Emily Fassi

List of Publications by Year in descending order

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933447 1281871 11 831 10 11 citations h-index g-index papers 12 12 12 2354 all docs docs citations times ranked citing authors

#	Article	IF	CITATIONS
1	Breast cancer polygenic risk scores in the clinical cancer genetic counseling setting: Current practices and impact on patient management. Journal of Genetic Counseling, 2021, 30, 588-597.	1.6	14
2	Widening of the genetic and clinical spectrum of Lamb–Shaffer syndrome, a neurodevelopmental disorder due to SOX5 haploinsufficiency. Genetics in Medicine, 2020, 22, 524-537.	2.4	21
3	Missense variants in $\langle i \rangle$ TAF1 $\langle j \rangle$ and developmental phenotypes: Challenges of determining pathogenicity. Human Mutation, 2020, 41, 449-464.	2.5	17
4	AMPA receptor GluA2 subunit defects are a cause of neurodevelopmental disorders. Nature Communications, 2019, 10, 3094.	12.8	150
5	CTCF variants in 39 individuals with a variable neurodevelopmental disorder broaden the mutational and clinical spectrum. Genetics in Medicine, 2019, 21, 2723-2733.	2.4	48
6	Mutations in ACTL6B Cause Neurodevelopmental Deficits and Epilepsy and Lead to Loss of Dendrites in Human Neurons. American Journal of Human Genetics, 2019, 104, 815-834.	6.2	59
7	Mutations in the $\langle scp \rangle PH \langle scp \rangle Domain of \langle i \rangle \langle scp \rangle DNM \langle scp \rangle 1 \langle i \rangle are associated with a nonepileptic phenotype characterized by developmental delay and neurobehavioral abnormalities. Molecular Genetics & amp; Genomic Medicine, 2018, 6, 294-300.$	1.2	19
8	De novo mutation screening in childhood-onset cerebellar atrophy identifies gain-of-function mutations in the CACNA1G calcium channel gene. Brain, 2018, 141, 1998-2013.	7.6	67
9	The Exome Clinic and the role of medical genetics expertise in the interpretation of exome sequencing results. Genetics in Medicine, 2017, 19, 1040-1048.	2.4	85
10	High Rate of Recurrent De Novo Mutations in Developmental and Epileptic Encephalopathies. American Journal of Human Genetics, 2017, 101, 664-685.	6.2	337
11	Genetic counseling in CHARGE syndrome: Diagnostic evaluation through follow up. American Journal of Medical Genetics, Part C: Seminars in Medical Genetics, 2017, 175, 407-416.	1.6	13