Aravindhan Veerapandiyan Mbbs

List of Publications by Year in descending order

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34 papers 456 citations

840585 11 h-index 713332 21 g-index

34 all docs

34 docs citations

times ranked

34

747 citing authors

#	Article	IF	CITATIONS
1	Subacute Liver Failure Following Gene Replacement Therapy for Spinal Muscular Atrophy Type 1. Journal of Pediatrics, 2020, 225, 252-258.e1.	0.9	79
2	The care of patients with Duchenne, Becker, and other muscular dystrophies in the <scp>COVID</scp> â€19 pandemic. Muscle and Nerve, 2020, 62, 41-45.	1.0	54
3	Nusinersen for older patients with spinal muscular atrophy: A realâ€world clinical setting experience. Muscle and Nerve, 2020, 61, 222-226.	1.0	51
4	Combination molecular therapies for type 1 spinal muscular atrophy. Muscle and Nerve, 2020, 62, 550-554.	1.0	51
5	Spectrum of COVIDâ€19 in children. Acta Paediatrica, International Journal of Paediatrics, 2020, 109, 1899-1900.	0.7	33
6	Spinal muscular atrophy care in the COVIDâ€19 pandemic era. Muscle and Nerve, 2020, 62, 46-49.	1.0	31
7	Cervical puncture to deliver nusinersen in patients with spinal muscular atrophy. Neurology, 2018, 91, e620-e624.	1.5	27
8	Pseudometabolic presentation of dystrophinopathy due to a missense mutation. Muscle and Nerve, 2010, 42, 975-979.	1.0	25
9	An expanded access program of risdiplam for patients with Type 1 or 2 spinal muscular atrophy. Annals of Clinical and Translational Neurology, 2022, 9, 810-818.	1.7	18
10	Implementing Pharmacogenomics Testing: Single Center Experience at Arkansas Children's Hospital. Journal of Personalized Medicine, 2021, 11, 394.	1.1	14
11	Child Neurology: Type 1 sialidosis due to a novel mutation in <i>NEU1</i> gene. Neurology, 2018, 90, 622-624.	1.5	11
12	Early-onset epileptic encephalopathy with myoclonic seizures related to 9q33.3-q34.11 deletion involving STXBP1 and SPTAN1 genes. Epileptic Disorders, 2018, 20, 214-218.	0.7	11
13	Molecular Dysregulation in Autism Spectrum Disorder. Journal of Personalized Medicine, 2021, 11, 848.	1.1	8
14	Semaphorin-Plexin Signaling: From Axonal Guidance to a New X-Linked Intellectual Disability Syndrome. Pediatric Neurology, 2022, 126, 65-73.	1.0	8
15	Variable penetrance of Andersen-Tawil syndrome in a family with a rare missense <i>KCNJ2</i> mutation. Neurology: Genetics, 2018, 4, e284.	0.9	6
16	Use of Head Computed Tomography (CT) in the Pediatric Emergency Department in Evaluation of Children With New-Onset Afebrile Seizure. Journal of Child Neurology, 2018, 33, 708-712.	0.7	6
17	COVID-19 in Pediatric Inpatients: A Multi-Center Observational Study of Factors Associated with Negative Short-Term Outcomes. Children, 2021, 8, 951.	0.6	4
18	Electronic health record cue identifies epilepsy patients at risk for obstructive sleep apnea. Neurology: Clinical Practice, 2018, 8, 468-471.	0.8	3

#	Article	IF	CITATIONS
19	BAG3 Myopathy Presenting With Prominent Neuropathic Phenotype and No Cardiac or Respiratory Involvement: A Case Report and Literature Review. Journal of Clinical Neuromuscular Disease, 2020, 21, 230-239.	0.3	3
20	Distal 1q21.1 and proximal 1q21.2 microduplication in a child with attention-deficit hyperactivity disorder. Acta Neurologica Belgica, 2019, 119, 289-290.	0.5	2
21	Dystrophinopathy in a Family Due to a Rare Nonsense Mutation Causing Predominant Behavioral Phenotype. Journal of Pediatric Neurology, 2020, 18, 210-213.	0.0	2
22	Pseudometabolic Presentation of Dystrophinopathy in a Family Due to a Rare Nonsense Mutation. Journal of Clinical Neuromuscular Disease, 2020, 21, 245-246.	0.3	2
23	A novel intronic homozygous mutation in the AMT gene of a patient with nonketotic hyperglycinemia and hyperammonemia. Metabolic Brain Disease, 2019, 34, 373-376.	1.4	1
24	A novel noncoding FKRP mutation in early onset limb-girdle muscular dystrophy. Neurology: Genetics, 2020, 6, e388.	0.9	1
25	Clinical Reasoning: A case of bilateral foot drop in a 74-year-old man. Neurology, 2020, 94, 405-409.	1.5	1
26	Exercise Intolerance and Rhabdomyolysis Due to Dystrophinopathy: A Pseudometabolic Presentation. Journal of Pediatric Neurology, 2022, 20, 080-082.	0.0	1
27	Congenital Myasthenic Syndrome due to a Novel Mutation in CHAT Gene. Journal of Clinical Neuromuscular Disease, 2021, 23, 54-55.	0.3	1
28	Epilepsy in hereditary spastic paraplegia associated with NIPA1 gene. Journal of Clinical Neuroscience, 2022, 100, 212-213.	0.8	1
29	Palliative Care in Duchenne Muscular Dystrophy: Goals of Care Discussions and Beyond. Muscle and Nerve, 2022, , .	1.0	1
30	Clinical Reasoning: A 6 year old boy with muscle twitching. Neurology, 2020, 96, 10.1212/WNL.00000000010746.	1.5	0
31	A Toddler With Bilateral Facial Weakness. Clinical Pediatrics, 2020, 59, 529-531.	0.4	O
32	Recovery of foot drop in chronic inflammatory demyelinating polyneuropathy (CIDP). Muscle and Nerve, 2021, 64, 59-63.	1.0	0
33	Limb-Girdle Muscular Dystrophy R9 due to a Novel Complex Insertion/Duplication Variant in <i>FKRP</i> Gene. Child Neurology Open, 2022, 9, 2329048X2210975.	0.5	0
34	Infantile-Onset Complex Hereditary Spastic Paraplegia Due to a Novel Mutation in SPAST Gene. Pediatric Neurology, 2022, 134, 71.	1.0	0