David J Picketts

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	Generation of a mouse model of the neurodevelopmental disorder with dysmorphic facies and distal limb anomalies syndrome. Human Molecular Genetics, 2022, 31, 3405-3421.	2.9	2
2	Sensory Experience Modulates Atrx-mediated Neuronal Integrity in the Mouse Retina. Neuroscience, 2021, 452, 169-180.	2.3	1
3	Transgenic mice with an R342X mutation in <i>Phf6</i> display clinical features of Börjeson–Forssman–Lehmann Syndrome. Human Molecular Genetics, 2021, 30, 575-594.	2.9	5
4	Impaired SNF2L Chromatin Remodeling Prolongs Accessibility at Promoters Enriched for Fos/Jun Binding Sites and Delays Granule Neuron Differentiation. Frontiers in Molecular Neuroscience, 2021, 14, 680280.	2.9	1
5	ATRX affects the repair of telomeric DSBs by promoting cohesion and a DAXX-dependent activity. PLoS Biology, 2020, 18, e3000594.	5.6	46
6	Neurodevelopmental Disorders Caused by Defective Chromatin Remodeling: Phenotypic Complexity Is Highlighted by a Review of ATRX Function. Frontiers in Genetics, 2020, 11, 885.	2.3	16
7	C3aR signaling and gliosis in response to neurodevelopmental damage in the cerebellum. Journal of Neuroinflammation, 2019, 16, 135.	7.2	5
8	Snf2h Drives Chromatin Remodeling to Prime Upper Layer Cortical Neuron Development. Frontiers in Molecular Neuroscience, 2019, 12, 243.	2.9	15
9	GENE-34. THERAPEUTICALLY TARGETING EPIGENOMIC AND TRANSCRIPTIONAL DYSFUNCTION IN ATRX-DEFICIENT GLIOMA. Neuro-Oncology, 2019, 21, vi104-vi105.	1.2	1
10	The role of ISWI chromatin remodeling complexes in brain development and neurodevelopmental disorders. Molecular and Cellular Neurosciences, 2018, 87, 55-64.	2.2	52
11	Atrx inactivation drives disease-defining phenotypes in glioma cells of origin through global epigenomic remodeling. Nature Communications, 2018, 9, 1057.	12.8	66
12	Recovery from impaired muscle growth arises from prolonged postnatal accretion of myonuclei in Atrx mutant mice. PLoS ONE, 2017, 12, e0186989.	2.5	8
13	The sub-nucleolar localization of PHF6 defines its role in rDNA transcription and early processing events. European Journal of Human Genetics, 2016, 24, 1453-1459.	2.8	21
14	Voluntary Running Triggers VGF-Mediated Oligodendrogenesis to Prolong the Lifespan of Snf2h-Null Ataxic Mice. Cell Reports, 2016, 17, 862-875.	6.4	39
15	Retinal interneuron survival requires non-cell-autonomous Atrx activity. Human Molecular Genetics, 2016, 25, ddw306.	2.9	3
16	A Notch-Gli2 axis sustains Hedgehog responsiveness of neural progenitors and Müller glia. Developmental Biology, 2016, 411, 85-100.	2.0	31
17	Forebrain neurogenesis: From embryo to adult. Trends in Developmental Biology, 2016, 9, 77-90.	1.0	3
18	PHF6 Degrees of Separation: The Multifaceted Roles of a Chromatin Adaptor Protein. Genes, 2015, 6, 325-352.	2.4	38

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19	Genome-wide characterisation of Foxa1 binding sites reveals several mechanisms for regulating neuronal differentiation in midbrain dopamine cells. Development (Cambridge), 2015, 142, 1315-1324.	2.5	25
20	Snf2h-mediated chromatin organization and histone H1 dynamics govern cerebellar morphogenesis and neural maturation. Nature Communications, 2014, 5, 4181.	12.8	71
21	Snf2l Regulates Foxg1-Dependent Progenitor Cell Expansion in the Developing Brain. Developmental Cell, 2012, 22, 871-878.	7.0	60
22	Loss of Atrx Sensitizes Cells to DNA Damaging Agents through p53-Mediated Death Pathways. PLoS ONE, 2012, 7, e52167.	2.5	31
23	PHF6 Interacts with the Nucleosome Remodeling and Deacetylation (NuRD) Complex. Journal of Proteome Research, 2012, 11, 4326-4337.	3.7	87
24	Compromised genomic integrity impedes muscle growth after Atrx inactivation. Journal of Clinical Investigation, 2012, 122, 4412-4423.	8.2	57
25	Matters of life and death: the role of chromatin remodeling proteins in retinal neuron survival. Journal of Ocular Biology, Diseases, and Informatics, 2011, 4, 111-120.	0.2	16
26	Tâ€cell acute lymphoblastic leukemia in association with Börjeson–Forssman–Lehmann syndrome due to a mutation in <i>PHF6</i> . Pediatric Blood and Cancer, 2010, 55, 722-724.	1.5	42
27	SCO-ping Out the Mechanisms Underlying the Etiology of Hydrocephalus. Physiology, 2009, 24, 117-126.	3.1	47
28	Altered visual function and interneuron survival in Atrx knockout mice: inference for the human syndrome. Human Molecular Genetics, 2009, 18, 966-977.	2.9	34
29	Börjeson-Forssman-Lehmann Syndrome Due to a Novel Plant Homeodomain Zinc Finger Mutation in the PHF6 Gene. Journal of Child Neurology, 2009, 24, 610-614.	1.4	17
30	Further clinical delineation of the Börjeson–Forssman–Lehmann syndrome in patients with <i>PHF6</i> mutations. American Journal of Medical Genetics, Part A, 2009, 149A, 246-250.	1.2	27
31	Characterization of novel isoforms and evaluation of SNF2L/SMARCA1 as a candidate gene for X-linked mental retardation in 12 families linked to Xq25-26. BMC Medical Genetics, 2008, 9, 11.	2.1	11
32	Patient mutations alter ATRX targeting to PML nuclear bodies. European Journal of Human Genetics, 2008, 16, 192-201.	2.8	40
33	The Imitation Switch Protein SNF2L Regulates Steroidogenic Acute Regulatory Protein Expression during Terminal Differentiation of Ovarian Granulosa Cells. Molecular Endocrinology, 2006, 20, 2406-2417.	3.7	41
34	Neuropeptide signaling and hydrocephalus: SCO with the flow. Journal of Clinical Investigation, 2006, 116, 1828-1832.	8.2	17
35	The chromatin-remodeling protein ATRX is critical for neuronal survival during corticogenesis. Journal of Clinical Investigation, 2005, 115, 258-267.	8.2	169
36	The chromatin-remodeling protein ATRX is critical for neuronal survival during corticogenesis. Journal of Clinical Investigation, 2005, 115, 258-267.	8.2	119

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37	A Tissue-specific, Naturally Occurring Human SNF2L Variant Inactivates Chromatin Remodeling. Journal of Biological Chemistry, 2004, 279, 45130-45138.	3.4	31
38	A Novel Transcription Regulatory Complex Containing Death Domain-associated Protein and the ATR-X Syndrome Protein. Journal of Biological Chemistry, 2004, 279, 20369-20377.	3.4	168
39	Isolation of human NURF: a regulator of Engrailed gene expression. EMBO Journal, 2003, 22, 6089-6100.	7.8	151
40	Increasing D4Z4 repeat copy number compromises C2C12 myoblast differentiation. FEBS Letters, 2003, 537, 133-138.	2.8	27
41	Neurodevelopmental defects resulting from ATRX overexpression in transgenic mice. Human Molecular Genetics, 2002, 11, 253-261.	2.9	58
42	Cloning and characterization of the murine Imitation Switch (ISWI) genes: differential expression patterns suggest distinct developmental roles for Snf2h and Snf2l. Journal of Neurochemistry, 2001, 77, 1145-1156.	3.9	95
43	Cell cycle-dependent phosphorylation of the ATRX protein correlates with changes in nuclear matrix and chromatin association. Human Molecular Genetics, 2000, 9, 539-547.	2.9	139
44	Mutations in a putative global transcriptional regulator cause X-linked mental retardation with α-thalassemia (ATR-X syndrome). Cell, 1995, 80, 837-845.	28.9	583