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OPEN Corrigendum: Cochlear gene therapy with ancestral AAV in adult mice: complete transduction of inner hair cells without cochlear dysfunction

Jun Suzuki, Ken Hashimoto, Ru Xiao, Luk H. Vandenberghe & M. Charles Liberman

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The authors forgot to cite previous studies relating to in vivo studies in newborn mice. These additional references are listed below as references1 and 2, and should appear in the text as below.

In the Introduction section,

"In order to maximize transduction efficiencies, a "designer" AAV vector, AAV2/Anc80L65, was tested based on promising results in organotypic explant cultures".

should read:

"In order to maximize transduction efficiencies, a "designer" AAV vector, AAV2/Anc80L65, was tested based on promising results in organotypic explant cultures and in vivo studies in newborn mice^{1,2}".

References

1. Landegger, L. D. et al. A Synthetic AAV vector enables safe and efficient gene transfer to the mammalian inner ear. Nat. Biotechnol. 35(3), 280-284, doi: 10.1038/nbt.3781 (2017).

2. Pan, B. et al. Gene therapy restores auditory and vestibular Function in a Mouse Model of Usher Syndrome Type 1c. Nat. Biotechnol. 35(3), 264-272, doi: 10.1038/nbt.3801 (2017).

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