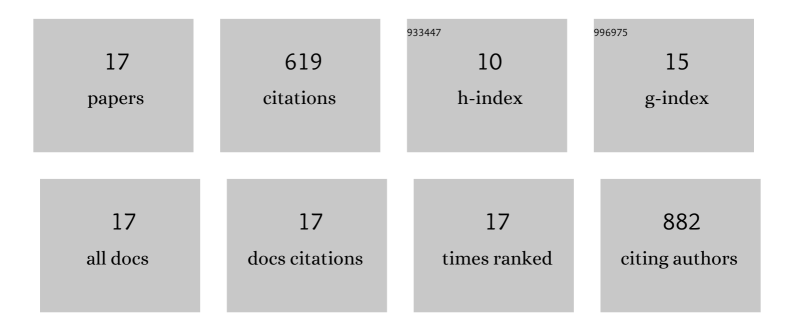
Wai Yan Yau

List of Publications by Year in descending order

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ΜΛΛΙ ΥΛΝ ΥΛΙΙ

#	Article	IF	CITATIONS
1	Biallelic expansion of an intronic repeat in RFC1 is a common cause of late-onset ataxia. Nature Genetics, 2019, 51, 649-658.	21.4	338
2	A novel RFC1 repeat motif (ACAGG) in two Asia-Pacific CANVAS families. Brain, 2020, 143, 2904-2910.	7.6	53
3	A MÄori specific RFC1 pathogenic repeat configuration in CANVAS, likely due to a founder allele. Brain, 2020, 143, 2673-2680.	7.6	45
4	Neuronal intranuclear inclusion disease is genetically heterogeneous. Annals of Clinical and Translational Neurology, 2020, 7, 1716-1725.	3.7	38
5	<scp><i>RFC1</i></scp> Intronic Repeat Expansions Absent in Pathologically Confirmed Multiple Systems Atrophy. Movement Disorders, 2020, 35, 1277-1279.	3.9	26
6	<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
7	Low Prevalence of NOTCH2NLC GGC Repeat Expansion in White Patients with Movement Disorders. Movement Disorders, 2021, 36, 251-255.	3.9	23
8	<scp>GGC</scp> Repeat Expansion in <scp><i>NOTCH2NLC</i></scp> Is Rare in European Leukoencephalopathy. Annals of Neurology, 2020, 88, 641-642.	5.3	14
9	Cerebellar ataxia, neuropathy and vestibular areflexia syndrome (CANVAS): genetic and clinical aspects. Practical Neurology, 2022, 22, 14-18.	1.1	14
10	GGC repeat expansion in NOTCH2NLC is rare in European patients with essential tremor. Brain, 2020, 143, e57-e57.	7.6	13
11	<scp>DNA</scp> repair in trinucleotide repeat ataxias. FEBS Journal, 2018, 285, 3669-3682.	4.7	12
12	<scp><i>NOTCH2NLC</i></scp> Intermediateâ€Length Repeat Expansion and Parkinson's Disease in Patients of European Descent. Annals of Neurology, 2021, 89, 633-635.	5.3	7
13	Fabry heterozygote mimicking multiple sclerosis. BMJ Case Reports, 2017, 2017, bcr-2017-220274.	0.5	6
14	The repeat variant in MSH3 is not a genetic modifier for spinocerebellar ataxia type 3 and Friedreich's ataxia. Brain, 2020, 143, e25-e25.	7.6	3
15	Questions on <i>NOTCH2NLC</i> Repeat Expansions in Parkinson Disease. JAMA Neurology, 2021, 78, 762.	9.0	2
16	Acute reversible seronegative cerebellar ataxia in a young woman with ovarian teratoma. Journal of the Neurological Sciences, 2016, 369, 227-228.	0.6	0
17	Reply: Genetic heterogeneity of neuronal intranuclear inclusion disease. What about the infantile variant?. Annals of Clinical and Translational Neurology, 2021, 8, 1002-1004.	3.7	0