Conrad C Weihl

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/4223278/publications.pdf

Version: 2024-02-01

70 papers 9,258 citations

108046 37 h-index 93651 72 g-index

78 all docs 78 docs citations

78 times ranked 18161 citing authors

| # | Article | IF | CITATIONS |
|----|--|------|-----------|
| 1 | Current and Future Approaches to Classify VUSs in LGMD-Related Genes. Genes, 2022, 13, 382. | 1.0 | 3 |
| 2 | Randomized phase 2 study of <scp>ACE</scp> â€083, a <scp>muscleâ€promoting</scp> agent, in facioscapulohumeral muscular dystrophy. Muscle and Nerve, 2022, 66, 50-62. | 1.0 | 8 |
| 3 | VCP suppresses proteopathic seeding in neurons. Molecular Neurodegeneration, 2022, 17, 30. | 4.4 | 15 |
| 4 | Progression to Loss of Ambulation Among Patients with Autosomal Recessive Limb-girdle Muscular Dystrophy: A Systematic Review. Journal of Neuromuscular Diseases, 2022, , 1-16. | 1.1 | 3 |
| 5 | Multisystem Proteinopathy Due to VCP Mutations: A Review of Clinical Heterogeneity and Genetic Diagnosis. Genes, 2022, 13, 963. | 1.0 | 28 |
| 6 | Clinical utility of antiâ€cytosolic 5'â€nucleotidase 1A antibody in idiopathic inflammatory myopathies. Annals of Clinical and Translational Neurology, 2021, 8, 571-578. | 1.7 | 18 |
| 7 | Neuronal VCP loss of function recapitulates FTLD-TDP pathology. Cell Reports, 2021, 36, 109399. | 2.9 | 25 |
| 8 | Molecular and cellular basis of genetically inherited skeletal muscle disorders. Nature Reviews Molecular Cell Biology, 2021, 22, 713-732. | 16.1 | 55 |
| 9 | Loss-of-function mutation in VCP mimics the characteristic pathology as in FTLD-TARDBP. Autophagy, 2021, 17, 4502-4503. | 4.3 | 5 |
| 10 | Mutations in the J domain of DNAJB6 cause dominant distal myopathy. Neuromuscular Disorders, 2020, 30, 38-46. | 0.3 | 20 |
| 11 | A cross-sectional study of hand function in inclusion body myositis: Implications for functional rating scale. Neuromuscular Disorders, 2020, 30, 200-206. | 0.3 | 4 |
| 12 | Optimizing hand-function patient outcome measures for inclusion body myositis. Neuromuscular Disorders, 2020, 30, 807-814. | 0.3 | 5 |
| 13 | Client processing is altered by novel myopathy-causing mutations in the HSP40 J domain. PLoS ONE, 2020, 15, e0234207. | 1.1 | 6 |
| 14 | High-protein diets increase cardiovascular risk by activating macrophage mTOR to suppress mitophagy. Nature Metabolism, 2020, 2, 110-125. | 5.1 | 85 |
| 15 | Inhibition of DNAJ-HSP70 interaction improves strength in muscular dystrophy. Journal of Clinical Investigation, 2020, 130, 4470-4485. | 3.9 | 14 |
| 16 | Phenotypic diversity in an international Cure VCP Disease registry. Orphanet Journal of Rare Diseases, 2020, 15, 267. | 1.2 | 11 |
| 17 | Loss of lipin 1â€mediated phosphatidic acid phosphohydrolase activity in muscle leads to skeletal myopathy in mice. FASEB Journal, 2019, 33, 652-667. | 0.2 | 30 |
| 18 | Desmin forms toxic, seeding-competent amyloid aggregates that persist in muscle fibers. Proceedings of the National Academy of Sciences of the United States of America, 2019, 116, 16835-16840. | 3.3 | 36 |

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|----|--|-----|-----------|
| 19 | Limbâ€girdle muscular dystrophy: A perspective from adult patients on what matters most. Muscle and Nerve, 2019, 60, 419-424. | 1.0 | 15 |
| 20 | VCP maintains lysosomal homeostasis and TFEB activity in differentiated skeletal muscle. Autophagy, 2019, 15, 1082-1099. | 4.3 | 44 |
| 21 | Estimating prevalence for limb-girdle muscular dystrophy based on public sequencing databases. Genetics in Medicine, 2019, 21, 2512-2520. | 1.1 | 56 |
| 22 | Lithium chloride corrects weakness and myopathology in a preclinical model of LGMD1D. Neurology: Genetics, 2019, 5, e318. | 0.9 | 15 |
| 23 | A metastable subproteome underlies inclusion formation in muscle proteinopathies. Acta Neuropathologica Communications, 2019, 7, 197. | 2.4 | 16 |
| 24 | Sporadic Inclusion Body Myositis and Other Rimmed Vacuolar Myopathies. CONTINUUM Lifelong Learning in Neurology, 2019, 25, 1586-1598. | 0.4 | 6 |
| 25 | CANOMAD and other chronic ataxic neuropathies with disialosyl antibodies (CANDA). Journal of Neurology, 2018, 265, 1402-1409. | 1.8 | 40 |
| 26 | Myopathy associated BAG3 mutations lead to protein aggregation by stalling Hsp70 networks. Nature Communications, 2018, 9, 5342. | 5.8 | 65 |
| 27 | Homozygous recessive MYH2 mutation mimicking dominant MYH2 associated myopathy. Neuromuscular Disorders, 2018, 28, 675-679. | 0.3 | 10 |
| 28 | TIA1 variant drives myodegeneration in multisystem proteinopathy with SQSTM1 mutations. Journal of Clinical Investigation, 2018, 128, 1164-1177. | 3.9 | 75 |
| 29 | Establishing prevalence in rare neuromuscular diseases. Neurology: Genetics, 2017, 3, e146. | 0.9 | 1 |
| 30 | Quantification of autophagy flux using LC3 ELISA. Analytical Biochemistry, 2017, 530, 57-67. | 1.1 | 43 |
| 31 | Cystinosis distal myopathy, novel clinical, pathological and genetic features. Neuromuscular Disorders, 2017, 27, 873-878. | 0.3 | 5 |
| 32 | Exploiting macrophage autophagy-lysosomal biogenesis as a therapy for atherosclerosis. Nature Communications, 2017, 8, 15750. | 5.8 | 258 |
| 33 | Keap1/Cullin3 Modulates p62/SQSTM1 Activity via UBA Domain Ubiquitination. Cell Reports, 2017, 19, 188-202. | 2.9 | 110 |
| 34 | Proteomics of rimmed vacuoles define new risk allele in inclusion body myositis. Annals of Neurology, 2017, 81, 227-239. | 2.8 | 59 |
| 35 | Comparisons of ELISA and Western blot assays for detection of autophagy flux. Data in Brief, 2017, 13, 696-699. | 0.5 | 8 |
| 36 | Regulation of SQSTM1/p62 via UBA domain ubiquitination and its role in disease. Autophagy, 2017, 13, $1615-1616$. | 4.3 | 47 |

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|----|---|------|-----------|
| 37 | <scp>VCP</scp> /p97 cooperates with <scp>YOD</scp> 1, <scp>UBXD</scp> 1 and <scp>PLAA</scp> to drive clearance of ruptured lysosomes by autophagy. EMBO Journal, 2017, 36, 135-150. | 3.5 | 259 |
| 38 | 215th ENMC International Workshop VCP-related multi-system proteinopathy (IBMPFD) 13–15 November 2015, Heemskerk, The Netherlands. Neuromuscular Disorders, 2016, 26, 535-547. | 0.3 | 38 |
| 39 | Rare Manifestation of a c.290 C>T, p.Gly97Glu <i>VCP</i> Mutation. Case Reports in Genetics, 2015, 2015, 1-5. | 0.1 | 16 |
| 40 | Autophagic vacuolar pathology in desminopathies. Neuromuscular Disorders, 2015, 25, 199-206. | 0.3 | 19 |
| 41 | Targeted sequencing and identification of genetic variants in sporadic inclusion body myositis. Neuromuscular Disorders, 2015, 25, 289-296. | 0.3 | 56 |
| 42 | Altered cofactor regulation with disease-associated p97/VCP mutations. Proceedings of the National Academy of Sciences of the United States of America, 2015, 112, E1705-14. | 3.3 | 87 |
| 43 | <i>SQSTM1</i> splice site mutation in distal myopathy with rimmed vacuoles. Neurology, 2015, 85, 665-674. | 1.5 | 74 |
| 44 | Myofibrillar disruption and RNA-binding protein aggregation in a mouse model of limb-girdle muscular dystrophy 1D. Human Molecular Genetics, 2015, 24, 6588-6602. | 1.4 | 31 |
| 45 | Myopathy-causing Mutations in an HSP40 Chaperone Disrupt Processing of Specific Client Conformers. Journal of Biological Chemistry, 2014, 289, 21120-21130. | 1.6 | 46 |
| 46 | Prion-like nuclear aggregation of TDP-43 during heat shock is regulated by HSP40/70 chaperones. Human Molecular Genetics, 2014, 23, 157-170. | 1.4 | 102 |
| 47 | Mutations in the Matrin 3 gene cause familial amyotrophic lateral sclerosis. Nature Neuroscience, 2014, 17, 664-666. | 7.1 | 398 |
| 48 | The VCP/p97 system at a glance: connecting cellular function to disease pathogenesis. Journal of Cell Science, 2014, 127, 3877-83. | 1.2 | 329 |
| 49 | A novel mutation in VCP causes Charcot–Marie–Tooth Type 2 disease. Brain, 2014, 137, 2897-2902. | 3.7 | 116 |
| 50 | Specific Inhibition of p97/VCP ATPase and Kinetic Analysis Demonstrate Interaction between D1 and D2 ATPase Domains. Journal of Molecular Biology, 2014, 426, 2886-2899. | 2.0 | 103 |
| 51 | Monitoring Autophagy in the Treatment of Protein Aggregate Diseases: Steps Toward Identifying Autophagic Biomarkers. Neurotherapeutics, 2013, 10, 383-390. | 2.1 | 11 |
| 52 | Mutations in prion-like domains in hnRNPA2B1 and hnRNPA1 cause multisystem proteinopathy and ALS. Nature, 2013, 495, 467-473. | 13.7 | 1,249 |
| 53 | Motor neuron involvement in multisystem proteinopathy. Neurology, 2013, 80, 1874-1880. | 1.5 | 85 |
| 54 | mTOR dysfunction contributes to vacuolar pathology and weakness in valosin-containing protein associated inclusion body myopathy. Human Molecular Genetics, 2013, 22, 1167-1179. | 1.4 | 58 |

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| 55 | Increased autophagy accelerates colchicine-induced muscle toxicity. Autophagy, 2013, 9, 2115-2125. | 4.3 | 35 |
| 56 | Guidelines for the use and interpretation of assays for monitoring autophagy. Autophagy, 2012, 8, 445-544. | 4.3 | 3,122 |
| 57 | Exome sequencing reveals <i>DNAJB6</i> mutations in dominantlyâ€inherited myopathy. Annals of Neurology, 2012, 71, 407-416. | 2.8 | 148 |
| 58 | Mdx mice have a defect in autophagy that is restored by rapamycinâ€loaded nanoparticle treatment. FASEB Journal, 2012, 26, 396.5. | 0.2 | 1 |
| 59 | Novel GNE mutations in two phenotypically distinct HIBM2 patients. Neuromuscular Disorders, 2011, 21, 102-105. | 0.3 | 15 |
| 60 | Endolysosomal sorting of ubiquitylated caveolin-1 is regulated by VCP and UBXD1 and impaired by VCP disease mutations. Nature Cell Biology, 2011, 13, 1116-1123. | 4.6 | 213 |
| 61 | Another VCP interactor: NF is enough. Journal of Clinical Investigation, 2011, 121, 4627-4630. | 3.9 | 7 |
| 62 | Sporadic inclusion body myositis: possible pathogenesis inferred from biomarkers. Current Opinion in Neurology, 2010, 23, 482-488. | 1.8 | 47 |
| 63 | Interaction with Polyglutamine Aggregates Reveals a Q/N-rich Domain in TDP-43. Journal of Biological Chemistry, 2010, 285, 26304-26314. | 1.6 | 138 |
| 64 | Valosin-containing protein (VCP) is required for autophagy and is disrupted in VCP disease. Journal of Cell Biology, 2009, 187, 875-888. | 2.3 | 444 |
| 65 | Valosin-containing protein disease: Inclusion body myopathy with Paget's disease of the bone and fronto-temporal dementia. Neuromuscular Disorders, 2009, 19, 308-315. | 0.3 | 205 |
| 66 | Inflammatory myopathies with mitochondrial pathology and protein aggregates. Journal of the Neurological Sciences, 2009, 278, 25-29. | 0.3 | 91 |
| 67 | Transgenic expression of inclusion body myopathy associated mutant p97/VCP causes weakness and ubiquitinated protein inclusions in mice. Human Molecular Genetics, 2007, 16, 919-928. | 1.4 | 99 |
| 68 | Motor neuron disease associated with copper deficiency. Muscle and Nerve, 2006, 34, 789-793. | 1.0 | 76 |
| 69 | Valproate may improve strength and function in patients with type III/IV spinal muscle atrophy. Neurology, 2006, 67, 500-501. | 1.5 | 111 |
| 70 | Inclusion body myopathy-associated mutations in p97/VCP impair endoplasmic reticulum-associated degradation. Human Molecular Genetics, 2006, 15, 189-199. | 1.4 | 169 |