Pau Pastor

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107 240 12,713 49 h-index g-index citations papers 268 16,727 7.6 5.17 avg, IF L-index ext. citations ext. papers

#	Paper	IF	Citations
240	Meta-analysis of 74,046 individuals identifies 11 new susceptibility loci for Alzheimer's disease. <i>Nature Genetics</i> , 2013 , 45, 1452-8	36.3	2714
239	Genetic meta-analysis of diagnosed Alzheimer's disease identifies new risk loci and implicates Alltau, immunity and lipid processing. <i>Nature Genetics</i> , 2019 , 51, 414-430	36.3	917
238	Identification of novel risk loci, causal insights, and heritable risk for Parkinson's disease: a meta-analysis of genome-wide association studies. <i>Lancet Neurology, The</i> , 2019 , 18, 1091-1102	24.1	562
237	Rare coding variants in PLCG2, ABI3, and TREM2 implicate microglial-mediated innate immunity in Alzheimer's disease. <i>Nature Genetics</i> , 2017 , 49, 1373-1384	36.3	508
236	Identification of common variants influencing risk of the tauopathy progressive supranuclear palsy. <i>Nature Genetics</i> , 2011 , 43, 699-705	36.3	386
235	Rare coding variants in the phospholipase D3 gene confer risk for Alzheimer's disease. <i>Nature</i> , 2014 , 505, 550-554	50.4	345
234	Structural brain changes in Parkinson disease with dementia: a voxel-based morphometry study. <i>Archives of Neurology</i> , 2005 , 62, 281-5		238
233	Frontotemporal dementia and its subtypes: a genome-wide association study. <i>Lancet Neurology, The</i> , 2014 , 13, 686-99	24.1	207
232	PINK1-linked parkinsonism is associated with Lewy body pathology. <i>Brain</i> , 2010 , 133, 1128-42	11.2	188
231	A novel Alzheimer disease locus located near the gene encoding tau protein. <i>Molecular Psychiatry</i> , 2016 , 21, 108-17	15.1	175
230	HDDD2 is a familial frontotemporal lobar degeneration with ubiquitin-positive, tau-negative inclusions caused by a missense mutation in the signal peptide of progranulin. <i>Annals of Neurology</i> , 2006 , 60, 314-22	9.4	174
229	Familial atypical progressive supranuclear palsy associated with homozigosity for the delN296 mutation in the tau gene. <i>Annals of Neurology</i> , 2001 , 49, 263-7	9.4	165
228	Genetic screening of Alzheimer's disease genes in Iberian and African samples yields novel mutations in presenilins and APP. <i>Neurobiology of Aging</i> , 2010 , 31, 725-31	5.6	162
227	Genetic analysis implicates APOE, SNCA and suggests lysosomal dysfunction in the etiology of dementia with Lewy bodies. <i>Human Molecular Genetics</i> , 2014 , 23, 6139-46	5.6	152
226	Convergent genetic and expression data implicate immunity in Alzheimer's disease. <i>Alzheimermand Dementia</i> , 2015 , 11, 658-71	1.2	146
225	Apolipoprotein Eepsilon4 modifies Alzheimer's disease onset in an E280A PS1 kindred. <i>Annals of Neurology</i> , 2003 , 54, 163-9	9.4	133
224	TREM2 is associated with the risk of Alzheimer's disease in Spanish population. <i>Neurobiology of Aging</i> , 2013 , 34, 1711.e15-7	5.6	121

(2016-2018)

223	Investigating the genetic architecture of dementia with Lewy bodies: a two-stage genome-wide association study. <i>Lancet Neurology, The</i> , 2018 , 17, 64-74	24.1	121
222	Significant association between the tau gene A0/A0 genotype and Parkinson's disease. <i>Annals of Neurology</i> , 2000 , 47, 242-245	9.4	111
221	Amygdalar and hippocampal MRI volumetric reductions in Parkinson's disease with dementia. <i>Movement Disorders</i> , 2005 , 20, 540-4	7	109
220	The unfolded protein response is activated in disease-affected brain regions in progressive supranuclear palsy and Alzheimer's disease. <i>Acta Neuropathologica Communications</i> , 2013 , 1, 31	7.3	104
219	Automated neuromelanin imaging as a diagnostic biomarker for Parkinson's disease. <i>Movement Disorders</i> , 2015 , 30, 945-52	7	93
218	Gene-wide analysis detects two new susceptibility genes for Alzheimer's disease. <i>PLoS ONE</i> , 2014 , 9, e94661	3.7	90
217	TREM2 and neurodegenerative disease. New England Journal of Medicine, 2013, 369, 1568-9	59.2	89
216	A novel mutation (K317M) in the MAPT gene causes FTDP and motor neuron disease. <i>Neurology</i> , 2005 , 64, 1578-85	6.5	88
215	Characterization of the repeat expansion size in C9orf72 in amyotrophic lateral sclerosis and frontotemporal dementia. <i>Human Molecular Genetics</i> , 2014 , 23, 749-54	5.6	84
214	Missense mutations in TENM4, a regulator of axon guidance and central myelination, cause essential tremor. <i>Human Molecular Genetics</i> , 2015 , 24, 5677-86	5.6	83
213	Rare mutations in SQSTM1 modify susceptibility to frontotemporal lobar degeneration. <i>Acta Neuropathologica</i> , 2014 , 128, 397-410	14.3	83
212	Assessing the role of the TREM2 p.R47H variant as a risk factor for Alzheimer's disease and frontotemporal dementia. <i>Neurobiology of Aging</i> , 2014 , 35, 444.e1-4	5.6	81
211	Pooled-DNA sequencing identifies novel causative variants in PSEN1, GRN and MAPT in a clinical early-onset and familial Alzheimer's disease Ibero-American cohort. <i>Alzheimer na Research and Therapy</i> , 2012 , 4, 34	9	78
210	Immune-related genetic enrichment in frontotemporal dementia: An analysis of genome-wide association studies. <i>PLoS Medicine</i> , 2018 , 15, e1002487	11.6	77
209	Genetic architecture of sporadic frontotemporal dementia and overlap with Alzheimer's and Parkinson's diseases. <i>Journal of Neurology, Neurosurgery and Psychiatry</i> , 2017 , 88, 152-164	5.5	76
208	Analysis of the C9orf72 gene in patients with amyotrophic lateral sclerosis in Spain and different populations worldwide. <i>Human Mutation</i> , 2013 , 34, 79-82	4.7	71
207	TBK1 Mutation Spectrum in an Extended European Patient Cohort with Frontotemporal Dementia and Amyotrophic Lateral Sclerosis. <i>Human Mutation</i> , 2017 , 38, 297-309	4.7	66
206	A comprehensive study of the genetic impact of rare variants in SORL1 in European early-onset Alzheimer's disease. <i>Acta Neuropathologica</i> , 2016 , 132, 213-224	14.3	62

205	Identification of Spanish familial Parkinson's disease and screening for the Ala53Thr mutation of the alpha-synuclein gene in early onset patients. <i>Neuroscience Letters</i> , 1997 , 235, 57-60	3.3	61
204	Novel haplotypes in 17q21 are associated with progressive supranuclear palsy. <i>Annals of Neurology</i> , 2004 , 56, 249-58	9.4	59
203	Heterozygous mutation causes familial ataxia with cognitive affective syndrome (SCA48). <i>Neurology</i> , 2018 , 91, e1988-e1998	6.5	59
202	Phosphorylated neurofilament heavy chain: A biomarker of survival for C9ORF72-associated amyotrophic lateral sclerosis. <i>Annals of Neurology</i> , 2017 , 82, 139-146	9.4	58
201	Genetic risk score predicting accelerated progression from mild cognitive impairment to Alzheimer's disease. <i>Journal of Neural Transmission</i> , 2013 , 120, 807-12	4.3	58
200	Molecular genetics of Alzheimer's disease. <i>Current Psychiatry Reports</i> , 2004 , 6, 125-33	9.1	58
199	Genome-wide association study in essential tremor identifies three new loci. <i>Brain</i> , 2016 , 139, 3163-31	69 1.2	57
198	Replication of MAPT and SNCA, but not PARK16-18, as susceptibility genes for Parkinson's disease. <i>Movement Disorders</i> , 2011 , 26, 819-23	7	55
197	Identification of evolutionarily conserved gene networks mediating neurodegenerative dementia. <i>Nature Medicine</i> , 2019 , 25, 152-164	50.5	55
196	Identification of a novel polymorphism in the promoter region of the tau gene highly associated to progressive supranuclear palsy in humans. <i>Neuroscience Letters</i> , 1999 , 275, 183-6	3.3	52
195	Moving beyond neurons: the role of cell type-specific gene regulation in Parkinson's disease heritability. <i>Npj Parkinson Disease</i> , 2019 , 5, 6	9.7	51
194	C9ORF72 repeat expansion in Australian and Spanish frontotemporal dementia patients. <i>PLoS ONE</i> , 2013 , 8, e56899	3.7	51
193	A nonsynonymous mutation in PLCG2 reduces the risk of Alzheimer's disease, dementia with Lewy bodies and frontotemporal dementia, and increases the likelihood of longevity. <i>Acta Neuropathologica</i> , 2019 , 138, 237-250	14.3	50
192	Genome-wide analysis of genetic correlation in dementia with Lewy bodies, Parkinson's and Alzheimer's diseases. <i>Neurobiology of Aging</i> , 2016 , 38, 214.e7-214.e10	5.6	49
191	Global investigation and meta-analysis of the C9orf72 (G4C2)n repeat in Parkinson disease. <i>Neurology</i> , 2014 , 83, 1906-13	6.5	49
190	Dementia in Parkinson disease: a proton magnetic resonance spectroscopy study. <i>Archives of Neurology</i> , 2002 , 59, 1415-20		48
189	Mitochondria function associated genes contribute to Parkinson's Disease risk and later age at onset. <i>Npj Parkinson Disease</i> , 2019 , 5, 8	9.7	47
188	Analysis of the CHCHD10 gene in patients with frontotemporal dementia and amyotrophic lateral sclerosis from Spain. <i>Brain</i> , 2015 , 138, e400	11.2	47

(2012-2002)

187	Further extension of the H1 haplotype associated with progressive supranuclear palsy. <i>Movement Disorders</i> , 2002 , 17, 550-6	7	47
186	Resequencing analysis of five Mendelian genes and the top genes from genome-wide association studies in Parkinson's Disease. <i>Molecular Neurodegeneration</i> , 2016 , 11, 29	19	46
185	COPPADIS-2015 (COhort of Patients with PArkinson's Disease in Spain, 2015), a globalclinical evaluations, serum biomarkers, genetic studies and neuroimagingprospective, multicenter, non-interventional, long-term study on Parkinson's disease progression. <i>BMC Neurology</i> , 2016 , 16, 26	3.1	46
184	Dopaminergic neuronal imaging in genetic Parkinson's disease: insights into pathogenesis. <i>PLoS ONE</i> , 2013 , 8, e69190	3.7	46
183	Genome-wide association analysis of dementia and its clinical endophenotypes reveal novel loci associated with Alzheimer's disease and three causality networks: The GR@ACE project. <i>Alzheimern</i> and Dementia, 2019 , 15, 1333-1347	1.2	45
182	Memory decline evolves independently of disease activity in MS. <i>Multiple Sclerosis Journal</i> , 2008 , 14, 947-53	5	45
181	Cortical atrophy and language network reorganization associated with a novel progranulin mutation. <i>Cerebral Cortex</i> , 2009 , 19, 1751-60	5.1	42
180	Shared genetic contribution to Ischaemic Stroke and Alzheimer's Disease. <i>Annals of Neurology</i> , 2016 , 79, 739-747	9.4	42
179	The endocytic membrane trafficking pathway plays a major role in the risk of Parkinson's disease. <i>Movement Disorders</i> , 2019 , 34, 460-468	7	40
178	Non-motor symptoms burden, mood, and gait problems are the most significant factors contributing to a poor quality of life in non-demented Parkinson's disease patients: Results from the COPPADIS Study Cohort. <i>Parkinsonism and Related Disorders</i> , 2019 , 66, 151-157	3.6	39
177	Automated analysis of FDG PET as a tool for single-subject probabilistic prediction and detection of Alzheimer's disease dementia. <i>European Journal of Nuclear Medicine and Molecular Imaging</i> , 2013 , 40, 1394-405	8.8	38
176	Update on genetics of essential tremor. Acta Neurologica Scandinavica, 2013, 128, 359-71	3.8	38
175	Tau phosphorylation and kinase activation in familial tauopathy linked to deln296 mutation. <i>Neuropathology and Applied Neurobiology</i> , 2003 , 29, 23-34	5.2	38
174	A novel mutation in the PSEN2 gene (T430M) associated with variable expression in a family with early-onset Alzheimer disease. <i>Archives of Neurology</i> , 2003 , 60, 1149-51		37
173	Enhanced gain of blink reflex responses to ipsilateral supraorbital nerve afferent inputs in patients with facial nerve palsy. <i>Clinical Neurophysiology</i> , 2001 , 112, 153-6	4.3	37
172	Polygenic risk and hazard scores for Alzheimer's disease prediction. <i>Annals of Clinical and Translational Neurology</i> , 2019 , 6, 456-465	5.3	36
171	CXCR4 involvement in neurodegenerative diseases. <i>Translational Psychiatry</i> , 2018 , 8, 73	8.6	36
170	A search for SNCA 3' UTR variants identified SNP rs356165 as a determinant of disease risk and onset age in Parkinson's disease. <i>Journal of Molecular Neuroscience</i> , 2012 , 47, 425-30	3.3	36

169	LINGO1 and risk for essential tremor: results of a meta-analysis of rs9652490 and rs11856808. Journal of the Neurological Sciences, 2012 , 317, 52-7	3.2	35
168	Deleterious ABCA7 mutations and transcript rescue mechanisms in early onset Alzheimer's disease. <i>Acta Neuropathologica</i> , 2017 , 134, 475-487	14.3	34
167	No evidence of association between common European mitochondrial DNA variants in Alzheimer, Parkinson, and migraine in the Spanish population. <i>American Journal of Medical Genetics Part B: Neuropsychiatric Genetics</i> , 2015 , 168B, 54-65	3.5	33
166	Investigation of the role of rare TREM2 variants in frontotemporal dementia subtypes. <i>Neurobiology of Aging</i> , 2014 , 35, 2657.e13-2657.e19	5.6	31
165	Frontobasal gray matter loss is associated with the TREM2 p.R47H variant. <i>Neurobiology of Aging</i> , 2014 , 35, 2681-2690	5.6	31
164	Age at onset in LRRK2-associated PD is modified by SNCA variants. <i>Journal of Molecular Neuroscience</i> , 2012 , 48, 245-7	3.3	31
163	Genetic variation in APOE cluster region and Alzheimer's disease risk. <i>Neurobiology of Aging</i> , 2011 , 32, 2107.e7-17	5.6	31
162	Forceps minor region signal abnormality "ears of the lynx": an early MRI finding in spastic paraparesis with thin corpus callosum and mutations in the spatacsin gene (SPG11) on chromosome 15. <i>Journal of Neuroimaging</i> , 2009 , 19, 52-60	2.8	31
161	Genome sequencing analysis identifies new loci associated with Lewy body dementia and provides insights into its genetic architecture. <i>Nature Genetics</i> , 2021 , 53, 294-303	36.3	31
160	Examination of motor output pathways in patients with corticobasal ganglionic degeneration using transcranial magnetic stimulation. <i>Brain</i> , 2001 , 124, 1131-7	11.2	30
159	Genetic variability in SQSTM1 and risk of early-onset Alzheimer dementia: a European early-onset dementia consortium study. <i>Neurobiology of Aging</i> , 2015 , 36, 2005.e15-22	5.6	29
158	Rare variants in EAmyloid precursor protein (APP) and Parkinson's disease. <i>European Journal of Human Genetics</i> , 2015 , 23, 1328-33	5.3	29
157	An association study between Heme oxygenase-1 genetic variants and Parkinson's disease. <i>Frontiers in Cellular Neuroscience</i> , 2014 , 8, 298	6.1	29
156	Ubiquitin associated protein 1 is a risk factor for frontotemporal lobar degeneration. <i>Neurobiology of Aging</i> , 2009 , 30, 656-65	5.6	29
155	Significant association between the tau gene A0/A0 genotype and Parkinson's disease. <i>Annals of Neurology</i> , 2000 , 47, 242-5	9.4	29
154	Heme Oxygenase-1 and 2 Common Genetic Variants and Risk for Restless Legs Syndrome. <i>Medicine</i> (<i>United States</i>), 2015 , 94, e1448	1.8	28
153	Penetrance of Parkinson's Disease in LRRK2 p.G2019S Carriers Is Modified by a Polygenic Risk Score. <i>Movement Disorders</i> , 2020 , 35, 774-780	7	27
152	Lack of association of LINGO1 rs9652490 and rs11856808 SNPs with familial essential tremor. European Journal of Neurology, 2011 , 18, 1085-9	6	27

151	SPG11 compound mutations in spastic paraparesis with thin corpus callosum. <i>Neurology</i> , 2008 , 71, 332-6	56.5	27
150	A novel presenilin 2 gene mutation (D439A) in a patient with early-onset Alzheimer's disease. <i>Neurology</i> , 2001 , 57, 1926-8	6.5	27
149	New insights into the genetic etiology of Alzheimer's disease and related dementias <i>Nature Genetics</i> , 2022 ,	36.3	27
148	TREM2 R47H variant and risk of essential tremor: a cross-sectional international multicenter study. <i>Parkinsonism and Related Disorders</i> , 2015 , 21, 306-9	3.6	26
147	Protective effect of LRRK2 p.R1398H on risk of Parkinson's disease is independent of MAPT and SNCA variants. <i>Neurobiology of Aging</i> , 2014 , 35, 266.e5-14	5.6	26
146	MAPT H1 Haplotype is Associated with Late-Onset Alzheimer's Disease Risk in APOEe4 Noncarriers: Results from the Dementia Genetics Spanish Consortium. <i>Journal of Alzheimer</i> Disease, 2016 , 49, 343-5	5 4 .3	26
145	A new mutation in the parkin gene in a patient with atypical autosomal recessive juvenile parkinsonism. <i>Neuroscience Letters</i> , 2000 , 289, 66-8	3.3	26
144	Different MAPT haplotypes are associated with Parkinson's disease and progressive supranuclear palsy. <i>Neurobiology of Aging</i> , 2011 , 32, 547.e11-6	5.6	25
143	Gamma-aminobutyric acid GABRA4, GABRE, and GABRQ receptor polymorphisms and risk for essential tremor. <i>Pharmacogenetics and Genomics</i> , 2011 , 21, 436-9	1.9	25
142	Gamma-aminobutyric acid (GABA) receptor rho (GABRR) polymorphisms and risk for essential tremor. <i>Journal of Neurology</i> , 2011 , 258, 203-11	5.5	25
141	Analysis of the Micro-RNA-133 and PITX3 genes in Parkinson's disease. <i>American Journal of Medical Genetics Part B: Neuropsychiatric Genetics</i> , 2010 , 153B, 1234-9	3.5	25
140	New insights on the genetic etiology of Alzheimer® and related dementia		25
139	A C6orf10/LOC101929163 locus is associated with age of onset in C9orf72 carriers. <i>Brain</i> , 2018 , 141, 2895-2907	11.2	25
138	5'-Upstream variants of CRHR1 and MAPT genes associated with age at onset in progressive supranuclear palsy and cortical basal degeneration. <i>Neurobiology of Disease</i> , 2009 , 33, 164-70	7.5	24
137	Discovering the 3' UTR-mediated regulation of alpha-synuclein. <i>Nucleic Acids Research</i> , 2017 , 45, 12888-	1229:03	23
136	Reversible oral-facial dyskinesia in a patient receiving ciprofloxacin hydrochloride. <i>Journal of Neurology</i> , 1996 , 243, 616-7	5.5	23
135	Common variants in Alzheimer's disease and risk stratification by polygenic risk scores. <i>Nature Communications</i> , 2021 , 12, 3417	17.4	23
134	Non-motor symptom burden is strongly correlated to motor complications in patients with Parkinson's disease. <i>European Journal of Neurology</i> , 2020 , 27, 1210-1223	6	22

133	Lack of evidence for a role of genetic variation in TMEM230 in the risk for Parkinson's disease in the Caucasian population. <i>Neurobiology of Aging</i> , 2017 , 50, 167.e11-167.e13	5.6	22
132	Rare variants in calcium homeostasis modulator 1 (CALHM1) found in early onset Alzheimer's disease patients alter calcium homeostasis. <i>PLoS ONE</i> , 2013 , 8, e74203	3.7	22
131	COPPADIS-2015 (COhort of Patients with PArkinson's Disease in Spain, 2015): an ongoing global Parkinson's disease project about disease progression with more than 1000 subjects included. Results from the baseline evaluation. <i>European Journal of Neurology</i> , 2019 , 26, 1399-1407	6	21
130	Fused in Sarcoma (FUS) gene mutations are not a frequent cause of essential tremor in Europeans. <i>Neurobiology of Aging</i> , 2013 , 34, 2441.e9-2441.e11	5.6	21
129	Nigrostriatal dopaminergic function in subjects with isolated action tremor. <i>Parkinsonism and Related Disorders</i> , 2012 , 18, 49-53	3.6	21
128	The effect of MAPT H1 and APOE 4 on transition from mild cognitive impairment to dementia. Journal of Alzheimerm Disease, 2010 , 22, 1065-71	4.3	21
127	Relative high frequency of the c.255delA parkin gene mutation in Spanish patients with autosomal recessive parkinsonism. <i>Journal of Neurology, Neurosurgery and Psychiatry</i> , 2002 , 73, 582-4	5.5	21
126	Aberrant mitochondria in a Bethlem myopathy patient with a homozygous amino acid substitution that destabilizes the collagen VI ½(VI) chain. <i>Journal of Biological Chemistry</i> , 2015 , 290, 4272-81	5.4	20
125	Rare Variants in PLD3 Do Not Affect Risk for Early-Onset Alzheimer Disease in a European Consortium Cohort. <i>Human Mutation</i> , 2015 , 36, 1226-35	4.7	20
124	Pathogenic Huntingtin Repeat Expansions in Patients with Frontotemporal Dementia and Amyotrophic Lateral Sclerosis. <i>Neuron</i> , 2021 , 109, 448-460.e4	13.9	20
123	Genetic variability and potential effects on clinical trial outcomes: perspectives in Parkinson's disease. <i>Journal of Medical Genetics</i> , 2020 , 57, 331-338	5.8	19
122	Dynamic Atlas-Based Segmentation and Quantification of Neuromelanin-Rich Brainstem Structures in Parkinson Disease. <i>IEEE Transactions on Medical Imaging</i> , 2019 , 38, 813-823	11.7	19
121	The Genetic Architecture of Parkinson Disease in Spain: Characterizing Population-Specific Risk, Differential Haplotype Structures, and Providing Etiologic Insight. <i>Movement Disorders</i> , 2019 , 34, 1851-	1863	18
120	Association Between Vitamin D Receptor rs731236 (Taq1) Polymorphism and Risk for Restless Legs Syndrome in the Spanish Caucasian Population. <i>Medicine (United States)</i> , 2015 , 94, e2125	1.8	18
119	No association of the SLC1A2 rs3794087 allele with risk for essential tremor in the Spanish population. <i>Pharmacogenetics and Genomics</i> , 2013 , 23, 587-90	1.9	18
118	Recruitment curve of the soleus H reflex in patients with neurogenic claudication. <i>Muscle and Nerve</i> , 1998 , 21, 985-90	3.4	18
117	Heme Oxygenase-1 and 2 Common Genetic Variants and Risk for Multiple Sclerosis. <i>Scientific Reports</i> , 2016 , 6, 20830	4.9	17
116	Heme Oxygenase 1 and 2 Common Genetic Variants and Risk for Essential Tremor. <i>Medicine (United States)</i> , 2015 , 94, e968	1.8	17

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115	Selective brain gray matter atrophy associated with APOE A and MAPT H1 in subjects with mild cognitive impairment. <i>Journal of Alzheimerm Disease</i> , 2013 , 33, 1009-19	4.3	17
114	Pleural and peritoneal leishmaniasis in an AIDS patient. <i>European Journal of Clinical Microbiology and Infectious Diseases</i> , 1997 , 16, 246-8	5.3	17
113	Progressive supranuclear palsy: clinical and genetic aspects. Current Opinion in Neurology, 2002 , 15, 429	9- 3 7	17
112	Orthostatic myoclonus: an underrecognized cause of unsteadiness?. <i>Parkinsonism and Related Disorders</i> , 2013 , 19, 1013-7	3.6	16
111	Early detection of patients in the pre demented stage of Alzheimer's disease: the Pre-Al Study. <i>Journal of Nutrition, Health and Aging</i> , 2009 , 13, 21-6	5.2	16
110	Heritability and genetic variance of dementia with Lewy bodies. <i>Neurobiology of Disease</i> , 2019 , 127, 497	2- <u>5.</u> @1	15
109	LRRK2 haplotype-sharing analysis in Parkinson's disease reveals a novel p.S1761R mutation. <i>Movement Disorders</i> , 2012 , 27, 146-51	7	15
108	H1-MAPT and the risk for familial essential tremor. <i>PLoS ONE</i> , 2012 , 7, e41581	3.7	15
107	LINGO1 gene analysis in Parkinson's disease phenotypes. <i>Movement Disorders</i> , 2011 , 26, 722-7	7	15
106	Analysis of the coding and the 5' flanking regions of the alpha-synuclein gene in patients with Parkinson's disease. <i>Movement Disorders</i> , 2001 , 16, 1115-9	7	15
105	Dopamine receptor D2 intronic polymorphism in patients with Parkinson's disease. <i>Neuroscience Letters</i> , 1999 , 273, 151-4	3.3	15
104	Analysis of neurodegenerative disease-causing genes in dementia with Lewy bodies. <i>Acta Neuropathologica Communications</i> , 2020 , 8, 5	7.3	15
103	Plasma levels of soluble TREM2 and neurofilament light chain in TREM2 rare variant carriers. <i>Alzheimerm Research and Therapy</i> , 2019 , 11, 94	9	15
102	Cerebrospinal fluid levels of coenzyme Q10 are reduced in multiple system atrophy. <i>Parkinsonism and Related Disorders</i> , 2018 , 46, 16-23	3.6	15
101	SNCA and mTOR Pathway Single Nucleotide Polymorphisms Interact to Modulate the Age at Onset of Parkinson's Disease. <i>Movement Disorders</i> , 2019 , 34, 1333-1344	7	14
100	LRP10 in Bynucleinopathies. <i>Lancet Neurology, The</i> , 2018 , 17, 1032	24.1	14
99	Alpha1-antichymotrypsin gene polymorphism and susceptibility to Parkinson's disease. <i>Neurology</i> , 1999 , 52, 297-301	6.5	13
98	Role for ATXN1, ATXN2, and HTT intermediate repeats in frontotemporal dementia and Alzheimer's disease. <i>Neurobiology of Aging</i> , 2020 , 87, 139.e1-139.e7	5.6	13

97	Cerebrospinal fluid cytokines in multiple system atrophy: A cross-sectional Catalan MSA registry study. <i>Parkinsonism and Related Disorders</i> , 2019 , 65, 3-12	3.6	12
96	Rare nonsynonymous variants in SORT1 are associated with increased risk for frontotemporal dementia. <i>Neurobiology of Aging</i> , 2018 , 66, 181.e3-181.e10	5.6	12
95	Common variation in the LRRK2 gene is a risk factor for Parkinson's disease. <i>Movement Disorders</i> , 2012 , 27, 1822-5	7	12
94	A polymorphism located at an ATG transcription start site of the heme oxygenase-2 gene is associated with classical Parkinson's disease. <i>Pharmacogenetics and Genomics</i> , 2011 , 21, 565-71	1.9	12
93	Lack of interaction of SNCA and MAPT genotypes in Parkinson's disease. <i>European Journal of Neurology</i> , 2011 , 18, e32	6	12
92	Common and rare TBK1 variants in early-onset Alzheimer disease in a European cohort. Neurobiology of Aging, 2018, 62, 245.e1-245.e7	5.6	12
91	Globular glial tauopathy caused by MAPT P301T mutation: clinical and neuropathological findings. <i>Journal of Neurology</i> , 2019 , 266, 2396-2405	5.5	11
90	Copy number variation analysis of the 17q21.31 region and its role in neurodegenerative diseases. <i>American Journal of Medical Genetics Part B: Neuropsychiatric Genetics</i> , 2016 , 171B, 175-80	3.5	11
89	Pleiotropic Effects of Variants in Dementia Genes in Parkinson Disease. <i>Frontiers in Neuroscience</i> , 2018 , 12, 230	5.1	11
88	No supportive evidence for TIA1 gene mutations in a European cohort of ALS-FTD spectrum patients. <i>Neurobiology of Aging</i> , 2018 , 69, 293.e9-293.e11	5.6	11
87	association with REM sleep behavior disorder. Neurology: Genetics, 2017, 3, e131	3.8	10
86	Gamma-aminobutyric acid (GABA) receptors genes polymorphisms and risk for restless legs syndrome. <i>Pharmacogenomics Journal</i> , 2018 , 18, 565-577	3.5	10
85	Analysis of C9orf72 repeat expansions in a large international cohort of dementia with Lewy bodies. <i>Neurobiology of Aging</i> , 2017 , 49, 214.e13-214.e15	5.6	10
84	A comprehensive screening of copy number variability in dementia with Lewy bodies. <i>Neurobiology of Aging</i> , 2019 , 75, 223.e1-223.e10	5.6	10
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