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# A murine model of hnRNPH2-related neurodevelopmental disorder reveals a mechanism for genetic compensation by Hnrnph1

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#### Erratum

Original citation: J Clin Invest. 2023;133(14):e160309. https://doi.org/10.1172/JCI160309 Citation for this erratum: J Clin Invest. 2024;134(8):e181331. https://doi.org/10.1172/JCI181331 During the preparation of this manuscript, a text error was introduced during copyediting by JCI staff. Specifically, the word "mice" was inadvertently introduced into a sentence in the introductory paragraph describing published reports of HNRNPH2 mutations in humans (references 5–7). The correct sentence is below: Although all 6 individuals in the initial report were female, subsequent studies have identified males carrying missense mutations in HNRNPH2 associated with a range of overlapping phenotypes (5–7). The JCI regrets the error.



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