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List of Publications by Year in descending order

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69 papers 1,126 citations

393982 19 h-index 454577 30 g-index

74 all docs

74 docs citations

74 times ranked 1484 citing authors

#	Article	IF	CITATIONS
1	An Exploratory Survey on the Care for Ataxic Patients in the American Continents and the Caribbean. Cerebellum, 2023, 22, 708-718.	1.4	1
2	Genetic Distribution of Five Spinocerebellar Ataxia Microsatellite Loci in Mexican Native American Populations and Its Impact on Contemporary Mestizo Populations. Genes, 2022, 13, 157.	1.0	1
3	ACE and ACE2 Gene Variants Are Associated With Severe Outcomes of COVID-19 in Men. Frontiers in Immunology, 2022, 13, 812940.	2.2	31
4	Cognitive Decline and White Matter Integrity Degradation in Myotonic Dystrophy Type I. Journal of Neuroimaging, 2021, 31, 192-198.	1.0	7
5	Increased risk of depression and impairment in quality of life in patients with lamellar ichthyosis. Dermatologic Therapy, 2021, 34, e14628.	0.8	7
6	Transcriptome Analysis Reveals Altered Inflammatory Pathway in an Inducible Glial Cell Model of Myotonic Dystrophy Type $1.$ Biomolecules, $2021,11,159.$	1.8	12
7	Association of TLR4 gene polymorphisms with sepsis after a burn injury: findings of the functional role of rs2737190 SNP. Genes and Immunity, 2021, 22, 24-34.	2.2	4
8	Non-lonic Surfactants for Stabilization of Polymeric Nanoparticles for Biomedical Uses. Materials, 2021, 14, 3197.	1.3	81
9	New Perspectives of Gene Therapy on Polyglutamine Spinocerebellar Ataxias: From Molecular Targets to Novel Nanovectors. Pharmaceutics, 2021, 13, 1018.	2.0	6
10	Therapeutic Applications of Terpenes on Inflammatory Diseases. Frontiers in Pharmacology, 2021, 12, 704197.	1.6	40
11	Nanoremediation: Nanomaterials and Nanotechnologies for Environmental Cleanup. Frontiers in Environmental Science, 2021, 9, .	1.5	30
12	Non-invasive methods for evaluation of skin manifestations in patients with ichthyosis. Archives of Dermatological Research, 2020, 312, 231-236.	1.1	6
13	Loss of Dystroglycan Drives Cellular Senescence via Defective Mitosis-Mediated Genomic Instability. International Journal of Molecular Sciences, 2020, 21, 4961.	1.8	8
14	Coexistence of Fragile-X Syndrome, 8p23.1 Deletion, and Balanced Translocation t(7;10)(p10;q24) in a Single Family. Genetic Testing and Molecular Biomarkers, 2020, 24, 527-531.	0.3	0
15	The Molecular Basis and Biologic Significance of the \hat{I}^2 -Dystroglycan-Emerin Interaction. International Journal of Molecular Sciences, 2020, 21, 5944.	1.8	5
16	Association of Genetic Polymorphisms in TLR3, TLR4, TLR7, and TLR8 with the Clinical Forms of Dengue in Patients from Veracruz, Mexico. Viruses, 2020, 12, 1230.	1.5	7
17	Controlled Transdermal Release of Antioxidant Ferulate by a Porous Sc(III) MOF. IScience, 2020, 23, 101156.	1.9	16
18	High prevalence of autosomal recessive congenital ichthyosis in a Mexican population caused by a new mutation in the TGM1 gene: epidemiological evidence of a founder effect. International Journal of Dermatology, 2020, 59, 969-977.	0.5	8

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19	<i>ALOX5</i> , <i>LPA</i> , <i>MMP9</i> and <i>TPO</i> gene polymorphisms increase atherothrombosis susceptibility in middle-aged Mexicans. Royal Society Open Science, 2020, 7, 190775.	1.1	2
20	Altered Plasma Acylcarnitines and Amino Acids Profile in Spinocerebellar Ataxia Type 7. Biomolecules, 2020, 10, 390.	1.8	8
21	Effect of UV and Gamma Irradiation Sterilization Processes in the Properties of Different Polymeric Nanoparticles for Biomedical Applications. Materials, 2020, 13, 1090.	1.3	35
22	The Need for Establishing a Universal CTG Sizing Method in Myotonic Dystrophy Type 1. Genes, 2020, 11, 757.	1.0	5
23	Founder Effects of Spinocerebellar Ataxias in the American Continents and the Caribbean. Cerebellum, 2020, 19, 446-458.	1.4	17
24	Involvement of the Auditory Pathway in Spinocerebellar Ataxia Type 7. Neurodegenerative Diseases, 2020, 20, 185-192.	0.8	2
25	Hyaluronic acid in wound dressings. Cellular and Molecular Biology, 2020, 66, 191-198.	0.3	39
26	From traditional biochemical signals to molecular markers for detection of sepsis after burn injuries. Burns, 2019, 45, 16-31.	1.1	27
27	Development and Evaluation of Alginate Membranes with Curcumin-Loaded Nanoparticles for Potential Wound-Healing Applications. Pharmaceutics, 2019, 11, 389.	2.0	36
28	Enhanced nuclear protein export in premature aging and rescue of the progeria phenotype by modulation of CRM1 activity. Aging Cell, 2019, 18, e13002.	3.0	19
29	Effects of Physical Rehabilitation in Patients with Spinocerebellar Ataxia Type 7. Cerebellum, 2019, 18, 397-405.	1.4	23
30	Wide Profiling of Circulating MicroRNAs in Spinocerebellar Ataxia Type 7. Molecular Neurobiology, 2019, 56, 6106-6120.	1.9	12
31	InÂvitro cell uptake evaluation of curcumin-loaded PCL/F68 nanoparticles for potential application in neuronal diseases. Journal of Drug Delivery Science and Technology, 2019, 52, 905-914.	1.4	33
32	Oropharyngeal dysphagia in early stages of myotonic dystrophy type 1. Muscle and Nerve, 2019, 60, 90-95.	1.0	3
33	New copolymers as hosts of ribosomal RNA. BMC Chemistry, 2019, 13, 33.	1.6	4
34	The intracellular domain of \hat{l}^2 -dystroglycan mediates the nucleolar stress response by suppressing UBF transcriptional activity. Cell Death and Disease, 2019, 10, 196.	2.7	3
35	Nonâ€invasive analysis of skin mechanical properties in patients with lamellar ichthyosis. Skin Research and Technology, 2019, 25, 375-381.	0.8	8
36	Comprehensive mapping of human body skin hydration: A pilot study. Skin Research and Technology, 2019, 25, 187-193.	0.8	7

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37	Oxidative Stress in Spinocerebellar Ataxia Type 7 Is Associated with Disease Severity. Cerebellum, 2018, 17, 601-609.	1.4	22
38	Voice Alterations in Patients With Spinocerebellar Ataxia Type 7 (SCA7): Clinical-Genetic Correlations. Journal of Voice, 2017, 31, 123.e1-123.e5.	0.6	6
39	Nanoparticulate strategies for the treatment of polyglutamine diseases by halting the protein aggregation process. Drug Development and Industrial Pharmacy, 2017, 43, 871-888.	0.9	15
40	A Complete Association of an intronic SNP rs6798742 with Origin of Spinocerebellar Ataxia Type 7 AG Expansion Loci in the Indian and Mexican Population. Annals of Human Genetics, 2017, 81, 197-204.	0.3	6
41	Origin of the myotonic dystrophy type 1 mutation in Mexican population and influence of Amerindian ancestry on CTG repeat allelic distribution. Neuromuscular Disorders, 2017, 27, 1106-1114.	0.3	4
42	Dysferlin quantification in monocytes for rapid screening for dysferlinopathies. Muscle and Nerve, 2016, 54, 1064-1071.	1.0	2
43	Verification of Inter-laboratorial Genotyping Consistency in the Molecular Diagnosis of Polyglutamine Spinocerebellar Ataxias. Journal of Molecular Neuroscience, 2016, 58, 83-87.	1.1	3
44	Interethnic variation of the MMP-9 microsatellite in Amerindian and Mexican Mestizo populations: considerations for genetic association studies. Genetics and Molecular Research, 2015, 14, 2929-2939.	0.3	4
45	Dopaminergic denervation switches dopamine D3 receptor signaling and disrupts its Ca2+ dependent modulation by CaMKII and calmodulin in striatonigral projections of the rat. Neurobiology of Disease, 2015, 74, 336-346.	2.1	14
46	Spinocerebellar Ataxia Type 7: A Neurodegenerative Disorder with Peripheral Neuropathy. European Neurology, 2015, 73, 173-178.	0.6	11
47	Nanoparticle technology for treatment of Parkinson's disease: the role of surface phenomena in reaching the brain. Drug Discovery Today, 2015, 20, 824-837.	3.2	77
48	A comprehensive clinical and genetic study of a large Mexican population with spinocerebellar ataxia type 7. Neurogenetics, 2015, 16, 11-21.	0.7	28
49	Altered nuclear structure in myotonic dystrophy type 1-derived fibroblasts. Molecular Biology Reports, 2015, 42, 479-488.	1.0	19
50	Association of vWA and TPOX Polymorphisms with Venous Thrombosis in Mexican Mestizos. BioMed Research International, 2014, 2014, 1-9.	0.9	8
51	Genetic Analysis of 17 Y-STRs in a Mestizo Population from the Central Valley of Mexico. Human Biology, 2014, 86, 289.	0.4	13
52	Recessive Spinocerebellar Ataxia with Paroxysmal Cough Attacks: A Report of Five Cases. Cerebellum, 2014, 13, 215-221.	1.4	1
53	Analysis of <scp>CAG</scp> repeats in five <scp>SCA</scp> loci in Mexican population: epidemiological evidence of a <scp>SCA7</scp> founder effect. Clinical Genetics, 2014, 85, 159-165.	1.0	27
54	Comprehensive Study of Early Features in Spinocerebellar Ataxia 2: Delineating the Prodromal Stage of the Disease. Cerebellum, 2014, 13, 568-579.	1.4	51

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55	Clinical and molecular effect on offspring of a marriage of consanguineous spinocerebellar ataxia type 7 mutation carriers: a family case report. International Journal of Clinical and Experimental Medicine, 2014, 7, 5896-903.	1.3	1
56	Origin of the Spinocerebellar Ataxia Type 7 Gene Mutation in Mexican Population. Cerebellum, 2013, 12, 902-905.	1.4	23
57	The association of single nucleotide polymorphisms in the calcitonin gene with primary osteoarthritis of the knee in Mexican mestizo population. Rheumatology International, 2013, 33, 2483-2491.	1.5	3
58	Spinocerebellar Ataxia Type 2: Clinical Presentation, Molecular Mechanisms, and Therapeutic Perspectives. Molecular Neurobiology, 2013, 47, 90-104.	1.9	83
59	P.18.11 Nerve conduction alterations in Mexican patients with myotonic dystrophy type 1. Neuromuscular Disorders, 2013, 23, 835.	0.3	0
60	The 46359CT polymorphism of DNMT3B is associated with the risk of cervical cancer. Molecular Biology Reports, 2013, 40, 4275-4280.	1.0	8
61	The relationship among IL-13, GSTP1, and CYP1A1 polymorphisms and environmental tobacco smoke in a population of children with asthma in Northern Mexico. Environmental Toxicology and Pharmacology, 2012, 33, 226-232.	2.0	17
62	Mexican mestizo population sub-structure: effects on genetic and forensic statistical parameters. Molecular Biology Reports, 2012, 39, 10139-10156.	1.0	16
63	Distribution of CTG repeats at the DMPK gene in myotonic distrophy patients and healthy individuals from the Mexican population. Molecular Biology Reports, 2011, 38, 1341-1346.	1.0	9
64	Perspectives on gene therapy in myotonic dystrophy type 1. Journal of Neuroscience Research, 2011, 89, 275-285.	1.3	24
65	Association of the calcitonin gene (CA) polymorphism with osteoarthritis of the knee in a Mexican mestizo population. Knee, 2010, 17, 157-160.	0.8	18
66	La electroforesis capilar como una nueva estrategia en la medicina y el diagnóstico clÃnico. Revista Medica De Chile, 2009, 137, .	0.1	0
67	Association of Interleukin-6 Gene Polymorphisms with Bone Mineral Density in Mexican Women. Archives of Medical Research, 2008, 39, 618-624.	1.5	20
68	Association of the estrogen receptor \hat{l}_{\pm} gene polymorphisms with osteoporosis in the Mexican population. Clinical Genetics, 2007, 72, 574-581.	1.0	23
69	Association of the CT gene (CA) polymorphism with BMD in osteoporotic mexican women. Clinical Genetics, 2006, 70, 402-408.	1.0	11