

Expression Variations of miRNAs and mRNAs in Rice (*Oryza sativa*)

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Abstract

Differences in expression levels are an important source of phenotypic variation within and between populations. MicroRNAs (miRNAs) are key players in post-transcriptional gene regulation that are important for plant development and stress responses. We surveyed expression variation of miRNAs and mRNAs of six accessions from two rice subspecies *Oryza sativa* L. ssp. *indica* and *Oryza sativa* L. ssp. *japonica* using deep sequencing. While more than half (53.7%) of the mature miRNAs exhibit differential expression between grains and seedlings of rice, only 11.0% show expression differences between subspecies, with an additional 2.2% differentiated for the development-by-subspecies interaction. Expression variation is greater for lowly conserved miRNAs than highly conserved miRNAs, whereas the latter show stronger negative correlation with their targets in expression changes between subspecies. Using a permutation test, we identified 51 miRNA–mRNA pairs that correlate negatively or positively in expression level among cultivated rice. Genes involved in various metabolic processes and stress responses are enriched in the differentially expressed genes between rice *indica* and *japonica* subspecies. Our results indicate that stabilizing selection is the major force governing miRNA expression in cultivated rice, albeit positive selection may be responsible for much of the between-subspecies expression divergence.

Key words: expression variation, microRNA, mRNA, rice.

Introduction

MicroRNAs (miRNAs) are short (~21 nt) non-coding RNAs that regulate target transcripts post-transcriptionally via translational repression or mRNA degradation (Jones-Rhoades et al. 2006; Filipowicz et al. 2008; Bartel 2009). Mature miRNAs are incorporated into RNA-induced silencing complexes (RISCs) and guide RISCs to specific targets through Watson–Crick base-pairing (Jones-Rhoades et al. 2006; Filipowicz et al. 2008; Bartel 2009). A large proportion of miRNAs that were initially discovered in plants regulate transcription factors (Zhang et al. 2006); some of them are highly conserved miRNAs and play important roles in plant development (Carrington and Ambros 2003; Chuck et al. 2009; Carlsbecker et al. 2010), stress adaptation (Lv et al. 2010, Ding et al. 2011), and hormone signaling (Liu et al. 2009). Sequence evolution of miRNAs and their target sites have been extensively explored between closely related plant species (Fahlgren et al. 2010; Ma et al. 2010). For example, rapid

sequence evolution of the miR482/2118 gene family has promoted the evolution of resistance genes in the Solanaceae (de Vries et al. 2015). However, little is known about how expression variation in miRNAs may affect variation in gene expression among individuals, which is an important source of phenotypic variation within species (Britten and Davidson 1971; King and Wilson 1975).

Rice is one of the most important crops in the world. The two rice subspecies, *Oryza sativa* L. ssp. *indica* (*indica*) and *Oryza sativa* L. ssp. *japonica* (*japonica*), show significant phenotypic and genetic differentiations (He et al. 2011). To date, 592 precursor miRNAs, representing 334 miRNA families, have been identified in rice (miRBase, v20.0). MiRNAs play a significant role in regulating rice development and stress responses (Campo et al. 2013). For example, regulation of *OsSPL14* by *OsmiR156* defines ideal plant architecture in rice (Jiao et al. 2010; Luo et al. 2012) and *OsmiR397* substantially enhance grain yield in rice through the repression of *OsLAC*

(Zhang et al. 2013). Moreover, differential expression of miRNAs was reported among *japonica* cv. Nipponbare and *indica* cv. 93-11, probably playing roles in heterosis (Chen et al. 2010). However, variation in miRNA expression among rice accessions and its effects on target and global gene expression remain largely unknown.

Here, we surveyed the expression variation in miRNAs and mRNAs among six rice accessions (three from each of *indica* and *japonica* subspecies) using RNA-seq. In contrast to extensive differential miRNA expression between developmental stages, only a few miRNAs showed expression differences between *indica* and *japonica* subspecies of rice. Highly conserved miRNAs exhibited less expression variation but repressed target gene expression more strongly than lowly conserved miRNAs. Our analysis also identified significantly correlated miRNA–mRNA pairs that differed in expression levels among *indica* and *japonica* cultivars, which may be candidates for adaptive regulatory evolution underlying *indica*–*japonica* differentiation in cultivated rice.

Results

Small RNA Data Processing and Expression Analysis of Rice miRNAs

Small RNA (sRNA) libraries of rice grains and seedlings were constructed and sequenced for three *indica* cultivars Khal Dawk Mali 105, Guangluai 4 and Rathuwee, and three *japonica* cultivars Taipei 309, Heukgyeong and Nipponbare individually. About 9–14 million short reads were obtained for each library, representing an average of 14,849,295 unique reads for *indica* grain library, 12,978,406 for *japonica* grain library, 9,924,065 for *indica* seedling library and 9,395,033 for *japonica* seedling library (supplementary table S1, Supplementary Material online). About 94% of the sRNAs were 20–24 nucleotides (nts) in length, with 21 and 24 nt as the two major size classes (supplementary fig. S1, Supplementary Material online). In plants, most miRNAs are 21-nt in length while 24-nt sRNAs consist mainly of sRNAs that are associated with repeats and transposable elements (TEs) (Axtell 2013). These results indicate that rice has a highly complex sRNA population to which repetitive sequences are the major contributors (Zhu et al. 2008). Small reads matching rice plastid DNA, structural noncoding RNAs (ncRNAs), and repetitive sequences were removed before miRNA annotation. On average, only a small portion of the total reads were mapped to the chloroplast (6.64%) or mitochondria (1.71%) genome, whereas short reads matching ncRNAs and repetitive sequences accounted for 49.3% and 14% of the total reads, respectively. While the sRNAs belonging to the above categories showed an uniform distribution among all 12 libraries, the proportion of signatures that matched miRNA precursors were consistently higher in seedlings (31.3%) than in grain (3.8%), consistent with a previous report (Xue et al. 2009).

To date, 592 rice miRNAs representing 334 families have been registered in the miRBase database (v20.0). Among them, 22 miRNA families were found conserved in both monocots and eudicots previously (Cuperus et al. 2011). We denoted these miRNA families as highly conserved miRNAs and the others as lowly conserved miRNAs in this study. MiRNAs were considered to be expressed by requiring at least ten raw reads in all three accessions from the same developmental stage and the same subspecies. As a result, a total of 272 mature miRNAs were expressed among the 12 libraries, comprising 111 highly conserved miRNAs and 161 lowly conserved miRNAs (table 1). The expression level of each mature miRNA was measured as Reads Per Million (RPM). A pairwise comparison of $\log_2(\text{RPM})$ of all miRNAs across the 12 libraries demonstrated the measurement of miRNA expression is highly reproducible between biological replicates in our data (Pearson's correlation, $r=0.86$ – 0.97 ; supplementary fig. S2, Supplementary Material online). In addition, we also predicted 20 novel miRNAs using the criteria for plant miRNA annotation (Meyers et al. 2008). Most of these novel miRNAs exhibited developmental stage- or subspecies-specific expression and were predicted to regulate 12 target genes (supplementary table S3, Supplementary Material online).

Highly Conserved miRNAs Exhibit More Stable Expression than the Lowly Conserved miRNAs

We first explored the miRNA expression variation across the 12 samples by hierarchical clustering (fig. 1A). Dendrogram classification grouped the samples first by subspecies (*indica*–*japonica*) and then by developmental stage (grain–seedling), indicating that rice miRNA expression, which is largely remodeled between the different developmental stages, exhibits a relatively small difference between the two subspecies. The mature miRNAs were clustered into several clades (fig. 1A). MiRNAs from the basal clades showed high expression levels with similar expression profiles across rice cultivars from different developmental stages and subspecies; most of them are highly conserved miRNAs, including miR156, miR164, miR166, miR167 and miR168, and two are lowly conserved miR535 and miR5794. In contrast, the majority of miRNAs showed grain- or seedling-biased expression between developmental stages. A number of miRNA families, including miR1861, miR1868, miR319, as well as miR444, exhibited higher expression in grains, whereas other miRNAs, such as members of the miR160, miR169, miR172, and miR529 families, mainly showed increased expression in seedlings. These stage-biased miRNAs are promising candidates for developmental regulation in rice (fig. 1A and supplementary table S2, Supplementary Material online).

Using a Log-linear Poisson Regression model (see Methods), 146 (53.7%), 30 (11.0%), and 6 (2.2%) of the 272 expressed mature miRNAs exhibited significantly differential expression for the factors developmental stage, subspecies

Table 1

Summary of Differentially Expressed miRNA Families

Effect	Highly Conserved (n = 111)	Lowly Conserved (n = 161)	Total (n = 272)
Developmental stage	64 (57.7%)	82 (50.9%)	146 (53.7%)
Subspecies	2 (1.8%)	28 (17.4%)	30 (11.0%)
Development × Subspecies	0 (0%)	6 (3.7%)	6 (2.2%)

NOTE.—MiRNAs with fold change ≥ 2 and $FDR \leq 0.05$ are defined as differentially expressed.

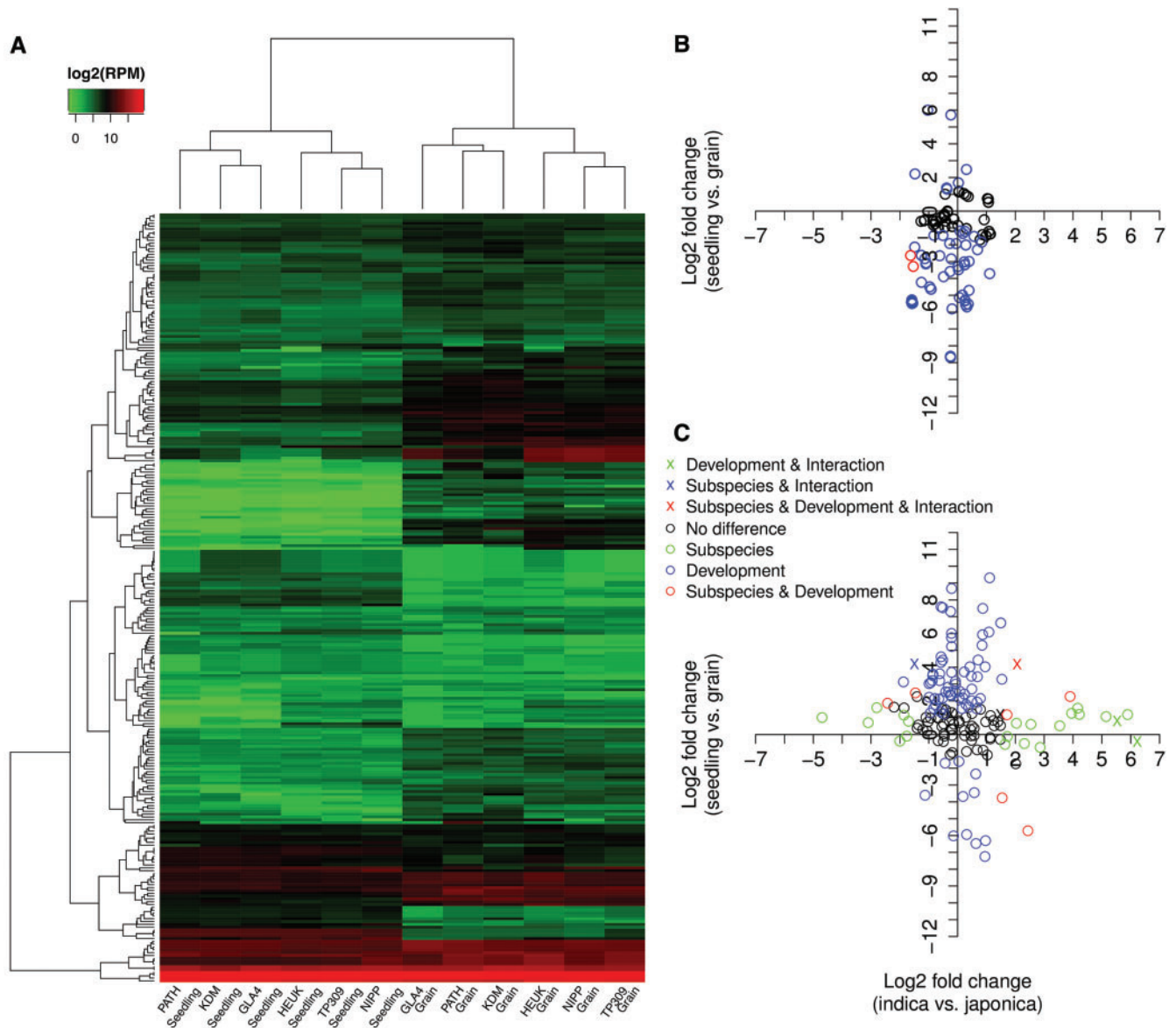


FIG. 1.—Expression variation of known miRNAs in rice. (A) Heat map and unsupervised hierarchical clustering of known miRNA expression. The color key represented the scale of the relative expression levels of the miRNAs (log₂ RPM). KDM: *indica* cv. Khal Dawk Mali 105; GLA4: *indica* cv. Guangluai 4; PATH: *indica* cv. Rathuwee; TP309: *japonica* cv. Taipei 309; HEUK: *japonica* cv. Heukgyeong; NIPP: *japonica* cv. Nipponbare. (B, C) Scatter plot of differentially expressed highly conserved (B) and lowly conserved (C) miRNAs. A generalized Poisson-regression linear model was used to identify the differentially expressed miRNAs for the factors of development, subspecies, and development-by-subspecies interaction. MiRNAs with fold change ≥ 2 and $FDR \leq 0.05$ are denoted as significantly differentially expressed. Mature miRNAs that show no differential expression (black) or show significant differential expression between subspecies (green), developmental stage (blue) and both (red) are indicated by circles in different colors, while those with differential expression for the additional factor of development-by-subspecies interaction are indicated by crosses with the same color setting.

and development-by-subspecies interaction (fold change ≥ 2 , $FDR \leq 0.05$), respectively (table 1). These miRNAs were considered to play important roles in regulating rice development or subspecies differentiation. For example, miR159a-f showed a grain-biased expression (supplementary table S2, Supplementary Material online) and is involved in ABA-mediated responses in the hormonal and abiotic stress signaling networks (Reyes and Chua 2007); members of miR156, miR166, miR169, miR171, and miR444 families showed a leaf-biased expression (supplementary table S2, Supplementary Material online) and are known to play defensive roles against stress by targeting transcription factors (Macovei et al. 2012).

When grouping the mature miRNAs according to their conservation, expression variation was less prominent in highly conserved miRNAs (fig. 1B) than lowly conserved miRNAs (fig. 1C). Only 1.8% (2/111) of highly conserved miRNAs were differentially expressed between subspecies while the proportion was 17.4% (28/161) for lowly conserved miRNAs. An additional 50.9% (82/161) and 3.7% (6/161) of lowly conserved miRNAs were also differentially expressed between developmental stages or development-by-subspecies interaction (table 1 and fig. 2A). Furthermore, magnitudes of the expression fold changes are much higher for lowly conserved miRNAs than highly conserved miRNAs (fig. 1B vs. C

and fig. 2B). These results strongly suggest that variation in expression is higher for the less conserved miRNAs than for the more conserved miRNAs found in both monocots and dicots. A possible explanation for this result is that conserved miRNAs are processed better from their foldback structures (Shen et al. 2011). Another explanation could also be that conserved miRNAs are under strong functional constraints (Cuperus et al. 2011). Using qRT-PCR, we validated the differential expression between subspecies for eight out of nine lowly conserved miRNAs in rice seedlings but not for the highly conserved miR166j-5p due to its low expression (supplementary fig. S3A and C).

Differential Expression of Transcriptome between *Indica* and *Japonica* Cultivars

To understand how miRNA expression may affect mRNA expression, we conducted RNA-seq for each of the seedling samples with 30 \times coverage (supplementary table S1, Supplementary Material online). We report the results of the whole transcriptome analysis here and analyzed the expression correlation between miRNAs and targets with the transcriptome as a control in the next two sections. All reads were mapped to the *japonica* cv. Nipponbare genome (Kawahara et al. 2013). Approximately 81% and 86% of the unique reads could be mapped to the reference genome for *indica*

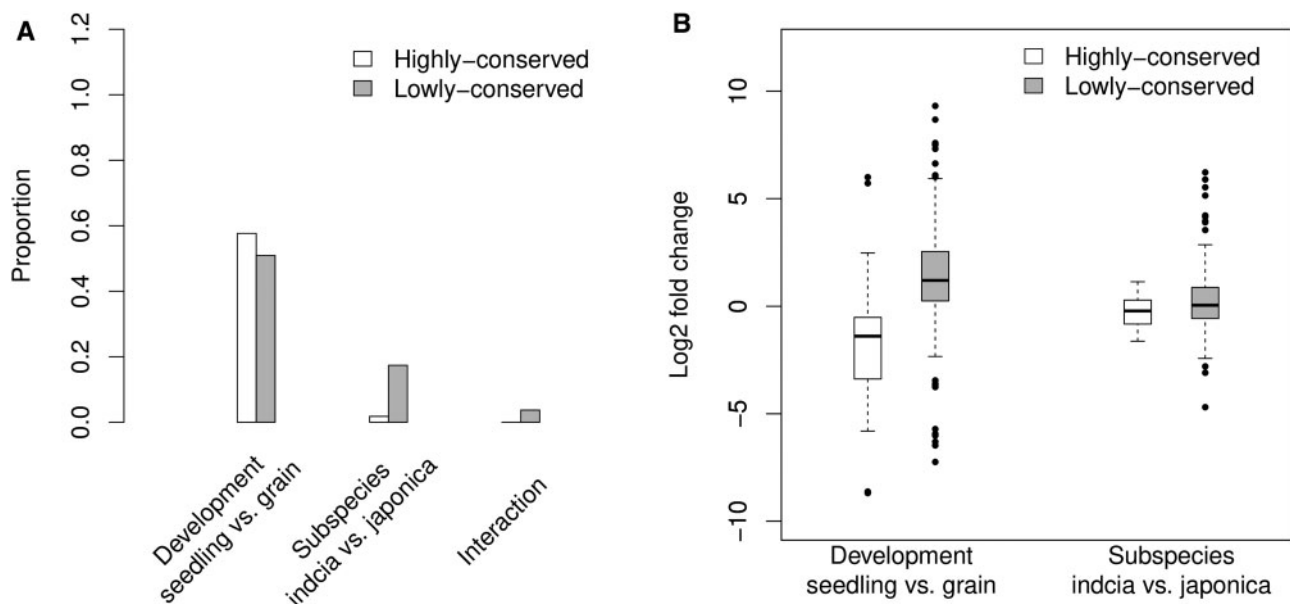


Fig. 2.—Expression variation of the highly conserved and lowly conserved miRNAs between subspecies or developmental stages. (A) The proportions of differentially expressed miRNAs in both sets of the highly conserved and lowly conserved miRNAs. A generalized Poisson-regression linear model was used to identify the differentially expressed miRNAs for the factors of development, subspecies, and development-by-subspecies interaction. MiRNAs with fold change ≥ 2 and $FDR \leq 0.05$ are denoted as significantly differentially expressed. MiRNAs that show significantly differential expression between subspecies or interactions are enriched in the lowly conserved miRNAs (Fisher's Exact Test, P -value $\ll 0.01$). (B) Fold changes in expression of the highly conserved and lowly conserved miRNAs between subspecies or developmental stages. The lowly conserved miRNAs exhibit significantly more variation in expression than the highly conserved miRNAs for both comparisons (Kolmogorov–Smirnov test, P -value $\ll 0.01$).

and *japonica* cultivars, respectively. To allow differential expression analysis between rice subspecies, 32,044 homologous genes were identified between the genomes of *japonica* cv. Nipponbare and *indica* cv. 93-11 (see Methods), of which 18,288 were considered to be expressed, i.e., having more than ten raw reads across all three accessions from the same developmental stage and the same subspecies (supplementary table S4, Supplementary Material online). A regression comparison was then performed based on the expression level of these 18,288 expressed homologous genes. Among them, 2,530 genes (13.8%) were differentially expressed (fold-change ≥ 2 and FDR $\leq 5\%$) between the two rice subspecies, with 1,452 up-regulated in *indica* and 1,078 in *japonica*, respectively (supplementary fig. S4, Supplementary Material online). To confirm the results of RNA-seq, we conducted qRT-PCR of ten experimentally validated or predicted miRNA targets and confirmed the differential expression patterns between rice subspecies in all the comparisons (supplementary fig. S3B and C, Supplementary Material online).

The differentially expressed genes (DEGs) are significantly enriched in Gene Ontology categories, including “plasma membrane,” “cell wall,” “endoplasmic reticulum,” and “extracellular region” for “Cellular Component”, categories of molecular binding (“Hydrocarbon binding”, “Oxygen binding”, “protein binding” etc.) and enzyme activities (“hydrolyase activity,” “catalytic activity,” “kinase activity” etc.) for “Molecular function,” and “cellular process,” “flower development,” and many categories of metabolic process (“Secondary metabolic process,” “lipid metabolic process,” etc.) or response to stimuli and stress (“endogenous stimulus,” “abiotic stimulus,” and “biotic stimulus”) for “Biological Process” (fig. 3A). Consistent with the GO analysis, DEGs are significantly enriched in the KEGG pathways related to secondary metabolites, such as “Glutathione metabolism” (supplementary fig. S5) and “starch and sucrose metabolism” (supplementary fig. S6, Supplementary Material online and fig. 3B). The former is known to play important roles in plant stress tolerance (Noctor et al. 1998), while the latter is presumably affecting the varietal difference of soluble sugar in different rice varieties (Yang et al. 2014). The over-representative KEGG pathways in DEGs also included “Plant–pathogen interaction” (supplementary fig. S7, Supplementary Material online) and “Plant hormone signal transduction” (supplementary fig. S8, Supplementary Material online and fig. 3B).

MicroRNAs, Particularly Highly Conserved miRNAs, Show a Negatively-Correlated Expression Pattern with Their Direct Targets

As miRNAs negatively regulated target gene expression, one may expect to see negative correlation between expression of miRNAs and target mRNAs. To test this speculation, we performed correlation analyses either for all the miRNAs and their targets globally using the log₂ fold changes of expression

between *indica* and *japonica* subspecies, or for individual miRNA-target pairs using their expression levels across six rice cultivars. We shall focus on the global analysis in this section and present the results of the individual analysis in the next. Such analyses were based on four target sets, including one predicted target set for all the expressed miRNAs (target set I; see “Methods” for rice miRNA target prediction) and three experimentally verified target sets by high-throughput degradome sequencing for *japonica* cv. Nipponbare (target sets II, III) or *indica* cv. 93-11 (target set IV) (Wu et al. 2009; Li et al. 2010; Zhou et al. 2010).

As for the global analysis, no correlation (Pearson’s correlation hereafter, P -value = 0.420, $r = -0.033$, $n = 609$) was observed for all the expressed miRNAs and their predicted targets (target set I) (supplementary fig. S9A, Supplementary Material online). However, when separating miRNAs according to their conservation, a weak but significant negative correlation was observed for the highly conserved miRNAs and their predicted targets (P -value = 0.019, $r = -0.119$, $n = 390$; fig. 4A), but not for the lowly conserved miRNAs and targets (P -value = 0.611, $r = 0.035$, $n = 219$; fig. 4B). The negative correlation in expression fold changes was highly significant for the co-expressed miRNAs and degradome-verified target set IV (Zhou et al. 2010) (P -value = 0.002, $r = -0.364$, $n = 68$; fig. 4C), and nearly significant for target set II (Wu et al. 2009) (P -value = 0.121, $r = -0.224$, $n = 49$; fig. 4D) and target set III (P -value = 0.119, $r = -0.161$, $n = 95$; supplementary fig. S9B, Supplementary Material online) (Li et al. 2010). The extent of negative correlation seemed to be associated with the proportion of targets of highly conserved miRNAs in the analyzed target sets, which is 86%, 81%, and 91% for target set II, III, and IV, respectively. Taken together, the overall expression of miRNAs correlates negatively with the expression of their targets in rice, which is consistent with miRNA functions in guiding mRNA cleavage in plants (Bartel 2004). Highly conserved miRNAs exhibit less variation in expression but repress targets more strongly than lowly conserved miRNAs.

We then compared the *indica*–*japonica* expression differentiation of miRNAs relative to that of target genes using transcriptome as a control. Interestingly, the extent of differential expression between rice subspecies was much greater for miRNAs (median absolute fold change = 0.630) than for the whole transcriptome (median absolute fold change = 0.263). Kolmogorov–Smirnov test (KS test) on the cumulative plot shows that the differences are highly significant (P -value $\ll 0.001$; supplementary fig. S10A, Supplementary Material online). In contrast, the extent of differential expression was comparable between miRNA targets (median absolute fold change = 0.279) and the transcriptome (median absolute fold change = 0.263, supplementary fig. S10B, Supplementary Material online). Although lowly conserved miRNAs (median absolute fold change = 0.677) were more variable in expression than highly conserved miRNAs (median absolute fold change

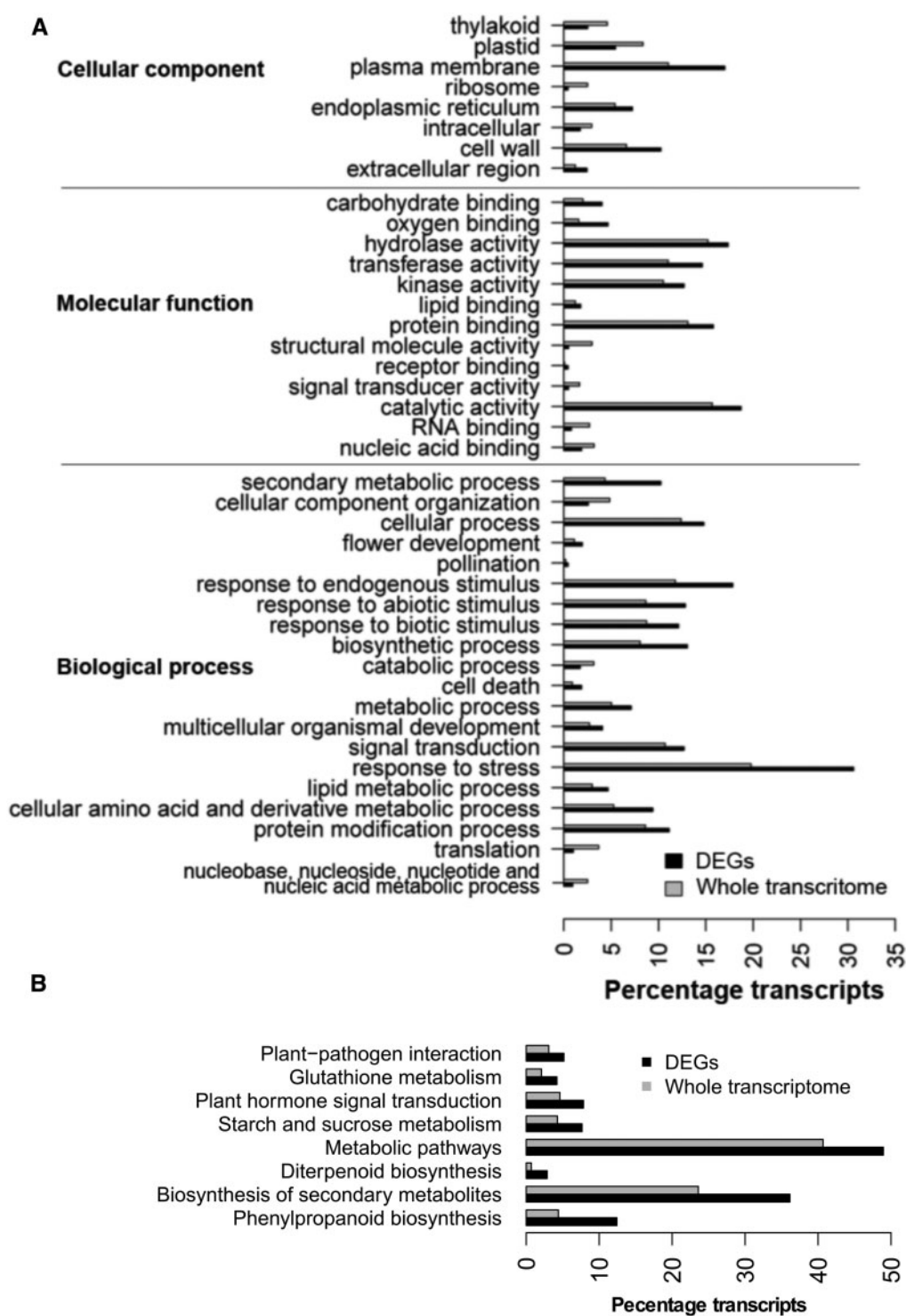


Fig. 3.—Gene ontology (GO) and KEGG pathway enrichment analyses of DEGs. The DEGs ($FDR \leq 0.05$) with a fold change larger than 2 or 1.5 were used for the enrichment analyses of GO terms and KEGG pathways, respectively. The significantly over-represented and under-represented GO terms (A) and KEGG pathways (B) with a $FDR \leq 0.05$ were presented. Grey and black bars indicate the percentages of DEGs and the whole transcriptome that were classified into different functional annotations, respectively.

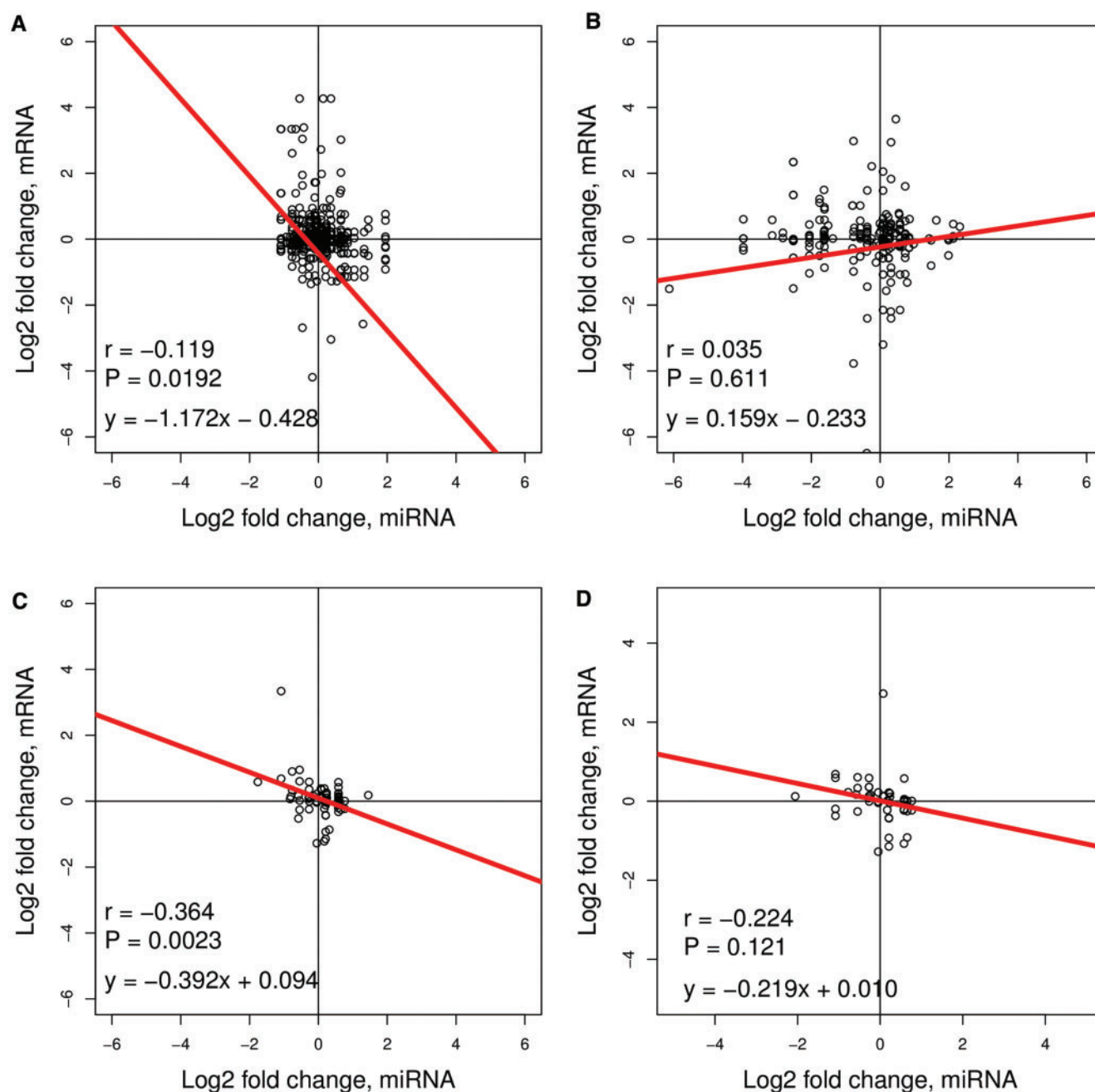


FIG. 4.—Correlation between the coexpressed miRNAs and their targets in seedlings. (A) highly conserved miRNAs and their predicted targets (390 pairs); (B) lowly conserved miRNAs and their predicted targets (219 pairs); (C) miRNAs and the degradome-verified targets in target set IV (Zhou et al. 2010) (68 pairs) and (D) miRNAs and the degradome-verified targets in target set II (Wu et al. 2009) (49 pairs). The log₂ fold changes of miRNA or mRNA expression between rice *indica* and *japonica* subspecies in seedlings were used for Pearson's correlation analysis.

=0.457, KS test P -value $\ll 0.001$, [supplementary fig. S10A](#), [Supplementary Material](#) online), no such expression difference was observed for their targets (KS test, P -value = 0.926, [supplementary fig. S10B](#), [Supplementary Material](#) online). It is thus evident that miRNA targets exhibit less variation in expression than miRNAs.

Identification of Individual miRNA-Target Pairs with Significant Expression Correlations

We further examined the individual correlations of the expression of miRNA–mRNA pairs across the six accessions. The average correlation coefficient for the 476 miRNA–target pairs (in target set I) was -0.11 . The highly conserved miRNAs tend

to be more negatively correlated with their corresponding target mRNAs than the lowly conserved miRNAs in expression variation among accession (KS test, P -value = 0.0008, fig. 5A) but no pair passed the 0.05 FDR level after the multiple-testing correction.

We then asked whether the mean expression correlation among all miRNA-target pairs shifted to the negative end in comparison with the random miRNA-gene pairs. Using a permutation test as previously described (Nunez-Iglesias et al. 2010), we identified 51 significantly correlated miRNA-target

pairs, including 29 negatively correlated and 22 positively correlated pairs (table 2). Indeed, the mean correlation of the real miRNA-mRNA pairs is more negative than the permuted pairs (empirical P -value \ll 0.001, fig. 5B), both for the highly conserved (empirical P -value \ll 0.001, fig. 5C) and lowly conserved miRNAs (empirical P -value = 0.019, fig. 5D). Therefore, the overall tendency in the individual miRNA-target correlations is towards the negative correlation, which is in contrast to a similar test performed in human brain samples (Nunez-Iglesias et al. 2010). Also, the degradome verified targets are significantly

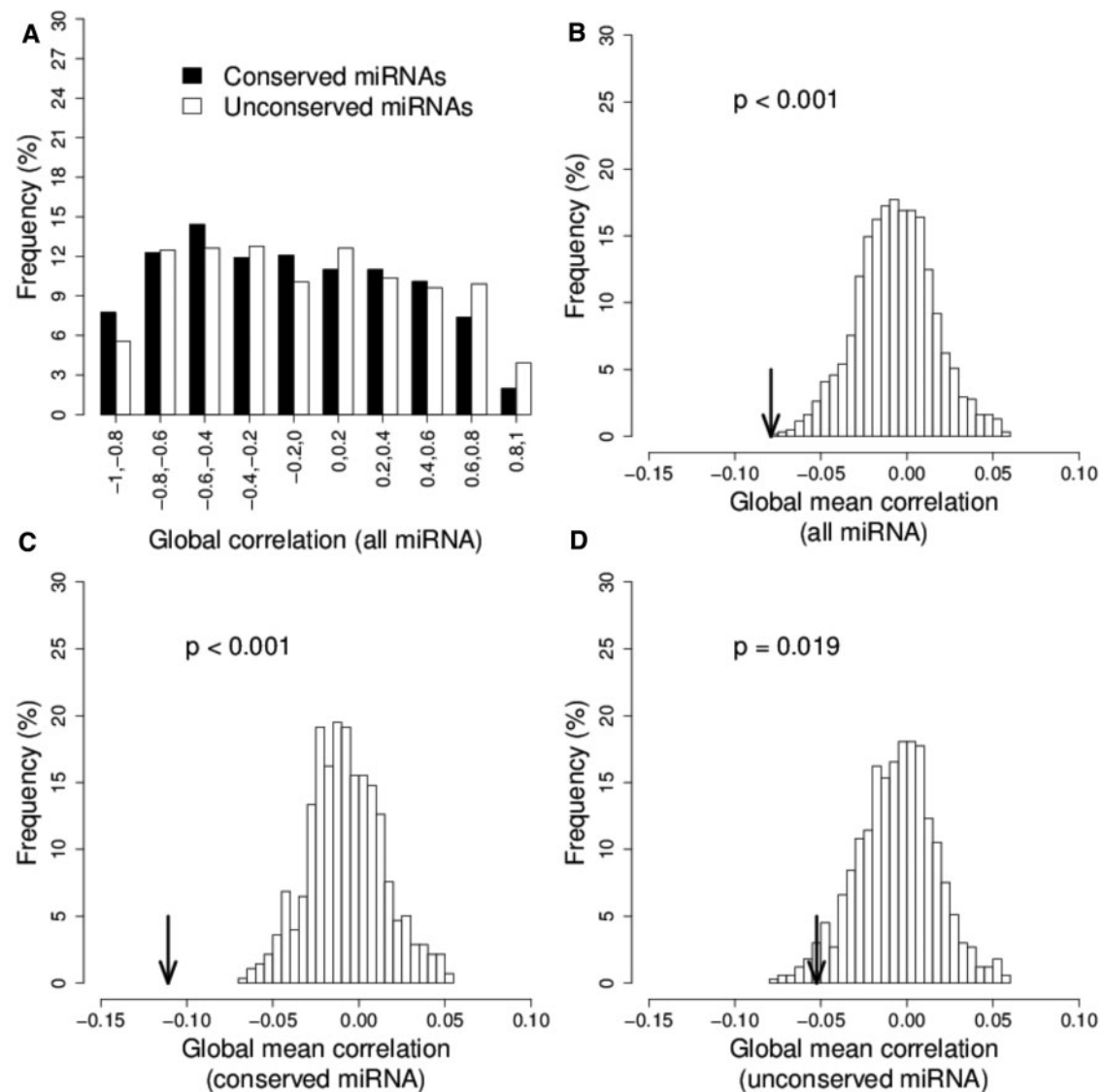


Fig. 5.—Permutation of miRNA-mRNA target relationships at the lineage level. (A) The empirical distribution of the Pearson's correlation coefficient values for 436 miRNA-mRNA pairs between expression levels of 96 miRNAs and those of their target mRNAs across 6 lineages. (B) The histogram plot represents the distribution of the global mean correlation values for the expression levels of all miRNA-mRNA pairs for 1,000 permutations, (C) for the highly conserved miRNAs and (D) for the lowly conserved miRNAs. The black arrowhead indicates the true value.

enriched in the correlated pairs (27 verified targets, Chi-squared test P -value $\ll 0.001$, table 2), suggesting that those genes are indeed under the control of miRNA regulation.

Significantly correlated miRNA-target pairs, both negatively and positively, participate in key biological processes such as reproduction, development, pigmentation, and stress responses (supplementary fig. S11, Supplementary Material online). The negatively-correlated pairs included the well-documented miRNA-target pairs, such as miR156:*SQUAMOSA promoter binding protein-like (SPL)* (Miura et al. 2010); miR169:*Nuclear transcription factor Y (NF-Y)* (Zhao et al. 2009), miR172:*Apetala2 (AP2)* (Zhu and Heliwell 2011), and miR396:*Growth-regulating factor (GRF)* (Debernardi et al. 2012) (table 2). The involvement of the positively correlated miRNA-target pairs in cell cycle, cell death, pollination and the transport process suggest that these pairs also play significant roles in rice development. Positive correlations between the miRNA and target expression levels across rice cultivars may be due to the miRNA-mediated regulatory circuits such as negative feedback loops or incoherent feedforward loops (Wu et al. 2009).

Discussion

We present here the first survey of miRNA expression variation in rice cultivars. Substantial miRNA expression changes were detected between rice grains and seedlings, consistent with the regulatory role of miRNAs in rice seed development (Zhu et al. 2008). In contrast, only a small fraction of miRNAs, mainly lowly conserved miRNAs, differed in expression level between rice *indica* and *japonica* subspecies. While miRNA genes are under strong purifying selection (Ehrenreich and Purugganan 2008; Wang et al. 2010), the single nucleotide polymorphisms (SNP) densities of pre-miRNAs and mature miRNAs are significantly higher than their flanking regions in rice (Liu et al. 2013), implying that *cis*-regulatory mutations affecting miRNA expression level are mostly deleterious and are thus quickly purged from the rice population. The low level of miRNA expression variation coupled with low level of *cis*-regulatory sequence polymorphism is consistent with the scenario that stabilizing selection commonly uses purifying selection to select against extreme values of characters, i.e., miRNA expression profiles in this study. Indeed, it has been demonstrated both empirically and theoretically that stabilizing selection is the major evolutionary force governing the evolution of gene expression (Denver et al. 2005; Rifkin et al. 2005; Hutter et al. 2008; Bedford and Hartl 2009; Hodgins-Davis et al. 2015). A recent study revealed that a vast majority of miRNAs are under stabilizing selection at the onset of *Drosophila* metamorphosis (Yeh et al. 2014). Using the same methodology, we estimated that the counts for rice miRNAs with expression variation compatible with particular evolutionary modes are 196, 29, 1, and 46 for stabilizing selection, directional selection, genetic drift and complex

scenarios, respectively (see supplementary text, Supplementary Material online). About 72% of miRNAs in this study are not significantly differentiated within or between rice subspecies, compatible with the evolutionary mode of stabilizing selection. These miRNAs under stabilizing selection are mostly conserved between eudicots and monocots, whereas more than half of the miRNAs under directional selection are species-specific to rice.

It is remarkable that lowly and highly conserved miRNAs showed sharply contrasting patterns in expression variation and regulation strength. The great expression variation of the lowly conserved miRNAs is largely coupled with their high level of sequence polymorphism among cultivated rice (Liu et al. 2013), suggesting they are under weak selection pressures. Such a coupling of expression variation and sequence polymorphism of rice miRNAs is compatible with the correlated divergences between gene sequences and expression patterns during organ evolution in angiosperms (Yang and Wang 2013). The negligible overall correlation between expression changes of the lowly conserved miRNAs, also the lowly expressed miRNAs, and their targets further indicates that these miRNAs exert very modest, if any, repression on target genes. Therefore, the lowly conserved miRNAs are more like young miRNAs, which are expressed lowly or in specialized tissues, evolve rapidly, and tend to be lack of targets (Rajagopalan et al. 2006; Fahlgren et al. 2010; Ma et al. 2010), rather than the old, deeply conserved miRNAs.

While most young miRNAs are evolutionarily transient (Fahlgren et al. 2010), they can occasionally be selectively favored. A good case in point is a new miRNA specific in *japonica* rice, *osa-miR7695*, which negatively regulates an alternatively spliced transcript of *OsNramp6* (Natural resistance-associated macrophage protein 6), conferring pathogen resistance (Campo et al. 2013). The differentially expressed miRNAs between *indica* and *japonica* subspecies, which are resulted from low expression polymorphism within subspecies and high expression divergence between subspecies, may represent a class of miRNAs that are favored by artificial selection in rice domestication and/or improvement. Previous evolutionary analyses have identified rice miRNA genes that are putative candidates of positive selection, including highly conserved *MIR164e*, *MIR395a/b*, and *MIR399d* (Wang et al. 2010), and lowly conserved or rice-specific *osa-miR5513*, *osa-miR818e*, *osa-miR1847*, *osa-miR1865*, *osa-miR160f*, *osa-miR5143*, and *osa-miR2118h-l* (Liu et al. 2013). Among them, *osa-miR395a-v,y* and *miR399a-d,i-j* (unadjusted P -value < 0.05 , FDR ≈ 0.2 , supplementary table S2, Supplementary Material online) showed significant differential expression between *indica* and *japonica* subspecies and the former correlated negatively with a putative target *LOC_Os03g53230* in this study. In the Solanaceae, differential expression of the *miR482/2188* gene family members that are under different evolutionary constraints also suggested

Table 2

Significantly Correlated miRNA–mRNA Pairs in the Permutation Test

miRNA	miRNA FC ^a	mRNA FC ^b	Correlation ^c	W	Target	TIGR Annotation ^d	Target Verification ^e
miR156k	0.080	0.142	0.953	2.082	LOC_Os01g69830	OsSPL2-SBP-box gene family member	Y ^{3,4}
miR160a–d	–0.048	–0.040	–0.815	–2.096	LOC_Os06g47150	Auxin response factor 18	Y ^{2,3,4}
miR160e	0.537	–0.040	–0.817	–1.923	LOC_Os06g47150	Auxin response factor 18	Y ^{2,3,4}
		0.222	0.873	1.573	LOC_Os04g43910	Auxin response factor	Y ^{2,3,4}
miR160f	0.379	0.222	0.702	1.593	LOC_Os04g43910	Auxin response factor	Y ^{2,3,4}
miR166a–d,f,n	0.073	2.722	0.888	1.845	LOC_Os04g48290	MATE efflux family protein	Y ²
miR166g,h	0.203	0.383	0.791	1.427	LOC_Os03g43930	START domain containing protein	Y ^{3,4}
miR166m	–0.252	–0.223	0.864	2.339	LOC_Os08g34740	SGT1 protein	N
miR168a	0.624	–1.165	–0.902	–1.639	LOC_Os11g44860	Cysteine-rich receptor-like protein kinase 28 precursor	N
miR169f–g	0.661	–0.930	–0.940	–1.922	LOC_Os03g29760	Nuclear transcription factor Y subunit	Y ^{2,3,4}
miR169h–m	0.846	–1.145	–0.948	–1.747	LOC_Os07g41720	Nuclear transcription factor Y subunit	Y ^{2,3,4}
miR171b–f	0.242	–0.126	–0.774	–1.880	LOC_Os05g34460	OsDegp7 - Putative Deg protease homologue	N
		0.098	0.668	1.649	LOC_Os02g44370	Myosin	Y ^{2,3,4}
miR171i	–0.426	–0.375	0.882	1.933	LOC_Os03g04300	Targeting protein-related	N
miR172b	–0.175	0.105	–0.867	–1.747	LOC_Os05g03040	AP2 domain containing protein	Y ^{2,3,4}
		0.321	–0.772	–1.593	LOC_Os03g44420	Tubulin/FtsZ domain containing protein	N
miR172c	0.072	–0.044	0.855	1.921	LOC_Os07g13170	AP2 domain containing protein	Y ^{2,3,4}
miR319a–b	0.171	–0.065	0.930	2.056	LOC_Os08g16660	Aspartic proteinase nepenthesin precursor	N
miR393	0.766	–1.128	–0.695	–1.645	LOC_Os03g36080	Expressed protein	Y ³
miR395b,d–e,g–n,p–s,y	1.449	–0.505	–0.972	–1.944	LOC_Os03g53230	Bifunctional 3-phosphoadenosine 5-phosphosulfate synthetase	N
miR396d–e	–0.035	0.053	–0.991	–2.419	LOC_Os06g02560	Growth-regulating factor	Y ^{2,3,4}
		0.129	–0.855	–1.999	LOC_Os03g47140	Growth regulating factor protein	Y ^{3,4}
		–0.325	–0.812	–1.887	LOC_Os02g53690	Ankyrin repeat domain containing protein	Y ^{3,4}
		0.241	–0.749	–1.690	LOC_Os04g51190	Growth-regulating factor	Y ^{3,4}
		–0.008	–0.693	–1.654	LOC_Os11g35030	Growth regulating factor protein	Y ^{2,3,4}
miR396f	–0.054	0.053	–0.987	–2.399	LOC_Os06g02560	Growth-regulating factor	Y ^{2,3,4}
		0.129	–0.870	–2.030	LOC_Os03g47140	Growth regulating factor protein	Y ^{3,4}
		–0.325	–0.795	–1.843	LOC_Os02g53690	Growth regulating factor protein	Y ^{3,4}
		0.241	–0.744	–1.671	LOC_Os04g51190	Growth-regulating factor	Y ^{3,4}
miR397a	–0.076	0.413	–0.812	–1.833	LOC_Os11g48060	Laccase-22 precursor	Y ³
miR397b	–0.108	0.413	–0.828	–1.875	LOC_Os11g48060	Laccase-22 precursor	Y ³
		1.263	0.731	1.745	LOC_Os09g27950	Galactosyltransferase	N
miR444b–c	–0.466	0.006	0.678	1.797	LOC_Os02g49840	Scarecrow	Y ^{2,3}
miR444f	1.947	–0.646	–0.941	–1.789	LOC_Os02g34080	Targeting protein for Xklp2	N
miR528	–0.581	0.136	–0.897	–1.583	LOC_Os08g44770	Copper/zinc superoxide dismutase	N
miR529b	–1.633	0.219	–0.876	–1.923	LOC_Os04g28420	Peptidyl-prolyl isomerase	N
		–0.038	–0.687	–1.529	LOC_Os12g08760	Carboxyvinyl-carboxyphosphonate phosphorylmutase	N
miR530	–0.900	–0.678	0.794	2.288	LOC_Os05g09650	Ubiquinone biosynthesis protein COQ4	N
miR1847	–0.376	–0.150	0.912	1.924	LOC_Os01g63190	Laccase precursor protein	N
miR1856	–6.129	–1.509	0.827	1.785	LOC_Os09g10274	Expressed protein	N
miR1860	0.174	–0.069	0.920	2.125	LOC_Os01g01030	Monocopper oxidase	N
miR1861a	1.984	0.021	0.796	1.590	LOC_Os03g40020	PPR repeat containing protein	N
miR1864	–1.514	0.414	–0.863	–1.374	LOC_Os01g14020	Expressed protein	N
miR1873	0.003	0.223	–0.717	–2.359	LOC_Os05g01790	Expressed protein	N
miR1884a	0.575	–1.074	–0.685	–1.786	LOC_Os11g34910	Expressed protein	Y ²

(continued)

Table 2 Continued

miRNA	miRNA FC ^a	mRNA FC ^b	Correlation ^c	W	Target	TIGR Annotation ^d	Target Verification ^e
miR1884b	0.083	0.435	0.952	1.782	LOC_Os06g14780	Expressed protein	N
		−0.039	−0.677	−2.294	LOC_Os12g01680	Macrophage migration inhibitory factor	N
		0.036	−0.573	−1.865	LOC_Os02g49870	Expressed protein	N
		−0.231	0.575	1.640	LOC_Os01g64520	Uricase	Y ²
		0.435	0.636	1.786	LOC_Os06g14780	Expressed protein	N
miR2097	0.543	0.787	0.779	1.585	LOC_Os08g43920	Carrier	N

^aFold change in the log₂ ratio of miRNA expression between *indica* and *japonica* subspecies in seedlings. The mean expression levels averaged from three accessions were used for the calculation. MiRNAs with significant differential expressions (unadjusted *P*-value ≤ 0.05 and Fold change ≥ 2) between rice subspecies are marked in bold.

^bFold change in the log₂ ratio of mRNA expression between *indica* and *japonica* subspecies in seedlings. The mean expression levels averaged from three accessions were used for the calculation. Genes with significantly differential expression are marked in bold.

^cPearson's correlation coefficient.

^dGene annotations from TIGR (Version 6).

^eThe Y^{2,3,4} Target is verified in the corresponding degradome target set II, III, or IV, respectively; N: not yet verified.

miRNA subfunctionalization between closely related plant species (de Vries et al. 2015).

Another pattern that we observed is the prevalent positive correlation between individual miRNA-target pairs in the cultivated rice, of which many target genes have been experimentally verified by degradome data. Given miRNA negatively regulated target expression, this observation may reflect the composite effect of miRNA-mediated circuits, such as negative feedback loops (FBLs) and incoherent feedforward loops (FFLs) (Alon 2007). miRNA-mediated FBLs and FFLs are recurrent network motifs in animals (Tsang et al. 2007; Wu et al. 2009), and also play significant roles in plants as evidenced by both experiments (Xie et al. 2003; Bari et al. 2006; Vaucheret et al. 2006) and computational analyses (Meng et al. 2009; Wu et al. 2009). From the network perspective, miRNAs can thus play a role in expression buffering and biological robustness, as well recognized in animal systems (Wu et al. 2009; Herranz and Cohen 2010; Ebert and Sharp 2012; Pelaez and Carthew 2012). We found that targets of rice miRNAs overall have less expression variation in comparison with the transcriptome, albeit miRNAs themselves are more variable in expression. These results are consistent with the notion that miRNAs act as dampers in buffering target expression variation (Fei et al. 2013) or target sequence diversity (de Vries et al. 2015). In contrast, miRNA targets compared with non-targets overall have higher level of expression variation in human, despite a small number of highly expressed targets with decreased expression variation, suggesting the dual function of miRNA regulation (Lu and Clark 2012).

Functional enrichment tests of the DEGs highlight that expression changes of genes involved in various metabolism processes and stress responses are important for the *indica-japonica* differentiation. Genes involved in plant-pathogen pathways are the most prominent examples, since the interaction between rice-host cultivar (genotype) and parasite population is shown to be critical in determining parasite affecting (Huang et al. 2012). Interestingly, we identified a

target (LOC_Os02g30900, *PBS1*) of a lowly conserved miRNA, miR1857, which is involved in this pathway (supplementary fig. S7, Supplementary Material online). The orthologous *PBS1* gene in *Arabidopsis* encodes a putative serine-threonine kinase, which is required for specific recognition of the bacterial protein *AvrPphB* (Swiderski and Innes 2001). Another related example has also been recently reported in *Arabidopsis*, where a peptide derived from the plant pathogen *Pseudomonas syringae* induces the expression of the host miR393 that targets auxin receptor (*TIR1*, *AFB2*, and *AFB3*), and consequently inhibits the growth of the bacteria (Navarro et al. 2006). It would be interesting to explore the role of miRNAs in mediating biotic stress response in the context of differentiation between *indica* and *japonica* subspecies.

Our permutation analysis provides a promising approach to further classify the important miRNA regulation pairs in rice. For example, besides the well documented miRNA-target pairs mentioned previously, recent study revealed that osa-miR171c controls the floral transition and maintenance of shoot apical meristem (SAM) indeterminacy in rice by targeting *GRAS* (*GAI-RGA-SCR*), a plant-specific transcription factors (LOC_Os02g44370, also for *OsHAM2*, table 2) (Fan et al. 2015); In addition, miR166-mediated post-transcriptional gene silencing of rice Class III HD-Zip genes (LOC_Os03g43930, also for *OsHB5*) is reported to be responsible for the auxin signals to regulate leaf and shoot development (Itoh et al. 2008; Toriba et al. 2010). There are also many miRNAs with un-verified targets in our list of correlated miRNA-target pairs (table 2). Further studies are necessary to confirm the functional contribution of these miRNAs to the process of rice development and/or differentiation. The joint miRNA-mRNA data and permutation analyses used in this study provide a novel idea to study miRNA regulatory relationships in rice. Our results may provide a valuable resource for further investigation of miRNA functions in rice developmental regulation, stress responses, and biomass yields under domestication.

Methods

Plant Materials

Seeds of *O. sativa* L. ssp. *indica* and ssp. *japonica* accessions as listed in table 3 were obtained from the International Rice Research Institute (IRRI, Manila, Philippines). For grains, the husks of the seeds were removed before RNA extraction. For seedlings, rice seeds were sterilized and germinated in Petri dishes containing distilled water at 37 °C under dark conditions for 2 days. The uniformly germinated seeds were transferred into Yoshida nutrient solution and grown under a 16-h light (28 °C)/8-h dark (25 °C) photoperiod for one week. The samples were collected and rinsed with double distilled water three times, and then immediately frozen in liquid nitrogen until use.

RNA Isolation and Preparation of Sequencing Libraries

Total RNA were extracted from rice grains and seedlings using TRIzol Reagent (Invitrogen), and evaluated using an Agilent 21100 Bioanalyzer (Agilent Technologies). Small RNA and transcriptome libraries were prepared using standard protocols of the Illumina Small RNA Sample Prep Kit or the Illumina TruSeq RNA Sample Prep kit, and sequenced using an Illumina Genome Analyzer (Illumina, San Diego, CA, USA) at BGI (Shenzhen, China). As we did not obtain enough quality RNA in grain, only the seedling samples were used for RNA-seq.

Expression Analysis of miRNAs

For all small RNA libraries, after trimming prime adaptors and filtering low quality or adaptor contaminated reads, clean reads within the length range of 19–30 nt were retained for further analysis. These reads were searched against the Rfam database (Griffiths-Jones et al. 2003) and the RepBase database (Jurka et al. 2005) using the SOAP software (Li et al. 2008) with two mismatches, in order to remove reads matching structural RNAs, including rRNA, tRNA, small nuclear RNA (snRNA), and small nucleolar RNA (snoRNA), and repeats/transposons. The remaining reads were mapped to the precursors of all known rice miRNAs registered in miRBase (Release 20, <http://www.mirbase.org/index.shtml>; last accessed 3 September 2015) using SOAP (Li et al. 2008). We grouped all the variants of rice mature miRNAs into 553 distinct mature miRNAs since some miRNA precursors produced more than one different mature sequences and some identical mature sequences were generated from distinct precursors. Read counts corresponding to individual mature miRNA sequences were normalized and rescaled as Reads Per Million (RPM), which divided the raw read count of each mature miRNA by the total mapped read count in each library and multiplied it by 1 million. Subsequent heatmap clustering and differential expression analysis were performed based on the average RPM of each distinct mature miRNA.

Table 3
Summary of the Reads Mapping of Small RNAs and Transcriptomes by RNA Sequencing

Tissue	Subspecies	Accession Name	Accession No. ^a	Rice variety			Small RNAs			Transcriptome				
				No. of Sequences Generated	No. of Sequences Matching the Rice Genome	No. of Unique Sequences	No. of Sequences Generated ^b	No. of Sequences Matching the Rice Genome ^c	Unique Mapped Reads ^c					
Seedling	<i>Indica</i>	Khao Dawk Mali 105	IRGC 27748	9,985,373	9,758,028 (97.7%)	2,786,744 (27.9%)	20,326,850	34,063,783 (83.79%)	33,569,936 (82.6%)	NA	NA			
		Guangluai 4	IRGC 114900	9,788,880	9,558,792 (97.7%)	3,048,520 (31.1%)	20,216,686	32,972,884 (81.54%)	32,494,623 (80.4%)					
	<i>Japonica</i>	Rathuwee	IRGC 8952	9,997,943	9,701,803 (97.0%)	3,403,862 (34.0%)	19,876,912	32,789,681 (82.48%)	32,328,952 (81.3%)					
		Taipei 309	IRGC 42576	8,221,032	8,051,214 (97.9%)	2,282,686 (27.8%)	19,094,551	33,286,973 (87.16%)	32,855,807 (86.0%)					
	Grain	<i>Indica</i>	Heukgyeong	IRGC 55530	9,371,607	8,988,554 (95.9%)	2,203,943 (23.5%)	19,889,198	34,863,682 (87.64%)			34,372,869 (86.4%)	NA	NA
			Nipponbare	IRGC 12731	10,592,461	10,416,128 (98.3%)	3,010,461 (28.4%)	18,818,192	32,995,013 (87.67%)			32,592,262 (86.6%)		
Total	<i>Indica</i>	Khao Dawk Mali 105	IRGC 27748	14,396,308	13,994,396 (97.2%)	5,408,929 (37.6%)	NA	NA	NA	NA	NA			
		Guangluai 4	IRGC 114900	14,433,295	14,137,205 (98.0%)	4,056,622 (28.1%)	NA	NA	NA					
	<i>Japonica</i>	Rathuwee	IRGC 8952	15,718,282	15,185,127 (96.6%)	5,740,388 (36.5%)	NA	NA	NA	NA	NA			
		Taipei 309	IRGC 42576	14,846,530	14,293,090 (96.3%)	2,786,744 (18.8%)	NA	NA	NA					
	Total	<i>Japonica</i>	Heukgyeong	IRGC 55530	10,225,740	9,512,313 (93.0%)	3,081,796 (30.1%)	NA	NA	NA	NA	NA		
			Nipponbare	IRGC 12731	13,862,947	13,356,435 (96.4%)	5,014,395 (36.2%)	NA	NA	NA				
Total				141,440,398	136,953,085 (96.8%)	42,825,090 (30.3%)	NA	NA	NA	NA	NA			

^aInternational Rice Germplasm Collection at IRRI in the Philippines (<http://archive.irri.org/GR/requests/requests.htm>).

^bNumber of pairs of 75-nt paired-ends sequencing reads.

^cThe proportion was calculated as the number of mapped reads versus the number of total reads. NA, Not applicable.

An unsupervised, two-dimensional, hierarchical clustering was conducted using R package *gplots* (Warnes et al. 2009). For differential expression analysis, a generalized Poisson-regression linear model was fitted in the R package *edgeR* (Nikolayeva and Robinson 2014) to identify the differentially expressed miRNAs for the factors of development, subspecies, and development-by-subspecies. MiRNAs with fold change ≥ 2 and $FDR \leq 0.05$ are denoted as significantly mi-expressed.

Expression Analysis of mRNAs

For the transcriptome libraries, after removing adaptors and low-quality reads from raw reads, clean reads were mapped to the reference genome of rice (*japonica* cv. Nipponbare) using the Bowtie software package (Langmead et al. 2009). Reads that could be mapped equally well to multiple locations without mismatch or with up to two mismatches were randomly assigned to one position and were retained for further analyses as previously described (Wang et al. 2009). According to the TIGR 6.0 gff3 file (Kawahara et al. 2013), reads matching gene or genomic region were recovered and expression level of each transcript was measured as numbers of reads per kilobase of exon region in a gene per million mapped reads (RPKM) as previously described (Mortazavi et al. 2008).

To identify homologous genes between rice subspecies, the longest isoform of each gene model of *japonica* cv. Nipponbare was aligned to the genome of *indica* cv. 93-11 by GMAP (Wu and Watanabe 2005). We considered a Nipponbare gene and its counterpart in the 93-11 genome as an homologous gene pair if they share at least 95% sequence identity in at least 95% of their longest isoforms (He et al. 2010). The resulting 32,044 homologous gene pairs were further filtered by requiring at least 10 raw reads in all the three accessions sequenced for at least one developmental stage of a subspecies. These expressed homologous genes were then used for the subsequent differential expression analysis. The generalized Poisson-regression linear model was fitted and the likelihood ratio test was performed with *lmtest* (Zeileis and Hothorn 2002) to identify the mRNAs with differential expression between rice subspecies. Genes with fold-change ≥ 2 and $FDR \leq 5\%$ were denoted as significantly differentially expressed.

Gene Functional Annotation

The gene ontology classification developed in the TIGR Rice Genome Annotation Project (<http://rice.plantbiology.msu.edu/>; last accessed 10 September 2015) was used to assign genes to a hierarchical biological process using the criteria of the Gene Ontology (GO) Consortium databases (Ashburner et al. 2000; Kawahara et al. 2013), and KEGG pathway annotation was identified according to KEGG database (<http://www.genome.jp/kegg/>; last accessed 10 September 2015) (Kanehisa and Goto 2000; Kanehisa et al. 2016). The GO and KEGG

enrichment test ($FDR \leq 0.05$) was performed by using the CORNA and *clusterProfiler* package, respectively (Wu and Watson 2009; Yu et al. 2012). KEGG pathway rendering of the expressed genes was conducted by the R package *Pathview* (Luo and Brouwer 2013). The DEGs with a fold change larger than 2 were used for the GO enrichment test and 1.5 for the KEGG pathway enrichment test, with a $FDR \leq 0.05$ was used as the significance threshold.

Identification of Novel miRNAs

After removing the reads mapped to the structural RNAs, the known miRNAs from miRBase and repeats/transposons, the remaining sequences were mapped on the rice genome and analyzed by an adjusted miRDeep script for novel plant miRNA prediction (Wen et al. 2012). The following criteria for miRNA annotation (Wang et al. 2009) were further applied to filter the novel miRNAs, including: (1) a secondary structure must have a hairpin with at least 18 paired nucleotides in its stem region; (2) the hairpin must have free energy less than or equal to -18 kcal/mol and no more than two central loops; (3) the miRNA and miRNA* form a duplex with a two-nucleotide overhang; (4) fewer than four mismatches exist in the miRNA/miRNA* duplex; and (5) $\geq 75\%$ of the sRNAs mapped onto a miRNA precursor are derived from the miRNA or miRNA* region. The candidate sequences were extracted and folded with Vienna RNAfold (Lorenz et al. 2011). The miRNAs with more than 10 raw reads matching the mature sequences in at least two replicate sRNA libraries were considered to be expressed.

MiRNA Target Prediction and Integration

A search for the miRNA target genes was performed for all known miRNAs and newly identified miRNA sequences on the *japonica* cv. Nipponbare cDNA dataset (TIGR 6.0) using *psRNAtarget* (Dai and Zhao 2011) with a maximum expectation score of 2.5 to reduce the false discovery rate (Klevebring et al. 2009). The predicted target set was referred to as target set I in this study. Three additional sets of miRNA targets that are verified by degradome analysis were also used, which include target set II (Wu et al. 2009) and target set III (Li et al. 2010) for rice *japonica* cv. Nipponbare, and target set IV for rice *indica* cv. 93-11 (Zhou et al. 2010).

miRNA Target Permutation Analysis

The permutation test and weighted shift for the miRNA-target correlations were processed as described previously (Nunez-Iglesias et al. 2010). In short, the miRNA-target pairs can be considered a bipartite graph, with nodes representing the miRNAs on one side and the targets on the other, and the edges represent the target prediction relationships. The nodes have associated expression measurements. Correlation statistics were then computed for each miRNA-target pair across the 6 rice lineages. Finally, the network was permuted by

shuffling the edges, re-computing the statistics, and then repeating this process 1000 times. From these, we can obtain an empirical P -value and a “weighted correlation shift”, W , which was defined as the difference between the true value and the mean permuted value divided by the standard deviation of the permuted values: $W = (r - r_0)/S_0$.

Quantitative miRNA and mRNA Analysis by qRT-PCR

For miRNA quantification, total RNA from 1 week old seedlings was subjected to stem-loop reverse transcription (RT) (Chen et al. 2005) followed by Taqman PCR (Applied Biosystems) using the miRNA UPL (Roche Diagnostics) probe assay as described previously (Varkonyi-Gasic et al. 2007; He et al. 2016). 5.8S rRNA was used as an internal control. Three biological replicates were examined. For mRNA quantification, the same total RNA was first treated with TURBO DNA-free kit (Ambion) to remove potential genomic DNA contamination and then used for reverse transcription with SuperScript III First-Strand Synthesis System (Invitrogen). qRT-PCR was performed with Platinum SYBR Green qPCR SuperMix (Invitrogen) according to the manufacturer’s instructions. *Actin* was used as an internal control. Three biological replicates with two technique repeats each were examined to ensure reproducibility. The relative levels of miRNA or mRNA were calculated using the $2^{-\Delta\Delta CT}$ method (Livak and Schmittgen 2001). The sequences of primers used are listed in [supplementary table S5, Supplementary Material](#) online.

Data Availability

The expression data generated by this study are available in the NCBI Gene Expression Omnibus (GEO) under accession GSE71925.

Supplementary Material

Supplementary text, tables S1–S5, and figures S1–S11 materials are available at *Genome Biology and Evolution* online (<http://www.gbe.oxfordjournals.org>).

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