## Liesbet Deprez

List of Publications by Year in descending order

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LIFERET NEDDEZ

#	Article	lF	CITATIONS
1	<i>KCNQ2</i> encephalopathy: Emerging phenotype of a neonatal epileptic encephalopathy. Annals of Neurology, 2012, 71, 15-25.	5.3	427
2	Paroxysmal exercise-induced dyskinesia and epilepsy is due to mutations in SLC2A1, encoding the glucose transporter GLUT1. Brain, 2008, 131, 1831-1844.	7.6	340
3	Earlyâ€onset absence epilepsy caused by mutations in the glucose transporter GLUT1. Annals of Neurology, 2009, 66, 415-419.	5.3	266
4	Molecular correlates of age-dependent seizures in an inherited neonatal-infantile epilepsy. Brain, 2010, 133, 1403-1414.	7.6	157
5	Microdeletions involving theSCN1A gene may be common inSCN1A-mutation-negative SMEI patients. Human Mutation, 2006, 27, 914-920.	2.5	114
6	The <i>SCN1A</i> variant database: a novel research and diagnostic tool. Human Mutation, 2009, 30, E904-E920.	2.5	112
7	Epilepsy as part of the phenotype associated with <i>ATP1A2</i> mutations. Epilepsia, 2008, 49, 500-508.	5.1	83
8	Genetics of epilepsy syndromes starting in the first year of life. Neurology, 2009, 72, 273-281.	1,1	63
9	Validation of a digital PCR method for quantification of DNA copy number concentrations by using a certified reference material. Biomolecular Detection and Quantification, 2016, 9, 29-39.	7.0	53
10	Towards an SI-Traceable Reference Measurement System for Seven Serum Apolipoproteins Using Bottom-Up Quantitative Proteomics: Conceptual Approach Enabled by Cross-Disciplinary/Cross-Sector Collaboration. Clinical Chemistry, 2021, 67, 478-489.	3.2	52
11	Assessment of Digital PCR as a Primary Reference Measurement Procedure to Support Advances in Precision Medicine. Clinical Chemistry, 2018, 64, 1296-1307.	3.2	50
12	Novel frameshift and splice site mutations in the neurotrophic tyrosine kinase receptor type 1 gene (NTRK1) associated with hereditary sensory neuropathy type IV. Neuromuscular Disorders, 2006, 16, 19-25.	0.6	30
13	Genome-wide linkage of febrile seizures and epilepsy to the FEB4 locus at 5q14.3-q23.1 and no MASS1 mutation. Human Genetics, 2006, 118, 618-625.	3.8	19
14	Faecal immunochemical tests for haemoglobin: Analytical challenges and potential solutions. Clinica Chimica Acta, 2021, 517, 60-65.	1.1	17
15	Commutability Assessment of Candidate Reference Materials for Pancreatic α-Amylase. Clinical Chemistry, 2018, 64, 1193-1202.	3.2	15