Emerging Role of microRNAs in Stroke Protection Elicited by Remote Postconditioning

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Remote ischemic conditioning (RIC) represents an innovative and attractive neuroprotective approach in brain ischemia. The purpose of this intervention is to activate endogenous tolerance mechanisms by inflicting a subliminal ischemia injury to the limbs, or to another "remote" region, leading to a protective systemic response against ischemic brain injury. Among the multiple candidates that have been proposed as putative mediators of the protective effect generated by the subthreshold peripheral ischemic insult, it has been hypothesized that microRNAs may play a vital role in the infarct-sparing effect of RIC. The effect of miRNAs can be exploited at different levels: (1) as transducers of protective messages to the brain or (2) as effectors of brain protection. The purpose of the present review is to summarize the most recent evidence supporting the involvement of microRNAs in brain protection elicited by remote conditioning, highlighting potential and pitfalls in their exploitation as diagnostic and therapeutic tools. The understanding of these processes could help provide light on the molecular pathways involved in brain protection for the future development of miRNA-based theranostic agents in stroke.

Keywords: microRNA, stroke, remote conditioning, tolerance, preconditioning

INTRODUCTION

Ischemic conditioning is a neuroprotective approach able to make the brain more resistant to an ischemic insult through the exposure to a subthreshold stimulus. This method provides neuroprotection when the conditioning stimulus is administered either before or after the detrimental ischemia, i.e., preconditioning or postconditioning.

Indeed, ischemic preconditioning is an endogenous defensive process triggered by a subclinical ischemic event that increases tissue resilience or, in other words, organ resistance to a subsequent, typically dangerous, ischemia episode. Non-ischemic conditioning cues can also promote neuroprotection against an ischemic insult, a phenomenon known as “cross-protection”(1). Surprisingly, when the subliminal boost is delivered after the ischemia insult, the neuroprotection achieved, referred to as postconditioning, is comparable to that shown in ischemic preconditioning models. Remarkably, combining preconditioning and postconditioning does not result in greater protection than either treatment alone (2).

We define preconditioning, perconditioning, and postconditioning from a strictly temporal standpoint, depending on whether the conditioning stimulus is delivered before, during, or after the detrimental ischemia (Figure 1).
It is now well recognized that stressing preconditioning or postconditioning stimuli elicit multiple endogenous defensive mechanisms in the brain, resulting in a latent protective phenotype. When the lethal ischemic insult is delivered inside this inactive protective phenotype, a partitioned set of reactions are triggered that are strikingly different from the phenotype of the unprimed or non-preconditioned brain, resulting in the so-called ischemia-tolerant phenotype (3).

Surprisingly enough, in the last years we and others produced data supporting the idea that preconditioning and postconditioning exert their effects also when applied to an anatomical site distant from the brain (4–8). In fact, remote ischemic conditioning (RIC) represents an innovative and attractive protective approach in brain ischemia. This method is designed to elicit the initiation of endogenous tolerance processes by providing a not-dangerous ischemic event in a distant tissue, i.e., arms or limbs, leading to a protective systemic response against stroke. Several studies have examined the effectiveness of RIC in reducing the effects of ischemic brain injury, as well as the potential pathophysiological pathways involved (4–8).

Among the multiple candidates that have been proposed as putative mediators of the protective effect generated by the subthreshold peripheral ischemic insult, it has been hypothesized that also in the case of remote conditioning, as it occurs in the case of direct conditioning (9–13), miRNAs may play a vital role in the infarct sparing effect (5, 7).

The effect of miRNAs during RIC can be exploited at different levels: (a) as transducers of protective messages to the brain or (b) as effectors of brain protection.

This review will summarize the most recent evidence supporting the involvement of microRNAs in the protection of the brain caused by remote conditioning, highlighting the potential and pitfalls in their exploitation as diagnostic and therapeutic tools.

**microRNAs AS THERAPEUTIC AND DIAGNOSTIC TOOLS**

For more than a century, the central view that has dominated molecular biology has been that protein production is mediated by the DNA–RNA–protein axis, which only involves transcription and translation mechanisms to enable decoding of the human genome for production of specific proteins. However, over the recent years, the discovery of non-coding RNAs has radically revolutionized this theory, defining new mechanisms involved in the modulation of protein expression. The human genome project discovered that protein-coding sequences make up only 1.5% of the genome, while introns, regulatory DNA sequences, interspersed elements, and non-coding RNA (ncRNA) molecules make up the remaining 98.5% (14). Without a doubt, the majority of mammalian genomes are translated into ncRNAs, many of which are spliced or processed into smaller products. Short ncRNAs and long ncRNAs are the two types of ncRNAs that have been identified and characterized so far. MicroRNA (miRNA), small interference RNA (siRNA), small nuclear RNA (snRNA), small nucleolar RNA (snoRNA), PIWI-interacting RNA (pi-RNA), transfer RNA (tRNA), circular miRNA (circRNA), ribosomal RNA (rRNA), and other uncharacterized tiny particles are among the short ncRNAs. As of now, miRNAs are characterized as RNA molecules of 18–24 nucleotides in length, transcribed from genes or from intronic regions of other genes, and may promote mRNA degradation or restrict protein translation to control gene expression. More than 60% of mammalian mRNAs are thought to be targeted by at least one miRNA, according to computational predictions (9, 15–19).

Later advances in the topic of miRNAs and their relationship to human diseases have revealed that miRNAs are useful biomarkers and possible disease-modifying agents (20). MiRNA expression profiles exhibit significant changes in response to disease, suggesting that miRNAs are important controllers of disease-related pathways (20).

MiRNAs are considered “downregulators” of gene expression by means of two primary mechanisms: (1) cleavage of mRNA and (2) repression of mRNA translation. MiRNAs interact with mRNA targets through partial sequence complementation, regularly inside the 3’ untranslated region of the mRNA target. In specific, nucleotides 2–7 of the miRNA (beginning from the 5’ end), named the “seed” sequence, are imperative for target binding. The extent of base pairing between miRNA and its target mRNA is currently thought to determine whether the mRNA is degraded or translationally repressed (21, 22).

miRNAs are differentially expressed among tissues, between male and female, during different developmental stages or in response to specific physiological or pathophysiological conditions (17, 23–25). Examples include miR-122, which is preferentially expressed in the liver (24), miR-133a and miR-133b, which are highly enriched in muscle (25), and the miR-302 family members which are specifically expressed in stem cells (26).

**microRNAs as Therapeutic Tools**

The development of precise and fast assays for miRNA target identification has played a considerable impact in the study of miRNA roles and in the characterization of the biologic activities in which they are implicated. Since the discovery of the potentialities of miRNAs as therapeutic agents, several effective algorithms have been developed for the prediction of miRNA targets (27–32).

There are various extending endeavors to create therapeutics that straightforwardly can target miRNAs, and agreeing to the sort of miRNA, its expression, and its work, diverse approaches are utilized to overexpress or restrain miRNAs (33).

A miRNA mimic is applied to replace the miRNA concentration, eventually reduced in the progression of a pathologic state. This approach was created in 2007 as gain-of-function instrument for particular miRNAs and comprises engineered double-stranded RNAs that are specifically recognized by RNA-induced silencing complex (RISC) (34). These miRNA-like RNA fragments have a 5’ end that gets a sequence with partial complementarity to the 3’UTR of the target genes, in this way imitating the miRNA working mode. This innovation has been further developed by pharmaceutical companies, and a library of miRNA mimics is accessible for
all human miRNAs found until presently. In addition, these particles experience modifications in backbone and ribose to advance steadiness in vivo and to overcome impediments related to pharmacodynamics. Over the years, other important changes have been made utilizing adeno-associated infection (AAV) vectors and tissue-specific promoters to improve the tissue specificity impact of miRNA mimics (35).

On the other hand, blocking a miRNA may represent a therapeutic option in some pathological conditions characterized by high levels of expression of those miRNAs involved in deleterious pathways. Anti-miRNAs are nucleotide sequences able to bind to the mature miRNA guide strand, causing its inhibition (20). Several methods are used to add modifications with the intent to generate stable and deliverable miRNA inhibitors, creating a class of antagonists known as antagomirs. Among these chemical modifications are included cholesterol conjugation and the use of locked nucleic acid (LNA) (36). LNA modification increases the stability and nuclease resistance of antisense oligonucleotides and, more importantly, improves the efficiency of hybridization to single-stranded RNA. Chemically modifying the miRNA using either LNAs or the 2′-O-methyl group (OME) can increase stability in vivo and is the basis for many antagonir therapies. Antagomirs are the most frequently used approach for therapeutic use of miRNA therapy (37).

An evolution of anti-miRNAs is represented by miRNA sponges, which are able to inhibit multiple miRNAs simultaneously (38). These molecules can contain a seed sequence for an entire miRNA family, however, they could be also used to target multiple miRNAs.

Finally, the usage of miR-Masks is an emerging strategy for altering miRNA activity (39). These masks are single-stranded 2′-O-methyl-modified antisense oligonucleotides that recognize the miRNA target site on the mRNA 3′UTR and hide it.

miRNAs as Diagnostic Tools
Beside their expression within tissues and organs, miRNAs are also present outside the cells (40). Several release mechanisms have been hypothesized including release through microvesicles, resulting from outward budding and plasma membrane separation (41). In particular, some vesicles called exosomes, typified by a specific process of biogenesis, are strongly involved in intercellular communication processes and are characterized by the presence of massive amounts of miRNAs inside them (42). Other miRNA transport systems in biological fluids include high-density lipoproteins (HDL) and apoptotic bodies, generated during the programmed cell death process.

Whatever the origins of miRNA, their presence in the blood and the ability to quantify their levels non-invasively cleared the path for the development of peripheral biomarkers for the diagnosis and prognosis of disorders such as brain ischemia. Undoubtedly, miRNA expression levels within the blood are reproducible and demonstrative of different pathologies (43). The interest in peripheral miRNA and their potential application as a biomarker for rapid diagnosis and prognosis are particularly relevant for ischemic patients (44, 45). In fact, several patient-based studies have already reported the occurrence of significant variations in the circulatory expression of miRNAs during cerebral ischemia, correlatable to the entity of the damage (46).

TRANSLATIONAL RELEVANCE OF miRNAs IN REMOTE CONDITIONING NEUROPROTECTION
Over the years, numerous studies demonstrated that stroke may trigger a re-arrangement of the miRNA profile within the cerebral tissue of animals exposed to brain ischemia (47, 48). The first miRNA expression profiling study in cerebral ischemia was performed in 2008, on the entire brains of rats subjected to
middle cerebral artery occlusion (MCAO) with reperfusion for 24 or 48 h (49). In this study, it has been shown that 106 transcripts were altered 24 h after stroke induction. However, the number of altered miRNAs was reduced to 82 at 48 h, thus indicating that miRNAs were differentially expressed after brain ischemia in a time-dependent manner. In fact, in the 48-h samples, only rno-miR-99a, −181 (a, b, and c), −195, −328, −379, and −539 were found to be exclusively expressed, whereas, in the 24-h samples, 32 miRNAs (rno-miR-16−17, −20a, −21, −24−25, −30a−3p, −34a, −92, −124a, −130a, −132, −134, −151a, −210, −215, −324−3p, −322, −329, −342, −361, −374, −382, −383, −422b, −433, −451, −497, −505, −664, let-7d, and let-7f) were found to be exclusively expressed. Notably, rno-miR-206, −214, −223, −290, −292−5p, −298, −327, and −494 were highly upregulated during both ischemia/reperfusion time intervals (49).

These pioneering data were partially confirmed later on in another miRNA profile study carried out in spontaneously hypertensive rats exposed to focal ischemia and reperfusion of different duration (50). In the brain samples obtained from these ischemic animals, it has been found that among the 238 miRNAs examined, 24 miRNAs were upregulated while 22 miRNAs were downregulated at one or more reperfusion time intervals. These data were obtained in the ipsilateral cortex of the ischemic core. Little changes in miRNA expression were detected in the contralateral cortex and in the peri-ischemic area adjacent to the ischemic core, compromised by the ischemic event but not completely damaged.

In this scenario, particular attention was focused to the possibility to restore ionic homeostasis in the penumbra region by using miRNAs to modulate ionic channels and transporters, since the tight correlation between brain damage due to brain ischemia and the disruption of ionic homeostasis is well known (8, 51−53). The plasma membrane sodium/calcium exchanger (NCX1), whose expression is controlled by miR-103 and whose activation ameliorates ischemic damage (51−54), appeared as one of the most promising candidates. Indeed, the capability of AntimiR-103 of inducing a brain-conditioning phenomenon in a rat model of transient brain ischemia has been evidenced (55). The mechanism of action of this LNA anti-miRNA consisted in blocking the detrimental increase of miR-103-1 responsible for the downregulation of NCX1, whose expression is necessary to counteract sodium and calcium imbalance occurring during stroke progression.

Interestingly, more recently, another miRNA, miR-223-5p, emerged as a possible modulator of the K+−dependent Na+/Ca2+ exchanger, NCKX2, a new promising stroke neuroprotective target. In fact, intracerebroventricular infusion of anti-miR-223-5p prevented NCKX2 downregulation occurring after ischemia in rats, thus promoting neuroprotection. Therefore, blocking miR-223-5p by anti-miRNA is a reasonable strategy to reduce the neurodetrimental effect induced by NCKX2 downregulation occurring during brain ischemia (56).

The translatability of miRNAs as biomarkers derives from their physical characteristics. Indeed, as anticipated above, miRNAs are found in a remarkably stable state in human plasma or serum and could be used as biomarkers for a variety of disorders. Chen et al. demonstrated that plasma miRNAs are resistant to RNases as well as other harsh settings such as low/high pHe, long-term storage, boiling, and repeated freezing/thawing cycles (57). The levels of miRNA expression in blood have been proven to be repeatable and predictive of illness condition. Although the mechanism of miRNA release into the circulation is unknown, their presence in the bloodstream and relationship with a variety of pathophysiological conditions is now widely established (58) and miRNAs produced by injured or circulating cells are thought to cause enhanced miRNA expression in peripheral biofluids (59).

Many studies demonstrated that this stability is related to the different carriers that mediate miRNA transport, including microvesicles, exosomes, apoptotic bodies, AGO protein complexes to form ribonucleoprotein, and HDL (60, 61). Exosomes, according to current thinking, can regulate the bioactivities of recipient cells by transporting lipids, proteins, and nucleic acids like miRNAs while circulating in the extracellular space, and several studies have shown that exosomes play important roles in immune response, tumor progression, and neurodegenerative disorders.

Taking into account all these considerations and in the attempt to find peripheral markers for stroke, several studies have been conducted to examine expression changes in circulating miRNAs following brain ischemia, at preclinical and clinical levels (10).

Contrasting results have been produced when the expression profiles of miRNAs in the brain were compared to those measured in the blood. In fact, some miRNAs such as miR-290 and miR-494 showed change in their expression in the same direction (upregulation or downregulation) in both tissues at 24 and 48 h, whereas expression levels of some other miRNAs, like miR-150, −195, and −320, exhibited an inverse trend (49). Once again, it should be underlined that brain tissue after ischemic stroke changes over the time, and the peri-ischemic region is included in the ischemic core in few hours. miRNA expression is surely influenced by the stage of the disease and the distance from the ischemic core.

Distinct miRNA patterns indicative of the stroke outcome have been found in blood samples of chronic stroke patients within 6–18 months from the stroke onset (62). Interestingly, several miRNAs have shown changes during disease progression. Notably, the number of miRNAs downregulated in all good-outcome stroke samples was usually higher than that of upregulated ones. However, miRNA expression profiles exhibited differential fold change values among the different stroke subtypes; for instance, patients affected by small-artery (SA) stroke show a distinctly different pattern from that
of patients affected by large-artery (LA) stroke. In specific, among the highly upregulated miRNAs measured in samples derived from patients affected by SA stroke, seven miRNAs, miR-130b, −29b, −301a, −339-5p, −532-5p, −634, and 886-5p, changed more than two-fold (62).

Recently, miR-107, miR-128b, and miR-153, three brain-enriched miRNAs, were identified within 24 h from hospital admission in plasma samples of patients affected by brain ischemia, demonstrating that their levels were upregulated after stroke onset and positively correlated with the severity of cerebral ischemic injury (63). Interestingly, no correlation was found between age and smoke with the levels of miRNAs, thus suggesting that this upregulation was mediated only by ischemic insult. In parallel, a further study examined the levels of two atherosclerosis-related miRNAs, miR-185, and −146a, in plasma of ischemic stroke patients in the acute phase, 1–5 days, or subacute phase, 6–30 days (10, 37). MiR-185 was found to be downregulated in both the acute and subacute phases, whereas miR-146a was found to be downregulated in the acute phase but upregulated in the subacute phase. Finally, eight miRNAs were found to be differentially expressed in blood collected 28 h after stroke onset (64). In fact, miR-122, miR-148a, let-7i, miR-19a, miR-320d, and miR-4429 were reduced, while miR-363 and miR-487b were upregulated in the bloodstream of acute ischemic patients. These miRNAs were predicted to be regulators of a number of genes and pathways linked to brain ischemia and involved in immune activation, leukocyte extravasation, and thrombus formation.

In the light of these premises and considering the versatility of miRNAs, the identification of pivotal miRNAs may represent a crucial step in defining new putative theranostic tools in stroke. Therefore, the research in the field has been attracted to the possibility of selecting peculiar miRNAs involved in endogenous neuroprotective phenomena including preconditioning and postconditioning.

The idea that ischemic preconditioning (IP) reprograms the response to ischemic injury and determines an altered expression of genes and proteins is well known and commonly accepted (3). This evidence drives researchers to examine whether cerebral ischemic preconditioning might be related to changes within the expression of miRNAs in brain tissue (9, 12, 13, 48). In specific, the impact of IP on miRNA expression profiles was basically assessed by three experimental works. In the first work, only eight miRNAs out of 360 analyzed were selectively upregulated in the brain of rats 3 h after IP (11). These miRNAs were categorized into the following groups: miR-200 family, including miR-200a, −200b, −200c, −429, and −141, and miR-182 family, including miR-182, −183, and −96. Notably, the increased expression of miR-200b, −200c, and −429 could explain the protective upregulation of HIF1α observed in the brain of rats subjected to IP. Indeed, these miRNAs target prolyl hydroxylase two gene (PHD2), an enzyme involved in HIF1α catabolism (11).

In another study, the effect of IP-only ischemia and post-IP tolerance to ischemia on brain miRNA expression profiles was examined; differences in terms of miRNA expression between male and female ischemic mice were also reported (17). In particular, a large subset of miRNAs was uniquely dysregulated in the IP group, including members of the mir-200 and −182 families, which were overexpressed. In addition, in silico prediction analysis allowed the identification of target miRNAs, dysregulated following ischemic tolerance. Notably, the predicted target methyl-CpG-binding protein 2 (MeCP2), which is a global regulator of transcription, appeared of particular interest in the development of the ischemic damage, and further confirmatory analysis demonstrated the relationship between miR-132 and MeCP2. Indeed, the overexpression of MeCP2 observed by immunohistochemical staining in brain tissue during ischemic tolerance was accompanied by the reduced expression of miR-132 (16, 17).

MeCP2 emerged as an important miRNA target also in another study carried out in spontaneously hypertensive rats exposed to ischemic preconditioning (50). Among the 265 miRNA screened, only the expression of 20 miRNAs was exclusively modified by preconditioning. In particular, 11 miRNAs were overexpressed, and nine miRNAs showed a reduced expression (16). Beside MeCP2, other major pathways targeted by stroke-regulated miRNAs and participating in cell regulation, proliferation, and apoptosis comprised MAP-kinase, miTOR, Wnt, and GnRh (50).

Few information is available on the role played by miRNAs in remote ischemic limb post-conditioning (RLIP), and studies in this field are in its infancy as testified by the scarcity of published papers. However, since this neuroprotection strategy presupposes a cross talk between the periphery and the central nervous system, it is possible to hypothesize that miRNAs play a role in this process of cell-to-cell communication.

Indeed, RLIP determines a marked neuroprotection by a not dangerous occlusion of the femoral artery (5). Although this method is being evaluated in clinical trials all over the world, the mechanisms activated by RLIP and implicated in the protection have yet to be fully understood (5, 65–67).

Seeing enormous potential, several research groups have focused their efforts on evidence that a short blockage of a distant artery, such as the femoral artery, can protect the brain against ischemic insults, a process known as RIC (4, 5, 65, 68–71).

In a clinical trial published in 2010, this procedure has led to excellent results in patients affected by myocardial infarction, when performed prior to percutaneous coronary intervention (72, 73).

Independently from tissue or organ affected by the harmful event, the mechanisms underlying the phenomenon of RLIP are classified in three steps:

1. The first events occurring in the remote organ or tissue, generated by the RIC stimulus (71). Indeed, blockage of the arm with a tourniquet or blood pressure cuff can trigger the release of autacoids like adenosine, Bradykinin, and calcitonin gene-related peptide, or likely of miRNAs, which protect the target organ or tissue (5, 65).

2. The type of the protection signal that is sent from a distant organ to a target organ. Neural and humoral mechanisms have been postulated as means to transport the peripheral signal from an organ such as a leg to a distant organ such as the brain or heart to explain how this phenomenon works (5, 65).
3. The event that takes place in the target organ and mediates the protective response (69).

Since miRNAs represent good candidates involved in all three phases above described, a putative translational strategy to induce stroke neuroprotection could consist in identifying miRNAs involved in these processes and in modulating these identified miRNAs by using either miRNA mimics or miRNA blockers, anti-miRNAs (9, 18, 19). In this regard, the use of miRNA mimics or anti-miRNAs appears to be a very promising method, given that a single miRNA can regulate the expression of multiple proteins at the same time and that cerebral ischemia is a multifactorial disorder with multiple potential therapeutic targets (9, 16, 18, 19).

Recently, microarray microfluidic analysis of 810 miRNAs in the ischemic rat brain revealed that let-7a-5p, miR-143-3p, miR-451-5p, and miR-485-3p had more significant expression alterations than the others. Their levels did increase significantly 24 h after ischemia induction, but when damaging ischemia was followed by remote ischemic postconditioning treatment, they were practically back to pre-ischemic levels (7). Further, more targeted investigations revealed that only let-7a and miR-143, out of the four miRNAs found and selected, would represent an important tool for stroke intervention. These findings are in line with earlier expression data and functional trials, as well as research on putative miRNA targets (49, 50). Indeed, knocking out the let-7a gene can protect against cerebral ischemia/reperfusion injury by reducing apoptosis and inflammatory reaction markers, as evidenced by a decrease in the number of p-p38 MAPK- and p-JNK-immunoreactive cells, as well as lower levels of TNF-α and IL-6 after treatment with a let-7a inhibitor (74). Let-7a has also been shown to target mitogen-activated protein kinase phosphatase 1 (MKP-1), which inactivates JNK1/2 and p38, suggesting a role for this miRNA during neuroinflammation and apoptosis (74).

Many membrane proteins involved in transduction pathways and solute transporters from different families are identified as probable targets of miR-143-3p (75). As a result, its function appears to be linked to cell responses to external stimuli. Recently, miR-143 has been found to play a role in increasing mitochondrial damage in myocardial ischemia via targeting PKCs (75). The activation of PKCs in cardiac myocytes is related to the opening of KATP channels. The latter occurs by preventing the opening of the mitochondrial permeability transition pore (mPTP) (75, 76). Notably, it has been demonstrated that a reduction of focal cerebral ischemic injury induced by delayed remote postconditioning may be achieved also through opening of KATP channels (77). Taken together, these data suggest that reduction of miR-143 expression in the brain after tMCAO plus RLIP might prevent PKCs downregulation, with subsequent opening of KATP channels, thus promoting mPTP closure. Although the action of these two miRNAs occurs at the neuronal level, miRNA-143 expression was also seen in the bodies of astrocytic GFAP-positive cells, validating the idea that astrocytes and neurons show different miRNA expression patterns after ischemia (78). Other investigations have corroborated the distinctive expression of miRNA in neurons and glial cells, and numerous hypotheses have been proposed to explain this phenomenon (78, 79). As a result, after remote postconditioning, astrocytes and neurons express different miRNAs; this could be due to differences not only in their miRNA repertoires but also in their cell-specific roles within the CNS.

It is worth noting that let-7a and miR-143 mimic therapy significantly reduces the neuroprotection afforded by RLIP. As a result, these two miRNAs and the proteins they target appear to be potential RLIP-induced neuroprotection mediators. More research is needed to understand which of the target proteins is involved in remote postconditioning-induced neuroprotection. As a result, altering the levels of expression of these miRNAs and their target proteins appears to be a promising stroke therapeutic strategy.

EXPLOITATION OF miRNAs AS STROKE THERANOSTIC TOOLS: PITFALLS AND CLINICAL PERSPECTIVES

The incessant work of recent years in studying the role of ncRNAs in stroke made important contributions that advance both the so-called miRNA revolution and our knowledge of the possibilities of exploitation of miRNA as theranostic tools in stroke.

A list of miRNAs responsible for the infarct-sparing effect of brain conditioning is now available (7, 9–13, 17). Furthermore, it is now clear that delivery of a class of the miRNA homologs may represent a novel avenue in therapy, and the changes in miRNA plasma concentrations could be used as biomarkers. However, the preliminary findings raise significant concerns and debates about their applicability in clinical practice.

To begin, it is critical to note that all conclusions are derived from preclinical models. Several interventions that had been shown to enhance outcomes in experimental animal stroke models, however, failed in clinical studies. Systematic assessments of experimental stroke studies have consistently found low-quality scores, unfavorable publication bias, and a lack of data from female, elderly, or comorbid animals, casting doubt on the robustness and predictive utility of single-laboratory preclinical investigations. The new idea of a multicenter preclinical randomized controlled trial (pRCT) is developing as a critical step before transitioning from animal modeling to clinical trial to improve the translation of therapy efficacy from bench to bedside. Notably, we have recently begun a multicenter preclinical study in rats and mice of both sexes to explore the efficacy of RIC in the experimental model of temporary middle cerebral artery (MCA) occlusion (trial registration number PCTE0000177).

An additional and arguably tangential issue that deserves investigation is the assessment that the majority of papers published in this field did not associate the described favorable effects with transport via exosomes, a finding that appears to contradict the emerging concept that extracellular vesicles serve as vectors for the RIPC-initiated humoral communication of protective signals to the brain.

The cellular source of the circulating miRNAs released in response to the RIPC stimulation is a third question that will be
difficult to answer: are miRNAs released from skeletal muscle or from a more conventional, blood-borne source?

Fourth is the importance of defining a temporal profile of expression. miRNAs are released in response to a stimulus in a time-related manner. Defining a precise timing of determination after stroke onset is fundamental in the case miRNAs are used as biomarkers, but is also extremely important when miRNA mimics are used as therapeutic tools.

In the vast majority of preclinical studies, MCAO was used to confirm alterations in specific miRNAs in young healthy male rats or mice. Furthermore, the majority of clinical stroke studies did not specify the number of male and female patients who participated in the trials, making it difficult to discuss the importance of gender-specific changes in miRNAs and their contributions to ischemic stroke and/or responses to antagonism of mimics. In future investigations, men and females animals must be used to confirm the upregulation or downregulation of particular miRNAs. Finally, because most stroke patients have comorbidities and are over the age of 50, animal stroke models with hypertension, hyperlipidemia, and diabetes mellitus, as well as older animals, should be included in confirmation investigations.

Lastly, it will be extremely important to define a delivery system. A successful miRNA therapy necessitates a precise and effective delivery method. To address the intrinsic instability of miRNA in circulation, off-target effects, and improper distribution, a delivery method is required, but it does not need to reduce cell permeability, increase excretion, or accumulate in off-target organs (10). MiRNAs can be injected intravenously or subcutaneously because they are small and water-soluble. Because miRNAs are single-stranded and open-ended, they are either destroyed by systemic nucleases in circulation or eliminated by the kidneys (56).

Accumulation in non-target organs such as liver and spleen, non-specific absorption, excretion, toxicity, and an immune-mediated response are among issues that might arise with various delivery systems (56). Systemic delivery must consider the delivery system’s stability in circulation as well as tissue-specific targeting. Some options for systemic delivery have been developed through ongoing research. In this regard, the viral capsid can be modified for tissue-specific delivery, and viral vectors can be used to boost circulatory stability during translocation (56). Adeno-associated viruses have exhibited great tissue specificity and acceptable safety profiles in clinical trials for gene therapy (37). Viral vectors, however, do have pitfalls including immune reactions and viral integration into the host genome (56). Other lipid-based vectors, like liposomes, are protective against nuclease, lysosomal, and endosomal degradation and can be effectively used (40). As natural transporters, exosome delivery systems are an attractive option. These offer specificity by binding in a receptor-mediated fashion, limiting off-target side effects (59). Nanoparticles are another possible strategy and have been used in an attempt to overcome excessive inflammatory reactions (8). An intriguing concept for drug delivery for IR injury is “passive drug targeting” where a colloidal-based drug delivery system is utilized. Because the endothelium at these places is weakened, these chemicals collect at sites of inflammation, allowing passive diffusion through the artery (40). Another option for systemic administration is mesenchymal stem cell-derived extracellular vesicles (MSC-EV), which have been investigated in vivo and found to be able to transport miRNAs (52). MSC-EV, like the other delivery systems, are currently being studied for clinical usage, with promising results.

CONCLUSIONS

Efforts will be needed in the next decade of miRNA research to enhance and evolve the tools for miRNA analysis and validation, as these technologies will be critical in establishing direct correlations between miRNA-mediated post-transcriptional gene expression and disease (9, 19). Furthermore, it is important to underline the need to speed up all those technical procedures capable of detecting miRNAs in the shortest and simplest possible way, to avoid running into the same problems currently present with the use of CT and MRI to make differentiated diagnosis of ischemic and hemorrhagic stroke.

AUTHOR CONTRIBUTIONS

The author confirms being the sole contributor of this work and has approved it for publication.

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