## GeneviÃ"ve Bernard

List of Publications by Year in descending order

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147801 106344 5,121 124 31 65 citations g-index h-index papers 130 130 130 7250 docs citations times ranked citing authors all docs

#	Article	IF	CITATIONS
1	RNA Polymerases I and III in development and disease. Seminars in Cell and Developmental Biology, 2023, 136, 49-63.	5.0	18
2	Distinguishing severe phenotypes associated with pathogenic variants in <i>POLR3A</i> . American Journal of Medical Genetics, Part A, 2022, 188, 708-712.	1.2	4
3	Experience of Parents of Children with Genetically Determined Leukoencephalopathies Regarding the Adapted Health Care Services During the COVID-19 Pandemic. Journal of Child Neurology, 2022, 37, 237-245.	1.4	2
4	Movement disorders in MCT8 deficiency/Allan-Herndon-Dudley Syndrome. Molecular Genetics and Metabolism, 2022, 135, 109-113.	1.1	17
5	Novel biallelic variants in NRROS associated with a lethal microgliopathy, brain calcifications, and neurodegeneration. Neurogenetics, 2022, 23, 151-156.	1.4	5
6	Association of Essential Tremor With Novel Risk Loci. JAMA Neurology, 2022, 79, 185.	9.0	17
7	Therapy Trial Design in Vanishing White Matter. Neurology: Genetics, 2022, 8, e657.	1.9	12
8	Clinical, neuroradiological, and molecular characterization of patients with atypical Zellweger spectrum disorder caused by PEX16 mutations: a case series. Neurogenetics, 2022, 23, 115-127.	1.4	0
9	Oculodentodigital Dysplasia. Neurology, 2022, 98, 675-677.	1.1	2
10	Endocannabinoid dysfunction in neurological disease: neuro-ocular DAGLA-related syndrome. Brain, 2022, 145, 3383-3390.	7.6	3
11	Response to Correspondence on "Stress in Parents of Children With Genetically Determined Leukoencephalopathies: A Pilot Study― Journal of Child Neurology, 2021, 36, 245-246.	1.4	0
12	Endocrine and Growth Abnormalities in 4H Leukodystrophy Caused by Variants in <i>POLR3A</i> , <i>POLR3B</i> , and <i>POLR1C</i> . Journal of Clinical Endocrinology and Metabolism, 2021, 106, e660-e674.	3.6	26
13	A Novel De Novo Variant in DYNC1H1 Causes Spinal Muscular Atrophy Lower Extremity Predominant in Identical Twins: A Case Report. Child Neurology Open, 2021, 8, 2329048X2110274.	1.1	1
14	POLR3-related leukodystrophy: How do mutations affecting RNA polymerase III subunits cause hypomyelination?. Faculty Reviews, 2021, 10, 12.	3.9	7
15	Adult Hereditary White Matter Diseases With Psychiatric Presentation: Clinical Pointers and MRI Algorithm to Guide the Diagnostic Process. Journal of Neuropsychiatry and Clinical Neurosciences, 2021, 33, 180-193.	1.8	4
16	Variants in LSM7 impair LSM complexes assembly, neurodevelopment in zebrafish and may be associated with an ultra-rare neurological disease. Human Genetics and Genomics Advances, 2021, 2, 100034.	1.7	3
17	The LORIS MyeliNeuroGene rare disease database for natural history studies and clinical trial readiness. Orphanet Journal of Rare Diseases, 2021, 16, 328.	2.7	4
18	RNA Polymerase III Subunit Mutations in Genetic Diseases. Frontiers in Molecular Biosciences, 2021, 8, 696438.	3.5	28

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19	Expanded phenotype of AARS1-related white matter disease. Genetics in Medicine, 2021, 23, 2352-2359.	2.4	8
20	ABHD16A deficiency causes a complicated form of hereditary spastic paraplegia associated with intellectual disability and cerebral anomalies. American Journal of Human Genetics, 2021, 108, 2017-2023.	6.2	9
21	De novo variants in POLR3B cause ataxia, spasticity, and demyelinating neuropathy. American Journal of Human Genetics, 2021, 108, 186-193.	6.2	19
22	A variant of neonatal progeroid syndrome, or Wiedemann–Rautenstrauch syndrome, is associated with a nonsense variant in POLR3GL. European Journal of Human Genetics, 2020, 28, 461-468.	2.8	16
23	Genome sequencing in persistently unsolved white matter disorders. Annals of Clinical and Translational Neurology, 2020, 7, 144-152.	3.7	26
24	<i>RARS1</i> å€related hypomyelinating leukodystrophy: Expanding the spectrum. Annals of Clinical and Translational Neurology, 2020, 7, 83-93.	3.7	18
25	4H leukodystrophy. Neurology: Genetics, 2020, 6, e409.	1.9	7
26	Expanding the phenotypic and molecular spectrum of RNA polymerase III–related leukodystrophy. Neurology: Genetics, 2020, 6, e425.	1.9	20
27	POLR3A variants with striatal involvement and extrapyramidal movement disorder. Neurogenetics, 2020, 21, 121-133.	1.4	24
28	Randomized Clinical Trial of <scp>Firstâ€Line</scp> Genome Sequencing in Pediatric White Matter Disorders. Annals of Neurology, 2020, 88, 264-273.	<b>5.</b> 3	17
29	POLR3-Related Leukodystrophy: Exploring Potential Therapeutic Approaches. Frontiers in Cellular Neuroscience, 2020, 14, 631802.	3.7	9
30	Increased Prevalence of Non-motor Symptoms in Essential Tremor. Tremor and Other Hyperkinetic Movements, 2020, 4, 162.	2.0	14
31	Classifying Hypomyelination: A Critical (White) Matter. Child Neurology Open, 2020, 7, 2329048X2098376.	1.1	1
32	Patient-Derived Stem Cells, Another in vitro Model, or the Missing Link Toward Novel Therapies for Autism Spectrum Disorders?. Frontiers in Pediatrics, 2019, 7, 225.	1.9	10
33	HSD10 mitochondrial disease: p.Leu122Val variant, mild clinical phenotype, and founder effect in Frenchâ€Canadian patients from Quebec. Molecular Genetics & Denomic Medicine, 2019, 7, e1000.	1.2	8
34	Postzygotic inactivating mutations of RHOA cause a mosaic neuroectodermal syndrome. Nature Genetics, 2019, 51, 1438-1441.	21.4	25
35	Leukodystrophy-associated POLR3A mutations down-regulate the RNA polymerase III transcript and important regulatory RNA BC200. Journal of Biological Chemistry, 2019, 294, 7445-7459.	3.4	39
36	Biallelic mutations in valyl-tRNA synthetase gene VARS are associated with a progressive neurodevelopmental epileptic encephalopathy. Nature Communications, 2019, 10, 707.	12.8	28

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37	Dystonia in RNA Polymerase Illâ€Related Leukodystrophy. Movement Disorders Clinical Practice, 2019, 6, 155-159.	1.5	14
38	Biallelic Loss-of-Function Variants in AIMP1 Cause a Rare Neurodegenerative Disease. Journal of Child Neurology, 2019, 34, 74-80.	1.4	9
39	Clinical spectrum of POLR3-related leukodystrophy caused by biallelic <i>POLR1C</i> pathogenic variants. Neurology: Genetics, 2019, 5, e369.	1.9	38
40	Health-Related Quality of Life for Patients With Genetically Determined Leukoencephalopathy. Pediatric Neurology, 2018, 84, 21-26.	2.1	9
41	Exosomes Induce Fibroblast Differentiation into Cancer-Associated Fibroblasts through TGF $\hat{l}^2$ Signaling. Molecular Cancer Research, 2018, 16, 1196-1204.	3.4	200
42	POLR3A variants in hereditary spastic paraplegia and ataxia. Brain, 2018, 141, e1-e1.	7.6	17
43	Bi-allelic Mutations in EPRS, Encoding the Glutamyl-Prolyl-Aminoacyl-tRNA Synthetase, Cause a Hypomyelinating Leukodystrophy. American Journal of Human Genetics, 2018, 102, 676-684.	6.2	58
44	Recessive mutations in <i><scp>NDUFA2</scp></i> cause mitochondrial leukoencephalopathy. Clinical Genetics, 2018, 93, 396-400.	2.0	10
45	4H Leukodystrophy: Lessons from 3T Imaging. Neuropediatrics, 2018, 49, 112-117.	0.6	12
46	Recessive Mutations in POLR3B Encoding RNA Polymerase III Subunit Causing Diffuse Hypomyelination in Patients with 4H Leukodystrophy with Polymicrogyria and Cataracts. Clinical Neuroradiology, 2017, 27, 213-220.	1.9	12
47	Longitudinal Outcomes in the 2014 Acute Flaccid Paralysis Cluster in Canada. Journal of Child Neurology, 2017, 32, 301-307.	1.4	50
48	4H Leukodystrophy: A Brain Magnetic Resonance Imaging Scoring System. Neuropediatrics, 2017, 48, 152-160.	0.6	20
49	Matchmaking facilitates the diagnosis of an autosomal-recessive mitochondrial disease caused by biallelic mutation of the tRNA isopentenyltransferase ( <i>TRIT1</i> ) gene. Human Mutation, 2017, 38, 511-516.	2.5	39
50	Pediatric leukodystrophies: The role of the otolaryngologist. International Journal of Pediatric Otorhinolaryngology, 2017, 101, 141-144.	1.0	4
51	A recessive ataxia diagnosis algorithm for the next generation sequencing era. Annals of Neurology, 2017, 82, 892-899.	<b>5.</b> 3	27
52	Tissue-engineered human 3D model of bladder cancer for invasion study and drug discovery. Biomaterials, 2017, 145, 233-241.	11.4	47
53	Revised consensus statement on the preventive and symptomatic care of patients with leukodystrophies. Molecular Genetics and Metabolism, 2017, 122, 18-32.	1.1	42
54	Absence of neurological abnormalities in mice homozygous for the Polr3a G672E hypomyelinating leukodystrophy mutation. Molecular Brain, 2017, 10, 13.	2.6	33

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55	RMND1-Related Leukoencephalopathy With Temporal Lobe Cysts and Hearing Loss—Another Mendelian Mimicker of Congenital Cytomegalovirus Infection. Pediatric Neurology, 2017, 66, 59-62.	2.1	12
56	The Movement Disorder of Brain-Lung-Thyroid Syndrome Can be Responsive to Methylphenidate. Tremor and Other Hyperkinetic Movements, 2017, 7, 508.	2.0	2
57	Whole exome sequencing in patients with white matter abnormalities. Annals of Neurology, 2016, 79, 1031-1037.	5.3	116
58	Diffuse hypomyelination is not obligate for POLR3-related disorders. Neurology, 2016, 86, 1622-1626.	1.1	65
59	Expert opinion and caution are imperative for interpretation of next generation sequencing data. European Journal of Medical Genetics, 2016, 59, 519-521.	1.3	1
60	Genome-wide association study in essential tremor identifies three new loci. Brain, 2016, 139, 3163-3169.	7.6	78
61	Utility of wholeâ€exome sequencing for those near the end of the diagnostic odyssey: time to address gaps in care. Clinical Genetics, 2016, 89, 275-284.	2.0	323
62	A unique pediatric case of radiation-induced parkinsonism. Journal of Pediatric Neurology, 2015, 09, 123-126.	0.2	0
63	Altered <i>PLP1</i> splicing causes hypomyelination of early myelinating structures. Annals of Clinical and Translational Neurology, 2015, 2, 648-661.	3.7	27
64	Large exonic deletions in POLRB gene cause POLR3-related leukodystrophy. Orphanet Journal of Rare Diseases, 2015, 10, 69.	2.7	18
65	Lysophosphatidic acid enhances collagen deposition and matrix thickening in engineered tissue. Journal of Tissue Engineering and Regenerative Medicine, 2015, 9, E65-E75.	2.7	21
66	Demonstration of the direct impact of ketamine on urothelium using a tissue engineered bladder model. Canadian Urological Association Journal, 2015, 9, 613.	0.6	16
67	Endocrine Aspects of 4H Leukodystrophy: A Case Report and Review of the Literature. Case Reports in Endocrinology, 2015, 2015, 1-6.	0.4	10
68	Disease specific therapies in leukodystrophies and leukoencephalopathies. Molecular Genetics and Metabolism, 2015, 114, 527-536.	1.1	45
69	Characterization of human disease phenotypes associated with mutations in <i>TREX1</i> , <i>RNASEH2A</i> , <i>RNASEH2B</i> , <i>RNASEH2C</i> , <i>SAMHD1</i> , <i>ADAR</i> , and <i>IFIH1</i> American Journal of Medical Genetics, Part A, 2015, 167, 296-312.	1.2	447
70	A clinical approach to the diagnosis of patients with leukodystrophies and genetic leukoencephelopathies. Molecular Genetics and Metabolism, 2015, 114, 501-515.	1.1	163
71	Consensus statement on preventive and symptomatic care of leukodystrophy patients. Molecular Genetics and Metabolism, 2015, 114, 516-526.	1.1	29
72	Recovery From Central Nervous System Acute Demyelination in Children. Pediatrics, 2015, 136, e115-e123.	2.1	40

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73	Neurogenic bladder and neuroendocrine abnormalities in Pol III-related leukodystrophy. BMC Neurology, 2015, 15, 22.	1.8	6
74	POLR3A and POLR3B Mutations in Unclassified Hypomyelination. Neuropediatrics, 2015, 46, 221-228.	0.6	22
75	Recessive mutations in POLR1C cause a leukodystrophy by impairing biogenesis of RNA polymerase III. Nature Communications, 2015, 6, 7623.	12.8	127
76	Mutations in RNF216 do not cause 4H syndrome. Parkinsonism and Related Disorders, 2015, 21, 1387-1388.	2,2	2
77	Novel SIL1 mutations cause cerebellar ataxia and atrophy in a French-Canadian family. Neurogenetics, 2015, 16, 315-318.	1.4	5
78	Adipose-derived stromal cells for the reconstruction of a human vesical equivalent. Journal of Tissue Engineering and Regenerative Medicine, 2015, 9, E135-E143.	2.7	28
79	Characterization of a psoriatic skin model produced with involved or uninvolved cells. Journal of Tissue Engineering and Regenerative Medicine, 2015, 9, 789-798.	2.7	18
80	Myelination Delay and Allan-Herndon-Dudley Syndrome Caused by a Novel Mutation in the <i>SLC16A2</i> Gene. Journal of Child Neurology, 2015, 30, 1371-1374.	1.4	20
81	Spastic Paraparesis and Marked Improvement of Leukoencephalopathy in Aicardi–Goutières Syndrome. Neuropediatrics, 2014, 45, 406-410.	0.6	9
82	Hypomyelinating leukodystrophies: Translational research progress and prospects. Annals of Neurology, 2014, 76, 5-19.	5.3	132
83	Brain Magnetic Resonance Imaging (MRI) Pattern Recognition in Pol III-Related Leukodystrophies. Journal of Child Neurology, 2014, 29, 214-220.	1.4	47
84	Clinical spectrum of 4H leukodystrophy caused by <i>POLR3A</i> and <i>POLR3B</i> mutations. Neurology, 2014, 83, 1898-1905.	1.1	170
85	<i>TUBB4A</i> de novo mutations cause isolated hypomyelination. Neurology, 2014, 83, 898-902.	1.1	52
86	A homozygous mutation in the NDUFS1 gene presents with a mild cavitating leukoencephalopathy. Neurogenetics, 2014, 15, 161-164.	1.4	12
87	Vanishing White Matter Disease in French-Canadian Patients From Quebec. Pediatric Neurology, 2014, 51, 225-232.	2.1	6
88	Increased Prevalence of Non-motor Symptoms in Essential Tremor. Tremor and Other Hyperkinetic Movements, 2014, 4, 162.	2.0	16
89	A treatable new cause of chorea: Betaâ€ketothiolase deficiency. Movement Disorders, 2013, 28, 1054-1056.	3.9	25
90	Assessment of interferon-related biomarkers in Aicardi-Goutières syndrome associated with mutations in TREX1, RNASEH2A, RNASEH2B, RNASEH2C, SAMHD1, and ADAR: a case-control study. Lancet Neurology, The, 2013, 12, 1159-1169.	10.2	473

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91	Strategies to Reconstruct a Functional Urethral Substitute by Self-assembly Method. Procedia Engineering, 2013, 59, 193-200.	1.2	4
92	Mutations in <i>POLR3A</i> and <i>POLR3B</i> are a major cause of hypomyelinating leukodystrophies with or without dental abnormalities and/or hypogonadotropic hypogonadism. Journal of Medical Genetics, 2013, 50, 194-197.	3.2	109
93	Teaching Neuro <i>Images</i> : Hypomyelinating leukodystrophy with hypodontia due to <i>POLR3B</i> . Neurology, 2013, 81, e145.	1.1	5
94	More Than Hypomyelination in Pol-III Disorder. Journal of Neuropathology and Experimental Neurology, 2013, 72, 67-75.	1.7	27
95	Diversity of ARSACS Mutations in French-Canadians. Canadian Journal of Neurological Sciences, 2013, 40, 61-66.	0.5	51
96	Adult-Onset Vanishing White Matter Disease Due to a Novel EIF2B3 Mutation. Archives of Neurology, 2012, 69, 765-68.	4.5	27
97	TACH Leukodystrophy: Locus Refinement to Chromosome 10q22.3-23.1. Canadian Journal of Neurological Sciences, 2012, 39, 122-123.	0.5	4
98	A Novel <i>PLP1</i> Mutation Further Expands the Clinical Heterogeneity at the Locus. Canadian Journal of Neurological Sciences, 2012, 39, 220-224.	0.5	3
99	4H Syndrome With Late-Onset Growth Hormone Deficiency Caused by POLR3A Mutations. Archives of Neurology, 2012, 69, 920-3.	4.5	56
100	Advances in the diagnosis of leukodystrophies. Future Neurology, 2012, 7, 595-612.	0.5	7
101	Tremorâ€ataxia with central hypomyelination (TACH): Dystonia as a new clinical feature. Movement Disorders, 2012, 27, 1831-1832.	3.9	6
102	Abnormal Myelination in Ring Chromosome 18 Syndrome. Journal of Child Neurology, 2012, 27, 1042-1047.	1.4	9
103	Ataxia-Telangiectasia Presenting With a Novel Immunodeficiency. Pediatric Neurology, 2012, 46, 322-324.	2.1	11
104	Exome Sequencing Identifies FUS Mutations as a Cause of Essential Tremor. American Journal of Human Genetics, 2012, 91, 313-319.	6.2	176
105	Mutations of POLR3A Encoding a Catalytic Subunit of RNA Polymerase Pol III Cause a Recessive Hypomyelinating Leukodystrophy p415. American Journal of Human Genetics, 2012, 91, 972.	6.2	1
106	Mutations in the Mitochondrial Methionyl-tRNA Synthetase Cause a Neurodegenerative Phenotype in Flies and a Recessive Ataxia (ARSAL) in Humans. PLoS Biology, 2012, 10, e1001288.	5.6	147
107	Effects of Serum-Free Culture at the Air–Liquid Interface in a Human Tissue-Engineered Skin Substitute. Tissue Engineering - Part A, 2011, 17, 877-888.	3.1	20
108	Surgical Option for the Correction of Peyronie's Disease: An Autologous Tissue-Engineered Endothelialized Graft. Journal of Sexual Medicine, 2011, 8, 3227-3235.	0.6	33

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109	Mutations of POLR3A Encoding a Catalytic Subunit of RNA Polymerase Pol III Cause a Recessive Hypomyelinating Leukodystrophy. American Journal of Human Genetics, 2011, 89, 415-423.	6.2	219
110	Recessive Mutations in POLR3B, Encoding the Second Largest Subunit of Pol III, Cause a Rare Hypomyelinating Leukodystrophy. American Journal of Human Genetics, 2011, 89, 652-655.	6.2	139
111	LINGO1 Variants in the French-Canadian Population. PLoS ONE, 2011, 6, e16254.	2.5	23
112	Tremor–ataxia with central hypomyelination (TACH) leukodystrophy maps to chromosome 10q22.3–10q23.31. Neurogenetics, 2010, 11, 457-464.	1.4	39
113	Chylomicron retention disease: Dystonia as a new clinical feature. Movement Disorders, 2010, 25, 1755-1756.	3.9	2
114	A Case of Secondary Dystonia Responding to Levodopa. Journal of Child Neurology, 2010, 25, 780-781.	1.4	13
115	Plasticity of locomotor sensorimotor interactions after peripheral and/or spinal lesions. Brain Research Reviews, 2008, 57, 228-240.	9.0	81
116	The Wobbly Child: An Approach to Inherited Ataxias. Seminars in Pediatric Neurology, 2008, 15, 194-208.	2.0	16
117	Simultaneous Guillain-Barré Syndrome and Acute Disseminated Encephalomyelitis in the Pediatric Population. Journal of Child Neurology, 2008, 23, 752-757.	1.4	25
118	Channelopathies: A Review. Pediatric Neurology, 2008, 38, 73-85.	2.1	69
119	Acute Combined Central and Peripheral Nervous System Demyelination in Children. Pediatric Neurology, 2008, 39, 307-316.	2.1	26
120	Dosage Effect of a Dominant CLCN1 Mutation: A Novel Syndrome. Journal of Child Neurology, 2008, 23, 163-166.	1.4	21
121	Refractory and lethal status epilepticus in a patient with ring chromosome 20 syndrome. Epileptic Disorders, 2008, 10, 254-259.	1.3	17
122	Neurotransmitter diseases and related conditions. Molecular Genetics and Metabolism, 2007, 92, 189-197.	1.1	3
123	Study of Cutaneous Reflex Compensation During Locomotion After Nerve Section in the Cat. Journal of Neurophysiology, 2007, 97, 4173-4185.	1.8	20
124	Insights on the Interactions of Synthetic Amphipathic Peptides with Model Membranes as Revealed by 31P and 2H Solid-State NMR and Infrared Spectroscopies. Biophysical Journal, 2006, 90, 4071-4084.	0.5	32