

Detection of 91 potential conserved plant microRNAs in *Arabidopsis thaliana* and *Oryza sativa* identifies important target genes

Eric Bonnet^{*†}, Jan Wuyts^{*†}, Pierre Rouzé^{*‡}, and Yves Van de Peer^{*§}

^{*}Department of Plant Systems Biology, Flanders Interuniversity Institute for Biotechnology, and [†]Laboratoire Associé de l'Institut National de la Recherche Agronomique, Ghent University, Technologiepark 927, B-9052 Ghent, Belgium

Communicated by Marc C. E. Van Montagu, Ghent University, Ghent, Belgium, June 17, 2004 (received for review April 19, 2004)

MicroRNAs (miRNAs) are an extensive class of tiny RNA molecules that regulate the expression of target genes by means of complementary base pair interactions. Although the first miRNAs were discovered in *Caenorhabditis elegans*, >300 miRNAs were recently documented in animals and plants, both by cloning methods and computational predictions. We present a genome-wide computational approach to detect miRNA genes in the *Arabidopsis thaliana* genome. Our method is based on the conservation of short sequences between the genomes of *Arabidopsis* and rice (*Oryza sativa*) and on properties of the secondary structure of the miRNA precursor. The method was fine-tuned to take into account plant-specific properties, such as the variable length of the miRNA precursor sequences. In total, 91 potential miRNA genes were identified, of which 58 had at least one nearly perfect match with an *Arabidopsis* mRNA, constituting the potential targets of those miRNAs. In addition to already known transcription factors involved in plant development, the targets also comprised genes involved in several other cellular processes, such as sulfur assimilation and ubiquitin-dependent protein degradation. These findings considerably broaden the scope of miRNA functions in plants.

comparative genomics | thale cress | rice | noncoding RNA

MicroRNAs (miRNAs) are small noncoding RNA gene products ≈ 22 nt long that are found in a variety of organisms, including animals and plants (1–3). The first miRNAs were discovered in *Caenorhabditis elegans* and control developmental timing by binding to specific target mRNAs (4–9). By pairing to mRNAs, documented miRNAs initiate cleavage of the mRNA or repress active translation (1–3, 10). miRNAs have similarities with small interfering (si)RNAs, which are short (21–24 nt) molecules involved, at least in plants, in posttranscriptional gene silencing (11). In this process, long, double-stranded RNAs are cleaved into siRNA fragments by DICER, an RNase III helicase (12). These fragments bind to complementary RNA, which activates the RNA interference silencing complex to cleave the target mRNA (13). One main difference between siRNAs and miRNAs is that siRNAs mediate the silencing of the genes at the locus from which they originate, and possibly their paralogs, whereas miRNAs regulate a wide variety of genes that differ from those of which they originated (1, 14–16).

In plants, as in animals, miRNAs are processed from transcripts that can fold into a stable hairpin (17, 18). However, the size of the potential hairpin precursors of the plant miRNAs is much more variable than that of animals. For example, *C. elegans* miRNAs are cleaved from precursors of ≈ 70 nt in length, with the mature miRNA located from 2 to 10 nt from the terminal loop of the stem-loop structure (19). Although some of the *Arabidopsis* precursors resemble those of *C. elegans*, others are much larger, such as the 190-nt-long precursor of mir169 (10). In addition, the shape of the predicted secondary structure of plant miRNAs appears to be more complex, sometimes with branched structures instead of a simple hairpin. Unlike animal miRNAs, plant miRNAs generally interact with their targets through near-perfect complementarity

(17–19), which greatly facilitates the computational identification of plant miRNA targets. For example, for described plant miRNAs (20, 21), 49 unique targets could already be identified. Later, many of these targets have been confirmed experimentally (15, 16, 22–24). Recent work (25, 26) also demonstrated the role of miRNAs in the control of leaf, stem, and flower development.

Different biochemical approaches were used to try to identify small RNAs in plants, but so far only a fraction of them could be verified. For example, Park *et al.* (21) identified 230 sequences of which only five appeared to be “true” miRNAs. With a protocol to preferentially clone DICER cleavage products, only 16 true miRNAs were isolated from a starting pool of 300 small RNAs of seedlings and flowers (18). From this set of 16, eight were also conserved in the rice (*Oryza sativa*) genome. Interestingly, the sequences adjacent to these miRNAs could form stem-loop structures analogous to those of *Arabidopsis*, with the miRNA sequence invariably on the same arm of the precursor in both species. Furthermore, although the *Arabidopsis* and rice sequences seemingly diverged considerably upstream and downstream of the miRNA, the miRNA itself differed in only a few base pairs. This conservation of secondary structure, despite the sequence variability observed in the precursor sequences, suggests that the secondary structure plays a major role, presumably in the processing of the mature miRNA from the precursor.

To estimate the numbers of potential miRNAs in organisms such as *Caenorhabditis*, *Drosophila*, fish, and human, different computational gene-finding strategies have been developed (27–30), all of them being based on a comparative approach. The core principle is to look for conserved sequences between different species that can fold into extended hairpins. The fact that there are much more conserved stem-loop structures than true miRNA genes stresses the importance of considering additional information to validate potential miRNA (29). Unfortunately, methods applied to detect animal miRNAs cannot be directly applied to plants. For example, the algorithms that used a fixed-length window to search for hairpins in intergenic sequences are justified in animals because most animal miRNA precursors have almost identical lengths (≈ 70 –80 nt). On the contrary, plants show a much greater length variability of miRNA precursors, invalidating this fixed-window approach. Additionally, whereas animal miRNAs are conserved for most of the complete precursor sequence, in plants only the mature miRNA is conserved (20).

Here, we propose a computational approach for detecting plant miRNAs, based on a comparison of the *Arabidopsis* and *Oryza* genomes, and considering the core features of experimentally determined plant miRNAs. By using this approach, we were able to identify many miRNA genes and their targets.

Abbreviations: IGR, intergenic region; miRNA, microRNA; TF, transcription factor; NAM, no apical meristem.

[†]E.B. and J.W. contributed equally to this work.

[§]To whom correspondence should be addressed. E-mail: yves.vandeppeer@psb.ugent.be.

© 2004 by The National Academy of Sciences of the USA

Materials and Methods

miRNA Reference Set. To derive a set of rules and parameters that describe and characterize known *Arabidopsis* miRNAs, we first defined a reference set of miRNA sequences, which consisted of 22 *Arabidopsis* miRNA sequences and their corresponding 43 precursor sequences. The difference between 22 and 43 is due to the fact that some identical miRNAs are found at different places in the genome and with different precursors. These sequences were obtained from different *in vivo* biochemical analyses, such as accumulation of miRNA in mutants and expression levels of miRNA sequences by northern analysis (18, 20, 21, 31), and were downloaded from the miRNA registry (32).

***Arabidopsis* Intergenic Sequences.** Sequences and annotations were downloaded from The Institute for Genomic Research (*Arabidopsis thaliana* release 3, July 2002). Because most known miRNA genes have been detected in intergenic regions (IGRs) (1) the study was limited to the *Arabidopsis* IGR sequences that were extracted, excluding known elements, such as protein-encoding genes plus their experimentally defined UTRs, pseudogenes, ribosomal RNA, small nucleolar RNA, and tRNAs. For protein-encoding genes without experimentally defined UTR, a 300-nt region, i.e., the average length for experimentally defined UTRs with a full-length cDNA sequence, was added both upstream and downstream. A total of 23,433 sequences were thus obtained with an average length of 2 ± 1.9 kb.

***O. sativa* Sequences.** In total, 3,601 rice bacterial artificial chromosome sequences from the International Rice Genome Sequencing Project (33) were downloaded from The Institute for Genomic Research assemblies (June 2003). The average length of the sequences was 138 ± 30 kb.

Conserved Intergenic Short Segments. *Arabidopsis* intergenic sequences were masked for repeat elements by using the program REPEATMASKER (<http://ftp.genome.washington.edu/RM/RepeatMasker.html>). Conserved IGRs between *Arabidopsis* and rice were obtained with BLAST (34), by using a bit-score low-cutoff value set at 30 bits. This cutoff value allows a limited dissimilarity between sequences, which is in agreement with what is observed in the miRNAs of the reference set (18). BLAST hits between *Arabidopsis* and rice with a length between 20 and 25 nt were retained for further analysis. To consider the secondary structure of the miRNAs precursors, conserved sequences were extended with 350 nt both upstream and downstream, which is the maximum length known for *Arabidopsis* miRNA sequences (18). Analyses of the reference set showed that the miRNA sequence could be located on either the 5' or 3' arm of the precursor sequence, without any preference for strand orientation. Consequently, the reverse complement of the sequences was also considered when the miRNAs were on the reverse strand.

Removing Vector Contamination and Known Noncoding RNAs. Potential miRNA precursors were carefully checked for repeat sequences, for vector contamination with the UNIVEC database (www.ncbi.nlm.nih.gov/VecScreen/VecScreen.html), for virus sequences (35), and also for homology with well known other types of noncoding RNAs, such as tRNA, ribosomal RNA, and small nucleolar RNA. The tRNA, ribosomal RNA, and small nucleolar RNA sequences were downloaded from the GtRDB database (36), the European ribosomal RNA database (37), and the small nucleolar RNA database (38), respectively. Sequence matches with a low *e*-value (<0.01) were discarded.

GC Content and Low-Complexity Filtering. We analyzed BLAST hits from IGRs of the genome, which are well known to contain a large amount of repeat and low-complexity sequences, thus giving many

false-positive conserved sequences. To reduce the number of false positives, a filter based on GC content and low complexity was applied to potential miRNA sequences. A Shannon entropy measure (39) was used as a low-complexity pattern filtering. The cutoff values were defined based on the distribution of the values obtained with the miRNA reference set. Sequences with a GC content ≥ 0.3 and ≤ 0.7 and with an entropy value ≥ 1.75 were retained for further analysis.

Potential miRNA Precursor and Precursor Secondary Structure. To define the potential stem-loop precursors within the extended sequences, we made use of the characteristic miRNA precursor molecules property that the mature miRNA is always excised from one side of an unbranched RNA helix (40). As a result, a sequence that corresponds to the reverse complement of the miRNA should be found on the complementary side of the mature miRNA. Therefore, for each potential miRNA, the reverse complement was aligned to the extended precursor molecule by using the local alignment algorithm implemented in the matcher program from the EMBOSS package (41). We modified the scoring matrix to allow G-U and U-G base pairs in the precursor pairing. These parameters were chosen so that all miRNAs in our reference set were identified. Each reverse complement was checked against the cutoff values that were extracted from the reference set. After this step, all sequences flanked by the miRNA and its reverse complement were extracted as potential miRNA precursors. Potential precursor sequences were all folded with the VIENNARNA package (42). The statistical significance of the folding of the miRNA precursor was assessed with a randomization test (see *Supporting Text*, which is published on the PNAS web site, for a detailed explanation).

Potential Targets. Because most plant miRNA sequences bind to their target with a near-perfect complementarity (22–24), this property was exploited to predict potential targets for miRNA sequences by using a computational approach (20). By using exactly the same procedure, potential target mRNAs were searched for by the PATSCAN software of Dsouza *et al.* (43). The validity was tested on the same data set as described in Rhoades *et al.* (20). The number of allowed mismatches varies with the length of the potential miRNA: two, three, and four mismatches for sequences of up to 21 nt, 23 nt, or more, respectively. This variable rule set took into account the possible diffuse nature of the boundaries of the miRNA sequence, which occurs more probably in longer sequences. Hits with four mismatches should be considered with caution because such hits have a great likelihood to occur by chance, although they might be genuine targets (20). The mRNA sequences (coding sequences including UTR sequences when available; version 04/17/2003) were downloaded from the *Arabidopsis* Information Resource database (44).

Clustering of miRNA Sequences. MiRNA sequence similarity was estimated with a procedure similar to that used by Grad *et al.* (30). A pairwise homology score between all of the sequences was computed with the matcher program from the EMBOSS package (41). All pairwise comparisons were converted to Euclidean distances and a hierarchical clustering was performed with the single-linkage method. A cutoff was set to group highly similar sequences. Statistical computations were carried out with the R package (www.r-project.org).

Expression Data. Potential miRNAs were checked against EST sequences (206,678 sequences downloaded from EMBL) and *Arabidopsis* Massively Parallel Signature Sequencing (mpss.udel.edu/at; ref. 45) for 20-nt signatures.

All supporting information cited in the text can be accessed at www.psb.ugent.be/bioinformatics.

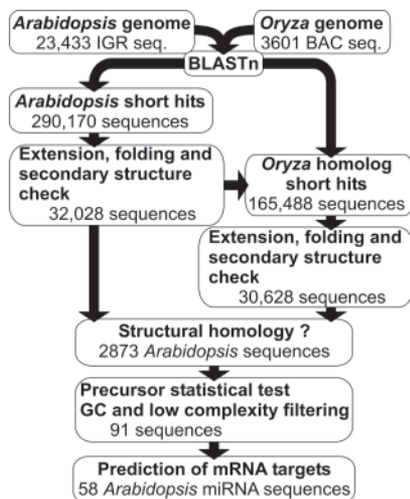


Fig. 1. Overview of the MIRFINDER computational pipeline to detect miRNAs in plants. For details, see text.

Results and Discussion

The MIRFINDER Computational Pipeline. A flowchart describing the general procedure of plant miRNA detection is shown in Fig. 1. The MIRFINDER computational pipeline was based on three major rules derived from the miRNA reference set: (i) the miRNA sequence is conserved between *Arabidopsis* and rice, whereas the rest of the precursor sequence has diverged; (ii) even though the precursor sequence has diverged, the ability of the precursor sequence to form a stem-loop secondary structure in both *Arabidopsis* and *Oryza* is conserved; and (iii) for two miRNA orthologs, the miRNA sequence is always located on the same arm of the stem-loop secondary structures in both species.

To select valid miRNA stem-loop structures, in addition to the three general rules described above, five characteristic features of the secondary structure of miRNA and their cutoff values were also derived from the reference set, of which one qualitative and five quantitative parameters. These parameters are (see Fig. 3, which is published as supporting information on the PNAS web site): (i) the miRNA should be part of a continuous helix; (ii) the minimum free energy value should be less than -30 kcal/mol; (iii) the minimum number of paired residues in the miRNA should be 15; (iv) the maximum number of unpaired residues in both the miRNA coding and complementary strand should be 5; and (v) the maximum number of G-U pairs in the miRNA should be 5.

The MIRFINDER pipeline (Fig. 1) starts by identifying short highly conserved sequences between the *Arabidopsis* and rice genomes. The 290,170 short sequences found in *Arabidopsis* with at least one corresponding hit in rice were extended to see whether they could form potential stem-loop secondary structures, which were determined by looking at the possible reverse complement of the miRNA within each extended sequence. Thus, for one potential miRNA sequence conserved between *Arabidopsis* and *Oryza*, multiple potential precursor sequences could be present. A total of 1,394,939 potential precursor *Arabidopsis* sequences were fold with the VIENNARNA package (42). From this pool, 32,648 sequences could be retained after filtering for typical miRNA precursor secondary structure features (see *Materials and Methods*). To verify whether these sequences have at least one homolog in rice, the same procedure was applied to the homologous sequences found in that genome. Every *Arabidopsis* stem-loop structure and its ortholog in *Oryza* were compared. The potential miRNA was retained only when an *Arabidopsis* miRNA sequence had an miRNA homolog in the rice genome that was located on the same arm (either 5' or 3') of the stem-loop structure in both species. A total of 2,873 *Arabi-*

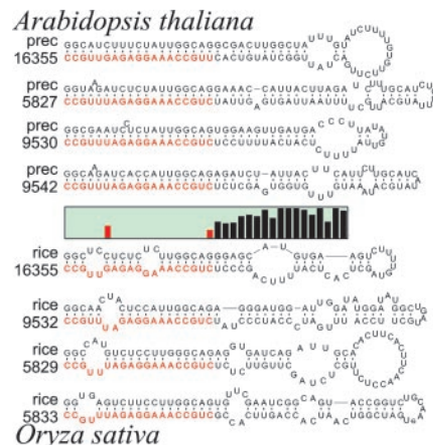


Fig. 2. RNA secondary structure models of the eight precursor molecules (four each from *Arabidopsis* and rice) of the miRNAs that target the 5' UTR of the *Arabidopsis* gene At2g33770.1 and its rice homolog. The precursor molecules are drawn with the miRNAs aligned and highlighted in red. (Middle) The graph represents the Shannon entropy of the nucleotide content of each corresponding position on the 3' strand of the precursors. A low value means that all molecules have the same nucleotide at that position, whereas a high value reflects little or no nucleotide conservation. The sequence divergence outside of the mature miRNA positions is clearly shown. RNA sequences are drawn from 5' to 3' in clockwise orientation. All precursors are truncated at the same position. RNA secondary structure drawings were made by using RNAVIZ (59).

dopsis sequences were found to have at least one such structural homolog in the *Oryza* genome. Examples of such miRNAs (Fig. 2) clearly show the conservation for the miRNA sequence, the non-conservation of the precursor sequence, and the conservation of the secondary structure. The use of GC content filtering (30) further reduced the set of 2,873 *Arabidopsis* potential miRNA to 852 sequences. A test study showed that $\approx 5\%$ of randomly selected genomic sequences from *C. elegans* could fold into a plausible miRNA precursor hairpin (19). To reduce the number of such false positives, all *Arabidopsis* precursor sequences were statistically tested with the randomization test (see *Supporting Text*). With this approach, the set of potential miRNA sequences was further reduced to 501. Last, an entropy-based filtering step removed miRNA sequences with a low-complexity pattern and further reduced our data set to 297 sequences.

Because an exhaustive approach was used, some potential miRNAs differed only by a few base pairs on either side, or were the reverse complement of each other. To avoid redundancy and overprediction, we clustered the overlapping miRNAs, with human supervision, and kept parent only when they complied with the features discussed above, also taking into account potential targets. After this clustering process, and after removing one potential miRNA because of unexpected sequence degeneracy (see below), 91 were kept as potential miRNAs. We could not observe any close cluster of miRNA sequences as it is the case for animal miRNAs (19). miRNA fasta files are available as Data Sets 1 and 2, which are published as supporting information on the PNAS web site.

A BLAST search against *Arabidopsis* Massively Parallel Signature Sequencing 20-nt signatures did not uncover any hits. The *Arabidopsis* precursor sequences gave two significant hits with ESTs, namely MIR30 and MIR36 and EST sequences BX838271 and AU239920, respectively, both with very high P value ($< 1e-45$). MIR30 (synonym of mir171) was already known to be expressed (18).

***Arabidopsis* miRNA Targets.** We could identify 58 *Arabidopsis* miRNA sequences that have at least one mRNA target sequence (see Table 1). One miRNA was excluded from this set because it

Table 1. List of the potential protein targets for the 91 potential miRNAs, grouped by annotated gene family and by biological function

Protein annotation	MiRNA	Common function/type
CCAAT box-binding factor	MIR13, MIR43, MIR47, MIR57, MIR81, MIR87	
CCAAT box-binding factor Hap2a	MIR77	Core transcription factor
CCAAT-binding factor B subunit homolog	MIR13, MIR43, MIR47, MIR57	
CCAAT-binding factor B subunit-related	MIR13, MIR43, MIR57, MIR77, MIR81, MIR87	
Auxin-response transcription factor (ARF)	MIR75	
ARF8	MIR54	
Transcription factor B3 family similar to ARF10	MIR40, MIR75	
Transcription factor B3 family	MIR40, MIR47	
NAC1/NAM protein family	MIR29, MIR82	Auxin signaling
NAM-related protein	MIR29, MIR82	
NAM-like protein	MIR29, MIR82	
NAM-related protein	MIR82	
NAM protein CUC2	MIR82	
Scarecrow transcription factor family	MIR30, MIR46, MIR58	Asymmetric cell division
Scarecrow-like transcription factor 14 (SCL14)	MIR14	Specify meristem quiescent Center and radial patterning
SCL6	MIR30, MIR46, MIR58	
Squamosa promoter-binding protein-related 2	MIR17, MIR85, MIR91	
Squamosa promoter-binding protein homologs	MIR17, MIR63, MIR85, MIR91	
Squamosa promoter-binding protein 4 (SPL4)	MIR17, MIR43, MIR63, MIR85, MIR91	Homeotic MADS box genes controlling floral development
AP2 domain transcription factor RAP2.7	MIR21, MIR84	
AP2 domain transcription factor, potential	MIR18, MIR21, MIR84	
Floral homeotic protein APETALA2	MIR18, MIR21, MIR84	
HD-Zip transcription factor Athb-15	MIR5, MIR70	Vascular development and leaf development
Phabulosa HD-Zip TF Athb-14	MIR70	
HD-Zip transcription factor Athb-9	MIR70	
PIL6 Myc-related bHLH transcription factor	MIR32	Circadian rhythm control by light
Myb family transcription factor MYB33	MIR89	
Myb family transcription factor MYB30, MYB120, MYB65	MIR88	Myb family transcription factor
Zinc finger (C3HC4-type RING finger) family	MIR12, MIR44, MIR56, MIR76, MIR80, MIR86	Zinc finger transcription factor
TIR1, E3 ubiquitin ligase SCF complex F-box subunit	MIR10	
TIR1-like genes, E3 ubiquitin ligase F-box subunit	MIR10, MIR20	Ubiquitination pathway
F-box protein GRR1-like protein 1, AtFBL18	MIR10	Controlling targeted
E2 UBC, ubiquitin-conjugating enzyme family	MIR16, MIR27, MIR67	Protein degradation (auxin-related for many)
CDC48 domain-containing AAA-type ATPase/NSF	MIR2	
TAZ RING BTB/POZ domain protein	MIR47	
F-box protein family	MIR63	
FLU, TPR-containing protein	MIR15	Chloroplast import of PORA
VQ-motif containing protein	MIR60	Control of plastid genes
Thioredoxin-like protein 3	MIR59	Redox control in chloroplast
ATP sulfurylase	MIR64	Sulfur metabolism
Sulfate transporter	MIR64	
Acyl transferase-containing multifunctional enzyme	MIR64	

Table 1. (continued)

Protein annotation	MiRNA	Common function/type
Copper/zinc superoxidase dismutase (CSD1)	MIR1	Cell death
Potential cytosine/deoxycytidine deaminase	MIR55	Purine salvage
Amine oxidase-related	MIR83	Overlaps auxin gene NAC1
Ypt/Rab GTPase-activating protein	MIR4	Intracellular trafficking
WASP domain containing-protein	MIR11	Signal transduction to actin
SAG101, leaf senescence-associated acyl hydrolase	MIR73	Senescence
Target of rapamycin, phosphatidylinositol 3-kinase	MIR34	Embryo development
Potential (glycerol) phosphate acyltransferase	MIR31, MIR72	Phospholipid synthesis
Laccase (diphenol oxidase), potential	MIR9	
Luciferin-rich repeat receptor kinase, potential	MIR55	
ABC transporter family protein	MIR51	Specific function unknown
Proline-rich protein family	MIR53	
Expressed protein	MIR21	
Expressed protein	MIR25	
Hypothetical protein	MIR6, MIR15	
Gypsy family retrotransposon gag protein	MIR88	Retrotransposon

An extended version of this table, which includes the gene loci (At codes) for each protein, appears as Table 2, which is published as supporting information on the PNAS web site.

had 76 targets, which is very unlikely. The highly repetitive motif corresponding to this sequence (CUUCAUCUUCAUCAUCAUCAG) might explain such a ubiquity, thus revealing a potential false positive.

The overall sensitivity of our procedure is demonstrated by the fact that we could identify six of the eight described miRNAs that are known to be conserved between *Arabidopsis* and rice, namely MIR156, MIR160, MIR164, MIR166, MIR167, and MIR171 (18, 46). The reason why we missed the two other miRNAs will be discussed below. If a hierarchical clustering approach were applied, the 91 *Arabidopsis* miRNAs would be clustered in 51 families (dendrogram and list are available as Fig. 4, which is published as supporting information on the PNAS web site).

The different mRNA targets were grouped into families (Table 1), of which 56% of all potential targets represent transcription factor (TF)-related proteins. Many of the TF targets listed were already known to be targets of miRNAs (1, 10). For example, we successfully identified both the previously described *Arabidopsis* mir171 (MIR30 in Table 1) and its targets, the Scarecrow TF family (At2g45160, At3g60630, and At4g00150). The function of mir171 is supported by both prediction and experimental evidence (17, 18, 20, 31). A few other TF targets, to our knowledge, are new, such as the Zn-finger protein family (At1g54150 in Table 1).

Interestingly, we discovered that almost half of the predicted targets are non-TF mRNAs. As shown in Table 1, the function of most of these targets is known, at least to some extent, and some can be clustered according to this function and/or their corresponding miRNAs into larger functional groups. The largest one, with *TIR1* as a typical member, comprises some of the many genes of the ubiquitination pathway, involved in proteasome-dependent degradation of targeted proteins (47), which plays an important role in plant development (48). *TIR1* is a component of the E3 ubiquitin ligase SCF^{TIR1} complex that mediates auxin response through the Aux/IAA proteins, which act as negative regulators. *TIR1* itself is responsible for the specificity of the complex and binds the Aux/IAA proteins that need to be destroyed (47, 48). This interaction is

promoted by auxin (49). The fact that *TIR1* mRNA is targeted by the potential miRNAs MIR10 and MIR20 adds a further layer of negative regulation to the auxin pathway. Also, besides the genes in the ubiquitination pathway, a large group of auxin-related TFs, including auxin-response transcription factors and no apical meristems (NAMs; refs. 50 and 51), are present that are also potential targets of miRNAs (Table 1). Altogether, there is strong suggestive evidence that miRNAs play a major role in auxin signaling. To validate prediction of MIR20, we found homologs with perfectly conserved miRNA sequences with a valid stem-loop precursor structure in three plant genomes (Fig. 5, which is published as supporting information on the PNAS web site), namely *Medicago truncatula* (working draft sequences downloaded from the EMBL database), *Populus trichocarpa* (public reads downloaded from <http://genome.jgi-psf.org>), and *Lotus corniculatus* (sequences downloaded from EMBL).

Another interesting example is the potential miRNA MIR64 that targets three different gene families (Table 1) involved in sulfur metabolism. The first target encodes ATP sulfurylase (APS), the first enzyme in the sulfate assimilation pathway, which is present both in the cytosol and the chloroplast. Of the four APS proteins present in *Arabidopsis* (52), MIR64 can pair with the mRNAs of only three of them, and its target sequence is indeed missing from the fourth, *APS2* (At1g19990). MIR64 also targets one of the three *Arabidopsis* sulfate transporters, namely that with low affinity induced in plant roots by sulfate starvation (53, 54). Altogether, MIR64 probably plays a specific role in the control of sulfur assimilation. The potential MIR1 targets a Cu/Zn superoxidase dismutase (CSD1; ref. 55), which is one of the main markers of programmed cell death in *Arabidopsis* (56). AtTOR, an ortholog of the yeast target of rapamycin involved in cell-cycle regulation during embryo development (57, 58) is also a miRNA target of interest.

False-Positives and -Negatives. We have tried to reduce the number of false-positives by combining multiple procedures and an overall

conservative approach. First, the number of potential miRNAs was strongly reduced by considering the miRNA secondary structure parameters based on the reference set. By comparing this data set with the rice genome, this number could be further lowered. The combination of a statistical test on the precursor sequences and GC and low-complexity filtering on the miRNA sequences further reduced the number of candidates by one order of magnitude.

The apparently missing targets for part of our candidates (58 of 91) could be explained by the less perfect than previously thought complementarity with mRNA target sequences for some plant miRNAs, as is the case for metazoan miRNAs (1–3). Recently, 20 miRNAs were experimentally identified in rice (46), and only seven had targets with near-perfect complementarity, which suggests that pairing between miRNAs and their target mRNAs can be less stringent in plants.

In contrast, a fraction of the true miRNA sequences might have been discarded although they are true miRNAs. A sequence that is conserved outside the strict limit of the miRNA sequence might have a hit length of more than the used upper threshold value (25 nt). As mentioned, we identified six of the eight miRNAs previously known to be conserved between *Arabidopsis* and rice (18). For these two missing miRNAs (mir162 and mir169), the conserved regions seem indeed longer than 25 nt and, as a result, are not reported in our approach. We hope that refinements of the computational pipeline used here for the prediction of miRNA may cope with this difficulty in the future.

In conclusion, the stringent search conditions that we applied have, in addition to most of the previously described miRNA genes conserved between *Arabidopsis* and *Oryza*, uncovered a considerable number of unknown miRNAs and their targets. If these miRNAs could be confirmed by experimental evidence, the number of different processes and pathways regulated by miRNAs in *Arabidopsis* would significantly increase. We would also like to stress that we used a deliberately conservative approach that retained only those potential miRNAs that complied with every feature of experimentally confirmed plant miRNAs. Many of the more recently reported rice miRNAs (46) do not fulfill all these criteria, suggesting that miRNAs in the *Arabidopsis* genome are probably more abundant than reported here and that additional experimental and *in silico* studies are needed.

Note Added in Proof. Upon acceptance of this paper, Jones-Rhoades and Bartel (60) reported several miRNAs identical to those we found, thereby confirming our approach. In addition, 34 miRNAs from our list matched perfectly with expressed small RNAs deposited in the *Arabidopsis* Small RNA Project Database, which can be accessed at <http://cgrb.orst.edu/smallRNA>.

We thank Ivo Hofacker for sharing parts of his software code and many members of our research group for helpful discussions. This work was supported by an Instituut voor de aanmoediging van Innovatie door Wetenschap en Technologie in Vlaanderen postdoctoral fellowship (to J.W.).

- Bartel, D. P. (2004) *Cell* **116**, 281–297.
- Carrington, J. C. & Ambros, V. (2003) *Science* **301**, 336–338.
- Lai, E. C. (2003) *Curr. Biol.* **13**, R925–R936.
- Lee, R. C., Feinbaum, R. L. & Ambros, V. (1993) *Cell* **75**, 843–854.
- Wightman, B., Ha, I. & Ruvkun, G. (1993) *Cell* **75**, 855–862.
- Moss, E. G., Lee, R. C. & Ambros, V. (1997) *Cell* **88**, 637–646.
- Reinhart, B. J., Slack, F. J., Basson, M., Pasquinelli, A. E., Bettinger, J. C., Rougvie, A. E., Horvitz, H. R. & Ruvkun, G. (2000) *Nature* **403**, 901–906.
- Abrahante, J. E., Daul, A. L., Li, M., Volk, M. L., Tennessen, J. M., Miller, E. A. & Rougvie, A. E. (2003) *Dev. Cell* **4**, 625–637.
- Lin, S. Y., Johnson, S. M., Abraham, M., Vella, M. C., Pasquinelli, A., Gamberi, C., Gottlieb, E. & Slack, F. J. (2003) *Dev. Cell* **4**, 639–650.
- Bartel, B. & Bartel, D. P. (2003) *Plant Physiol.* **132**, 709–717.
- Hamilton, A. J. & Baulcombe, D. C. (1999) *Science* **286**, 950–952.
- Bernstein, E., Caudy, A. A., Hammond, S. M. & Hannon, G. J. (2001) *Nature* **409**, 363–366.
- Hannon, G. J. (2002) *Nature* **418**, 244–251.
- Mallory, A. C. & Vaucheret, H. (2004) *Curr. Opin. Plant Biol.* **7**, 120–125.
- Vazquez, F., Gascioli, V., Cr  t  , P. & Vaucheret, H. (2004) *Curr. Biol.* **14**, 346–351.
- Boutet, S., Vazquez, F., Liu, J., Beclin, C., Fagard, M., Grati  s, A., Morel, J. B., Crete, P., Chen, X. & Vaucheret, H. (2003) *Curr. Biol.* **13**, 843–848.
- Llave, C., Kasschau, K. D., Rector, M. A. & Carrington, J. C. (2002) *Plant Cell* **14**, 1605–1619.
- Reinhart, B. J., Weinstein, E. G., Rhoades, M. W., Bartel, B. & Bartel, D. P. (2002) *Genes Dev.* **16**, 1616–1626.
- Lau, N. C., Lim, L. P., Weinstein, E. G. & Bartel, D. P. (2001) *Science* **294**, 858–862.
- Rhoades, M. W., Reinhart, B. J., Lim, L. P., Burge, C. B., Bartel, B. & Bartel, D. P. (2002) *Cell* **110**, 513–520.
- Park, W., Li, J., Song, R., Messing, J. & Chen, X. (2002) *Curr. Biol.* **12**, 1484–1495.
- Kasschau, K. D., Xie, Z., Allen, E., Llave, C., Chapman, E. J., Krizan, K. A. & Carrington, J. C. (2003) *Dev. Cell* **4**, 205–217.
- Tang, G., Reinhart, B. J., Bartel, D. P. & Zamore, P. D. (2003) *Genes Dev.* **17**, 49–63.
- Llave, C., Xie, Z., Kasschau, K. D. & Carrington, J. C. (2002) *Science* **297**, 2053–2056.
- Emery, J. F., Floyd, S. K., Alvarez, J., Eshed, Y., Hawker, N. P., Izhaki, A., Baum, S. F. & Bowman, J. L. (2003) *Curr. Biol.* **13**, 1768–1774.
- Aukerman, M. J. & Sakai, H. (2003) *Plant Cell* **15**, 2730–2741.
- Lim, L. P., Glasner, M. E., Yekta, S., Burge, C. B. & Bartel, D. P. (2003) *Science* **299**, 1540.
- Lim, L. P., Lau, N. C., Weinstein, E. G., Abdelhakim, A., Yekta, S., Rhoades, M. W., Burge, C. B. & Bartel, D. P. (2003) *Genes Dev.* **17**, 991–1008.
- Lai, E. C., Tomancak, P., Williams, R. W. & Rubin, G. M. (2003) *Genome Biol.* **4**, R42.1–R42.20.
- Grad, Y., Aach, J., Hayes, G. D., Reinhart, B. J., Church, G. M., Ruvkun, G. & Kim, J. (2003) *Mol. Cell* **11**, 1253–1263.
- Mette, M. F., van der Winden, J., Matzke, M. & Matzke, A. J. (2002) *Plant Physiol.* **130**, 6–9.
- Griffiths-Jones, S. (2004) *Nucleic Acids Res.* **32**, D109–D111.
- Sasaki, T. & Burr, B. (2000) *Curr. Opin. Plant Biol.* **3**, 138–141.
- Altschul, S. F., Gish, W., Miller, W., Myers, E. W. & Lipman, D. J. (1990) *J. Mol. Biol.* **215**, 403–410.
- Peterson-Burch, B. D. & Voytas, D. F. (2002) *Mol. Biol. Evol.* **19**, 1832–1845.
- Lowe, T. M. & Eddy, S. R. (1997) *Nucleic Acids Res.* **25**, 955–964.
- Wuyts, J., Perri  re, G. & Van de Peer, Y. (2004) *Nucleic Acids Res.* **32**, D101–D103.
- Brown, J. W., Echeverria, M., Qu, L. H., Lowe, T. M., Bachellerie, J. P., Huttenhofer, A., Kastenmayer, J. P., Green, P. J., Shaw, P. & Marshall, D. F. (2003) *Nucleic Acids Res.* **31**, 432–435.
- Shannon, C. E. (1948) *Bell Syst. Tech. J.* **27**, 379–423.
- Ambros, V., Bartel, B., Bartel, D. P., Burge, C. B., Carrington, J. C., Chen, X., Dreyfuss, G., Eddy, S. R., Griffiths-Jones, S., Marshall, M., et al. (2003) *RNA* **9**, 277–279.
- Rice, P., Longden, I. & Bleasby, A. (2000) *Trends Genet.* **16**, 276–277.
- Hofacker, I. L., Fontana, W., Stadler, P. F., Bonhoeffer, L. S., Tacker, M. & Schuster, P. (1994) *Monatsh. Chem.* **125**, 167–188.
- Dsouza, M., Larsen, N. & Overbeek, R. (1997) *Trends Genet.* **13**, 497–498.
- Huala, E., Dickerman, A. W., Garcia-Hernandez, M., Weems, D., Reiser, L., LaFond, F., Hanley, D., Kiphart, D., Zhuang, M., Huang, W., et al. (2001) *Nucleic Acids Res.* **29**, 102–105.
- Brenner, S., Williams, S. R., Vermaas, E. H., Storck, T., Moon, K., McCollum, C., Mao, J. I., Luo, S., Kirchner, J. J., Eletr, S., et al. (2000) *Proc. Natl. Acad. Sci. USA* **97**, 1665–1670.
- Wang, J. F., Zhou, H., Chen, Y. Q., Luo, Q. J. & Qu, L. H. (2004) *Nucleic Acids Res.* **32**, 1688–1695.
- Vierstra, R. D. (2003) *Trends Plant Sci.* **8**, 135–142.
- Hellmann, H. & Estelle, M. (2002) *Science* **297**, 793–797.
- Dharmasiri, N., Dharmasiri, S., Jones, A. M. & Estelle, M. (2003) *Curr. Biol.* **13**, 1418–1422.
- Xie, Q., Frugis, G., Colgan, D. & Chua, N. H. (2000) *Genes Dev.* **14**, 3024–3036.
- Hagen, G. & Guilfoyle, T. (2002) *Plant Mol. Biol.* **49**, 373–385.
- Hatzfeld, Y., Lee, S., Lee, M., Leustek, T. & Saito, K. (2000) *Gene* **248**, 51–58.
- Lappartient, A. G., Vidmar, J. J., Leustek, T., Glass, A. D. & Touraine, B. (1999) *Plant J.* **18**, 89–95.
- Takahashi, H., Yamazaki, M., Sasakura, N., Watanabe, A., Leustek, T., Engler, J. A., Engler, G., Van Montagu, M. & Saito, K. (1997) *Proc. Natl. Acad. Sci. USA* **94**, 11102–11107.
- Kliebenstein, D. J., Monde, R. A. & Last, R. L. (1998) *Plant Physiol.* **118**, 637–650.
- Swidzinski, J. A., Sweetlove, L. J. & Leaver, C. J. (2002) *Plant J.* **30**, 431–446.
- Vespa, L., Vachon, G., Berger, F., Perazza, D., Faure, J.-D. & Herzog, M. (2004) *Plant Physiol.* **134**, 1283–1292.
- Menand, B., Desnos, T., Nussaume, L., Berger, F., Bouchez, D., Meyer, C. & Robaglia, C. (2002) *Proc. Natl. Acad. Sci. USA* **99**, 6422–6427.
- De Rijk, P., Wuyts, J. & De Wachter, R. (2003) *Bioinformatics* **19**, 299–300.
- Jones-Rhoades, M. W. & Bartel, D. P. (2004) *Mol. Cell* **14**, 787–799.