The pervasive role of microRNAs in arrhythmia: Animal models and novel discoveries

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Introduction

Cardiac arrhythmia and in particular atrial fibrillation (AF), which is the most common sustained arrhythmia, is one of the main cause of morbidity and mortality worldwide [1]. The aetiology is attributable to multiple factors such as genetic, lifestyle and other pathological stress [2]. For pathologies as long QT syndrome (LQTS), Brugada syndrome, Timothy syndrome, "torsade de pointes" phenotype and cardiac sudden death, a genetic aetiology was recognized. However, even for arrhythmia syndromes with genetic basis, prediction of the phenotypic expression was not simple because of the extreme variability in the severity of disease manifestation also among family members presenting the same mutation [3]. Indeed, the arrhythmic phenotype has features of a "complex phenotype" which probably reflects the intricate and interconnected events that occur in cardiac cells, from the action potential activation to the regulation of excitation contraction coupling event. For these reasons, cardiac arrhythmia remains a big challenge for modern drug treatment and the comprehension of how DNA variation and other concomitant factors might influence disease is of great importance. In this panorama, microRNAs (miRNAs), small molecules of 22-25 nucleotides negatively regulating gene expression [4] might represent an interesting field of investigation. Although they are no longer a new discovery, the interest in their functions in heart pathologies and in their possible use as therapeutic molecular drugs, continues to be under study, thanks also to their unique capacity to contemporarily modulate more genes and whole pathways. Following this prospective, several clinical trials have been initiated using miRNA- and siRNA-based therapeutics [5]. Even if we are far to create a miRNA-based treatment in arrhythmia, different evidences demonstrated the possibility to pursue this aim for cardiac pathologies as suggested by the action of 2 molecular drugs MGN-9103 [6] and MGN-1374 [7] developed by an American company, used to treat heart failure and cardiac infarction respectively.

A deeper knowledge of roles played by miRNAs in the development of cardiac arrhythmias is a crucial prerequisite to enable the use of these small molecules as new drugs or drug targets. Although in the literature is present an extensive characterization of miRNAs targeting genes involved in heart physiology, there are few papers describing miRNAs with a causative role in arrhythmogenesis. This review is focused on miRNA genes deeply involved in arrhythmia and on the principal animal models, which allowed to decipher the functional miRNA role in arrhythmic cardiac pathologies.

Genome wide analyses and miRNAs

The advent of genome wide approach allowed to discover association among genetic loci and pathological states and represents the classic genetic analyses to find signature for heritable risk factors. During last years the attention focused on mutation located in coding sequence (CDS) of genes [3]. A genome wide analyses unveiled a strong association between two sequence variants on chromosome 4q25 and AF. Both variants were in close proximity to PITX2 gene [8] and this observation leads to discover a direct regulation of miR-17-92 and miR106b-25 clusters [9] by PITX2 and successively the involvement of these miRNA clusters in arrhythmia.

Besides "indirect" approaches to identify miRNAs with a role in arrhythmia, the number of studies focused on discovering human miRNA sequence variants are rather limited. In the analyses of Hedley et al. DNA sequence variations in genes coding for miR-1 and miR-133a in LQT probands were inspected, but the results showed that these miRNA variants were not correlated to LQTS in this cohort [10]. On the contrary, a sequencing of 2600 individual DNAs allowed to find a naturally occurring mutation in the mi-r-499. Overexpression of wild-type and mutant miR-499 in murine heart showed that the presence of the miR-499 c17 mutation alters myocardial protein levels of the miR-499 targets leading to a less severe cardiac remodelling after heart failure [11].

Sequencing the regulatory regions of genes, not only the CDS, should be of particular importance in miRNA investigation. Indeed, functional microRNAs bind the miRNA targets through RNA-RNA base pairing generally in the 3‘-UTR. The canonical recognition requires usually a perfect pairing between roughly a 6–8 nucleotide stretch of the 5’-end of the miRNA, known as "seed" [4,12,13] and the corresponding complementary sequence in the target 3’-UTR, called "seed-match".

Therefore, mutations in miRNA seed-match outside the CDS of genes related to arrhythmic pathologies, might be the causative effect in cases of arrhythmias with yet-unknown genetic aetiology. The literature describes several examples of polymorphisms in the 3’-UTR of key genes associated to heart diseases corroborating this hypothesis [14-19]. However as mentioned before, these analyses are mainly association studies which do not investigate the functional meaning of these variants. Insightful was the work of Hoffman et al. focused on the analysis of SHOX2 in AF patients. The authors showed in particular that patients carrying the 3’UTR variant c.*28T>C present significantly longer PR intervals. Mechanistically, the variant creates a new functional
binding site for hsa-miR-92b-5p, which may reduce SHOX2 expression and thereby facilitate proarrhythmogenic remodelling leading to AF [20].

MiRNA effectiveness is based on thermodynamic forces and affinity to targets, which are parameters used in several algorithms to predict RNA-RNA recognition. Changes in this affinity affects the balance of gene expression and the contribution of each miRNA in regulating shared miRNA targets or the possibility for a miRNA to interact with competitors in a big net of communication [21,22]. The amount of functional miRNA results from the balance among competitive targets and the final result is a combination of RNA abundance and phenomena of redundancy with palliative action. In this very complex panorama, emerged long non-coding RNA [lncRNA] as new players in arrhythmic disease, because they can behave as miRNA sponges. Evidences of their possible role emerged from the observation of a different expression profiles of lncRNA in AF patients [23]. An example is the lncRNA Kcna2 antisense RNA [24] whose expression increases in rats with congestive heart failure [CHF] and that has been recently shown to contribute to ventricular arrhythmia via silencing Kcna2. Another interesting case of direct relation among miRNAs and lncRNAs in arrhythmic pathologies is the lncRNA TCONS_00075467 that modulates atrial electrical remodelling by sponging miR-328 which regulates the downstream CACNA1C protein [25]. Moreover, a complex molecular mechanism was discovered by Zhu et al. for LncRNA MALAT1 that was found upregulated in cardiomyocytes from arrhythmic model rats [26]. It was suggested that MALAT1 might act as a competing endogenous RNA for miR-200c that targets the high-mobility group box 1 (HMGB1) protein, a highly conserved nuclear protein that operates as a chromatin-binding factor. The upregulation of HMGB1 consequent to the downregulation of miR-200 might mediate the downregulation of the transient outward potassium current [26].

The interplay between miRNAs and Transcription Factors

Since miRNAs and transcription factors (TFs) are trans-acting factors that interact with cis-regulatory elements, they potentially generate a complex combinatorial code. This is clearly shown by the observation that genes with many TF binding sites have a higher probability to be controlled by miRNAs than genes with few TF binding sites [27]. Emerging evidences indicate a crucial miRNA/TF interplay in many biological processes and in arrhythmia. PITX2, TBX5 and MYOCOD are among the TFs principally involved in heart morphogenesis and especially studied for AF [28]. Classic analysis, based on in silico approach and in vitro assays to determine miRNA/ mRNA functional interaction, revealed miR-21, miR-10b, and miR-1 as mayor regulator of PITX2C, TBX5, and MYOCOD respectively in AF patients [29]. A more sophisticated molecular screening method, the PAR-Clip method, found several other miRNAs targeting PITX2 such as miR-153, miR-5692, miR-369, miR-4517, miR-5583, miR-377, miR-374 [30] whose role in pathogenesis is still unknown.

By high throughput comparison of zebrafish embryos depleted or not for Tbx5 [31] the miR-17-92 cluster was identified as directly regulated by Tbx5. In zebrafish, cardiac defects generated by Tbx5 depletion can be partially reverted by modulation of miR-19a, a member of this cluster, in line with a role of this miRNA as Tbx5 functional effectors. Furthermore, miR-19a is able to affect heart rate control in Holt-Oram syndrome [HOS] fish by targeting the Sphingosine-1-Phosphate Receptor 1 which is highly expressed in mammalian cardiomyocytes and implicated in numerous cardiovascular processes [32]. Moreover miR-17-92 cluster has been also shown to directly repress genes, such as SHOX2 and TBX3, that are required for sinoatrial node development while miR-1 also targets the zinc finger homeobox 3 gene (ZFHX3) which is another important AF susceptibility-conferring gene [33-35].

A potentially interesting factor that, differently from the previous we mentioned, is considered as transcriptional repressor, is the neuron-restrictive silencer factor (NRSF) [36], that selectively regulates the expression of multiple fetal cardiac genes [37]. In transgenic mice the expression of a NRSF dominant-negative mutant (dnNRSF) leads to cardiomyopathy, high susceptibility to arrhythmias and sudden death [38]. Although a direct relationship between NRSF gene and regulating miRNAs is not still discovered in arrhythmias, intriguingly, a PAR-clip extensive analysis of miRNA-RNA-targets interaction revealed miR-106 and miR-17-5p, known to be involved in arrhythmias [see next chapter], among the miRNAs able to recognize NRSF.

Functional characterization of miRNAs in animal models of arrhythmia

The "OMIC" studies, which are fascinating and nowadays indispensable for genetic investigation, can be tricky in the phases of matching and interpretation for miRNA data analysis. Slight modification in miRNA expression might lead to a pathological state and methods used to evaluate miRNA abundance in a sample might be critical, with a risk to lose information. An example of a big effort ended in a poor result was a study of 2016 aimed to find a strong net of regulation among deregulated miRNAs and related validated targets in AF [40]. From this study based on microarray expression profiles and bioinformatics analysis, emerged only generic classes of gene ontology on "metabolism" and "regulation of cellular/biological process" which does not improve our knowledge about the regulatory network in AF.

Significant steps forward in the characterization of miRNAs causative of arrhythmias have been done by exploiting transgenic/mutant animal models, which can also allow to evaluate miRNA therapeutic potential.

The miR-1, maybe the most cited cardiac miRNA, is highly expressed in cardiac tissue and found deregulated in the major part of heart pathology studies. MiR-1 is involved both in cardiogenesis and cardiac conduction system development [41]. A recent study based on transgenic mice with cardiomyocytes-specific over-expression of miRNA-1, revealed a new contribution in arrhythmias. This study showed that miR-1 overexpression dysregulates intracellular trafficking and, as a consequence, alters calcium handling by targeting Syntaxin 6, which indirectly decreased Cav1.2 expression [42]. Due to the huge amount of miR-1 target genes in cardiac cells [43], the use of this miRNA, or of similar highly multitasking miRNAs, in gene therapy is challenging. A potential therapeutic role of miR-1 is shown only in models of myocardial infarction [44] and cardiac reprogramming [45].

The miR-17-92 cluster is known to be involved in heart development and its deregulation was associated to pathological cardiac states, but the physiological involvement of each cluster member is not completely understood and is still under study. MiR-19a/b members of the cluster were associated to arrhythmias because they target cx43 gene [46]. Exploiting the simplified but conserved zebrafish model, the involvement of miR-19 in heart rate was uncovered. Zebrafish lacking miR-19b by CRISPR-mediated knockout show severe bradycardia as well as susceptibility to arrhythmias and cardiomyopathy. Moreover miR-19b affects action potential duration [APD], in part, by directly targeting KCNQ4 [47].
To gain insight into miR-17-92 cluster involvement in arrhythmia, miR-17-92\textsuperscript{null/+} mice were studied since miR-17-92\textsuperscript{null/null} mutants show post-natal lethality. This study included also miR-106b-25 cluster that shears with miR-17-92 cluster miRNAs belonging to the same family and that, together to miR-17-92 cluster, is controlled by PITX2 [see also previous chapter of this review]. Although neither miR-17-92\textsuperscript{null/+} nor miR-106b-25\textsuperscript{null/+} or miR-106b-25\textsuperscript{null/null} mice show spontaneous episodes of AF, in condition of pacing-induced AF, the loss of miR-17-92 and miR-106b-25 increases susceptibility to AF. Interestingly, mice with miR-17-92 cardiac-specific knockout or miR-106b-25 haploinsufficiency showed arrhythmogenic events and senoatrial node dysfunction [9]. In a different study, miR-106b-25\textsuperscript{null/null} mice exhibited enhanced sarcoplasmic reticulum Ca\textsuperscript{2+}-leak and increased ryanodine receptor type-2 [RyR2] expression [34] in line with studies showing that the level of RyR2 protein was elevated in atria of paroxysmal AF patients [48].

In a different manner, cardiac specific overexpression of miR-206 in adult mice was shown to suppress cx43 expression and to induce abnormal heart rate and PR interval causing also shortening of life-span in the experimental mice [49]. Moreover, upregulation of miR-206 cluster induced by overexpression of HMG1B, was shown to induce downregulation of TimP3, an inhibitor of metalloproteinases 2 and 9 [MMP-2 and MMP-9] [50,51]. This data suggests that miR-206 modulation, affecting collagen deposition activity and enhancing collagenolytic activity, can potentially have not only a causative role in arrhythmogenesis but also impact AF remodelling phenomena. Another miRNA potentially involved in AF remodelling is miR-208a in the heart affects both cardiac growth and conduction. In zebrafish where its role in cardiac development and function has been confirmed. Upregulation of miR-182 induces downregulation of several calcium channel proteins and alteration of calcium handling. In line with this observation, stable overexpression of miR-182 in zebrafish myocardium is able to induce spontaneous events of arrhythmia at 3 months of age and die suddenly between 6 and 12 months of age [57].

A recent mRNA revealed to play a possible important role in arrhythmia is miR-182 identified in mouse model of HOS as downregulated by Tbx5. This mRNA has been functionally studied in zebrafish where its role in cardiac development and function has been confirmed. Upregulation of miR-182 induces downregulation of several calcium channel proteins and alteration of calcium handling. In line with this observation, stable overexpression of miR-182 in zebrafish myocardium is able to induce spontaneous events of arrhythmia at 3 months of age and die suddenly between 6 and 12 months of age [57].

To a further mRNA which is deregulated in AF patients is miR-208 [52], considered a cardiac specific miRNA. Transgenic overexpression of miR-208a in the heart affects both cardiac growth and conduction. On one side miR-208 targets thyroid hormone–associated protein 1 and myostatin, which are negative regulators of muscle growth and hypertrophy and, on the other side, this miRNA affects the cardiac conduction system by controlling GATA4 which in turn regulates the expression of the cardiac Tfs homeodomain-only protein [Hop] and the gap junction protein cx40 [53].

Deregulation of miR-499 and miR-27b have been also shown to increase AF vulnerability although studies in animal model clearly demonstrated that these miRNAs are not strictly arrhythmogenic. MiR-499 is overexpressed in patients with permanent AF [54] and its constitutive overexpression in transgenic murine hearts caused hypertrophy. MiR-499 deregulation seems to primarily influence cardiac gene expression and predispose to cardiac stress-induced dysfunction [55]. MiR-27b, identified because up-regulated in a mouse model of high fat diet [56], increases vulnerability to atrial arrhythmia by targeting cx43. Transgenic mice with cardio-specific overexpression of miR-27b born normally and have normal cardiac structure, although 30% of them develop significant cardiac hypertrophy at 3 months of age and die suddenly between 6 and 12 months of age [57].

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**New miRNAs discovered targeting gene involved in Arrhythmia**

Besides the microRNAs just described, the following table presents a short list of validated miRNA-mRNA targets derived from literature analysis published from 2016 to 2018 and generated by searching for genes known to be involved in arrhythmia or for new discoveries underlie arrhythmic disorders (Table 1).

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