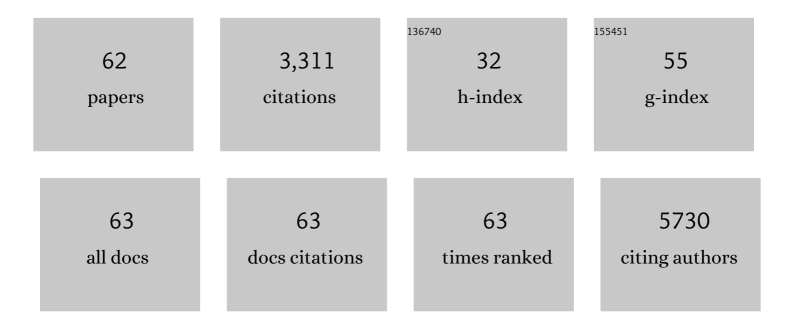
Dinesh C Soares

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	Energy Research at ACS in the Age of Open Access. ACS Omega, 2021, 6, 7967-7969.	1.6	1
2	Celebrating 5 Years of Open Access with <i>ACS Omega</i> . ACS Omega, 2020, 5, 16986-16986.	1.6	2
3	Translation elongation factor 1A2 is encoded by one of four closely related <i>eef1a</i> genes and is dispensable for survival in zebrafish. Bioscience Reports, 2020, 40, .	1.1	4
4	Missense Mutations in the Human Nanophthalmos Gene <i>TMEM98</i> Cause Retinal Defects in the Mouse. , 2019, 60, 2875.		16
5	Characterization of a novel RP2-OSTF1 interaction and its implication for actin remodeling. Journal of Cell Science, 2018, 131, .	1.2	6
6	ACS Omega 2017: A Year-End Expression of Appreciation for the Fundamental Contributions of Our Reviewers. ACS Omega, 2018, 3, 595-607.	1.6	2
7	HIV-1 Uncoating and Reverse Transcription Require eEF1A Binding to Surface-Exposed Acidic Residues of the Reverse Transcriptase Thumb Domain. MBio, 2018, 9, .	1.8	18
8	Rare disruptive variants in the DISC1 Interactome and Regulome: association with cognitive ability and schizophrenia. Molecular Psychiatry, 2018, 23, 1270-1277.	4.1	37
9	Pathogenic mutations in retinitis pigmentosa 2 predominantly result in loss of RP2 protein stability in humans and zebrafish. Journal of Biological Chemistry, 2017, 292, 6225-6239.	1.6	16
10	A structural organization for the Disrupted in Schizophrenia 1 protein, identified by high-throughput screening, reveals distinctly folded regions, which are bisected by mental illness-related mutations. Journal of Biological Chemistry, 2017, 292, 6468-6477.	1.6	22
11	PLAA Mutations Cause a Lethal Infantile Epileptic Encephalopathy by Disrupting Ubiquitin-Mediated Endolysosomal Degradation of Synaptic Proteins. American Journal of Human Genetics, 2017, 100, 706-724.	2.6	37
12	A recurrent de novo mutation in <i>ACTG1</i> causes isolated ocular coloboma. Human Mutation, 2017, 38, 942-946.	1.1	21
13	SAF-A Regulates Interphase Chromosome Structure through Oligomerization with Chromatin-Associated RNAs. Cell, 2017, 169, 1214-1227.e18.	13.5	166
14	Ablation of EYS in zebrafish causes mislocalisation of outer segment proteins, F-actin disruption and cone-rod dystrophy. Scientific Reports, 2017, 7, 46098.	1.6	52
15	Novel pathogenic mutations in C1QTNF5 support a dominant negative disease mechanism in late-onset retinal degeneration. Scientific Reports, 2017, 7, 12147.	1.6	30
16	HpARI Protein Secreted by a Helminth Parasite Suppresses Interleukin-33. Immunity, 2017, 47, 739-751.e5.	6.6	130
17	A structurally distinct TGF-β mimic from an intestinal helminth parasite potently induces regulatory T cells. Nature Communications, 2017, 8, 1741.	5.8	159
18	Condensin II mutation causes T-cell lymphoma through tissue-specific genome instability. Genes and Development, 2016, 30, 2173-2186.	2.7	41

DINESH C SOARES

#	Article	IF	CITATIONS
19	Novel de novo <i><scp>EEF</scp>1A2</i> missense mutations causing epilepsy and intellectual disability. Molecular Genetics & Genomic Medicine, 2016, 4, 465-474.	0.6	44
20	Atypical Femoral Fracture in Osteoporosis Pseudoglioma Syndrome Associated with Two Novel Compound Heterozygous Mutations in <i>LRP5</i> . Journal of Bone and Mineral Research, 2015, 30, 615-620.	3.1	25
21	Evolutionary Characterization of the Retinitis Pigmentosa GTPase Regulator Gene. , 2015, 56, 6255.		16
22	Specific Interaction between eEF1A and HIV RT Is Critical for HIV-1 Reverse Transcription and a Potential Anti-HIV Target. PLoS Pathogens, 2015, 11, e1005289.	2.1	16
23	Targeted sequencing of the Paget's disease associated 14q32 locus identifies several missense coding variants in RIN3 that predispose to Paget's disease of bone. Human Molecular Genetics, 2015, 24, 3286-3295.	1.4	29
24	RPGR: Its role in photoreceptor physiology, human disease, and future therapies. Experimental Eye Research, 2015, 138, 32-41.	1.2	98
25	DISC1 as a genetic risk factor for schizophrenia and related major mental illness: response to Sullivan. Molecular Psychiatry, 2014, 19, 141-143.	4.1	62
26	DISC1 complexes with TRAK1 and Miro1 to modulate anterograde axonal mitochondrial trafficking. Human Molecular Genetics, 2014, 23, 906-919.	1.4	84
27	Monoallelic and Biallelic Mutations in MAB21L2 Cause a Spectrum of Major Eye Malformations. American Journal of Human Genetics, 2014, 94, 915-923.	2.6	79
28	708 Common and 2010 rare DISC1 locus variants identified in 1542 subjects: analysis for association with psychiatric disorder and cognitive traits. Molecular Psychiatry, 2014, 19, 668-675.	4.1	59
29	Highly homologous eEF1A1 and eEF1A2 exhibit differential post-translationalmodification with significant enrichment around localised sites of sequence variation. Biology Direct, 2013, 8, 29.	1.9	26
30	DISC1 genetics, biology and psychiatric illness. Frontiers in Biology, 2013, 8, 1-31.	0.7	102
31	The intermediate filament protein, vimentin, is a regulator of NOD2 activity. Gut, 2013, 62, 695-707.	6.1	71
32	NDE1 and NDEL1: twin neurodevelopmental proteins with similar â€~nature' but different â€~nurture'. Biomolecular Concepts, 2013, 4, 447-464.	1.0	40
33	The Mitosis and Neurodevelopment Proteins NDE1 and NDEL1 Form Dimers, Tetramers, and Polymers with a Folded Back Structure in Solution. Journal of Biological Chemistry, 2012, 287, 32381-32393.	1.6	38
34	Solution Structure of CCP Modules 10–12 Illuminates Functional Architecture of the Complement Regulator, Factor H. Journal of Molecular Biology, 2012, 424, 295-312.	2.0	24
35	A t(1;11) translocation linked to schizophrenia and affective disorders gives rise to aberrant chimeric DISC1 transcripts that encode structurally altered, deleterious mitochondrial proteins. Human Molecular Genetics, 2012, 21, 3374-3386.	1.4	61
36	Structural Analysis of the C-Terminal Region (Modules 18–20) of Complement Regulator Factor H (FH). PLoS ONE, 2012, 7, e32187.	1.1	39

DINESH C SOARES

#	Article	IF	CITATIONS
37	DISC1: Structure, Function, and Therapeutic Potential for Major Mental Illness. ACS Chemical Neuroscience, 2011, 2, 609-632.	1.7	109
38	PKA Phosphorylation of NDE1 Is DISC1/PDE4 Dependent and Modulates Its Interaction with LIS1 and NDEL1. Journal of Neuroscience, 2011, 31, 9043-9054.	1.7	72
39	In silico structure-function analysis of pathological variation in the <i>HSD11B2</i> gene sequence. Physiological Genomics, 2010, 42, 319-330.	1.0	24
40	The structure of the KlcA and ArdB proteins reveals a novel fold and antirestriction activity against Type I DNA restriction systems in vivo but not in vitro. Nucleic Acids Research, 2010, 38, 1723-1737.	6.5	50
41	Oxovanadium(IV) Cyclam and Bicyclam Complexes: Potential CXCR4 Receptor Antagonists. Inorganic Chemistry, 2010, 49, 1122-1132.	1.9	39
42	The Central Portion of Factor H (Modules 10–15) Is Compact and Contains a Structurally Deviant CCP Module. Journal of Molecular Biology, 2010, 395, 105-122.	2.0	51
43	Structural Models of Human eEF1A1 and eEF1A2 Reveal Two Distinct Surface Clusters of Sequence Variation and Potential Differences in Phosphorylation. PLoS ONE, 2009, 4, e6315.	1.1	60
44	Solution Structure of Factor I-like Modules from Complement C7 Reveals a Pair of Follistatin Domains in Compact Pseudosymmetric Arrangement. Journal of Biological Chemistry, 2009, 284, 19637-19649.	1.6	15
45	An interrupted beta-propeller and protein disorder: structural bioinformatics insights into the N-terminus of alsin. Journal of Molecular Modeling, 2009, 15, 113-122.	0.8	11
46	A Cytogenetic Abnormality and Rare Coding Variants Identify ABCA13 as a Candidate Gene in Schizophrenia, Bipolar Disorder, and Depression. American Journal of Human Genetics, 2009, 85, 833-846.	2.6	102
47	NDE1 and NDEL1: Multimerisation, alternate splicing and DISC1 interaction. Neuroscience Letters, 2009, 449, 228-233.	1.0	41
48	eEF1A2 and neuronal degeneration. Biochemical Society Transactions, 2009, 37, 1293-1297.	1.6	31
49	The DISC locus in psychiatric illness. Molecular Psychiatry, 2008, 13, 36-64.	4.1	554
50	The evolution of TEP1, an exceptionally polymorphic immunity gene in Anopheles gambiae. BMC Evolutionary Biology, 2008, 8, 274.	3.2	47
51	Structure of the N-terminal Region of Complement Factor H and Conformational Implications of Disease-linked Sequence Variations. Journal of Biological Chemistry, 2008, 283, 9475-9487.	1.6	58
52	A common variant in the 3'UTR of the GRIK4 glutamate receptor gene affects transcript abundance and protects against bipolar disorder. Proceedings of the National Academy of Sciences of the United States of America, 2008, 105, 14940-14945.	3.3	50
53	Structure-based Mapping of DAF Active Site Residues That Accelerate the Decay of C3 Convertases. Journal of Biological Chemistry, 2007, 282, 18552-18562.	1.6	27
54	Molecular evolution of the human SRPX2 gene that causes brain disorders of the Rolandic and Sylvian speech areas. BMC Genetics, 2007, 8, 72.	2.7	25

DINESH C SOARES

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55	Disease-Associated Sequence Variations in Factor H: A Structural Biology Approach. , 2006, 586, 313-327.		5
56	Opportunities for New Therapies Based on the Natural Regulators of Complement Activation. Annals of the New York Academy of Sciences, 2005, 1056, 176-188.	1.8	12
57	Large-scale modelling as a route to multiple surface comparisons of the CCP module family. Protein Engineering, Design and Selection, 2005, 18, 379-388.	1.0	45
58	Functional Insights from the Structure of the Multifunctional C345C Domain of C5 of Complement. Journal of Biological Chemistry, 2005, 280, 10636-10645.	1.6	25
59	ARABIDOPSIS CRINKLY4 Function, Internalization, and Turnover Are Dependent on the Extracellular Crinkly Repeat Domain. Plant Cell, 2005, 17, 1154-1166.	3.1	94
60	Complement Control Protein Modules in the Regulators of Complement Activation. , 2005, , 19-62.		27
61	Complement Control Protein Modules in the Regulators of Complement Activation. , 2005, , 19-62.		5
62	Structural Analysis of the Complement Control Protein (CCP) Modules of GABAB Receptor 1a. Journal of Biological Chemistry, 2004, 279, 48292-48306.	1.6	59