Keywords: microRNA; Regulation; Tumorigenesis; Metastasis; Mechanism

Introduction

Small non-coding RNAs, identified so far including small interference RNAs (siRNAs), microRNAs (miRNAs), piwi-interacting RNAs (piRNAs), promoter-associate (paRNAs), transcription initiation tiRNA, and tRNA-derived RNA fragments (tRFs), are 18-30 nucleotides in length. Although they account for only a very small fraction of the expressed genome, these small RNAs have been acknowledged as pivotal regulators at post-transcriptional level to control diverse biological pathways relating to chromosome architecture and segregation behavior, transcription, and RNA processing and stability, and thereby to phenotypically coordinate development, growth control, apoptosis, self-defense, and stem cell maintenance [1-6]. Accordingly, as expected, increasingly obvious evidences demonstrate that alteration of miRNAs expression profile could contribute to the pathogenesis of a wide range of human diseases as well as phenotypic abnormality among the eukaryotes [7]. Moreover, siRNA/miRNA based knockdown has been technically adopted in a wide range as powerful tools to elucidate gene functions, to discover and validate the drug targets, and to develop therapeutic intervention. Given the pivotal roles of siRNA/miRNA in diverse biological pathways as well as the broad application of RNAi, it is essential to understand the siRNA/miRNA pathway and its regulation mechanism. Since the identification of the major components of the RNAi/miRNA pathway [2], focus has been switched to the regulation mechanism of the pathway itself in recent years, resulting in discovery and characterization of co-factors functioning as regulators of the RNAi/miRNA pathway at different levels from pre-transcription to post-transcription.

miRNA Processing Pathway

In mammals, majority of miRNA genes are transcribed initially by RNA polymerase II as primary transcripts (pri-miRNAs) range from hundreds to thousands of nucleotides in length that bear one or more hairpin structures [8]. The pri-miRNAs are 5'-capped, spliced and polyadenylated [5,9,10]. Subsequently, two sequential biochemical processing steps lead to generation of mature form of miRNAs. In most cases, the first step takes place inside the nucleus where the pri-miRNAs were cropped by Drosha, a RNA endonuclease III in mammals and its partner DGCIR8 [10] to produce a ~70 nt stem-loop-structured precursor (pre-miRNAs) that harbors the miRNA in the 5’ or 3’ half of the stem [8,11]. However, this is not the only way of pre-miRNAs biogenesis. Alternatively, generation of some pre-miRNAs can bypass the Drosha processing step. One example is “mirtrons,” the pre-miRNA-like hairpins which are generated by splicing and debranching of the short hairpin introns [12-14]. Besides “mirtron”, some small nucleolar RNAs (snoRNAs), tRNAs, and endogenous short hairpin RNAs (shRNAs) are processed into miRNA-like molecules without relying on Drosha processing either [15-18]. More recently another source of miRNAs, terminal hairpins of endogenous siRNA long-stem-loop precursors, has been identified as being able to bypass the Drosha processing step [19].

The pre-miRNAs are transported into cytoplasm by exportin-5/RanGTP for the secondary processing [8,10,20,21]. The second processing step occurs in the cytoplasm where the pre-miRNAs are further dic to mature form of miRNAs with length of 18-25 nt by Dicer, another member of RNase III family in collaboration with its cofactors TRBP and PACT [8,22,23]. Once maturated, the guide/functinal strand and the passenger strand of the miRNA duplex are facing different fates. It is generally
believed that the guide strand is sorted to load into RISC complex to regulate gene expression at post-transcriptional levels, while the passenger strand is finally degraded [24]. However, recent deep sequencing results in Drosophila have indicated that a large number of passenger strands are functional and associated with AGO1 or AGO2 instead of being degraded [25-28]. Of the sequence features determining the selection of guide strands, the thermodynamic asymmetry of the miRNA duplex contributes greatly to the selecting process [29,30], while other features such as the central mismatches and the identity of the 5’ nucleotide affect sorting as well. The mature form of miRNAs function as recognizer once they are loaded into the RNA induced silencing complex (RISC) to match their target mRNAs through base-pairing. The RISC consists of several components with AGO protein as core for RNA silencing [5,31,32].

Regulation of microRNA Pathway

Any essential regulators must be tightly regulated to maintain normal cellular homeostasis. Similarly, misregulation of miRNAs/siRNAs expression levels alters their target levels, severely affecting normal cellular metabolism. To precisely regulate the expression levels of individual miRNA/siRNA, a serial of regulatory mechanisms along the miRNA/siRNA pathway have been discovered at levels of genome, epigenome, and transcriptome.

Regulation at Level of miRNA Gene Sequences

Several events at the DNA level contribute to alteration of miRNA biogenesis such as copy number change or mutations in the miRNA genes. So far very few reports are available at this level and most of them are found in human cancer cells [33]. Mutations were identified in 5 of 42 tested miRNA genes in patients with chronic lymphocytic leukemia (CLL), but not in the normal healthy controls. More specifically, mutation of 7 bp downstream from pre-miR15a–6-1 genomic DNA in germline, inhibited processing. A more common form of point mutations is the single nucleotide polymorphism (SNP) which alters the miRNA processing and in some cases the miRNA functions as well. It has been shown that 315 SNPs were associated with 265 human miRNA genes and only 12 of these SNPs were found in miRNA precursor sequences, and one SNP was located in the miR125a seed sequence [34]. Some SNPs could confer the differences in the Drosha-mediated processing such as miR125a precursor SNPs, miR-502, miR-510, miR-890, and miR-892b [34-36], while differences were observed in the Dicer-conferred processing in some other SNPs such as miR-196a G/T SNPs as well [37]. However, mutations including SNPs are passive consequences of the mutations occurred in the miRNA genes. Therefore, it is not an active/auto miRNA processing regulation mechanism but rather pathologically associated with some human diseases. For example, miR-125a SNP is highly associated with breast cancer tumorigenesis [16]. Thus, SNPs could serve as pathological markers clinically.

Regulation at Epigenetic Level

Some epigenetic factors that affect transcription such as histone deacetylation and hypermethylation of promoter region of miRNA gene may also significantly inhibit or even completely silence the biogenesis of miRNAs. An example is that hypermethylation of miR-127 gene promoter region results in either dramatic inhibition or completely silencing of the miR-127 expression in many kinds of tumors more evidenced by the fact that treatments of these tumor cells with agents of demethylation and histone deacetylase inhibitor dramatically increase the miR-127 levels [38].

Regulation at Transcriptional/Posttranscriptional Levels

Regulations of miRNA processing at post-transcriptional level are thought to be the most important part of the regulation mechanisms. Several factors contribute to the regulations at RNA level. One of the factors is the modification of pri-/pre-miRNA sequences which affect miRNA processing but do not alter activity of the RNAi/miRNA processing machinery. A more common mechanism is the alteration of the activity of the RNAi/miRNA processing machinery. Modulations of the RNAi/miRNA processing machinery activity are conferred by protein-protein interactions such as the co-factors that interact with Drosha and Dicer, or by protein-RNA interactions such as the co-factors that interact with the terminal loops of the pre- and/or pre-miRNAs. Indeed, more and more protein components are being identified to enhance or inhibit the miRNA processing efficiency at post transcription level.

Modifications of pri-/pre-miRNA sequences

Pri-/pre-miRNA sequence modifications, typically RNA editing could alter the miRNA processing efficiency. The editing characterized so far is conversion of adenosine (A) to inosine (I) by adenosine deaminases on RNA (ADARs) in dsRNA structures [39,40]. Two isoforms of ADAR, ADAR1 and ADAR2 are responsible for the editing. The editing takes place in both pri-miRNAs and pre-miRNAs and most likely in nuclei due to the predominant nuclear localization of ADARs. In addition, the pre-miRNAs may also be edited in cytoplasm, caused by ADARs shuttle in and out of nucleus [29,40]. The edited pri-miRNAs may block the Drosha-mediated processing such as primiR-22 and primiR-142 because the editing occurred at the position surrounding the Drosha cleavage site [41,42]. However, some edited primiRNAs such as pri-miR-142 and pri-miR-151 do not affect the Drosha-mediated processing but rather hinder the Dicer-mediated processing [29]. The editing does not only affect the miRNA processing, but also interfere with the miRNA functions due to alterations of the “seed” region crucial for the recognition and binding of the RISC to targets. This interference has been confirmed in the edited miR-376 [29]. Even though the edited pri-miRNAs could not be appropriately processed by Drosha or Dicer, their accumulation was not detectable in nuclei, suggesting that specific degradation mechanism must be adopted to maintain homeostasis. Under in vitro conditions, the edited pri-miR142 could be cleaved by Tudor-SN [42]. It has been shown that Tudor-SN is a component of RISC and has ribonuclease activity specific to inosine-containing dsRNAs [43]. However, so far it is still lacking evidence for in vivo relevance of the degradation of the edited pri-miRNA by Tudor-SN. Since the ADARs-based editing is not a pathological event at least based on the current research data, this represents an auto-regulation mechanism. However, the physiological significances are not clear and the fate of the edited miRNAs remains to be elusive.

Co-factors interaction with major components of the RNAi/miRNA pathway

Of the co-factors regulating miRNA processing, the DEAD-box RNA helicase p68 and p72, central tumor suppressor p53, nuclear factors NF90 and NF45, Ars2 and CBC, and the splicing factors SF2/ASF have been shown to modulate the Drosha-mediated primiRNA processing [44-50]. More specifically, the p68/p72 complex has dual functions: enhancement or inhibition in regulations of the Drosha-mediated processing of a subset of miRNAs depending on the protein components with which the p68 interacts [45,48,51]. Interaction of p68 with p53 or SMAD, induced by transforming growth factor β (TGF-β) and bone morphogenetic proteins (BMPs), could facilitate the
10 processing through promoting recruitment of Drosha complex to the target pri-miRNAs. On the other hand, interaction with estrogen receptor alpha (ERα) inhibits the pri-miRNA processing. In addition to p68/p72 complex, Ars2 and CBC are believed to serve as recruitment factors to enhance the pri-miRNA processing as well by binding the microprocessor [46]. More recently, it was found that the splicing factors SF2/ASF could directly interact with Drosha to facilitate Drosha-mediated processing of a subset of pri-miRNAs such as miR-7, miR-215, miR-215, and miR-222, and so on [52].

In addition to the positive regulators for the Drosha-mediated processing, a negative regulator complex NF90/NF45 was found to inhibit the processing step by reducing the accessibility of Drosha-DGCR8 to pri-miRNAs rather than by interaction between NF90/NF45 and the endogenous Drosha-DGCR8, evidenced by the high affinity of the NF90/NF45 to pri-miRNAs in Drosophila versus the lack of available data to show its interaction with Drosha-DGCR8 complex at present. Unexpectedly, while overexpression of NF90/NF45 in 293 cells enhanced accumulation of pri-let-71-1, primiR-21 and pri-miR-15a-16-1, respectively, the mature miRNA levels were not significantly altered [47], suggesting that the unknown mechanism(s) is in compensation with decreased pri-miRNA processing efficiency induced by NF90/NF45 overexpression. Given the dual regulations of the Drosha-mediated primiRNA processing by positive and negative regulators, it is rational to believe that the balance between contributions made by the positive and negative co-factors involved in the processing could maintain the homeostasis of the pri-miRNAs, pre-miRNAs, and the mature miRNAs.

Like in the first step processing, more co-factors/signaling pathways are also involved in the Dicer-mediated pre-miRNA processing. TRBP and PACT were first characterized as co-factors not essential to the processing but facilitate the RISC assembly. However, further study showed that TRBP phosphorylation mediated by the mitogen activated protein kinase (MAPK) signaling pathway could stabilize the Dicer-containing complex [53]. This is the first report regarding post-translational regulation of miRNA/RNAi pathway. Another co-factor, Wig-1, a p53-induced zinc finger protein can interact directly with Dicer to increase the activity of Dicer [54].

Regulation by Argonaute protein family members

Argonaute protein family members (AGO1–4), particularly AGO2 in mammals are confirmed to function as RISC-based miRNA cleavage or translational repression. Surprisingly, these AGO members themselves also enhance the miRNA processing, evidenced by the fact that ectopic overexpression of Agol-4 significantly elevated levels of some mature miRNAs such as miR-215, miR-17-5p, miR-23b, and miR-92 [55]. What is more, using its intrinsic endonuclease activity in mammals, like Dicer, AGO2 could cleave the pre-miRNAs to an additional processing intermediate which may facilitate miRNA duplex dissociation and formation of RISC complex [55]. More recently, two groups found a Dicer-independent but Argonaute 2 catalytic activity-dependent novel miRNA processing pathway [56,57]. In vivo and in vitro data indicated that pre-miR-451 was loaded into RISC complex and cleaved by AGO2 followed by uridylation and trimming to generate functional miRNAs. This novel pathway might not be a main pathway for miRNA biogenesis, but it links the conservation of Argonaute catalysis to a conserved mechanism of miRNA biogenesis [56,57].

Co-factors interaction with RNA stem loop structure

Some co-factors regulate miRNA processing not only through interaction with Drosha or Dicer but also through binding with terminal loop structures in the pri-miRNAs or premiRNAs. One RNA terminal loop binding protein involved in pri- and pre-miRNA processing is the KH-type splicing regulatory protein (KSRP) [58], a component of both Drosha and Dicer complexes. KSRP binds to the terminal loop of its target pri-and pre-miRNAs (eg. miR-21, miR-16, miR-155, and let-7a) and enhances Drosha and Dicer-mediated processing through interactions with Drosha and Dicer [58]. In contrast to these positive cofactors, Lin28, a developmentally regulated RNA binding protein was identified to selectively inhibit the processing of pri-let-7 by competing for binding to the conserved nucleotides in the loop of pri-let-7 [59–62]. Another mechanism that Lin28 negatively modulates processing of pre-let-7 is adding an oligouridinyl tail to the pre-let-7 in cytoplasm [63]. It has been proven that Lin28 itself does not uridylate pre-let-7 but it could recruit terminal uridylyl transferase 4 (TU/Tase 4, or TUT4), which has been demonstrated to be responsible for the uridylation [63–65].

Another RNA binding protein that so far has been proven to be involved in miRNA processing is heterogeneous nuclear ribonucleoprotein A1 (hnRNPA1). hnRNPA1 has been indicated to facilitate processing of pri-miR-18a, a member of miR17-92 cluster by binding to both the terminal loop and a region in the stem [66,67]. However, so far no report is available for the effect of hnRNPA1 on processing of pre-miR-18a. In addition to pri-miRNA-18a, other pri-miRNAs such as pri-let-7a-1 and pri-miR-101-1 are also found to be the binding targets of hnRNPA1, but no evidences have shown the hnRNPA1 conferred enhancement or inhibition of the processing in let-7a-1 and miR-101-1. Since only few RNA terminal loop binding proteins and few pri- and pre-miRNAs were tested for processing efficiency alterations, it is too early to draw a conclusion on whether the terminal–loop RNA binding protein-mediated modulation of the miRNA processing efficiency is a common regulation mechanism.

Regulation of mature miRNA stability

Accumulation of mature form of miRNAs depends on regulations at levels of transcription, processing, recycling and turning over. While it has been acknowledged that regulation at any of these processing steps is a key determinant, the stability of mature miRNAs after processing serves as an important element for the homeostasis of functional miRNAs. Several recent findings address the mechanisms for controlling the decay of mature miRNAs. The first factor that affects the stability of mature miRNAs is Argonaute protein levels [55,68–70]. In addition to functioning as enhancement of biogenesis from pre-miRNAs to mature miRNAs in some cases, Argonaute proteins can protect miRNAs from degradation by taking guide strand as "refugees". The second factor affecting mature miRNAs is the protective modifications such as methylation [71], uridylation [72], adenylation [73–76], diverse nucleotide substitutions, additions, and deletions [73–75,77]. The methylation catalyzed by HEN1 methyltransferase to add methyl group to the 3'ends of miRNAs is the first characterized mechanism for mature miRNAs modification in Arabidopsis [71]. The HEN1 mutation conferred substantially decreased miRNA levels and the enhanced uridylation of the 3'ends, suggesting the methylation of the 3'ends could prevent uridylation and degradation in plants [71]. But the enzymes responsible for uridylation have not yet been identified. By contrast, as far as the substrate is concerned in animals, the HEN1 methyltransferase only methylated piRNAs, and in some cases, siRNAs [78–80], no miRNAs have been found to be methylated by it. The uridylation has dual effects on the stability of the miRNAs [72] in plants. It seems that the uridylation of the passenger strand equals to addition of a tag to promote degradation, instead the uridylation of the guide strand could serve as a protective modification [72] as
the uridylated miRNAs favors the Argonaute bindings. However, so far no uniform modification has been yet found in animals. 3' adenylation by the cytoplasmic poly(A) polymerase GLD-2 (also known as TUtax2) could stabilize miRNAs in animals. However, of the animal miRNAs identified to be adenylated, only miR-122 in liver has been characterized for functional consequence [51]. It seems that the GLD-2 non-specifically adds adenine residues to miRNAs, but only a fraction of these adenylated miRNA species could be stabilized. Similarly, adenylation also occurred in both full length and truncated plant miRNAs [81], but the factors responsible for the adenylation and the related functional consequences are still waiting for elucidation. While diverse nucleotide substitutions, additions and deletions have been detected in animals, and the types and the extents could affect their target recognitions, possible effects on miRNA stability are less predictable [73-75,77].

The third factor is the degradation factor for mature miRNAs. The small RNA degrading nuclease (SDN) is the first exonuclease identified in plants to catalyze 3'- to 5'- cleavage of mature miRNAs [82]. The methylated and uridylated miRNAs are more resistant to degradation by SDN in vitro conditions. By contrast, the 5'- to 3'-exonuclease XRN-2 is responsible for degradation of the single strand and nonincorporated miRNAs in animals [83]. While the functional miRNAs take refuge in RISC complex to avoid degradation, cells apparently have specific mechanisms to release the miRNAs from Argonaute to expose the released miRNAs for degradation. More specifically, the XRN-2-mediated degradation is in target availability dependent manner [83]. More recently, it was found that the XRN1 instead of XRN2 could be responsible for miR-382 which contributes to HIV provirus latency and is very unstable in human cells [84]. At normal physiological conditions, the rates of biogenesis and turnover determine the accumulation of functional miRNAs. The appropriate ratio of miRNAs to their targets levels could be maintained to guide RISC reprogramming or turnover. The tight control of the functional miRNA homeostasis is the key to prevent detrimental overexpression or deficiency in miRNAs.

Co-factors enhancing RISC loading or RISC-mediated Silencing

Although many co-factors were demonstrated to be involved in processing of miRNAs, few co-factors have been identified to be associated with AGO proteins in mammals or in flies to enhance RISC loading or RISC-mediated silencing. The protein components so far identified include putative RNA helicase, RNA binding, heat shock protein, P-body component, endoribonuclease, type I collagen prolyl-4-hydroxylase, and so on. MOV10, a putative RNA helicase, and TNRC6 (-A, -B, -C) are associated with Ago1 and Ago2 and functionally required to mediate miRNA-guided mRNA cleavage [85,86]. RBM 4, a putative RNA-binding motif protein can efficiently bind miRNAs and interact with AGO1 and AGO2 in RNA dependent manner, and is required for microRNA-guided gene regulation [87]. Hsp90, a heat shock protein, is associated with AGO2 to efficiently target hAgo2 to P-bodies and stress granules and thereby enhance microRNA- and siRNA-confereed gene silencing [88]. GW182, a P-body component in Drosophila, and TNGW1, a 210 kDa isoform of the GW182 in human, both interact with AGO1 and the interaction is essential for miRNA-mediated translation repression and miRNA turn over [89,90]. The α-(P4H-α(I)) and β-(P4H-β) subunits of type I collagen prolyl-4-hydroxylase (C-P4H(I)) interact with Ago2 and get Ago2 and Ago4 more efficiently hydroxylated than Ago1 and Ago3. The hydroxylation maintains stability of Ago2 and efficiency of RNAi activity [91]. Imp8 functions as a gene silencing factor to stimulate RISC-mediated silencing by enhancing association of AGO2 complexes to target miRNAs [92]. C3PO, an endoribonuclease, promotes RNAi by facilitating RISC activation in Drosophila [93]. However, it seems that some of them are not essential for RISC loading or target cleavage.

miRNA Intercellular Transportation

Both siRNAs and miRNAs are trafficking intercellularly instead of being completely restricted in specific cell from which the miRNAs are generated. Recent studies showed that RISC formation and turnover depends on the endosomal pathway [6,94], suggesting a link between RNA silencing and membrane trafficking. Endosome maturation involves a gradual accumulation of internal vesicles which will be finally released into the extracellular milieu called exosomes. The exosome containing the ribonuclease complexes were recently found to carry miRNAs and miRNAs and can transfer [95,96] these nucleic acids into a recipient cell. The purified exosome-like vesicles from the cultured monocytes contained a large number of miRNAs, low levels of AGO2, and high levels of GW182 [94]. P-body component required for miRNAs functions by binding to AGO proteins [96]. Microvesicles-based transfers of RNAs and proteins from cell to cell are believed to play important roles in intercellular communications. Recent studies have demonstrated intercellular trafficking of miRNAs by a variety of cell types such as normal somatic cells, cancer cells, and stem cells [97-101]. The mechanism for miRNA secretion and transfer is just being elucidated. The miRNAs are secreted by a ceramide-dependent secretory pathway and the secreted miRNAs are transferable and functional in the recipient cells [99]. Growing evidences have demonstrated the variety of physiological roles in which the secretory miRNAs may play [102,103], and one of such suggests that the secretory miRNAs could mainly serve in chronic biological events such as the formation of a tumor microenvironment.

MicroRNA and Tumorigenesis

Global deregulation of miRNAs biogenesis and cancers

MicroRNAs are predicted to regulate over 50% of all human protein-coding genes, enabling them function in many physiological and developmental processes [38]. Generally speaking, cancer is partially featured by global down regulation of miRNA expression in accordance with the accumulation of pri- and pre-miRNAs compared to normal tissue [59], while the specific deregulation of certain miRNAs is seen in specific tumor types [104,105]. Aberration of any key components in the microRNA pathway as summarized above may contribute to pathology of cancer genesis, progress, and development. Evidences have emerged to link the deregulation of these key components expression or activity and several malignancies, suggesting that the tight control of the microRNA pathway itself at any stages is essential in maintenance of normal metabolism.

Abnormality of key components involved in microRNA pathway and cancer

Global deregulation of microRNA expression levels has been linked to almost all the type of cancers identified so far, whereas the abnormality of the main protein components in the microRNA pathway accounts for at least one of the main causes for the global deregulation. Copy number abnormalities of some key coding genes such as Dicer, Ago2, Exportin 5 (XPO5), Drosha, TRBP and DGCGR that are essential for microRNA biogenesis in human genome have been detected in breast cancer, ovarian cancer, and in melanoma [11,106]. In addition to copy number abnormalities, abnormal expression levels of mature
form of microRNAs have been observed in some types of cancers due to deregulation of the expression levels of these components involved in the miRNA pathway or their activity alteration. Disruption of miRNAs biogenesis by depletion of any key components in miRNA processing pathway such as Drosha, DGC8R, Dicer, and TRBP, has been linked to promotion of oncogenesis [107-112]. Evidences have arisen in recent years from different animal models or cancer cell lines [107-112]. Under in vitro conditions, mutation of XPO5 inactivation enhances retention of microRNA precursors in the nuclei and accordingly decreases the levels of mature miRNAs in several human tumors [113]. More direct evidences arise from the fact that XPO5 knockdown enhances tumorigenicity of cells injected into mice in accordance with the reverse effect that overexpression of wild type of XPO5 in colorectal cancer cells expressing mutant XPO5 [113]. A serial of other evidences have suggested the essential roles of the normal microRNA biogenesis in inhibition of oncogenesis. In a mouse model of retinoblastoma, monoallelic loss of Dicer1 leads to accumulated tumor formation [108], consistent with the situations in different human tumors in a way that heterozygous instead of homozygous genetic deletion of Dicer 1 frequently occurs, suggesting that Dicer1 functions as a haploinsufficient tumor suppressor [109,110]. TRBP is one of the essential components in microRNA pathway in ways that maintain the stability of Dicer1 and microsatellite, and its concurrent impairment has been indicated in weakness or eradication of miRNA biogenesis and the resulted colon tumors formation/development [112,114,115]. Some small molecule compounds specifically or non-specifically up-regulate expression of miRNAs through direct or indirect interaction with key components involved in miRNA pathway. For example, enoxacin promotes miRNAs biogenesis through enhancing both processing of miRNA precursors to mature form of miRNAs and loading of miRNAs into RISC to increase the efficiency of RNAi [116]. Later on it is found that this compound could significantly inhibit cancer cell growth and metastasis in mouse model [117] due to elevation of a substantial set of miRNAs levels. Recently, a set of iron chelators have been found to interact with Dicer, a key enzyme for the biogenesis of miRNAs with the mechanism that cellular iron chelation enhances the interaction of DICER1 and Polyc binding protein 2 (PCBP2), binding of PCBP2 with miRNA precursors, and processing of precursors into mature miRNAs [118].

Specific deregulation of miRNAs and cancers

Global deregulation of miRNAs, due to alteration of the key components involved in miRNA pathway, is pretty common in cancers. Another important issue in miRNA mediated cancer is the deregulation of the specific miRNAs.

Specific miRNAs in tumorigenesis and tumor development

Tumorigenesis initiates from regulation imbalance of cell proliferation and apoptosis in ways that overexpression of oncogenes and/or silencing of tumor suppressor genes. MicroRNAs contribute to regulation of tumorigenesis and tumor development through negatively regulating expression of genes involved in the processes of cell proliferation and apoptosis. At molecular levels, some microRNAs target tumor suppressor genes, while some others are believed to regulate expression of oncogenes. In case of down regulation or null expression of oncogenes-specific miRNAs, the expression of the related target genes will be enhanced, causing or promoting tumorigenesis. Accordingly, overexpression of tumor suppressor genes-specific miRNAs leads to insufficient expression of targets functioning as tumor suppressors, causing tumorigenesis and tumor development. As a general phenomenon, reduced expression levels of miRNAs were detected in tumors relative to that in normal tissues [107]. Although it is not conclusive whether this phenomenon is exactly the cause or the result of the tumorigenesis, based on the previous studies it could suggest that while the majority of miRNAs may be involved in tumor suppression, alteration of miRNAs expression levels might be both the causes and results. As evidence, differentiation state of HL60 cells is maintained partially by enhanced miRNAs expression, and after differentiation treatment, HL60 cells display significantly increased expression of miRNAs [107]. Consistently, activation of the c-Myc oncogenic transcription factor (Myc) leads to the widespread repression of miRNA expression due to the fact that Myc directly binds to miRNA promoters, causing tumorigenesis [119], whereas enforced expression of repressed miRNAs diminishes the tumorigenic potential of lymphoma cells. Other study reports that attenuation of miRNAs biogenesis by genetic or RNAi approach promotes tumorigenesis and tumor development in lung cancer model [107]. All together it is demonstrated that extensive reprogramming of the miRNA transcriptome contributes to tumorigenesis.

At physiological levels, some miRNAs function as regulation of angiogenesis, while some miRNAs are thought to be involved in tumor metastasis. In most cases, tumorigenesis originates from deficiency of miRNA generation. Typically, miR-15a and miR-16-1 target Bcl-2, an essential anti-apoptosis factor that tightly regulates proliferation. Deficiency of both miRNA biogenesis leads to overexpression of Bcl-2 and consequently promotes tumorigenesis and tumor development. On the other hand, overexpression of some miRNAs leads to tumorigenesis as well. For example, miR-221 and miR-222 target KIT protein, proto-oncogene c-Kit or tyrosine-protein kinase Kit or CD117. Overexpression of both miR-221 and miR-222 dedifferentiates thryocytes and leads to tumorigenesis [104]. Similarly, miR-150 that targets a transcription factor c-Myb [120] and tumor suppressor EGR2 [52], is dramatically overexpressed in gastric cancer tissues and overexpression of miR-150 significantly leads to deficiency of EGR2 expression and therefore promotes gastric cancer cell proliferation [52].

miRNAs and Tumor Metastasis

Deregulation of some miRNAs leads to tumorigenesis and promotes progression of primary tumors. More importantly, some miRNAs have been linked to tumor metastasis as reviewed already [121-126]. Essentially, abnormal expression of miRNAs including mutation, down-regulation or overexpression may affect expression levels of their target genes involved in tumor metastasis such as cell migration, invasion, and anoikis resistance in ways that some cancer cell metastasis suppressors are down-regulated due to overexpression of the specific regulatory miRNAs or some metastasis enhancers are overexpressed due to the down regulation of some specific miRNAs. So far a set of miRNAs involved in metastasis have been identified (Table 1).

Metastasis–promoting miRNAs

Some miRNAs target to suppressors for metastasis named as metastasis-promoting miRNAs, their up-regulation is beneficial to tumor metastasis. miR-221 and miR-222 overexpression induced a down regulation of PTPμ that is believed to suppress cell migration. Phenotypically, miR-222 and miR-221 induce an increase in cell migration and growth in soft agar in glioma cells. Consistently, the re-expression of PTPμ gene is able to revert the miR-222 and -221 effects on cell migration. The miR-221/222 overexpression and invasiveness increase is further confirmed in human glioma cancer samples [127], suggesting that the miR-221 and -222 regulate glioma tumorigenesis at least in part through the control of PTPμ protein expression.

Citation: Jiang R, Li Y (2013) Regulation of Mirna Pathway and Roles of Micrornas in Tumorigenesis and Metastasis. Human Genet Embryol S2: 007. doi:10.4172/2161-0436.S2-007
Transcriptionally induced by Twist 1, a transcription factor, miR-10b has been known for its promotion to breast cancer metastasis and epithelial-mesenchymal transition evidenced by the fact that the miR-10b is highly expressed in metastatic breast cancer cells and positively regulates cell migration and invasion in a way that miR-10b inhibits translation of its direct target mRNA encoding for homeobox D10 and in turn activates another well-characterized pro-metastatic gene, RHOC [128]. Some miRNAs target metastasis suppressors, up-regulation of such miRNAs such as miR-182 will be beneficial to tumor metastasis. Metastasis suppressor 1 (MTSS1), one target gene of miR-182, plays an important role in the metastasis of cancers. Evidences from both in vivo and in vitro suggest elevated expression of miR-182 enhances metastasis through the down-regulation of its target gene MTSS1 [129]. Thus, miR-182 and MTSS1 were potential prognostic markers and/or therapeutic targets in HCC. By activation of signal transducer and activator of transcription 3 (STAT3), overexpression of PRL-3 induces expression of miR-21, miR-17 and miR-19a. Subsequently, induced expression of these miRNAs is involved in the proliferation and metastasis of colon cancer [130].

miR-10a functions as enhancement of the colony formation activity, migration and invasion of HeLa and C33A cells through targeting to CHL1. Independent experiment demonstrates that CHL1 serves as a strong repressor for colony formation activity, migration and invasion. Finally, overexpression of CHL1 lacked the 3'UTR abolished the effects of miR-10a, providing an alternative strategy for blocking tumor metastasis [131]. Similarly, induced expression by hypoxia, miR-103/107 target the known metastasis suppressors DAPK and KLF4 in CRC cells, resulting in increased cell motility and cell-matrix adhesion and decreased cell-cell adhesion and epithelial marker expression, and suggesting that this regulatory circuit may contribute in part to hypoxia stimulated tumor metastasis. Clinically, a miR-103/107 high, DAPK low, and KLF4 low expression profile has become a featured signature to serve as a prognostic marker for metastasis recurrence and poor survival, strategically suggesting the possibility that disruption of this circuit in blocking the CRC metastasis [132].

miR-9 targets MMP-14 and thereby downregulate expression of MMP14 and its downstream gene vascular endothelial growth factor (VEGF) in cultured NB cell
lines SH-SY5Y and SK-N-SH, inhibiting the invasion, metastasis and angiogenesis of NB [133].

Metastasis-inhibiting miRNAs

Some miRNAs target to metastasis enhancers named as metastasis-inhibiting miRNAs accordingly. Down-regulation of these miRNAs initiates or enhances tumor metastasis. MiR-335 suppresses lung and bone metastasis by human cancer cells in vivo through regulation of a set of genes whose collective expression in a large cohort of human tumors is associated with risk of distal metastasis. Thus, miR-335 is identified as metastasis suppressor microRNAs in human breast cancer [134]. MiR-126 suppresses metastatic endothelial recruitment, metastatic angiogenesis and metastatic colonization through coordinate targeting of IGFBP2, PITPN1 and MERTK- novel pro-angiogenic genes and biomarkers of human metastasis [135]. Human miR-373 and miR-520c have been characterized to stimulate tumor cell migration and invasion in vitro and in vivo mechanistically due to suppression of CD44. miR-373 and miR-520c are implicated as metastasis-promoting miRNAs [20]. MiR-7 is linked to significantly suppress metastasis in highly metastatic GC cell lines and metastatic tissues both in vivo and in vitro through translational inhibition of the insulin-like growth factor-1 receptor (IGF1R) oncogene. Thus, targeting this novel miR-7/IGF1R/Snail axis would be helpful as a therapeutic approach to block GC metastasis [136]. Through a Slit-miR-218-Robol regulatory circuit, miR-218 functions as suppressor for tumor cell invasion and metastasis in vitro and in vivo. MiR-218 targets Robol, one of several Slit receptors interaction, whereas decrease of miR-218 activates the Slit-Robol pathway through Robol and Slit 2 interaction, triggering tumor metastasis. Thus, targeting miR-218 may provide a strategy for blocking tumor metastasis. [137]. Targeting ILK, miR-625 significantly inhibits invasion and metastasis of gastric cancer cells both in vitro and in vivo [138]. MiR-372 and miR-373, each permits proliferation and tumorigenesis of primary human cells that harbor both oncogenic RAS and active wild-type p53. These miRNAs neutralize p53-mediated CDK inhibition, possibly through directly inhibiting the expression of the tumor-suppressor LAT2S. These miRNAs are potential novel oncogenes involved in the tumorigenesis of human testicular germ cells by malfunctioning the p53 pathway, rendering tumorigenic growth continual in the presence of wild-type p53 [139,140].

Perspectives on Regulation of RNAi/miRNA Pathway and Roles of miRNAs in Tumorigenesis and Metastasis

Significant achievements have been made in understanding regulation of microRNA pathway as well as functions of miRNAs in tumorigenesis and metastasis in recent years, laying down the foundation for further uncovering regulation of gene expression at posttranscriptional level particularly in response to stress and disease initiation such as tumorigenesis and cancer development. Since chemical biology offers a more dynamic way to monitor the activity of specific pathways compared to traditional forward or reverse genetic approach, its emergence to the RNAi/miRNA pathway although still in its infancy, has lead to further understanding on the regulation mechanisms by identification of a number of small molecule modulators targeting to the specific steps in RNAi/miRNA pathway. Further identification of small molecule identifiers and characterization of targets of these modifiers will be essential to further dissect the regulation mechanism. Processing of some specific miRNAs could be regulated in the cell type or physiological condition dependent manners [59,62], while the underlying mechanism of regulation remains to be elucidated. The specific regulation of some miRNAs, particularly those that have been linked to human diseases, is of importance in terms of mechanism of study and development of new therapeutic strategy. Thus, it would be essential to identify small molecules modifiers that could regulate the processing and activity of specific miRNAs. While technically challenged, the success in identification of miR-21-specific inhibitor could shed light on the feasibility of the chemical biology approach in this regard [141]. Besides potential targeting of the protein(s) involved in the regulation of the processing of selective miRNAs, several small molecules could directly bind to RNAs, including miRNAs [9,119,142]. Given the diversity of RNA secondary structures among miRNA precursor population, it is plausible to speculate that small molecules targeting to specific pri-, pre- or mature miRNA could be identified. Substantial type of miRNAs has been identified in cancer cells with significant deregulation compared to normal adjacent tissues, caused by transcription factors and chromatin modulators as well as RNA binding proteins and their interacting partners. With development and application of high throughput technologies, more cancer-closely-related miRNAs will be identified and further study on metabolic pathways regulated by these cancer-related miRNAs may lead to discovery of unknown pathological mechanisms for tumorigenesis and cancer development. Considering the fact that miRNAs are involved in nearly every cellular process, the identification and characterization of small molecule modulators of RNAi/miRNA pathway will provide novel insights into the fundamentally pathological mechanisms of human diseases particularly cancer. Furthermore, these RNAi modulators, particularly RNAi enhancers, could potentially facilitate the development of RNA interference as a tool for biomedical research and therapeutic interventions.

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Citation: Jiang R, Li Y (2013) Regulation of Mima Pathway and Roles of Micromas in Tumorigenesis and Metastasis. Human Genet Embryol S2: 607. doi:10.4172/2161-0436.S2-007

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