Engy Asem Ashaat

List of Publications by Year in Descending Order

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The third column is the impact factor (IF) of the journal, and the fourth column is the number of citations of the article.

12
papers62
citations4
h-index7
g-index14
ext. papers103
ext. citations2.2
avg, IF0.92
L-index

#	Paper	IF	Citations
12	Chromosome 9p terminal deletion in nine Egyptian patients and narrowing of the critical region for trigonocephaly. <i>Molecular Genetics & Enomic Medicine</i> , 2021 , 9, e1829	2.3	2
11	Clinical and genetic characterization of ten Egyptian patients with Wolf-Hirschhorn syndrome and review of literature. <i>Molecular Genetics & Egyptian Patients Wolf-Hirschhorn Syndrome and Review of literature.</i>	2.3	1
10	First Report of Two Egyptian Patients with Desbuquois Dysplasia due to Homozygous Mutations. <i>Molecular Syndromology</i> , 2021 , 12, 279-288	1.5	О
9	Assessment of Multiplex Ligation-Dependent Probe Amplification (MLPA) as a diagnostic test for Egyptian patients with Williams-Beuren syndrome. <i>Gene Reports</i> , 2020 , 20, 100767	1.4	
8	A novel homozygous variant in the TRAPPC9 gene causing intellectual disability and autism Spectrum disorder. <i>Meta Gene</i> , 2020 , 26, 100783	0.7	1
7	Turner syndrome in diverse populations. American Journal of Medical Genetics, Part A, 2020, 182, 303-37	13 .5	8
6	Phenotypic and Molecular Cytogenetic Analysis of a Case of Monosomy 1p36 Syndrome due to Unbalanced Translocation. <i>Molecular Syndromology</i> , 2020 , 11, 284-295	1.5	
5	Rubinstein-Taybi syndrome in diverse populations. <i>American Journal of Medical Genetics, Part A</i> , 2020 , 182, 2939-2950	2.5	4
4	Blepharophimosis-ptosis-intellectual disability syndrome: A report of nine Egyptian patients with further expansion of phenotypic and mutational spectrum. <i>American Journal of Medical Genetics, Part A</i> , 2020 , 182, 2857-2866	2.5	1
3	The potential impact of COMT gene variants on dopamine regulation and phenotypic traits of ASD patients. <i>Behavioural Brain Research</i> , 2020 , 378, 112272	3.4	7
2	Study of C677T variant of methylene tetrahydrofolate reductase gene in autistic spectrum disorder Egyptian children. <i>American Journal of Medical Genetics Part B: Neuropsychiatric Genetics</i> , 2019 , 180, 305	5-³₹ 0 9	5
1	Williams-Beuren syndrome in diverse populations. <i>American Journal of Medical Genetics, Part A</i> , 2018 , 176, 1128-1136	2.5	31