Vandana A Gupta

List of Publications by Year in descending order

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361045 414034 32 1,426 20 32 citations h-index g-index papers 33 33 33 2568 docs citations times ranked citing authors all docs

#	Article	IF	CITATIONS
1	246th ENMC International Workshop: Protein aggregate myopathies 24–26 May 2019, Hoofddorp, The Netherlands. Neuromuscular Disorders, 2021, 31, 158-166.	0.3	5
2	<scp>hnRNP L /scp> is essential for myogenic differentiation and modulates myotonic dystrophy pathologies. Muscle and Nerve, 2021, 63, 928-940.</scp>	1.0	7
3	Prednisolone rescues Duchenne muscular dystrophy phenotypes in human pluripotent stem cell–derived skeletal muscle in vitro. Proceedings of the National Academy of Sciences of the United States of America, 2021, 118, .	3.3	32
4	Biâ€allelic loss of function variants in <scp>GOLGA2</scp> are associated with a complex neurological phenotype: Report of a second family. Clinical Genetics, 2021, 100, 748-751.	1.0	8
5	Novel Recessive <i>TNNT1</i> Congenital Coreâ€Rod Myopathy in French Canadians. Annals of Neurology, 2020, 87, 568-583.	2.8	19
6	ACTN2 mutations cause "Multiple structured Core Disease―(MsCD). Acta Neuropathologica, 2019, 137, 501-519.	3.9	31
7	Homozygous <i>TRPV4</i> mutation causes congenital distal spinal muscular atrophy and arthrogryposis. Neurology: Genetics, 2019, 5, e312.	0.9	15
8	Dysregulation of NRAP degradation by KLHL41 contributes to pathophysiology in nemaline myopathy. Human Molecular Genetics, 2019, 28, 2549-2560.	1.4	22
9	Transgenic zebrafish model of DUX4 misexpression reveals a developmental role in FSHD pathogenesis. Human Molecular Genetics, 2019, 28, 320-331.	1.4	14
10	An open source microcontroller based flume for evaluating swimming performance of larval, juvenile, and adult zebrafish. PLoS ONE, 2018, 13, e0199712.	1.1	13
11	An integrated clinical program and crowdsourcing strategy for genomic sequencing and Mendelian disease gene discovery. Npj Genomic Medicine, 2018, 3, 21.	1.7	24
12	RNA helicase, DDX27 regulates skeletal muscle growth and regeneration by modulation of translational processes. PLoS Genetics, 2018, 14, e1007226.	1.5	34
13	A novel early onset phenotype in a zebrafish model of merosin deficient congenital muscular dystrophy. PLoS ONE, 2017, 12, e0172648.	1.1	9
14	Mutationâ€specific effects on thin filament length in thin filament myopathy. Annals of Neurology, 2016, 79, 959-969.	2.8	54
15	GOLGA2, encoding a master regulator of golgi apparatus, is mutated in a patient with a neuromuscular disorder. Human Genetics, 2016, 135, 245-251.	1.8	38
16	Mutation of the mitochondrial carrier SLC25A42 causes a novel form of mitochondrial myopathy in humans. Human Genetics, 2016, 135, 21-30.	1.8	34
17	Bridging integrator 1 (Bin1) deficiency in zebrafish results in centronuclear myopathy. Human Molecular Genetics, 2014, 23, 3566-3578.	1.4	28
18	POMK mutations disrupt muscle development leading to a spectrum of neuromuscular presentations. Human Molecular Genetics, 2014, 23, 5781-5792.	1.4	72

#	Article	IF	CITATIONS
19	ZBTB42 mutation defines a novel lethal congenital contracture syndrome (LCCS6). Human Molecular Genetics, 2014, 23, 6584-6593.	1.4	25
20	Kelch proteins: emerging roles in skeletal muscle development and diseases. Skeletal Muscle, 2014, 4, 11.	1.9	119
21	Leiomodin-3 dysfunction results in thin filament disorganization and nemaline myopathy. Journal of Clinical Investigation, 2014, 124, 4693-4708.	3.9	153
22	Identification of KLHL41 Mutations Implicates BTB-Kelch-Mediated Ubiquitination as an Alternate Pathway to Myofibrillar Disruption in Nemaline Myopathy. American Journal of Human Genetics, 2013, 93, 1108-1117.	2.6	147
23	Loss of Catalytically Inactive Lipid Phosphatase Myotubularin-related Protein 12 Impairs Myotubularin Stability and Promotes Centronuclear Myopathy in Zebrafish. PLoS Genetics, 2013, 9, e1003583.	1.5	22
24	A Network of Transcription Factors Operates during Early Tooth Morphogenesis. Molecular and Cellular Biology, 2013, 33, 3099-3112.	1.1	19
25	αâ€Actininâ€2 deficiency results in sarcomeric defects in zebrafish that cannot be rescued by αâ€actininâ€3 revealing functional differences between sarcomeric isoforms. FASEB Journal, 2012, 26, 1892-1908.	0.2	34
26	Myotubularin-Deficient Myoblasts Display Increased Apoptosis, Delayed Proliferation, and Poor Cell Engraftment. American Journal of Pathology, 2012, 181, 961-968.	1.9	37
27	Exome Sequencing and Functional Validation in Zebrafish Identify GTDC2 Mutations as a Cause of Walker-Warburg Syndrome. American Journal of Human Genetics, 2012, 91, 541-547.	2.6	167
28	A Splice Site Mutation in Laminin- $\hat{l}\pm 2$ Results in a Severe Muscular Dystrophy and Growth Abnormalities in Zebrafish. PLoS ONE, 2012, 7, e43794.	1.1	48
29	The zebrafish dag1 mutant: a novel genetic model for dystroglycanopathies. Human Molecular Genetics, 2011, 20, 1712-1725.	1.4	101
30	Mullerian-Inhibiting Substance Induces Gro-l ² Expression in Breast Cancer Cells through a Nuclear Factor-l ² B–Dependent and Smad1-Dependent Mechanism. Cancer Research, 2007, 67, 2747-2756.	0.4	12
31	Modification of Msx1 by SUMO-1. Biochemical and Biophysical Research Communications, 2006, 345, 74-77.	1.0	25
32	Role of water structure on phase separation in polyelectrolyte–polyethyleneglycol based aqueous two-phase systems. Polymer, 2002, 43, 3387-3390.	1.8	58