

# Identification of miniature inverted-repeat transposable elements (MITEs) and biogenesis of their siRNAs in the Solanaceae: New functional implications for MITEs

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Small RNAs regulate the genome by guiding transcriptional and post-transcriptional silencing machinery to specific target sequences, including genes and transposable elements (TEs). Although miniature inverted-repeat transposable elements (MITEs) are closely associated with euchromatic genes, the broader functional impact of these short TE insertions in genes is largely unknown. We identified 22 families of MITEs in the Solanaceae (*MiS1-MiS22*) and found abundant *MiS* insertions in Solanaceae genomic DNA and expressed sequence tags (EST). Several Solanaceae MITEs generate genome changes that potentially affect gene function and regulation, most notably, a *MiS* insertion that provides a functionally indispensable alternative exon in the tobacco mosaic virus *N* resistance gene. We show that MITEs generate small RNAs that are primarily 24 nt in length, as detected by Northern blot hybridization and by sequencing small RNAs of *Solanum demissum*, *Nicotiana glutinosa*, and *Nicotiana benthamiana*. Additionally, we show that stable RNAi lines silencing *DICER-LIKE3* (*DCL3*) in tobacco and *RNA-dependent RNA polymerase 2* (*RDR2*) in potato cause a reduction in 24-nt MITE siRNAs, suggesting that, as in *Arabidopsis*, TE-derived siRNA biogenesis is *DCL3* and *RDR2* dependent. We provide evidence that *DICER-LIKE4* (*DCL4*) may also play a role in MITE siRNA generation in the Solanaceae.

[Supplemental material is available online at [www.genome.org](http://www.genome.org).]

Small interfering RNA (siRNA) and microRNA (miRNA) play fundamental roles in eukaryotic gene regulation, antiviral defense, genome organization, and maintenance of genome integrity. These small RNAs are processed from RNA precursors by type III RNase enzymes, referred to as DICER or DICER-LIKE proteins (DCL), and incorporated into RNA induced silencing complexes (RISC) to mediate post-transcriptional gene silencing (PTGS) by translational arrest or mRNA cleavage, or into RNA-induced transcriptional silencing complexes (RITS) to mediate transcriptional gene silencing (TGS) by repressive DNA and histone modifications (Baulcombe 2006; Li and Ding 2006; Ding and Voinnet 2007; Henderson and Jacobsen 2007; Slotkin and Martienssen 2007). In *Arabidopsis*, small RNA biogenesis enzymes are encoded by gene families in which the individual members have specialized functions in parallel pathways. *DCL1* and hyponastic leaves 1 (*HYL1*) produce miRNAs that have a major role in developmental regulation (Carrington and Ambros 2003; Chen 2005; Poethig et al. 2006). *DCL4*, *DCL2*, and *DCL3* act hierarchically in response to RNA virus infection by generating viral siRNAs (Ding

and Voinnet 2007). *DCL4*, RNA-dependent RNA polymerase 6 (*RDR6*), and suppressor of gene silencing 3 (*SGS3*) are involved in *trans*-acting siRNA (ta-siRNA) accumulation and regulate juvenile-adult transition (Poethig et al. 2006). *DCL2*, *DCL1*, *RDR6*, and Nuclear RNA polymerase D 1A (*NRPD1A*) act on natural antisense transcripts and produce nat-siRNA in response to abiotic stress (Borsani et al. 2005). More recently, *RDR2*, *NRPD1A*, and *SNF2*, in conjunction with *DCL4*, have been implicated in non-cell-autonomous silencing (Dunoyer et al. 2007; Smith et al. 2007).

Small RNA pathways also regulate the activity of transposable elements—both DNA transposons and retrotransposons—via a class of transposable element (TE)-derived 24-nt siRNAs that guide transcriptional gene silencing (TGS), generated by a pathway involving *DCL3*, *RDR2*, *NRPD1A*, and *ARGONAUTE 4* (*AGO4*) (Henderson and Jacobsen 2007). Most TEs are concentrated in highly repetitive regions near centromeres and telomeres, where siRNA-dependent chromatin modifications establish vast stretches of heterochromatin that dominate the genomic makeup of many organisms (Bennetzen and Kellogg 1997). However, isolated examples have also uncovered the co-option of TE siRNAs in controlling the expression of genes bearing TE insertions. For example, the expression of *FLC* and *FWA*, which govern flowering time, is regulated by siRNA-guided TGS targeting of a *Mutator*-family TE and directly repeated SINE elements

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inserted in the *FLC* intron (Liu et al. 2004) and *FWA* promoter (Chan et al. 2004), respectively. It is thought that TE mobility may provide the evolutionary capacity to alter patterns of gene regulation via the insertion of TEs within “gene islands.”

A particular class of truncated DNA transposons, miniature inverted-repeat transposable elements (MITEs), may play a role in the co-option of TE-derived siRNAs in gene regulation. MITEs, which are generally <600 bp, lack open reading frames, and thus their mobility depends on the activity *in trans* of transposases encoded by cognate full-length autonomous transposons (Jiang et al. 2003, 2004b). The full-length DNA transposons are hypothesized to be the evolutionary progenitors of MITEs, based on sequence conservation between miniature inverted-repeat transposable elements (MITEs) and autonomous transposon terminal inverted repeats (TIRs) and target site duplications (TSDs). MITEs have amplified to a high copy number in several plant genomes, as in rice, which contains more than 90,000 MITEs grouped into approximately 100 different families (Feschotte et al. 2003; Jiang et al. 2004b; Juretic et al. 2004). In the rice and *Arabidopsis* genomes, the majority of MITEs are inserted in the euchromatin arms of chromosomes (Feng et al. 2002; Santiago et al. 2002; Wright et al. 2003), rather than in the heterochromatin. This observed close physical association between MITEs and plant genes (Wessler 1998; Mao et al. 2000) has provoked the hypothesis that MITEs play a role in gene regulation, as predicted of “controlling elements” (McClintock 1951). However, prior to the recent discovery of small RNA pathways, a plausible explanation for mechanisms of MITE-based gene regulation, and for functional roles of TEs in general, has been incomplete. The recent proposal that several families of miRNAs in humans, *Arabidopsis*, and rice are derived from MITEs (Piriyapongsa and Jordan 2007, 2008; Piriyapongsa et al. 2007) gives further, tantalizing evidence that MITEs may serve a major evolutionary role in co-opting TE silencing pathways for gene regulation, not only in plants, but also across the eukaryotic spectrum. However, investigation of biogenesis of small RNAs derived from MITEs has not been thoroughly studied in plants, and the roles of MITEs in general, particularly when associated with plant genes, remain largely unexplored.

Research in the Solanaceae family, which comprises more than 3000 species including the important crop plants tomato (*Solanum lycopersicum*), potato (*Solanum tuberosum*), eggplant (*Solanum melongena*), pepper (*Capsicum annuum*), and tobacco (*Nicotiana tabacum*), has produced comparative genomic resources—including the sequencing of multiple alleles of several resistance gene loci (Chang et al. 2002; Song et al. 2003; Schornack et al. 2004; Huang et al. 2005; Kuang et al. 2005) that facilitate identification and analysis of TEs. Using comparative sequence analysis and the related empty site (RESite) method (Le et al. 2000), we examined homeologous *R-gene* regions from different haplotypes of the same or closely related Solanaceae species to identify 22 MITE families. Using Northern blot hybridization analysis, we show that all MITE families tested generate small RNAs in Solanaceae species, and we identified specific MITE-derived siRNAs by small RNA sequencing. We determined that MITE small RNA are generated by a pathway similar to that required for TE-derived siRNA biogenesis in *Arabidopsis* by silencing each of the four *DICER-LIKE* (*DCL*) homologs in *N. tabacum* and *RDR2* in *S. tuberosum*, demonstrating that silencing of *DCL3* and *RDR2* causes a reduction in MITE-derived small RNAs. However, in contrast to *Arabidopsis*, our findings suggest a role for *DCL4* in MITE small RNA biosynthesis in the Solanaceae. These results reveal

the presence of MITE-derived small RNAs in the Solanaceae and provide a foundation for a systematic and in-depth analysis of the roles of MITEs in small RNA-mediated gene regulation and evolution in Solanaceae.

## Results

### Identification and analysis of MITE families in Solanaceae

#### Identification and classification of MITEs

We identified 22 families of MITEs in Solanaceae (*MiS1–MiS22*) within six syntenic tomato, potato, and tobacco *R-gene* regions based on shared structural similarities and the analysis of related empty sites (RESite) (Le et al. 2000). Fifteen of these *MiS* families are novel, while seven have been described previously (Table 1). Based on their similarity to the characteristic features of known MITEs, including TIR and TSD sequence composition, *MiS2–MiS4*, *MiS6–MiS11*, *MiS12–MiS17*, and *MiS18–MiS19* were grouped into *hAT*-like, *Mutator*-like, *Stowaway*-like, and *Tourist*-like superfamilies, respectively (Table 1). *MiS1*, *MiS5*, and *MiS20–22* share no similarity with any known MITE superfamily. Eight *MiS* families have members that exceed the length limitations (<600 bp) that arbitrarily define MITEs, but retain MITE properties, including high copy number, lack of an encoded transposase, association with genes, and conserved structures (TIRs, TSDs); hence, we identified these families as *MiS* and included them in our analysis. We extended our Solanaceae MITE sequence data set using the representative member of each *MiS* family as a query in BLASTN searches of Solanaceae sequences in the GenBank nonredundant database and retrieved *MiS* sequences with *E*-values <  $e^{-5}$ , the minimal stringency necessary to detect a 30-nt stretch of identical sequence within an 80-nt MITE. The full-length sequence of the original *MiS* data set retrieved from GenBank is provided in Supplemental Data S1.1. Consistent with previous reports on MITE association with genes, we found several MITE insertions located near genes and within introns, exons, 5'- and 3'-untranslated regions (UTRs), and promoters. The location of 198 members of *MiS1–22* and their positions within a gene or location near a gene is provided in Supplemental Table S1.1. Notably, Solanaceae MITEs are located upstream and downstream of genes, within promoters, 5'- and 3'-untranslated regions, introns, and exons.

#### Estimation of genome-wide copy number

MITEs often exist in high copy number, and we hypothesized that Solanaceae genomes would contain abundant *MiS* sequences. To assess the abundance and amplification of *MiS*, we used the sequence of a representative *MiS* from each family *MiS(X)-I* (Supplemental Data S1.1) as queries in BLASTN searches of tomato and potato BAC-end ([http://www.tigr.org/tdb/sol/sol\\_ma\\_blast.shtml](http://www.tigr.org/tdb/sol/sol_ma_blast.shtml)) and tobacco genomic DNA (NCBI *Nicotiana tabacum* gss) databases and extrapolated the number of hits retrieved to estimate genome-wide copy numbers as described in Supplemental Table S1.2. An extended data set of *N. tabacum* *MiS1*, *MiS5*, and *MiS17* sequences is provided in Supplemental Data S1.2. Several *MiS* families from the *hAT*-like (*MiS2*, *MiS4*), *Stowaway*-like (*MiS12*, *MiS13*, *MiS16*, *MiS17*), and *Tourist*-like superfamilies (*MiS18*, *MiS19*) are represented in all three species, ranging from an estimated 100 to several thousand copies, which suggests an origin that predates the divergence of these species. In contrast, the estimated copy numbers of the

**Table 1. Structural features of MITE families in Solanaceae**

Family	Superfamily	Size (bp)	TSD	TIR (size, in bp)	Reference
<i>MiS1</i>		~200	2 bp	TGTTAY (6)	This study
<i>MiS2</i>	<i>hAT</i> -like	317–1404	8 bp	CAGDGGCGA (9)	This study
<i>MiS3</i>	<i>hAT</i> -like	165–177	8 bp	CAGRGGCGGA (10)	<i>NtToya1</i> (Schenke et al. 2003)
<i>MiS4</i>	<i>hAT</i> -like	798–972	8 bp	CAGGGGTGTA (12)	(Guyot et al. 2005)
<i>MiS5</i>		~355	5 bp	TGTNAAA (7)	This study
<i>MiS6</i>	<i>Mutator</i> -like	326–935	9 bp, A/T rich	GGGAAAAGG (150–250)	This study
<i>MiS7</i>	<i>Mutator</i> -like	980–1200	9 bp, A/T rich	GGCTTAAGTC (84–127)	This study
<i>MiS8</i>	<i>Mutator</i> -like	300–350	9 bp, A/T rich	GGGATAANGG (~150)	This study
<i>MiS9</i>	<i>Mutator</i> -like	332–1320	9 bp, A/T rich	GGGAAAAGG (100–180)	This study
<i>MiS10</i>	<i>Mutator</i> -like	855–1111	9 bp, A/T rich	CGGAAAAGG (260–410)	<i>Sol3</i> (Oosumi et al. 1995)
<i>MiS11</i>	<i>Mutator</i> -like	~300	9 bp, A/T rich	GGGATAATGC (129–137)	<i>TAPIR</i> (Mao et al. 2001)
<i>MiS12</i>	<i>Stowaway</i> -like	~260	TA	CTYCYTCYGT (50)	This study
<i>MiS13</i>	<i>Stowaway</i> -like	209–250	TA	CTCCCTCCGT (27)	This study
<i>MiS14</i>	<i>Stowaway</i> -like	246–310	TA	CTCCCTCNGT (26)	This study
<i>MiS15</i>	<i>Stowaway</i> -like	~300	TA	CTCCCTCCGT (32)	This study
<i>MiS16</i>	<i>Stowaway</i> -like	250–336	TA	CTYYCTYGT (50)	This study
<i>MiS17</i>	<i>Stowaway</i> -like	~250	TA	CTCCCTYCGTT (30)	<i>NtStowaway101</i> (Schenke et al. 2003)
<i>MiS18</i>	<i>Tourist</i> -like	436–1113	3 bp	GGGGTYGTTT (22)	<i>Alien</i> (Pozueta-Romero et al. 1996)
<i>MiS19</i>	<i>Tourist</i> -like	190–240	3 bp	GGGGTYGTTT (13)	This study
<i>MiS20</i>		235–508	12 bp	GKGGCGTAGC (10)	This study
<i>MiS21</i>		717–797	10 bp	GAGTTAATAC (340)	<i>Foldback</i> (Rebatchouk and Narita 1997)
<i>MiS22</i>		1.6–3.4 kb	5 bp	TGTMAC (6)	This study

(TSD) Target site duplication; (TIR) terminal inverted repeat.

*MiS1* family are much higher in tobacco (>1000) than in tomato and potato (<60), while most *Mutator*-like *MiS* (*MiS6*–*MiS11*) have higher estimated copy numbers in tomato and potato (approximately 400–3200) than in tobacco (0–32), suggesting that amplifications have occurred after the divergence of *Solanum* and *Nicotiana*. The most abundant family was *MiS22*, which retrieved 2000 and 1874 hits, respectively, in the tomato and potato BAC-end sequence databases, extrapolating to an estimated 3516 and 9802 copies in the respective genomes. While we expect errors in our extrapolation, owing to incomplete genome sequence data sets for all three species and the bias of the MITE query sequences to identify more *MiS* hits in their cognate genomes as compared to diverged genomes, our results nonetheless indicate that many MITE families have amplified to high copy numbers in Solanaceae genomes, a property consistent with other genomes.

### Structural, functional, and evolutionary implications of MITE insertions

In addition to their characteristic TIR and TSD sequence composition, and high copy number, MITEs have features with evolutionary and functional implications. For example, MITEs can capture and move gene sequences to new locations; MITE genetic and structural properties implicate cognate autonomous elements as both progenitors and activators of MITE transposition, and MITEs are often associated with genes. While analyzing *MiS* insertions, we discovered three families of elements—*MiS5*, *MiS2*, and *MiS1*—that are of particular interest because features of some family members provide insight into their evolutionary and functional properties.

#### Tandem MITE complex

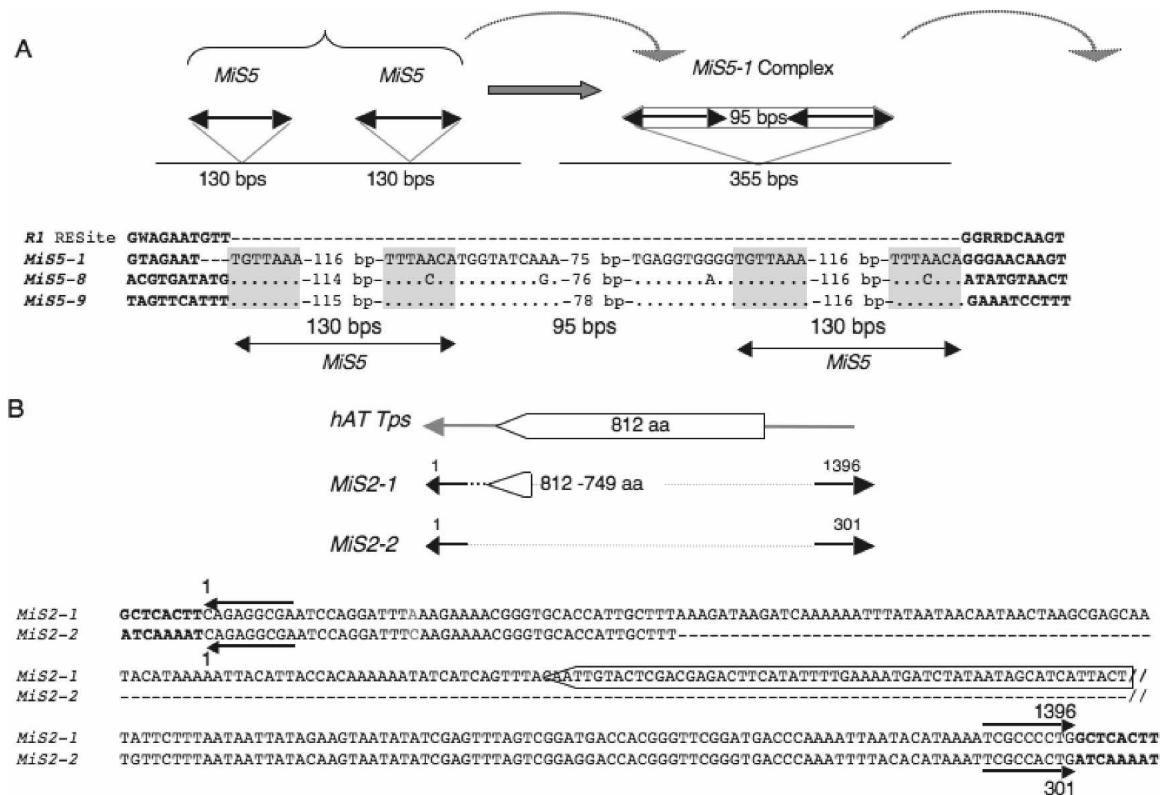
Analysis of *MiS5-1*, a 355-bp insertion in an *R1* resistance gene homolog (*R1b-11*, AC149266, 1364–1718 bp) in the *Solanum demissum* Chromosome 5 *R1*-gene cluster, revealed that it is a complex comprising two tandem, nearly identical 130-bp *MiS5* elements (with identical TIRs and two mismatches overall) separated by 95 bp of intervening sequence (Fig. 1A). Additional *MiS5*

complexes were identified in *S. demissum* and other *Solanum* species (Supplemental Data S1.1). For example, *MiS5-8*, a 355-bp complex (*S. lycopersicum*, AF411809, 126,063–126,418 bp), and *MiS5-9*, a 357-bp complex (*S. tuberosum*, TA45582\_4113, 250–606 bp), share 83% and 91% overall identity with *MiS5-1*, respectively, but have unrelated sequences flanking each insertion. Analyses of 51 RESites for *MiS5-1* identified among *S. demissum* *R1* homologs lacking *MiS5-1* suggest that *MiS5* does not generate a TSD but rather a 3-bp target-site deletion. We found no insertions or deletions within the 51 RESites, while the *R1b-11* homolog containing the complex *MiS5-1* insertion showed a 3-bp deletion at the target site (Fig. 1A).

Numerous *MiS5* (~130 bp) sequences were also found in Solanaceae BAC-end, GenBank nonredundant databases, and EST databases (Supplemental Tables S1 and S2.1). Retention of nucleotide identity in the intervening sequences of distinct *MiS5* complexes indicates that tandem MITEs have the capacity to be excised and transpose as a complex, thereby providing a mechanism that “captures” and carries intervening sequence to a new genomic location.

#### Autonomous transposon progenitors of MITEs

MITEs are activated by and hypothesized to be derived from full-length autonomous DNA transposons. Consistent with this hypothesis, we identified two highly homologous MITEs in *S. demissum*, *MiS2-1* (AC149265, 50,964–52,375) and *MiS2-2* (AC149488, 68,757–69,073), which share ~85% overall nucleotide identity and contain nearly identical *hAT* superfamily TIR sequences. *MiS2-1* is 1396 bp long and contains a 192-bp sequence homologous to a putative *hAT* transposase (AAT39314, 749–812 amino acids), while *MiS2-2* is only 301 bp long and has no predicted transposase coding sequence (Fig. 1B; see Supplemental Data S1.1 for sequences of *MiS2-1* and *MiS2-2*). It appears that *MiS2-1* and *MiS2-2* are different derivatives of a progenitor *hAT* superfamily transposon. We also identified a candidate progenitor of the *Mutator*-like *MiS10* family, an autonomous *Mutator*-like TE in *S. demissum* (AC149302, 44,107–50,193), which has a



**Figure 1.** Structures of *MiS5* and *MiS2*. (A) Several *MiS5* members are found as complex with an intervening sequence. A model is presented in which tandem insertions of the same *MiS5* element generate a *MiS* complex that captures and transposes intervening sequence. (Bold black lines flanked by arrowheads, TIRs) *MiS*. The sequences of the *MiS5-1* RESite in *S. demissum* (*R1*; AC149265.1) and three *MiS5* complexes, *MiS5-1* in *S. demissum* (AC149266, 1364–1718), *MiS5-8* in *S. lycopersicum* (AF411809, 126,063–126,418), and *MiS5-9* in *S. tuberosum* (TA45582\_4113, 250–606), are shown. (Dots) Nucleotide identity; (dashes in *R1*) the RESite lacking *MiS5* insertion; (dashes in *MiS5-1*) a 3-bp deletion at its insertion site; (gray) TIRs; (bold) flanking sequence. (B) *MiS2-1* (1396 bp) and *MiS2-2* (301 bp) are *hAT*-type elements in GenBank accessions AC149265, 50,964–52,375, and AC149488, 68,757–69,073, respectively. (Open arrowhead) The *MiS2-1* sequence is predicted to encode a truncated product with similarity to the C terminus (749–812 amino acids) of candidate *S. demissum* *hAT* transposase (AAT39314). Conserved sequences between *MiS2-1* and *MiS2-2* are shown aligned. (Horizontal arrows above and below the aligned sequences) The 9-bp *MiS2* TIRs. (Bold) The 8-bp TSDs generated by the *MiS2* insertions. The *MiS2-1* sequence enclosed in the open arrow is predicted to encode amino acids 794–812 of the candidate *S. demissum* *hAT* transposase.

characteristic 9-bp A/T-rich TSD and ~350-bp *MiS10*-like TIRs (Supplemental Data S1.1) and encodes a predicted MuDR-like transposase.

#### A MITE serves as an alternative exon

*MiS1-1*, a 186-bp insertion, was identified in intron 3 of the tobacco mosaic virus (TMV) resistance gene, *N* (U15605), by sequence comparison between the *N* gene of *Nicotiana glutinosa* and the *NH* gene (A535010) of *N. tabacum*, a candidate *N* ortholog, with a predicted product sharing 88% amino acid identity with *N* (Fig. 2A; Stange et al. 2004). The *MiS1* inserted sequence was used as a BLASTN query of GenBank, leading to the discovery of a second sequence, 87 bp in length, in the third intron of the *N* gene, as well as a third *MiS1* sequence, 193 bp in length, located 387 bp upstream of the *Nicotiana sylvestris* *RAN-1* gene (AY563050). These three elements exhibit 80%–85% nucleotide identity (data not shown). They share the 6-bp terminal inverted-repeat (IR) sequences but are flanked by unrelated sequences, indicating that the elements are members of a novel MITE family, which we have named *MiS1* with members *MiS1-1*, -2, and -3, corresponding to the 186-, 87-, and 193-bp elements, respectively (Table 1; Supplemental Data S1.1).

*N* is transcribed into two mRNA isoforms ( $N_S$  and  $N_L$ ), and

expression of  $N_L$  containing a 70-bp alternative exon is required for full resistance to TMV. Approximately 8 h following TMV infection, the relative abundance of the alternatively spliced  $N_L$  isoform increases (Whitham et al. 1994; Dinesh-Kumar and Baker 2000). *MiS1-1* provides the 70-bp alternative exon for the  $N_L$  mRNA isoform, and the alternative splice sites are located within the MITE sequence (Fig. 2A). The *MiS1* element is not found in other closely related *N* homologs in GenBank (AF211528, AY438027, AY438028, AY535010, AC151803, AJ300266, AC154033), and the deduced amino acid sequence encoded by the *N* alternative exon has no significant similarity to any known protein. Based on BLASTN searches of the tobacco genomic sequence database (NCBI *Nicotiana tabacum* gss), the *MiS1* family is present in high copy numbers in the tobacco genome (Supplemental Table S1.2). We conclude that the *N* alternative exon was generated through the insertion of *MiS1-1* into intron 3 of the *N* gene and provides an example of a MITE insertion contributing coding sequence to a functional alternative-splicing product.

#### MITES are transcribed

Our original *MiS* data set revealed the close association of members of each MITE family with genes. We hypothesized that, in



([http://tigrblast.tigr.org/euk-blast/plantta\\_blast.cgi](http://tigrblast.tigr.org/euk-blast/plantta_blast.cgi); Childs et al. 2007) using a representative *MiS* sequence (*MiSX-1*) from each family (Supplemental Data S1.1) as a query. We identified approximately 666 ESTs containing *MiS* insertions with *E*-values of  $<e-5$  and found that each *MiS* family is represented in at least one of the three EST databases (Supplemental Fig. S1; Supplemental Tables S1.2 and S2.1). Generally, there was a positive correlation between the number of hits detected in EST databases and the estimated copy number in genomic DNA (Supplemental Fig. S1; Supplemental Table S1.2), for example, high copy number *MiS* families, such as *MiS3*, *MiS17*, and *MiS18* in tobacco, and *MiS13*, *MiS17*, and *MiS22* in tomato and potato, also retrieved a high number of EST hits. As revealed by analysis of genomic sequence, *MiS22* yielded the most hits in EST databases, being present in 101 sequences (0.05% of the total) in the potato EST database.

In 104 of the 666 TAs containing a Solanaceae MITE-related sequence, there was sufficient information to resolve the position of the insertion within the predicted gene. Of these, 46 of the insertions were located in the 3'-UTR, 11 in the 5'-UTR, one serving as an alternative exon located in an intron (in the *N* gene, discussed above), and 46 in coding regions, disrupting ORFs (Supplemental Table S2.1). Some insertions of a MITE family were observed in both "+" and "-" (complementary) orientations. Interestingly, five of the 10 *MiS1* elements found in the EST/TA databases were inserted in the 3'-UTR of the predicted corresponding transcripts. The 3'-UTRs of these transcripts are terminated within the *MiS1* sequences, before the TIR, regardless of the orientation of the insertion, raising the possibility that *MiS1* family members may encode polyadenylation signals for transcript termination and poly(A) addition. We identified candidate polyadenylation signal sequences, near upstream elements (NUE), far upstream elements (FUE), and cleavage elements (CE), FUE.1 and NUE.3 (AATAAA) in 93% and CE.1 (TAATTA) (Shen et al. 2008) in 11% of the 76 *MiS1* tobacco sequences surveyed, whereas the FUE.1/NUE.3 and CE.1 signal sequences were identified in only 33% and 7% of the 1007 surveyed *MiS17* tobacco sequences. A sequence alignment of two of *N. benthamiana* transcript assemblies containing *MiS1* elements in complementary orientations in their 3'-UTRs is shown in Figure 2B and described further below.

### Detection and cloning of *MiS* small RNA

Recently, transposon-derived small RNA has been shown to play an important role in silencing retrotransposon and *Mutator*-type elements via TGS and PTGS mechanisms (Xie et al. 2004; Slotkin et al. 2005; Sridhar et al. 2007). Our discovery that Solanaceae genomes carry candidate autonomous transposon progenitors and high copy numbers of MITEs led us to hypothesize that *MiS* small RNAs may play a role in transcriptional silencing of MITEs as described for transposon regulation in *Arabidopsis* and other species. Our finding that Solanaceae MITEs are transcribed and co-transcribed with adjacent genes, presumably by RNA polymerase II (Pol II), and may form predicted double-stranded RNA (dsRNA) or single-stranded hairpin precursors of small RNAs, suggested that post-transcriptional silencing may also play a role in MITE regulation. A putative dsRNA structure formed from "complementary insertions" of *MiS1* is depicted in Figure 2B. *MiS1* family members were found in sense and antisense orientations in the 3'-UTRs of two *N. benthamiana* EST transcript assemblies, TA10655\_4100 ([http://plantta.tigr.org/cgi-bin/plantta\\_report.pl?ta=TA10655\\_4100](http://plantta.tigr.org/cgi-bin/plantta_report.pl?ta=TA10655_4100)) and TC10011 ([\[dcfci.harvard.edu/tgi/cgi-bin/tgi/tc\\\_report.pl?tc=TC10011&species=n\\\_benthamiana\]\(http://compbio.dcfci.harvard.edu/tgi/cgi-bin/tgi/tc\_report.pl?tc=TC10011&species=n\_benthamiana\)\) \(Fig. 2B; Supplemental Table S2.1\), which are predicted to encode a hypothetical protein and a product homologous to WRKY transcription factor 22, respectively. If transcripts bearing these complementary \*MiS\* sequences are co-expressed, we hypothesize that stretches of dsRNA form through pairing of 3'-UTR \*MiS1\* sequences and may functionally mimic natural antisense transcripts precursors of regulatory small RNAs \(Borsani et al. 2005\).](http://compbio.</a></p>
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An example of a hairpin-like structure formed from an *MiS9*-containing transcript (DN938187) with a long TIR (~100 nt or more) is shown in Figure 2C.

### *MiS* small RNA detection

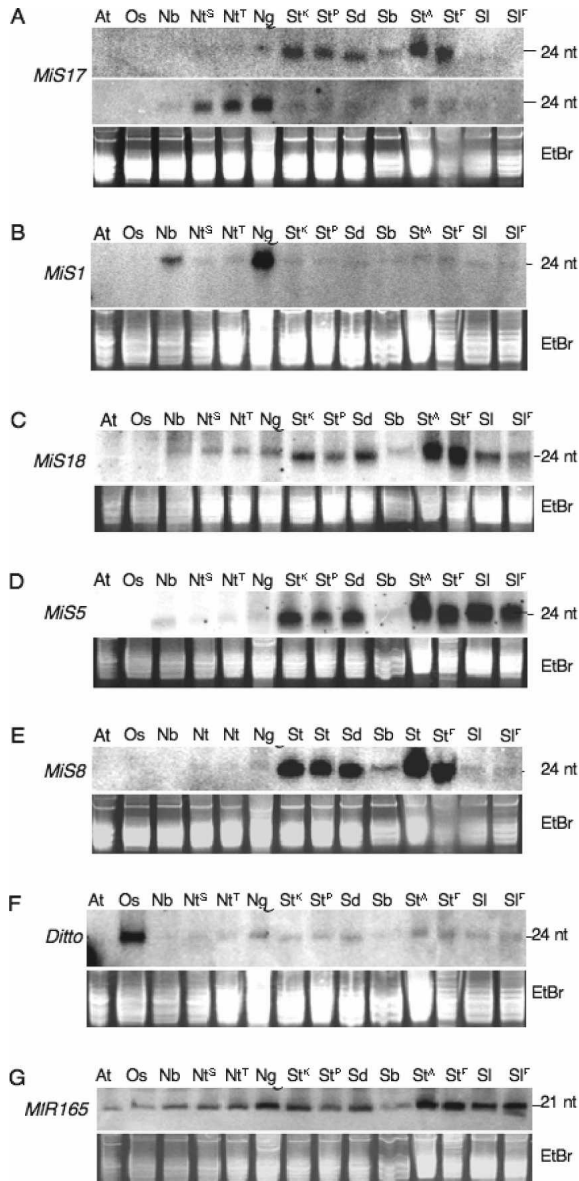
As a first step to test our hypothesis that Solanaceae MITEs generate small regulatory RNAs, we performed Northern blot hybridization analysis using probes for specific *MiS* families to analyze the low-molecular-weight RNA fractions isolated from *Nicotiana benthamiana*, *N. tabacum* SR1, *N. tabacum* SR1:NN, *N. glutinosa*, *S. demissum*, *S. tuberosum* Kennebec, *S. lycopersicum*, *Oryza sativa*, and *Arabidopsis thaliana* (Fig. 3; probes described in Supplemental Table S3.1). Each *MiS* probe detected small RNAs in several Solanaceae species, with 24-nt RNA constituting the predominant size class (Fig. 3A–E).

The *MiS* probes detected small RNAs most strongly in their species of origin and in closely related species. For example, the *MiS17* probe, amplified from a pool of *S. demissum* and *S. tuberosum* genomic DNA, only detected a strong signal in *Solanum* species (Fig. 3A, top panel), while an *MiS17* probe amplified from a pool of *N. benthamiana*, *N. tabacum*, and *N. glutinosa* genomic DNA only detected a strong signal in *Nicotiana* species (Fig. 3A, middle panel). Similarly, the *MiS1* probe, amplified from *N. glutinosa* genomic DNA, displayed a strong signal in *N. glutinosa* and in *N. benthamiana*, but a much weaker signal in *N. tabacum* and in *Solanum* species (Fig. 3B), while the *MiS18*, *MiS5*, and *MiS8* probes, amplified from *S. demissum* genomic DNA, only displayed strong signals in *Solanum* species (Fig. 3C–E, respectively). None of the *MiS* probes detected small RNA in *Arabidopsis* or rice. As a control, a probe made from the rice *Tourist*-superfamily MITE *Ditto* displayed a much stronger hybridization signal in rice than in other species, and a probe for the conserved miRNA MIR165 (Supplemental Table S3.2; Reinhart et al. 2002) displayed a strong signal in all species (Fig. 3, F and G, respectively).

### *MiS* small RNA sequence

We hypothesized that the ~24-nt *MiS*-related small RNAs detected in Northern blots were generated by a NRPD1A–RDR2–DCL3 pathway, similar to that previously described for biogenesis of 24-nt repeat-associated siRNA for transcriptional silencing of transposons in *Arabidopsis*. Based on our predictions of paired transcripts and hairpin structures, we postulated that small RNA pathways associated with post-transcriptional gene silencing could also produce *MiS* small RNAs.

The genomic origin, mechanism of biogenesis, and potential function of small RNAs can be inferred from their sequence, length, and 5'-terminal base composition. To test our hypotheses on the origin, biogenesis, and potential function of MITE small RNAs detected by Northern blot hybridization analyses, we cloned and sequenced small RNAs from *S. demissum*, *N. benthamiana*, and *N. glutinosa* to obtain a total of 4421, 1408, and 1806



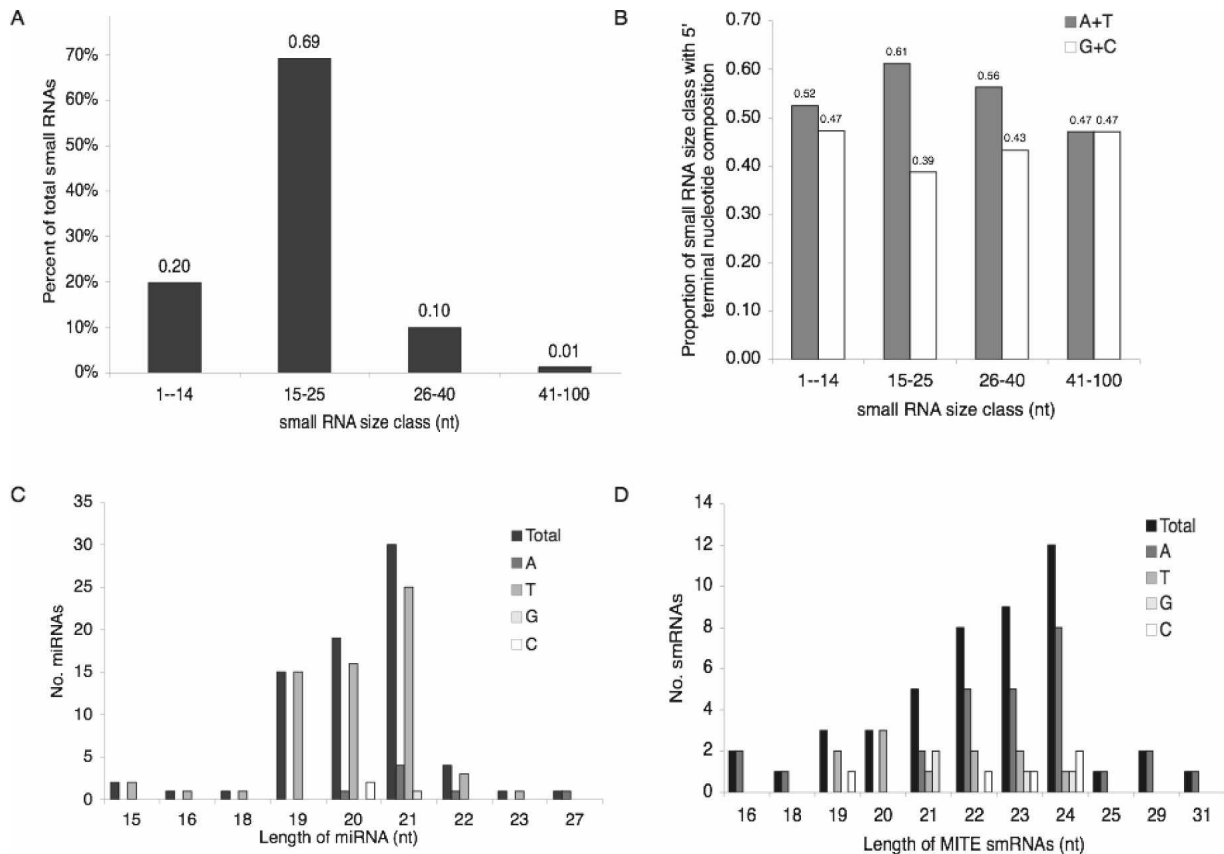
**Figure 3.** *MiS* small RNAs detected in *Solanum* and *Nicotiana* species by Northern blot hybridization. Forty micrograms of small RNA-enriched from total RNA isolated from Solanaceae species was size separated using a 20% acrylamide gel, blotted, and hybridized with *MiS*-derived probes (see Supplemental Table S3.1 for a description of probes). (A) Hybridization probe for (upper panel) *Solanum-MiS17* probe; (lower panel) *Nicotiana-MiS17*; (B) *Nicotiana-MiS1* probe; (C) *Solanum-MiS15* probe; (D) *Solanum-MiS5* probe; (E) *Solanum-MiS8* probe; (F) *Os-Ditto* probe; (G) *MIR165*. An EtBr stain is shown for each gene as a sample loading control. (At) *A. thaliana*, Col-0; (Os) *O. sativa* Nipponbare; (Nb) *N. benthamiana*; (Nt<sup>S</sup>) *N. tabacum*, SR1; (Nt<sup>G34</sup>) *N. tabacum*, SR1:NN; (Ng) *N. glutinosa*; (St<sup>K</sup>) *S. tuberosum*, Kennebec; (St<sup>P</sup>) *S. tuberosum*, Pentland Ivory; (Sd) *S. demissum*; (Sb) *S. bulbocastanum*; (St<sup>L</sup>) *S. tuberosum*, Katahdin leaves; (St<sup>T</sup>) *S. tuberosum*, Katahdin flowers; (Sl<sup>F</sup>) *S. lycopersicum*, VF36 leaves; (Sl<sup>F</sup>) *S. lycopersicum*, VF36 flowers.

small RNA reads of good quality, respectively (Supplemental Data S2.1, S2.2, and S2.3). Sequenced *S. demissum* small RNAs between 15 and 25 nt account for 69.2% of the total *S. demissum* data set; those below 15 nt comprise the second largest group and account for 19.8%, and small RNAs between 26 and 40 nt account for

9.9% (Fig. 4A). 5'-Terminal nucleotide composition analysis showed an enrichment for 5' A and T residues in the 15- to 25-nt size class of small RNAs, with this enrichment diminishing in the larger and smaller size classes (Fig. 4B).

We attempted to establish the identity of each nonredundant small RNA between 15 and 40 nt. To identify known miRNAs in this data set, we performed BLASTN searches of the miRbase (<http://microrna.sanger.ac.uk/sequences/index.shtml>), which identified 38 unique candidate miRNA sequences corresponding to 74 reads within the *S. demissum* small RNA data set and 30 unique candidate miRNA sequences within the combined *N. benthamiana* and *N. glutinosa* data set (Supplemental Table S2.2; Griffiths-Jones 2004; Griffiths-Jones et al. 2006). Analysis of candidate miRNA sequences confirmed the preference of a 5'-terminal T residue for 35 of 38 *S. demissum* and 27 of 30 *Nicotiana* candidate miRNAs described for AGO1-associated miRNAs (the *S. demissum* results are summarized in Fig. 4C; Supplemental Table 2.2; Pilcher et al. 2007; Mi et al. 2008). Candidate miRNA targets were identified in Solanaceae EST/TA databases for 14 of the 38 unique *S. demissum* miRNAs and for eight of the 30 unique *Nicotiana* miRNAs (Supplemental Table S2.2). The *Nicotiana* miR403 target was validated by 5'-RACE analysis (Supplemental Fig. S2.3).

To identify *MiS* small RNAs, we used small RNAs as queries in BLASTN searches of Solanaceae MITE sequences. Notably, 43 nonredundant *MiS* small RNA sequences from *S. demissum* and 11 nonredundant *MiS* small RNA sequences from *N. benthamiana* and *N. glutinosa* were identified (Table 2). These small RNAs matched members of the *MiS1-MiS3*, *MiS5-MiS15*, *MiS17*, *MiS20*, and *MiS22* families, and some were found in both plus and minus orientations (Table 2). Consistent with the size classes detected in the small RNA blots, the size distribution of cloned *MiS* small RNA peaked at 24 nt. The majority of other *MiS* small RNAs were 21 nt, 22 nt, or 23 nt in length (Fig. 4D). In contrast to cloned candidate miRNAs, a preference for a 5'-terminal A residue was observed for *MiS* small RNAs (Fig. 4D). Based on recent findings showing AGO-mediated sorting of small RNAs based on 5'-terminal nucleotide composition, we hypothesize that the 24-nt *MiS* small RNAs with a 5'-terminal adenosine may be recruited by AGO4 and involved in siRNA-mediated DNA methylation (RdDM) and transcriptional silencing, whereas AGO2 may recruit 21-nt *MiS* smRNAs with 5' A residues (Mi et al. 2008). BLASTN searches of Solanaceae nr GenBank databases identified potential MITE progenitors and/or targets of *MiS* smRNA-mediated transcriptional silencing within gene promoters (*Asc1* and *S11*, the *Alternaria* stem canker locus and the *Solanum chacoense* self-incompatibility *S11* allele, respectively), introns (*LeCTR3*, *ure*, *R1-like*), or upstream regions of genes (*R1-like*) (Table 2). BLASTN searches against the TIGR Plant Transcript Assembly database identified candidate hits with *E*-values < 0.01 for 25 of the 43 *MiS* small RNAs (Table 2). Some *MiS* small RNAs retrieved multiple hits. One *MiS1-1* small RNA from *N. glutinosa*, sNg\_317\_22\_1 (Table 2), matched (+/-) a sequence within the 70-bp *MiS1-1*-derived alternative exon of the *N* gene (Fig. 2A; Table 2). The characteristic sequence, length, and terminal base composition of *S. demissum*, *N. benthamiana*, and *N. glutinosa* small RNAs support the hypothesis that the 24-nt *MiS* small RNAs are generated from MITEs by an NRPD1A-RDR2-DCL3 pathway. The finding that adenosine is the 5'-terminal nucleotide of most 24-nt *MiS* small RNAs supports the hypothesis that these RNAs interact with AGO4 for siRNA-dependent DNA methylation and transcriptional silencing. DCL2- and DCL4-dependent biogenesis



**Figure 4.** Characteristics of cloned small RNAs. (A) Length distribution of small RNAs of the total *S. demissum* data set (4421 smRNAs). (Black bars) Indicate the percentage of small RNAs in each size class among the total data set. (B) Proportion of the total *S. demissum* small RNAs within each of the four size classes with A or T or G or C as the 5'-terminal nucleotide. Within each size class, (gray bar) the portion of small RNAs with a 5'-terminal A or T; (white bars) the portion with a 5'-terminal G or C. (C) Size distribution and 5'-terminal nucleotide composition of miRNAs ranging from 15 to 27 nt in length. (Black bar) The total number of cloned miRNA in each size class. The number of miRNAs with a 5' (dark gray bar) A, (gray bar) T, (light gray bar) G, or (white bar) C residue. (D) Size distribution and 5'-terminal nucleotide composition of MITE small RNAs ranging from 15 to 31 nt in length. The 5' nucleotide composition is indicated as in C.

pathways may also generate 21–22-nt MITE small RNAs that mediate post-transcriptional regulation of MITE expression.

#### Genetic requirements for MITE small RNA biogenesis

To test the hypothesis that *MiS* 21–24-nt small RNAs are products of DCL2-, DCL3-, or DCL4-dependent siRNA pathways, we investigated the genetic requirements for their biogenesis. We generated RNAi lines targeting candidate orthologs of *DCL2*, *DCL3*, and *DCL4* in *N. tabacum*, and an RNAi line targeting the *RDR2* ortholog of *S. tuberosum*. Supplemental Tables S4.1, S4.2, and S4.3 and Supplemental Figures S2.1 and S2.2 provide details on vector construction and characterization of *N. tabacum* RNAi lines. Characterization of transgenic NtDCL RNAi lines demonstrated that *N. tabacum* orthologs of *Arabidopsis* *DCL2-4* loci were nonredundantly silenced (Supplemental Fig. S2.2). Low-molecular-weight RNA fractions were isolated from mature leaf tissue of tobacco NtDCL and potato RDR2 RNAi lines and wild-type (WT) controls, and blotted and hybridized with *MiS5-1* small RNA probe, sSd\_3090\_20, or the *MiS17* probe (Supplemental Table S3.2), to assess *MiS* small RNA accumulation in the RNAi lines.

Accumulation of 24-nt *MiS5-1*-related small RNA, detected with the sSd\_3090\_20\_2 (Table 2) hybridization probe, was sig-

nificantly reduced, with a relative abundance of 0.13 in the potato *RDR2* RNAi line compared to wild-type (WT) levels. This reduction was accompanied by the appearance of several longer RNA species of ~60 to 100 nt (Fig. 5A). The sSd\_3090\_20 probe detected *MiS5* small RNA in WT potato and tomato, but not in tobacco, *Arabidopsis*, or rice, which may reflect either the low copy number of *MiS5* in the tobacco genome or divergence of *Solanum* and *Nicotiana* *MiS5* sequences (Supplemental Fig. S1; Supplemental Table S.1). In contrast, the MIR171 probe detected conserved MIR171 at similar levels in the potato *RDR2* RNAi line and wild-type potato (Fig. 5A).

In tobacco, four *DCL2* RNAi lines—NtDCL2-1-14, NtDCL2-12, NtDCL2-14, and NtDCL2-3-4—showed a slight increase in MIR171 accumulation but no detectable change in *MiS17* small RNA accumulation compared to wild type (Fig. 5B). In contrast, NtDCL3 and NtDCL4 lines showed a threefold to 20-fold reduction in *MiS17* small RNA, but no significant change in the accumulation of MIR171 (Fig. 5B) compared to wild type. Intriguingly, shorter small RNAs of 21 to 22 nt as well as longer small RNAs of ~30 nt and greater appear in the NtDCL3 lines, but not in the NtDCL4 lines.

These data indicate that *MiS* small RNA biogenesis involves RDR2, DCL3, and possibly DCL4. In *Arabidopsis*, RDR2 and DCL3 operate in a pathway generating ~24-nt siRNAs that guide TGS;





Table 2. Continued

MiS-siRNA ID	MITE-smRNA sequence	MiS <sup>a</sup>	E-value	Strand	Gene <sup>b</sup>	EST/TA <sup>c</sup>	Function	E-value	Strand
sNg_1411_17_1	CTACCCCTTAATGAGAAG	MiS17.Nt740	3.00E-04	+/-					
sNb_116_21_1	AGCTTATAAGGTGAGGATTGC	MISS.Nt109	4.00E-04	+/+					
sNb_1356_24_1	AATCCTCACCTCATAAGCTAGCTT	MISS.Nt109	4.00E-08	+/-					
sNb_1961_19_1	CTTTTGGGGTTGAGTTAGG	MISS.Nt114	2.00E-05	+/-					
sNg_1166_23_1	GGGGTTGAGTTAAACCCAATGTC	MISS.Nt21	3.00E-05	+/-					

Potential progenitors and targets of cloned *S. demissum*, *N. glutinosa*, or *N. benthamiana* small RNAs were identified using small RNAs (Supplemental Tables S2.1, S2.2, and S2.3 respectively) as queries in BLASTN searches of the *MiS* data set, *Solanum* and *Nicotiana* GenBank nr databases, and the Solanaceae Transcript Assembly Databases (<http://plantta.tigr.org/search.shtml>).

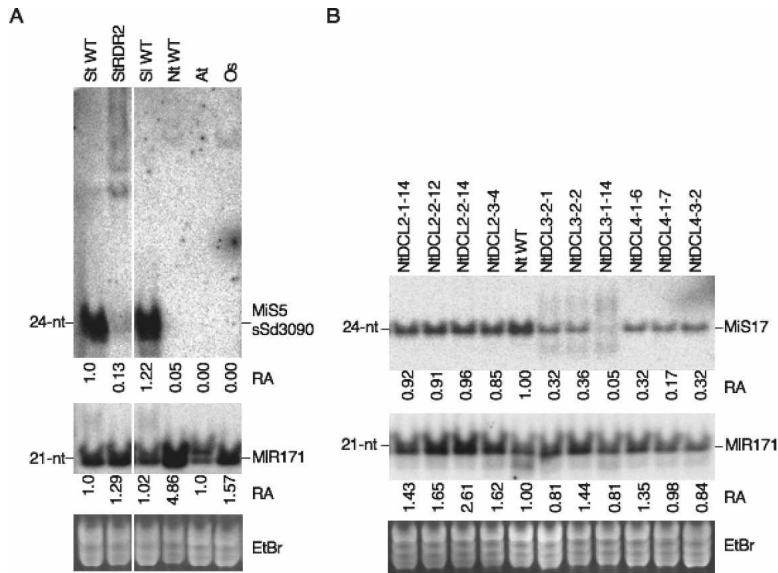
<sup>a</sup>Listed are the top hits obtained by BLASTN searches of the *MiS* data set using *S. demissum*, *N. glutinosa*, or *N. benthamiana* small RNAs as queries. The *S. demissum* *MiS* sequences are listed in Supplemental Data S1.1. The *N. glutinosa* or *N. benthamiana* *MiS* sequences are listed in Supplemental Data S1.2.

<sup>b</sup>Small RNA matches to MITEs located within or near genes identified by BLASTN searches of the Solanaceae GenBank nr database. All small RNA hits match identified MITEs, and locations are indicated with (U) upstream; (P) promoter; (I) intron; (E) exon; (AE) alternative exon; (5') 5'-UTR; (3') 3'-UTR. The GenBank accession number is listed below each gene.

<sup>c</sup>TIGR Plant EST and Transcript Assembly (TA) databases (<http://plantta.tigr.org/index.shtml>).

<sup>d</sup>(*Asc1*) *Solanum lycopersicon* *Alternaria* stem canker locus.

<sup>e</sup>(*ST1*) *Solanum chacoense* self-incompatibility *ST1* allele.



**Figure 5.** RNA-dependent RNA polymerase (RDR2) and DICER-LIKE (DCL) are involved in the biogenesis of *MiS* small RNAs. Small RNA-enriched fractions isolated from wild type and RNAi lines were prepared and analyzed by Northern blot hybridization as described in Figure 3. Blots were hybridized with *MiS* and miRNA probes (see Supplemental Table S3.2 for a description of probes). (A) Detection of *MiS5* (Sd3090 probe) small RNA and MIR171 in (St WT) wild-type *S. tuberosum* Katahdin; (St *rdr2*) *S. tuberosum* Katahdin RDR2 RNAi line; (SI WT) wild-type *S. lycopersicum* VF36; (Nt WT) wild-type *N. tabacum* SR1; (At) wild-type *Arabidopsis* Col; and (Os) wild-type *Oryza sativa* Nipponbare. (B) Detection of *MiS17* plus strand small RNA and MIR171 in wild-type tobacco SR1 and *N. tabacum* RNAi lines targeting (*DCL2*) DCL2-1, DCL2-2, and DCL2-3; (*DCL3*) DCL3-2 and DCL3-2; and (*DCL4*) DCL4-1 and DCL4-3 (for construction and characteristics of tobacco RNAi lines, see Supplemental Figs. S2.1 and S2.2; Supplemental Tables S4.1–S4.4). (A, B, bottom panels) The ethidium bromide (EtBr) staining of gels as loading controls. The numbers below the hybridization panels in A and B indicate the relative abundance (RA) of siRNA detected by Northern blot hybridization in each sample compared to StWT in A and NtWT in B. Relative siRNA abundance has been normalized (WT = 1.0) according to EtBr staining of gels in A and B.

however, a role for DCL4 in generating endogenous 24-nt small RNA is unexpected and has not been observed in *Arabidopsis*.

## Discussion

### Solanaceae MITEs contribute to evolution of gene complexity

MITEs are closely associated with genes (Wessler 1998; Mao et al. 2000), but the functional impact of these genic insertions has not been clearly established. Our investigation revealed a novel mechanism by which MITEs may assert gene regulation—via small RNA pathways. However, we also identified several MITEs that have the capacity to alter gene structure, thus playing the traditionally appreciated role of TEs in the evolution of new genes and transcriptome diversity. In future studies of MITE function, it will be important to account for the variety of mechanisms, both structural and small RNA-mediated, by which MITEs have an impact on gene regulation and genome evolution.

Our analysis of *MiS* demonstrates that MITEs can create new gene structures in several ways. We found that the *N* alternative exon, which generates the *N<sub>L</sub>* mRNA isoform, is encoded by *MiS1-1*. It is possible that *MiS1* contributes to regulation of *N* expression at the transcriptional and/or post-transcriptional level. TMV infection might induce transient modification of TGS mechanisms that lead to changes in *N* transcription. Virus infection could also cause changes in post-transcriptional regulation

of *N* expression by temporarily inhibiting PTGS and *MiS1* siRNA-guided cleavage of the *N<sub>L</sub>* mRNA. The premature translation termination codon induced by the *MiS1* alternative exon could render *N<sub>L</sub>* mRNA a target of nonsense-mediated decay, a post-transcriptional surveillance pathway. The predicted product of *N<sub>L</sub>* mRNA, a truncated TIR-NB protein (*N<sup>tr</sup>*), may play a functional role in TMV resistance and participate in TIR domain-deduced recognition and oligomerization of *N* (Mestre and Baulcombe 2006; Burch-Smith et al. 2007). Additionally, we found evidence that *MiS1* may provide a polyadenylation signal when located in the 3'-UTR of *Nicotiana* transcripts, indicating that *MiS1* may have a general capacity to generate novel 3'-truncated polyadenylated transcripts. Furthermore, as with other TEs, insertion of *MiS* in exons can introduce premature stop codons, and insertion and subsequent excision can introduce short insertions and deletions, which contribute to the diversity of *R* genes in particular (Kuang et al. 2005). We also found that *MiS5-1* has the capacity to capture and move host sequence (Fig. 1A), similar to Pack-MULEs reported in *Arabidopsis*, rice, and maize (Jiang et al. 2004a). All of these mechanisms are expected to contribute to the complexity of the genome.

### A model for MITE small RNAs in gene regulation

Our identification of *MiS* small RNAs and multiple genes bearing *MiS* insertions, which could serve as both origins of small RNA and targets for small RNA-mediated silencing, suggests that *MiS* silencing pathways may be co-opted for regulation of genes bearing *MiS*, similar to the regulation imposed by small RNA-mediated regulation via the SINE-like direct repeats in the *FWA* promoter (Chan et al. 2004). This hypothesis can be extended by proposing that the amplification and diversification of *MiS* and other mobile elements may contribute to evolution of networks of coordinately regulated genes via insertion and subsequent selection of homologous elements in multiple genes, thus providing target sites for co-regulation by RISC or RITS complexes loaded with target-specific *MiS* and other TE small RNAs. Indeed, two studies have implicated MITEs as negative transcription regulators that may be targets of TGS: investigation of the *LAT59* gene-defined *Stowaway-Le1*, which is located in the promoter, as a negative regulator of expression (Twell et al. 1991; Bureau and Wessler 1994), and inhibition of DNA methylation of the rice ubiquitin2 promoter, harboring *Kiddo* and *MDM1*, resulted in an increase in the ubiquitin2 transcript (Yang et al. 2005). *Arabidopsis* small RNAs are sorted into distinct AGO complexes, and this sorting has important functional implications. AGO1 preferentially recruits small RNAs with a 5'-terminal uridine, and AGO2 and AGO4 recruit small RNAs with a 5'-terminal adenosine (Mi et al. 2008). Consistent with these findings, we determined that the 5'-terminal residue of the majority of our candidate miRNAs is

uridine. The 5'-terminal residue of the majority of *MiS* small RNAs was determined to be adenosine, suggesting that *MiS* smRNAs interact with AGO4 as described for repeat-associated siRNAs (rasiRNAs) involved in RNA-directed DNA methylation, histone modification, and transcriptional gene silencing of transposons and other repeats in *Arabidopsis*. AGO2 may recruit *MiS* 21-nt RNAs with 5'-terminal A residues.

Secondary siRNA generated from PTGS may also play a role in these hypothetical *MiS* regulatory networks. The identification of multiple conserved *MiS* sequences in ESTs databases, and their potential to form siRNA precursors from dsRNA via sense-antisense pairing or hairpin structures from RNA transcribed from inverted repeats suggests that *MiS* may have the capacity to generate small RNAs similar to nat-siRNAs or NRPD1A-independent siRNAs derived from single-stranded hairpin precursors. These siRNAs potentially target gene transcripts bearing *MiS* insertions for PTGS. The identification of *Nicotiana MiS1* small RNAs matching paired 3'-UTRs containing complementary *MiS1* insertions (Fig. 2B; Table 2) suggests a potential role for *MiS1* small RNAs in post-transcriptional gene silencing. The presence of a transcribed *MiS9* insertion in the 3'-UTR of tomato and potato ESTs/TAs encoding putative DCL2 products (Supplemental Table S2.1) and identification of *MiS9* small RNA with perfect matches to the predicted *MiS9* ssRNA hairpin also supports the hypothesized role of MITE-derived siRNAs in post-transcriptional silencing. In this regard, it is interesting to note that deep sequencing of tomato small RNAs revealed that TAPIR fold-back elements produce small RNAs in a miRNA-like pattern (Moxon et al. 2008). In the cases of pseudogenes with *MiS* insertions, PTGS may provide the ability to generate secondary siRNAs that regulate functional homologs.

### The biogenesis of MITE siRNA

Using RNAi knockdown lines, we showed that RDR2, DCL3, and possibly DCL4 contribute to *MiS* small RNA accumulation in different ways. As expected, knockdown of *RDR2* and *DCL3* significantly reduced 24-nt *MiS* small RNA accumulation and, in addition, resulted in the appearance of novel small RNA species. In the StRDR2 RNAi line, several larger *MiS5* RNAs, some with a length of ~100 nt, were observed. In the NtDCL3-1 and NtDCL3-2 RNAi lines, novel size classes of ~21–22-nt novel small RNAs and >30 nt were detected. These species were not observed in the NtDCL4, NtDCL2, or StRDR2 RNAi lines.

Our results support the hypothesis that biogenesis of the majority of *MiS* small RNA involves an NRPD1A/RDR2/DCL3/AGO4-dependent core pathway involved in transcriptional silencing and DNA methylation in *Arabidopsis* (Henderson and Jacobsen 2007). The smaller 21–22-nt RNAs detected in the NtDCL3 RNAi lines could result from DCL4 and DCL2 cleavage of accumulated *MiS* dsRNA precursor. The absence of these small RNAs in potato RDR2 RNAi lines indicates that their biogenesis may depend on RDR2. Processing of RDR2 products by DCL4 has been observed in plants expressing ssRNA hairpin structures from transgenes for induced silencing (Henderson and Jacobsen 2007; Smith et al. 2007). Nevertheless, a role for DCL4 in *MiS* small RNA accumulation is interesting and awaits further investigation.

Understanding the interactions of different silencing pathway components in production of TE-derived small RNAs will be resolved with genetic analysis that incorporates lines carrying mutations in multiple silencing RNA pathway loci. These lines will be required in experimental systems to investigate the potential functional roles of Solanaceae small RNAs. Given the high

copy number of gene-associated transposons in Solanaceae, their potential to generate both RNA polymerase IV-dependent and -independent substrates for small interfering RNAs, and the significant functional redundancies documented at multiple steps in the silencing pathway (Henderson et al. 2006; Smith et al. 2007; Zhang et al. 2007), it will be important to systematically account for the different mechanisms of TE-derived small RNAs and their potential functional roles in gene silencing.

## Methods

### MITE analyses

Transposable elements were investigated at six resistance gene loci: the *R1* locus on Chromosome 5 of *S. demissum* (Kuang et al. 2005), the major late blight resistance cluster (*MLB*) on Chromosome 11 of *S. demissum* (H. Kuang and B. Baker, unpubl.), the *N* locus in tobacco (Whitham et al. 1994), the *Bs4* locus in tomato (Schornack et al. 2004), the *Prf/Pto* locus in tomato (Chang et al. 2002), and the *RB* locus on Chromosome 8 of *Solanum bulbocastanum* (Song et al. 2003). Sequences of two or three haplotypes of *R1*, *Bs4*, *N*, *Prf/Pto*, and *RB* were obtained from GenBank accessions (*R1*: AC135288, AC139840, AC142505, AC144791, AC145120, AC146506, AC149265, AC149266, AC149267, AC149287, AC149288, AC149290, AC149291, AC149301, AC149487, AC150162, AC151803, AC151815, AC151957, AC154033; *Bs4*: AY438027, AY438028; *N*: U15605, A535010 (*NH*); *Prf/Pto*: AF220602, AF220603; *RB*: AY303170, AY303171).

MITEs were detected using the related empty site (RESite) method, which confirms and delimits a MITE sequence by comparing its flanking sequence with a homologous region (Le et al. 2000). Sequences from syntenic regions or homologs of *R*-genes were aligned using ClustalX (Thompson et al. 1994). All insertions >60 bp were investigated for transposable element structural features such as TIRs and TSDs. To obtain additional MITEs, the sequences of all identified MITEs were used to perform BLASTN queries ( $E < e^{-5}$ ) in GenBank and in the TIGR Plant Transcript Assembly databases of tomato, potato, tobacco, pepper, *N. benthamiana*, and *Petunia hybrida* ([http://tigrblast.tigr.org/euk-blast/plantta\\_blast.cgi](http://tigrblast.tigr.org/euk-blast/plantta_blast.cgi)) (Childs et al. 2007). Copy numbers were estimated by performing BLASTN searches with a representative member of each *MiS* family (Supplemental Data S1.1) in potato and tomato bacterial artificial chromosome (BAC) clone end and tobacco genomic sequence ([http://www.tigr.org/tdb/sol/sol\\_ma\\_blast.shtml](http://www.tigr.org/tdb/sol/sol_ma_blast.shtml); [http://www.tigr.org/tdb/sol/sol\\_ma\\_blast.shtml1](http://www.tigr.org/tdb/sol/sol_ma_blast.shtml1); NCBI *Nicotiana tabacum* gss; Supplemental Table S1.2). MITEs may generally be considered to be shorter than 600 bp; however, we identified MITE families that included longer members, which we classified with their shorter counterparts based on the following criteria: We define MITEs with significantly similar internal sequences ( $< e^{-5}$  for BLASTN) to be members of the same MITE family, and MITEs with no significant similarity to be members of different MITE families, and further define MITE families with related TIR sequences as superfamilies. Each MITE was given a unique, two-number designation *MiS(X)-X*, with the first number identifying the family and the second identifying the individual member. All MITEs identified previously in Solanaceae species were included in this study for further analysis (Oosumi et al. 1995; Pozueta-Romero et al. 1996; Rebatchouk and Narita 1997; Mao et al. 2001; Schenke et al. 2003; Guyot et al. 2005).

### Small RNA cloning and analyses

Total RNA (600 µg) was resolved by electrophoresis on a denaturing 15% polyacrylamide gel. A gel slice containing RNAs from

the range of ~10–60 nt was excised, and RNA was eluted by incubating in 0.3 M NaCl overnight at 4°C and ethanol-precipitated. Eluted RNAs were ligated to RNA/DNA chimeric adaptors (5'-ATCGTAGGCACCUGAAA-3' and 5'-pUUUCTGTA GGCACCATCAATT-3'). The final ligation product was resolved through denaturing polyacrylamide gel electrophoresis, gel-eluted, ethanol-precipitated, and reverse-transcribed using an antisense reverse primer and superscript III RT (Invitrogen). The resulting cDNA was PCR-amplified using sense and antisense primers, and the products were resolved on a native 15% polyacrylamide gel, eluted, and BanI-digested. Digestion products were ligated to the TOPO-TA cloning vector (Invitrogen), and transformed plasmid DNA was purified and sequenced from clones using standard sequencing methods.

A Perl program was written to extract small RNA sequence from the raw ABI files by recognizing the adaptor sequences with a two-edit distance (insertion, deletion, and mismatch, with each count as one edit distance). All small RNAs between 1 and 100 nt in length were collected in FASTA format and numbered numerically from 1 to 4421. For example, the identifier sSd\_1915\_24 refers to *S. demissum* small RNA number 1915 and is 24 nt in length. Statistics on the small RNA size and end composition were compiled from the total small RNA data set contained in the FASTA file. A nonredundant small RNA FASTA formatted file was generated by removing redundant small RNA sequences and appending a read number to the small RNA identifier (Supplemental Data S2.1). MITE small RNAs (47 reads) and miRNAs (74 reads) were identified by BLASTN searches against a database of all *MiS* sequences (*MiS1-22*) and against the miR database <http://microrna.sanger.ac.uk/sequences/index.shtml> (Griffiths-Jones 2004; Griffiths-Jones et al. 2006), using a cut-off *E*-value of  $1e-3$ . Small RNAs derived from rRNA and ribosomal genes (812 reads), tRNA (25 reads), and non-*MiS* transposable elements (18) were identified by BLASTN search against the TIGR Plant Repeat database (Ouyang and Buell 2004) and NCBI nr nucleotide database. Small RNAs that remained unidentified by the above searches (3323 reads) were placed into an unknown small RNA file. Candidate targets were predicted for all small RNAs by BLASTN against TIGR Plant Transcript Assembly databases ([http://tigrblast.tigr.org/euk-blast/plantta\\_blast.cgi](http://tigrblast.tigr.org/euk-blast/plantta_blast.cgi)).

### Small RNA detection

To detect *MiS* small RNA accumulation in various Solanaceae species, RNA was extracted from young leaf tissue of *O. sativa* Nipponbare, *A. thaliana* Columbia, *N. benthamiana*, *N. tabacum* SR1, *N. tabacum* SR1:NN (TG34), *N. glutinosa*, *S. demissum*, *S. tuberosum* Kennebec and Katahdin, *S. lycopersicum* VF36, and from floral tissues of *S. demissum* and *S. lycopersicum* plants, using TRIzol reagent (Sigma-Aldrich) as described previously (Chellappan et al. 2005), and blots carrying 40 µg of total RNA per lane were hybridized with random-primed DNA probes using amplified MITE sequences as templates (Supplemental Table S3.1). Hybridization and subsequent washes were performed as described (Chellappan et al. 2005).

To determine the genetic requirements for *MiS* small RNA accumulation, low-molecular-weight RNA was extracted from the mature leaf tissues of *O. sativa*, *A. thaliana*, *S. tuberosum* Katahdin, *S. lycopersicum* VF36, *N. tabacum* SR1, and from the RNAi knockdown lines in *S. tuberosum* Katahdin and *N. tabacum* SR1, as described previously (Diaz-Pendon et al. 2007), and blots carrying 10 µg per lane of low-molecular-weight enriched RNA were hybridized with end-labeled DNA oligonucleotides (Supplemental Table S3.2) to detect sSd3090\_20\_1 (*MiS5*), MIR171, and *MiS17* small RNAs. Hybridization and subsequent washes were

done as described (Diaz-Pendon et al. 2007). Blots were exposed to phosphor storage screens and scanned using a PhosphorImager (Molecular Dynamics). The quantification of band density was determined by Quantity One software (Bio-Rad, version 4.6.5) using the Volume Rectangle Tool.

### Generation of transgenic DCL and RDR2 RNAi lines

To generate tobacco RNAi lines targeting the different *DCL* genes, *Arabidopsis* DCL2, DCL3, and DCL4 protein sequences were used as queries for tBLASTN searches against the NCBI *Nicotiana tabacum* gss database. All *Nicotiana* sequence hits were collected and categorized as one of the *DCL* candidate homologs according to their BLASTX scores against the *Arabidopsis* reference protein (Supplemental Table S4.2). *Nicotiana* DNA fragments corresponding to each *DCL* homolog were amplified using the primers described in Supplemental Table S4.1 and cloned into a pENTR/D/TOPO vector (Invitrogen). For each *DCL* homolog, unique constructs were generated to target different regions of the *DCL* gene; these constructs are distinguished by a numerical appendix, for example, NtDCL2-1, NtDCL2-2, and NtDCL2-3 (Supplemental Fig. S2.1; Supplemental Table S4.2). The DNA fragments (Supplemental Table S4.3) were introduced into a pHELLSGATE8 vector by LR reaction (Invitrogen) to produce a hairpin construct, sequenced, and transferred to *Agrobacterium* strain LBA4404 for transformation of *N. tabacum* SR1. Different lines generated with the same construct are distinguished by a line number, for example, Nt DCL2-1-14 (see Supplemental Fig. S2.2). Transgenic T0 generation plants were screened for silencing of *DCL* transcript expression by RT-PCR (Supplemental Fig. S2.2; Supplemental Table S4.4), and positive transformants were selected for small RNA analyses. A 413-bp cDNA fragment corresponding to the *N. benthamiana* RDR2 cDNA sequence (AY722009) was amplified from potato *cv.* Katahdin cDNA. The primers used for amplification included 5'-CACCCGCCAGAAGTCTGTGTATC-3' (RC13-forward) and 5'-TCACAGTGATCGACAACAGATTGC-3' (RC14-reverse). Construct design and transformation procedures were performed essentially as described by Bhaskar et al. (2008). Transgenic potato (*cv.* Katahdin) plants silenced for RDR2, *St* RDR2, were verified by RT-PCR.

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