MicroRNAs: SMALL RNAs WITH A BIG ROLE IN GENE REGULATION

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MicroRNAs are a family of small, non-coding RNAs that regulate gene expression in a sequence-specific manner. The two founding members of the microRNA family were originally identified in *Caenorhabditis elegans* as genes that were required for the timed regulation of developmental events. Since then, hundreds of microRNAs have been identified in almost all metazoan genomes, including worms, flies, plants and mammals. MicroRNAs have diverse expression patterns and might regulate various developmental and physiological processes. Their discovery adds a new dimension to our understanding of complex gene regulatory networks.

RNA INTERFERENCE (RNAi). A form of posttranscriptional gene silencing, in which dsRNA induces degradation of the homologous mRNA, mimicking the effect of the reduction, or loss, of gene activity.

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Non-coding RNAs participate in a surprisingly diverse collection of regulatory events, ranging from copynumber control in bacteria1 to X-chromosome inactivation in mammals². MicroRNAs (miRNAs) are a family of 21-25-nucleotide small RNAs that, at least for those few that have characterized targets, negatively regulate gene expression at the post-transcriptional level³⁻⁵. Members of the miRNA family were initially discovered as small temporal RNAs (stRNAs) that regulate developmental transitions in Caenorhabditis elegans6. Over the past few years, it has become clear that stRNAs were the prototypes of a large family of small RNAs, miRNAs, that now claim hundreds of members in worms, flies, plants and mammals. The functions of miRNAs are not limited to the regulation of developmentally timed events. Instead, they have diverse expression patterns and probably regulate many aspects of development and physiology^{3,4,7-9}. Although the mechanisms through which miRNAs regulate their target genes are largely unknown, the finding that at least some miRNAs feed into the RNA INTERFERENCE (RNAi) pathway has provided a starting point in our journey to understand the biological roles of miRNAs.

In this review, we revisit the history of miRNAs and summarize recent findings in miRNA biogenesis, translational repression and biological function. We conclude by highlighting the continuing genome-wide efforts to identify novel miRNAs and to predict their targets.

The discovery of miRNAs

The founding member of the miRNA family, *lin-4*, was identified in C. elegans through a genetic screen for defects in the temporal control of post-embryonic development^{10,11}. In *C. elegans*, cell lineages have distinct characteristics during 4 different larval stages (L1–L4). Mutations in *lin-4* disrupt the temporal regulation of larval development, causing L1 (the first larval stage)specific cell-division patterns to reiterate at later developmental stages¹⁰. Opposite developmental phenotypes - omission of the L1 cell fates and premature development into the L2 stage — are observed in worms that are deficient for *lin-14* (REF. 12). Even before the molecular identification of lin-4 and lin-14, these loci were placed in the same regulatory pathway on the basis of their opposing phenotypes and antagonistic genetic interactions¹¹. Most genes identified from mutagenesis screens are protein-coding, but lin-4 encodes a 22-nucleotide non-coding RNA that is partially complementary to 7 conserved sites located in the 3'-untranslated region (UTR) of the *lin-14* gene (FIG. 1b)^{13,14}. *lin-14* encodes a nuclear protein, downregulation of which at the end of the first larval stage initiates the developmental progression into the second larval stage^{13,15}. The negative regulation of LIN-14 protein expression requires an intact 3' UTR of its mRNA14, as well as a functional lin-4 gene¹³. These genetic interactions inspired a series of molecular and biochemical studies demonstrating that

the direct, but imprecise, base pairing between *lin-4* and the *lin-143'* UTR was essential for the ability of *lin-4* to control LIN-14 expression through the regulation of protein synthesis^{16–18}. Through an analogous mechanism, *lin-4* also negatively regulates the translation of *lin-28*, a cold-shock-domain protein that initiates the developmental transition between the L2 and L3 stages¹⁹. Compared with *lin-14*, *lin-28* has fewer *lin-4* binding sites, which might lead to its translational repression being delayed following *lin-4* expression owing to less efficient *lin-4* binding^{6,19}.

The discovery of lin-4 and its target-specific translational inhibition hinted at a new mechanism of gene regulation during development. In 2000, almost 7 years after the initial identification of lin-4, the second miRNA, let-7, was discovered, also using forward genetics in worms. let-7 encodes a temporally regulated 21-nucleotide small RNA that controls the developmental transition from the L4 stage into the adult stage²⁰⁻²². Similar to *lin-4*, *let-7* performs its function by binding to the 3' UTR of lin-41 and *hbl-1* (*lin-57*), and inhibiting their translation²⁰⁻²⁴. The identification of let-7 not only provided another vivid example of developmental regulation by small RNAs, but also raised the possibility that such RNAs might be present in species other than nematodes. Unlike lin-4, the orthologues of which in flies and mammals initially escaped bioinformatic searches, and were only recognized recently^{25,26}, both *let-7* and *lin-41* are evolutionarily conserved throughout metazoans, with homologues that were readily detected in molluscs, sea urchins, flies, mice and humans²⁷. This extensive



Figure 1 | The molecular hallmarks of *lin-4*, the founding member of the microRNA family. a | The precursor structure and mature microRNA (miRNA) sequence of *lin-4*. b | Sequence complementarity between *lin-4* (red) and the 3'-untranslated region (UTR) of *lin-14* mRNA (blue). *lin-4* is partially complementary to 7 sites in the *lin-14* 3' UTR; its binding to these sites of complementarity brings about repression of LIN-14 protein synthesis^{13,18}. RISC, RNA-induced silencing complex.

conservation strongly indicated a more general role of small RNAs in developmental regulation, as supported by the recent characterization of miRNA functions in many metazoan organisms.

miRNAs and siRNAs - what's the difference

Hundreds of miRNAs have now been identified in various organisms, and the RNA structure and regulatory mechanisms that have been characterized in *lin-4* and *let-7* still provide unique molecular signatures as to what defines miRNAs. miRNAs are generally 21–25nucleotide, non-coding RNAs that are derived from larger precursors that form imperfect stem-loop structures (FIG. 1a)^{4,5}. The mature miRNA is most often derived from one arm of the precursor hairpin, and is released from the primary transcript through stepwise processing by two ribonuclease-III (RNase III) enzymes^{28,29}. At least in animals, most miRNAs bind to the target-3' UTR with imperfect complementarity and function as translational repressors (see below for a discussion of plant miRNAs)⁴.

Almost coincident with the discovery of the second miRNA, let-7, small RNAs were also characterized as components of a seemingly separate biological process, RNA interference (RNAi). RNAi is an evolutionarily conserved, sequence-specific gene-silencing mechanism that is induced by exposure to dsRNA³⁰. In many systems, including worms, plants and flies, the stimulus that was used to initiate RNAi was the introduction of a dsRNA (the trigger) of ~500 bp. The trigger is ultimately processed in vivo into small dsRNAs of ~21-25 bp in length, designated as small interfering RNAs (siRNAs)^{31,32}. It is now clear that one strand of the siRNA duplex is selectively incorporated into an effector complex (the RNA-induced silencing complex; RISC). The RISC directs the cleavage of complementary mRNA targets, a process that is also known as post-transcriptional gene silencing (PTGS) (FIG. 2)33. The evolutionarily conserved RNAi response to exogenous dsRNA might reflect an endogenous defense mechanism against virus infection or parasitic nucleic acids³⁰. Indeed, mutations of the RNAi components greatly compromise virus resistance in plants, indicating that PTGS might normally mediate the destruction of the viral RNAs³⁴. In addition, siRNAs can also regulate the expression of target transcripts at the transcriptional level, at least in some organisms. Not only can siRNAs induce sequence-specific promoter methylation in plants^{35,36}, but they are also crucial for heterochromatin formation in fission yeast^{37,38}, and transposon silencing in worms^{39,40}.

Fundamentally, siRNAs and miRNAs are similar in terms of their molecular characteristics, biogenesis and effector functions (see below for details). So, the current distinctions between these two species might be arbitrary, and might simply reflect the different paths through which they were originally discovered. miRNAs and siRNAs share a common RNase-III processing enzyme, Dicer, and closely related effector complexes, RISCs, for post-transcriptional repression (FIG. 2). In fact, much of our current knowledge of the biochemistry of miRNAs stems from what we know about siRNAs and the RNAi pathway.





On the other hand, miRNAs differ from siRNAs in their molecular origins and, in many of the cases that have been characterized so far, in their mode of target recognition. miRNAs are produced as a distinct species from a specific precursor that is encoded in the genome. The structure of the primary miRNA transcript and the recognition of this precursor by a nuclear processing machinery probably determines the sequence and structure of mature miRNAs⁴. By contrast, siRNAs are sampled more randomly from long dsRNAs that can be introduced exogenously or produced from bi-directionally transcribed endogenous RNAs that anneal to form dsRNA³⁰. In many cases, miRNAs bind to the target 3' UTRs through imperfect complementarity at multiple sites, and therefore negatively regulate target expression at the translational level. By contrast, siRNAs often form a perfect duplex with their targets at only one site, and therefore direct the cleavage of the target mRNAs at the site of complementarity (FIG. 2).

The artificial nature of current distinctions between siRNA and miRNA is highlighted by numerous recent findings. The extent of complementarity between the siRNA/miRNA and its target can determine the mechanism of silencing. For example, with the exception of miR-172, which acts as a translational repressor, all characterized plant miRNAs anneal to their targets with nearly complete complementarity at a single site, either in the coding region or in the UTRs, therefore condemning their target mRNAs to destruction by cleavage and degradation⁴¹⁻⁴³. A similar situation has also been described for one mammalian miRNA, miR-196: the nearly perfect base pairing between miR-196 and Hoxb8 directs cleavage of Hoxb8 mRNA both in mouse embryos and in cell culture⁴⁴. Conversely, when siRNAs pair with their targets imperfectly, siRNAs can trigger translational repression rather than mRNA cleavage in mammalian tissue culture⁴⁵. So, we are left with the question of whether miRNAs are truly different from siRNAs or whether our current understanding fails to functionally distinguish these two species under physiological conditions.

Both miRNAs and siRNAs depend on Dicer for their maturation^{46,47}, and both have been shown to be part of similar RISCs (FIG. 2)^{48,49}. However, the effector complexes have only been studied for a few miRNAs, and in no case has there been biochemical data to confirm that most miRNA-containing complexes have been accounted for^{50,51}. So, as we get beyond the superficial similarities of the structure and functions of these small RNAs, we must now begin to focus on the details that distinguish the modes of action of siRNAs and miRNAs *in vivo* to understand their true biological functions.

Biogenesis of miRNAs

Two processing events lead to mature miRNA formation in animals. In the first, the nascent miRNA transcripts (pri-miRNA) are processed into ~70-nucleotide precursors (pre-miRNA); in the second event that follows, this precursor is cleaved to generate ~21–25-nucleotide mature miRNAs²⁹. Little is known about the transcriptional regulation of pri-miRNAs, except that certain primiRNAs are located within introns of host genes, including both protein-coding genes and non-coding genes, and might therefore be transcriptionally regulated through their host-gene promoters⁵². In addition, certain miRNAs are clustered in polycistronic transcripts, indicating that these miRNAs are coordinately regulated during development⁵².

The sequential cleavages of miRNA maturation are catalysed by two RNase-III enzymes, Drosha and Dicer (FIG. 2)^{28,47}. Both are dsRNA-specific endonucleases that generate 2-nucleotide-long 3' overhangs at the cleavage site. Drosha is predominantly localized in the nucleus and contains two tandem RNase-III domains, a dsRNA binding domain and an amino-terminal segment of unknown function²⁸. Regardless of the diverse primary sequences and structures of pri-miRNAs, Drosha cleaves these into ~70-bp pre-miRNAs that consist of an imperfect stem-loop structure (FIG. 2)²⁸. Although the precise mechanisms that Drosha uses to discriminate miRNA precursors remain unknown, several studies





BOOTSTRAP SAMPLING As applied to molecular phylogenies, nucleotide or amino-acid sites are sampled randomly, with replacement, and a new tree is constructed. This is repeated many times and the frequency of appearance of a particular node among the bootstrap trees is viewed as a support (confidence) value for deciding on the significance of that node. have addressed the features of pri-miRNAs that contribute to Drosha cleavage both *in vitro* and *in vivo*^{28,53}. The efficiency of Drosha processing depends on the terminal loop size, the stem structure and the flanking sequence of the Drosha cleavage site, because shortening of the terminal loop, disruption of complementarity within the stem and removal or mutation of sequences that flank the Drosha cleavage site significantly decrease, if not abolish, the Drosha processing of pri-miRNAs^{28,53}.

After the initial cleavage by Drosha, pre-miRNAs are exported from the nucleus into the cytoplasm by Exportin 5 (Exp5), a Ran-GTP dependent nucleo/cytoplasmic cargo transporter (FIG. 2)⁵⁴. Once inside the cytoplasm, these hairpin precursors are cleaved by Dicer into a small, imperfect dsRNA duplex (miRNA: miRNA*) that contains both the mature miRNA strand and its complementary strand (miRNA*)47,55,56. Dicer contains a putative helicase domain, a DUF283 domain, a PAZ (Piwi-Argonaute-Zwille) domain, two tandem RNase-III domains and a dsRNA-binding domain (dsRBD) (FIG. 3a)⁴⁶. Recent structural analysis of the PAZ domain revealed a variant of the OB fold, a module that allows a low-affinity interaction with the 3' end of ssRNAs57-59. This association also allows the PAZ domain to interact with dsRNAs that present 2-nucleotide 3' overhangs, such as those that result from Drosha cleavage^{51,57,58}. In addition, efficient Dicer cleavage also requires the presence of the overhang and a minimal stem length (D. Siolas and G.J.H., unpublished observations), indicating a model in which the Dicer PAZ domain might recognize the end of the Drosha cleavage product, and therefore position the site of the second RNase-III cleavage on the stem of the miRNA precursors⁶⁰. Dicer cleavage generates mature miRNAs that range from 21 to 25 nucleotides, and such differences in size possibly result from the presence of bulges and mismatches on the pre-miRNA stem.

During Dicer processing, efficient cleavage of dsRNA requires dimerized RNase-III domains, because, on the basis of known RNAse-III structures, functional catalytic sites can only be formed at the interface of the RNase-III dimer⁶¹. As Dicer might only catalyse one cleavage event during miRNA processing, it is probable that Dicer functions as a monomer with its two RNase-III domains forming an intramolecular dimer and that it generates an active catalytic site for dsRNA cleavage⁶⁰. Similar to Dicer, Drosha also contains two tandem RNase-III domains and carries out a single cleavage event to generate pre-miRNA. Given our lack of knowledge about the precise structural elements that guide Dicer and Drosha cleavage, it is difficult to speculate about their exact biochemical mechanisms. However, it is probable that Drosha and Dicer share closely related mechanisms for the processing of miRNAs.

Although plant miRNAs seem to be produced from long, primary transcripts, the maturation of miRNAs in plants must, by necessity, occur differently from miRNA maturation in animals, as no plant Drosha homologue has yet been found. However, the Dicer family in Arabidopsis thaliana has a level of complexity that has not been observed in other organisms studied so far (FIG. 3b). There are four Dicer homologues in A. thaliana - DCL1, DCL2, DCL3 and DCL4 — two of which (DCL1 and DCL4) contain nuclear localization signals (FIG. 3a). Therefore, it seems possible that Drosha function might be carried out by one or more specialized Dicer enzymes in plants. The Dicer homologue DCL1 contains two nuclear localization signals, and predominates the nuclei when expressed as a GFP-fusion protein in transient-transfection studies⁶². In plants that are deficient for DCL1, the production of mature miRNAs is reduced for all miRNAs examined, yet no accumulation is detected for the corresponding pre-miRNAs^{62,63}. These findings imply that DCL1 might catalyse both the Drosha- and Dicer-like cleavages for the maturation of some, if not most, miRNAs inside the nucleus.

Consistent with this model, mature miRNAs seem to be produced within the nucleus in plants, because nuclear expression of P19, a viral protein that represses the accumulation of siRNAs and miRNAs, results in a significant reduction in mature miRNAs, whereas the cytoplasmic expression of P19 has no such effects62. It is not clear whether other Dicer homologues in plants are also involved in miRNA maturation. However, it has been speculated that the presence of multiple plant Dicer enzymes might explain the complexity of siRNA species in plants. Unlike siRNAs in animals, which are usually 21-22 nucleotides long, plants generate two classes of siRNA: 21-22- and ~25-bp-long siRNAs⁶⁴. Given the different classes of small dsRNAs found in plants, it is possible that each Dicer homologue regulates separate cleavage events to generate specific species of dsRNA duplexes65.

The functional specificity of different Dicer enzymes in organisms with multiple Dicer homologues has recently been indicated by a series of genetic and biochemical studies in Drosophila melanogaster^{66,67}. Two Dicer homologues have been identified in flies: Dicer1 and Dicer2 (FIG. 3a)67. Deficiency in Dicer1 disrupts the processing of pre-miRNAs, whereas loss of Dicer2 affects the production of siRNAs, but not miRNA maturation67. These findings are consistent with the fact that the PAZ domain is only present in Dicer1, but absent in Dicer2 (FIG. 3a), given a model in which the PAZ domain recognizes the staggered ends of pre-miRNAs67 and mediates their cleavage. Both Dicer1 and Dicer2 seem to function downstream of miRNA and siRNA production to facilitate the RISC-mediated gene silencing. Originally indicated by physical interactions between Dicer and RISC in D. melanogaster⁶⁸, this model has recently gained concrete biochemical support. Dicer2 forms a complex with R2D2, a dsRNA-binding protein, and the formation of the Dicer2/R2D2 complex with siRNAs enhances sequence-specific mRNA degradation that is mediated by the RISC complex69,70. In addition, Dicer1 deficiency affects siRNA-mediated gene silencing without disrupting siRNA production, indicating a possible role of Dicer1 in enhancing RNAi effector activity67. Therefore, the functions of Dicer might be broader than previously suspected, involving both the initiation and the effector steps of miRNA/siRNA-mediated gene silencing.

The effector complex for miRNAs shares core components with that of siRNAs, so much so that both are collectively referred to as the RNA-induced silencing complex (RISC). The target specificity, and probably also the functional efficiency, of a miRNA requires that the mature miRNA strand from the miRNA:miRNA* duplex be selectively incorporated into the RISC for target recognition71,72. The miRNA* strand, on the other hand, is probably degraded rapidly on its exclusion from the RISC, as the recovery rate of miRNA*s from endogenous tissues is ~100-fold lower than that of miRNAs71. As Dicer processes the pre-miRNA into the miRNA: miRNA* duplex, the stability of the 5' ends of the two arms of the miRNA:miRNA* duplex is usually different. Although mature miRNAs can reside on either strand of the hairpin stem, it is almost always derived from the

strand with the less stable 5' end compared with the miRNA* strand71,72. These findings indicate that the relative instability at the 5' end of the mature miRNA might facilitate its preferential incorporation into the RISC. The selective assembly of the mature miRNA into the RISC probably reflects the relative ease of unwinding from one end of the miRNA: miRNA* duplex (FIG. 2)^{71,72}. Therefore, the thermodynamic properties of the miRNA precursor determine the asymmetrical RISC assembly, and therefore, the target specificity for posttranscriptional inhibition. However, in rare cases in which miRNA and miRNA* have similar 5'-end stability, each arm of the miRNA precursor is predicted to be assembled into the RISC at similar frequencies. This prediction has been confirmed by similar recovery rates for such miRNAs and miRNA*s from endogenous tissues⁷¹. This thermodynamic model also applies to the asymmetrical assembly of the siRNA duplex, in which the strand of siRNA with the less stable 5' end is preferentially assembled into the RISC complex to target mRNA cleavage^{71,72}. Altogether, there seems to be a common thermodynamic mechanism that regulates the asymmetric assembly of siRNA or miRNA from the dsRNA duplexes, which safeguards specificity towards corresponding targets.

Post-transcriptional repression by miRNAs

The precise molecular mechanisms that underlie posttranscriptional repression by miRNAs still remain largely unknown. One of the best-studied examples is *lin-4*, which negatively regulates its target, *lin-14*, by repressing its translation¹⁶. Base pairing between *lin-4* and lin-14 has proved to be crucial for their interaction in vivo, as mutations that affect their complementarity compromise or abolish this negative regulation¹³. Interestingly, *lin-4* only inhibits the synthesis of the LIN-14 protein but fails to affect the synthesis, polyadenylation state or abundance of lin-14 mRNA16. Furthermore, the initiation of lin-14 translation seems to occur normally in the presence of *lin-4*, because *lin-14* mRNAs are efficiently incorporated into polyribosomes regardless of lin-4 expression¹⁶. Therefore, it is reasonable to speculate that the translational repression by lin-4 occurs after translational initiation, probably during translational elongation and/or the subsequent release of the LIN-14 protein16.

Translational repression of target genes is not specific to *lin-4*; in fact, it turns out to be the predominant mechanism by which miRNAs negatively regulate their targets throughout the animal kingdom. *let-7* and *Bantam* (which encodes an anti-apoptotic miRNA) bind to the 3' UTRs of their targets and negatively regulate their translation in worms and flies, respectively^{20,73}. In addition, *miR-30*, a mammalian miRNA, inhibits protein synthesis of a reporter gene that bears an artificial 3' UTR with *miR-30* complementary sites⁵³.

Although most animal miRNAs repress target translation, one miRNA, *mir-196*, was found recently to direct mRNA cleavage of its target, *Hoxb8* (REF. 44). In plants, however, most miRNAs that have been studied so far mediate the destruction of their target mRNAs⁷⁴. These

Table 1 Expression studies on mammalian microRNAs					
Expression pattern	microRNA	References			
Tissue-specific expression patterns of mammalian microRNAs					
ES-cell specific	miR-296	86			
Expressed in ES cells, but upreguated on differentiation	<i>miR-21</i> and <i>miR-22</i>	86			
Expressed in both ES cells and various adult tissues	miR-15a, miR-16, miR-19,b, miR-92, miR-93, miR-96, miR-130 and miR-130b	86			
Enriched during mouse brain development	miR-128, miR-19b, miR-9, miR-125b, miR-131, miR-178, miR-124a, miR-266 and miR-103	26,90			
Enriched in adult brain	miR-9*, miR-125a, miR-125b, miR-128, miR-132, miR-137, miR-139, miR-7, miR-9, miR-124a, miR-124b, miR-135, miR-153, miR-149, miR-183, miR-190 and miR-219	26			
Enriched in lung	miR-18, miR-19a, miR-24, miR-32, miR-130, miR-213, miR-20, miR -141, miR-193 and miR-200b	26			
Enriched in spleen	miR-99a, miR-127, miR-142-a, miR-142-s, miR-151, miR-189 and miR-212	26			
Haemetopoietic tissues	miR-181, miR-223 and miR-142	26			
Enriched in liver	miR-122a, miR-152, miR-194, miR-199 and miR-215	26			
Enriched in heart	miR-1b, miR-1d, miR-133, miR-206, miR-208 and miR-143	26			
Enriched in kidney	miR-30b, miR-30c, miR-18, miR-20, miR-24, miR-32, miR-141, miR-193 and miR-200b	26			
Ubiquitously expressed	miR-16, miR-26a, miR-27a, miR143a, miR-21, let-7a, miR-7b, miR-30b and miR-30c	26			
Abnormal microRNA expression during tumorigenesis					
Downregulated in chronic lymphocytic leukaemias	miR-15 and miR-16	102			
Downregulated in lung cancer cell lines	miR-26a and miR-99a	89			
Downregulated in colon cancers	miR143/miR-145 cluster	103			
Upregulated in Burkitt lymphoma	miR-155	88			

ES cells, embryonic stem cells.

plant miRNAs differ from animal miRNAs in that their base pairing with the corresponding targets is nearly perfect, and that their complementary sites are located throughout the transcribed regions of the target gene, instead of being limited to the 3' UTRs. So far, only one plant miRNA, miR-172, has been shown to act as a translational repressor during A. thaliana flower development. Unlike most characterized animal miRNAs, the site of miR-172 complementarity (which is high) within the target gene APETALA 2 (AP2) was located in the coding region instead of the 3' UTR41. An important caveat to these observations is that those plant miRNAs that act through RNA cleavage might be overrepresented in the current studies owing to the ease of their target identification through computational analysis (see below).

Despite progress in identifying protein and RNA components of the RISC, the biochemical mechanism by which this complex functions still remains unknown. Genetic screens combined with biochemical purification have pinpointed several important components. Argonaute (AGO) proteins belong to an evolutionarily conserved family that is defined by the presence of a PAZ domain and a Piwi domain⁷⁵. AGO-family proteins have been consistently co-purified with RISC activity in many organisms⁴⁸, and mutations in AGO homologues have been associated with distinct developmental

and/or RNAi phenotypes⁷⁵. As a core component of the RISC, the AGO family has multiple homologues in each metazoan species (24 for *C. elegans*, 5 for *D. melanogaster* and 8 for mammals)⁷⁵. Given the diverse mutant phenotypes and expression patterns of AGO homologues, RISCs might come in different flavours and act in a tissue-specific or a developmentally regulated manner.

In addition to AGO homologues, several other proteins have also been co-purified with the RISC. It is not clear, however, whether these RISC-associated proteins are core RISC components or whether they act as accessory proteins that provide functional specificity for the RISCs under different developmental and/or physiological contexts. The RNA-binding proteins VIG and Fragile X-related protein49,68, and the nuclease Tudor-SN, have been co-purified with RISC activity from D. melanogaster S2 CELL extracts⁷⁶. Such components also associate with siRNAs and miRNAs in mammalian cells and C. elegans⁷⁶. Alternative RISCs must also exist, as the helicases Gemin3 and Gemin4 have been co-purified with miRNAs and the human Argonaute homologue eIF2C2 (AGO2)50 from mammalian cells. These observations are consistent with the model according to which RISCs exist in different subtypes, perhaps specialized to mediate different post-transcriptional repression events in vivo.

S2 CELL

RNAi.

POLYSOME

A cell line that is isolated from

line is phagocytic, which might

contribute to its susceptibility to

A functional unit of protein

ribosomes that are attached

along the length of a single

synthesis that consists of several

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dissociated *Drosophila* melanogaster embryos. The cell

microRNA	Target gene(s)	Function	Mode of repression	References	
Caenorhabditis elegans					
lin-4	lin-14, lin-28	Regulation of developmental transition between the first two larval stages, L1 and L2	Translational repression	13,18,19	
let-7	lin-41, hbl-1	Regulation of developmental transition between the last larval stage (L4) and the adult stage	Translational repression	20–23	
lsy-6	cog-1	Determination of left/right asymmetry of neuronal development	Unknown	9	
Drosophila melanogaster					
Bantam	hid	Promotion of cell proliferation and suppression of apoptosis	Translational repression	73	
miR-14	Unknown	Suppression of apoptosis and regulation of fat metabolism	Unknown	97	
Mus musculus					
miR-181	Unknown	Promotion of haematopoietic differentiation towards the B-cell lineage	Unknown	99	
miR-196	Hoxb8	Unknown	PTGS	44	
Arabidopsis thaliana					
miR-165, miR-166	PHB, PHV and REV	Regulation of leaf morphogenesis	PTGS	104,105	
miR-172	AP2	Regulation of flowering time and floral-organ identity	Translational repression	41	
miR-JAW	TCP transcription factors	Regulation of leaf development and embryogenesis	PTGS	98	
miR-39	SCL family proteins	Unknown	PTGS	42	
miR-159	MYB33 family transcription factors	Regulation of leaf morphogenesis	PTGS	43,63,98	
Zea mays					
miR-166	rld1	Regulation of leaf morphogenesis	PTGS	106	

Table 2 | microRNAs and their targets: examples of microRNAs with experimentally validated functions/targets

AP2, APETALA 2; Hid, head involution defective; Hoxb8, homeobox B8; PHB, PHABULOSA; PHV, PHAVOLUTA; PTGS, post-transcriptional gene silencing; REV, REVOLUTA; rld1, rolled leaf1; SCL, SCARECROW-LIKE; TCP, teosinte branched 1-cycloidea-PCF.

Genome-wide efforts for miRNA identification

The extension of miRNA-mediated regulation from being a curiosity in C. elegans to a potentially general mechanism for gene regulation began with the work of the Tuschl, Bartel and Ambros groups, who collectively identified more than 100 novel miRNAs by cloning and sequencing endogenous small RNAs of 21-25 bp long from worms, flies and mammals77-79. miRNAs that were isolated in these studies show the characteristics of Dicer cleavage, temporally and spatially regulated expression and, in many cases, evolutionary conservation, either throughout metazoans or within closely related species77-79. Similar to lin-4 and let-7, these miRNAs mostly map to one arm of their predicted stem-loop precursors. The existence of miRNA precursors with stem-loop structure is also indicated by northern-blot detection of ~70-nucleotide precursors, and by cloning of the rare reverse-strand miRNA77-79. In addition to the continued cloning efforts, novel miRNAs have been isolated through their association with POLYSOMES⁸⁰ and ribonucleoprotein complexes^{50,51}. So far, several hundred miRNAs have been identified in mammals, fish, flies, worms and plants, corresponding to ~1% of the protein-coding genes in each organism⁸¹. As the current miRNA cloning technology clearly favours highly expressed miRNAs, more exhaustive cloning efforts are needed that must include more tissue types and

developmental stages. In addition to experimental approaches, bioinformatic predictions have helped to identify novel miRNAs in various organisms, mostly on the basis of pre-miRNA hairpin structures and sequence conservation throughout evolution^{82–84}. With the growing number of experimentally confirmed miRNAs, refined bioinformatic approaches will undoubtedly increase in power for identifying miRNAs that have escaped experimental searches.

A comprehensive miRNA registry

The global efforts for miRNA cloning and characterization have led to the establishment of an important collection of miRNA data. The miRNA Registry (see online links box) contains up-to-date annotation for all published miRNAs⁸¹. Most database entries require experimental validation of mature miRNA expression and computational prediction of the corresponding hairpin precursor. In cases in which homologous miRNAs can be easily identified from closely related species, such as mouse and human, computationally predicted entries are also included in the registry to provide a more comprehensive list⁸¹. At the time of writing, the database contains 116 C. elegans miRNAs, 50 miRNAs from C. briggsae, 78 D. melanogaster miRNAs, 43 A. thaliana miRNAs, 28 miRNA from rice, 207 mouse miRNAs, 185 miRNAs from rat and 191 human miRNAs.

Following the identification of hundreds of miRNAs in various organisms, large-scale studies on miRNAexpression profiles were carried out in many model organisms using northern-blot analysis, microarrays and miRNA cloning^{25,85-87}. miRNAs show dynamic temporal and spatial expression patterns, disruption of which is associated with developmental/physiological abnormalities (TABLE 1). For example, the miR-155 precursor is enriched in Burkitt lymphoma tissues; the location of miR-155 correlates with a specific chromosomal translocation⁸⁸. miR-26a and miR-99a expression is reduced in human lung cancer cell lines, which correlates with their location in regions of loss of heterozygosity in lung tumours/cell lines89. miR-9 and miR-131 levels are reduced in Presenilin-1-deficient mice that show lethal developmental defects90. These findings indicate that miRNAs might have a general role in regulating gene expression in diverse developmental and physiological processes, and provide substantial hints that misregulation of miRNA function might contribute to human disease.

Functional characterization of miRNAs

Although the studies of *lin-4* and *let-7* shaped our understanding of miRNA molecular structures and functional mechanisms, their roles in temporal regulation of development only revealed one of many possible aspects of miRNA function. Mutations in Dicer homologues disrupt the biogenesis of miRNAs, and cause diverse developmental defects, including germline defects in *C. elegans*⁹¹, abnormal embryogenesis in *A. thaliana*⁶³, developmental arrest in zebrafish⁹² and depletion of stem cells in mice⁹³. Mutations in AGO family proteins are also associated with pleiotropic developmental phenotypes, such as premature MERISTEM differentiation in *A. thaliana zwille* mutants⁹⁴, defective embryogenesis and larval development in *C. elegans alg-1* and *alg-2* mutants⁵⁵ and decreased stem-cell self-renewal during oogenesis in *Drosophila piwi* mutants⁹⁵. Because Dicer and AGO are essential components in miRNA and siRNA biogenesis and function, these defects might reflect the collective functions of multiple miRNAs and/or siRNAs that are expressed during early development.

So far, functional studies have only touched on a handful of miRNAs. Following the classic studies on the role of *lin-4* and *let-7* in worm developmental timing, functional characterization of miRNAs continues to benefit from the loss-of-function mutations of miRNA genes. The fly miRNA, Bantam, was originally characterized from a fly P-ELEMENT screen for genes that promote cell proliferation and suppress apoptosis during tissue growth%. Its sequence complementarity with the 3' UTR and its functional antagonism of the pro-apoptotic hid (head involution defective) gene were the first clues for the translational repression of hid by Bantam^{73,96}. In addition, fly miR-14 was identified through a P-element screen for inhibitors of apoptotic cell death. Deficiency in miR-14 enhances cell death that is induced by the celldeath activator, Reaper, and results in defective stress responses and fat metabolism⁹⁷. Loss of *miR-14* also leads to an elevated level of Drice, an apoptotic effector caspase, indicating a direct or indirect repression of Drice by miR-14. Perhaps one of the most interesting miRNAs identified so far is *lsy-6*, a miRNA that is asymmetrically expressed in worm taste-receptor neurons to promote specific cell fate in left-right patterning. ASE left (ASEL) and ASE right (ASER) are two taste-receptor neurons, each with specific

Box1 | Bioinformatic predictions of microRNA targets

The functional characterization of microRNAs (miRNAs) heavily relies on the identification of miRNA target genes. In plants, because certain miRNAs are nearly perfectly complementary to their targets, bioinformatic predictions of miRNA targets have proved to be relatively straightforward, and in fact, powerful⁴³. Interestingly, many predicted miRNA targets are transcription factors, highlighting possible miRNA functions in regulating diverse developmental processes in plants⁴³. However, the computational prediction of animal miRNA targets is more difficult, both because the miRNAs are only ~21-25 nucleotides long and because the miRNA-target pairings are not entirely complementary¹⁰⁰. As a result, bioinformatic prediction has to rely on rules that are built on a few known miRNA-target interactions, which are generalized for genome-wide searches. Starting from 3 animal miRNAs, lin-4, let-7 and Bantam, the targets of which have been experimentally validated, the Cohen group screened the fly genome for miRNA targets on the basis of the following 3 criteria: perfect complementarity or G•U pairing between the target 3'-untranslated region (UTR) and the first 8 nucleotides of miRNA; favourable structural and thermodynamic heteroduplex formation between miRNA and its putative targets; and evolutionary conservation of miRNA target sites between Drosophila melanogaster and D. pseudoobscura¹⁰¹. To experimentally validate potential targets, they fused a GFP reporter gene upstream of the predicted target 3' UTR and examined GFP expression in WING DISC with and without overexpression of the corresponding miRNA¹⁰¹. Using this approach, they gained experimental support for the informatically predicted targets of three miRNAs, including Notch genes for miR-7, pro-apoptotic genes for miR-2 and genes that encode metabolic enzymes for miR-277 (REF. 101).

The Burge group developed the TargetScan algorithm for predicting vertebrate miRNA targets on the basis of miRNAtarget complementarity (particularly at the 5' region of the miRNA) and evolutionary conservation among vertebrates. In their first study, they experimentally validated 11 predicted targets out of 15 tested, using a HeLa cell reporter system that contains a luciferase reporter gene fused to the predicted target 3'-UTR fragments¹⁰⁰. Among their predicted targets, 29% have unknown functions, whereas 71% have diverse functions, such as DNA binding, transcriptional regulation, signaltransducer and kinase activity¹⁰⁰. Although the prediction success rate is difficult to determine on a genome-wide scale, these studies are an important step towards our understanding of miRNA function in animals.

MERISTEM

The undifferentiated tissue at the tips of stems and roots in which new cell division is concentrated.

P-ELEMENTS

A family of transposable elements that are widely used as the basis of tools for mutating and manipulating the *Drosophila* genome.

WING DISC

A sac-like structure of a mature third instar fly larva, which will give rise to the adult wing.

REVIEWS

INFLORESCENCE TISSUE The reproductive backbone that displays the flowers. expression patterns of chemoreceptor genes. The specific *lsy-6* expression in the ASEL neuron suppresses the expression of an *Nkx*-type homeobox gene, *cog-1*, thereby promoting the ASEL cell fate by specifying ASEL-specific expression of chemoreceptors.

The functional characterization of plant miRNAs has also benefited from genetic mutations in miRNA genes. The *miR-JAW* gene was originally identified as a gain-of-function mutation causing the uneven leaf curvature and shape⁹⁸. The identification of *miR-JAW* targets — the *TCP* genes — was first indicated by their opposing biological functions during leaf morphogenesis and their sequence complementarity⁹⁸.

For most of the miRNAs for which genetic mutations are unavailable, functional characterization might have to begin with expression studies and/or bioinformatic predictions. For example, *miR-181* is enriched in B-lymphoid cells of mouse bone marrow; this unique expression pattern led to the discovery of its function in promoting haematopoietic differentiation towards the B-cell lineage⁹⁹. *miR-39*, which accumulates predominantly in the INFLORESCENCE TISSUE of A. thaliana⁴² is another example. Using the sequence complementarity and overlapping expression pattern as the target search criteria, Llave *et al.* predicted the *miR-39* targets as Scarecrow-like (SCL) family proteins and further confirmed their hypothesis through biochemical studies (TABLE 2)⁴².

Most of the miRNAs that have been characterized so far seem to regulate aspects of development, including larval developmental transitions and neuronal development in *C. elegans*, growth control and apoptosis in *D. melanogaster*, haematopoietic differentiation in mammals and leaf development in *A. thaliana* (TABLE 2). But, only a handful of miRNAs have been carefully studied so far, and the diverse expression patterns of miRNAs and altered miRNA expression under certain physiological conditions, such as tumorigenesis, indicate a range of unknown functions that might extend beyond developmental regulation. In addition, bioinformatic predictions of miRNA targets (BOX 1) have revealed a diversity of regulatory pathways that might be subject to miRNA-mediated regulation. Overall, no specific indication has emerged that miRNA-mediated regulation is restricted to one biological process. Instead, the emerging picture is that miRNAs have the potential to regulate almost all aspects of cellular physiology.

Conclusions

Since the discovery of miRNAs as stRNAs in C. elegans^{13,18}, remarkable advances in the characterization of this gene family have not only demonstrated that these small, non-coding RNAs are a prevalent class of regulatory RNAs, but they have also indicated the outlines of a biochemical mechanism for their functions in gene regulation. However, with relatively few exceptions, we know little about the precise roles of the vast majority of miRNAs in regulating gene expression. Furthermore, the precise mechanisms by which miRNA- and siRNAmediated repression might differ remain to be explained. For example, miRNAs and siRNAs mature in a similar way and join structurally related, if not identical, effector complexes. However, subtle differences between these classes of small RNA are beginning⁴, and will no doubt continue, to emerge as we understand more of the precise relationships between miRNAs, siRNAs and the protein components of the RNAi machinery.

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Competing interests statement

The authors declare that they have no competing financial interests.

Online links

DATABASES

The following terms in this article are linked online to: Entrez: http://www.ncbi.nih.gov/Entrez

AP2 | cog-1 | hbl-1 (lin-57) | Hoxb8 | let-7 | lin-4 | lin-14 | lin-28 | lin-41 | lys-6 | miR-99 | miR-99a | miR-155 | miR-181 | miR-196 TAIR: http://www.arabidopsis.org DCL1 | DCL2 | DCL3

FURTHER INFORMATION

miRNA Registry: http://www.sanger.ac.uk/Software/Rfam/mirna/ PHYLIP programs:

http://evolution.genetics.washington.edu/phylip.html Access to this links box is available online.

CORRECTION

MicroRNAs: SMALL RNAs WITH A BIG ROLE IN GENE REGULATION

Lin He and Gregory J. Hannon

Nature Reviews Genetics 5, 522–531 (2004); doi: 10.1038/nrg1379

In figure 2, the orientation of some RNA structures was incorrect. The corrected version is shown below. This correction has been made to the online enhanced text and PDF version of this review.

