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Post-transcriptional small RNA pathways in plants: mechanisms and regulations

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Small RNAs are riboregulators that have critical roles in most eukaryotes. They repress gene expression by acting either on DNA to guide sequence elimination and chromatin remodeling, or on RNA to guide cleavage and translation repression. This review focuses on the various types of post-transcriptional small RNA-directed pathways in plants, describing their roles and their regulations.

For a long period of time, DNA, the support of heredity, and proteins, the actors of the cellular machinery, have been considered the most important components of biological systems. By contrast, RNA was originally considered as an intermediate molecule, bridging the gap between DNA and protein (mRNA), or serving functional roles during splicing (snRNA) and translation (tRNA and rRNA). The recent discovery of an increasing number of large and small non-protein-coding RNAs with specific regulatory roles has changed our view of gene expression. In particular, 20- to 27-nucleotide (nt) small RNAs belonging to two classes, microRNAs (miRNAs) and short interfering RNAs (siRNAs), are known to play essential roles in the four Eukaryote kingdoms (protists, fungi, plants, animals), with the surprising exception of the yeast *Saccharomyces cerevisiae*. Small RNAs are involved in a variety of phenomena that are essential for genome stability, development, and adaptive responses to biotic and abiotic stresses. Their mode of action also is diverse. They guide DNA elimination during the formation of the macronucleus in protists and heterochromatin assembly in fungi and plants. They target endogenous mRNAs for cleavage and translational repression in plants and animals, and protect both plant and animal cells against virus infection through an RNA-based immune system. They also control the movement of transposable elements at the transcriptional and post-transcriptional level in plants and animals.

**Cis-acting siRNAs as an RNA-based immune mechanism and a silencing tool**

Because small RNAs are repressors of gene expression, small RNA-mediated regulation is often referred to as RNA silencing, gene silencing, or RNA interference (RNAi). RNA silencing was discovered in plants more than 15 years ago during the course of transgenic experiments that eventually led to silencing of the introduced transgene and, in some cases, of homologous endogenous genes or resident transgenes [Matzke et al. 1989; Linn et al. 1990; Napoli et al. 1990; Smith et al. 1990; van der Kroef et al. 1990]. Gene silencing results from transcription inhibition (transcriptional gene silencing [TGS]) or from RNA degradation (post-transcriptional gene silencing [PTGS]), and correlates with the accumulation of siRNAs corresponding to the silenced promoter or to the degraded RNA, respectively (Hamilton and Baulcombe 1999; Mette et al. 2000). The production of virus-derived siRNAs was observed in response to virus infection [Hamilton and Baulcombe 1999], and plant mutants defective in PTGS were hypersusceptible to virus infection [Mourrain et al. 2000; Dalmay et al. 2001; Qu et al. 2005; Schwach et al. 2005]. Studies in worm revealed that mutants defective in RNAi (the animal counterpart of PTGS) lose control of their transposable elements [Ketting et al. 1999]. Altogether, these results suggested that the PTGS/RNAi pathway corresponds to an RNA-based immune system that allows cells to control endogenous (transposons) or exogenous (virus, transgenes) nucleic acid invaders through the action of *cis*-acting siRNAs, which derive from and target the invaders [Vance and Vaucheret 2001; Bartel 2004; Baulcombe 2004; Dunoyer and Voinnet 2005].

Biochemical dissection of the RNAi pathway in animals has revealed that long double-stranded RNAs [dsRNAs] are potent inducers of RNAi and that the pathway involves RNase III enzymes, which dice dsRNAs into siRNAs, and Argonaute enzymes, which slice single-stranded RNAs complementary to siRNAs [for review, see Du and Zamore 2005; Tomari and Zamore 2005]. In *Drosophila* [Fig. 1], dsRNAs are processed into double-stranded siRNAs by DCR-2, an RNase III enzyme of the Dicer family. DCR-2 interacts with the dsRNA-binding domain protein R2D2, resulting in asymmetric...
siRNA duplexes, with Dicer positioned at the less stable end and R2D2 at the other end. This asymmetry orients the loading of siRNA duplexes onto AGO2, a protein of the Argonaute family, which interact with Dicer enzymes. The strand of the siRNA duplex referred to as the passenger strand is cleaved by the RNA slicer activity of AGO2, allowing the other siRNA strand to pair with homologous mRNAs and guide AGO2-catalyzed mRNA cleavage. In Arabidopsis, Dicer enzyme that processes siRNAs during S-PTGS has not been fully elucidated, but it is likely to be DICER-LIKE4 (DCL4), which processes siRNAs from long dsRNAs during IR-PTGS (Dunoyer et al. 2005). Transgene-derived siRNAs are methylated by HUA ENHANCER1 (HEN1) (Boutet et al. 2003; Li et al. 2005; Yu et al. 2005) and guide sequence-specific ARGONAUTE1 (AGO1)-catalyzed mRNA cleavage (Morel et al. 2002; Baumberger and Baulcombe 2005; Qi et al. 2005). Additional components have been identified by forward and reverse genetic screens (Table1), including NUCLEAR RNA POLYMERASE IVa [NRPD1a], RNA-DEPENDENT RNA POLYMERASE2 [RDR2], SILENCING DEFECTIVE3 [SDE3], and WERNER EXONUCLEASE [WEX] (Dalmay et al. 2000; Glazov et al. 2003; Herr et al. 2005). The steps at which these proteins act in the PTGS pathway are still not fully understood. Interestingly, NRPD1a and SGS3 appear to be plant specific.

Part of the PTGS machinery [AGO1, HEN1, RDR6, and SGS3] also participates in the RNA-based immune response against cucumber mosaic virus (CMV) infection, a widely distributed virus that infects >800 plant species (Mourrain et al. 2000; Morel et al. 2002; Boutet et al. 2003), suggesting that transgene-derived RNAs produced during S-PTGS likely mimic viral RNAs during CMV infection and activate the corresponding RNA-based immune response. However, it is unlikely that all viruses activate the same response. Indeed, rdr6 mutants show hypersusceptibility to CMV, a cucumovirus, and potato virus X [PVX], a potexvirus, but not to turnip mosaic virus [TuMV], a potyvirus; turnip crinkle virus [TCV], a carmovirus; turnip vein clearing virus [TVCV] and tobacco mosaic virus [TMV], tobamoviruses; or tobacco rattle virus [TRV], a tobravirus (Dalmay et al. 2000, 2001; Mourrain et al. 2000; Qu et al. 2005; Schwach et al. 2005). In contrast, rdr1 mutants impaired in a paralog of RNA-DEPENDENT RNA POLYMERASE6 show hypersusceptibility to TMV, suggesting that different post-transcriptional siRNA-mediated pathways are activated in response to infection by different viruses (Yu et al. 2003). Given the existence of 10 AGOs, four DCLs, and six RDRs in the plant model Arabidopsis (Morel et al. 2002; Schauer et al. 2002, Yu et al. 2003), the varying responses of different viruses to these mutations presumably reflect a diversification of siRNA pathways.

MicroRNAs as endogenous regulators of gene expression

Because transposon-, virus-, and transgene-derived siRNAs target the degradation of RNAs from which they derive, they are cis-acting siRNAs, and the immune phenomenon that relies on their action is referred to as auto-

![ Figure 1. miRNA and siRNA pathways in animals. The pathways in Drosophila are shown as exampled, owing to the remarkable dissection of the pathway in this organism. A color code is used to indicate members of the same gene family. Translation repression and mRNA cleavage steps involve additional components that are not represented in this figure.](image-url)
lencing (Bartel 2004). In contrast, miRNAs are endogenous small RNAs that are produced by genes distinct from the genes that they regulate, and, as such, miRNA-mediated regulation is referred to as heterosilencing (Bartel 2004). The first miRNAs discovered were the small temporal RNAs (*stRNAs*) *lin-4* and *let-7*, which were identified from a screen of *Caenorhabditis elegans* mutants exhibiting timing defects during larval development (Lee et al. 1993; Reinhart et al. 2000). These two single-stranded small RNAs appeared to be processed from imperfectly paired stem-loop precursor RNAs that are ∼70 nt long, and the mature 21-nt stRNAs repressed gene expression by binding to multiple copies of partially complementary sites within the 3’ untranslated region (UTR) of their target mRNAs and repressing translation (Ambros 2004; Bartel 2004; Du and Zamore 2005; Kim 2005). After the discovery of stRNAs and siRNAs, the stage was set for the search for more noncoding small RNA and several laboratories successfully cloned numerous 21- to 24-nt RNAs from worm, fly, mouse, human, and plants. The cloned small RNAs that displayed the characteristics of the original *lin-4* and *let-7* stRNAs—i.e., they are processed from partially folded stem-loop precursor RNAs that derive from genes that are distinct from the genes that they regulate—were collectively termed microRNAs. Most other endogenous small RNAs that do not fit this definition are collectively termed siRNAs and are assumed to derive from long dsRNAs resulting from convergent transcription or from the action of RNA-dependent RNA polymerases. As a complement to cloning efforts, genetic approaches have made important contributions to the identification of miRNAs and their targets. For example, the first miRNAs functionally characterized in flies (*bantam*) and plants (*miR319/JAW*) were both identified from genetic screens (Brennecke et al. 2003; Palatnik et al. 2003). Although both plant and animal miRNAs derive from partially folded stem-loop precursor RNAs, their biogenesis shows some differences [see Figs. 1, 2]. Different animal miRNAs are often clustered within a single pre-

### Table 1. Plant proteins involved in small RNA-mediated post-transcriptional gene silencing

<table>
<thead>
<tr>
<th>Code</th>
<th>Protein name [alternative name]</th>
<th>Discovery</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>AGO1</td>
<td>ARGONAUTE1 (DND)</td>
<td>Developmental screen (Bohmert et al. 1998) PTGS-deficient (Morel et al. 2002)</td>
<td>RNA slicer [Vaucheret et al. 2004; Baumberger and Baulcombe 2005; Qi et al. 2005]</td>
</tr>
<tr>
<td>AGO7</td>
<td>ARGONAUTE7 (ZIP)</td>
<td>Developmental screen (Hunter et al. 2003)</td>
<td>RNA slicer [Adenot et al. 2006]</td>
</tr>
<tr>
<td>DCL1</td>
<td>DICER-LIKE1 (CAF, EMB76, SIN1, SUS1)</td>
<td>Developmental screen (Schauer et al. 2002)</td>
<td>RNase III [Park et al. 2002; Reinhart et al. 2002]</td>
</tr>
<tr>
<td>DCL2</td>
<td>DICER-LIKE2</td>
<td>Reverse genetics (Xie et al. 2004)</td>
<td>RNase III [Xie et al. 2004; Borsani et al. 2005]</td>
</tr>
<tr>
<td>DCL3</td>
<td>DICER-LIKE3</td>
<td>Reverse genetics (Xie et al. 2004)</td>
<td>RNase III [Xie et al. 2004]</td>
</tr>
<tr>
<td>DRB4</td>
<td>dsRNA-BINDING PROTEIN4</td>
<td>Reverse genetics (Vazquez et al. 2004a)</td>
<td>dsRNA-binding protein [Hiraguri et al. 2005]</td>
</tr>
<tr>
<td>SDE3</td>
<td>SILENCING DEFECTIVE3</td>
<td>Developmental screen (Peragine et al. 2004)</td>
<td>RNA helicase [Dalmay et al. 2001]</td>
</tr>
<tr>
<td>SGS3</td>
<td>SUPPRESSOR OF GENE SILENCING3 (SDE2)</td>
<td>PTGS-deficient (Mourrain et al. 2000)</td>
<td>RNA stabilizer [Yoshikawa et al. 2005]</td>
</tr>
</tbody>
</table>

Proteins involved in small RNA-mediated TGS are reviewed in Chan et al. (2005).
cursor RNA, whereas plant miRNAs derive from individual precursors (Ambros 2004; Bartel 2004; Du and Zamore 2005; Kim 2005). Animal miRNA precursors (pri-miRNAs) adopt fold-back stem-loops that are first cleaved by the RNase III Drosha on each arm of the stem-loop to liberate the pre-miRNA (Ambros 2004; Bartel 2004; Du and Zamore 2005; Kim 2005), which is subsequently exported to the cytoplasm by Exp5 and cleaved by the RNase III Dicer to liberate the loop and the miRNA/miRNA* duplex that typically has two nucleotide 3' overhangs, similar to siRNA duplexes (Ambros 2004; Bartel 2004; Du and Zamore 2005; Kim 2005). Both Drosha and Dicer are assisted in the cleavage processes by specific dsRNA-binding proteins; for example, Pasha and Loquacious, respectively, in Drosophila [Du and Zamore 2005]. Both Drosa and Dicer are assisted in the cleavage processes by specific dsRNA-binding proteins; for example, Pasha and Loquacious, respectively, in Drosophila [Du and Zamore 2005]. In plants, the length of fold-back stem-loops generally is longer than that of animal stem-loops. DCL1 has both Drosha and Dicer functions and was found in the nucleus (Park et al. 2002; Reinhart et al. 2002; Papp et al. 2003; Kurihara and Watanabe 2004). DCL1 interacts with HYL1 to make both cuts within the miRNA precursor to liberate the miRNA/miRNA* duplex (Han et al. 2004; Vazquez et al. 2004a; Hiraguri et al. 2005). The mature miRNA is methylated by HEN1 [Yu et al. 2005], which also methylates siRNAs [Li et al. 2005]. Methylation likely protects small RNAs from degradation and polyuridylation. Whether small RNAs are methylated in other kingdoms is not known. Like DCL1, HYL1 and HEN1 are found in the nucleus, suggesting that miRNA processing is essentially nuclear. HASTY is an ortholog of EXP5 and likely exports miRNA duplexes or mature miRNAs to the cytoplasm [Park et al. 2005]. The accumulation of some miRNAs is not affected in hst mutants, suggesting either that these miRNAs are exported by other ways—for example, in association with AGO1—or that they act in the nucleus by targeting pre-mRNAs, such as proposed for the targeting of partially spliced DCL1 mRNA by miR162 in Arabidopsis [Xie et al. 2003; Park et al. 2005].

The action of plant and animal miRNAs also shows striking differences [see Figs. 1, 2]. Both associate with an Argonaute protein that is part of a ribonucleoprotein complex called RNA-induced silencing complex (RISC) (Bartel 2004). However, most animal miRNAs guide translational repression of their targets, which is accompanied by mRNA destabilization [Bagga et al. 2005; Lim et al. 2005], whereas all tested plant miRNAs guide AGO1-catalyzed mRNA cleavage [Llave et al. 2002; Kasschau et al. 2003], similar to AGO2-catalyzed mRNA cleavage by siRNAs in animals. MicroRNA-guided
miRNA cleavage is not limited to plants. Indeed, miR196 guides the cleavage of HoxB8 mRNA in mice, presumably due to the unusually high degree of complementarity between the miRNA and its target (Yekta et al. 2004). On the other hand, it is unclear if miRNAs can guide translation repression in plants. Indeed, the initial proposal that miR172 guides translation repression of AP2 mRNA in plants [Aukerman and Sakai 2003; Chen 2004] has recently been revisited, and the unchanged level of AP2 mRNA in plants overexpressing miR172 has been attributed to increased AP2 gene transcription, which results from reduced negative feedback regulation by the AP2 protein [Schwab et al. 2005].

As opposed to animal miRNAs that only show limited complementarity with their targets at the 5’ end of the miRNA [the seed region], plant miRNAs show a high degree of complementarity with their targets [Ambros 2004; Bartel 2004; Du and Zamore 2005; Kim 2005]. Indeed, the SCL genes were the first miRNA targets identified in plants because they exhibit full complementarity to miR171 [Llave et al. 2002; Rhoades et al. 2002]. The high degree of complementarity together with conservation between monocots and dicots allowed a confident prediction of miRNA targets [Rhoades et al. 2002; Jones-Rhoades and Bartel 2004]. In addition, plant miRNAs cleave their targets, allowing a simple validation of putative targets by 5’ RACE [Llave et al. 2002; Kasschau et al. 2003]. Most plant miRNAs exhibit four or less mismatches with their targets, with the exception of TCP targets that have up to five mismatches with miR319 [Palatnik et al. 2003], and these mismatches are usually located in the 3’ region of the miRNA [Mallory et al. 2004; Parizotto et al. 2004; Schwab et al. 2005]. Consistently, biochemical analyses using wheat germ extracts revealed that mutated targets with mismatches between positions 3 and 11 of the miRNAs were poorly cleaved, whereas mutated targets with mismatches at other positions were more efficiently cleaved [Mallory et al. 2004]. The importance of near-perfect complementarity between miRNAs and their targets in plants was used to gain insight into their roles by expressing miRNA-resistant targets carrying silent mutations that disrupt the pairing but not the protein sequence, and looking at the changes induced by this loss of miRNA-mediated regulation [Emery et al. 2003; Palatnik et al. 2003; Chen 2004; Mallory et al. 2004]. The definitive demonstration of the relationship between a miRNA [miR168] and its target [AGO1] was achieved by rescuing miR168-resistant AGO1 plants with an artificial miRNA carrying compensatory mutations that restored pairing [Vaucheret et al. 2004].

As opposed to animal miRNAs that control a variety of phenomena [Carrington and Ambros 2003; Ambros 2004; Bartel 2004], about two-thirds of the known plant miRNAs control the expression of transcription factors that regulate crucial steps during plant development [Rhoades et al. 2002; Jones-Rhoades and Bartel 2004]. Clues to the importance of miRNAs during development were revealed when mutants originally isolated in genetic screens for developmental defects turned out to be generally impaired at different points in the miRNA pathway (Table 1). For example, the carpel factory (caf), short integuments1 (sin1), suspensor (sus1), and embryo defective76 (emb76) mutants all have mutations in the DCL1 gene [Schauer et al. 2002], and the first argonaute mutants (ago1) were identified in plants and named for the squid tentacle-like appearance of their leaves [Bohmer et al. 1998]. Similarly, hyponastic leaves1 (hyl1), hua enhancer1 (hen1), and hasty (hst) were originally isolated as mutants impaired in hormone response, flower development, and the transition from juvenile to adult vegetative phase, respectively [Telfer and Poethig 1998; Lu and Fedoroff 2000; Chen et al. 2002]. For recent reviews on the specific developmental roles of miR156, miR159, miR160, miR164, miR165/166, miR167, miR172, and miR319 in phase changes, leaf morphogenesis and polarity, root initiation, vascular development, transition from vegetative to reproductive growth, or floral differentiation, see Chen [2005], Jones-Rhoades et al. [2006], and Zhang et al. [2006a]. Among the nontranscription factor targets of miRNAs, there are ATP-sulfurylases, superoxide dismutases, laccases, and ubiquitin-conjugating enzymes [Jones-Rhoades and Bartel 2004]. Two miRNAs—miR395, which regulates an ATP-sulfurylase, and miR399, which regulates a ubiquitin-conjugating enzyme—are induced by sulfate starvation and phosphate starvation, respectively, pointing to an adaptive role of miRNAs in stress response [Jones-Rhoades and Bartel 2004; Fujii et al. 2005].

Trans-acting siRNAs as secondary regulators of gene expression in plants

Intensive cloning efforts in plants have revealed that the great majority of endogenous small RNAs are DCL3-processed 24-nt cis-acting siRNAs that derive from heterochromatin, transposons and repeat elements [Reinhart et al. 2002; Xie et al. 2004; Gustafson et al. 2005; Lu et al. 2005]. Another class corresponds to DCL1-processed miRNAs that derive from MIR genes, and a third class corresponds to DCL4-processed trans-acting siRNAs (ta-siRNAs) that derive from TAS genes [Peragine et al. 2004; Vazquez et al. 2004b; Allen et al. 2005]. TAS genes transcribe long primary RNAs that are not predicted to encode proteins and seem to function by serving as the precursors for ta-siRNA production (Fig. 2). TAS primary RNAs are cleaved by specific miRNAs so the production of ta-siRNAs requires AGO1, DCL1, HEN1, and HYL1 [Peragine et al. 2004; Vazquez et al. 2004b; Allen et al. 2005]. The production of ta-siRNAs also requires SGS3 and RDR6, which transform one of the two single-stranded TAS cleavage products into dsRNA [Peragine et al. 2004; Vazquez et al. 2004b; Allen et al. 2005]. DMR6-derived TAS dsRNAs are sequentially processed into 21-nt ta-siRNAs by DCL4 [Gascioli et al. 2005; Xie et al. 2005b; Yoshikawa et al. 2005], which associates with the dsRNA-binding protein DRB4 [Hiraguri et al. 2005; Adenot et al. 2006], starting from the miRNA-cleaved end of the TAS precursors, to generate clusters of ta-siRNAs [Vazquez et al. 2004b; Allen et al. 2005; Yoshikawa et al. 2005].
Like miRNAs, ta-siRNAs are methylated by HEN1 [Li et al. 2005], and like miRNAs, the 5’ half of ta-siRNAs shows a high level of complementarity with endogenous mRNAs. ta-siRNAs regulate the expression of their target mRNAs by guiding mRNA cleavage [Peragine et al. 2004; Vazquez et al. 2004b; Allen et al. 2005]. Interestingly, different members of the same gene family can be targeted by either miRNAs or ta-siRNAs. For example, members of the same PPR subfamily are targeted by either mir161 or TAS2 ta-siRNAs [Rhoades et al. 2002; Allen et al. 2005; Yoshikawa et al. 2005]. In addition, members of the ARF family are targeted by either mir160, mir167, or TAS3 ta-siRNAs [Rhoades et al. 2002; Allen et al. 2005]. Like some miRNAs, some ta-siRNAs may act in the nucleus, as suggested by the complementarity of one of the TAS1a-derived ta-siRNAs with the intron of the At2g46740 pre-mRNA, which encodes an FAD-binding domain-containing protein, and the elevated level of unspliced At2g46740 RNA in tasiRNA-deficient mutants [Vazquez et al. 2004b]. It is likely that AGO1 is responsible for cleaving mRNAs targeted by TAS1 ta-siRNAs because TAS1 ta-siRNAs associate with AGO1 in vitro [Baumberger and Baulcombe 2005; Qi et al. 2005]. AGO1 also seems to be involved in TAS2-mediated regulation, whereas AGO7 seems to be involved in TAS3-mediated regulation. Indeed, ago7 mutants show reduced levels of TAS3 ta-siRNAs, but not of TAS1 or TAS2 ta-siRNAs, and elevated levels of TAS3 ta-siRNA targets, but not of TAS1 or TAS2 ta-siRNA targets [Peragine et al. 2004; Allen et al. 2005; Adenot et al. 2006]. Whether ta-siRNAs also exist in animals has not been reported.

Natural cis-acting siRNAs as specific regulators of gene expression in plants

An additional class of endogenous siRNAs [nat-siRNAs], which derives from pairs of natural cis-antisense transcripts, was recently discovered in Arabidopsis, revealing an endogenous function for DCL2 [Borsani et al. 2005]. In the case examined, one transcript is expressed constitutively whereas the other is induced by salt treatment. When both transcripts are present, a stable 24-nt nat-siRNA deriving from the region of complementary and corresponding to the induced transcript is produced through the action of DCL2, NRPD1a, RDR6, and SGS3 [Fig. 2]. This 24-nt nat-siRNA guides the cleavage of the constitutive transcript and establishes a phase for the sequential production of 21-nt nat-siRNAs by DCL1 and further cleavage of the constitutive transcript. However, the production of these 21-nt nat-siRNAs does not appear to be required for down-regulating the constitutive transcript because dcl1 mutants that accumulate the 24-nt nat-siRNA but not the 21-nt nat-siRNAs exhibit transcript levels similar to those of wild-type plants. Therefore, unlike 21-nt ta-siRNAs that have a functional role in targeting unlinked transcripts, 21-nt nat-siRNAs may only be produced to ensure cleavage in cis. There are -2000 pairs of natural cis-antisense transcripts in Arabidopsis [Wang et al. 2005]. nat-siRNAs corresponding to other cis-antisense transcript pairs exist and are detected only under inducible conditions, suggesting that this type of regulation may not be restricted to a single example of overlapping transcripts [Borsani et al. 2005].

Regulation of the miRNA and ta-siRNA pathways by miRNAs and ta-siRNAs

Two miRNAs, miR162 and miR168, are involved in the regulation of the miRNA pathway itself by guiding cleavage of DCL1 and AGO1 mRNAs, respectively [Rhoades et al. 2002; Xie et al. 2003; Vazquez et al. 2004a]. Such feedback regulation of the two major actors of the miRNA pathway is probably essential to maintain a proper balance of miRNA steady-state levels, which depends on both miRNA production by DCL1 and miRNA turn-over during AGO1-catalyzed mRNA cleavage. Indeed, analysis of miRNA levels in ago1 mutants has revealed that most miRNAs were unstable in the absence of AGO1 protein, pointing to a stabilization role of AGO1 in addition to its catalytic role [Vaucheret et al. 2004]. Expression of a miR168-resistant AGO1 mRNA results in an increased AGO1 level and in increased accumulation of some miRNAs, in particular miR168, suggesting that AGO1 is normally limiting, and that the level of AGO1 is maintained in equilibrium by the dual effect of miR168 on AGO1 mRNA and of AGO1 protein on mir168 [Vaucheret et al. 2006].

Because ta-siRNAs are sequentially processed from long dsRNAs, which result from the transformation of the primary TAS single-stranded RNA [ssRNA] into dsRNA by RDR6, ta-siRNA duplexes contain both the strand corresponding to the primary TAS RNA and the strand complementary to the primary TAS RNA [Peragine et al. 2004; Vazquez et al. 2004b; Allen et al. 2005]. Like miRNAs, ta-siRNAs obey the asymmetry rule [the strand of the duplex that has the lowest stability at its 5’ end accumulates], whereas the passenger strand is degraded [Vazquez et al. 2004b; Allen et al. 2005]. Like miRNAs, mature ta-siRNAs deriving from the (+) strand [which corresponds to the primary TAS RNA] target genes that show little resemblance with the TAS genes [Peragine et al. 2004, Vazquez et al. 2004b, Allen et al. 2005]. However, mature ta-siRNAs deriving from the (−) strand [which is complementary to the primary TAS RNA] have the potential to target and regulate the primary TAS RNA. An overaccumulation of primary TAS RNA was observed in ta-siRNA-deficient mutants, pointing to a locus-specific feedback regulation that allows the maintenance of ta-siRNA steady-state levels [Vazquez et al. 2004b].

Open questions

How many miRNAs, ta-siRNAs, and nat-siRNAs exist?

MIR and TAS genes, as well as the cis-antisense gene pairs that generate nat-siRNAs, have some of the char-
characteristics of other plant genes; i.e., they are transcribed by DNA-dependent RNA polymerase II into capped and polyadenylated RNAs (Parizotto et al. 2004b; Allen et al. 2005; Xie et al. 2005a; Yoshikawa et al. 2005). However, because MIR and TAS genes do not contain obvious signatures such as the ORF of protein coding genes, they cannot be predicted with confidence from the genome sequence using this criterion alone. Although the minimal folding free energy index for miRNA precursors is significantly higher than tRNAs, rRNAs, or mRNAs (Zhang et al. 2006b), searching for genomic sequences that have the potential to form fold-back stem-loop structures similar to that of miRNA precursors is insufficient, as 138,864 inverted repeat structures exist in the Arabidopsis genome (Jones-Rhoades and Bartel 2004). Bioinformatic predictions that rely on the conservation of fold-back stem-loop structures between monocot and dicot species have been successful at identifying MIR genes that have been conserved after the divergence of monocot and dicot species (Jones-Rhoades and Bartel 2004). Young MIR genes that are species specific are more difficult to predict using bioinformatics tools. However, because MIR genes and their targets are assumed to derive from the same ancestor gene by inverted-repeat duplication, young MIR genes exhibit greater homology with their targets beyond the miRNA and miRNA complementary sequences (Allen et al. 2004), a criterion that could be helpful for identifying species-specific MIR genes. In addition, TAS genes have not been predicted successfully from the genome sequence because ta-siRNAs derive from ssRNAs that do not exhibit any known signatures that indicate how they are transformed into dsRNAs by RNA-dependent RNA polymerases. Cis-antisense gene pairs can be identified from annotated genomes; however, the rules that regulate the production of nat-siRNAs are not known. Therefore, the most reliable way to identify miRNAs, ta-siRNAs, and nat-siRNAs is the intensive cloning of small RNAs isolated from samples representing all tissues, all stages of life of the organism, and all conditions of growth, followed by the computational analysis of their arrangement within the genome. MIR genes can be confidently predicted when two collinear 21-nt small RNAs that are partially complementary are found 50–250 nt from each other, which is the size range of most MIR loops in plants, with a 10:1 ratio, which is the usual ratio between miRNA and miRNA*, owing to the degradation of miRNA* sequences, which are not stabilized in RISC. TAS genes can be confidently predicted when multiple small RNAs are found in 21-nt increments, originating from both DNA strands, including pairs of perfectly complementary molecules. Genetic confirmation of the nature of the small RNAs can be obtained in model species such as Arabidopsis or rice (Table 2). Intensive small RNA sequencing has been performed in Arabidopsis using MPSS; however, MPSS only provides a 17-nt sequence, which does not always allow the assignment of a confident genomic location to the small RNA because of the uncertainty about the last 4–7 nt. Nevertheless, 70,000 unique sequences were identified from 1,500,000 MPSS reads, revealing complexity to the small RNA repertoire in plants (Lu et al. 2005). However, this analysis did not reveal an abundant number of additional MIR or TAS genes. In addition, this set of small RNAs did not contain the nat-siRNAs because they are only expressed under particular conditions of growth (Borsani et al. 2005). It is likely that the sequencing of a larger number of small RNAs from other sources of material will reveal rare or tissue-specific miRNAs, ta-siRNAs, nat-siRNAs, and heterochromatin/repeat sequence siRNAs that have been missed so far.

### How are MIR and TAS genes regulated?

MicroRNAs and ta-siRNAs impose a regulation on their targets that has important consequences on gene expression and often determines changes in cell fate (Bartel 2004). Although the regulation of MIR and TAS genes is critical for proper regulation of their targets, little is known about the expression patterns of MIR and TAS genes in plants, whereas the tissue-specific expression of animal miRNAs has been extensively analyzed (Wienholds et al. 2005). In situ hybridizations have revealed that miR165/166 and miR172 accumulate in a tissue-specific manner (Chen 2004; Kidner and Martienssen 2004), however, the basis of this tissue-specific accumulation is unknown. Fusions between MIR164c and MIR171 promoters and reporter genes also have revealed tissue-specific expression, suggesting that part of the regulation of MIR164c and MIR171 genes occurs at the transcription level (Parizotto et al. 2004; Baker et al. 2005). Post-transcriptional regulations likely play additional roles. Indeed, mutations in the SERRATE (SE) gene encoding a zinc-finger protein result in miR165/166 precursor overaccumulation and reduced mature miR165/166 accumulation, indicating that SE is required for post-transcriptional regulation of MIR165 and MIR166 genes (Grigg et al. 2005). Clearly, most regulations of MIR and TAS genes remain to be identified.

### Are miRNAs, ta-siRNAs, and nat-siRNAs mobile?

Transgene PTGS mediated by siRNAs in plants is not cell autonomous. After local triggering, either spontaneously or artificially, PTGS spreads systemically from cell

| Table 2. Genetic requirements of endogenous small RNAs in plants |
|------------------------|------------------------|------------------------|------------------------|------------------------|------------------------|
| **Small RNA** | **Size (nt)** | **DCL1** | **DCL2** | **DCL3** | **DCL4** | **RDR2** | **RDR6** |
| miRNA | 21 | + | – | – | – | – | – |
| ta-siRNA | 21 | + | – | – | + | – | – |
| nat-siRNA | 24 | – | – | + | – | – | – |
| nat-siRNA | 21 | + | + | – | – | – | – |
| hc-siRNA | 24 | – | – | + | + | – | – |

The size indicated corresponds to the major accumulating species of small RNA. miRNA size ranges from 20 to 24 nt. ta-siRNA size ranges from 20 to 22 nt. hc-siRNA size ranges from 21 to 24 nt. A plus sign (+) indicates that the enzyme is required. A minus sign (−) indicates that the enzyme is dispensable.
to cell via plasmodesmata and at long distance through the vascular tissues (Palauqui et al. 1997; Voinnet et al. 1998). Although siRNAs have not been formerly identified as the mobile silencing signal, the 21-nt transgene-derived siRNAs produced by DCL4 are likely to be mobile and responsible for at least cell-to-cell propagation of PTGS because dcl4 mutants fail to exhibit short distance spreading of PTGS triggered by a vascular-specific promoter (Dunoyer et al. 2005). If transgene-derived 21-nt siRNAs are mobile, are endogenous 21-nt miRNAs, 21-nt ta-siRNAs, and 21-nt nat-siRNAs also mobile? The tissue-specific accumulation of miR165/166 and miR172 revealed by in situ hybridizations suggests that these miRNAs are not mobile (Chen 2004; Kidner and Martienssen 2004) or at least do not accumulate at distances far from their production, although there could be limited mobility within a given tissue. However, some miRNAs—miR156, miR159, and miR167—have been cloned from vascular tissues, suggesting that some miRNAs could move within certain parts of the plant (Yoo et al. 2004). It is possible that miRNAs are transported from a distant source and only accumulate within tissues where stabilizing factors are present. In addition, it is likely that small RNAs are not mobile as free molecules and that their mobility depends on their association with particular proteins, which itself could depend on their mode of biogenesis. Whether the tissue-specific accumulation of miRNAs, ta-siRNAs, or nat-siRNAs is modified in mutants impaired in the movement of transgene-derived siRNAs will likely shed light on this important question.

What is the contribution of translation repression versus mRNA cleavage?

MicroRNA-mediated translational repression in animals was originally proposed to have no effect on mRNA stability; however, recent reports have shown that mRNA accumulation is also affected (Bagga et al. 2005; Lim et al. 2005). In plants, miRNA-mediated regulation is assumed to occur almost exclusively by cleavage (Llave et al. 2002; Kasschau et al. 2003). However, protein levels have only been analyzed in the case of miR172-mediated regulation of AP2, and there was no correlation between AP2 mRNA and AP2 protein levels. Rather, the amount of AP2 mRNA was unchanged when the level of miR172 was either decreased (in dcl1 or hen1 mutants) or elevated (in plants overexpressing miR172 under the control of the strong 35S promoter), whereas the amount of AP2 protein inversely correlated with the amount of miR172 (Aukerman and Sakai 2003; Chen 2004). This prompted the proposal that miR172-mediated regulation of AP2 was occurring by translational repression, although AP2 had no distinguishing features when compared with the other miRNA targets, and although 5’ RACE indicated that, like other miR172 targets, AP2 mRNA was cleaved in the middle of the region of complementarity with miR172 (Aukerman and Sakai 2003; Kasschau et al. 2003; Lauter et al. 2005). Qualitative PCR using an AP2 cleavage-specific adapter showed that, although AP2 mRNA remained constant, cleavage actually was increased in 35S-MIR172 plants (Schwab et al. 2005). This led to the suggestion that AP2 was regulating its own transcription and that the unchanged AP2 mRNA level in 35S-MIR172 plants resulted from increased cleavage, which led to reduced AP2 protein levels and increased transcription of the AP2 gene. However, this hypothesis does not rule out the possibility that translational repression occurs in plants. Additional analyses of both mRNA and protein levels of other miRNA targets are required before this question can be answered. In addition, it is possible that plant miRNA targets that are regulated by translational repression have not been identified yet. Indeed, translational repression in animals is mediated by small segments of complementarity corresponding to the “seed” region of miRNAs; i.e., between bases 2 and 8 (Bartel 2004). Since the prediction of miRNA targets in plants is based on longer segments of complementarity with miRNAs, the existing list of miRNA targets does not contain genes with shorter segments of complementarity because they are not distinguishable from background noise in bioinformatics analyses (Rhoades et al. 2002; Jones-Rhoades and Bartel 2004). Genome-wide transcriptome analyses of plants overexpressing several miRNAs did not reveal an abundance of additional targets (Schwab et al. 2005). However, this may be because translational repression has only a subtle effect on mRNA accumulation. Comparing the transcriptome to the proteome in plants overexpressing miRNAs or in miRNA-deficient mutants may help identify targets regulated by translational repression.

Why are TAS cleavage products transformed into dsRNA after miRNA-mediated cleavage, whereas cleavage products deriving from other miRNA targets are not?

A unique feature of TAS transcripts is that after miRNA-guided cleavage, one of the two cleavage products is transformed into dsRNA by RDR6, and sequentially processed into 21-nt ta-siRNAs by DCL4 (Peragine et al. 2004; Vazquez et al. 2004b; Allen et al. 2005; Gasciolli et al. 2005; Xie et al. 2005b; Yoshikawa et al. 2005). Large-scale cloning of small RNAs from Arabidopsis has not revealed an abundance of small RNAs deriving from miRNA targets, except for the TAS transcripts targeted by miR173 and miR390, indicating that not all miRNA-derived cleavage products enter this pathway (Lu et al. 2005). The basis of the specificity of RDR6 for TAS-derived cleavage products is unclear, as is unclear the reason why only one TAS-derived cleavage products is transformed into dsRNA and why, depending on the TAS locus considered, either the 5’ or 3’ cleavage product is used as substrate. The fact that TAS RNAs are non-protein-coding RNAs, whereas other miRNA targets are translated into proteins does not seem to be the reason. Indeed, 35S-driven transgenes transcribing a translatable GFP RNA with a miRNA complementary site produces
siRNAs after miRNA-guided cleavage that lead to consistent PTGS of the *GFP* transgenes in wild-type transformants, whereas only miRNA-guided cleavage of *GFP* RNAs occurs in *rdr6* transformants (Parizotto et al. 2004). The production of GFP siRNAs does not seem to result exclusively from high levels of transcription due to the 35S promoter. Indeed, whereas introduction of an extra-copy of a wild-type gene usually does not trigger PTGS of the endogenous copy, introduction of an extra-copy of a wild-type gene usually does not trigger PTGS of the endogenous copy, but never occurs in *rdr6* transformants or in wild-type plants transformed with an *AGO1* gene mutated in the miR168 complementary site (Vaucheret et al. 2004). The subcellular location of the target genes may play a role by determining the ability of transcribed RNAs to become a substrate for RDR6. The accumulation of miR173 and miR390, the two miRNAs that target *TAS* primary RNAs, is unchanged in *hst* mutants (Xie et al. 2005a), suggesting that these miRNAs act in the nucleus. It is likely that the entire ta-siRNA pathway occurs in the nucleus because DCL4, which processes RDR6-derived dsRNAs, is located in the nucleus (Hiraguri et al. 2005) and because the accumulation of ta-siRNAs is unchanged in *hst* mutants (Allen et al. 2005; Park et al. 2005), suggesting that *TAS* primary transcripts are not exported to the cytoplasm. Other miRNA target mRNAs may be efficiently exported to the cytoplasm and translated into proteins or cleaved by miRNA-guided AGO1. If RDR6 is in the nucleus while cleavage products are in the cytoplasm, no ta-siRNAs can be produced. Only in ectopic cases where transgene-derived RNAs may not be efficiently exported to the cytoplasm due to their particular genomic location or high abundance does the probability that miRNA-guided cleavage occurs in the nucleus increase, leading to PTGS of homologous transgene and endogenous RNAs.

**How does DCL2 produce the initial 24-nt nat-siRNA?**

Cis-antisense gene pairs produce transcripts that have a long region of complementarity. In the example analyzed by Zhu and colleagues (Borsani et al. 2005), the processing of this long dsRNA by DCL2 produces only one 24-nt nat-siRNA, which subsequently guides the cleavage of one of the two transcripts to set the frame for the production of 21-nt nat-siRNAs by DCL1. DCL1-mediated production of 21-nt nat-siRNAs in frame with the 24-nt nat-siRNA-guided cleavage site resembles DCL4-mediated production of 21-nt ta-siRNAs in frame with the miRNA-guided cleavage site on the *TAS* transcripts. However, it is unclear how the cis-antisense gene pair-derived long dsRNA is processed into a single 24-nt nat-siRNA by DCL2. It is possible that DCL2 has the ability to perform sequence-specific endonucleolytic cleavages of partially folded miRNA precursors to produce miRNA duplexes. The sequence and structural requirements for DCL-mediated endonucleolytic cleavages remains to be determined.

**What is the role of siRNA- and miRNA-directed methylation?**

Endogenous or transgene-derived siRNAs that correspond to promoter sequences guide DNA methylation in homologous promoter sequences, which is assumed to be, at least in part, responsible for TGS (Mette et al. 2000; Matzke et al. 2001; Chan et al. 2004). In addition to guiding mRNA degradation during PTGS, transgene-derived siRNAs that correspond to transcribed sequences also guide DNA methylation in the homologous transgene sequence (Matzke et al. 2001; Vance and Voinnet 2001; Baulcombe 2004). However, methylation in the transcribed sequence does not seem to influence transcription because transgene transcription rates have been found similar in wild-type (silenced) plants and *ago1, rdr6*, or *sgs3* (nonsilenced) mutants (Mourrain et al. 2000; Morel et al. 2002). Therefore, the incidence and the role of methylation associated with PTGS remains unclear. Interestingly, *PHB* and *PHV* genes, which are regulated by miR165/166, are specifically methylated downstream of the miRNA complementary site (Bao et al. 2004). Indeed, *PHB* and *PHV* are not methylated in *serate* mutants that are defective in the production of miR165/166 and in *phb* and *phv* mutants carrying *PHB* or *PHV* genes mutated in the miRNA complementary site (Bao et al. 2004; Grigg et al. 2005). Heterozygous *PHB/phb* or *PHV/phv* plants that carry both a wild-type allele and a mutant target allele are only methylated in the wild-type allele, suggesting that DNA methylation occurs in *cis* and depends on the ability of the miRNA to bind to the transcribed RNAs (Bao et al. 2004). The incidence and the role of methylation associated with miRNAs also are unclear, as is unclear why methylation occurs downstream of the miRNA complementary site. Methylation-deficient mutants exist in Arabidopsis (Chan et al. 2005), but these mutants have not been reported to have developmental abnormalities that could result from defects in *PHB* or *PHV* expression; however, the methylation status of *PHB* and *PHV* has not been examined in these mutants. Whether other genes that are repressed by miRNAs are methylated is unknown. It also is unknown if ta-siRNAs or nat-siRNAs, which resemble transgene-derived siRNAs, trigger the methylation of the *TAS* genes or the cis-antisense gene pairs from which they derive, respectively, and if such methylation could play a role in their regulation.

**Why do viruses inhibit the miRNA pathway?**

Most viruses express proteins that are able to inhibit PTGS to some extent (Vance and Voinnet 2001; Baulcombe 2004; Dunoyer and Voinnet 2005; Li and Ding 2005). Coinoculation of a virulent virus (i.e., a virus that
strongly inhibits PTGS] with a mild virus [i.e., a virus that weakly inhibits PTGS] results in increased symptoms, a phenomenon known as synergism (Pruss et al. 1997). This observation has led to the hypothesis that virus-mediated counteraction of the plant PTGS defenses is necessary to establish efficient systemic infections (Vance and Vaucheret 2001; Baulcombe 2004; Dounoyer and Voinnet 2005; Li and Ding 2005), consistent with the accentuation of viral symptoms observed in PTGS-deficient mutants (Mourrain et al. 2000; Dalmay et al. 2001; Qu et al. 2005; Schwach et al. 2005). Viruses also inhibit transgene-induced PTGS because the production and action of transgene-derived siRNAs is likely similar to the production and action of virus-derived siRNAs (Vance and Vaucheret 2001; Baulcombe 2004; Dounoyer and Voinnet 2005; Li and Ding 2005). Viruses also inhibit the miRNA pathway to some extent [Mallooy et al. 2002; Kasschau et al. 2003; Chapman et al. 2004; Dounoyer et al. 2004]. Whether viruses inhibit the miRNA pathway because the miRNA and siRNA pathways share common components or because the miRNA pathway directly or indirectly impacts virus infection is unknown. Several near-perfect homologies between plant small RNAs and viral genomes have been reported, suggesting that viruses could be targeted by endogenous small RNAs [Llave 2004]; however, there are no experimental data supporting this hypothesis. Studies in human cells have revealed that an endogenous miRNA, miR32, targets the retrovirus PVF-1, suggesting that miRNAs may have an antiviral role [Lecellier et al. 2005]. On the other hand, attempts to control virus proliferation by introduction of artificial siRNAs into virus-infected human cells resulted in the appearance of mutations in the siRNA complementarity site of the virus that are compatible with the life cycle of the viruses [Li and Ding 2005]. Therefore, if it happened by chance that plant miRNAs were able to target viral RNAs, it is likely that the corresponding viruses would mutate to escape targeting in order to successfully infect plants [Simon-Mateo and Garcia 2006]. It also is possible that viruses inhibit the miRNA pathway because the up-regulation of some miRNA targets provides favorable conditions for virus proliferation. So far, none of the known miRNA targets have been shown to directly control virus proliferation. However, many miRNA targets encode transcription factors that regulate a large number of unknown genes (Rhoades et al. 2002; Jones-Rhoades and Bartel 2004), among which could be a gene controlling virus proliferation.

The questions asked in this review only represent a small fraction of the questions that researchers in the RNAi field are currently trying to answer. Given the formidable expansion of investigation in this recent subject, it is likely that many questions will find their answers in the near future through the crossing approaches of developmental biologists, biochemists, and virologists. However, the apparent complexity of small RNA-related pathways also increases as we gain further insights into their mechanisms, raising more and more questions.

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