

Smrithi Salian

List of Publications by Year in descending order

Source: <https://exaly.com/author-pdf/9174725/publications.pdf>

Version: 2024-02-01

14
papers

96
citations

1478505

6
h-index

1474206

9
g-index

14
all docs

14
docs citations

14
times ranked

172
citing authors

#	ARTICLE	IF	CITATIONS
1	PIGF deficiency causes a phenotype overlapping with DOORS syndrome. <i>Human Genetics</i> , 2021, 140, 879-884.	3.8	2
2	Epileptic encephalopathy caused by <sc>ARV1</sc> deficiency: Refinement of the genotypeâ€“phenotype spectrum and functional impact on <sc>GPI</sc>â€“anchored proteins. <i>Clinical Genetics</i> , 2021, 100, 607-614.	2.0	6
3	A variant of neonatal progeroid syndrome, or Wiedemannâ€“Rautenstrauch syndrome, is associated with a nonsense variant in POLR3GL. <i>European Journal of Human Genetics</i> , 2020, 28, 461-468.	2.8	16
4	De Novo KAT5 Variants Cause a Syndrome with Recognizable Facial Dysmorphisms, Cerebellar Atrophy, Sleep Disturbance, and Epilepsy. <i>American Journal of Human Genetics</i> , 2020, 107, 564-574.	6.2	14
5	Further evidence for causation of ischiopspinal dysostosis by a pathogenic variant in <i>BMPER</i> and expansion of the phenotype. <i>Congenital Anomalies (discontinued)</i> , 2019, 59, 26-27.	0.6	3
6	Seven additional families with spondylocarpotarsal synostosis syndrome with novel biallelic deleterious variants in <i>FLNB</i>. <i>Clinical Genetics</i> , 2018, 94, 159-164.	2.0	10
7	Pycnodysostosis: Novel Variants in CTSK and Occurrence of Giant Cell Tumor. <i>Journal of Pediatric Genetics</i> , 2018, 07, 009-013.	0.7	6
8	Additional three patients with Smithâ€“McCort dysplasia due to novel <i>RAB33B</i> mutations. <i>American Journal of Medical Genetics, Part A</i> , 2017, 173, 588-595.	1.2	17
9	Cover Image, Volume 173A, Number 3, March 2017. <i>American Journal of Medical Genetics, Part A</i> , 2017, 173, i.	1.2	0
10	Severe Form of Brachydactyly Type A1 in a Child with a c.298Gâ€“>â€“A Mutation in IHH Gene. <i>Journal of Pediatric Genetics</i> , 2017, 06, 177-180.	0.7	5
11	Variable presentation of Fraser syndrome in two fetuses and a novel mutation in <i>FRAS1</i>. <i>Congenital Anomalies (discontinued)</i> , 2017, 57, 83-85.	0.6	8
12	Focal dermal hypoplasia with a de novo mutation p.e300FNx01 of porcn gene in a male infant. <i>Indian Journal of Dermatology</i> , 2016, 61, 700.	0.3	3
13	Novel ALOX12B mutation identified in parents following single nucleotide polymorphism microarray testing of banked DNA from a fatal case of congenital ichthyosis. <i>Indian Journal of Dermatology</i> , 2016, 61, 122.	0.3	0
14	Familial 7q11.23 duplication with variable phenotype. <i>American Journal of Medical Genetics, Part A</i> , 2015, 167, 2727-2730.	1.2	6