Mauricio Arcos-Burgos

List of Publications by Year in descending order

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46799 57758 8,481 127 44 89 citations h-index g-index papers 133 133 133 9612 docs citations times ranked citing authors all docs

#	Article	IF	CITATIONS
1	Machine Learning Prediction of ADHD Severity: Association and Linkage to <i>ADGRL3</i> , <i>DRD4</i> , and <i>SNAP25</i> . Journal of Attention Disorders, 2022, 26, 587-605.	2.6	11
2	Executive function deficit in bipolar offspring: A neurocognitive endophenotype?. Journal of Affective Disorders, 2022, 297, 246-249.	4.1	2
3	Human Genetic Host Factors and Its Role in the Pathogenesis of Chikungunya Virus Infection. Frontiers in Medicine, 2022, 9, 654395.	2.6	2
4	Ataxia with Ocular Apraxia Type 1 (AOA1) (APTX, W279* Mutation): Neurological, Neuropsychological, and Molecular Outlining of a Heterogenous Phenotype in Four Colombian Siblings. Molecular Neurobiology, 2022, , 1.	4.0	0
5	MejorÃa del ACV con estimulación magnética recapitula la ontogenia. Acta Medica Colombiana: AMC: Organo De La Asociacion Columbiana De Medicina Interna, 2022, 47, .	0.0	O
6	Structural Protein Effects Underpinning Cognitive Developmental Delay of the PURA p.Phe233del Mutation Modelled by Artificial Intelligence and the Hybrid Quantum Mechanics–Molecular Mechanics Framework. Brain Sciences, 2022, 12, 871.	2.3	4
7	Frequency of actionable Exomic secondary findings in 160 Colombian patients: Impact in the healthcare system. Gene, 2022, 838, 146699.	2.2	1
8	A multifactorial model of pathology for age of onset heterogeneity in familial Alzheimer's disease. Acta Neuropathologica, 2021, 141, 217-233.	7.7	33
9	Rare Functional Variants Associated with Antidepressant Remission in Mexican-Americans. Journal of Affective Disorders, 2021, 279, 491-500.	4.1	3
10	The role of psychosocial adversity in the aetiology and course of attention deficit hyperactivity disorder. Revista Colombiana De Psiquiatr $ ilde{A}$ a, 2021, , .	0.3	1
11	A Comprehensive Machine Learning Framework for the Exact Prediction of the Age of Onset in Familial and Sporadic Alzheimer's Disease. Diagnostics, 2021, 11, 887.	2.6	2
12	ADGRL3, FGF1 and DRD4: Linkage and Association with Working Memory and Perceptual Organization Candidate Endophenotypes in ADHD. Brain Sciences, 2021, 11, 854.	2.3	4
13	Psychopathological Risk in Siblings of Subjects with Attention-Deficit/Hyperactivity Disorder: A cross-Sectional Study. Revista Colombiana De PsiquiatrÃa, 2021, , .	0.3	O
14	Utility of a Short Neuropsychological Protocol for Detecting HIV-Associated Neurocognitive Disorders in Patients with Asymptomatic HIV-1 Infection. Brain Sciences, 2021, 11, 1037.	2.3	2
15	Impulsive and Omission Errors: Potential Temporal Processing Endophenotypes in ADHD. Brain Sciences, 2021, 11, 1218.	2.3	4
16	ADHD Endophenotypes in Caribbean Families. Journal of Attention Disorders, 2020, 24, 2100-2114.	2.6	7
17	Familial Alzheimer's Disease and Recessive Modifiers. Molecular Neurobiology, 2020, 57, 1035-1043.	4.0	7
18	Mutations in sphingolipid metabolism genes are associated with ADHD. Translational Psychiatry, 2020, 10, 231.	4.8	7

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19	Chikungunya outbreak (2015) in the Colombian Caribbean: Latent classes and gender differences in virus infection. PLoS Neglected Tropical Diseases, 2020, 14, e0008281.	3.0	10
20	Calpainopathy: Description of a Novel Mutation and Clinical Presentation with Early Severe Contractures. Genes, 2020, 11, 129.	2.4	5
21	The Mendelian Legacy to Mental and Behavioral Disorders. International Journal of Psychological Research, 2020, 13, 6-8.	0.6	2
22	Targeting Neuroplasticity, Cardiovascular, and Cognitive-Associated Genomic Variants in Familial Alzheimer's Disease. Molecular Neurobiology, 2019, 56, 3235-3243.	4.0	7
23	Myalgic Encephalomyelitis/Chronic Fatigue Syndrome: A Comprehensive Review. Diagnostics, 2019, 9, 91.	2.6	140
24	Genetic Variation Underpinning ADHD Risk in a Caribbean Community. Cells, 2019, 8, 907.	4.1	14
25	ADGRL3 (LPHN3) variants predict substance use disorder. Translational Psychiatry, 2019, 9, 42.	4.8	29
26	Uveitis and Multiple Sclerosis: Potential Common Causal Mutations. Molecular Neurobiology, 2019, 56, 8008-8017.	4.0	18
27	Congenital Leptin Deficiency and Leptin Gene Missense Mutation Found in Two Colombian Sisters with Severe Obesity. Genes, 2019, 10, 342.	2.4	21
28	Generation of one iPSC line (IMEDEAi006-A) from an early-onset familial Alzheimer's Disease (fAD) patient carrying the E280A mutation in the PSEN1 gene. Stem Cell Research, 2019, 37, 101440.	0.7	4
29	Neural Plasticity during Aging. Neural Plasticity, 2019, 2019, 1-3.	2.2	8
30	Low-frequency and rare variants may contribute to elucidate the genetics of major depressive disorder. Translational Psychiatry, 2018, 8, 70.	4.8	25
31	Role of the IL-1 Pathway in Dopaminergic Neurodegeneration and Decreased Voluntary Movement. Molecular Neurobiology, 2017, 54, 4486-4495.	4.0	38
32	Genetics of non-syndromic childhood obesity and the use of high-throughput DNA sequencing technologies. Journal of Diabetes and Its Complications, 2017, 31, 1549-1561.	2.3	43
33	A latent genetic subtype of major depression identified by whole-exome genotyping data in a Mexican-American cohort. Translational Psychiatry, 2017, 7, e1134-e1134.	4.8	19
34	Young adult outcomes in the followâ€up of the multimodal treatment study of attentionâ€deficit/hyperactivity disorder: symptom persistence, source discrepancy, and height suppression. Journal of Child Psychology and Psychiatry and Allied Disciplines, 2017, 58, 663-678.	5.2	207
35	The PHF21B gene is associated with major depression and modulates the stress response. Molecular Psychiatry, 2017, 22, 1015-1025.	7.9	56
36	Rare DNA variants in the brain-derived neurotrophic factor gene increase risk for attention-deficit hyperactivity disorder: a next-generation sequencing study. Molecular Psychiatry, 2017, 22, 580-584.	7.9	30

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37	Neural Plasticity in Obesity and Psychiatric Disorders. Neural Plasticity, 2016, 2016, 1-3.	2.2	1
38	A Mutation in <i>DAOA</i> Modifies the Age of Onset in <i>PSEN1</i> E280A Alzheimer's Disease. Neural Plasticity, 2016, 2016, 1-7.	2.2	25
39	A multi-ethnic genome-wide association study identifies novel loci for non-syndromic cleft lip with or without cleft palate on 2p24.2, 17q23 and 19q13. Human Molecular Genetics, 2016, 25, ddw104.	2.9	163
40	Mutations modifying sporadic Alzheimer's disease age of onset. American Journal of Medical Genetics Part B: Neuropsychiatric Genetics, 2016, 171, 1116-1130.	1.7	20
41	An Ultraconserved Brain-Specific Enhancer Within ADGRL3 (LPHN3) Underpins Attention-Deficit/Hyperactivity Disorder Susceptibility. Biological Psychiatry, 2016, 80, 943-954.	1.3	48
42	Retrospective assessment of childhood ADHD symptoms for diagnosis in adults: validity of a short 8-item version of the Wender-Utah Rating Scale. ADHD Attention Deficit and Hyperactivity Disorders, 2016, 8, 215-223.	1.7	4
43	Distinctive adaptive response to repeated exposure to hydrogen peroxide associated with upregulation of DNA repair genes and cell cycle arrest. Redox Biology, 2016, 9, 124-133.	9.0	47
44	<i> <scp>ADGRL</scp> 3 (<scp>LPHN</scp> 3) </i> variants are associated with a refined phenotype of <scp>ADHD</scp> in the <scp>MTA</scp> study. Molecular Genetics & Genomic Medicine, 2016, 4, 540-547.	1.2	35
45	Polyautoimmunity in Sjögren Syndrome. Rheumatic Disease Clinics of North America, 2016, 42, 457-472.	1.9	65
46	Definition of mutations in polyautoimmunity. Journal of Autoimmunity, 2016, 72, 65-72.	6. 5	19
47	Linkage and association analysis of ADHD endophenotypes in extended and multigenerational pedigrees from a genetic isolate. Molecular Psychiatry, 2016, 21, 1434-1440.	7.9	19
48	A Genome-wide Association Study of Nonsyndromic Cleft Palate Identifies an Etiologic Missense Variant in GRHL3. American Journal of Human Genetics, 2016, 98, 744-754.	6.2	146
49	APOE*E2 allele delays age of onset in PSEN1 E280A Alzheimer's disease. Molecular Psychiatry, 2016, 21, 916-924.	7.9	89
50	Reproductive success is predicted by social dynamics and kinship in managed animal populations. F1000Research, 2016, 5, 870.	1.6	1
51	<i>LPHN</i> 3 and attentionâ€deficit/hyperactivity disorder: a susceptibility and pharmacogenetic study. Genes, Brain and Behavior, 2015, 14, 419-427.	2.2	58
52	Novel and rare functional genomic variants in multiple autoimmune syndrome and Sjögren's syndrome. Journal of Translational Medicine, 2015, 13, 173.	4.4	30
53	Temporal Gene Expression in the Hippocampus and Peripheral Organs to Endotoxin-Induced Systemic Inflammatory Response in Caspase-1-Deficient Mice. NeuroImmunoModulation, 2015, 22, 263-273.	1.8	4
54	Response to Uher et al American Journal of Psychiatry, 2015, 172, 396-398.	7.2	1

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55	Candidate gene discovery in autoimmunity by using extreme phenotypes, next generation sequencing and whole exome capture. Autoimmunity Reviews, 2015, 14, 204-209.	5.8	29
56	Cross validation of pooling/resampling GWAS using the WTCCC data. Molecular Biology and Genetic Engineering, 2015, 3, 1.	0.8	O
57	Whole Exome Sequencing of Extreme Morbid Obesity Patients: Translational Implications for Obesity and Related Disorders. Genes, 2014, 5, 709-725.	2.4	19
58	Clinical Outcomes and Genome-Wide Association for a Brain Methylation Site in an Antidepressant Pharmacogenetics Study in Mexican Americans. American Journal of Psychiatry, 2014, 171, 1297-1309.	7.2	33
59	Mistargeting of Peroxisomal EHHADH and Inherited Renal Fanconi's Syndrome. New England Journal of Medicine, 2014, 370, 129-138.	27.0	99
60	Origin of the <i>PSEN1</i> E280A mutation causing earlyâ€onset Alzheimer's disease. Alzheimer's and Dementia, 2014, 10, S277-S283.e10.	0.8	39
61	An ¹ Hâ€MRS framework predicts the onset of Alzheimer's disease symptoms in <i>PSEN1</i> mutation carriers. Alzheimer's and Dementia, 2014, 10, 552-561.	0.8	26
62	A New Method for Detecting Significant p-values with Applications to Genetic Data. Revista Colombiana De Estadistica, 2014, 37, 69.	0.4	13
63	Influence of a Latrophilin 3 (LPHN3) risk haplotype on event-related potential measures of cognitive response control in attention-deficit hyperactivity disorder (ADHD). European Neuropsychopharmacology, 2013, 23, 458-468.	0.7	35
64	Pooling/bootstrap-based GWAS (pbGWAS) identifies new loci modifying the age of onset in PSEN1 p.Glu280Ala Alzheimer's disease. Molecular Psychiatry, 2013, 18, 568-575.	7.9	37
65	What is next after the genes for autoimmunity?. BMC Medicine, 2013, 11, 197.	5 . 5	23
66	Prediction of susceptibility to major depression by a model of interactions of multiple functional genetic variants and environmental factors. Molecular Psychiatry, 2012, 17, 624-633.	7.9	79
67	The Multiple Autoimmune Syndromes. A Clue for the Autoimmune Tautology. Clinical Reviews in Allergy and Immunology, 2012, 43, 256-264.	6.5	64
68	Exploratory data from complete genomes of familial alzheimer disease age-at-onset outliers. Human Mutation, 2012, 33, 1630-1634.	2.5	18
69	Analysis of brain metabolism by proton magnetic resonance spectroscopy (1H-MRS) in attention-deficit/hyperactivity disorder suggests a generalized differential ontogenic pattern from controls. ADHD Attention Deficit and Hyperactivity Disorders, 2012, 4, 205-212.	1.7	20
70	GWAS reveals new recessive loci associated with non-syndromic facial clefting. European Journal of Medical Genetics, 2012, 55, 510-514.	1.3	37
71	A common genetic network underlies substance use disorders and disruptive or externalizing disorders. Human Genetics, 2012, 131, 917-929.	3.8	60
72	A cooperative interaction between LPHN3 and 11q doubles the risk for ADHD. Molecular Psychiatry, 2012, 17, 741-747.	7.9	52

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73	Contribution of LPHN3 to the genetic susceptibility to ADHD in adulthood: a replication study. Genes, Brain and Behavior, 2011, 10, 149-157.	2.2	103
74	Potential cognitive endophenotypes in multigenerational families: segregating ADHD from a genetic isolate. ADHD Attention Deficit and Hyperactivity Disorders, 2011, 3, 291-299.	1.7	17
7 5	From the black widow spider to human behavior: Latrophilins, a relatively unknown class of G proteinâ€coupled receptors, are implicated in psychiatric disorders. American Journal of Medical Genetics Part B: Neuropsychiatric Genetics, 2011, 156, 1-10.	1.7	29
76	Screening of human <i>LPHN3</i> for variants with a potential impact on ADHD susceptibility. American Journal of Medical Genetics Part B: Neuropsychiatric Genetics, 2011, 156, 11-18.	1.7	49
77	A two-locus genetic interaction between LPHN3 and $11q$ predicts ADHD severity and long-term outcome. Translational Psychiatry, 2011, 1, e17-e17.	4.8	37
78	Risk HLA-DQA1 and PLA ₂ R1 Alleles in Idiopathic Membranous Nephropathy. New England Journal of Medicine, 2011, 364, 616-626.	27.0	442
79	Linear clinical progression, independent of age of onset, in Niemann–Pick disease, type C. American Journal of Medical Genetics Part B: Neuropsychiatric Genetics, 2010, 153B, 132-140.	1.7	145
80	Toward a better understanding of ADHD: LPHN3 gene variants and the susceptibility to develop ADHD. ADHD Attention Deficit and Hyperactivity Disorders, 2010, 2, 139-147.	1.7	36
81	A common variant of the latrophilin 3 gene, LPHN3, confers susceptibility to ADHD and predicts effectiveness of stimulant medication. Molecular Psychiatry, 2010, 15, 1053-1066.	7.9	245
82	A genome-wide association study of cleft lip with and without cleft palate identifies risk variants near MAFB and ABCA4. Nature Genetics, 2010, 42, 525-529.	21.4	518
83	Pharmacogenetic Impact of VKORC1 and CYP2C9 Allelic Variants on Warfarin Dose Requirements in a Hispanic Population Isolate. Clinical and Applied Thrombosis/Hemostasis, 2010, 16, 83-90.	1.7	18
84	Gray platelet syndrome: natural history of a large patient cohort and locus assignment to chromosome 3p. Blood, 2010, 116, 4990-5001.	1.4	137
85	Association of PDE11A global haplotype with major depression and antidepressant drug response. Neuropsychiatric Disease and Treatment, 2009, 5, 163.	2.2	24
86	Genome Scan, Fine-Mapping, and Candidate Gene Analysis of Non-Syndromic Cleft Lip with or without Cleft Palate Reveals Phenotype-Specific Differences in Linkage and Association Results. Human Heredity, 2009, 68, 151-170.	0.8	113
87	Epilepsy, Ataxia, Sensorineural Deafness, Tubulopathy, and <i>KCNJ10</i> Mutations. New England Journal of Medicine, 2009, 360, 1960-1970.	27.0	518
88	Planning in borderline personality disorder: Evidence for distinct subpopulations. World Journal of Biological Psychiatry, 2009, 10, 512-517.	2.6	8
89	FOXE1 association with both isolated cleft lip with or without cleft palate, and isolated cleft palate. Human Molecular Genetics, 2009, 18, 4879-4896.	2.9	136
90	Polymorphisms in the neural nicotinic acetylcholine receptor $\hat{l}\pm 4$ subunit (CHRNA4) are associated with ADHD in a genetic isolate. ADHD Attention Deficit and Hyperactivity Disorders, 2009, 1, 19-24.	1.7	19

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91	ADHD latent class clusters: DSM-IV subtypes and comorbidity. Psychiatry Research, 2009, 170, 192-198.	3.3	42
92	Locus homogeneity between syndactyly type 1A and craniosynostosis Philadelphia type?. American Journal of Medical Genetics, Part A, 2008, 146A, 2308-2311.	1.2	6
93	Metaâ€analysis of genomeâ€wide linkage scans of attention deficit hyperactivity disorder. American Journal of Medical Genetics Part B: Neuropsychiatric Genetics, 2008, 147B, 1392-1398.	1.7	160
94	Latent Class Subtyping of Attention-Deficit/Hyperactivity Disorder and Comorbid Conditions. Journal of the American Academy of Child and Adolescent Psychiatry, 2008, 47, 797-807.	0.5	73
95	Early Interstitial Lung Disease in Familial Pulmonary Fibrosis. American Journal of Respiratory and Critical Care Medicine, 2007, 176, 698-705.	5.6	157
96	CRISPLD2: a novel NSCLP candidate gene. Human Molecular Genetics, 2007, 16, 2241-2248.	2.9	78
97	Tuning major gene variants conditioning human behavior: the anachronism of ADHD. Current Opinion in Genetics and Development, 2007, 17, 234-238.	3.3	35
98	Attention-Deficit/Hyperactivity Disorder and Comorbid Disruptive Behavior Disorders: Evidence of Pleiotropy and New Susceptibility Loci. Biological Psychiatry, 2007, 61, 1329-1339.	1.3	69
99	Environmental influences that affect attention deficit/hyperactivity disorder. European Child and Adolescent Psychiatry, 2007, 16, 337-346.	4.7	69
100	Support for association between ADHD and two candidate genes:NET1andDRD1. American Journal of Medical Genetics Part B: Neuropsychiatric Genetics, 2005, 134B, 67-72.	1.7	180
101	A multigenerational pedigree of late-onset Alzheimer's disease implies new genetic causes. Brain, 2005, 128, 1707-1715.	7.6	14
102	Attention deficit/hyperactivity disorder (ADHD): Complex phenotype, simple genotype?. Genetics in Medicine, 2004, 6, 1-15.	2.4	65
103	Chagas' disease susceptibility/resistance: linkage disequilibrium analysis suggests epistasis between major histocompatibility complex and interleukin-10. Tissue Antigens, 2004, 64, 18-24.	1.0	18
104	Mutations in SLC6A19, encoding BOAT1, cause Hartnup disorder. Nature Genetics, 2004, 36, 999-1002.	21.4	272
105	Pedigree disequilibrium test (PDT) replicates association and linkage between DRD4 and ADHD in multigenerational and extended pedigrees from a genetic isolate. Molecular Psychiatry, 2004, 9, 252-259.	7.9	61
106	A novel alteration in metaxin 1, F202L, is associated with N370S in Gaucher disease. Journal of Human Genetics, 2004, 49, 220-222.	2.3	7
107	Genetic analysis of candidate loci in non-syndromic cleft lip families from Antioquia-Colombia and Ohio. American Journal of Medical Genetics Part A, 2004, 125A, 135-144.	2.4	51
108	Attention-Deficit/Hyperactivity Disorder and Comorbidities in 18 Paisa Colombian Multigenerational Families. Journal of the American Academy of Child and Adolescent Psychiatry, 2004, 43, 1506-1515.	0.5	52

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109	Interferon Regulatory Factor 6 (<i>IRF6</i>) Gene Variants and the Risk of Isolated Cleft Lip or Palate. New England Journal of Medicine, 2004, 351, 769-780.	27.0	534
110	Meta-Analysis of 13 Genome Scans Reveals Multiple Cleft Lip/Palate Genes with Novel Loci on 9q21 and 2q32-35. American Journal of Human Genetics, 2004, 75, 161-173.	6.2	200
111	Attention-Deficit/Hyperactivity Disorder in a Population Isolate: Linkage to Loci at 4q13.2, 5q33.3, 11q22, and 17p11. American Journal of Human Genetics, 2004, 75, 998-1014.	6.2	192
112	TAP1 and TAP2 polymorphisms analysis in northwestern Colombian patients with systemic lupus erythematosus. Annals of the Rheumatic Diseases, 2003, 62, 363-365.	0.9	28
113	D6S439 microsatellite identifies a new susceptibility region for primary Sjögren's syndrome. Journal of Rheumatology, 2003, 30, 2152-6.	2.0	2
114	Multiple sclerosis in the tropics: genetic association to STR's loci spanning the HLA and TNF. Multiple Sclerosis Journal, 2002, 8, 249-255.	3.0	19
115	TAP, HLA-DQB1, and HLA-DRB1 polymorphism in Colombian patients with primary SjA¶gren's syndrome. Seminars in Arthritis and Rheumatism, 2002, 31, 396-405.	3.4	31
116	Vitiligo: complex segregation and linkage disequilibrium analyses with respect to microsatellite loci spanning the HLA. Human Genetics, 2002, 110, 334-342.	3.8	81
117	Genetics of population isolates. Clinical Genetics, 2002, 61, 233-247.	2.0	227
118	Attention-deficit/hyperactivity disorder (ADHD): feasibility of linkage analysis in a genetic isolate using extended and multigenerational pedigrees. Clinical Genetics, 2002, 61, 335-343.	2.0	29
119	Rheumatoid arthritis association in Colombian population is restricted to HLA-DRB1*04 QRRAA alleles. Genes and Immunity, 2002, 3, 56-58.	4.1	31
120	Mutations in IRF6 cause Van der Woude and popliteal pterygium syndromes. Nature Genetics, 2002, 32, 285-289.	21.4	784
121	Clinical features of multiple sclerosis in a genetically homogeneous tropical population. Multiple Sclerosis Journal, 2001, 7, 227-229.	3.0	1
122	Multiple Sclerosis: Association to HLA DQ \hat{l} ± in a Tropical Population. Experimental and Clinical Immunogenetics, 1999, 16, 131-138.	1.2	11
123	Neurocysticercosis in Persons with Epilepsy in Medellin, Colombia. Epilepsia, 1998, 39, 1334-1339.	5.1	47
124	Clonal Population Structure of Colombian Sylvatic Trypanosoma cruzi. Journal of Parasitology, 1998, 84, 1143.	0.7	11
125	Complex segregation analysis of nonsyndromic cleft lip/palate in a Chilean population. Genetics and Molecular Biology, 1998, 21, 139-144.	1.3	7
126	Complex segregation analysis of schizophrenia in Santiago, Chile. Schizophrenia Research, 1997, 26, 65-69.	2.0	3

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127	Complex segregation analysis of non-myoclonic idiopathic generalized epilepsy in families ascertained from probands affected with idiopathic epilepsy with tonic-clonic seizures in Antioquia, Colombia. Human Genetics, 1996, 98, 214-218.	3.8	5