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Strand-specific RNA-seq reveals widespread occurrence of novel *cis*-natural antisense transcripts in rice

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Abstract

Background: *Cis*-natural antisense transcripts (*cis*-NATs) are RNAs transcribed from the antisense strand of a gene locus, and are complementary to the RNA transcribed from the sense strand. Common techniques including microarray approach and analysis of transcriptome databases are the major ways to globally identify *cis*-NATs in various eukaryotic organisms. Genome-wide *in silico* analysis has identified a large number of *cis*-NATs that may generate endogenous short interfering RNAs (nat-siRNAs), which participate in important biogenesis mechanisms for transcriptional and post-transcriptional regulation in rice. However, the transcriptomes are yet to be deeply sequenced to comprehensively investigate *cis*-NATs.

Results: We applied high-throughput strand-specific complementary DNA sequencing technology (ssRNA-seq) to deeply sequence mRNA for assessing sense and antisense transcripts that were derived under salt, drought and cold stresses, and normal conditions, in the model plant rice (*Oryza sativa*). Combined with RAP-DB genome annotation (the Rice Annotation Project Database build-5 data set), 76,013 transcripts corresponding to 45,844 unique gene loci were assembled, in which 4873 gene loci were newly identified. Of 3819 putative rice *cis*-NATs, 2292 were detected as expressed and giving rise to small RNAs from their overlapping regions through integrated analysis of ssRNA-seq data and small RNA data. Among them, 503 *cis*-NATs seemed to be associated with specific conditions. The deep sequence data from isolated epidermal cells of rice seedlings further showed that 54.0% of *cis*-NATs were expressed simultaneously in a population of homogenous cells. Nearly 9.7% of rice transcripts were involved in one-to-one or many-to-many *cis*-NATs formation. Furthermore, only 17.4-34.7% of 223 many-to-many *cis*-NAT groups were all expressed and generated nat-siRNAs, indicating that only some *cis*-NAT groups may be involved in complex regulatory networks.

Conclusions: Our study profiles an abundance of *cis*-NATs and nat-siRNAs in rice. These data are valuable for gaining insight into the complex function of the rice transcriptome.

Keywords: *Oryza sativa*, *Cis*-NATs, Nat-siRNAs, SsRNA-seq, Transcriptome

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Background

Cis-natural antisense transcripts (*cis*-NATs) are endogenous RNA molecules that are transcribed from the opposite DNA strand of the same genomic locus and overlap partly with sense RNA by convergent or divergent orientation. It has been clearly demonstrated that *cis*-NATs are an important biogenesis mechanism to generate endogenous short interfering RNAs (siRNAs) known as 'natural antisense siRNAs' (nat-siRNAs) [1,2]. Numerous evidence indicates that NATs use diverse transcriptional and post-transcriptional gene regulatory mechanisms to carry out different biological roles [3], including RNA interference [4], gene silencing [5-7], RNA masking-induced alternative splicing [8] and RNA editing [9].

Since a large number of natural antisense transcripts were first identified to be widespread in the human genome [10], computational analysis of data generated from large-scale sequencing projects has been widely used to globally identify *cis*-NATs in various eukaryotic organisms. Common techniques and databases, including *in silico* analyses of expressed sequence tag (EST) databases [11,12], genomic annotation of large transcript sets [13-17], large-scale sequencing of full-length complementary DNAs (cDNAs) [18-20] and tiling arrays [21-24], have been applied to identify NATs. Recently, more efforts were made to characterize nat-siRNAs and NATs at a genome-wide level: massively parallel signature sequencing (MPSS) data [25,26], combining pyrophosphate-based high-throughput sequencing and computational analyses of genomic annotation datasets [27,28] and asymmetric strand-specific analysis of gene expression (ASSAGE) [29]. It has been reported that about 5-70% in mammals and 7-9% in plants of all transcripts are overlapped as *cis*-NATs. These studies have demonstrated four major characteristics of *cis*-NATs. (i) Both *cis*-NAT pairs can encode proteins or be non-protein-coding transcripts. In the mammalian genome, a non-protein-coding antisense RNA partner of a protein-coding transcript is considered the most prominent form [13,15]. In *Arabidopsis*, ~88% of the sense-antisense transcripts (6858 of 7805) were shown to be pairs of protein-coding genes (AGI code genes) and non-protein-coding RNAs from non-AGI transcriptional units [21,24]; however, it has been reported that > 86% of rice bidirectional transcript pairs included a coding sequence in both strands [19]. (ii) The distribution of antisense transcripts in mammals was found to be non-random across the genome [29]. (iii) Expression levels of sense and antisense transcripts can be either positively or negatively correlated [15,21,30,31]. (iv) In *Arabidopsis*, most of the *cis*-NATs are arranged in convergent orientation. Several genes were found to be involved in two *cis*-NATs as a network: one pair is convergent, another is divergent [20,27].

However, there are still some limitations to comprehensively identifying *cis*-NATs. Firstly, the percentage of *cis*-NATs in different eukaryotic genomes, especially in plants, is estimated mainly by the alignment of full-length cDNAs, ESTs and predicted coding sequences to the genome. The transcriptomes are still not sequenced deeply enough to provide all transcripts, including low copy number and non-coding RNAs (ncRNAs). Secondly, as important evidence to determine the regulation of *cis*-NATs by RNA interference, nat-siRNAs are still far from saturated, despite many efforts in plants (rice and *Arabidopsis*) in recent years [26-28,32-35]. Thirdly, it is also essential to accurately quantify the expression levels of sense and antisense transcripts at a global level.

The advent of second-generation sequencing technology enables deep sequencing of transcripts. The paired-end tag sequencing strategy of strand-specific cDNA sequencing technology (ssRNA-seq) has the potential to globally produce abundant and novel transcripts with clear polarity and to accurately assess gene activity [36,37]. Moreover, each cell type has its unique transcriptome, so a single-cell-level description of gene expression and regulation can be instructive concerning cell populations [38,39]. The latest RNA-seq applied to single cells gives more precise transcriptome quantifications than a PCR-based amplification method [40]. Meanwhile, high-throughput sequencing techniques have made it feasible to obtain all small RNAs species, genome-wide, as it can generate hundreds of millions reads in a single sequencing run.

Here, we took advantage of ssRNA-seq technology to deeply sequence cDNAs with clear transcriptional orientations in the model plant species rice (*Oryza sativa* L.). All mRNAs were derived from seedlings grown under normal (seedling mixture, and only epidermal cells as well) and abiotic stressed conditions for assessing rice *cis*-NATs at the best possible resolution. We also deeply sequenced small RNAs to investigate nat-siRNAs from rice seedlings under normal and several stresses conditions. In addition, we tried to sequence transcriptome of rice leaf epidermal cells and evaluate the expression of *cis*-NATs identified in this research. We identified 2292 rice *cis*-NATs with both evidence of gene expression and nat-siRNAs from their overlapping regions. About 54.0% of them were shown to be simultaneously expressed in epidermal cells. Some *cis*-NATs gave rise to nat-siRNAs exclusively in the overlap regions, and some *cis*-NATs seemed to be expressed under specific abiotic stress conditions. This study was the first attempt of applying ssRNA-seq to deeply investigate novel transcripts and revealed widespread occurrence of *cis*-NATs in rice.

Results

ssRNA-seq and assembly of rice transcripts

In order to comprehensively identify rice *cis*-NATs, we applied ssRNA-seq to deeply sequence rice cDNAs for assessing transcripts with clear transcriptional orientations. Rice seedlings grown under normal and three abiotic stress conditions (salt, cold and drought treatments) were collected for preparations of mRNA and small RNAs (see Methods), and four strand-specific cDNA libraries were then constructed according to the empirical protocol [37]. In general, a modified RNA-seq method is used for incorporation of deoxy-UTP during second-strand cDNA synthesis and subsequent destruction of the uridine-containing strand in the sequencing library. This enables the identification of transcript orientation. The high-throughput sequencing of strand-specific cDNAs was performed on the Illumina GAIIX. In total, 14.7, 11.5, 14.2 and 13.2 million paired-end reads of 2×120 bp with high-quality, which were generated from the untreated and salt, cold and drought stress treatments libraries, respectively, matched unambiguously and uniquely to the rice reference genome [41] (Additional file 1). We estimated the accuracy of transcriptional orientation by comparing the sequencing reads with the annotated gene datasets [42]. About 89.4-95.5% of the mapped reads appeared to be aligned with correct transcriptional orientation, providing strong evidence for ssRNA-seq data in a strand-specific manner. In contrast to ssRNA-seq data, only half of the mapped reads from the previous RNA-seq data were consistent with the gene models at the same transcriptional orientations. In addition, we also deeply sequenced epidermal cells of normal rice seedlings, and about 10.4 million paired-end reads of 2×100 bp were generated as high-quality data.

Using the software TopHat and Cufflinks [43-46], we assembled ssRNA-seq data combined with the rice genome annotation [42] into 76,013 transcripts corresponding to 45,844 unique gene loci (including 4873 novel gene loci). Of them, 25,924 were identified as novel transcripts, which were composed of 5063 ncRNA, 16,494 CDS with protein hits and 4367 CDS without any protein hits (Additional file 2).

Identification of putative *cis*-NATs in rice

Based on renewed assembled transcripts, we identified putative *cis*-NATs which overlapped, but were opposite, from the same or adjacent gene locus. In total, 5813 pairs of rice *cis*-NATs (Additional file 3) were preliminarily screened out. After excluding those in which either of the pairs encoded a transposon, rRNA, tRNA, snRNA, snoRNA or miRNA, we obtained 3819 putative *cis*-NATs with mean overlapped length of 785 nt (Table 1). According to the directions of the involved transcripts, 2149 (56.3%) *cis*-NATs were categorized in enclosed, 898

(23.5%) in convergent (3'-3' overlap), and 772 (20.2%) in divergent (5'-5' overlap) orientations. Of rice *cis*-NATs, 36.1% (1378 of 3819) were pairs of protein-coding genes and non-protein-coding RNAs (Table 1). Another 33.4% (1275 of 3819) were a PFAM domain-containing transcript partner of a predicted CDS without any PFAM domain. In general, the majority of *cis*-NATs (3358 or 87.9%) were one-to-one type, i.e. one transcript in a *cis*-NAT pair had only one antisense partner. The remaining 461 *cis*-NATs (composed of 685 transcripts) were involved in networks of 223 *cis*-NAT groups.

Small RNAs and nat-siRNAs in rice

To investigate the complexity of small RNAs in rice, we generated four small RNA libraries from rice seedlings under normal conditions and three abiotic stress treatments (salt, cold and drought), and deeply sequenced the libraries on the Illumina GAIIX. The small RNA libraries were made with an RNA ligation method which produces strand-specific libraries. After removing low-quality reads and those mapped to rRNA, tRNA, sn/snoRNA, mitochondria and chloroplasts, 4,843,040, 3,973,627, 2,894,255 and 5,492,145 distinct small RNAs (representing 17,632,759, 12,923,509, 8,720,251 and 20,069,157 of 48,683,191, 49,254,272, 25,705,840 and 50,192,805 raw reads, respectively) from corresponding untreated, salt, cold and drought conditions, were identified (Additional file 1). Small RNAs of 24-nt were the predominant size class (Additional file 4A). We found that the majority were located in transposon-related regions, followed by upstream, intergenic, downstream, and intron regions (Additional file 4C).

To determine the amount of nat-siRNAs, we searched in the four small RNA libraries that matched uniquely and perfectly to the overlapping regions of the 3819 *cis*-NATs. A total of 180,239 reads corresponding to 90,977 unique small RNAs (normal: 25,420; salt: 18,598; cold: 18,152 and drought: 28,807) were derived from overlapped regions (Additional files 1 and 5). Here, nat-siRNAs showed wide ranges in size of 18-34 nt, with 21-25 nt the most common size (Additional file 4B). The 5'-first nucleotide of nat-siRNAs was predominantly adenosine (Additional file 4F), which differed from that of the total small RNAs (Additional file 4E).

cis-NATs and nat-siRNAs with gene expression evidence under normal or abiotic stress conditions

To gain further insight into regulation of *cis*-NATs, we examined the expression levels of 3358 one-to-one type *cis*-NATs. The 2292 *cis*-NAT pairs were detected as expressed [i.e. Fragments Per Kilobase of exon Models (FPKM) > 0] with both sense and antisense transcripts under normal (1789 pairs), cold (1668), salt (1572) or drought (1668) conditions, and also with nat-siRNAs in an overlapping region (Table 2 and Additional file 6).

Table 1 Statistics of 3819 *cis*-NATs identified in rice

Chr.	Transcripts	<i>cis</i> -NATs ^a	Enclosed ^b	3'-3' ^c	5'-5' ^d	CDS-p ^e vs. ncRNA	CDS-p vs. CDS-n ^f	CDS-p vs. CDS-p	CDS-n vs. ncRNA	CDS-n vs. CDS-n	ncRNA vs. ncRNA
1	10,536	558	301	146	11	205	176	92	39	30	16
2	8,527	440	239	115	86	159	141	63	36	30	11
3	9,197	491	296	120	75	175	156	90	29	22	19
4	6,762	361	185	95	81	100	132	55	27	27	20
5	5,940	290	161	71	58	107	108	34	21	11	9
6	6,166	299	170	69	60	111	100	38	23	13	14
7	5,862	278	163	63	52	90	96	30	24	19	19
8	5,290	239	138	51	50	103	74	27	19	10	6
9	4,256	189	100	34	55	70	71	10	16	10	12
10	4,144	221	128	41	52	85	74	24	19	10	9
11	4,861	231	139	47	45	92	75	25	23	7	9
12	4,472	222	129	46	47	81	72	24	20	15	10
Total	76,013	3,819	2,149	898	772	1,378	1,275	512	296	204	154

^a *cis*-NAT pairs without transposons.

^b One transcript being entirely reverse-complementarily overlapped by the other.

^c Convergent *cis*-NAT (with 3'-ends overlapped).

^d Divergent *cis*-NAT (with 5'-ends overlapped).

^e Coding sequence with PFAM domain-containing.

^f Predicted as coding sequence but without any PFAM domain hits.

We found that 166 pairs produced siRNAs exclusively and had more than five unique small RNA reads in the overlapping regions (Additional file 7). To explore whether small RNAs were more enriched in the overlapping compared to non-overlapping regions, we calculated small RNA densities of *cis*-NATs in these two regions. The siRNA densities of 13 *cis*-NATs were > 5 times that in the non-overlapping regions (Additional file 8). We further investigated *cis*-NATs which generated nat-siRNAs with strand bias: 25-28% of *cis*-NAT pairs exhibited strong strand bias in spawning nat-siRNAs with > 5-fold change in both normal and abiotic stresses (Additional file 9A). More than 75% of *cis*-NATs generated small RNAs from different directions with > 2-fold change (Additional file 9B). Interestingly, *cis*-NAT pairs, which were composed of the protein-coding gene partner of non-protein-coding RNAs, overwhelmingly produced small RNAs from the strand of protein-coding transcripts (Additional file 9).

Of 2292 *cis*-NATs, 1072 (46.7%) *cis*-NATs were expressed in both normal and abiotic stresses (Figure 1A). The *cis*-NAT pairs could be divided into five subgroups based on a scatter plot comparing transcripts expressional ratio trends of *cis*-NAT pairs between normal and cold stressed conditions (Figure 1B). We also investigated the functional bias among each group of *cis*-NAT pairs through functional domain and expressional profiling analyses. We found that Protein kinase domains were commonly identified in *cis*-NAT pairs among the five subgroups. However, other functional domains and expression levels of sense/antisense

transcripts were detected to be associated with different subgroups of *cis*-NAT pairs.

Subgroup-1 *cis*-NATs were predominantly associated with the pentatricopeptide repeat (PPR) and Protein tyrosine kinase domains. We also found that the expression levels of the *cis*-NAT pairs of Subgroup-1 under stress conditions were similar to that under normal conditions. Subgroup-2 *cis*-NATs were identified to contain domains of leucine rich repeats and glycosyltransferase. In this subgroup, the expression levels of sense-transcripts of the *cis*-NAT pairs were significantly increased under stress conditions, while expression levels of antisense transcripts were greatly reduced. Thus, expression levels were higher for sense than for antisense transcripts under stress conditions. Subgroup-3 can be classified as the family of alpha/beta hydrolase fold. The expression levels of sense-transcripts of the Subgroup-3 *cis*-NAT pairs were significantly reduced under stress conditions, while expression levels of antisense transcripts significantly increased. Thus, expression levels were lower for sense than for antisense transcripts under stress conditions. Subgroup-4 *cis*-NATs can be classified as the families of the eukaryotic aspartyl protease and sugar (and other) transporter. Subgroup-5 *cis*-NATs were identified to be the families of glycosyl hydrolases and ubiquitin-conjugating enzyme. The remaining four *cis*-NAT pairs were not classified into any subgroups, as the expression levels of sense transcripts were not correlated with that of anti-sense partner under normal and abiotic conditions. We identified these four *cis*-NAT pairs as abnormal values (Figure 1B;

Table 2 Numbers of *cis*-NATs with expression evidence and nat-siRNAs, under four different conditions and in epidermal cells

	Both expressed <i>cis</i> bsgNAT pairs	Validated in epidermal cells	Pairs with siRNAs only in overlap region ^f	Pairs with siRNAs enriched in overlap region ^g
sd ^a	1,789	1,043	72	9
ST ^b	1,572	949	58	5
CD ^c	1,668	1,003	66	5
DT ^d	1,668	986	75	5
Co-	1,072 (46.7%) ^e	725 (58.6%) ^e	10 (5.9%) ^e	2 (15.4%) ^e
Total	2,292	1,238	166	13

^a 14-d-old-seedling.

^b 14-d-old-seedling treated with 200 mM NaCl.

^c 14-d-old-seedling grown under cold stress at 4°C for 24 h in darkness.

^d 14-d-old-seedling treated with 20% PEG-6000.

^e The percentage of co-*cis*-NATs in the corresponding total *cis*-NATs was calculated.

^f Small RNAs were only enriched in the overlapping regions with more than five reads.

^g The density of siRNAs in overlapping regions was > 5 times that in non-overlapping regions.

indicated as the brown spots). Subgroup-1 was shown to be represented the major part. Ratio values of 913 *cis*-NAT pairs (Subgroup-1) between normal and cold stressed conditions were highly correlated, with $R^2 > 0.85$. Similar results were also observed between normal and salt/drought stressed conditions with $R^2 > 0.93$ of 917 pairs and $R^2 > 0.93$ of 898 pairs, respectively (Additional file 10). In total, of 1072 *cis*-NAT pairs, 95.5% (1024) showed positive correlations between sense and antisense transcripts. One *cis*-NAT pair (Os05t0500000-00 vs. Os05t0500101-01) was taken as an example to compare expression changes under normal and cold stressed conditions (Figure 2). Northern blot analyses also demonstrated their differential expression levels under abiotic stresses (Figure 2C). Sense transcript Os05t0500000-00 is annotated as an UDP-glucuronosyl and an UDP-glucosyl transferase, which is an important enzyme for catalyzing transportation of sugars. Antisense transcript Os05t0500101-01 had no hits to any protein families.

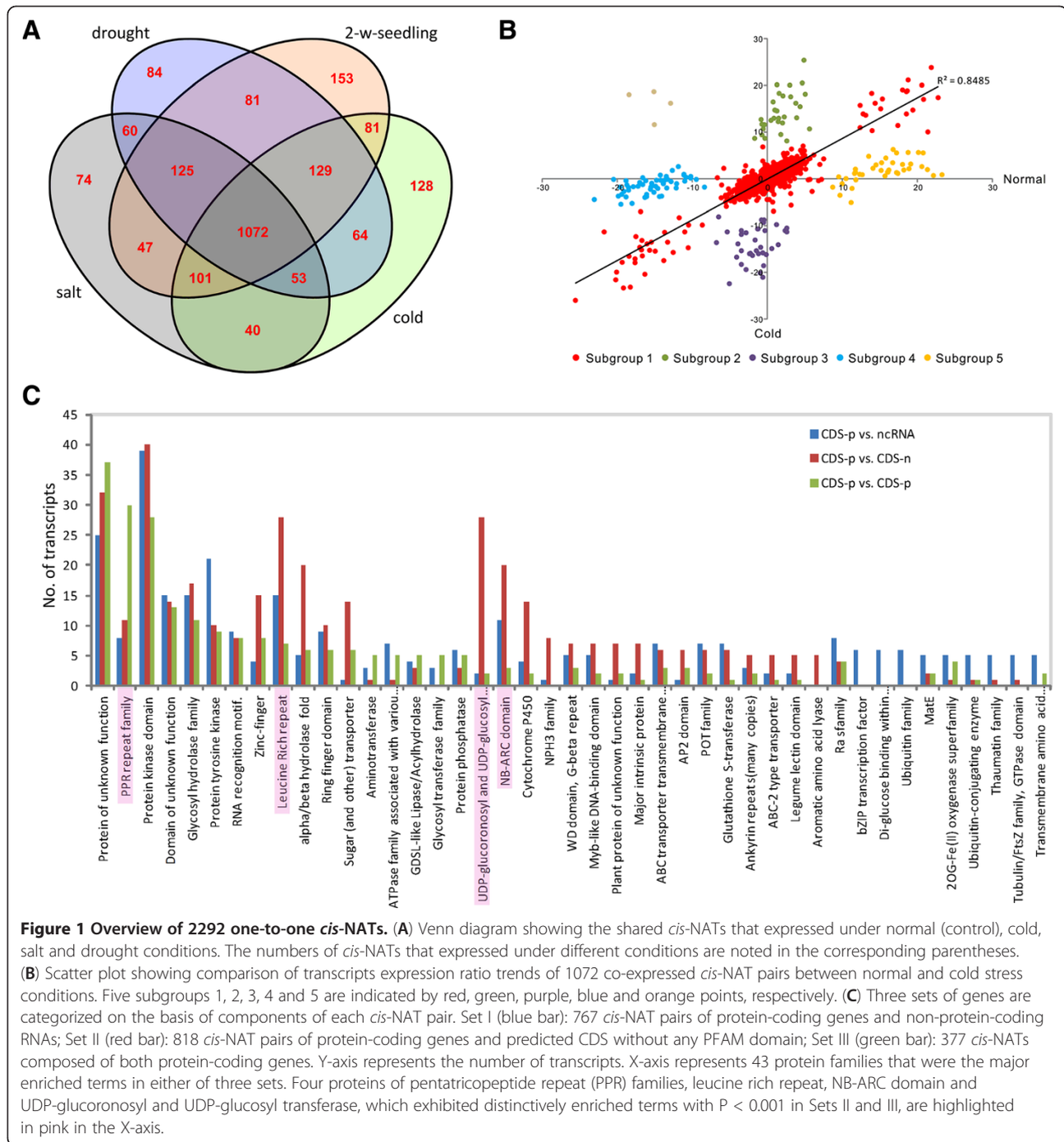
The numbers of *cis*-NATs expressed under different conditions were quite different (Figure 1A). The shared *cis*-NATs expressed under normal (control), cold, salt and drought conditions were also identified. Among them, 503 (21.9%) *cis*-NATs were detected preferentially under abiotic stresses compared to normal conditions. There were 84, 74 and 128 *cis*-NAT pairs clearly expressed under drought, salt and cold stresses, respectively. As an example, under cold treatment, sense and antisense transcripts of *cis*-NAT pair Os09t0482800-02 vs. CUFF.14823.1 were both much more expressed than under normal or other stressed conditions (Figure 3). Semi-quantitative RT-PCR and reverse transcription PCR (RT-PCR) were used to validate the expression of the *cis*-NAT pair under normal and cold conditions (Figure 3B). Functional annotation of this *cis*-NAT pair was composed of EF-hand protein (Os09t0482800-02) and ncRNA (CUFF.14823.1). Moreover, 1238 of 2292 (54.0%) *cis*-NATs were expressed in leaf epidermal cells. Of 1238

cis-NATs, 725 belonged to co-expressed pairs (Table 2). We further performed analysis of differentially expressed genes (DEGs). In total, we identified 112 *cis*-NATs with DEGs of at least either of each pair, which belonged to different subgroups (Additional file 11). Of them, expression levels of 69 *cis*-NATs (either sense or antisense transcripts) were up-regulated under abiotic stresses, while 46 *cis*-NATs were down-regulated under abiotic stresses. Moreover, the results showed no clear expression pattern between transcripts and nat-siRNAs (Figure 4). The number of nat-siRNAs notably increased along with higher transcriptional expression level under cold-stress conditions (Figure 4A and B); in contrast, some *cis*-NATs generated less nat-siRNAs, along with higher expression levels (Figure 4C and D). In addition, some *cis*-NATs generated nat-siRNAs with more complex expression patterns (Figure 4E-H).

We used an enrichment analysis of PFAM protein families [47] to functionally characterize three sets that could be annotated from 2292 *cis*-NATs. The three sets were 767 *cis*-NAT pairs of protein-coding genes and non-protein-coding RNAs (Set I), 818 *cis*-NAT pairs of protein-coding genes and predicted CDS without any PFAM domain-containing (Set II), and 377 *cis*-NATs composed of both protein-coding genes (Set III), respectively. Roughly, 43 protein families were the major enriched terms in either of three sets (Figure 1C). Of them, four proteins exhibited distinctively enriched terms with P -values < 0.001. PPR families were enriched in Set III, leucine rich repeat and NB-ARC domain were significantly reduced in Set III, and UDP-glucuronosyl and UDP-glucosyl transferase were significantly enriched in Set II (Figure 1C).

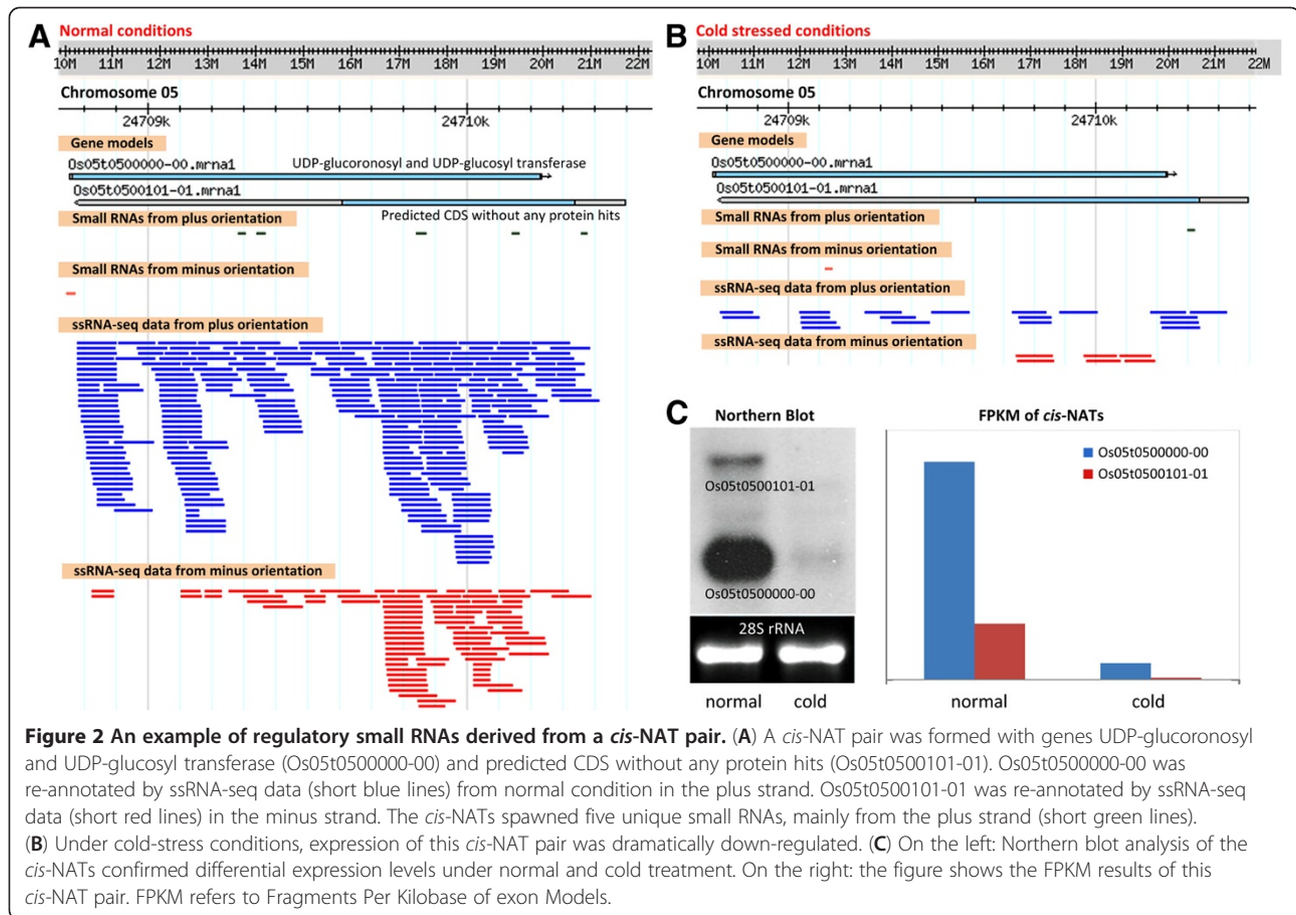
Networks formed by *cis*-NAT groups

We further investigated networks formed by *cis*-NATs in rice, named 'many-to-many' *cis*-NATs, i.e. one transcript



in a *cis*-NAT had more than one antisense partner. Networks formed by *cis*-NATs reflect the complexity of their post-transcriptional regulation. It has been reported that in *Arabidopsis*, several genes are involved in two *cis*-NATs, one is convergent and the other divergent [20,27]. In rice, we found 461 *cis*-NATs (composed of 685 transcripts) involved in 223 *cis*-NAT groups (Additional file 12). Of them, 209 groups belonged to one-to-two type, nine belonged to one-to-three type, four belonged to two-

to-two groups, and one belonged to one-to-four group, respectively. Five types were identified in one-to-two *cis*-NAT groups (Additional file 12). Except 46 groups which were composed of convergent together with divergent *cis*-NATs, it also included 74 enclosed together with enclosed, 40 enclosed together with divergent, 48 enclosed together with convergent, and one convergent together with convergent. Interestingly, we found only 17.4-34.7% *cis*-NAT groups could be detected with



simultaneous expression evidence and nat-siRNAs in the same sample from normal, cold, salt or drought conditions; another 29.2-48.6% cis-NAT groups were either expressed (Additional file 13). Similar to one-to-one type cis-NATs, protein kinase domain, leucine rich repeats, NB-ARC domain and the PPR repeat family were the major enriched PFAM protein domains.

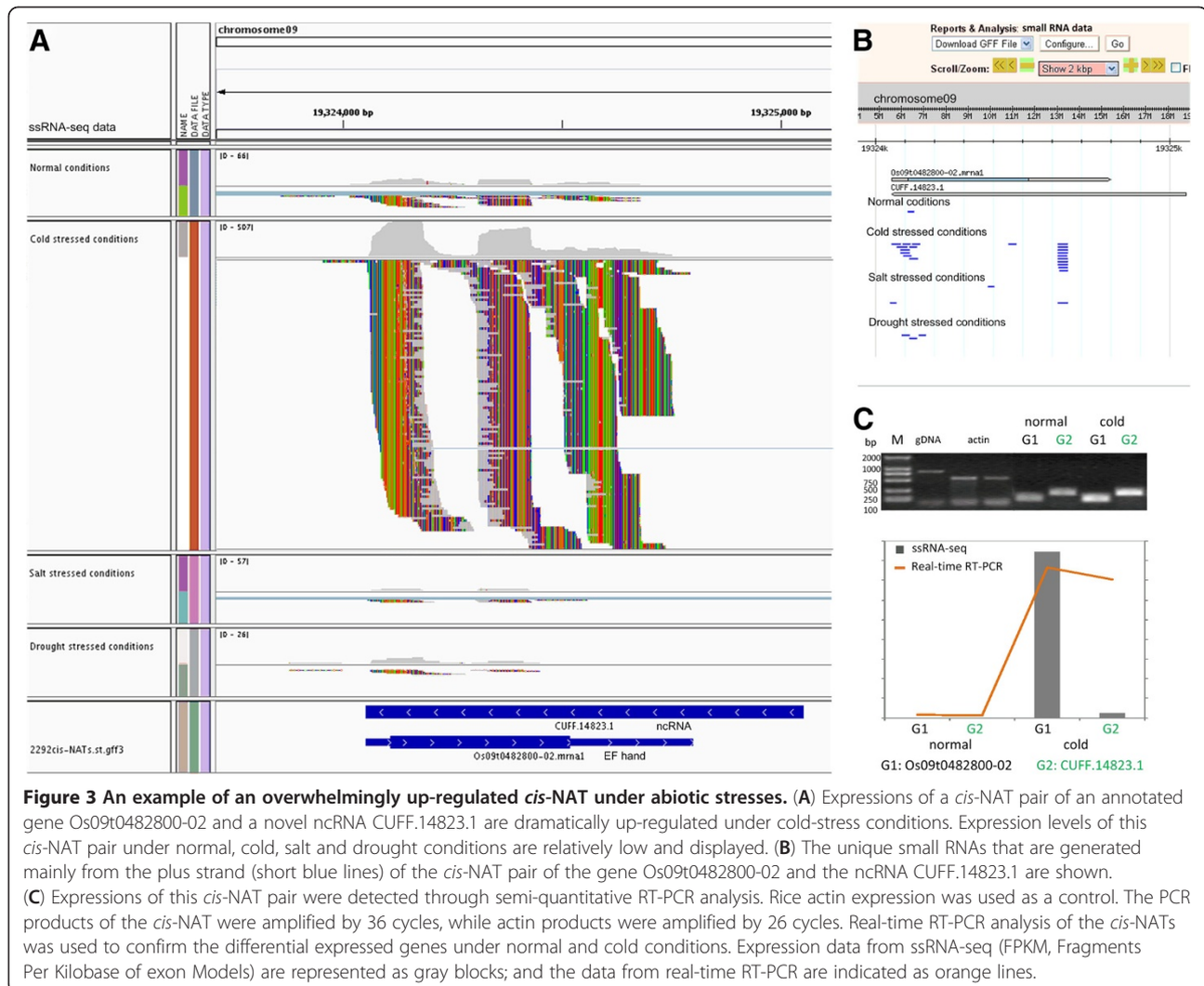
Discussion

Increasing amounts of evidence indicate that antisense transcripts are frequently functional and play various biological roles using different transcriptional and post-transcriptional gene regulatory mechanisms [1,2,48-53]. Consequently, we tried to advance rice cis-NATs analysis through an integrated approach of ssRNA-seq of rice transcriptome and genome-wide *in silico* transcriptome data analysis.

There have been five previous studies using global investigation of rice cis-NATs. An analysis of 32,127 full-length rice cDNA sequences showed 687 bidirectional transcript pairs [19]. Comprehensive expression profiles of rice using MPSS technology identified 11,001 antisense signatures corresponding to 8023 annotated genes with highly specific expression patterns [26]. A

comparative analysis of cis-NATs in eukaryotes identified 1088 rice cis-NATs referring to MSU and KOME gene datasets [17]. Combining pyrophosphate-based high-throughput sequencing of rice small RNA and computational analysis of the MSU rice gene models identified 344 cis-NATs formed by protein-coding genes [28]. Recently, 767 pairs of cis-NATs were identified using the rice genome annotation data (MSU TGAP 6.1) [35]. In this study, we identified 3819 pairs of rice cis-NATs. Of them, 2292 were identified as expressed and gave rise to small RNAs from their overlapping regions. The pairs of cis-NATs were confirmed to be formed in laser microdissection-captured rice seedling leaf epidermal cells, which were developed from a single-cell tissue. This indicated that antisense transcripts were more widespread and of more complex occurrence than previously found in the rice genome.

Moreover, qRT-PCR and Northern blot analyses confirmed the presence of cis-NATs and differential expression levels under normal and abiotic stresses. Nearly 9.7% of rice transcripts were involved in one-to-one or many-to-many cis-NATs formation. An overwhelming majority of cis-NATs (3358 of 3819, i.e. 87.9%) were of one-to-one type. Small RNAs generated from cis-NATs



were broadly detected under either normal developmental conditions or stresses. Similar to the situation for mammals [15], we found that the most prominent form of antisense transcription in rice was a non-coding RNA partner of a protein-coding transcript.

We identified 4873 novel gene loci in this study. Compared with previous research, more novel transcribed activity regions with reliable transcriptional orientation were identified by ssRNA-seq. This also indicated that exploration of the rice transcriptome atlas is far from finished. Along with further progress of high-throughput sequencing technology and development of more efficient software for *de novo* transcriptome assembly, more accurate transcriptional units (TUs) could be defined in the future.

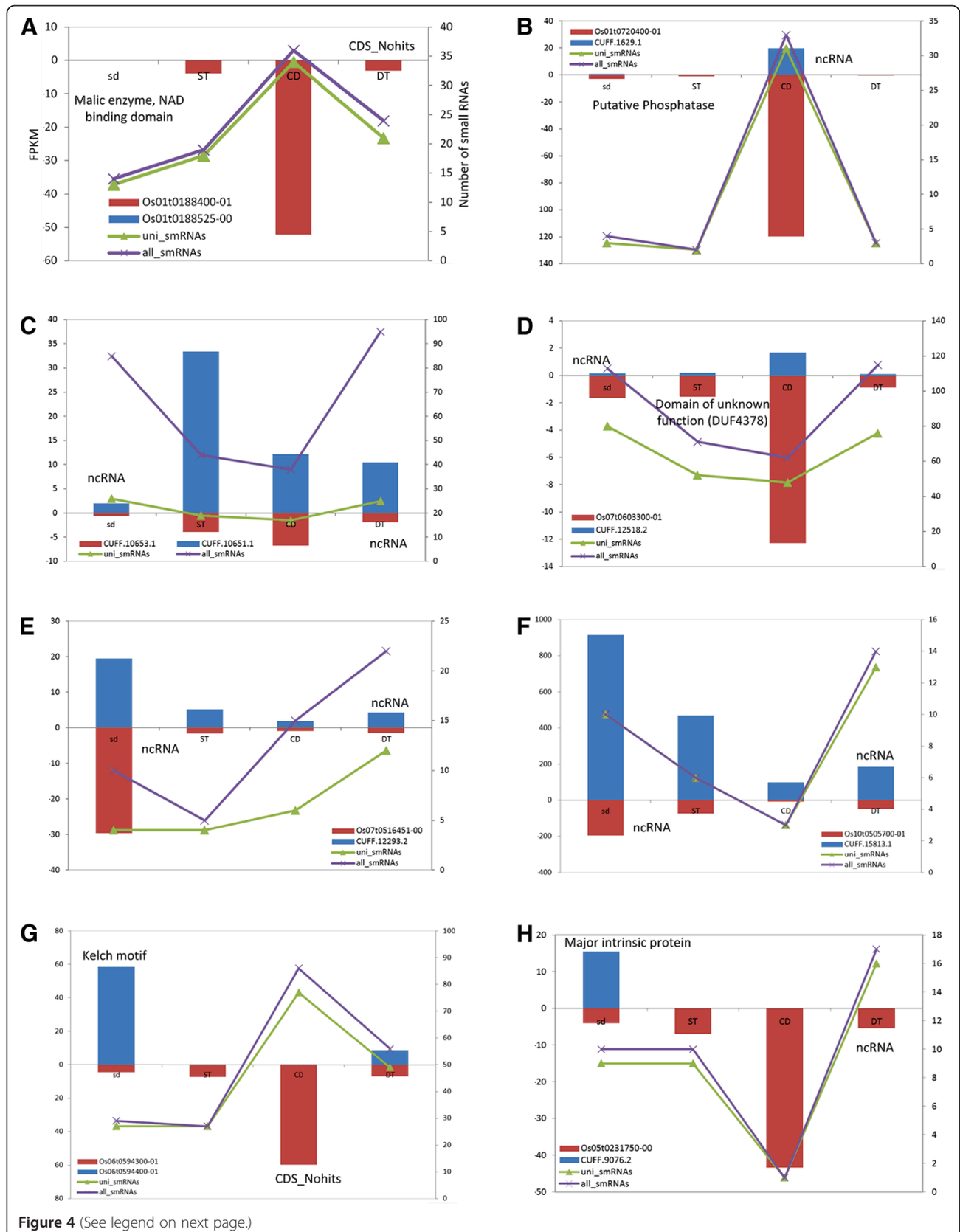
It should be pointed out that in our research about 4.5-10.6% of reads appeared to be aligned to antisense transcripts in error (Additional file 1). This is higher than the 3.88% rate reported by Wang et al. [54], who

used a slightly modified version of this dUTP method by increasing the incubation time with UDG to enforce the complete degradation of dUTPs.

In consideration of the limited amount of RNAi-related small RNAs (mainly including microRNAs and small interfering RNAs) in public databases, we proposed the following: (i) more microRNAs and siRNAs need to be identified and (ii) more classes of small RNAs that are engaged in RNAi-related machinery or those that are not will be found in future.

Conclusions

By applying a strand-specific RNA-seq approach, we systematically identified rice *cis*-natural antisense transcripts and putative nat-siRNAs. Our study profiled the most abundant of transcriptional active regions and revealed widespread occurrence of *cis*-NATs in rice, suggesting that regulation through *cis*-NATs and nat-siRNAs could be a common biological phenomenon.



(See figure on previous page.)

Figure 4 Expression patterns of eight one-to-one *cis*-NATs and the distribution of small RNAs located in overlapped regions of these gene pairs. (A-H) Expression patterns of eight *cis*-NATs are displayed. The left Y-axis of each graph represents FPKM of each transcript. Blue bars indicate FPKM of plus- transcripts (above the X-axis); red bars indicate FPKM of its partner of minus- transcripts (below the X-axis). The right Y-axis indicates the number of small RNAs located in overlapped regions. The green curves represent unique nat-siRNAs, and the purple curves represent all nat-siRNAs. FPKM refers to Fragments Per Kilobase of exon Models.

Both ssRNA-seq data and small RNA data obtained by high-throughput sequencing technology would supply important resources for further rice transcriptome analysis. It also indicated the feasibility of global investigations of *cis*-NATs by ssRNA-seq in eukaryotic genomes.

Methods

Materials and growth conditions

Rice seeds (*O. sativa* ssp. *japonica* cv. Nipponbare) were used in all experiments. Seeds germinated at 28°C in darkness for 2 d were transferred to a plant growth chamber to grow for 14 d under controlled conditions (12/12 h and 30/24°C of light/dark cycles) to produce seedlings and epidermal cells. For drought and salt stress treatments, 14-d-old seedlings were incubated in solutions containing 20% PEG-6000 and 200 mM NaCl, for 4 h at 30°C, respectively. For cold treatment, seedlings at the same developmental stage were treated at 4°C for 24 h in darkness.

Strand-specific cDNA library construction and sequencing

We prepared the strand-specific cDNA libraries according to a protocol [37]. The ssRNA-seq is a simple modification of the RNA-Seq method that incorporates deoxy-UTP during second-strand cDNA synthesis and subsequent destruction of the uridine-containing strand in the sequencing library. Thus it enables identifying the orientation of transcripts. Total RNA was isolated using the TRIzol reagent (Invitrogen), then total genomic DNA was removed from tissues using DNase (New England Biolabs), which was examined by gel electrophoresis. The OligoTex mRNA midi kit (Qiagen) was used to purify poly(A) mRNA from the total RNA samples. Next, mRNAs were fragmented using the RNA fragmentation kit (Ambion). The first cDNA strand was synthesized using random hexamer primers and second-strand cDNA was synthesized where dUTP was used instead of dTTP. In this step, Actinomycin D was used to increase strand specificity by inhibiting second-strand cDNA synthesis. At 15°C 0.5 µl of actinomycin D solution (120 ng/µl), 0.5 µl of RNase OUT (40 units/µl, Invitrogen) and 0.5 µl of SuperScript III polymerase (200 units/µl, Invitrogen) were added to the reaction. Then EB (20 µl) (10mM Tris-Cl, pH 8.5, Qiagen) was added to the reaction, and the dNTPs were removed by purification of the first strand mixture on a self-made 200 µl G-50 gel filtration spin-column equilibrated with

1mM Tris-Cl, pH 7.0. After second strand synthesis and DNA fragmentation process, the sequencing libraries were further constructed by following the manufacturer's instructions (Illumina). Fragments of 300-400 bp were recovered and purified, and then enriched by PCR for 15 cycles. Each library was loaded into one lane of the Illumina GA IIX for 2 × 120 bp pair-end sequencing at a concentration of 2 pM, except that library of normal seedlings was loaded into two lanes. Image analysis and base calling were finished using the Illumina GA processing pipeline v1.4.

Laser microdissection (LM)-captured rice seedling leaf epidermal cells and aRNA preparation

Leaves of 15-d-old seedlings of rice variety TP309 grown in a growth chamber at 12/12h and 25/22°C of day/night cycle were used. Seedling leaves were cut into pieces about 5 mm long and immediately processed by microwave-accelerated acetone fixation (BP-111-RS laboratory microwave, Microwave Research & Applications Inc.) and paraffin-embedded as described by Tang et al. [55]. Cross-sections of 10 µm thickness, parallel to leaf vascular bundles, were obtained using a Leica RM2235 rotary microtome. Paraffin-tape transfer system (Instrumedics) and Veritas Microdissection Instrument (Acturus Bioscience) were used for LM-capturing epidermal cells.

We carried out microdissection of epidermal cells, which are a group of cells (a homogenous population cells). There were about 200-500 cells per sample. The total RNAs of epidermal cells were extracted by PicoPure RNA isolation kit (Acturus, CA, USA) with the DNase (RNase-free, Qiagen) treatment. The integrity of the total RNAs was evaluated by Agilent 2100 Bioanalyzer using RNA-6000 Pico LabChips (Agilent Technologies). Because the total RNA quantity of LM-captured epidermal cells was about 10 ng, we used a TargetAmp two-round aminoallyl-aRNA (antisense RNA) amplification kit (Epicentre Biotechnologies, Madison, WI, U.S.A.) with Super-Script III and SuperScript II reverse transcriptases (Invitrogen, Carlsbad, CA U.S.A.) to amplify laser-microdissected RNA [56]. For each amplification, approximately 0.5 ng of total RNA (in a 2-µl volume) was used as starting material and, typically, 5 to 10 µg aminoallyl cRNA was recovered. For evaluating the fidelity of two-round RNA amplification, the TargetAmp one-round aminoallyl-aRNA amplification

kit (Epicentre) was used as a one-round amplification control. For each one-round amplification, 400 ng of total RNA was used as starting material and, typically, more than 10 µg of aminoallyl cRNA was recovered. The quality of amplified RNA was also evaluated by Agilent 2100 Bioanalyzer, and only those with a 'bell-shaped' curve with peak size > 300 nucleotides were used for RNA-seq.

Small RNA sequencing

The same materials described above were used for small RNA library construction using the Illumina small RNA preparation kit (v1.5) following the manufacturer's instructions. Small RNAs of 18-34 bp were enriched by polyacrylamide gel electrophoresis, and ligated to 5' and 3' adapters. The ligation product was reverse-transcribed into cDNA, which was then amplified by 15 PCR cycles and subjected to Illumina's Solexa proprietary sequencing. Here, each library was loaded into two lanes of the Illumina GA IIX for 35-bp sequencing, except that the library of cold treatment was loaded into one lane.

Northern analysis and qRT-PCR

The PCR primers and Northern probes were designed for *cis*-NAT validation (Additional file 14). Northern blot analysis was carried out as described [57]. To ensure that each pair of primers had specificity of its transcript, one was selected from the overlapping region of *cis*-NATs, and the other from the flanking region. Total RNA was extracted using the Trizol reagent (Invitrogen) according to the manufacturer's instructions. After treated with DNaseI (NEB), 5 µg of total RNA was used to synthesize the oligo (dT) primed first-strand cDNA using SuperScript™ II reverse transcriptase (Invitrogen). For semi-quantitative RT-PCR, rice actin expression was used as a control, *cis*-NAT PCR products were amplified by 36 cycles, while actin products were amplified by 26 cycles. Real-time RT-PCR was performed on the Applied Biosystems 7500 real-time PCR System. Diluted cDNA was amplified using SYBR Premix Ex Taq™ (TaKaRa). The expression levels of transcripts were normalized by endogenous eEF-1α (AK061464) transcripts. Three technical replicates were taken for each set.

Assembly of rice transcripts

All ssRNA-seq data obtained from normal and cold, salt and drought stress conditions were respectively mapped to the rice reference genome [41] using software TopHat [43]. Tolerances were set to allow at most two mismatches for paired-end reads in each alignment; and reads with multiple alignments were ignored. The corresponding outputs together with RAP-DB genome

annotation data [42] were subsequently used for transcript assembly to detect known and unannotated transcripts and isoforms by another software package, Cufflinks [44,46]. Then, we integrated these transcripts together according to another command, Cuffmerge, from the Cufflinks package. Finally, referring to the renewed gene models, we calculated FPKM (Fragments Per Kilobase of exon Models) of each transcript under normal, abiotic stressed conditions, and leaf epidermal cells as well [45]. Open reading frames (ORFs) of novel transcripts were predicted using the 'getorf' program of EMBOSS package [58], with the longest ORF extracted for each transcript. For functional annotation, all transcripts were searched against the PFAM database [47] using HMMER v3.0 (E-value < 0.0001) [59]. The MSU Oryza Repeat Database [60] was used to determine transposable element coordinates on the rice pseudomolecules and all transcribed regions, which were annotated using RepeatMasker [61].

Identification of *cis*-NATs and *nat*-siRNAs

The renewed gene annotation results were applied as models. Those assembled genes with uncertain transcriptional orientation were filtered out first. To profile all potential *cis*-NAT pairs in the rice genome, transcripts that originated from the same locus but from opposite strands and with non-redundant overlapped length > 25 nt were chosen. From each group of sense-antisense pairs extracted, we selected as representative pairs those with the longest exonic overlap. We wrote perl scripts to classify *cis*-NAT types: convergent, divergent or enclosed. Furthermore, we examined expression levels of all assembled genes using ssRNA-seq data. Thus, we could identify expressed transcripts from putative *cis*-NATs. According to the number of antisense partners in each *cis*-NAT pair, two types of *cis*-NATs were defined. One was one-to-one type, i.e. one transcript in a *cis*-NAT pair has only one antisense partner. Another is many-to-many *cis*-NATs, i.e. one transcript in a *cis*-NAT has more than one antisense partner.

Raw data of small RNA reads from four corresponding small RNA libraries were aligned to the rice genome [41] by SMALT v0.5.7 with default parameters [62]. Mapping scores below the threshold were not reported. After removing the 5'- and 3'-adapters, only those continuously and perfectly matched reads with length of 18-34 bp were extracted for further analysis. Reads with multiple alignments or with any mismatches were completely filtered out. Next, we discarded those sequences that could be unambiguously mapped to rRNA, tRNA, sn/snoRNA, mitochondria and chloroplasts. We calculated small RNA densities according to a published method [28]. Small RNAs (at least one unique read) which fully located in overlapped regions of *cis*-NATs

were extracted as putative nat-siRNAs. The *cis*-NATs were identified from the set of DEGs across normal and three abiotic-stress libraries. The analysis was carried out based on the following criteria: FPKM of transcripts were used for comparison by computing fold changes (with absolute value ≥ 2) and Fisher's exact test ($p < 0.001$) according to an 'R' statistical package named 'DEGseq' [63].

In addition, R^2 is the square of the Pearson product-moment correlation coefficient relating the regressor and the response variable. R^2 gives some information about the goodness-of-fit of a model. In regression, R^2 is a statistical measure of how well the regression line approximates the real data points. 'R' software was used to calculate R^2 [64] of *cis*-NATs between normal and stressed conditions.

Data release

The raw sequences were deposited in the EBI European Nucleotide Archive with accession number E-MTAB-721 (<http://www.ebi.ac.uk/arrayexpress/browse.html?keywords=E-MTAB-721>) and ERP001962 (<http://www.ebi.ac.uk/ena/data/search?query=ERP001962>). The assembled transcripts can be freely downloaded from http://www.ncgr.ac.cn/scientific_data.asp.

Additional files

Additional file 1: Summary of pair-end reads of ssRNA-seq and small RNAs from normal and three abiotic stress conditions.

Additional file 2: Statistics of rice transcripts.

Additional file 3: Summary of 5813 putative *cis*-NATs in rice. This table lists all *cis*-NATs identified from rice renewed assembled transcripts. Each column represents plus transcripts (transcriptional orientations are the same as the reference), its length, minus transcripts, length, overlapped length and detailed overlapped locations in rice genome (IRGSP v5.0) as indicated.

Additional file 4: Overview of small RNAs in rice. (A) Distribution of the lengths of all unique small RNAs generated by high-throughput sequencing from rice under four different conditions. (B) Length distribution of nat-siRNAs. (C) Distribution of all unique small RNAs located in different sequence components. (D) Distribution of nat-siRNAs located in different sequence components. The blue, red, green and purple bars represent small RNAs or nat-siRNAs from normal, salt stressed, cold stressed and drought stressed conditions, respectively (A-D). (E) First-nucleotide distribution of all unique small RNAs under four different conditions. (F) First-nucleotide distribution of nat-siRNAs under four different conditions.

Additional file 5: Sequences of small RNAs mapped to the overlapped regions of *cis*-NATs. The 25,420, 18,598, 18,152 and 28,807 unique small RNAs from normal, salt, cold and drought conditions, respectively, are shown to be perfectly mapped to the overlapped regions of *cis*-NATs.

Additional file 6: Details of 2292 expressed *cis*-NATs. The detailed information of 2292 *cis*-NAT pairs with expression evidence (i.e. FPKM > 0) of both sense and antisense transcripts and with nat-siRNAs in the overlapping region under normal (1789 pairs), cold (1668), salt (1572) and drought (1668) conditions are shown. Columns 1-4 and 9-11 represent *cis*-NATs type, transcripts, protein domain and the length of transcripts, respectively. Columns 5-8 and 12-15 represent FPKM of transcripts

under normal, salt, cold and drought conditions, respectively. Column 16 shows the overlapped length. Columns 17-20 show the number of corresponding nat-siRNAs located in the overlapped regions. The rest of the columns are the detailed strand bias of nat-siRNAs from four different conditions.

Additional file 7: Lists of 166 *cis*-NATs that produced nat-siRNAs exclusively in the overlapping regions with more than five unique small RNAs from four different conditions. Of 166 *cis*-NATs, 72, 58, 66 and 75 *cis*-NATs were obtained from normal, salt, cold and drought treatments, respectively. These *cis*-NATs are listed separately in four tables, respectively indicated by blue, orange, red and green colour.

Additional file 8: Lists of 13 *cis*-NATs that produced nat-siRNAs more enriched in the overlapping regions than the non-overlapping regions. Of 13 *cis*-NATs, nine, six, five and five *cis*-NATs obtained from normal, salt, cold and drought treatments, respectively. These *cis*-NATs are listed separately in four tables, respectively indicated by blue, orange, red and green colour.

Additional file 9: Strand bias of *cis*-NATs that gave rise to nat-siRNAs. The strand bias is computed among different types of *cis*-NAT pairs under four different conditions. A total of nine types are listed. Here, plus indicates that the orientation of transcripts are the same as the reference genome, minus indicates opposite orientation of transcripts. One strand means only one strand of the *cis*-NATs gave rise to small RNAs. The plus/minus or minus/plus means one strand of the *cis*-NATs spawns at least five (part A) or two (part B) fold more small RNAs than the other.

Additional file 10: Scatter plots of expression of 1072 *cis*-NATs. The two scatter plots compare transcripts expression ratio trends of 1072 co-expressed *cis*-NAT pairs between normal and drought-stress conditions (A), and between normal and salt-stress conditions (B). The results show five subgroups: red, green, purple, blue and orange spots represent subgroups 1-5, respectively.

Additional file 11: Lists of 112 *cis*-NATs with DEGs. Differentially Expressed Genes (DEGs) were identified in one-to-one *cis*-NATs under salt, drought and cold treatments by comparing their FPKM with that under normal conditions. The gene names, FPKM, fold change (with absolute value > 2) and p-value (< 0.001) are listed.

Additional file 12: All many-to-many *cis*-NAT groups. In total, 223 *cis*-NAT groups were identified as many-to-many type. Of them, 209 groups belonged to one-to-two type, nine belonged to one-to-three type, four belonged to two-to-two groups, and one belonged to the one-to-four group. The meaning of each column is as given in Additional file 6.

Additional file 13: Statistics of 209 networks formed by *cis*-NAT groups. The numbers of one-to-two *cis*-NAT groups with expression evidence under normal, salt, cold and drought conditions are shown. These one-to-two *cis*-NAT groups were divided into five classes according to their *cis*-NATs types. Here, 'All EXP' indicates all *cis*-NAT groups with expression evidence (FPKM > 0) of both sense and antisense transcript, and with nat-siRNAs (number of small RNAs > 1) in overlapping region as well. 'Either EXP' indicates either of *cis*-NAT groups with expression evidence of both sense and antisense transcript, and with nat-siRNAs in the overlapping region as well.

Additional file 14: Primers designed for real-time RT-PCR and Northern Blots in this research.

Competing interests

The authors declare that they have no competing interests.

Authors' contributions

TL and BH initiated the project; TL and BH designed the experiments; GL, YG, YZ, ZZ, WL, WT and QF carried out the experiments; TL, CZ and YZ performed the computational analyses; TL, YZ and BH participated in the experimental analyses. TL and BH coordinated the research. TL and BH wrote the paper. All authors have read and approved the manuscript for publication.

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