

Rare *EIF4A2* variants are associated with a neurodevelopmental disorder characterized by intellectual disability, hypotonia, and epilepsy

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(The American Journal of Human Genetics 110, 120–145; January 5, 2023)

In the originally published version of this article, the stock number of two fly lines, UAS-eIF4A and Nubbin-GAL4, are incorrectly mentioned. The correct stock number for UAS-eIF4A is FlyORF:F000979 and for Nubbin-GAL4 is BDSC #86108. This has now been fixed online. The authors regret this inadvertent error.

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<https://doi.org/10.1016/j.ajhg.2023.02.010>.

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