level of miR156 and miR168 were high in leaves and roots under drought stress in comparison to control samples while the expression of miR164 and miR395 was down-regulated in root and leaf tissues of drought-stressed samples. The expression of miR169 was induced in leaf but inhibited in root tissues under drought stress in comparison to control samples while the expression of miR164 and miR395 was down-regulated in root and leaf tissues of drought-stressed samples. The expression of miR169 was induced in leaf but inhibited in root tissues after drought treatment, whereas the expression level of miR171 was induced in root but inhibited in root tissues under drought. As for miR166, although its expression was down-regulated in root tissues in response to drought, the expression level of miR166 were not changed between control and drought stressed leaves of peach (Figure 6). The relative expression profile of miR156, miR164, miR168, miR171 and miR395 using qRT-PCR had a good correlation with deep sequencing. However, there is a discrepancy between the results obtained from deep sequencing and qRT-PCR experiment. Deep sequencing results showed that the expression of miR169 was down-regulated in leaf tissue while qPCR experiment revealed that its expression was up-regulated in leaf tissue after treatment. Rather than experimental methods, duration of drought probably caused expression level differences.

qRT-PCR was also used for detection and quantification of predicted targets of six drought-responsive miRNAs (miR156, miR164, miR166, miR169, miR171 and miR395). Our results...
revealed a negative correlation between the levels of miRNAs and those of their target messages (Figures 6A and 6B). Thus, down-regulation of miRNA could lead to increased expression of its target gene. For instance, the decreased expression of miR164 and miR395 promoted the expression of their targets genes in both root and leaf tissues. Conversely, drought-induced up-regulation of miR156 led to down-regulation of its target gene.

**Discussion**

Among various abiotic stresses, drought is considered to be one of the most detrimental factors to agriculture and adversely influence crop productivity and quality due to its high scale of impact and wide distribution [73]. As land plants are sessile organisms, they cannot escape from unfavourable environmental stress conditions surrounding them. Thus, land plants have to develop various mechanisms at the physiological and molecular
Table 5. The expression level of drought-responsive miRNAs (also, evolutionary conserved) in both leaf and root libraries of *P. persica*.

| miRNA | LC-expressed | LS-expressed | Up/Down          | RC-expressed | RS-expressed | Up/Down          |
|-------|--------------|--------------|-------------------|--------------|--------------|-------------------|
| miR156 | 934112       | 1346985      | ↓ (slightly up-regulated) | 68360        | 81033        | ↓ (slightly up-regulated) |
| miR157 | 262981       | 465762       | ↑ (Up-regulated)   | 606233       | 87444        | ↓ (down-regulated)   |
| miR159 | 3033         | 1889         | Not significantly changed | 2176         | 489          | ↓ (down-regulated)   |
| miR160 | 2969         | 2235         | Not significantly changed | 2118         | 13332        | ↑ (up-regulated)     |
| miR165 | 476          | 261          | ↓ (down-regulated)  | 405          | 119          | ↓ (down-regulated)   |
| miR167 | 92812        | 71576        | ↓ (down-regulated)  | 83715        | 5571         | ↓ (down-regulated)   |
| miR168 | 20085        | 21168        | Not significantly changed | 14751        | 16549        | Not significantly changed |
| miR169 | 907          | 804          | Not significantly changed | 1420         | 94           | ↓ (down-regulated)   |
| miR171 | 262          | 128          | Not significantly changed | 90           | 393          | ↑ (up-regulated)     |
| miR390 | 4098         | 5471         | Not significantly changed | 1862         | 2659         | Not significantly changed |
| miR393 | 91           | 48           | Not significantly changed | 83           | 20           | ↓ (down-regulated)   |
| miR395 | 120          | 42           | ↓ (down-regulated)  | Not detected in root library | Not detected in root library |
| miR396 | 2993         | 3033         | Not significantly changed | 3388         | 539          | ↓ (down-regulated)   |
| miR397 | 10464        | 11428        | Not significantly changed | 36291        | 4151         | ↓ (down-regulated)   |
| miR398 | 1741         | 1569         | Not significantly changed | 10259        | 586          | ↓ (down-regulated)   |
| miR408 | 77621        | 55268        | Not significantly changed | 71947        | 34715        | ↓ (down-regulated)   |

As seen in the table, most of the drought-responsive miRNAs were markedly down-regulated except of miR156. Comprehensive information about the fold-change (log2), p-value, expression level of libraries with normalized value can be found in Table S6.

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Figure 5. miRNA target transcripts molecular function and biological process categories. (A) The pie diagrams demonstrating the significant number of putative peach miRNA targets within the molecular function categories based on the Blast2Go data mining. As shown in Figure 3, the GO hits pertaining to binding and catalytic activity function was overwhelmingly dominant component of all hits. (B) Pie chart illustrating the composition of miRNA-target transcripts (GO term) of each library in the biological-processes categories.

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levels in order to cope with stress. Recently, miRNAs have turned out to be new players in plant tolerance to environmental stresses like drought, cold, heat and high salinity [74] and much effort has been devoted to understanding their role in the responses of drought stress in various plants including; * Hordeum vulgare* [9], *Medicago truncatula* [66], *Populus euphratica* [68], *Vigna unguiculata* [69] and *Oryza sativa* [75]. In the present study, deep sequencing technology was used for quantitative determination of genome-wide miRNA expression patterns of *P. persica* in response to drought. A total of 535 known miRNAs were detected in peach, although 126, 256, 293, 329, 197, 157 and 126 known miRNAs were identified in *M. truncatula* [66], *P. euphratica* [68], *V. unguiculata* [69], *A. hypogaea* [76], *G. max* [77], *P. aphrodite* [78] and *V. amurensis* [79], respectively. Hence, it can be possible to deduce that the small RNA repertoire of peach is relatively richer than other plant species. By comparing the expression level of individual peach miRNAs in drought-stressed tissues to control, the expression level of 453 miRNAs significantly changed in leaf tissue, whereas 368 (221 up-regulated, 147 down-regulated) of the 465 miRNAs had expression levels that significantly changed in root tissue (Table S6). Among these miRNAs, drought-responsive miRNAs (Table 5, Figure 2) were differentially expressed and showed fluctuations in their expression in both peach leaf and root. The expression of miR165 and miR167 was found to be significantly down-regulated in leaf and especially root under drought stress, whereas miR156 was slightly but not significantly up-regulated in drought-stressed tissues. It should be noted that expression of miR165/166 was induced in leaf but inhibited in root tissues of *H. vulgare* after dehydration stress [9] while miR166 was down-regulated in both leaf and root tissues of *P. persica*. Similarly, miR171 expression was up-regulated in leaves of barley while transcript level of miR171 was only increased in the root, but not in the leaf of peach in response to drought. Although miR167 was significantly down-regulated in leaf and root libraries of peach, its up-regulation was observed in the *Arabidopsis thaliana* [42] and *P. euphratica* [68]. Some miRNAs in different plant species display different expression patterns in response to drought; for example, miR168 was up-regulated and down-regulated in *A. thaliana* [42] and *O. sativa* [80], respectively while its expression level was not significantly affected by drought. As another drought-responsive miRNA, miR395 was down-regulated and its expression was restricted to root tissue and was not detected in leaves of peaches (Table 5). As the expression levels of miR159, miR396 and miR397 were down-regulated in peach after treatment, this finding is inconsistent with previous reports suggesting that expression of these miRNAs was up-regulated in both *A. thaliana* and *P. euphratica* [42,68]. Although the measured expression level of the miR166 did not change in leaf tissue of peach under drought stress, the qPCR results indicate that its expression level decreased in root tissue after treatment (Figure 6). This result is consistent with the previous finding that the miR166 was downregulated in roots of barley after drought stress [71].

Because the function of miRNAs appears to be in gene regulation by targeting specific mRNAs for degradation or translation inhibition, identifying the potential target transcripts of miRNAs is crucially important for understanding miRNA-mediated processes such as drought tolerance in plants. Therefore, target prediction analyses were particularly conducted for drought-responsive miRNAs, mentioned above, whose targets generally encode transcription factors and transporters. Among them, miR159 was up-regulated in response to water limitation and was confirmed to target MYB transcription factors (myb33 and myb101) in *Arabidopsis* under drought stress in response to ABA accumulation [81]. However, in contrast to *Arabidopsis*, miR159 was down-regulated in rice [80] and peach root tissue. Although some members of MYB-family transcription factors were found in peach transcriptome libraries (at GDR; Genome database for Rosaceae), we could not determine the Myb
transcription factors as targets for miR159 and this result may be consistent with previous findings that miR139 target was not related to MYB in tomato [92]. Another miRNA, miR160 is known to target three Auxin Response Factors (ARF 10, ARF 16 and ARF 17) in Arabidopsis [83] and it was reported that ARFs are transcription factors binding to auxin-responsive promoter elements to induce or repress auxin-regulated transcripton [84] during the plant development such as root development and branching. After drought treatment, miR160 was up-regulated in peach roots and this miRNA (miR160a and miR160b) was also up-regulated in drought-tolerant cowpea cultivar in response to drought [69]. Thus, the upregulation of miR160 could be important in drought responses among different plant species. It was revealed that the miR169 family members were associated with drought response and high salt stress [52,55]. The miR169 targets a gene family encoding the alpha subunit of CCAAT-binding NFY transcription factors (NFYA) requiring for adaption to drought stress [8]. Two members, expression level of miR169a and miR169c were substantially down-regulated in Arabidopsis via ABA-dependent way [52] and also showed that level of miR169 was decreased under drought stress in M. truncatula by using high-throughput sequencing and qRT-PCR methods [66]. In addition, expression level of miR169 increased in two cowpea genotypes [69]. These results show good concordance with our findings that miR169 was down-regulated in peach after treatment. However, miR169g was up-regulated in rice during drought stress because the miR169g promoter contains two putative DRE (dehydration-responsive) cis-elements, causing the upregulation in response to drought and cold [53]. As an abiotic stress, drought disturbs the balance between ROS production and ROS elimination and thus leads to ROS accumulation in plant cells, which damages nucleic acids, oxidizes proteins and causes lipid peroxidation [85]. The detoxification of ROS radicals were carried out by Superoxide dismutases (SODs). The miR398 regulates the expression level of two Cu/Zn superoxide dismutases (cytosolic CSD1 and chloroplastic CSD2) under drought stress and level of miR398 was down-regulated in both M. truncatula [66] and maize [70], whereas its up-regulation was found in Triticum dicoccoides after 8-h stress [71]. In this study, we found that the level of miR398 was down-regulated in peach after stress treatment. A recently published paper [62] showed that miR408 are up-regulated in response to water deficit in M. truncatula by targeting plantacyanin, and this miRNA was also induced in leaf tissue of H. vulgare under dehydration stress [9]. Contrary to M. truncatula and H. vulgare, expression level of miR408 decreased in peach and this result is consistent with previous finding [67] where they also detected the induction in expression of miR408 upon drought stress in O.sativa. However, it is necessary to specify that plantacyanin, putative target of miR408, was not found in peach transcriptome library during the computational target prediction process, therefore experimental methods such as RLM-RACE or degradome sequencing may be used for accurate determination of miR408 target.

Conclusions

In the present study, genome-wide identified miRNAs and their expression pattern of drought-responsive miRNAs in roots and leaves of P. persica by using high-throughput sequencing. The expression level of 262 (104 up-regulated, 158 down-regulated) of the 435 miRNAs for LC/LS and 368 (221 up-regulated, 147 down-regulated) of the 465 miRNAs for RC/RS changed significantly in response to drought. Among them, drought responsive miRNAs (miR156, miR164, miR166, miR169, miR171, and miR395) were detected and their expression levels were measured by qRT-PCR. Our research also represents the first concerted effort to determine the large-scale small RNA datasets of peach. After sequencing, we identified a total of 531, 471, 535 and 407 known miRNAs and a total of 197, 221, 239 and 265 novel miRNA candidates for LC, LS, RC and RS tissues, respectively. Most of putative target transcripts for these miRNAs in biological process were related to nucleic acid binding (transcription factors) and catalytic activity. These results will greatly contribute to the understanding of post transcriptional gene regulation response to drought stress in peach.

Materials and Methods

Plant materials and stress treatment

P. persica cultivar Françoise plants obtained from in vitro culture clones were grown in plastic pots for one month [86]. Drought-stress treatments were applied to the plants with similar stem length and leaf area for one week by withholding the water until leaves-wrinkle in greenhouse conditions as +24/18°C day/night 16/8 h light/dark. Then, root and leaf tissue samples were collected and immediately frozen in liquid nitrogen and stored at −80°C until RNA isolation.

Total RNA Isolation, small RNA library construction and sequencing

Total RNAs from leaves and roots of control samples and plants exposed to drought stress used in this study was isolated using an TriPure Isolation Reagent (Roche) according to the manufacturer’s protocol. The quality and quantity of purified RNA were assessed by using a Nanodrop ND-2000c spectrophotometer (Nanodrop Technologies, Wilmington, DE, USA). Then, DNase treatment was carried out as described before [9] and all samples were stored at −20°C for miRNA quantification. For each control and stress treatment samples, equal amount of total RNA was pooled from three biological replicates to generate enough RNA (approximately 1000 ng) for deep sequencing. P. persica small RNA library construction, cluster generation and deep sequencing were carried out by the BGI (Beijing Genomics Institute, Hong Kong). Briefly, the isolated total RNAs of each sample was resolved on a denaturing 15% polyacrylamide gel for size selection and these small RNAs (≤30 bases) were ligated to a pair of Solexa adaptors at the 5’ and 3’ ends using T4 RNA ligase. After ligation and purification, adapter-ligated small RNAs were reverse transcribed and 15-cycles amplified (cDNA RT-PCR) with a pair of adapter complementary primers in order to produce sequencing libraries. Then, PCR products were purified and were directly sequenced using Illumina HiSeq 2000 instrument according to manufacturer’s recommendation (BGI, Shenzhen, China).

Bioinformatics analysis of sequencing data and novel miRNA prediction

After the sequencing reactions were complete, the high-quality small RNA reads ranging from 18 to 30 nucleotides were obtained from raw data analysis pipeline including; removing the low quality tags and trimming adaptor sequences so as to identify conserved and novel miRNAs in P. persica. Then, small RNA reads were used as queries to search against the Rfam family database and NCBI Genbank database to remove non-coding RNAs such as rRNA, tRNA, snRNA, snoRNA; the remaining sequences were searched against the miRBase database v18.0 with up to two mismatches to identify “conserved” mature miRNA orthologs. small RNAs not mapped to any miRNAs in miRBase were subsequently analyzed for potential novel miRNAs by the program
MIREAP (developed by BGI) with default parameters for mapping the peach genome and obtaining all candidate precursors with hairpin-like structures of novel miRNA candidates. Secondary structures of novel miRNAs were also checked using MiFold 3.2 [87].

Identifying miRNAs responsive to drought treatment

In order to identify drought-responsive miRNAs and determine their genome-wide expression changes in response to drought, we first compared the gene expression patterns of miRNAs in control and the drought-treated leaf and root tissues in peach. Towards this purpose, we considered the following criteria: (i) adjusted p-value should be less than 0.01 (p-value < 0.01) in at least one data set, (ii) fold change or log₂ ratio of normalized counts between drought and control libraries was greater than 1 or less than −1 in one of the libraries. The frequency of miRNA read counts was normalized as transcripts per million (TPM) and normalization of one of the libraries. The frequency of miRNA read counts was normalized as transcripts per million (TPM) and normalization of one of the libraries. The frequency of miRNA read counts was normalized as transcripts per million (TPM) and normalization of one of the libraries. The frequency of miRNA read counts was normalized as transcripts per million (TPM) and normalization of one of the libraries.

**Normalization formula:**

\[
\frac{\text{Actual miRNA count}}{\text{Total count of clean reads}} \times 10^6
\]

Actual read counts and normalized counts for each miRNA in each library are provided (Table S6). Afterwards, the fold-change between treatment and control and P-value were calculated from the normalized expression using the formula shown below:

**Fold-change formula:**

\[
\text{Fold change} = \log_2(\text{treatment/control})
\]

\[
P \text{- value formula:}
\]

\[
\left(\frac{N_2}{N_1}\right)^{x/y} \times \frac{(x+y)!}{x!y!} \frac{C(y \leq y_{\text{min}} | x) = \sum_{y=0}^{y_{\text{min}}} p(y/x)}{D(y \geq y_{\text{max}} | x) = \sum_{y=1}^{y_{\text{max}}} p(y/x)}
\]

Poisson distribution model was used for estimating the statistical significance of miRNA expression changes under control and treatment conditions. Up-regulation of any miRNA expression levels was considered a positive value while negative values indicate down-regulation.

**miRNA validation and quantification by quantitative stem-loop RT-PCR**

For miR156, miR164, miR166, miR168, miR169, miR171 and miR395, the miRNA stem-loop reverse transcription reaction was performed in a volume of 10 µL containing 2, 20, and 200 ng of total RNA samples of leaf and root samples (1 µL), 0.5 µL 10 mM dNTP mix, 1 µL stem-loop RT primer (1 µM) and 7.5 µL nuclease free water. All those components of the reaction were synthesized 2 µL RT stem-loop cDNA products, quantitative PCR reactions were performed as followed: 10 µL 2× PCR Master mix, 1 µL forward (10 pmol), 1 µL reverse (10 pmol) primers, 0.3 µL (30 nM) reference dye and 10.7 µL nuclease-free water were mixed. With specifically designed forward primers for each individual miRNAs, the universal reverse primer (5’-GTGCAGGGTGCCGAGGT-3’) [30] (Table S8) was designed for all the quantifications. Specified qRT-PCR thermal setup was adjusted as follows: 95°C for 15 min, followed by 40 cycles of 95°C for 5 s, 56°C for 10 s and 72°C for 30 s. All PCR products were denatured at 95°C and cooled to 65°C and the fluorescence signals were accumulated consistently from 65°C to 95°C as the temperature increased at 0.2°C per second. The reactions were repeated at least three times for credible statistical analysis.

**Target transcript validation**

In order to validate and detect expression level of predictedmiRNA target genes which are related to drought stress, qRT-PCR was performed with a number of gene-specific primers. The target transcripts of Ppe-mir156, Ppe-mir166, Ppe-mir168, Ppe-mir169, Ppe-mir171, and Ppe-mir395 were obtained using psRNATarget (user-submitted transcripts and miRNA option) and BlastN algorithms. Specific PCR primers were designed using Primer3Plus software and also, primer dimers and hairpin formations were checked with the Autodimer program (Table S8). At first, complementary DNA (cDNA) was generated from 1500 ng RNA using Superscript III First-Strand Synthesis System (reverse transcriptase, Invitrogen) according to manufacturer's instructions. In brief, the qPCR was performed in a 96-well plate instrument (LightCycler® 480 Instrument II) and in 20 µl reactions that contained 1 µl of this cDNA, 300 nM each of specific forward and reverse primer, and FastStart SYBR Green I master mix (Roche). Each sample was run in biological and technical triplicates for each gene and relative quantity of these target transcripts calculated based on the housekeeping gene 18s rRNA (forward: GTGACGGGTGACGGAGAATT/reverse: GA-GACTAATGGCGCGGGTG) as a normalizer. The qRT-PCR conditions were as follows: preheating for 10 min at 95°C before 40 cycles of 95°C for 30 s, 55°C for 1 min followed by 72°C for 10 min. To eliminate false-positive results, the melting curves of the gained real-time PCR data were analysed for each run and the data of the fluorescence signal were obtained from 55°C to 95°C as the temperature increased at 0.5°C per second.

**Target Sequence annotation, Gene Ontology (GO) classification and KEGG pathway mapping**

Because the majority of plant miRNAs have perfect or near-perfect complementarity with their target sites, the computational methods for finding the putative targets of miRNAs is the preferred way for prediction of conserved and novel peach miRNAs. Therefore, putative mature miRNA sequences were used as query to search against the Prunus persica EST database and high quality cDNA sequence by using BlastN search (http://www.plantgdb.org/XGDB/phplib/download.php?GDB = Pe). Alignments between each miRNA and its putative miRNA target(s) should meet certain criteria as follows: (i) No more than four
mismatches between miRNA and its target (G-U bases count as 0.5 mismatches), (ii) No more than two adjacent mismatches in the miRNA/target duplex, (iii) No adjacent mismatches in in positions 2–12 of the miRNA/target duplex (5’ of miRNA), (iv) No mismatches in positions 10–11 of miRNA/target duplex, (v) No more than 2.5 mismatches in positions 1–12 of the of the miRNA/ target duplex (5’ of miRNA) as noted by Allen [88] and Schwab [11]. The functional annotation and categorization of identified putative miRNA targets were determined using the Blast2Go (B2G) software suite v2.3.1 with the default parameters (http://www.blast2go.com/b2ghome) [89]. Beside, these putative miRNA target sequences were used as query against the KEGG (Kyoto Encyclopedia of Genes and Genomes) database using the KEGG automatic annotation server [90] in order to reveal their biological function in various cellular metabolic pathways. With the aid of KAAS annotation tool, an orthology number (KO) in database was assigned to the genes within KEGG Genes database based on the sequence similarity comparisons and then, the KO numbers associated with the corresponding unique KEGG gene were used for mapping one of the KEGG’s reference metabolic pathways.

Supporting Information

Figure S1 Reads abundance of various categories of small RNAs in each libraries from Prunus persica. (a) Leaf control library, (b) Drought-stressed leaf, (c) Root control library, (d) Drought-stressed root.

Table S1 Summary of data cleaning of small RNA reads produced by Illumina sequencing.

Table S2 Summary of known (conserved and non-conserved) miRNAs in libraries.

Table S3 The graphs representing the nucleotide bias at each position of novel mature miRNA candidates.

Table S4 Novel Prunus miRNA sequences, locations and alignments.

Table S5 Prunus miRNA targets.

Table S6 Read counts and normalized counts for each miRNA in each library.

Table S7 KEGG biochemical pathway analysis of Prunus miRNA target genes.

Table S8 List of primers used for quantification and validation of P. persica miRNAs and their targets.

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Author Contributions

Conceived and designed the experiments: TU. Performed the experiments: VE YB EO. Analyzed the data: UCA SU TU. Contributed reagents/materials/analysis tools: YB. Wrote the paper: VE TU.

References

1. Takhtajan A (1997) Diversity and classification of flowering plants. New York: Columbia University Press.
2. Tani E, Tsaballa A, Stedel G, Kallioniemi C, Papatheophious D, et al. (2011) The study of a SPATULA-like HHLH transcription factor expressed during (Prunus persica) fruit development. Plant Physiol Biochem 49: 6. 654–63.
3. Zhebentyayeva TN, Swire-Clark G, Georgi LL, Garay L, Jung S, et al. (2008) A framework physical map for peach, a model Rosaceae species. Tree Genet Genomes 4: 745–756.
4. Baird WV, Estager AS, Wells J (1994) Estimating nuclear DNA content in peach and related diploid species using laser flow cytometry and DNA hybridization. J Am Soc Hort Sci 119: 1312–1216.
5. Chalmers DJ, Wilson LB (1978) Productivity of peach trees: tree growth and water stress in relation to fruit growth and assimilate demand. Ann Bot 42: 283–294.
6. Rieger M, Duummel MJ (1992) Comparison of drought resistance among Prunus species from divergent habitats. Tree Physiol 11: 369–380.
7. Eldem V, Okey S, Unver T (2012) Plant microRNAs: new players in functional genomics. Turk J of Agr and Forestry, doi:10.3906/tar-1206-50.
8. Sunkar R, Li YF, Jagadeeswaran G (2012) Functions of microRNAs in plant stress responses. Trends Plant Sci 17: 196–203.
9. Kantar M, Unver T, Badalak H (2010) Regulation of barley miRNAs upon dehydration stress correlated with target gene expression. Funct Integr Genomics 10: 493–507.
10. Unver T, Badalak H (2009) Conserv microRNAs and their targets in model grass species Brachypodium distachyon. Planta 230: 659–69.
11. Schwab R, Palumik JF, Riester M, Schmid M, et al. (2005) Specific effects of microRNAs on the plant transcriptome. Dev Cell 8: 317–327.
12. Bartel DP (2004) MicroRNAs: genomics, biogenesis, mechanism, and function. Cell 116:281–297.
13. Carrington JC, Ambros V (2003) Role of microRNAs in plant and animal development. Science 301: 336–338.
14. Zhang B, Pan XP, Cobb GP, Anderson TA (2006b) Plant microRNA: a small regulatory molecule with big impact. Dev Biol 295: 3–16.
15. Chen X (2005) MicroRNA biogenesis and function in plants. FEBRS Lett 579: 5923–5931.
16. Lee Y, Kim M, Han J, Yoon KH, Lee S, et al. (2004) MicroRNA genes are transcribed by RNA polymerase II. EMBO J 23: 4051–4060.
17. Kurihara Y, Watanabe Y (2004) Arabidopsis micro-RNA biogenesis through Dicer-like 1 protein functions. Proc Natl Acad Sci USA 101: 12753–12758.
18. Song L, Han MH, Lesicka, J, Fedoroff N (2007) Arabidopsis primary microRNA processing proteins HYL1 and DCL1 define a nuclear body distinct from the Cajal body. Proc Natl Acad Sci USA 104: 5437–5442.
19. Fang Y, Spectre DL (2007) Identification of nuclear bodies containing proteins for microRNA biogenesis in living Arabidopsis plants.Curr Biol 17: 818–823.
20. Yang L, Liu Z, Lu F, Dong A, Huang H (2006) SERRATE is a novel nuclear regulator in primary microRNA processing in Arabidopsis. Plant J 47: 841–850.
21. Lobbes D, Kallapi G, Schmid DD, Martin C, Clarke J (2006) SERRATE: a novel player on the plant microRNA scene. EMBO Rep 7: 1052–1057.
22. Lin SL, Chang D, Ying SY (2005) Asymmetry of intronic pre-miRNA structures in functional RISC assembly. Gene 356: 32–38.
23. Kosomara A, Griffiths-Jones S (2011) miRBase: integrating microRNA annotation and deep-sequence data. Nucleic acids research 39: D152–D157.
24. Zhang Z, Yu J, Li D, Zhang Z, Liu F, et al. (2010) PMRD: plant microRNA database. Nucleic Acids Res 38: D806–813.
25. Park-W, Li J, Song R, Messing J, Chen X (2002) CARPELI FACTORY, a Dicer Homolog, and HEN1, a Novel Protein, Act in microRNA metabolism in Arabidopsis thaliana. Curr Biol 12: 1484–1495.
26. Reinhardt RJ, Weinstrin EG, Rhoades MW, Bartel B, Bartel DP (2002) MicroRNAs in plants. Genes & Dev 16: 1061–1066.
27. Jagadeeswaran G, Zheng Y, Li YF, Shakal LI, Matts J, et al. (2006) Cloning and characterization of small RNAs from Medicago truncatula reveals four novel legume-specific microRNA families. New Phytol 174: 85–98.
28. Yao Y, Guo G, Ni Z, Sunkar R, Du J, et al. (2007) Cloning and characterization of microRNAs from wheat (Triticum aestivum L.). Genome Biol 8: R96.
29. Ambros V, Lee RC (2004) Identification of microRNAs and other tiny RNA molecules by cDNA cloning. Methods Mol Biol 265: 131–158.
30. Unver T, Parmaksiz I, Dündar E (2008) Identiﬁcation of conserved micro-RNAs and their target transcripts in opium poppy (Papaver somniferum L.). Plant Cell Reports 29:757–769.
31. Unver T, Bakar M, Shearman RC, Budak H (2010) Genome-wide proﬁling and analysis of Festuca arundinacea miRNAs and transcription in response to foliar glyphosate application. Mol Genet Genomics 283: 397–413.
32. Chen C, Ridzon DA, Broome AJ, Zhou Z, Lee DH, et al. (2005) Real-time quantiﬁcation of microRNAs by stem-loop RT-PCR. Nucleic Acid Res 33.
33. Zhang J, Xu Y, Huan Q, Chong K (2009) Deep sequencing of Brachypodium small RNAs at the global genome level identifies microRNAs involved in cold stress response. BMC Genomics 10: 449.

34. Gonzalez-Becas D, Blanco J, Donaire L, Saladé M, Mascarell-Creu A, et al. (2011) Analysis of the melon (Cucumis melo) small RNAome by high-throughput pyrosequencing. BMC Genomics 12:939.

35. Li H, Dong Y, Sun Y, Zhu E, Yang J, et al. (2011) Investigation of the microRNAs in safflower seed, leaf, and petal by high-throughput sequencing. Planta 233: 611–619.

36. Wei B, Cai T, Zhang R, Li A, Hsu N, et al. (2009) Novel microRNAs uncovered by deep sequencing of small RNA transcription in bread wheat (Triticum aestivum L.) and Brachypodium distachyon (L.) Beauv. Funct Integr Genomics 9: 499–511.

37. De Paula D, Cattonaro F, Pignone D, Sonnante G (2012) The miRNAome of globe artichoke: conserved and novel micro RNAs and target analysis. BMC Genomics 13:41.

38. Varaljay E, Burgaya J, Havelda Z (2008) MicroRNA detection by northern blotting using locked nucleic acid probes. Nat Protoc 3: 190–196.

39. Havelda Z (2010) In situ detection of miRNAs using LNA probes. Methods Mol Biol 592: 127–135.

40. Nielsen BS (2012) MicroRNA in situ hybridization. Methods Mol Biol 822: 67–84.

41. Ding Y, Chen Z, Zhu C (2011) Microarray-based analysis of cadmium-responsive microRNAs in rice (Oryza sativa). J Exp Bot 62: 5363–5373.

42. Liu HH, Tian X, Li YJ, Wu CA, Zheng CC (2008) Microarray-based analysis of drought-induced microRNAs in rice. Biochem Biophys Res Commun 354: 585–590.

43. Willenbrock H, Salomon J, Sokilde R, Barken KB, Hansen TN, et al. (2009) Systematic comparison of microarray profiling, real-time PCR, and next-generation sequencing technologies for measuring differential microRNA expression. RNA 16: 991–1006.

44. Willenbrock H, Salomon J, Sokilde R, Barken KB, Hansen TN, et al. (2009) Quantitative miRNA expression analysis comparing microarrays with next-generation sequencing. RNA 15: 2028–2034.

45. Zhang BH, Pan XP, Cox SB, Cobb GP, Anderson TA (2006) Evidence that miRNAs are different from other RNAs. Cell Mol Life Sci 63: 246–254.

46. Axtell MJ, Bartel DP (2005) Antiquity of microRNAs and their targets in land plants. Science 307: 1540–1541.

47. Willenbrock H, Salomon J, Sokilde R, Barken KB, Hansen TN, et al. (2009) Systematic comparison of microarray profiling, real-time PCR, and next-generation sequencing technologies for measuring differential microRNA expression. RNA 16: 991–1006.

48. Axtell MJ, Bartel DP (2005) Antiquity of microRNAs and their targets in land plants. Plant Cell 17: 1658–1673.

49. Lagos-Quintana M, Rauhut R, Lendeckel W, Tuschl T (2001) Identification of novel genes coding for small expressed RNAs. Science 294: 853–858.

50. Lu S, Sun YH, Chiang VL (2008) Stress-responsive microRNAs in Populus. Proc Natl Acad Sci USA 105: 19220–19225.

51. Zhao B, Ge L, Liang R, Li W, Ruan K, et al. (2009) Members of miR-169 family are induced by high salinity and transiently inhibit the NF-YA transcription factor. BMC Mol Biol 10:29.

52. Zhang L, Chia JM, Kumari S, Stein JC, Liu Z, et al. (2009) A genome-wide characterization of microRNA genes in maize. PLoS Genet 5(11):e1000716.

53. Kantar M, Lucas SJ, Budak H (2011) miRNA expression patterns of Triticum dicoccoides in response to shock drought stress. Planta 235: 471–484.

54. Olivier V (2009) Origin, Biogenesis, and Activity of Plant MicroRNAs. Cell 136: 787–799.

55. Barrera-Figueroa BE, Gao L, Dlop NN, Wu Z, Elders JD, et al. (2011) Identification and comparative analysis of drought-associated microRNAs in two coexpressed genotypes. BMC Plant Biol 11:127.

56. Zhao B, Ge L, Liang R, Li W, Ruan K, et al. (2009) Members of miR-169 family are induced by high salinity and transiently inhibit the NF-YA transcription factor. BMC Mol Biol 10: 29.

57. Zhang L, Chia JM, Kumari S, Stein JC, Liu Z, et al. (2009) A genome-wide characterization of microRNA genes in maize. PLoS Genet 5(11):e1000716.

58. Shu J, Yang Q, Chen X, Wang J, Pan L, et al. (2011) Identification and Characterization of microRNAs from Peanut (Arachis hypogaea L.) by High-Throughput Sequencing. PLoS ONE 6(11): e27530.

59. Li H, Dong Y, Yin H, Wang N, Yang J, et al. (2011) Characterization of the stress associated microRNAs in Glycine max by deep sequencing. BMC Plant Biol 11:170–81.

60. An F-M, Hsiao S-R, Chan M-T (2011) Sequencing-Based Approaches Reveal Low Ambient Temperature-Responsive and Tissue-Specific MicroRNAs in Phalaenopsis Orchid. PLoS ONE 6(5): e18937.

61. Wang C, Han J, Liu C, Korir NK, Kayesh E, et al. (2012) Identification of microRNAs from Amur grapes (Vitis amurensis Rupr.) by deep sequencing and analysis of microRNA variations with bioinformatics. BMC Genomics 13: 122.

62. Zhou L, Liu Y, Liu Z, Kong D, Duan M, et al. (2010) Genome-wide identification and analysis of drought-responsive microRNAs in Oryza sativa. J Exp Bot 61: 4157–4168.

63. Jones-Rhoades MW, Bartel DP (2004) Computational identification of plant microRNAs. Plant Cell 16: 621–635.

64. Liu HH, Tian X, Li YJ, Wu CA, Zheng CC (2008) Microarray-based analysis of drought-induced microRNAs in Oryza sativa. Biochem Biophys Res Commun 354: 385–390.

65. Chi X, Yang Q, Chen X, Wang J, Pan L, et al. (2011) Identification and Characterization of microRNAs from Peanut (Arachis hypogaea L.) by High-Throughput Sequencing. PLoS ONE 6(11): e27530.

66. Li H, Dong Y, Yin H, Wang N, Yang J, et al. (2011) Characterization of the stress associated microRNAs in Glycine max by deep sequencing. BMC Plant Biol 11:170–81.

67. Zhang L, Chia JM, Kumari S, Stein JC, Liu Z, et al. (2009) A genome-wide characterization of microRNA genes in maize. PLoS Genet 5(11):e1000716.

68. Zhou L, Liu Y, Liu Z, Kong D, Duan M, et al. (2010) Genome-wide identification and analysis of drought-responsive microRNAs in Oryza sativa. J Exp Bot 61: 4157–4168.

69. Reyes JL, Chua NH (2007) ABA induction of miR159 controls transcript levels of two MYB factors during Arabidopsis seed germination. Plant J 49: 592–606.

70. Zhang L, Chia JM, Kumari S, Stein JC, Liu Z, et al. (2009) A genome-wide characterization of microRNA genes in maize. PLoS Genet 5(11):e1000716.

71. Buxdorf K, Hendelman A, Stav R, Lapidot M, Ori N, et al. (2010) Genome-wide identification and analysis of drought-responsive microRNAs in Oryza sativa. J Exp Bot 61: 4157–4168.

72. Buxdorf K, Hendelman A, Stav R, Lapidot M, Ori N, et al. (2010) Genome-wide identification and analysis of drought-responsive microRNAs in Oryza sativa. J Exp Bot 61: 4157–4168.

73. Liu HH, Tian X, Li YJ, Wu CA, Zheng CC (2008) Microarray-based analysis of drought-induced microRNAs in Oryza sativa. Biochem Biophys Res Commun 354: 385–390.

74. Liu HH, Tian X, Li YJ, Wu CA, Zheng CC (2008) Microarray-based analysis of drought-induced microRNAs in Oryza sativa. Biochem Biophys Res Commun 354: 385–390.

75. Liu HH, Tian X, Li YJ, Wu CA, Zheng CC (2008) Microarray-based analysis of drought-induced microRNAs in Oryza sativa. Biochem Biophys Res Commun 354: 385–390.