

Purvi Majethia

List of Publications by Year in descending order

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Version: 2024-02-01

8
papers

34
citations

2257833

3
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2053595

5
g-index

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all docs

8
docs citations

8
times ranked

22
citing authors

#	ARTICLE	IF	CITATIONS
1	Further evidence of affected females with a heterozygous variant in FGF13 causing X-linked developmental and epileptic encephalopathy 90. <i>European Journal of Medical Genetics</i> , 2022, 65, 104403.	0.7	4
2	Second report of SHMT2 related neurodevelopmental disorder with cardiomyopathy, spasticity, and brain abnormalities. <i>European Journal of Medical Genetics</i> , 2022, 65, 104481.	0.7	2
3	Further evidence of muscle involvement in neurodevelopmental disorder with epilepsy, spasticity, and brain atrophy. <i>Annals of Human Genetics</i> , 2022, 86, 94-101.	0.3	3
4	Wiedemannâ€“Rautenstrauch syndrome in an Indian patient with biallelic pathogenic variants in POLR3A. <i>American Journal of Medical Genetics, Part A</i> , 2021, 185, 1602-1605.	0.7	3
5	Biallelic start loss variant, c.<sc>1A</sc>â€“G in <sc><i>GCSH</i></sc> is associated with variant nonketotic hyperglycinemia. <i>Clinical Genetics</i> , 2021, 100, 201-205.	1.0	4
6	Multilocus disease-causing genomic variations for Mendelian disorders: role of systematic phenotyping and implications on genetic counselling. <i>European Journal of Human Genetics</i> , 2021, 29, 1774-1780.	1.4	7
7	NAD(P)HX dehydratase (NAXD) deficiency due to a novel biallelic missense variant and review of literature. <i>European Journal of Medical Genetics</i> , 2021, 64, 104266.	0.7	6
8	Expanding the electro-clinical phenotype of CARS2-associated neuroregression. <i>Epilepsy and Behavior Reports</i> , 2021, 16, 100485.	0.5	5