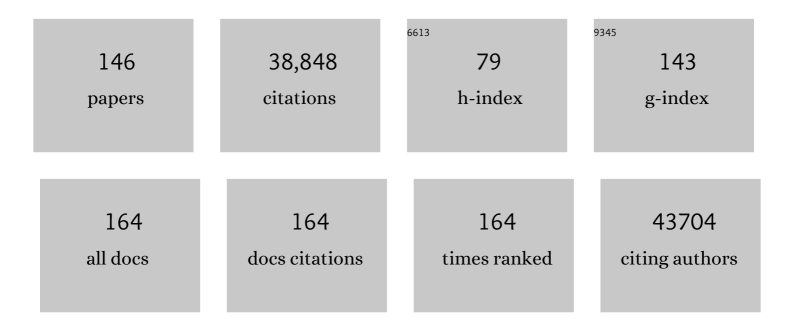
## J Paul Taylor

List of Publications by Year in descending order

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Ι ΡΛΙΙΙ ΤΛΥΙΩΡ

#	Article	IF	CITATIONS
1	Guidelines for the use and interpretation of assays for monitoring autophagy (3rd edition). Autophagy, 2016, 12, 1-222.	9.1	4,701
2	Guidelines for the use and interpretation of assays for monitoring autophagy. Autophagy, 2012, 8, 445-544.	9.1	3,122
3	Guidelines for the use and interpretation of assays for monitoring autophagy in higher eukaryotes. Autophagy, 2008, 4, 151-175.	9.1	2,064
4	Phase Separation by Low Complexity Domains Promotes Stress Granule Assembly and Drives Pathological Fibrillization. Cell, 2015, 163, 123-133.	28.9	2,053
5	Decoding ALS: from genes to mechanism. Nature, 2016, 539, 197-206.	27.8	1,533
6	Mutations in prion-like domains in hnRNPA2B1 and hnRNPA1 cause multisystem proteinopathy and ALS. Nature, 2013, 495, 467-473.	27.8	1,249
7	Toxic Proteins in Neurodegenerative Disease. Science, 2002, 296, 1991-1995.	12.6	1,103
8	Exome Sequencing Reveals VCP Mutations as a Cause of Familial ALS. Neuron, 2010, 68, 857-864.	8.1	1,100
9	HDAC6 rescues neurodegeneration and provides an essential link between autophagy and the UPS. Nature, 2007, 447, 860-864.	27.8	1,068
10	G3BP1 Is a Tunable Switch that Triggers Phase Separation to Assemble Stress Granules. Cell, 2020, 181, 325-345.e28.	28.9	697
11	Tau protein liquid–liquid phase separation can initiate tau aggregation. EMBO Journal, 2018, 37, .	7.8	696
12	GGGGCC repeat expansion in C9orf72 compromises nucleocytoplasmic transport. Nature, 2015, 525, 129-133.	27.8	692
13	HDAC6 controls autophagosome maturation essential for ubiquitin-selective quality-control autophagy. EMBO Journal, 2010, 29, 969-980.	7.8	660
14	Eukaryotic Stress Granules Are Cleared by Autophagy and Cdc48/VCP Function. Cell, 2013, 153, 1461-1474.	28.9	600
15	C9orf72 Dipeptide Repeats Impair the Assembly, Dynamics, and Function of Membrane-Less Organelles. Cell, 2016, 167, 774-788.e17.	28.9	577
16	Altered Ribostasis: RNA-Protein Granules in Degenerative Disorders. Cell, 2013, 154, 727-736.	28.9	543
17	Axonal Transport of TDP-43 mRNA Granules Is Impaired by ALS-Causing Mutations. Neuron, 2014, 81, 536-543.	8.1	521
18	Genome-wide Analyses Identify KIF5A as a Novel ALS Gene. Neuron, 2018, 97, 1268-1283.e6.	8.1	517

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19	CREB-binding protein sequestration by expanded polyglutamine. Human Molecular Genetics, 2000, 9, 2197-2202.	2.9	496
20	TIA1 Mutations in Amyotrophic Lateral Sclerosis and Frontotemporal Dementia Promote Phase Separation and Alter Stress Granule Dynamics. Neuron, 2017, 95, 808-816.e9.	8.1	493
21	Disease-causing mutations in Parkin impair mitochondrial ubiquitination, aggregation, and HDAC6-dependent mitophagy. Journal of Cell Biology, 2010, 189, 671-679.	5.2	483
22	Phase Separation of C9orf72 Dipeptide Repeats Perturbs Stress Granule Dynamics. Molecular Cell, 2017, 65, 1044-1055.e5.	9.7	437
23	Repeat expansion disease: progress and puzzles in disease pathogenesis. Nature Reviews Genetics, 2010, 11, 247-258.	16.3	425
24	Global Analysis of TDP-43 Interacting Proteins Reveals Strong Association with RNA Splicing and Translation Machinery. Journal of Proteome Research, 2010, 9, 1104-1120.	3.7	422
25	CGG Repeat-Associated Translation Mediates Neurodegeneration in Fragile X Tremor Ataxia Syndrome. Neuron, 2013, 78, 440-455.	8.1	422
26	Therapeutic reduction of ataxin-2 extends lifespan and reduces pathology in TDP-43 mice. Nature, 2017, 544, 367-371.	27.8	422
27	Mutations in the Matrin 3 gene cause familial amyotrophic lateral sclerosis. Nature Neuroscience, 2014, 17, 664-666.	14.8	398
28	VCP/p97 is essential for maturation of ubiquitin-containing autophagosomes and this function is impaired by mutations that cause IBMPFD. Autophagy, 2010, 6, 217-227.	9.1	389
29	Aggresomes protect cells by enhancing the degradation of toxic polyglutamine-containing protein. Human Molecular Genetics, 2003, 12, 749-757.	2.9	378
30	Nuclear-Import Receptors Reverse Aberrant Phase Transitions of RNA-Binding Proteins with Prion-like Domains. Cell, 2018, 173, 677-692.e20.	28.9	376
31	Measuring Friedreich ataxia: Interrater reliability of a neurologic rating scale. Neurology, 2005, 64, 1261-1262.	1.1	316
32	Stress Granule Assembly Disrupts Nucleocytoplasmic Transport. Cell, 2018, 173, 958-971.e17.	28.9	303
33	Autophagy and the ubiquitin-proteasome system: Collaborators in neuroprotection. Biochimica Et Biophysica Acta - Molecular Basis of Disease, 2008, 1782, 691-699.	3.8	302
34	TDP-43 in the Ubiquitin Pathology of Frontotemporal Dementia With VCP Gene Mutations. Journal of Neuropathology and Experimental Neurology, 2007, 66, 152-157.	1.7	295
35	Valproic acid increases SMN levels in spinal muscular atrophy patient cells. Annals of Neurology, 2003, 54, 647-654.	5.3	269
36	Ubiquitin Modulates Liquid-Liquid Phase Separation of UBQLN2 via Disruption of Multivalent Interactions. Molecular Cell, 2018, 69, 965-978.e6.	9.7	257

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37	Deficiency of ATP13A2 Leads to Lysosomal Dysfunction, α-Synuclein Accumulation, and Neurotoxicity. Journal of Neuroscience, 2012, 32, 4240-4246.	3.6	245
38	TDP-43 Mediates Degeneration in a Novel <i>Drosophila</i> Model of Disease Caused by Mutations in VCP/p97. Journal of Neuroscience, 2010, 30, 7729-7739.	3.6	243
39	Beyond aggregation: Pathological phase transitions in neurodegenerative disease. Science, 2020, 370, 56-60.	12.6	231
40	VCP Is Essential for Mitochondrial Quality Control by PINK1/Parkin and this Function Is Impaired by VCP Mutations. Neuron, 2013, 78, 65-80.	8.1	209
41	Lost in Transportation: Nucleocytoplasmic Transport Defects in ALS and Other Neurodegenerative Diseases. Neuron, 2017, 96, 285-297.	8.1	208
42	The Role of Dipeptide Repeats in C9ORF72-Related ALS-FTD. Frontiers in Molecular Neuroscience, 2017, 10, 35.	2.9	207
43	Overexpression of IGF-1 in Muscle Attenuates Disease in a Mouse Model of Spinal and Bulbar Muscular Atrophy. Neuron, 2009, 63, 316-328.	8.1	205
44	Motor neuron disease-associated loss of nuclear TDP-43 is linked to DNA double-strand break repair defects. Proceedings of the National Academy of Sciences of the United States of America, 2019, 116, 4696-4705.	7.1	203
45	Molecular heterogeneity and CXorf67 alterations in posterior fossa group A (PFA) ependymomas. Acta Neuropathologica, 2018, 136, 211-226.	7.7	199
46	Chronic optogenetic induction of stress granules is cytotoxic and reveals the evolution of ALS-FTD pathology. ELife, 2019, 8, .	6.0	184
47	Sarcoplasmic redistribution of nuclear TDPâ€43 in inclusion body myositis. Muscle and Nerve, 2009, 40, 19-31.	2.2	179
48	A Drosophila model of FUS-related neurodegeneration reveals genetic interaction between FUS and TDP-43. Human Molecular Genetics, 2011, 20, 2510-2523.	2.9	177
49	Higherâ€order oligomerization promotes localization of <scp>SPOP</scp> to liquid nuclear speckles. EMBO Journal, 2016, 35, 1254-1275.	7.8	172
50	Transgenic mice expressing mutant forms VCP/p97 recapitulate the full spectrum of IBMPFD including degeneration in muscle, brain and bone. Human Molecular Genetics, 2010, 19, 1741-1755.	2.9	171
51	TDP-43 and RNA form amyloid-like myo-granules in regenerating muscle. Nature, 2018, 563, 508-513.	27.8	163
52	Ubiquitination of G3BP1 mediates stress granule disassembly in a context-specific manner. Science, 2021, 372, eabf6548.	12.6	151
53	Native Functions of the Androgen Receptor Are Essential to Pathogenesis in a Drosophila Model of Spinobulbar Muscular Atrophy. Neuron, 2010, 67, 936-952.	8.1	150
54	Bridging biophysics and neurology: aberrant phase transitions in neurodegenerative disease. Nature Reviews Neurology, 2019, 15, 272-286.	10.1	150

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55	Protein-RNA Networks Regulated by Normal and ALS-Associated Mutant HNRNPA2B1 in the Nervous System. Neuron, 2016, 92, 780-795.	8.1	137
56	Aberrant histone acetylation, altered transcription, and retinal degeneration in a Drosophila model of polyglutamine disease are rescued by CREB-binding protein. Genes and Development, 2003, 17, 1463-1468.	5.9	130
57	C9orf72 Poly(PR) Dipeptide Repeats Disturb Biomolecular Phase Separation and Disrupt Nucleolar Function. Molecular Cell, 2019, 74, 713-728.e6.	9.7	128
58	Profilin 1 Associates with Stress Granules and ALS-Linked Mutations Alter Stress Granule Dynamics. Journal of Neuroscience, 2014, 34, 8083-8097.	3.6	126
59	Polyglutamines Placed into Context. Neuron, 2003, 38, 681-684.	8.1	124
60	ULK1 and ULK2 Regulate Stress Granule Disassembly Through Phosphorylation and Activation of VCP/p97. Molecular Cell, 2019, 74, 742-757.e8.	9.7	123
61	Cancer-associated DDX3X mutations drive stress granule assembly and impair global translation. Scientific Reports, 2016, 6, 25996.	3.3	121
62	Flightless flies: <i>Drosophila</i> models of neuromuscular disease. Annals of the New York Academy of Sciences, 2010, 1184, e1-20.	3.8	120
63	Regulatory Role of RNA Chaperone TDP-43 for RNA Misfolding and Repeat-Associated Translation in SCA31. Neuron, 2017, 94, 108-124.e7.	8.1	114
64	HDAC6 at the Intersection of Autophagy, the Ubiquitin-proteasome System, and Neurodegeneration. Autophagy, 2007, 3, 643-645.	9.1	107
65	Novel mutations expand the clinical spectrum of <i>DYNC1H1</i> -associated spinal muscular atrophy. Neurology, 2015, 84, 668-679.	1.1	106
66	Activation of expression of genes coding for extracellular matrix proteins in Tat-producing glioblastoma cells Proceedings of the National Academy of Sciences of the United States of America, 1992, 89, 9617-9621.	7.1	101
67	Exome sequencing in amyotrophic lateral sclerosis implicates a novel gene, DNAJC7, encoding a heat-shock protein. Nature Neuroscience, 2019, 22, 1966-1974.	14.8	101
68	Rescue of polyglutamine-mediated cytotoxicity by double-stranded RNA-mediated RNA interference. Human Molecular Genetics, 2002, 11, 175-184.	2.9	100
69	Disease mutations in Rab7 result in unregulated nucleotide exchange and inappropriate activation. Human Molecular Genetics, 2010, 19, 1033-1047.	2.9	99
70	Targeting protein homeostasis in sporadic inclusion body myositis. Science Translational Medicine, 2016, 8, 331ra41.	12.4	99
71	Neurotoxic microglia promote TDP-43 proteinopathy in progranulin deficiency. Nature, 2020, 588, 459-465.	27.8	98
72	Histone Deacetylases Suppress CGG Repeat–Induced Neurodegeneration Via Transcriptional Silencing in Models of Fragile X Tremor Ataxia Syndrome. PLoS Genetics, 2010, 6, e1001240.	3.5	93

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73	Linking hnRNP Function to ALS and FTD Pathology. Frontiers in Neuroscience, 2018, 12, 326.	2.8	92
74	MFN1 deacetylation activates adaptive mitochondrial fusion and protects metabolically challenged mitochondria. Journal of Cell Science, 2014, 127, 4954-63.	2.0	91
75	RAN translation at CGG repeats induces ubiquitin proteasome system impairment in models of fragile X-associated tremor ataxia syndrome. Human Molecular Genetics, 2015, 24, 4317-4326.	2.9	91
76	The Role of Autophagy in Age-Related Neurodegeneration. NeuroSignals, 2008, 16, 75-84.	0.9	89
77	Protein Arginine Methyltransferase 6 Enhances Polyglutamine-Expanded Androgen Receptor Function and Toxicity in Spinal and Bulbar Muscular Atrophy. Neuron, 2015, 85, 88-100.	8.1	89
78	Valosin-containing protein (VCP) mutations in sporadic amyotrophic lateral sclerosis. Neurobiology of Aging, 2012, 33, 2231.e1-2231.e6.	3.1	86
79	Ubiquitination is essential for recovery of cellular activities after heat shock. Science, 2021, 372, eabc3593.	12.6	86
80	The tumor suppressor protein p53 strongly alters human immunodeficiency virus type 1 replication. Journal of Virology, 1994, 68, 4302-4313.	3.4	86
81	Motor neuron involvement in multisystem proteinopathy. Neurology, 2013, 80, 1874-1880.	1.1	85
82	Multisystem proteinopathy. Neurology, 2015, 85, 658-660.	1.1	85
83	FOXO3a Is Broadly Neuroprotective In Vitro and In Vivo against Insults Implicated in Motor Neuron Diseases. Journal of Neuroscience, 2009, 29, 8236-8247.	3.6	83
84	Safety, Tolerability, and Pharmacokinetics of High-Dose Idebenone in Patients With Friedreich Ataxia. Archives of Neurology, 2007, 64, 803.	4.5	77
85	Convergence of Parkin, PINK1, and α-Synuclein on Stress-induced Mitochondrial Morphological Remodeling. Journal of Biological Chemistry, 2015, 290, 13862-13874.	3.4	76
86	TIA1 variant drives myodegeneration in multisystem proteinopathy with SQSTM1 mutations. Journal of Clinical Investigation, 2018, 128, 1164-1177.	8.2	75
87	Hsp70 dynamics in vivo: effect of heat shock and protein aggregation. Journal of Cell Science, 2004, 117, 4991-5000.	2.0	72
88	Fragile X protein mitigates TDP-43 toxicity by remodeling RNA granules and restoring translation. Human Molecular Genetics, 2015, 24, ddv389.	2.9	72
89	Evidence that a sequence similar to TAR is important for induction of the JC virus late promoter by human immunodeficiency virus type 1 Tat. Journal of Virology, 1992, 66, 7355-7361.	3.4	68
90	Interactome Mapping Provides a Network of Neurodegenerative Disease Proteins and Uncovers Widespread Protein Aggregation in Affected Brains. Cell Reports, 2020, 32, 108050.	6.4	64

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91	Selective Accumulation of Aggregation-Prone Proteasome Substrates in Response to Proteotoxic Stress. Molecular and Cellular Biology, 2009, 29, 1774-1785.	2.3	61
92	A functional deficiency of TERA/VCP/p97 contributes to impaired DNA repair in multiple polyglutamine diseases. Nature Communications, 2013, 4, 1816.	12.8	60
93	A screen for drugs that protect against the cytotoxicity of polyglutamine-expanded androgen receptor. Human Molecular Genetics, 2003, 13, 437-446.	2.9	58
94	Valosin-containing protein and the pathogenesis of frontotemporal dementia associated with inclusion body myopathy. Acta Neuropathologica, 2007, 114, 55-61.	7.7	56
95	Polyglutamine-Expanded Androgen Receptor Truncation Fragments Activate a Bax-Dependent Apoptotic Cascade Mediated by DP5/Hrk. Journal of Neuroscience, 2009, 29, 1987-1997.	3.6	56
96	High-fidelity reconstitution of stress granules and nucleoli in mammalian cellular lysate. Journal of Cell Biology, 2021, 220, .	5.2	56
97	TDP-43 suppresses CGG repeat-induced neurotoxicity through interactions with HnRNP A2/B1. Human Molecular Genetics, 2014, 23, 5036-5051.	2.9	55
98	A small-molecule Nrf1 and Nrf2 activator mitigates polyglutamine toxicity in spinal and bulbar muscular atrophy. Human Molecular Genetics, 2016, 25, 1979-1989.	2.9	55
99	Dynein light chain 1 is required for autophagy, protein clearance, and cell death in <i>Drosophila</i> . Proceedings of the National Academy of Sciences of the United States of America, 2010, 107, 742-747.	7.1	50
100	Central nervous system-derived cells express a kappa B-binding activity that enhances human immunodeficiency virus type 1 transcription in vitro and facilitates TAR-independent transactivation by Tat. Journal of Virology, 1994, 68, 3971-3981.	3.4	48
101	hnRNPDL Phase Separation Is Regulated by Alternative Splicing and Disease-Causing Mutations Accelerate Its Aggregation. Cell Reports, 2020, 30, 1117-1128.e5.	6.4	47
102	TAR-independent replication of human immunodeficiency virus type 1 in glial cells. Journal of Virology, 1992, 66, 7522-7528.	3.4	46
103	Network Analyses Reveal Novel Aspects of ALS Pathogenesis. PLoS Genetics, 2015, 11, e1005107.	3.5	45
104	Translational Repression of G3BP in Cancer and Germ Cells Suppresses Stress Granules and Enhances Stress Tolerance. Molecular Cell, 2020, 79, 645-659.e9.	9.7	40
105	Disease mutations in the prion-like domains of hnRNPA1 and hnRNPA2/B1 introduce potent steric zippers that drive excess RNP granule assembly. Rare Diseases (Austin, Tex ), 2013, 1, e25200.	1.8	38
106	Clinical and neuropathological features of ALS/FTD with TIA1 mutations. Acta Neuropathologica Communications, 2017, 5, 96.	5.2	38
107	Characterization of HNRNPA1 mutations defines diversity in pathogenic mechanisms and clinical presentation. JCI Insight, 2021, 6, .	5.0	38
108	Selective modulation of the androgen receptor AF2 domain rescues degeneration in spinal bulbar muscular atrophy. Nature Medicine, 2018, 24, 427-437.	30.7	35

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109	Mutational analysis of the VCP gene in Parkinson's disease. Neurobiology of Aging, 2012, 33, 209.e1-209.e2.	3.1	31
110	Effects of Mutations on the Aggregation Propensity of the Human Prion-Like Protein hnRNPA2B1. Molecular and Cellular Biology, 2017, 37, .	2.3	31
111	Transcription of intermediate filament genes is enhanced in focal cortical dysplasia. Acta Neuropathologica, 2001, 102, 141-148.	7.7	29
112	G-quadruplex poses quadruple threat. Nature, 2014, 507, 175-177.	27.8	29
113	TAR-Independent Activation of HIV-1 Requires the Activation Domain but Not the RNA-Binding Domain of Tat. Virology, 1993, 195, 780-785.	2.4	28
114	Archetypal and New Families With Alexander Disease and Novel Mutations in <emph type="ital"&gt;GFAP. Archives of Neurology, 2012, 69, 208.</emph 	4.5	28
115	Pediatric Cancer Variant Pathogenicity Information Exchange (PeCanPIE): a cloud-based platform for curating and classifying germline variants. Genome Research, 2019, 29, 1555-1565.	5.5	28
116	B2 attenuates polyglutamineâ€expanded androgen receptor toxicity in cell and fly models of spinal and bulbar muscular atrophy. Journal of Neuroscience Research, 2010, 88, 2207-2216.	2.9	26
117	Abnormal distribution of heterogeneous nuclear ribonucleoproteins in sporadic inclusion body myositis. Neuromuscular Disorders, 2014, 24, 611-616.	0.6	26
118	No mutations in hnRNPA1 and hnRNPA2B1 in Dutch patients with amyotrophic lateral sclerosis, frontotemporal dementia, and inclusion body myopathy. Neurobiology of Aging, 2014, 35, 1956.e9-1956.e11.	3.1	26
119	Rare Inherited forms of Paget's Disease and Related Syndromes. Calcified Tissue International, 2019, 104, 501-516.	3.1	26
120	Genetic interaction of hnRNPA2B1 and DNAJB6 in a <i>Drosophila</i> model of multisystem proteinopathy. Human Molecular Genetics, 2016, 25, 936-950.	2.9	25
121	Activation of HIV-1 transcription by Tat in cells derived from the CNS: Evidence for the participation of NF-κB — A review. Advances in Neuroimmunology, 1994, 4, 291-303.	1.8	22
122	Heterozygous frameshift variants in HNRNPA2B1 cause early-onset oculopharyngeal muscular dystrophy. Nature Communications, 2022, 13, 2306.	12.8	20
123	TDP-43 and PINK1 mediate CHCHD10S59L mutation–induced defects in Drosophila and in vitro. Nature Communications, 2021, 12, 1924.	12.8	19
124	Altered acetylation in polyglutamine disease: an opportunity for therapeutic intervention?. Trends in Molecular Medicine, 2002, 8, 195-197.	6.7	17
125	Dissection and Imaging of Active Zones in the <em>Drosophila</em> Neuromuscular Junction. Journal of Visualized Experiments, 2011, , .	0.3	15
126	Identification of compound heterozygous variants in <i>OPTN</i> in an ALS-FTD patient from the CReATe consortium: a case report. Amyotrophic Lateral Sclerosis and Frontotemporal Degeneration, 2018, 19, 469-471.	1.7	15

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127	Pathological phase transitions in ALS-FTD impair dynamic RNA–protein granules. Rna, 2022, 28, 97-113.	3.5	15
128	A Novel Conserved Isoform of the Ubiquitin Ligase UFD2a/UBE4B Is Expressed Exclusively in Mature Striated Muscle Cells. PLoS ONE, 2011, 6, e28861.	2.5	13
129	Machine learning suggests polygenic risk for cognitive dysfunction in amyotrophic lateral sclerosis. EMBO Molecular Medicine, 2021, 13, e12595.	6.9	13
130	Revealing the Mutational Spectrum in Southern Africans With Amyotrophic Lateral Sclerosis. Neurology: Genetics, 2022, 8, e654.	1.9	10
131	Hemolytic anemia presenting as idiopathic intracranial hypertension. Neurology, 2002, 59, 960-961.	1.1	9
132	RNA-binding proteins in neurological disease. Brain Research, 2012, 1462, 1-2.	2.2	9
133	RNA That Gets RAN in Neurodegeneration. Science, 2013, 339, 1282-1283.	12.6	9
134	Sexual Reassignment Fails to Prevent Kennedy's Disease. Journal of Neuromuscular Diseases, 2016, 3, 121-125.	2.6	9
135	Exome Sequencing Reveals VCP Mutations as a Cause of Familial ALS. Neuron, 2011, 69, 397.	8.1	7
136	A PR plug for the nuclear pore in amyotrophic lateral sclerosis. Proceedings of the National Academy of Sciences of the United States of America, 2017, 114, 1445-1447.	7.1	6
137	A case of familial ALS due to multi-system proteinopathy 1 and Huntington disease. Amyotrophic Lateral Sclerosis and Frontotemporal Degeneration, 2015, 16, 124-126.	1.7	5
138	Huntingtin Fragments and SOD1 Mutants Form Soluble Oligomers in the Cell. PLoS ONE, 2012, 7, e40329.	2.5	5
139	Repeat expansion and neurological disease. , 2002, , 32-54.		4
140	VCP Is Essential for Mitochondrial Quality Control by PINK1/Parkin and this Function Is Impaired by VCP Mutations. Neuron, 2013, 78, 403.	8.1	4
141	RNA metabolism in neurological disease. Brain Research, 2014, 1584, 1-2.	2.2	4
142	Genome-Wide Analyses Identify KIF5A as a Novel ALS Gene. SSRN Electronic Journal, 0, , .	0.4	4
143	The Role of Protein Disorder and Self-Association in the Formation of Cellular Bodies. Biophysical Journal, 2015, 108, 6a.	0.5	1
144	Error in Figure in: Archetypal and New Families With Alexander Disease and Novel Mutations in GFAP. Archives of Neurology, 2012, 69, 643.	4.5	0

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145	Spinal Muscular Atrophy. Neurological Disease and Therapy, 2005, , 209-226.	0.0	Ο
146	Autophagy and the Ubiquitin-Proteasome System - Protein Catabolism Comes Full Circle. , 2012, , 136-147.		0