

Gene therapy restores hearing and balance in Usher syndrome

25 September 2017



The cochlear sensory epithelium (organ of Corti), and the vestibular sensory epithelia, ampullar cristas (inset) of the inner ear injected with the AAV8 virus producing GFP and the SANS protein. The cochlear and vestibular sensory cells were immunostained for GFP and myosin VI and analyzed with a confocal microscope. The green and orange cells produce the protein of the therapeutic gene. Credit: © Institut Pasteur

Scientists from the Institut Pasteur, Inserm, the CNRS, Collège de France, University Pierre et Marie Curie, and University Clermont Auvergne, have recently restored hearing and balance in a mouse model of Usher syndrome type 1G, characterized by profound congenital deafness and vestibular disorders caused by severe dysmorphogenesis of the mechanoelectrical transduction apparatus of the inner ear's sensory cells. These findings open up new possibilities for the development of gene therapy treatments for hereditary forms of deafness.

Hearing loss, sometimes associated with other disorders such as balance defects, is the most common sensory deficit, affecting more than 280

million people worldwide, according to WHO. In France, one child in 700 is born with severe or profound hearing loss, and one in every 1,000 will lose their sense of <u>hearing</u> before adulthood.

Over the past 20 years, scientists have made remarkable progress in deciphering the genetic origins of congenital hereditary hearing loss, which is usually caused by inner ear dysfunction. The inner ear comprises the hearing organ or cochlea, together with the five balance organs (the saccule, utricle and three semicircular canals), which contain the sensory cells, or hair cells, that detect mechanical vibrations and convert them into electrical signals. To date, mutations in more than 100 genes have been associated with inner ear defects, and it is estimated that mutations in more than 100 genes can cause genetic forms of deafness.

The various hereditary forms of hearing loss include Usher syndrome type 1 (USH1), a particularly severe clinical form of deaf-blindness, and specifically the USH1G genetic form. USH1G patients are profoundly deaf and have no balance function at birth, and they subsequently suffer from prepubertal-onset sight loss leading to blindness. USH1G syndrome is due to mutations in the gene encoding the scaffold protein sans, which is essential for the cohesion of the hair bundle of the inner ear hair cells.

Patients with hearing loss and balance dysfunction are currently fitted with auditory prostheses and may be given balance rehabilitation therapy, but the outcomes are variable. One possible alternative for treating such hereditary inner ear defects is gene therapy. This approach entails transferring a healthy (non-mutant) copy of the defective gene to restore the expression of the missing protein. So far, gene therapy attempts have only resulted in partial improvements of hearing in mouse models of specific human deafness forms that did not include severe anomalies in hair cell structure.



Hair bundles



Hair bundles of vestibular sensory cells analyzed using scanning electron microscopy. The image shows a normal hair bundle with its characteristic "staircase" pattern (in yellow), a defective Usher type 1G hair bundle (in pink) and a treated Usher type 1G hair bundle (in green), whose normal/characteristic form was restored with gene therapy. Credit: © Institut Pasteur

In this context, scientists from the Institut Pasteur, Inserm, the CNRS, Collège de France, University Pierre et Marie Curie, and University Clermont Auvergne, have now succeeded in restoring hearing and balance in a mouse model of USH1G syndrome using gene therapy. With a single local injection of the USH1G gene just after birth, the scientists observed a restoration of the structure and mechanosensory function of the inner ear hair bundles - profoundly damaged before birth -, resulting in a long-term partial recovery of hearing, and complete recovery of vestibular function in these mice. These results unexpectedly establish that inner ear defects due to major morphogenetic abnormalities of the hair bundle can be reversed even after birth, with durable efficacy, by gene therapy.

The scientists injected the USH1G gene into the inner ear using the innocuous AAV8 virus, which enabled them to specifically target the hair cells. The expression of the therapeutic gene was detected 48 hours after injection. The team demonstrated that a single injection to restore the

production and localization of the missing protein in hair cells successfully improved hearing and balance functions in the young mice. These findings suggest that the therapeutic protein was able to interact normally with its binding partners among the USH1 molecular complex (the proteins cadherin 23, protocadherin 15, myosin VIIA and harmonin), as required for the mechanoelectrical transduction apparatus of the hair bundle to function correctly.

As Saaïd Safieddine, CNRS Director of Research at the Institut Pasteur and co-senior author of the study with Prof. Christine Petit (head of the Genetics & physiology hearing unit at the Institut Pasteur), says, "We have just shown that it is possible to partially correct a specific form of hereditary <u>hearing loss</u> accompanied by balance problems using local gene therapy performed after the embryogenesis of the ear, which is primarily affected by the mutation responsible for the disorder. This suggests that the time window for effectively treating USH1 syndrome using gene therapy may be larger than initially thought."

This study represents a significant step towards the development of clinical trials in gene therapy for the curative treatment of hereditary deafness and balance loss in humans.

More information: Alice Emptoz et al, Local gene therapy durably restores vestibular function in a mouse model of Usher syndrome type 1G, *Proceedings of the National Academy of Sciences* (2017). DOI: 10.1073/pnas.1708894114

Provided by Institut Pasteur



APA citation: Gene therapy restores hearing and balance in Usher syndrome (2017, September 25) retrieved 23 June 2022 from <u>https://medicalxpress.com/news/2017-09-gene-therapy-usher-syndrome.html</u>

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