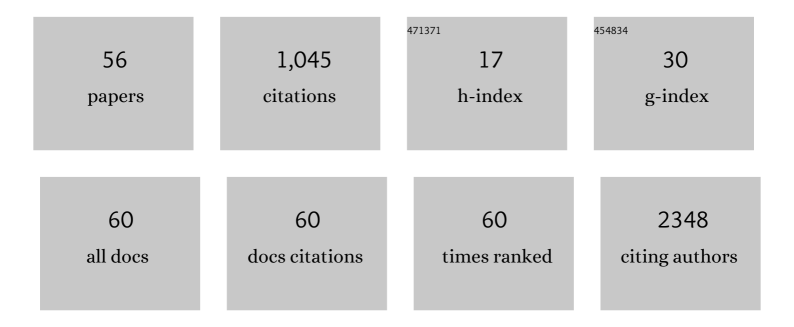
Jodi Warman Chardon

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/2484409/publications.pdf

Version: 2024-02-01



#	Article	IF	CITATIONS
1	Laryngospasm in amyotrophic lateral sclerosis. Muscle and Nerve, 2022, 65, 400-404.	1.0	6
2	Temporal evolution of nerve conduction study abnormalities in antiâ€myelinâ€associated glycoprotein neuropathy. Muscle and Nerve, 2021, 63, 401-404.	1.0	10
3	Autologous Hematopoietic Stem Cell Transplantation for Chronic Inflammatory Demyelinating Polyradiculoneuropathy. Canadian Journal of Neurological Sciences, 2021, , 1-7.	0.3	3
4	Whole genome sequencing reveals biallelic <scp><i>PLA2G6</i></scp> mutations in siblings with cerebellar atrophy and cap myopathy. Clinical Genetics, 2021, 99, 746-748.	1.0	3
5	A Canadian Adult Spinal Muscular Atrophy Outcome Measures Toolkit: Results of a National Consensus using a Modified Delphi Method. Journal of Neuromuscular Diseases, 2021, 8, 579-588.	1.1	7
6	Pseudohypertrophy of the extensor digitorum brevis in diabetic polyneuropathy. Muscle and Nerve, 2021, 64, E20-E22.	1.0	0
7	MuSK not MNGIE: Atypical MuSK-antibody myasthenia presenting as a genetic disorder. Neuromuscular Disorders, 2021, , .	0.3	0
8	Intrafamilial variability of limb-girdle muscular dystrophy, LGMD1D type. European Journal of Medical Genetics, 2020, 63, 103655.	0.7	10
9	Impact of disuse muscular atrophy on the compound muscle action potential. Muscle and Nerve, 2020, 61, 58-62.	1.0	5
10	247th ENMC International Workshop: Muscle magnetic resonance imaging - Implementing muscle MRI as a diagnostic tool for rare genetic myopathy cohorts. Hoofddorp, The Netherlands, September 2019. Neuromuscular Disorders, 2020, 30, 938-947.	0.3	11
11	Myofibrillar Myopathy Mimicking Polyneuropathy. Case Reports in Neurology, 2020, 12, 97-102.	0.3	1
12	Advances in the diagnosis of inherited neuromuscular diseases and implications for therapy development. Lancet Neurology, The, 2020, 19, 522-532.	4.9	36
13	A National Spinal Muscular Atrophy Registry for Real-World Evidence. Canadian Journal of Neurological Sciences, 2020, 47, 810-815.	0.3	6
14	Novel Recessive <i>TNNT1</i> Congenital Coreâ€Rod Myopathy in French Canadians. Annals of Neurology, 2020, 87, 568-583.	2.8	19
15	Abnormal fatty acid metabolism is a core component of spinal muscular atrophy. Annals of Clinical and Translational Neurology, 2019, 6, 1519-1532.	1.7	72
16	Intermittent undulating tongue as an involuntary movement in early amyotrophic lateral sclerosis. Parkinsonism and Related Disorders, 2019, 67, 1-2.	1.1	1
17	A Survey of Cerebrospinal Fluid Total Protein Upper Limits in Canada: Time for an Update?. Canadian Journal of Neurological Sciences, 2019, 46, 283-286.	0.3	2
18	The Added Value of Cardiac Magnetic Resonance in Muscular Dystrophies. Journal of Neuromuscular Diseases, 2019, 6, 389-399.	1.1	10

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19	MYO-MRI diagnostic protocols in genetic myopathies. Neuromuscular Disorders, 2019, 29, 827-841.	0.3	46
20	Age matters. Neurology: Neuroimmunology and NeuroInflammation, 2019, 6, e576.	3.1	7
21	<i>PDXK</i> mutations cause polyneuropathy responsive to pyridoxal 5′â€phosphate supplementation. Annals of Neurology, 2019, 86, 225-240.	2.8	54
22	Dataset for worldwide survey of cerebrospinal total protein upper reference values. Data in Brief, 2019, 23, 103760.	0.5	0
23	Neurolymphomatosis of the lumbosacral plexus and its branches: case series and literature review. BMC Cancer, 2019, 19, 1149.	1.1	16
24	Distal Cervical Spondylotic Amyotrophy: Case Reports Demonstrating Clinical/Imaging Segmental Discrepancy. Journal of Clinical Neuromuscular Disease, 2019, 21, 107-111.	0.3	2
25	Targeted exome analysis identifies the genetic basis of disease in over 50% of patients with a wide range of ataxia-related phenotypes. Genetics in Medicine, 2019, 21, 195-206.	1.1	65
26	Neurolymphomatosis of the Brachial Plexus and its Branches: Case Series and Literature Review. Canadian Journal of Neurological Sciences, 2018, 45, 137-143.	0.3	26
27	Novel <i>ELOVL4</i> mutation associated with erythrokeratodermia and spinocerebellar ataxia (SCA) Tj ETQq1	1 0,7,8431	14 rgBT /Overl
28	Teaching Video Neurolmages: Rippling muscle disease with caveolin myopathy. Neurology, 2018, 91, e1726-e1727.	1.5	1
29	Intraneural Ganglion Cysts of the Fibular Nerve: A Cause of Fluctuating Painful Foot Drop. Canadian Journal of Neurological Sciences, 2018, 45, 601-603.	0.3	Ο
30	Whole-transcriptome sequencing in blood provides a diagnosis of spinal muscular atrophy with progressive myoclonic epilepsy. Human Mutation, 2017, 38, 611-614.	1.1	25
31	Cardiac Amyloidosis Phenotype Associated With a Glu89Lys Transthyretin Mutation. Canadian Journal of Cardiology, 2017, 33, 830.e5-830.e7.	0.8	2
32	Clinical and genetic study of hereditary spastic paraplegia in Canada. Neurology: Genetics, 2017, 3, e122.	0.9	82
33	Survival Motor Neuron Protein is Released from Cells in Exosomes: A Potential Biomarker for Spinal Muscular Atrophy. Scientific Reports, 2017, 7, 13859.	1.6	13
34	Teaching Video Neuro <i>Images</i> : Trapezius muscle hypertrophy in multifocal motor neuropathy. Neurology, 2017, 89, e81-e82.	1.5	0
35	The Role of Muscle Imaging in the Diagnosis and Assessment of Children with Genetic Muscle Disease. Neuropediatrics, 2017, 48, 233-241.	0.3	11
36	Combined isolated trigeminal and facial neuropathies from perineural invasion by squamous cell carcinoma: A case series and review of the literature. Journal of Clinical Neuroscience, 2017, 35, 5-12.	0.8	7

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37	Autosomal dominant cerebellar ataxia, deafness, and narcolepsy (ADCA-DN) associated with progressive cognitive and behavioral deterioration Neuropsychology, 2017, 31, 292-303.	1.0	4
38	A crucial first randomized controlled trial of thymectomy in non-thymomatous myasthenia gravis. Journal of Thoracic Disease, 2016, 8, E1375-E1378.	0.6	2
39	Supramaximal Stimulus Intensity as a Diagnostic Tool in Chronic Demyelinating Neuropathy. Neuroscience Journal, 2016, 2016, 1-5.	2.3	5
40	Diffuse leukoencephalopathy with spheroids presenting as primary progressive aphasia. Neurology, 2016, 86, 1464-1465.	1.5	3
41	Identification of a methylation profile for DNMT1-associated autosomal dominant cerebellar ataxia, deafness, and narcolepsy. Clinical Epigenetics, 2016, 8, 91.	1.8	66
42	Marked enlargement of neck circumference from nerve hypertrophy in CIDP. Neurology, 2016, 87, 442-442.	1.5	1
43	Systematic analysis of clinical deficits in unilateral hypoglossal nerve palsy. Muscle and Nerve, 2016, 54, 1055-1058.	1.0	0
44	Syndrome disintegration: Exome sequencing reveals that Fitzsimmons syndrome is a coâ€occurrence of multiple events. American Journal of Medical Genetics, Part A, 2016, 170, 1820-1825.	0.7	19
45	Amyloid Neuropathy Following Domino Liver Transplantation. JAMA Neurology, 2016, 73, 477.	4.5	4
46	Very lateâ€onset Sandhoff disease presenting as Kennedy Disease. Muscle and Nerve, 2015, 52, 1135-1136.	1.0	4
47	LIMS2 mutations are associated with a novel muscular dystrophy, severe cardiomyopathy and triangular tongues. Clinical Genetics, 2015, 88, 558-564.	1.0	23
48	Autoimmune peripheral neuropathies. Clinica Chimica Acta, 2015, 449, 37-42.	0.5	26
49	Axons to Exons: the Molecular Diagnosis of Rare Neurological Diseases by Next-Generation Sequencing. Current Neurology and Neuroscience Reports, 2015, 15, 64.	2.0	29
50	Exome Sequencing as a Diagnostic Tool for Pediatricâ€Onset Ataxia. Human Mutation, 2014, 35, 45-49.	1.1	91
51	The utility of exome sequencing for genetic diagnosis in a familial microcephaly epilepsy syndrome. BMC Neurology, 2014, 14, 22.	0.8	18
52	Deletion of <i>AFG3L2</i> associated with spinocerebellar ataxia type 28 in the context of multiple genomic anomalies. American Journal of Medical Genetics, Part A, 2014, 164, 3209-3212.	0.7	11
53	Recent Advances in the Genetic Etiology of Brain Malformations. Current Neurology and Neuroscience Reports, 2013, 13, 364.	2.0	20
54	Missense mutations in ITPR1 cause autosomal dominant congenital nonprogressive spinocerebellar ataxia. Orphanet Journal of Rare Diseases, 2012, 7, 67.	1.2	124

#	Article	IF	CITATIONS
55	Deletion of filamin A in two female patients with periventricular nodular heterotopia. American Journal of Medical Genetics, Part A, 2012, 158A, 1512-1516.	0.7	6
56	Does Diabetes Alter CSF Total Protein Levels? A Retrospective Cohort Study. Neurohospitalist, The, 0, , 194187442110393.	0.3	0