MicroRNA-183 Family in Inner Ear: Hair Cell Development and Deafness

Mohammad Reza Mahmoodian sani¹, Morteza Hashemzadeh-Chaleshtori², Massoud Saidijam¹, Mohammad-Saeid Jami², and Payam Ghasemi-Dehkordi²

¹Research Center for Molecular Medicine, Hamadan University of Medical Sciences, Department of Genetics and Molecular Medicine, Hamadan University of Medical Sciences, Hamadan, ²Cellular and Molecular Research Center, Shahrekord University of Medical Sciences, Sharekord, Iran

miRNAs are essential factors of an extensively conserved post-transcriptional process controlling gene expression at mRNA level. Various biological processes such as growth and differentiation are regulated by miRNAs. Web of Science and PubMed databases were searched using the Endnote software for the publications about the role miRNA-183 family in inner ear: hair cell development and deafness published from 2000 to 2016. A triplet of these miRNAs particularly the miR-183 family is highly expressed in vertebrate hair cells, as with some of the peripheral neurosensory cells. Point mutations in one member of this family, miR-96, underlie DFNA50 autosomal deafness in humans and lead to abnormal hair cell development and survival in mice. In zebrafish, overexpression of the miR-183 family induces extra and ectopic hair cells, while knockdown decreases the number of hair cell. The miR-183 family (miR-183, miR-96 and miR-182) is expressed abundantly in some types of sensory cell in the eye, nose and inner ear. In the inner ear, mechanosensory hair cells have a robust expression level. Despite much similarity of these miRs sequences, small differences lead to distinct targeting of messenger RNAs targets. In the near future, miRNAs are likely to be explored as potential therapeutic agents to repair or regenerate hair cells, cell reprogramming and regenerative medicine applications in animal models because they can simultaneously down-regulate dozens or even hundreds of transcripts.

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KEY WORDS: miRNA · miR-183 family · Hair cell · Hearing loss.

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family in inner ear: hair cell development and deafness.

**History of miRNA and Biogenesis**

The first miRNA (lin-4) was discovered in 1993 [1]. Generally, miRNAs are small RNAs originated from a stem-loop shaped precursor RNA that mainly intervenes in the regulation of gene expression. This regulation occurs after targeting and leading particular mRNAs to degradation [2]. The seed region of the 5' mature miRNA molecule including seven nucleotides contribute to binding to the 3'-UTR of complementary mRNA [3]. The miRNA molecule can completely bind to the mRNA target leading to full degradation of the target mRNA or partially bind to the target mRNA and cause the inhibition of mRNA translation. A wide variety of miRNAs exist in different cells and tissues which play an important role in many biological processes such as cell cycle, apoptosis, tumorigenesis and neurogenesis [4,5]. In addition based on bioinformatic predictions, almost one third of protein-coding genes in the human genome are regulated by miRNA [6]. To date, more than 100 gene loci have been demonstrated to be involved in deafness, 44 of which have been reported to possess at least one mutation; however, there are still a large number of unexplained cases. Approximately 98% of RNA in mammalian cells do not code protein [7] and non-coding RNA such as miRNAs may be involved in the development of the inner ear and thus have a role in hearing loss [8]. In 2005 miRNAs were identified in the vestibular system for the first time. Since the discovery of a tissue-specific miR-183 family and its correlation with deafness in mice and humans, miRNAs have been shown to be involved in some other diseases such as cholesteatoma, schwannomas and otitis media.

**The miR-183 Family**

miR-96, detected in human cancer cell lines, was the first discovered member of human miRNA of this family [9]. Bioinformatics tools allowed the discovery of miR-182 and miR-183 through comparison between human or mouse RNAs with Fugu rubripes genome (a type of Japanese puff-fish) [10]. Considering the same expression pattern in retina and very close chromosomal loci, both miR-182 and miR-183 are categorized in the family [11]. In addition, miR-96 was added to this family due to its sequence homology with miR-182 and miR-183, and its chromosomal locus that is between miR-182 and miR-183 [12,13]. miR-183 family consists of three homologous miRNA, miR-183, miR-182, and miR-96 (Fig. 1) which are expressed in sensory neurons and hair cells in vertebrates and sensory cells of all species of animals [14]. Among vertebrates, members of miR-183 family are expressed in the olfactory epithelium, eye, neuromast and ear. In the inner ear of mice, miRNAs are expressed in sensory neurons and hair cells of the corti organ and end organs of vestibule (Table 1) [15]. Minor differences in seed sequence among this family members causes them to have independent target RNAs (Fig. 1). This miRNAs is expressed in a completely synchronous manner throughout development and is needed for maturation of sensory organs [16]. miR-96, mir-182, and miR-183 comprise a fully conserved family of miRNAs and localize in intergenic loci of chromosome 7 long arm (Fig. 1) [17]. In vertebrates, the expression of miR-183 family seems to be limited to ciliated sensory epithelial cells and certain cranial and spiral ganglia. In zebrafish, these miRNAs have been detected in eye, nasal epithelium, sensory hair cells in ear, and neuromasts [18,19]. miR-182, alongside miR-183 and miR-96, are from a polycistronic miRNA cluster which localize in a 4 kb region on the long arm of chromosome 6. They are expressed particularly in certain organs such as eye, nose, and inner ear [12].

**miR-96**

miR-96 could indirectly cause increase in the expression of significant developmental genes such as Gfi1, Ptprq, and Tmc1 [21]. In olfactory system, this miRNA family is vital for differentiation of neural precursor into mature olfactory neurons, but has no contribution to onset of specific differentiation or maturation steps [22]. miR-96 is likely to play a similar role in auditory system. Understanding miR-96 mechanism may help to explain the complicated expression of the genes required for cochlear activity and contribute to advances in non syndromic progressive deafness. Studies of diminuendo mice (a mice model to investigate the causes of
hearing loss and its progression) have demonstrated that a mutation in miR-96 gene causes a mutation at locus 7 of miR-96 seed sequence [23]. Functional activity of miR-96 as an miRNA-183 family member was confirmed in a mice model with a mutation at seed regions loci. The mutation was caused by N-ethyl-Nitrosurea and led to partial hearing loss in heterozygous mice and absolute hearing loss in homozygous mice. The mechanism through which miRNAs affect inner ear development has not been yet explained fully. In a study to determine the miR-96 target genes, 12 genes were predicted by miRanda software with strong filtering, five of which (Aqp5, Celsr2, Myrip, Odj2, and Ryk) were confirmed as miR-96 target by analysis of luciferase. In a study to compare the genes expression in inner ear between wild and mutant diminuendo mice using microarray, 96 transcripts were affected significantly and the expression of five genes (Pitpnm1, oncomodulin, Preston, Ptprq, and Gf11) was found to decline. None of these genes have a binding site for miR-96. They are likely to affect downstream targets indirectly and the changes in expression could phenotype diminishing mice. Studies of diminuendo mice have indicated that miR-96 is responsible for maturation of stereocilia bundle in inner and external ear [24]. In addition, synaptic morphology of mutant mice remained immature and miR-96 is speculated to contribute to developing cochlear nerve. C1ic5 is an inner chloride cell canal alongside stereocilia in inner ear and has been known as a target for both miR-96 and miR-182 [25]. Mutant mice for C1ic5 have a similar stereocilial morphology to ENU mice which has been already described. C1ic5 has a binding site for miR-96, which has been confirmed by luciferase. Transfecting miR-96-contained liposomes into HEICO1-derived hearing cell leads to decline of C1ic5 at surfaces of both mRNA and protein [26].

**miR-182**

miR-182 is expressed as hair cell-like at differentiation of fundamental precursor cell of inner ear. Differentiation of fundamental precursor cell of inner ear into hair cell-like is accelerated. miR-182 activity could be associated with targeting Tbx1, which is a transcription factor playing role in development and differentiation of hair cell [27]. Increased expression of miR-182 causes increase in the number of ectopic hair cells in the inner ear. Ectopic expression of miR-182 in supporting cell could suppress and convert their gene into hair cell. Therefore if deafness is due to losing hair cells, direct transfection of miR-182 into SC could facilitate their transforming into hair cells (hair cell regeneration). Previous studies have shown that miR-183 family members are likely to suppress prosensory or SC genes to help to differentiate hair cell specifically. SOX2 as a marker prosensory cells which is also targeted by miR-182 [28,29]. In several genome wide studies on autism, the susceptibility gene to this disease, which is near 7q32/2, has been detected close to miR-182 chromosomal locus. Defective activity of miR-182 family could cause defect in sensory development steps and contribute to autism incidence [30-35]. C>A mutation has been detected at a binding site of miR-182 on radixin (RDX) 3'UTR in a human family with recessive autosomal non syndromic deafness [36]. Moreover in addition to the mutation at protein-coding loci, mutation at 3'UTR non coding loci, where miRNAs usually bind, could cause disorder in gene activity. Although in vitro investigation of luciferase did not confirm the mutual effect between RDX and miR-182 [29]. In vitro studies have indicated that Microphthalmia-associated transcription factor is a transcription factor required for retinal development and maintenance. A gene is directly targeted by miR-182. miR-182 is likely to contribute to regulating visual circadian rhythm, which is probably done by changing ADCY6 adenylyl cyclase expression [12].

**Expression and Activity of the miR-183 Family in the Developing Hearing Organ of Chicken**

Basillar papilla is similar to mammals auditory organ (corti organ) in various dimensions [37-39]. In studies, chicken embryo was used to investigate whether this family was expressed. Basillar papilla, an auditory organ, exhibited a mild increase in expression (mostly on peripheral neuron) in prosensory cells at seventh embryonic day and an increase in expression (mostly in apical hair cells) nine years later for this family. However transfection had no effect on phenotype. No detectable change in hair cell or bundle morphology was seen due to increased expression by miR-183 family vector [40].

**Expression and Activity of the miR-183 Family in Hearing Organ of Mouse**

Throughout development of mice embryos, miR-183 family is initially expressed across the entire otic and spiral vesicle and atrial ganglionic neurons, and then only in hair cells and neurons accompanying them. miR-183 family has been also seen in greater epithelial ridge at 15.5–17.5 embryonic days, and is expressed in inner sulcus and spiral lumbus 4–11 days after birth [41]. Although miR-183 family in the in-
miR-183 Family Activity in Hair Cell

The significance of miR-183 family for hair cell development has been recently confirmed in human, mice, and zebrafish (Fig. 3). In genetic investigation of deafness locus of human autosomal the two families had a common point mutation in seed sequence which was transferred to mature miRNA and cause loss of miRNA function. Knocking down miR-183 family members cause defect in semicircular canals and inner ear [51].

The Functional Role of miR-183 Family in the Development of the Inner Ear

The functional role of miR-183 family members has been explored in inner ear development in zebrafish embryo [52]. Synthetic injection of miR-182 and miR-96 into embryos accelerates hair cell ectopic growth while mir-183 leads to hair cell normal development (Fig. 4). Although this family are a scended from similar initial transcript, their role varies probably due to difference in their downstream targets. miR-183 family inhibition using morpholino antisense oligonucleotides affects ear normal development at frontier and posterior macula. The severity of hair cell loss depends on the number of inhibited miRNA. All miR-183 family members are involved in controlling sensory cell. Since both the inhibition of this family proliferation and increase in its expression

Table 1. Validation of miRNA-gene targets found in the inner ear

| miRNA  | Gene target                                   | Experimental system used                                   | Ref.                                      |
|--------|----------------------------------------------|-----------------------------------------------------------|------------------------------------------|
| miR-183| TAO kinase 1 (Taok1) Early growth response 1 (Egr1) | Rat cochlear organotypic cultures transfected with antisense morpholinos. | Patel, et al. [49]                       |
|        | (Ir1) Insulin receptor substrate 1 (Irs1)     | In situ hybridization: luciferase assay in HEK293 cells.  | Weston, et al. [50]                     |
| miR-182| SRY-box containing transcription factor 1 (Sox2) | Luciferase assay in COS1 cells: overexpression of miRNA in cultured olc progenitor/stem cells. | Wang, et al. [27]                        |
| miR-182| T-box 1 (Tbx1)                                | Co-expression in mouse auditory HE-OCC1 cells: luciferase assay in A549 cells: down-regulation of target | Liberman, et al. [26]                   |
| miR-96 | Chloride intracellular channel 5 (Clic5)      |                                                           |                                          |
| miR-182|                                             |                                                           |                                          |
cause disorder in development, the balance in expression of this family could be critical to normal development of hair cell.

**miRNAs and Regeneration of Hair Cells**

Treatment of hair cell regeneration is still a big challenge. Due to hair cell inability to regenerate, the stem cells with hair cell-like phenotype have been genetically manipulated [54]. A challenge of this approach is that how cells proliferation and differentiation should be regulated in reprogramming and differentiating. miRNAs could contribute greatly to initiating cell differentiation, maintaining differentiation steps, and terminating differentiation. To date, hundreds of miRNAs have been identified in cultured cochlear precursor cells. Expression patterns are different in differentiated and undifferentiated cells [55]. Therefore, the role of any involved miRNAs in cochlear (Fig. 2) cells differentiation has been yet explained fully. Understanding miRNAs regulatory roles could elucidate the hair cell regeneration in mammals inner ear.

**Family miR-183 and Age-Related Hearing Loss**

The above-mentioned miR-183 expression in hair cells could help to differentiate and maintain the apical structures of these cells. At older ages, the hair cells gradually lose their apical structure and hearing threshold increases. By increasing miR-183 family level in auditory cell, the apical structures and hearing ability may be recovered.

**Regulatory Factor Atoh1 and miR-183 Family**

Atoh1 is an important transcription factor which can initiate hair cell differentiation and begins to express approximately at E12.5-E14.5 in hair cells [50]. Studies have indicated that miR-183 in developing cochlea did not express at E12.5 and expressed partially at E14.5. At E16.5, miR-183 family is expressed in all sensory ganglia and 17.5 entire inner ear epithelium. These findings confirm no evidence of miR-183 family in vivo expression prior to Atoh1. Therefore, this family has no contribution to onset of development although it may play a significant role in development. The expression level of miR-183 family ranges within the initial steps of inner ear development. However miR-183 family expression is specific to hair cells in the subsequent steps of development [55].

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**Fig. 4.** miR-183 regulates the number of hair cells. In developing zebrafish, injecting miR-183 family reagents into an embryonic cell and then analyzing the inner ear were done 1–2 days after fertilization. Injecting miR-96 or miR-182 at monocellular embryonic step causes production of a large number of hair cells after one day. Declined level of any members of miR-183 family after morpholino antisense injection leads to producing few hair cells two days after injection [29,52]. Adapted from Li, et al. Curr Opin Otolaryngal Head Neck Surg 2010:18:459-65. [29]. Adapted from Li, et al. J Neurosci 2010;30:3254-63. [52].

**Fig. 5.** Shows the time table of miRNAs expression at development and early steps after birth. miR-183 family is a subgroup of miRNAs that has been already investigated more frequently. miR-183 are expressed initially in otic vesicle at E9.5, which continues till reaching into cochlear-vestibular ganglion and neural tube at E11.5 and E17.5. The above expression has been reported in cochlear hair cells till p30 in some studies. Other miRNAs were identified by ISH. *represents the miRNAs that were tested only at this step [56]. Adapted from Rudnicki, et al. EMBO Mol Med 2012:4:849-59. [56].
miRNA-183 Family and Deafness

Because of the limitations of studying miRNAs and their targets in human inner ear, animal models are very important in this regard. Therefore miRNAs role in development and activity of inner ear has been investigated in zebrafish and mice. Transcriptomic and proteomic investigations help to figure out miRNAs role and targets. miRNAs are thought to be used for regeneration of inner ear hair cells and finally contribute to treatment [56]. Normal expression of miR-183 family is specific to sensory organs and essential for their development. Knocking down miR-183 causes increase in the expression of the genes that are through the signaling axis PI3K/AKT/mTOR as they undergo apoptosis and dysregulation. Migration and invasion are upregulated by miR-183 family and mainly mediated by miR-182. miR-183 expression has been confirmed in various cancers [57-69]. The association of miR-183 family with development and diseases could be also explained by beta-catenin being able to regulate this family expression [70,71]. Beta-catenin is activated in downstream Wnt signaling pathway and is a survived transcription factor which plays a role in embryonic period and contributes to regulation of organ development. Under normal biological conditions, miR-183 expression takes place only in ciliated sensory cells, where the length and durability of survival activities are important for the entire organism. These findings make the assumption that the miR-183 family is a constituent of developmental mechanism for maintenance and longevity of sensory neuron. Obviously when miRNAs are removed in rodents, neurodegeneration takes place, confirming the hypothesis stating that this family plays a critical role in maintaining neurons health. These findings have been obtained by combined in vitro/bioinformatic studies [72]. Studies have speculated that miR-183 family is likely to suppress prosensory cells genes or to suppress SC genes to assist hair cells in specific differentiation, and SOX2 as a prosensory marker has been predicted to be a target for miR-182 [28]. Recent studies of nervous system in several organism models have demonstrated the critical role of miRNAs in development and function of nervous system. miRNAs have a great potential to help to develop a new generation of drugs but potential problems continue to raise about them, including high dose-dependent side effects and toxicity (if used in vivo) [73]. However researchers argue that miRNAs-based therapeutic interventions will be definitely helpful for the patients in the near future thanks to rapid developments in science and technology.

Conflicts of interest
The authors have no financial conflicts of interest.

Conclusions

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