Emerging paradigms of regulated microRNA processing

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MicroRNAs (miRNAs) modulate a broad range of gene expression patterns during development and tissue homeostasis, and in the pathogenesis of disease. The exquisite spatio–temporal control of miRNA abundance is made possible, in part, by regulation of the miRNA biogenesis pathway. In this review, we discuss two emerging paradigms for post-transcriptional control of miRNA expression. One paradigm centers on the Microprocessor, the protein complex essential for maturation of canonical miRNAs. The second paradigm is specific to miRNA families, and requires interaction between RNA-binding proteins and cis-regulatory sequences within miRNA precursor loops.

The microRNA (miRNA) biogenesis pathway is a series of biochemical steps that converts the primary miRNA transcript (pri-miRNA) to the biologically active, mature miRNA [Kim et al. 2009]. The characterized first step is the recognition of the canonical stem–loop structure of the miRNA by the Microprocessor complex. This leads to ribonucleolytic cleavage of the pri-miRNA, removal of the flanking sequences, and liberation of the stem–loop precursor miRNA (pre-miRNA). The pre-miRNA is exported to the cytoplasm via the Exportin5 pathway. The precursor is then cleaved by the ribonuclease enzyme Dicer. This leads to an ~22-nucleotide duplex, one strand of which is loaded into the Argonaute-containing RNA-induced silencing complex (RISC). This represents the endpoint of biogenesis; the single-stranded mature miRNA is ready to guide mRNA binding, which leads to translational repression and/or mRNA destabilization (Kim et al. 2009). As an exception to this rule, the less abundant “mirtrons” circumnavigate Drosha processing and are processed only by Dicer (Okamura et al. 2008).

In principle, miRNA abundance could be controlled at transcription of the pri-miRNA, during any of the biogenesis steps, or at turnover of the mature miRNA. Control at transcription is well established. Most miRNAs are transcribed by RNA polymerase II, and their upstream regulatory regions contain canonical core promoters and enhancers [Lee and Dutta 2009]. Early studies by several laboratories, however, found that mature miRNA expression does not always correlate with expression of the pri-miRNA [Obernosterer et al. 2006; Thomson et al. 2006; Blenkiron et al. 2007; Wulczyn et al. 2007]. Thus, miRNAs themselves must be post-transcriptionally regulated. In fact, regulation at multiple biogenesis steps and at turnover of the mature miRNA has now been established [Hwang et al. 2007; for review, see Pawlicki and Steitz 2009]. This review focuses on the regulation of miRNA production during biogenesis. The majority of discoveries on miRNA regulation can be distilled down to two contrasting paradigms based on the biochemical point of regulation: at the Microprocessor complex, and at the terminal loop of specific miRNA precursors.

Multiple regulatory events converge on the Microprocessor complex

As the first processing step in miRNA biogenesis, the Microprocessor is positioned to play a pivotal role in the regulation of mature miRNA abundance. This complex is minimally composed of two proteins: the dsRNA-binding protein DGCR8 [Pasha], and the RNase III enzyme Drosha [Lee et al. 2003; Denli et al. 2004; Gregory et al. 2004; Han et al. 2004; Landthaler et al. 2004]. These two proteins represent the essential requirements for the first processing step, defined by reconstitution of activity with purified recombinant proteins. In human cell extracts, however, Drosha has been described to reside in multiple complexes [Gregory et al. 2004]. A complex that is comprised solely of Drosha and DGCR8 has robust pri-miRNA processing activity. A second, larger complex has multiple accessory proteins, yet also has pri-miRNA processing activity. It is not clear whether this large complex is assembled around RNAs, since many of the accessory factors contain RNA interaction motifs, and RNase treatment of extracts has been shown to shift the size of the Drosha complex [Han et al. 2004]. The accessory proteins in this large Drosha complex include the EWSR1, Fus, numerous heterogenous nuclear RNA complex (hnRNP) proteins, and the DEAD-box helicases p68/DDX5 and p72/DDX17. While the exact biochemical composition of the cellular Microprocessor is unknown,
a recent collection of studies implicates p68 and p72 as important cofactors of the Microprocessor in vivo.

**Integrating signal transduction with Drosha processing: the role of p68 and p72**

The p68 and p72 helicases are conserved across eukaryotes and are implicated in diverse RNA processing pathways (for review, see Fuller-Pace 2006). In cell-free assays, both proteins have moderately processive ATP-dependent RNA unwinding and annealing activities. In cells, the proteins have been linked to several pathways—most notably splicing and transcription. p68 was isolated as a component of the spliceosome, and specifically is required for proper assembly–disassembly of the U1 5′ splice site. Interestingly, p68 and p72 interact with several components of the transcriptional machinery, including RNA polymerase II and CBP/p300. This raises the interesting possibility that one role of p68 is to coordinate/localize the splicing machinery with the transcription apparatus. Independent of this, p68 acts as a transcriptional cofactor for several factors, including CBP/p300, p53, and the estrogen receptor α [ERα]. For example, etoposide-mediated induction of p53-responsive genes is dependent on p68. Remarkably, this requirement is not dependent on p68 helicase activity, as an ATPase mutant allele of p68 rescued induction by p53. RNA binding, however, was still required. Similarly, ERα coactivation is not dependent on helicase activity. This suggests a model whereby p68 and possibly p72 mediate assembly of transcription complexes at promoters and recruit the splicing machinery to the promoter, thus facilitating cotranscriptional splicing events.

Recently, p68 and p72 were implicated as regulators of processing for many miRNAs [Fukuda et al. 2007]. This was first demonstrated in studies on mice that are homozygous null for either p68 or p72. For both genes, loss of function is lethal; p68-null mice die at embryonic day 11.5, and p72-null mice are postnatal lethal. miRNA expression analysis of null embryos indicated reduced mature miRNA levels. Whether the reduction of these miRNAs was due to loss of function in the respective helicase, or whether the mutant phenotype led to aberrant miRNA expression due to developmental alterations, remains to be addressed. However, mouse embryonic fibroblasts (MEFs) derived from p72-null embryos had a reduction in at least two miRNAs [miR-16 and mir-145], but not all miRNAs, and this reduction could be rescued by introduction of ectopic p72. pri-miRNA expression was unaffected, demonstrating an alteration in processing of the miRNA precursors. Interestingly, ATPase mutant alleles of p72 did not rescue production of either miR-16 or mir-145. This is in contrast to the transcriptional coactivation role for p68, which was not dependent on helicase activity. Therefore, for at least these two miRNAs, a role for helicase activity (or possibly ATP-dependent conformation change) is required. Furthermore, cell-free extracts that lack either p68 or p72 are inefficient at conversion of pri-miR-16 to the corresponding precursor miRNA. RNA immunoprecipitation [RIP] experiments in the same study showed that Drosha recruitment to pri-miR-199a and pri-miR-214 was lost upon RNAi-mediated depletion of p68 or in p72 knockout cells, consistent with the observation that mature miR-199a and mir-214 levels are lower in p68/p72-null embryos. Thus, in vivo, these helicases seem to be required to properly recruit the Microprocessor to some pri-miRNAs. As discussed below, it is now clear that multiple critical cellular signaling pathways use the p68 and p72 association with Microprocessor to effect regulation of pri-miRNA processing.

**Transforming growth factor β/bone morphogenic protein (TGF-β/BMP)**

The first study to report a signal transduction-mediated change in miRNA biogenesis demonstrated that the TGF-β and BMP pathways specifically promote processing of pri-miR-21 [Davis et al. 2008]. The contractile phenotype of human smooth muscle cells treated with TGF-β or BMP was found to be dependent on the rapid up-regulation of miR-21. Interestingly, this rapid increase in miR-21 occurred post-transcriptionally; quantitative real-time PCR [qRT–PCR] experiments revealed that pre-miR-21 and mature miR-21 levels quickly increased upon TGF-β and BMP treatment, while levels of pri-miR-21 were unchanged. Davis et al. [2008] then reasoned that SMAD proteins might be the intermediaries responsible for connecting these signaling pathways with miR-21 processing, as they are translocated into the nucleus upon TGF-β or BMP signaling and are known to bind DNA. Furthermore, a previous study reported that SMAD1 interacts directly with the RNA helicase p68 [Warner et al. 2004]. Accordingly, depletion of receptor-specific SMADs [SMAD1 and SMAD5] or p68 abrogated the BMP and TGF-β-driven increase in pri-miR-21 processing. Some initial mechanistic insight was also provided, GST pull-downs with recombinant SMADs in nuclear extracts indicated that p68 binds to the MH2 domain of R-SMADs while the MH1 domain binds to pri-miR-21 either directly or through other unknown factors.

The full complement of miRNAs that are regulated by this pathway is presently unknown. Since the TGF-β and BMP signaling pathways regulate many biological phenomena, it is possible that these pathways stimulate processing of other pri-miRNAs; indeed, Davis et al. [2008] also observed that pri-miR-199a processing is regulated in this manner. This is not surprising, since these signaling pathways converge on miRNA processing via p68, and this helicase has been shown to modulate a number of miRNAs.

**The p53 DNA damage pathway**

Unexpectedly, a subsequent study demonstrated that the tumor suppressor p53 also promotes pri-miRNA processing. It was noted that many miRNAs depleted in p72 knockout mice are coincidentally also up-regulated by the DNA-damaging agent Doxorubicin [Dox]. As Dox strongly stimulates p53 activity, leading to induction of p53-responsive genes, it was perhaps not surprising that
p53 might also induce the expression of miRNAs. In fact, miR-34 had been well established as a direct p53 transcriptional target (He et al. 2007). However, after measuring pri-miRNA expression levels of induced miRNAs, Suzuki et al. (2008) realized that these miRNAs were not being induced transcriptionally. Rather, processing at the Drosha step was triggered. In hindsight, this was not so surprising: p68 was known to regulate pri-miRNA processing, and the helicase had been shown to interact with p53 as a coactivator; ergo, p53 might be able to directly modulate miRNA processing (Bates et al. 2005). Accordingly, coimmunoprecipitation experiments as well as in vitro Drosha processing assays demonstrated that p53 associated with the large Drosha complex upon treatment of cells with Dox. Intriguingly, p53 alleles with known oncogenic activity decreased processing efficiency of p53-targeted miRNAs by disrupting p68’s association with Drosha, thus displacing the Drosha complex from selected pri-miRNAs. Again, the biochemical basis of p53’s association with p68 and the subsequent increase in target pri-miRNA processing remains obscure. Detailed structure–function studies will be needed to understand how p68 binding to p53 (both wild-type and oncogenic alleles) modulates Drosha’s ability to process pri-miRNAs.

ERα/estrogen

Another interesting report revealed how a signaling cascade can negatively regulate pri-miRNA processing. Kato and colleagues (Endoh et al. 1999) noticed that ERα was shown previously to interact with p68 and p72. Augmentation of the ERα signaling pathway produced a striking pattern of changes in certain miRNAs, many miRNAs were increased in ERα−/− mice (Yamagata et al. 2009). Furthermore, administration of estradiol [E2] to ovariectomized mice reduced the same miRNAs within the uterus. Importantly, this effect of E2/ERα could be recapitulated in the human breast cancer cell line MCF7. The physiological importance of ERα-regulated miRNA biogenesis was evident from the observation that the 3’ untranslated region [UTR] of VEGF, an ERα target gene, is targeted by ERα-repressed miRNAs. Yamagata et al. (2009) then demonstrated in vitro that E2-bound ERα could directly inhibit Drosha processing of ERα target pri-miRNAs. This study delved into more biochemical detail than the reports mentioned above: It was determined that p68 and/or p72 bridge the interaction between Drosha and E2-bound ERα; this interaction requires the C-terminal domain of Drosha and the N terminus of ERα. Thus, when ERα is recruited to the large Drosha complex in an E2-dependent manner, Drosha is dissociated from ERα-targeted pri-miRNA loci. It will be interesting to determine whether ERα/E2 weakens the overall integrity of the large Drosha complex, or simply its affinity for certain pri-miRNAs.

A pri-miRNA processing holoenzyme? Facts and future directions

As mentioned above, Drosha isolated from mammalian extracts exists in a large complex with many different RNA-binding proteins. The role of most of these Drosha-associated factors in pri-miRNA processing is not clear. It seems entirely possible that many of the factors purified in the large cellular Drosha complex may be the result of a nonphysiological RNA–protein complex assembled upon cell lysis. In support of this, while the dsRNA-binding proteins NFAT-90 and NFAT-45 were shown previously to be Drosha-associated factors (Gregory et al. 2004), one recent study showed that this heterodimer does not interact with Drosha and, in fact, inhibits processing of certain pri-miRNAs (Gregory et al. 2004; Sakamoto et al. 2009). This study underscores the need to more carefully analyze the functional interactions between Drosha–DGCR8 and the many auxiliary factors that are contained within the large complex. As with p68 and p72, there may indeed be important roles for other Drosha-associated factors, as general Microprocessor cofactors or as factors that integrate specific cell signaling pathways with pri-miRNA processing.

So, does Drosha/DGCR8 function within the context of a larger holoenzyme in vivo? The studies mentioned above suggest that, of the Drosha-associated factors initially identified, p68 and p72 are the first to be assigned as important cofactors in Drosha processing of certain pri-miRNAs. Taken together, the aforementioned studies suggest a model shown in Figure 1. Drosha processing is known to occur cotranscriptionally (Kim et al. 2009), p68 has been proposed to link mRNA transcription to splicing (Fuller-Pace and Ali 2008). Therefore, p68 and/or p72 might perform a similar function with miRNAs. That is, p68 might act as a bridge between the Microprocessor and specific miRNAs, at the RNA polymerase holoenzyme during transcription. This effect might be independent of helicase activity, as is the role of p68 in mRNA transcriptional events. However, p72 activation of miR-16 and miR-145 processing is ATP-dependent, suggesting a second, helicase-dependent role for p72 (Fukuda et al. 2007). This is further supported by the fact that p72 modulates Microprocessor function in cell-free extracts that is unrelated to transcription. It is important to note that all of these cell-free assays are based on crude extracts or on partially purified components. Without an exact understanding of the protein composition of the assays, it is difficult to fully understand the role of these accessory proteins.

It’s all in the loop: specific interactions between regulatory proteins and the terminal loop of miRNA precursors

As discussed above, one mode of regulated miRNA biogenesis occurs through modulation of Microprocessor activity. A second regulatory phenomenon occurs when RNA-binding proteins bind directly to specific sequences within pri-miRNAs, diverting the RNA away from biogenesis events, and often leading to degradation of the RNA.

Lin28 as a master regulator of let-7 production

let-7 is one of the most abundant miRNA families in mammals, with high expression in essentially all adult
tissues. In embryonic cells, in contrast, mature let-7 is present at 1000-fold lower levels. Unexpectedly, pri-let-7 expression is constant throughout development, thus providing a dramatic illustration of post-transcriptional regulation of a miRNA (Thomson et al. 2006). Research published over the past 2 years has demonstrated that the RNA-binding protein Lin28 blocks let-7 miRNA maturation in early embryonic cells, and the related Lin28B contributes to let-7 reduction in cancer (for review, see Viswanathan and Daley 2010). Initial studies demonstrated that Lin28 represses Drosha and Dicer processing of let-7 in vitro by binding to conserved sequences in the precursor loop. Furthermore, overexpression of Lin28 in differentiated cells and knockdown of Lin28 in embryonic cells resulted in the depletion and accumulation of mature let-7, respectively. The exact mechanism whereby Lin28 blocks processing steps is unknown. What was clear is that the blocked pri-miRNA/pre-miRNA was not accumulating in cells, suggesting a turnover mechanism. We now know, in the case of the pre-let-7, that Lin28 directly recruits a poly(U) polymerase, terminal (U) transferase (TUT4), to the precursor RNA (Hagan et al. 2009; Heo et al. 2009; Lehrbach et al. 2009). Oligo-uridylation of the precursor occurs, which triggers degradation. The fate of the blocked pri-let-7 is still unknown (see Fig. 2).

The story of Lin28 thus comes full circle. This embryonic stem (ES) cell-specific RNA-binding protein was one of the earliest-recognized let-7 targets (Reinhart et al. 2000). In embryonic development, its expression is reciprocal to that of mature let-7. Concordantly, Lin28 overexpression, in combination with several ES-specific transcription factors, is sufficient to reprogram somatic cells into induced pluripotent stem (iPS) cells (Yu et al. 2007).

KH-type splicing regulatory factor (KSRP) binds let-7 loop sequences and activates miRNA processing

Recently, a role was uncovered for the RNA-binding protein KSRP in promoting miRNA biogenesis in mammals (Trabucchi et al. 2009). This protein binds specifically to 5´guanosine-rich patches on the loop region of several miRNA precursors, including let-7. Knockdown of KSRP significantly decreases the expression of these miRNAs, in part by preventing efficient recruitment of Drosha and Dicer to the pri-miRNA and pre-miRNA, respectively. The obvious question is: How can Lin28 and KSRP bind to the let-7 loop, but with opposing terminal outcomes? Interestingly, while recombinant fragments of KSRP bind to the let-7 terminal loop with high affinity in vitro, KSRP cannot bind to pri-let-7 in embryonic cells, which have abundant Lin28. This suggests a model whereby Lin28 and KSRP bind in a mutually exclusive manner. Specifically, Lin28 may sterically hinder KSRP binding even though the two proteins bind to unique sequences on opposite sides of the let-7 loop. Alternatively, an unidentified factor in embryonic cells could bind cooperatively with Lin28 to let-7 loops, thereby preventing KSRP binding.

Regulation of miRNA processing by hnRNP proteins

While the above examples point to regulators with specific functions, there are other examples of RNA-binding proteins with a pleiotropic function in RNA biogenesis. One such example is the highly abundant RNA-binding protein hnRNP A1. This protein has been well established as a component of many hnRNPs and performs essential functions in many RNA processing and transport pathways (He and Smith 2009). Using cross-linking
immunoprecipitation (CLIP) (Ule et al. 2005), Guil and Caceres (2007) demonstrated that hnRNP A1 interacts with specific nucleotides of the loop and stem of pri-miR-18a. This miRNA is one of six miRNA stem–loops on the oncogenic polycistron miR-17-92. The role for hnRNP A1 in miRNA processing was demonstrated when HeLa extracts depleted of hnRNP A1 by RNAi failed to process pri-miR-18a from pri-miR-17-92, while the other pri-miRNAs in the polycistron were still processed. Interestingly, when recombinant hnRNP A1 was added back to the depleted extract, pri-miR-18a processing was not restored; this suggests that hnRNP A1 might have to be “precomplexed” with other RNA-binding factors in order to act specifically on pri-miR-18a. Of course, it is also possible that bacterially purified hnRNP A1 is missing post-translational modifications that are required for the positive regulation of pri-miR-18a processing.

Further studies demonstrated that hnRNP A1 binding to the loop and stem regions of pri-miR-18a relaxes the stem region and facilitates cleavage by Drosha. Bioinformatic analysis and RNA affinity purification using miRNA loop sequences, as well as in vitro Drosha assays, revealed that many hnRNP proteins could potentially form an extensive interaction network with miRNA loops (Michlewski et al. 2008). Thus, hnRNPs may broadly control the levels of miRNAs at the post-transcriptional level. The general physiological functions of hnRNPs are largely unknown, although their interactions with miRNAs and non-miRNA-containing substrates may play critical roles in gene expression regulation.

Contrasting mechanisms allow fine-tuned expression of miRNAs

We discussed two paradigms for regulated miRNA biogenesis. Control at the Microprocessor allows integration of signaling events from many pathways, leading to changes in expression of divergent miRNA families. All of the discussed events are mediated by the helicases p68 and p72. In many cases, the regulated miRNA families are similar, but there are notable exceptions. So how does this pathway allow specificity? Some interaction with the pri-miRNA must confer specificity. In the case of SMAD regulation of miR-21 processing, specificity could arise from interaction of the pri-miRNA with the SMAD-interacting protein SNIP1. This coactivator protein had independently been shown to affect miRNA processing, and its Arabidopsis homolog has direct RNA-binding activity (Yu et al. 2008). In other cases, the key regulatory protein might itself interact with the pri-miRNA. As an example, p53 can immunoprecipitate pri-miR-16; but is there a direct interaction between p53 and the pri-miRNA, or is it bridged by p68? Interestingly, many transcription factors, including p53, have been shown to have RNA-binding activity (Riley and Maher 2007). Such interactions may provide the critical specificity. Of course, unidentified proteins may be part of the Microprocessor, possibly as a transient complex assembled after signal reception. Clearly, to fully understand this complex, we need better characterization of components, and ultimately reconstitution in defined activity assays. In contrast with the Microprocessor paradigm, the loop-binding regulators clearly have a mechanism for specificity. In the
case of Lin28, the expression of that regulator is confined to the stem cell compartment, providing an obvious mechanism of control. KSRP and the various hnRNP proteins, in contrast, are widely expressed. Their biological function in the miRNA pathway is still poorly understood.

In addition to these paradigms, other regulatory events during miRNA biogenesis have been described, but could not be included in this review. For example, the metabolite heme can modulate Microprocessor function, although the biological role of this effect is unclear [Faller et al. 2007]. The nuclear cap-binding complex component ARS2 is important for fidelity of Microprocessor cleavage [Gruber et al. 2009]. The miRNA primary sequence can be altered by A–I editing, leading to altered processing and function [Yang et al. 2006]. When taken into consideration, it is clear that miRNA processing is regulated in a complex manner, and we are only beginning to understand its true nature.

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