De Novo Mutations in Synaptic Transmission Genes Including DNM1 Cause Epileptic Encephalopathies

EuroEPINOMICS-RES Consortium,* Epilepsy Phenome/Genome Project, and Epi4K Consortium

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In the list of consortium members for the Epilepsy Phenome/Genome Project, member Dina Amrom’s name was misspelled as Amron. The authors regret the error.

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De Novo Truncating Variants in ASXL2 Are Associated with a Unique and Recognizable Clinical Phenotype


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In the originally published version of this article, Table 1 unfortunately contained two errors: c.2472delC (p.Ser825Valfs*18) should have been c.2472delC (p.Ser825Valfs*16) (as correctly notated in the main text), and the title should have mentioned six individuals instead of four. The errors have been corrected online, and the authors apologize for any confusion.

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