Sentence Count = 10 (including title)

21 January 2011

# Researchers reveal function of novel molecule that underlies human deafness

**New research from the University of Sheffield has revealed that a novel molecular mechanism that underlies deafness is caused by a mutation of a specific microRNA called miR-96.**

**The discovery could provide the basis for treating progressive hearing loss and deafness.**

The research team, led by Dr Walter Marcotti, Royal Society University Research Fellow from the University's Department of Biomedical Science, in collaboration with Professor Karen Steel at the Sanger Institute in Cambridge, discovered that the mutation in miR-96 prevents development of the auditory sensory hair cells.

These cells are located in the inner ear and are essential for encoding sound as electrical signals that are then sent to the brain.  
  
The research has been published this week in the Proceedings of the National Academy of Sciences journal and was based on studies of mice, which do not normally hear until about 12 days after birth.

Prior to this age their immature hair cells must execute a precise genetic program that regulates the development of distinct types of sensory hair cell, namely inner and outer hair cells.  
  
The research teams found that in a strain of mice called diminuendo - which carry a single base mutation in the miR-96 gene - hair cell development is arrested around birth.  
  
The study shows that miR-96 normally regulates hair cell development by influencing the expression of many different genes associated with a wide range of developmental processes at a specific stage.

The researchers discovered that the mutation hinders the development not only of the mechanically sensitive hair bundle on the cell apex but also the synaptic structures at the base that govern transfer of electrical information to the sensory nerves.

These new findings suggest that miR-96 is a master regulator responsible for coordinating the development of the sensory cells that are vital to hearing.  
  
Since the mutation in miR-96 is known to cause human deafness and microRNA molecules can be targeted by drugs, the work also raises new opportunities for developing treatments to treat hearing loss.  
  
Dr Walter Marcotti said: "Progressive hearing loss affects a large proportion of the human population, including new born and young children.

Despite the relevance of this problem, very little is currently known regarding the genetic basis of progressive hearing loss.

Our research has provided new and exciting results that further our understanding of auditory development as well as possible molecular targets for the development of future therapies."

**Notes for Editors:** The work was supported by the Royal National Institute for Deaf People (RNID), The Wellcome Trust and the University of Sheffield.  
  
To read the research paper, entitled miR-96 regulates the progression of differentiation in mammalian cochlear inner and outer hair cells, visit the link below.

**For further information please contact: Shemina Davis, Media Relations Officer, on 0114 2225339 or email shemina.davis@sheffield.ac.uk**

* [miR-96 regulates the progression of differentiation in mammalian cochlear inner and outer hair cells](http://www.pnas.org/content/early/2011/01/13/1016646108.abstract)