Identification of Novel *Oryza sativa* miRNAs in Deep Sequencing-Based Small RNA Libraries of Rice Infected with *Rice Stripe Virus*

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Abstract

MicroRNAs (miRNAs) play essential regulatory roles in the development of eukaryotes. Methods based on deep-sequencing have provided a powerful high-throughput strategy for identifying novel miRNAs and have previously been used to identify over 100 novel miRNAs from rice. Most of these reports are related to studies of rice development, tissue differentiation, or abiotic stress, but novel rice miRNAs related to viral infection have rarely been identified. In previous work, we constructed and pyrosequenced the small RNA (sRNA) libraries of rice infected with *Rice stripe virus* and described the character of the small interfering RNAs (siRNA) derived from the RSV RNA genome. We now report the identification of novel miRNAs from the abundant sRNAs (with a minimum of 100 sequencing reads) in the sRNA library of RSV-infected rice. 7 putative novel miRNAs (pn-miRNAs) whose precursor sequences have not previously been described were identified and could be detected by Northern blot or RT-PCR, and were recognized as novel miRNAs (n-miRNAs). Further analysis showed that 5 of the 7 n-miRNAs were up-expressed while the other 2 n-miRNAs were down-expressed in RSV-infected rice. In addition, 23 pn-miRNAs that were newly produced from 19 known miRNA precursors were also identified. This is first report of novel rice miRNAs produced from new precursors related to RSV infection.

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Introduction

MicroRNAs (miRNAs) are small 19-24 nt RNAs that play essential roles in eukaryotes by targeting complementary miRNAs for degradation or translational repression [1,2]. In plants, primary miRNA (pri-miRNA) is first transcribed by polymerase II, and then processed by Dicer-like 1 (DCL1) into the precursor miRNA (pre-miRNA), normally of about 70–300 nucleotides (nt). The pre-miRNA is further processed into the mature miRNA-miRNA* duplex [3,4,5]. These processes occur in the nucleus. In the next stage, the duplex is transferred into the cytoplasm and unwound [3,4]. The miRNA is then assembled into and RNA-induced silencing complex (RISC) and guides the RISC to cleave or suppress the target mRNA [3,4,6].

miRNAs in plants regulate leaf morphogenesis, the development of roots and flowers and other key processes, and are recognized as important regulators of plant development [7,8,9,10,11,12]. Recent research has revealed that miRNAs also play roles in plant defense against pathogens by regulating the expression of resistance (R) genes directly or indirectly, or targeting the viral genome to impair viral replication [13,14,15,16,17]. Hence, the miRNA pathway also plays a key role during pathogen-plant interactions.

In plants, over 4000 miRNAs have been identified from over 50 species (miRBase version 18.0, http://www.mirbase.org/cgi-bin/browse.pl) [18]. *Medicago truncatula*, *Oryza sativa* and *Glycine max* are the three plants that have the most identified miRNAs (respectively 674, 661 and 395 miRNAs) [18]. Some miRNA families have functions that are conserved across the plant kingdom and thus their sequences are similarly conserved (e.g. miR156, miR159, miR160 and miR165). Other miRNA families are specific to particular plants, and are not found elsewhere, indicating that they have novel and specific functions [19,20].

With the development of next generation sequencing technologies, deep-sequencing has provided a powerful high-throughput strategy for identifying novel miRNAs. In this way, hundreds of miRNAs have been identified from Arabidopsis, Brassica rapa, rice, wheat, barley, peanuts, grapevine and other plants [21,22,23,24,25,26,27,28,29,30,31,32]. In rice, Sunkar et al. identified 23 new miRNAs from three small RNA (sRNA) libraries of control rice seedlings and seedlings exposed to drought or salt.
stress; six of the new miRNAs are conserved in monocots [27]. Chen et al identified 24 novel microRNA families from rice embryogenic callus, some of which were suggested to function in meristem development [33]. Li et al investigated the H$_2$O$_2$-regulated miRNAs in rice seedlings and discovered 32 new miRNAs [34]. Peng et al identified 43 novel miRNAs from the sRNA libraries of rice spikelets [35], while Wang et al identified 75 novel miRNAs from the developing pollen of rice [36].

Until now, most reports of novel rice miRNAs have been related to studies of rice development, tissue differentiation, or abiotic stress, while the novel rice miRNAs related to viral infection have rarely been identified. Identifying the novel miRNAs related to viral infection in rice would be helpful to broaden our understanding of the miRNA response of rice to virus infection in general and is likely to provide a model for other cereal plants in particular. *Rice stripe virus* (RSV) is the type member of the genus *Tenuivirus*. It is transmitted by the brown planthopper *Laodelphax striatellus* in a persistent manner and causes rice stripe disease, which is severe in rice fields in East Asia [37]. In a recent study, Du et al reported the changed expression of the known miRNAs during the RSV infection, and found that the known osa-miR168, 156, 396, 159, 535, 166, 172, 167, 528 and 444 were the ten most...
abundant miRNA families in RSV-infected rice, and that RSV infection induced the expression of novel miRNAs in a phased pattern from several conserved miRNA precursors, and enhanced the accumulation of some rice miRNA*s, but not their corresponding miRNAs [38].

In previous work, we constructed and pyrosequenced sRNA libraries of rice inoculated with RSV and mock-inoculated controls, and described the character of the small interfering RNAs (siRNA) derived from the RSV RNA genomes [39]. Here, we report the identification of novel miRNAs from the abundant

| sRNA No. | sRNA sequence | length | Sequencing reads | The corresponding known miRNA of sRNA |
|----------|---------------|--------|-----------------|-------------------------------------|
| Seq1     | UUUGGAUUGAAGGGAGCUCUG | 21     | 140             | osa-miR159a.1                       |
| Seq2     | UGCUGGCUCCCCUGUAUGCAG | 21     | 121             | osa-miR160a                         |
| Seq3     | UGCUGGCUCCCCUGUAUGCAG | 21     | 118             | osa-miR160f                         |
| Seq4     | UCAGAUAACCUCUGCAUGCCAG | 21     | 572             | osa-miR162a                         |
| Seq5     | UGGAGAGGACGGGAGCUGUGCA | 21     | 1851            | osa-miR164a                         |
| Seq6     | UCAGACCAAGCUCUAUUCCCC | 21     | 3647            | osa-miR166a                         |
| Seq7     | UCAGCAAGCUCUAUUCUCUC | 21     | 601             | osa-miR166g                         |
| Seq8     | UCCAGACCAAGCUCUAUUCUCC | 21     | 318             | osa-miR166k                         |
| Seq9     | UGAAGCUGCAGCUGAUGCUA | 21     | 118012          | osa-miR167d                         |
| Seq10    | UGAAGCUGCAGCUGAUGCUA | 21     | 24678           | osa-miR167a                         |
| Seq11    | UCCUGUGUGCAGAUGCGGCAC | 21     | 83988           | osa-miR168a                         |
| Seq12    | CAGCAAGAUGACUGCUCCGG | 21     | 738             | osa-miR169b                         |
| Seq13    | UAGCAAGAUGACUGCUCCUG | 21     | 342             | osa-miR169f.1                       |
| Seq14    | CAGCAAGAUGACUGCUCCGA | 21     | 269             | osa-miR169a                         |
| Seq15    | UGAUUGAGCUGCGAUGCAUAUC | 21     | 350             | osa-miR171b                         |
| Seq16    | AGAAUCUUGAUGAUCGCAU | 21     | 94444           | osa-miR172a                         |
| Seq17    | UUCCACAGCUUUUCUGAUCUU | 21     | 127             | osa-miR396c                         |
| Seq18    | UCCACAGCUCUUUCUGAUCUG | 21     | 1662            | osa-miR396d                         |
| Seq19    | UCCAAAGAAGAGUCUGCCUG | 21     | 286             | osa-miR399d                         |
| Seq20    | UUGACACUCUCUCUCCUGGC | 21     | 157             | osa-miR408                          |
| Seq21    | UGCAGUGUGGCUCCUCAAGCUU | 21     | 6725            | osa-miR444b                         |
| Seq22    | UGCAGUGUGGCUCUCAGCAGCU | 21     | 333             | osa-miR444a                         |
| Seq23    | UUGUGUCUCAGCGUGUGCC | 21     | 439             | osa-miR444b.1                       |
| Seq24    | UGGAGGGCAUGCGAGAGAG | 21     | 19789           | osa-miR528                          |
| Seq25    | UGACAGCAGAGAGACCGC | 21     | 2553            | osa-miR535                          |
| Seq26    | AAGACGCGAUGAUUAAAGUGGACA | 24     | 387             | osa-miR812g                         |
| Seq27    | UAUGAUUUGUGGCAAUUGCAAA | 21     | 167             | osa-miR819a                         |
| Seq28    | UUGAAGACCAUGCAUGCAAA | 21     | 950             | osa-miR827a                         |
| Seq29    | UGAUAAUCUUGGCUUCAUCCAA | 21     | 328             | osa-miR1320-3p                       |
| Seq30    | UAGGGACUAUCUCUUGUGCC | 21     | 722             | osa-miR1425-5p                       |
| Seq31    | CAGCAAGAAGAGUAUUGCUUAAU | 21     | 414             | osa-miR1425-3p                       |
| Seq32    | UAGGGACUAUGCGAUGAUAUGCU | 21     | 140             | osa-miR1428e-3p                      |
| Seq33    | GUUGCACGGGGGUUGUUGUGUCG | 24     | 520             | osa-miR1429-3p                       |
| Seq34    | AUCAGGAGAUGACACACGAC | 21     | 528             | osa-miR1432                         |
| Seq35    | UGGAAGUUGGGAGAUUGGG | 21     | 131             | osa-miR1850.1                       |
| Seq36    | CUAGAUAUUUGGUAUUGGAG | 24     | 763             | osa-miR1862e                        |
| Seq37    | AGUCUGUAAUCAGCUAAUGUAG | 24     | 253             | osa-miR1863                         |
| Seq38    | UGGCAUGAUUGAUGCCUGGCAC | 24     | 114             | osa-miR1870                         |
| Seq39    | AAAGCUACGGGUUGCUAUAUUUA | 24     | 103             | osa-miR1884b-3p                      |
| Seq40    | UAUGAUGAGUUGCAUUGUCAC | 21     | 194             | osa-miR2871-3p                       |
| Seq41    | UUCUGUGUGUGCUAGAAGAC | 21     | 299             | osa-miR5144-5p                       |
| Seq42    | AGCUUCUGACCGUGACGUUCUC | 24     | 299             | osa-miR5150-5p                       |
| Seq43    | UUGAGAAGGUAUUGAUCGAGAU | 21     | 163             | osa-miR5542                         |

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Figure 2. sRNAs that were recognized as newly produced miRNAs from known precursors and their location in the secondary structures of their precursors. The sequences of the known mature miRNAs in precursors are colored with blue. Names, sequences and sequencing reads (bracketed) of sRNAs identified here are colored with green. The location of sRNA is shown by a black line. Mismatched nucleotides are shown in red.

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Figure 3. sRNAs that were recognized as newly produced miRNAs from known precursors and their location in the secondary structures of their precursors. The sequences of the known mature miRNAs in precursors are colored with blue. Names, sequences and sequencing reads (bracketed) of sRNAs identified here are colored with green. The location of sRNA is shown by a black line. Mismatched nucleotides are shown in red.

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sRNAs (with a minimum of 100 sequencing reads) in the reported sRNA library of RSV-infected rice. 7 putative novel miRNAs (pmiRNAs) whose precursor sequences have not been described before were identified. Transcription analysis revealed that 7 pmiRNAs were produced in rice, and that their expression levels changed in RSV-infected rice. This is first report on novel rice miRNAs produced from new precursors related to RSV infection, and will deepen our understanding of miRNA functions during RSV infection.

Materials and Methods

sRNA libraries used here

In our previous work, we constructed sRNA libraries of RSV-infected and non-infected rice by the Illumina Solexa sequencing system [39]. A total of 917,776 and 1,009,009 unique sequences of 18–30 nt long small RNAs were contained in the respective sRNA libraries of RSV-infected and non-infected rice. About 23% of within these two sets was similar. Almost 40% of sequences were 24 nt long, 23% of sequences were 21 nt and 15% were 22 nt [39]. Here, we attempted to identify novel miRNAs in the RSV-infected sRNA library. For optimum identification and detectability by gel blot in the following experimental confirmation, we only chose to analyze the highly abundant sRNAs (total 239 sequences) that had a minimum of 100 sequencing reads (Table S1). Three sRNA libraries of RSV-infected rice reported by Du et al were also used to search for the putative miRNAs identified here [38].

Identification of putative novel miRNAs

The analysis was done in three steps. In the first step, we aligned the 239 chosen sRNAs with all the mature miRNAs in miRBase to identify the known miRNAs. The recognized miRNA sequences in sRNAs were excluded from the next step. In the second step, the remaining sRNAs were aligned with all the reported miRNA precursor sequences to identify putative miRNAs newly produced from the known precursors. These sRNAs were excluded from the following step. In the third step, the precursor sequences of all remaining sRNAs were predicted using a modification of the reported method [27]. Briefly, searches were made for the sRNAs in the rice genomic sequences [http://www.ncbi.nlm.nih.gov/blast/Blast.cgi] and their loci in the genome were recorded. The sRNAs that had more than 26 loci in the rice genome were considered in the following analysis for two reasons: 1) the highest number of loci of known miRNAs in the rice genome is 26 (os-miR395); and 2) the higher number of loci more often occurs when the loci are located in the repeat-rich regions of the rice genome; sRNAs but not miRNAs can be produced from the repeat-rich regions that have predicted fold-back structures (miRBase). This largely excludes the contaminative effect of siRNA on miRNA identification.

For each locus, two sequences respectively extending 200 nt upstream and 20 nt downstream, or 20 nt upstream and 200 nt downstream of the sRNA were extracted for secondary structure prediction. The secondary structures were predicted by the Mfold RNA folding platform [http://mfold.rna.albany.edu/?q=mfold/RNA-Folding-Form]. A sequence that has a secondary structure with at least 16 paired nucleotides in its stem region and has free energy less than or equal to −20 kCal/mol was considered to be a putative precursor sequence.

Plant materials

RSV-infected rice were prepared as described [39]. Briefly, viruliferous adult brown planthoppers (Laodelphax striatellus Fallen) (carrying the RSV-Zhejiang isolate) were transferred onto healthy rice seedlings (Oryza sativa L. japonica. cv. Nipponbare) at the three-leaf stage for virus inoculation. Control seedlings were inoculated with non-viruliferous planthoppers. After 72 h, the planthoppers were removed. Systemic infections were confirmed by RT-PCR specific for RSV Zhejiang isolate [40]. One week after inoculation, leaves were collected from the infected and control (Mock) plants, frozen and stored at −80°C until used. All rice plants were grown in a glasshouse at 28–30°C day/25°C night, with a 12 h day/night light cycle under well-watered conditions.

sRNAs gel blot analysis

Total RNA was isolated from the frozen plant materials with Trizol (Invitrogen, USA) according to the manufacturer’s instructions. 50 μg of DNase-treated total RNA was separated on a 15% polyacrylamide gel, and transferred electrophoretically to Hybrid-N+ membranes (Amersham Bioscience) using 20×SSC. Membranes were baked at 80°C for 2 hours. DNA oligonucleotides complementary to the putative miRNA sequences were end-labeled with DIG using the DIG Oligonucleotide 3′-end labeling Kit (Roche). Membranes were pre-hybridized for at least 1 h and hybridized overnight at 42°C using DIG High Prime Labeling and Detection Starter Kit II (Roche). The hybridization signals were visualized by exposure to X-ray film (Kodak).

Reverse transcription (RT)-PCR and real-time PCR for sRNAs

We used the published method for RT-PCR of sRNAs with modifications [41]. Briefly, for each sRNA, the specific stem-loop RT primer was used for reverse transcribing from purified total RNA. Then the RT product was used for PCR with the specific forward primer and the universal reverse primer. For SYBR green-based real-time PCR analysis, the reactions were incubated in a 384-well plate at 95°C for 10 min, followed by 40 cycles of 95°C for 15 s and 60°C for 1 min. U6 was used as the internal control. All reactions were run in triplicate, and the results were analyzed by the ΔΔCt method. The primers used are listed in Table S2.

Target prediction

Putative targets for the novel miRNAs were predicted by psRNATarget [http://plantgrn.noble.org/psRNATarget/] [42], using the DFCI Oryza sativa gene index release 18.0 as a reference set. The parameters for prediction were default, namely, maximum expectation was set at 3.0; length for complementarity scoring was set at 20 bp; target accessibility (range 0–100, less is better) was set at 25.0; Flanking length around target site for target accessibility analysis was set at 17 bp upstream and 13 bp downstream.

Statistical analysis

The number of reads of each sRNA from RSV-infected rice was divided by the number of reads in the non-infected controls of the same experiment. A two-tailed t-test was used to evaluate whether the means of these ratios from several experiments differed significantly from 1 (equal numbers of reads).

Results

Identification of known miRNAs

To identify the novel miRNAs from the library, we chose 239 sRNAs that had a minimum of 100 sequencing reads for analysis according to the method reported by Sunkar et al [27]. By searching in the miRBase database (Version 18.0, http://www.mirbase.org/index.shtml), 43 sRNAs that completely matched with the deposited rice miRNAs were identified (Table 1).
Meanwhile, 52 sRNA that did not completely match with the deposited rice miRNAs but contained identical or similar (>95% identity) seed sequences to the deposited rice miRNAs were also found (Figure 1). Considering that the isoforms of miRNAs exist more often in pyrosequenced sRNAs [43,44,45], we here recognized these 52 sRNAs as known rice miRNAs. Thus, a total of 95 known miRNAs were identified. These miRNAs belong to 46 miRNA families, of which the five with the highest frequency in the library were osa-miR167, 168, 528, 172 and 444. Other well-known miRNA families, osa-miR 156, 169, 170 and 397 also existed in the library (Table 1 and Figure 1).

Identification of miRNAs newly produced from known miRNA precursors

One miRNA precursor can produce several mature miRNAs with different sequences. Du et al (2011) recently reported that RSV infection induced different miRNAs to be produced from a single precursor in a phased pattern [38]. Hence, it was possible that the remaining 144 sRNAs contained some miRNAs that had been produced from a known conserved precursor but did not match known mature miRNAs. To identify these miRNAs, all 144 sRNAs were aligned with the known rice miRNA precursor stem-loop sequences. 23 of the sRNA sequences were identified among 19 known miRNA precursors, and were therefore considered to be novel miRNAs newly produced from known miRNA precursors. Their locations on the precursors are shown in Figure 2 and Figure 3.

Among these 23 sRNAs, 12 were produced from the 5’ arm and 11 were produced from the 3’ arm of their respective precursor. Seq114 and 115 were located at the miRNA* region of osa-miR400; Seq116 was located at the miRNA* region of osa-miR167 h; Seq117 was located at the miRNA* region of osa-miR398a; Seq118 was located at the miRNA* region of osa-miR444c. These sRNAs may be the miRNA* or miRNA* isoforms of the corresponding miRNAs.

We also searched for these 23 sRNA in the control library and other reported RSV and non-infected rice sRNA libraries, and found that they existed in the libraries with different sequencing reads (Table 2). Statistical analysis showed that Seq99, Seq102 and Seq107 were significantly down-expressed in RSV-infected rice, indicating that they may have a function in response to RSV infection (Table 2).

### Table 2. The putative novel miRNAs newly produced from known miRNA precursors.

| sRNA No. | sRNA sequence | Length | SR in Exp1 | SR in Exp2 | SR in Exp3 | SR in Exp4 | Mean (ln(i/n)) | SD | t (3DF) | Significance (alpha) |
|----------|---------------|--------|------------|------------|------------|------------|---------------|----|---------|----------------------|
| Seq96    | AAGGATGACTTCGCGATGGAGCA 24 285(231) 39(81) 0(0) 2(11) 0.63 | 0.542 | -1.17 | 0.325 |
| Seq97    | ATGACGACTGTCGGGCTTATAC 24 223(214) 42(69) 19(128) 27(38) 0.63 | 0.369 | -2.02 | 0.137 |
| Seq98    | AGTGGCCATTGTATGCAACAAC 24 182(224) 36(82) 7(44) 8(6) 0.69 | 0.508 | -1.24 | 0.304 |
| Seq99    | TTTATCTGTTGTCTTCTCAAA 24 153(195) 5(14) 5(26) 9(51) 0.38 | 0.283 | -4.39 | 0.022* |
| Seq100   | CAGCATTCTACGTCGAGGCC 21 547(472) 13(18) 0(0) 0(12) 0.06 | 0.627 | -1.54 | 0.221 |
| Seq101   | GATGAGTTCGTAAGGCTTCAGCAC 21 323(279) 59(281) 17(118) 15(158) 0.39 | 0.481 | -2.54 | 0.084 |
| Seq102   | CAGGATGACCTTACCGGATGATCGA 21 126(153) 32(67) 20(29) 14(74) 0.54 | 0.277 | -3.29 | 0.046* |
| Seq103   | TTGATCAGCTTTGAAAGGCCAG 21 118(187) 14(26) 1(8) 1(16) 0.62 | 0.788 | -0.96 | 0.408 |
| Seq104   | AACGAGTGCATGACCTACTATTCTCTTG 24 224(262) 32(35) 41(68) 26(335) 0.61 | 0.381 | -2.03 | 0.135 |
| Seq105   | ATGTCGATGACTTACCTGTAGGA 24 138(164) 27(34) 15(38) 17(73) 0.57 | 0.299 | -2.90 | 0.062 |
| Seq106   | TAAGGATGACTTCGACATTTTGGA 21 375(350) 8(17) 6(60) 8(70) 0.44 | 0.455 | -2.47 | 0.090 |
| Seq107   | AGAAGTCTTCTACAGTTGGAACCCG 24 134(155) 51(142) 24(75) 21(93) 0.44 | 0.287 | -3.89 | 0.030* |
| Seq108   | AAAATTTCCAAATAGACGGCAGG 24 110(81) 56(75) 29(9) 1(11) 0.60 | 0.577 | -1.37 | 0.264 |
| Seq109   | AAGGGCTCATATGGTTGGGACGAG 21 132(131) 37(7) 5(15) 4(16) 0.50 | 0.343 | -2.89 | 0.063 |
| Seq110   | AAAATTTCTATAATAGACGGGACGAC 24 307(278) 73(64) 51(19) 4(28) 0.66 | 0.533 | -1.26 | 0.295 |
| Seq111   | ATCTTATATATGCACTGCTGAAC 24 169(112) 17(18) 0(0) 0(2) 0.82 | 0.762 | -0.41 | 0.707 |
| Seq112   | CTTGATATGAGTGCAGTGCAC 21 118(94) 24(20) 2(0) 7(9) 1.08 | 0.261 | 0.59 | 0.594 |
| Seq113   | TTGACCTTTAGGACATGTTTGAC 24 145(133) 13(8) 17(7) 6(10) 1.18 | 1.196 | 0.30 | 0.787 |
| Seq114   | ACGAGCTGAGCGBGGACTGG 21 170(83) 24(30) 9(7) 11(1) 3.78 | 4.838 | 1.15 | 0.333 |
| Seq115   | CAGCACGATGCGACGAGCTGG 21 951(382) 126(133) 82(42) 105(53) 1.85 | 0.647 | 2.62 | 0.079 |
| Seq116   | AGGTATCAGATGTTTACCT 21 204(5) 37(2) 340(22) 364(63) 20.13 | 14.807 | 2.58 | 0.081 |
| Seq117   | AGTGTAGCTGAACAGACAGT 22 251(101) 0(0) 2(4) 2(6) 1.11 | 1.197 | 0.15 | 0.888 |
| Seq118   | CGGCGAGTGTTGGACACGAC 21 355(350) 1800(222) 252(146) 357(121) 3.45 | 3.211 | 1.52 | 0.225 |

a: Sequencing reads (SR) were normalized to one million with the unique sequence reads of each library.
b: exp1 is the experiment reported by us. Sequencing reads in the non-infected sRNA library are bracketed, while those in the RSV-infected sRNA library are not.
c: exp2-4 are the three repeats of Du et al (2011). Sequencing reads in non-infected sRNA libraries are bracketed, while those in RSV-infected sRNA libraries are not.
d: mean of the number of infected divided by number of non-infected reads; the following columns show the standard deviation and the t-test value to test whether the mean differs significantly from 1 (equal numbers of reads).

* indicating the sRNAs that have the significant changes in RSV-infected rice (p value<0.05).
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Identification of novel miRNAs (n-miRNAs) produced from potential new precursor sequences not previously described

The precursor sequences and corresponding stem-loop structures were predicted for each of the remaining 121 sRNAs that did not match the deposited rice miRNAs or miRNA precursors. Seven sRNAs (Seq119-125) that have precursor sequences capable of forming a stem-loop structure with free energies ranging from −23.5 to −67.9 kcal/mol, and that have no more than 26 hit loci in the rice genome were identified (Table 3, Figure 4), while the other 114 sRNAs had no precursor sequence that could form a stem-loop structure (Table 3). Among these 7 sRNAs, Seq125 had 23 loci in the rice genome and the other sRNAs had only one hit locus each.

All 7 sRNAs could also be found with 1–5517 sequencing reads in the non-infected sRNA library reported by us and in 6 rice sRNA libraries reported by Du et al [38]. The relative high expression level, the stable precursor structures and their stable appearance in rice sRNA libraries indicated that the 7 sRNAs might be putative novel miRNAs (n-miRNAs).

Detection of the n-miRNAs by Northern blots and Reverse transcription (RT)-PCR

To confirm that the 7 n-miRNAs were actually produced in plants, we detected them in RSV-infected rice by Northern blots. In three independent experimental repeats, 4 of the 7 n-miRNAs (Seq119, 120, 124, and 125) could be detected in total RNAs of RSV-infected rice, but the other three n-miRNAs could not (Figure 5A). Considering the low resolving power of Northern blots, we then detected the n-miRNAs by RT-PCR. All 7 n-miRNAs were cloned and their sequences were verified (Figure S1). These results experimentally demonstrate that the 7 n-miRNAs are actually produced in RSV-infected rice, and can be recognized as novel miRNAs (n-miRNAs).

Expression analysis of the n-miRNAs in RSV-infected rice

The 7 n-miRNAs analyzed here were identified from the RSV-infected rice sRNA library. To know whether their expression levels were affected by RSV infection, we tried to compare the number of sequencing reads in our reported RSV-infected and non-infected rice sRNA libraries (experiment1, exp1 in table 3) [39], and in the three pairs of RSV-infected and non-infected rice sRNA libraries recently reported by Du et al. (three replicates, exp2–4 in table 3) [38]. The statistical analysis showed only that Seq125 was down-expressed with significant changes in RSV-infected rice, while there were no statistically significant changes for the other n-miRNAs (Table 3). This may reflect the big differences between the four experiments and indicates the fallibility of results taken only from sequencing reads.

To investigate this further, we checked the expression changes in RSV-infected and non-infected rice by real-time PCR. In three independent repeats, Seq120 and Seq125 were down-expressed in RSV-infected rice, while Seq119, Seq121, Seq122, Seq123 and 124 were up-expressed after RSV infection (Figure 5B).

Predicted targets of the n-miRNAs

The targets of n-miRNAs were predicted by using the web-based psRNA Target Server (http://plantgrn.noble.org/psRNATarget/). Considering that plant miRNA target sites are predominantly located in ORFs, we here only focused on finding target sites in coding regions. Seq119, 120, 121 and 123 had one or two targets, while Seq122, 124 and 125 had three or more targets each (Table 4). Predicted targets were involved in gene
expression, signal transduction, energy metabolism and even resistance to stress. The broad range of targets suggests that the identified n-miRNAs play roles during rice development. Moreover, two putative disease resistant proteins were targets of Seq124, suggesting a potential function during RSV infection.

Discussion

The challenge for identification of the novel miRNAs is to judge whether sRNAs satisfy the criteria of miRNAs or not. Generally, three criteria are considered: 1) the miRNA and antisense miRNA should be derived from the opposite stem arms of the precursor stem-loop structure; 2) the mismatched bases between the miRNA and anti-sense miRNA are restricted to four or fewer; and 3) the frequency of asymmetric bulges in miRNA/miRNA* duplex is restricted to less than one and the size should be less than two bases [45]. Several ancillary criteria: conservation, targets, DCL1 independence and RNA-dependent RNA polymerase (RDR) independence of sRNAs, are also considered when annotating miRNAs, but are not required [45]. Here, 23 pn-miRNAs that were newly produced from known miRNA precursors satisfy the three primary criteria, supporting the case that these 23 are true miRNAs. However, among the 7 pn-miRNAs that were potentially produced from the new precursors, Seq119 and 120 did not satisfy the second and third criteria. Seq119 had six mismatched bases and Seq120 had five mismatched bases. Moreover, Seq120 had an asymmetric bulge with four bases in the miRNA/miRNA* duplex. However, in the miRBase, not all of the known miRNAs are satisfying all of the criteria. For example, the known osa-miR419 has 14 mismatched bases; osa-miR415 has 10 mismatched bases; osa-miR416 has 7 mismatched bases; osa-miR414 has 7 mismatched bases; osa-miR139a.2 has 6 mismatched bases; osa-miR413 has 5 mismatched bases; osa-miR172a has 5 mismatched bases; osa-miR319a-5p has 5 mismatched bases. Moreover, osa-miR419 has an asymmetric bulge with 7 bases in the miRNA/miRNA* duplex; osa-miR416 has an asymmetric bulge with 6 bases in the miRNA/miRNA* duplex. Hence, it is possible that an sRNA which does not satisfy the three primary criteria is nevertheless a miRNA. Since Seq119 and 120 have precursors with a stable secondary structure, and have potential targets, we recognized them as pn-miRNAs, although they do not satisfy the second and third primary criteria.

In the pyrosequenced sRNA libraries, small interfering RNA from double stranded RNAs or decay products of large precursors are usually found at low frequency [45]. We supposed that siRNA without function might not be specifically accumulated. Hence, to further decrease the effect of siRNA on miRNA identification, we only chose the highly abundant sRNAs with a minimum of 100 sequencing reads in the library for identification, while limiting the locus numbers of sRNAs in the rice genome. This largely eliminates the side effect of siRNA on identification. However, under such conditions, some novel miRNAs present in low

![Figure 4](https://example.com/figure4.png)

**Figure 4.** sRNAs that were recognized as pn-miRNAs and the secondary structures of their precursors. The free energy of each structure is indicated as dG (kcal/mol). Sequences of sRNAs are colored green within precursors.

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Novel miRNA in Rice Stripe Virus-Infected Rice

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abundance might be excluded and it is therefore probable that some further novel miRNAs remain to be discovered.

Deep sequencing has been a powerful method for identifying miRNAs in many plants. In addition, the sequencing reads are thought to reflect the levels of expression of the sequences. Among the n-miRNAs identified here, results from sequencing reads showed that only Seq123 was down-expressed with significant changes in RSV-infected rice. However, real-time PCR results showed that all n-miRNAs had altered expression patterns in the RSV-infected rice: Seq120 and Seq125 were down-expressed in RSV-infected rice, while Seq119, Seq121, Seq122, Seq123 and 124 were up-expressed after RSV infection (Figure 5B). Some of this inconsistency may be because the time after RSV infection when the rice was sampled differed between our research and that of Du et al. We harvested the RSV-infected rice one week after inoculation when RSV infection was systemic, but before symptoms had appeared. In the work reported by Du et al, rice plants were harvested after three weeks when typical symptoms of RSV had appeared. If the expression levels of different n-miRNAs kept changing as the RSV infection developed, such inconsistency is easily explained. It would therefore be interesting to investigate further the exact responses of the n-miRNAs and their functions during the whole process of RSV infection.

In the 46 known miRNA families identified, osa-miR167, 168, 528, 172, 444, 166, 535, 164, 396 and 827 were the ten most abundant miRNA families. This is nearly completely consistent with the results of Du et al., in which osa-miR168, 156, 396, 159, 535, 166, 172, 167, 528 and 444 were the ten most abundant miRNA families, indicating stable functions for these miRNAs in RSV-infected rice. Production of new miRNAs from known precursors was also detected here. One

Table 4. Predicted targets for the identified n-miRNAs in rice.

| pn-miRNAs No. | Target Acc. | Maximum expectation | Target Description |
|---------------|------------|---------------------|--------------------|
| Seq119        | EE591526   | 1.0                 | hypothetical protein |
| Seq120        | NC_008405  | 3.0                 | Glycosyl transferase, group 1 domain containing protein |
| Seq121        | CF953168   | 3.0                 | Transposon protein  |
| Seq122        | CI534848   | 3.0                 | Transposon protein  |
| Seq123        | CI316474   | 2.0                 | Predicted protein   |
|               | CA762086   | 2.5                 | Os11g0311300 protein |
| Seq124        | CT862337   | 3.0                 | Hatching enzyme     |
| Seq125        | CX050672   | 3.0                 | Os06g0677700 protein |
| Seq124        | CI455154   | 1.0                 | Similar to SEC1-family transport protein SLY1 (AtSLY1) |
| Seq125        | CT848706   | 1.5                 | Conserved hypothetical protein |
| Seq126        | CI309171   | 1.5                 | TGF-beta receptor   |
| Seq127        | EG112217   | 1.5                 | ABC transporter permease protein |
| Seq128        | CT736378   | 1.5                 | LMBR1-like conserved region domain containing protein |
| Seq129        | CI247402   | 2.0                 | CHCH domain containing protein |
| Seq130        | CI277601   | 2.0                 | 2OG-Fe(II) oxygenase domain containing protein |
| Seq131        | CA754682   | 2.0                 | Protein of unknown function DTF516 family protein |
| Seq132        | CT855745   | 2.5                 | Conserved hypothetical protein |
| Seq133        | CI317745   | 2.5                 | Disease resistance protein family protein |
| Seq134        | AK067 489  | 2.5                 | Disease resistance protein family protein |
| Seq135        | FG954690   | 3.0                 | Mitochondrial carrier protein |
| Seq136        | CI741022   | 3.0                 | Conserved hypothetical protein |
| Seq137        | CA763859   | 3.0                 | Mitochondrial carrier protein |
| Seq138        | CR281443   | 3.0                 | Similar to ATP sulfurylase |

Figure 5. Transcription analyses of n-miRNAs and their changed expression in RSV-infected rice. A, 4 of 7 pn-miRNAs were detectable in Northern blot analysis. B represents the expression analysis of 7 n-miRNAs in RSV-infected rice and non-infected rice through real-time PCR. Expression levels of the 7 n-miRNAs in non-infected rice were assigned as 1.0.

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or more new miRNAs were identified from each of 19 known miRNAs. Among these 19 known miRNA precursors, these are the first reports of a second mature miRNA from osa-MIR169a, 187a, 183b, 180a, 206g, 807b, 806, 809b, 812a, 1439, 812k and 2123c. Most of the newly produced miRNAs have predicted targets, indicating their potential function in rice (Table S3).

Moreover, sequencing reads of Seq99, 102, 104, 105, 107 and 116 showed a stable change between RSV-infected and non-infected rice sRNA libraries, indicating a potential role in RSV-induced rice disease.

RSV-induced rice disease is serious in China, Japan and Korea. Virus variation and the functions of the RSV-encoded proteins have been well studied in recent years [40,46,47,49,50,51,52]. Now we report 7 novel miRNAs related to RSV infection. This is the first report of novel miRNAs related to viral infection in rice. miRNAs play key roles during rice development and contribute to the plant defense against several bacterial, fungal or viral diseases. We are not sure whether the novel miRNAs identified here play some roles in plant defense against viral infection. However, it was found that Seq124 had two potential targets that were putative disease resistance proteins, indicating the possibility that these novel miRNAs function in the rice plant defense against RSV. We intend to do further work to clarify the roles of these novel miRNAs during RSV infection.

Supporting Information

Figure S1 Alignments of each n-miRNA and its cloned sequence. Sequence 1 represents the n-miRNA sequence; Sequence 2 represents the cloned sequence in pGEM-T vector for sequencing. Sequence 3 represents the expected sequence for cloning. (TIF)

Table S1 sRNAs used for analysis here. (XLS)

Table S2 Primers used for RT-PCR and real-time PCR (DOC)

Table S3 Predicted targets for the identified miRNAs newly produced from known miRNA precursors. (DOC)

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Author Contributions

Conceived and designed the experiments: FY JC. Performed the experiments: WG GW FY YL HZ. Analyzed the data: FY LL HC JC. Wrote the paper: WG GW FY JC.

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