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## **Research Article**

# miRNAs in endothelial cell signaling: The endomiRNAs

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#### ABSTRACT

microRNAs (miRNAs) have a pivotal role during the formation and function of the cardiovascular system. More than 300 miRNAs have been currently found within the mammalian genome, however only few specific miRNAs, named endomiRNAs, showed conseved endothelial cell expression and function. In this review we present an overview of the currently known endomiRNAs, focusing on their genome localization, processing and target gene repression during vasculogenesis and angiogenesis.

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## miRNAs biogenesis and regulation

microRNAs (miRNAs) are highly conserved small non-coding RNAs (~22 nucleotide—nt) that play an important role in the regulation of gene expression at the post-transcriptional level. miRNAs are located within introns and exons of protein coding genes or in intergenic regions. They are transcribed by RNA polymerase II as long primary miRNA transcripts (pri-miRNA) containing one or several hairpin structures with 5'CAP and polyadenylated tails [1]. In the nucleus, the pri-miRNA is cleaved by a microprocessor complex, the RNAse III enzymes Drosha and DGR8 [2]. This cleavage step results in an  $\sim$ 65 nt precursor miRNA (pre-miRNA) which is exported to the cytoplasm in association with Exportin-5 and cleaved by Dicer to 18-24 nt duplex. Finally, this miRNA duplex is loaded into the RNAinduced silencing complex (RISC) together with the Argonaute (Ago) proteins. RISC can bind to the 3'-untranslated region (UTR) of the target mRNA on a partial miRNA-mRNA complementarity to the first 8 nt within the 5' miRNA end called the seed sequence [3]. This binding causes the degradation of the target mRNA and consequent translational inhibition [4]. miRNAs derived from intronic hairpins are called mirtrons and they bypass the cleavage step from Drosha while nuclear transport and cleavage are common to the canonical miRNA biogenesis pathway. The processing of the miRNAs is a tightly regulated mechanism involving regulatory proteins, editing of miRNA transcripts and leading to either elevated or decreased miRNA levels (Fig. 1) [3]. These regulatory proteins can be subdivided into three groups: Drosha binding/associated proteins, Dicer binding proteins and proteins that bind to the terminal loop of the pri- and/or premiRNAs. For example, the DEAD box helicase p68/p72 complex facilitates the processing of a set of miRNAs by binding Drosha [5]. This complex can be activated upon interaction with Smad factors after Transforming Growth Factor  $\beta$  (TGF- $\beta$ ) and Bone Morphogenesis Protein (BMP) stimulus [6]. Similarly, TAR RNA binding protein (TRBP) can interact and stabilize Dicer after phosphorylation mediated by the mitogen-activated protein kinase (MAPK) signaling pathway [7,8]. Therefore, alteration

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03 Fig. 1 – miRNAs processing and regulation. miRNA precursors 107 (pri-miRNA) are processed by Drosha and DGCR8 proteins into 108 smaller RNA hairpins named pre-miRNAs. Protein-protein or 109 RNA binding proteins are able to inhibit (red) or promote 110 (green) pri-in to pre-miRNA processing with the nucleus. 111 After export to the cytoplasm, pre-miRNAs are associated with 112 the endonuclease DICER and other regulatory protein such as 113 TRBP. DICER cleaves the pre-miRNAs into a  $\sim$ 22 nt duplex 114 miRNA which is incorporated in the RNA-inducing-silencing-115 complex (RISC) where the mature miRNA associates with 116 AGO2 to induce translation repression of the target mRNA. 117 This process can be regulated by several growth factor 118 singling pathways through ERK-MAP kinase or SMAD 119 activation. hnRNP=heterogeneous nuclear riboprotein; 120 KSRP = KH-type splicing regulatory protein; SMAD = mothers 121 against decapentaplegic homolog; ADAR = adenosine 122 deamenases acting in RNA. (For interpretation of the 123 references to color in this figure, the reader is referred to the 124 web version of this article.) 125 126

128 of miRNA processing by ERK may result in a pro-growth factor 129 phenotype. Finally, many other RNA binding proteins such as LIN28 [8], hnRNP [9] and KSRP [10] can directly bind to the 130 131 terminal loop of different pre-miRNAs and modulate the cleavage 132 of their mature sequence in different cell types or different stages 133 of developments. The most recent mechanism discovered on 134 miRNA editing and destabilization is the conversion of adenosine to inosine within the pri- or pre-miRNA by the adenosine 135 136 deaminase ADAR [11]. Splice variants of ADAR have been 137 identified in the cardiac tissue suggesting a specific ADAR/ 138 miRNAs regulation in different cellular contexts. However, the 139 expression and function of this protein in the cardiovascular 140 system remain uninvestigated.

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# 143 Targeting miRNAs144

145To probe the in vitro and in vivo function of miRNAs many146approaches have been developed to block the miRNAs activity.

Currently, three strategies are used in miRNA loss-of function studies: genetic knockouts, miRNAs sponge, and antisense oligonucleotides. miRNA knockout allows the generation of animal systems with the whole mount or tissue specific deletion of candidate miRNA genes. The generation of knockouts in Caenor- Q2 habditis elegans and Drosophila as well in mouse model has been extensively used to unravel the function of miRNAs in early embryonic development [12]. Recently, a genome-wide knockout resource covering 476 mouse miRNA genes was described and embryonic stem cells repertories have been made available to improve the studies of the miRNAs in vivo [13]. So far, the majority of the ~25 miRNAs mouse knockouts do not exhibit severe defects during embryonic development. Indeed, many miRNA families exist as duplicates or have identical seed regions, thus there remains the question of functional redundancy. In addition, genome deletion of a single miRNA encoded within a gene cluster is particularly difficult without affecting the expression of the other miRNAs. Alternative methods to perform gene knockout may overcome family complexity and redundancy of miRNA gene function. Recently improvements in artificial transcription activator-like effector nucleases (TALENs) provide a promising and powerful new approach for precise genome targeting of miRNAs genes [14].

miRNA sponges are transcripts with repeated miRNA antisense sequences that can sequester miRNAs from endogenous mRNA targets. Specific design of sponges can bind and sequester all miRNAs seed family members or a single miRNA within a cluster offering additional advantages in the analysis of complex miR-NAs activity. Sponge has been proven to induce transient, long term and tissue specific miRNAs inhibition in several animal models [15]; however, the success of their activity is strongly dependent on the endogenous miRNA expression levels. Chemically modified antisense nucleotides are the most common approach to perform loss of miRNA function studies. A variety of commercial chemical modifications such as the 2'-O-Methil (2'-O-ME), 2'-O-methoxyethyl (2'-MOE) or locked nucleic acid (LNA) and morpholino oligonucleotides are now available. These modifications confer nuclease resistance and high binding affinity to target miRNAs [16]. These oligos named antimiR can target miRNAs at different stages of their biogenesis or block miRNAs activity by binding the mature miRNA sequence or by targeting the miRNA binding site within the target mRNA [17]. Several studies evaluated the efficiency of different modified antimiR mediated inhibitions in both vitro and in vivo [16]. This strategy has been used to block the activity of individual miRNAs while new strategies still have to be developed to block miRNA families. A recent publication has shown the efficiency of eight nucleotides LNA modified phophotiorate nucleotide, named tiny LNA, complementary to the seed region of the target miRNAs. Tiny LNA inhibits individual miRNA and/or miRNA families in culture cells and several tissues when delivered to the adult mouse [18]. AntimiR or tiny LNA oligonucleotides' disadvantages are related to their transitory and partial inhibitory effect in addition to potential off-target effects.

## **Target prediction tools**

Open-access bioinformatic databases have been developed to facilitate the analysis of miRNAs and their target prediction [19].

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207 Prediction of miRNA-mRNA interactions is a challenging task, 208 due to the short length of miRNAs, the requirement of only 209 partial homology for binding, the redundancy among members of 210 an miRNA family, and the existence of multiple putative miRNA 211 recognition sites [20]. The majority of the computational target 212 prediction programs are based on several features, such as 213 complementarity between the 5'-seed of the miRNA and the 214 3'-UTR of the target mRNA, thermodynamic stability of the 215 miRNA-mRNA duplex, conservation among species, and the 216 presence of several miRNA target sites. Several different in silico 217 target prediction programs exist to identify miRNA-target pre-218 diction. Such prediction programs must be used carefully since 219 (i) the genome is not fully sequenced, (ii) only a limited number 220 of miRNA targets have been experimentally validated, and 221 (iii) there is low conservation among species regarding mature 222 microRNAs sequence as well as the target prediction. Experi-223 mental high-throughput studies (such as chromatin marks and 224 poly(A)-site mapping followed by RNA-Seq) are needed to allow 225 for generation of precise prediction algorithms [21,22].

## miRNA-target validation tools

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230 Efficient experimental strategies are needed to validate computa-231 tionally predicted miRNA target genes. To validate predicted 232 miRNA-mRNA interactions, several experimental approaches have 233 been used, such as biochemical methods (luciferase assays, qRT-234 PCR, western blotting, and RNA seq), "omics" approaches (SILAC, 235 LAMP), and RISCome analysis (quantification of mRNAs in the RNA-236 induced sequencing complex; RISC RNA sequencing, RIP-chip) [23]. 237 Experiments performed in cultured cell lines are usually useful to 238 validate microRNA effects on target mRNA. Recently, animal models 239 have also been used to validate target mRNA in vivo. Novel 240 techniques have been implemented in the zebrafish model such 241 as fluorescent miRNA sensor and morpholino (MO) target protector. 242 In the first case, validation of the target prediction is tested using a 243 reporter assay based on monitoring GFP/mCherry fluorescence in 244 zebrafish embryos microinjected with mRNA encoding the fluor-245 escent reporter fused to the 3'-UTR of the target gene, in the 246 presence or absence of specific miRNA duplex. In this assay, a 247 decrease in fluorescence in the presence of miRNA duplex indicates 248 miRNA-mediated repression and then the confirmation of the 249 target in vivo. In the second case, MOs are designed to be 250 complementary to specific miRNA-binding sites in target mRNAs. 251 In this way MOs have been shown to efficiently protect the target 252 mRNA from translational inhibition or degradation. To study the 253 regulation of a particular target, it is important to establish first that 254 the 3'-UTR is regulated by a particular miRNA, miRNAs may speed-255 up degradation or slow-down translation of their targets, but 256 repression caused by either mechanisms can be assessed by 257 measuring protein output of a reporter. The efficiency of "target 258 protectors" to block the interaction of miRNAs with a particular 259 binding site in a target mRNA can also be investigated by coinjections with reporter mRNAs containing a GFP/mCherry coding 260 261 sequence and a 3'-UTR region with the miRNA-binding site. 262 Subsequently, a tested target protector can be used to study 263 whether protection of a specific target mRNA from silencing has 264 any biological effect in a tissue-specific manner (by generation of a 265 cell autonomous miRNA sensor). Recently, a proteomics approach 266 to validate predicted miRNA targets in C. elegans by using

quantitative targeted proteomics via selected reaction monitoring (RIP-chip-SRM) has been reported [21]. This technique can be applied to validate candidate lists generated by computational methods or in large-scale experiments, and the described strategy should be readily adaptable to other organisms.

# miRNAs function in endothelial cells: the "endomiRNAs"

The mammalian genome encodes ~300 highly conserved miRNAs  $(\sim 1-2\%$  of the genes) able to regulate, at least by bioinformatics prediction, over 60% of protein coding genes [24]. In the past five years this magnitude of gene regulation by miRNAs, has been revised based upon numerous evidences : (1) in animal system miRNAs are usually expressed in a non-correlated manner with their predicted mRNA targets [25]; (2) majority of the miRNAs produce subtle proteins reduction (<2 fold change) [26]; (3) many miRNAs can be deleted in vivo without causing evident phenotypes [27]. If so, what is the real inhibitory potential of miRNAs? Increasing evidence suggested that miRNAs confer robustness to complex signaling pathways, functioning as "buffers" to gene fluctuations. They can reinforce transcriptional programs and attenuate aberrant transcripts thereby conferring accuracy and uniformity to developmental transitions, cell fate switches and stress-responses [28]. The remodeling of the vascular system is a sophisticated result of the balance between stimulators and inhibitors pathway. These pathways are fed by a web of vascular growth factors and downstream proteins signaling tightly controlled by feedback loops and redundant components. For these properties the cardiovascular system is particularly sensitive to the regulation of miRNAs. In vitro and in vivo studies have shown that Dicer has an important function in angiogenesis. Dicer mutant mice embryos and yolk sack manifest angiogenic defects [29]. Accordingly, mice carrying an endothelial tissue specific deletion of Dicer shows defects in postnatal angiogenic response to a variety of stimuli, including exogenous VEGF, tumors, limb ischemia and wound healing [30]. Finally, genetic silencing of Drosha expression in endothelial cells also results in a reduction in capillary sprouting and tube formation [31]. In this review, we will summarize the state-of-art on the major conserved families of miRNAs expressed in endothelial cells, endomiRNAs miR-126 and themiR-17-92, miR-23-27-24 and miR-222-221 clusters (Fig. 2).

## miR-126

In 2008, three independent groups analyzed and reported defects in miR-126 deficient vascular cells, varying from zebrafish to human and mice [32,33]. miR-126 is the most important and possibly the only endothelial-specific miRNA in vertebrates. It is expressed in the vascular tree of vertebrate throughout development. In addition, miR-126 has been found as one of the most enriched microRNAs in cultured human endothelial cells and in endothelial cells derived from embryonic stem cells or isolated from mouse embryos. miR-126 has been clearly shown to be a master miRNA in vascular function working as a key positive regulator of angiogenic signaling in vitro and in vivo [32]. miR-126 is encoded in an intron of EGF-like domain 7 (Egfl7) gene. Egfl7 encodes a secreted matrix component that is produced by

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339 Fig. 2 - Schematic representation of the endomiRNAs and 340 their targets. Four different microRNAs families have been 341 implicated in controlling endothelial cell behavior and 342 signaling. Endothelial microRNAs-endomiRNAs-include miR-343 126 and miR-23-27-24, miR-17-92 and miR-222-221 clusters. 344 These families of microRNAs have been found to be expressed 345 in endothelial cells (or cells that interact with the 346 endothelium) and thus regulate endothelial cell responses. 347 See text for detailed information. 348

351 angiogenic stimuli and induces endothelial migration [34]. Since 352 miR-126 is located within an intron of the Egfl7 gene, questions 353 arise whether the vascular abnormalities seen in miR-126 studies 354 are due to the loss of miR-126 or Egfl7 activity. Comparison 355 between miR-126 and Egfl7 knock out models indicates that the 356 severe vascular defects are only observable in animal deficient in 357 miR-126, highlighting the requirement of miR-126 for the 358 developing and postnatal vasculature. Remarkably, deletion of 359 miR-126 in mice results in vascular developmental defects such 360 as delayed angiogenic sprouting, widespread hemorrhaging, and 361 partial embryonic lethality [33]. In addition, miR-126 mutant 362 mice that successfully complete embryogenesis display dimin-363 ished angiogenesis and increased mortality after coronary liga-364 tion, a model for myocardial infarction. Also endothelial cells deficient in miR-126 fail to respond to angiogenic factors, 365 366 including VEGF, epidermal growth factor (EGF), and bFGF. In 367 zebrafish, knockdown of miR-126 by MO injections induced 368 collapsed blood vessels and cranial hemorrhages in the develop-369 ing organism suggesting its primary role for maintaining vascular 370 structure during development [32]. Mechanistically, two direct 371 targets of miR-126 that seem to explain these angiogenic defects 372 are Sprouty-related EVH1 domain-containing protein 1 (Spred1) 373 and a regulatory subunit of PI3K, PIK3R2 (also known as  $p85\beta$ ) 374 [33]. Because Spred1 and PIK3R2 are negative regulators of 375 cellular signaling cascades, affecting the MAPK and PI3K signal-376 ing pathways, respectively, miR-126 promotes VEGF and other 377 growth factors signaling by targeting multiple signaling pathways, 378 miR-126 may fine-tune angiogenic responses. Spred1, which is a 379 negative regulator of cell survival, was validated as a direct miR-380 126 target being expressed in miR-126 deficient zebrafish. Indeed, 381 downregulation of miR-126 in endothelial progenitor cells from 382 diabetes human patients impairs their functional properties, via 383 target gene Spred-1 [35]. Recently, VEGF-A has been proposed as 384 another important target of miR-126 [36]. More recent studies on 385 zebrafish embryos suggested that miR-126 expression is under 386 control of the mechano-sensitive zinc finger transcription factor klf2a. In this model, pulsatile flow induces expression of miR-126 within the endothelia cells to modulate VEGF signaling during the remodeling of the aortic arches [37]. Many endothelial miRNAs can also be involved in vascular inflammation particularly in leukocyte activation and their infiltration into the vascular wall, miR-126 is a good example. Indeed, a recent study provides first evidence that miRNAs control vascular inflammation since miR-126 inhibits the expression of vascular cell adhesion molecule 1 (VCAM-1), which mediates leukocyte adherence to endothelial cells. Thus, decreasing miR-126 in endothelial cells it is possible to increase TNF $\alpha$ -stimulated VCAM-1 expression and enhances leukocyte adherence to endothelial cells. Using the embryonic stem differentiation system to model primitive erythropoiesis and miR-126 null embryos, it has been found that miR-126 regulates the termination of EryP-CFC development in vivo by targeting VCAM-1 [38].

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#### miR-17-92 cluster

The miR-17-92 cluster is a polycistronic miRNA gene and encodes for six mature miRNAs namely miR-17, miR-18a, miR-19a, miR-20a, miR-19b and miR-92a. All the members of this cluster are highly expressed in endothelial cells regulating vascular integrity and angiogenesis [39]. Interestingly recent evidence showed that members of the miR-17-92 family are differentially expressed during endothelial cells differentiation of pluripotent stem cells. While miR-17. miR18a and miR-19a were increased upon induction of endothelial differentiation, miR-92a was decreased. Even so, the inhibition of each of these miRNAs did not affect endothelial cell differentiation similarly to miR-126 activity which is required for endothelial cell function but not for endothelial differentiation [40]. Numerous studies focus on the endothelial cell function of miR-17-92 cluster that revealed complex and controversial results. The entire cluster is highly up-regulated in multiple human tumor cell types, however only miR-18a and miR-19a have a pro-angiogenic function during tumor angiogenesis. The overexpression of these miRNAs in tumor cell, repressed the secreted anti-angiogenic factors thrombospondin (TSP-1) and connective tissue growth factor (CTGF), inducing an increase of neovascularization in a paracrine fashion [41]. Interestingly, these miRNAs showed an anti-angiogenic function when over expressed directly in endothelial cells. miR-17a, miR18a, miR19a and miR-20b inhibited endothelial cells sprouting when ectopically expressed, while individual miRNA knockdown promotes angiogenesis. In these experiments the protein kinase Jak1 was identified as the major target mRNA responsible for the pro-angiogenic effect [25]. Accordingly, the over expression of another member of the cluster, miR-92a inhibits vascular network formation in matrigel assay via targeting integrin  $\alpha$  5 (ITGA5) and indirectly suppressing eNOS production [42]. The anti- or pro-angiogenic function of the miR-17-92 cluster seems strictly related to the cellular context. Interesting, the miR-17-92 members are also differentially regulated in endothelial cells exposed to hemodynamic force [43]. miR-92a is downregulated in endothelial cells exposed to laminar flow while its mature sequence is upregulated by oscillatory flow consistently with its atheroprotective function in vivo [43]. The differential expression and therefore function of the single miRNAs derived from miR-17-92 cluster are mostly due to post-transcriptional regulation of the primiRNA [44]. Consequently, physiological and pathological conditions as well as tissue dependent miRNA editing events can modulate the expression of a single miRNA belonging to the miR-17-92 cluster.

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447 Further investigations need to be performed to elucidate this aspect. 448 Mouse targeted deletion of the miR-17-92 cluster reveals essential 449 functions of these miRNAs during embryonic development. The 450 functional analysis of the miR-17-92 cluster is complicated by the 451 existence of two paralog clusters miR-106a-363 and miR-106b-25 452 generated by duplication event during evolution [45]. Mice lacking 453 miR-17-92 or both miR-17-92 and miR106b-25, died shortly after 454 birth showing lung hypoplasia and impaired B cell development due 455 to an increased level of the pro-apoptotic Bim protein [45]. The 456 characterization of the vascular development in these mice is 457 currently missing. In addition, the endothelial specific deletion of 458 miR-17-92 cluster will be necessary to elucidate the function of 459 miR17-92 cluster during the embryonic vascular morphogenesis.

## miR-23-27-24 cluster

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464 The miR-23/27/24 cluster exists in the mammalian genome as duplicate gene located in different genomic locations: miR-23b, 465 466 miR-27b and miR-24-1 are located within the intron region at 467 the human chromosome 9q22.32 while the intergenic cluster 468 miR-23a, miR-27a and miR-24-2 gene are localized at the 469 chromosome 19q13.13. The mature sequence of the respective 470 miR-23a/b, 27a/b and miR-24 1 and 2 are identical and highly 471 conserved among vertebrates. The dissection of miR-23-27-24 472 gene expression and function is intrinsically complicated by their 473 genome structure. For example, miR-24 pri-miRNA and the 474 mature sequence is up-regulated after BMP2 stimulation in 475 mesenchymal cells, while miR-27b and 23b are unaffected by 476 this treatment suggesting that even if they are in close proximity 477 within the genome the pri-miRNA of miR-24 and miR-23-27 are 478 transcribed independently [45]. All the members of this cluster 479 are highly enriched in vascularized organs and endothelial cells. 480 Silencing of miR-23-27 inhibits the angiogenesis in response to 481 VEGF by impairing the activation of MAP and PI3K-AKT kinase 482 signaling pathway. Similarly, silencing of miR-23-27 suppresses 483 choroid revascularization after laser injury [46,47]. miR-23 and 484 miR-27 sequences possess different seed regions, therefore they 485 can target diverse mRNA targets. However, in endothelia cells 486 both miR-23-27 bind the Sprouty2 and sema6A 3'-UTR reinfor-487 cing the reciprocal inhibitory activity. The repressions of these 488 two target genes by miR-23-27 serve to maintain an active RAC/ RAF/ERK pathway after VEGF stimulation [46,47]. Intriguing, 489 490 miR-24 expression is highly induced in endothelial cells after 491 stress condition such as oxidative stress. Indeed, miR-24 acti-492 vates pro-apoptotic signaling within the vasculature of the 493 myocardium after myocardial infarction and in vivo treatment 494 with antimiR against miR-24 improves vascularization and pre-495 serves cardiac function after myocardial infarction. GATA2 and 496 p21 activated kinase Pak1 seem to be the major gene targets 497 assigned to the miR-24/pro-apoptotic function during vascular 498 remodeling [48]. Currently there are no mouse knockout models 499 for the miR-23-27-24 cluster and therefore no data that can 500 suggest their function during embryonic vascular development. 501 However, the downregulation of miR-27 in zebrafish embryos 502 induced venous remodeling and angiogenesis of intersegmental 503 vessel (ISV) defects. The miR-27 loss of function phenotype can 504 be rescued by the repression of either Sprouty2 or Dll4 genes 505 proposing these two genes as major miR-27 targets during 506 zebrafish vascular development [49].

### miR-222-221 cluster

miR-222 and 221 are highly conserved miRNAs transcribed from a common polyadenylated pri-miRNA located on the human chromosome X. Both miR-221 and miR-222 have a proproliferative effect on cancer cells [50]. miR-222 and miR-221 are also highly expressed in endothelial cells after growth factor stimulation or in the quiescent state [26]. While both miRNAs share the same seed region, numerous studies identified different target genes controlled by miR-221 and miR-222. For example, compared with miR-222, miR-221 is not biologically relevant for inflammation mediated by vascular growth factors [51] where during zebrafish embryonic development miR-222 is dispensable for vascular remodeling [52]. Similar observations were reported in a model of liver tumorigenesis, in which miR-221, but not miR-222 was able to accelerate tumor growth in mice [53]. The regulatory activities of this miRNA cluster seem to be cell type dependent. The inhibition of miR-222-221 decreases neointimal lesion formation but increases re-endotheliazation during hyperplasia following vascular injury. Indeed, miR-222-221 are both highly expressed in endothelial and vascular small muscle cells (VSMC). In VSMS this cluster have a pro-proliferative and promigratory effect by targeting two cell cycle dependent inhibitors p27 (cdkn1b/Kip1) and p57 (kip2) [54]. In contrast exogenous miR-222-221 show anti-proliferative activity in human venous and lymphatic endothelial cells by targeting numerous target mRNAs depending on the experimental setting. These targets include ETS1 transcription factor [55], the stem cells factor receptor cKit [56] and the transcriptional repressor ZEB2 [57]. However, the molecular mechanism responsible for miR-222-221 mediated opposite cellular effects is currently unclear. Importantly, miR-221 expression varies significantly in response to both serum and VEGF treatment [25] generating discrepancies between different experimental conditions. The function of the miR-222-221 cluster during the mouse embryonic development has not been investigated yet. Recent studies using the zebrafish model showed that miR-221 is a NOTCH dependent miRNA required for endothelial tip cells proliferation and migration. miR-221 promotes endothelial cells sprouting by targeting two distinct target genes, PIK3R1and p27. miR-221/p27 target regulation is required to coordinate proliferation, while tuning PI3K output by miR-221/PIK3R1 repression is fundamental for tip cell migration upon VEGFC-Flt4 activation [52].

## Conclusion

This review summarizes current research progress and knowledge on the roles of miRNAs in regulating endothelial cell function and signaling. The discovery of miRNAs as regulators of vascular specific signaling pathways has created new options for the design of therapeutic agents that could modify gene expression in vascular-associated disease. Recently, it was discovered that extracellular miRNAs circulate in the bloodstream and that such circulating miRNAs are remarkably stable [58]. This has raised the possibility that miRNAs may be probed in the circulation and can serve as novel diagnostic markers for vascular-related diseases such as myocardial infarction, heart failure, atherosclerosis, hypertension, and type 2 diabetes. These

discoveries are expected to present opportunities for clinical diagnostic and therapeutic approaches in miRNA-based vascular diseases.

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