## Viorica Chelban

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/5180317/publications.pdf

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42 papers

2,756 citations

20 h-index 276875 41 g-index

42 all docs

42 docs citations

42 times ranked 4714 citing authors

#	Article	IF	CITATIONS
1	Combining biomarkers for prognostic modelling of Parkinson's disease. Journal of Neurology, Neurosurgery and Psychiatry, 2022, 93, 707-715.	1.9	9
2	Low Prevalence of NOTCH2NLC GGC Repeat Expansion in White Patients with Movement Disorders. Movement Disorders, 2021, 36, 251-255.	3.9	23
3	Expanding the Spectrum of <scp><i>AP5Z1â€</i></scp> Related Hereditary Spastic Paraplegia ( <scp>HSPâ€6PG48</scp> ): A Multicenter Study on a Rare Disease. Movement Disorders, 2021, 36, 1034-1038.	3.9	9
4	<i>RFC1</i> -related ataxia is a mimic of early multiple system atrophy. Journal of Neurology, Neurosurgery and Psychiatry, 2021, 92, 444-446.	1.9	25
5	Mitochondrial <scp>D</scp> <scp>NA</scp> Analysis from Exome Sequencing Data Improves Diagnostic Yield in Neurological Diseases. Annals of Neurology, 2021, 89, 1240-1247.	5.3	12
6	Identification of Candidate Parkinson Disease Genes by Integrating Genome-Wide Association Study, Expression, and Epigenetic Data Sets. JAMA Neurology, 2021, 78, 464.	9.0	95
7	Investigation of Autosomal Genetic Sex Differences in Parkinson's Disease. Annals of Neurology, 2021, 90, 35-42.	5.3	29
8	Spastic paraplegia preceding PSEN1 â€related familial Alzheimer's disease. Alzheimer's and Dementia: Diagnosis, Assessment and Disease Monitoring, 2021, 13, e12186.	2.4	7
9	Drug-resistant epilepsy: modern concepts, integrative mechanisms, and therapeutic advances. The Moldovan Medical Journal, 2021, 64, 72-85.	0.1	0
10	Genetic and phenotypic characterization of NKX6â€2 â€related spastic ataxia and hypomyelination. European Journal of Neurology, 2020, 27, 334-342.	3.3	16
11	Diagnosis Across the Spectrum of Progressive Supranuclear Palsy and Corticobasal Syndrome. JAMA Neurology, 2020, 77, 377.	9.0	94
12	An update on MSA: premotor and non-motor features open a window of opportunities for early diagnosis and intervention. Journal of Neurology, 2020, 267, 2754-2770.	3.6	25
13	Genetic and Clinical Heterogeneity in Thirteen New Cases with Aceruloplasminemia. Atypical Anemia as a Clue for an Early Diagnosis. International Journal of Molecular Sciences, 2020, 21, 2374.	4.1	25
14	Hereditary polyneuropathy with optic atrophy due to PDXK variant leading to impaired Vitamin B6 metabolism. Neuromuscular Disorders, 2020, 30, 583-589.	0.6	7
15	<i>MYORG</i> -related disease is associated with central pontine calcifications and atypical parkinsonism. Neurology: Genetics, 2020, 6, e399.	1.9	13
16	Bi-allelic JAM2 Variants Lead to Early-Onset Recessive Primary Familial Brain Calcification. American Journal of Human Genetics, 2020, 106, 412-421.	6.2	47
17	Penetrance of Parkinson's Disease in <i>LRRK2</i> p.G2019S Carriers Is Modified by a Polygenic Risk Score. Movement Disorders, 2020, 35, 774-780.	3.9	57
18	<scp><i>RFC1</i></scp> Intronic Repeat Expansions Absent in Pathologically Confirmed Multiple Systems Atrophy. Movement Disorders, 2020, 35, 1277-1279.	3.9	26

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19	Automated Brainstem Segmentation Detects Differential Involvement in Atypical Parkinsonian Syndromes. Journal of Movement Disorders, 2020, 13, 39-46.	1.3	16
20	A novel frameshift deletion in autosomal recessive SBF1-related syndromic neuropathy with necklace fibres. Journal of Neurology, 2020, 267, 2705-2712.	3.6	3
21	Identification of novel risk loci, causal insights, and heritable risk for Parkinson's disease: a meta-analysis of genome-wide association studies. Lancet Neurology, The, 2019, 18, 1091-1102.	10.2	1,414
22	The Genetic Architecture of Parkinson Disease in Spain: Characterizing Populationâ€Specific Risk, Differential Haplotype Structures, and Providing Etiologic Insight. Movement Disorders, 2019, 34, 1851-1863.	3.9	47
23	The endocytic membrane trafficking pathway plays a major role in the risk of Parkinson's disease. Movement Disorders, 2019, 34, 460-468.	3.9	66
24	<i>PDXK</i> mutations cause polyneuropathy responsive to pyridoxal 5′â€phosphate supplementation. Annals of Neurology, 2019, 86, 225-240.	5.3	54
25	Proximity extension assay testing reveals novel diagnostic biomarkers of atypical parkinsonian syndromes. Journal of Neurology, Neurosurgery and Psychiatry, 2019, 90, 768-773.	1.9	29
26	Investigation of somatic CNVs in brains of synucleinopathy cases using targeted SNCA analysis and single cell sequencing. Acta Neuropathologica Communications, 2019, 7, 219.	5.2	35
27	An update on advances in magnetic resonance imaging of multiple system atrophy. Journal of Neurology, 2019, 266, 1036-1045.	3.6	47
28	Autonomic dysfunction in genetic forms of synucleinopathies. Movement Disorders, 2018, 33, 359-371.	3.9	17
29	<i><scp>GLS</scp></i> loss of function causes autosomal recessive spastic ataxia and optic atrophy. Annals of Clinical and Translational Neurology, 2018, 5, 216-221.	3.7	13
30	Genotypeâ€phenotype correlations, dystonia and disease progression in spinocerebellar ataxia type 14. Movement Disorders, 2018, 33, 1119-1129.	3.9	26
31	Mutations in XRCC1 cause cerebellar ataxia and peripheral neuropathy. Journal of Neurology, Neurosurgery and Psychiatry, 2018, 89, 1230-1232.	1.9	18
32	LRP10 in α-synucleinopathies. Lancet Neurology, The, 2018, 17, 1032.	10.2	15
33	LRP10 in α-synucleinopathies. Lancet Neurology, The, 2018, 17, 1033-1034.	10.2	11
34	Lysosomal storage disorder gene variants in multiple system atrophy. Brain, 2018, 141, e53-e53.	7.6	11
35	Mutations in NKX6-2 Cause Progressive Spastic Ataxia and Hypomyelination. American Journal of Human Genetics, 2017, 100, 969-977.	6.2	38
36	Analysis of the prion protein gene in multiple system atrophy. Neurobiology of Aging, 2017, 49, 216.e15-216.e18.	3.1	6

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37	Genotype-phenotype correlations and expansion of the molecular spectrum of AP4M1-related hereditary spastic paraplegia. Orphanet Journal of Rare Diseases, 2017, 12, 172.	2.7	17
38	Truncating mutations in <i>SPAST</i> patients are associated with a high rate of psychiatric comorbidities in hereditary spastic paraplegia. Journal of Neurology, Neurosurgery and Psychiatry, 2017, 88, 681-687.	1.9	30
39	Triple trouble: a striking new phenotype or competing genes in a family with hereditary spastic paraplegia. Journal of Neurology, 2016, 263, 1232-1233.	3.6	4
40	Severe axonal neuropathy is a late manifestation of SPG11. Journal of Neurology, 2016, 263, 2278-2286.	3.6	11
41	A genome-wide association study in multiple system atrophy. Neurology, 2016, 87, 1591-1598.	1.1	139
42	Genetic and phenotypic characterization of complex hereditary spastic paraplegia. Brain, 2016, 139, 1904-1918.	7.6	170