Tim Becker

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/5541198/publications.pdf

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55	2,827	25	52
papers	citations	h-index	g-index
56	56	56	5572 citing authors
all docs	docs citations	times ranked	

#	Article	IF	Citations
1	Assessment of significance of conditionally independent GWAS signals. Bioinformatics, 2021, 37, 3521-3529.	1.8	1
2	Consistent Assignment of Risk and Benign Allele at rs2303153 in the CF Modifier Gene SCNN1B in Three Independent F508del-CFTR Homozygous Patient Populations. Genes, 2021, 12, 1554.	1.0	2
3	Prior fluid and electrolyte imbalance is associated with COVID-19 mortality. Communications Medicine, 2021, 1, .	1.9	12
4	The exhaustive genomic scan approach, with an application to rare-variant association analysis. European Journal of Human Genetics, 2020, 28, 1283-1291.	1.4	3
5	Genetic information from discordant sibling pairs points to ESRP2 as a candidate trans-acting regulator of the CF modifier gene SCNN1B. Scientific Reports, 2020, 10, 22447.	1.6	4
6	Meta-analysis identifies novel risk loci and yields systematic insights into the biology of male-pattern baldness. Nature Communications, 2017, 8, 14694.	5.8	58
7	Single nucleotide polymorphisms in the angiogenic and lymphangiogenic pathways are associated with lymphedema caused by Wuchereria bancrofti. Human Genomics, 2017, 11, 26.	1.4	17
8	Low-level <i>APC</i> mutational mosaicism is the underlying cause in a substantial fraction of unexplained colorectal adenomatous polyposis cases. Journal of Medical Genetics, 2016, 53, 172-179.	1.5	51
9	Haplotype synthesis analysis reveals functional variants underlying known genome-wide associated susceptibility loci. Bioinformatics, 2016, 32, 2136-2142.	1.8	2
10	TREM2 rare variant p.R47H is not associated with Parkinson's disease. Parkinsonism and Related Disorders, 2016, 23, 109-111.	1.1	17
11	Alzheimer's disease risk variants modulate endophenotypes in mild cognitive impairment. Alzheimer's and Dementia, 2016, 12, 872-881.	0.4	50
12	Association of age-of-onset groups with GWAS significant schizophrenia and bipolar disorder loci in Romanian bipolar I patients. Psychiatry Research, 2015, 230, 964-967.	1.7	11
13	The influence of genetic variants in SORL1 gene on the manifestation of Alzheimer's disease. Neurobiology of Aging, 2015, 36, 1605.e13-1605.e20.	1.5	27
14	Novel genetic matching methods for handling population stratification in genome-wide association studies. BMC Bioinformatics, 2015, 16, 84.	1.2	8
15	PLD3 in non-familial Alzheimer's disease. Nature, 2015, 520, E3-E5.	13.7	58
16	Convergent genetic and expression data implicate immunity in Alzheimer's disease. Alzheimer's and Dementia, 2015, 11, 658-671.	0.4	173
17	Immunochip-Based Analysis: High-Density Genotyping of Immune-Related Loci Sheds Further Light on the Autoimmune Genetic Architecture of Alopecia Areata. Journal of Investigative Dermatology, 2015, 135, 919-921.	0.3	7
18	METAINTER: meta-analysis of multiple regression models in genome-wide association studies. Bioinformatics, 2015, 31, 151-157.	1.8	18

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19	SUCLG2 identified as both a determinator of CSF A \hat{l}^2 1 \hat{a} \in "42 levels and an attenuator of cognitive decline in Alzheimer's disease. Human Molecular Genetics, 2014, 23, 6644-6658.	1.4	45
20	The CF-modifying gene EHF promotes p.Phe508del-CFTR residual function by altering protein glycosylation and trafficking in epithelial cells. European Journal of Human Genetics, 2014, 22, 660-666.	1.4	26
21	Genome-wide association study reveals two new risk loci for bipolar disorder. Nature Communications, 2014, 5, 3339.	5.8	294
22	Genome-wide association study and mouse expression data identify a highly conserved 32 kb intergenic region between WNT3 and WNT9b as possible susceptibility locus for isolated classic exstrophy of the bladder. Human Molecular Genetics, 2014, 23, 5536-5544.	1.4	19
23	Investigation of the role of rare TREM2 variants in frontotemporal dementia subtypes. Neurobiology of Aging, 2014, 35, 2657.e13-2657.e19.	1.5	34
24	Gene-Wide Analysis Detects Two New Susceptibility Genes for Alzheimer's Disease. PLoS ONE, 2014, 9, e94661.	1.1	155
25	Androgenetic Alopecia: Identification of Four Genetic Risk Loci and Evidence for the Contribution of WNT Signaling to Its Etiology. Journal of Investigative Dermatology, 2013, 133, 1489-1496.	0.3	83
26	CLCA4 variants determine the manifestation of the cystic fibrosis basic defect in the intestine. European Journal of Human Genetics, 2013, 21, 691-694.	1.4	11
27	A One-Degree-of-Freedom Test for Supra-Multiplicativity of SNP Effects. PLoS ONE, 2013, 8, e78038.	1.1	2
28	Six Novel Susceptibility Loci for Early-Onset Androgenetic Alopecia and Their Unexpected Association with Common Diseases. PLoS Genetics, 2012, 8, e1002746.	1.5	92
29	Integrated Genome-Wide Pathway Association Analysis with INTERSNP. Human Heredity, 2012, 73, 63-72.	0.4	11
30	Quick, "Imputation-free―meta-analysis with proxy-SNPs. BMC Bioinformatics, 2012, 13, 231.	1.2	7
31	Follow-Up Study of the First Genome-Wide Association Scan in Alopecia Areata: IL13 and KIAA0350 as Susceptibility Loci Supported with Genome-Wide Significance. Journal of Investigative Dermatology, 2012, 132, 2192-2197.	0.3	107
32	Significance Levels in Genome-Wide Interaction Analysis (GWIA). Annals of Human Genetics, 2011, 75, 29-35.	0.3	21
33	Genetic Variants in CTLA4 Are Strongly Associated with Alopecia Areata. Journal of Investigative Dermatology, 2011, 131, 1169-1172.	0.3	43
34	An association study on contrasting cystic fibrosis endophenotypes recognizes KRT8 but not KRT18 as a modifier of cystic fibrosis disease severity and CFTR mediated residual chloride secretion. BMC Medical Genetics, 2011, 12, 62.	2.1	15
35	Genes that determine immunology and inflammation modify the basic defect of impaired ion conductance in cystic fibrosis epithelia. Journal of Medical Genetics, 2011, 48, 24-31.	1.5	62
36	Initial interrogation, confirmation and fine mapping of modifying genes: STAT3, IL1B and IFNGR1 determine cystic fibrosis disease manifestation. European Journal of Human Genetics, 2011, 19, 1281-1288.	1.4	15

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37	Hierarchical fine mapping of the cystic fibrosis modifier locus on 19q13 identifies an association with two elements near the genes CEACAM3 and CEACAM6. Human Genetics, 2010, 127, 383-394.	1.8	12
38	Evaluation of Potential Power Gain with Imputed Genotypes in Genome-Wide Association Studies. Human Heredity, 2009, 68, 23-34.	0.4	3
39	INTERSNP: genome-wide interaction analysis guided by a priori information. Bioinformatics, 2009, 25, 3275-3281.	1.8	129
40	Joint analysis of tightly linked SNPs in screening step of genome-wide association studies leads to increased power. European Journal of Human Genetics, 2009, 17, 1043-1049.	1.4	17
41	Susceptibility variants for male-pattern baldness on chromosome 20p11. Nature Genetics, 2008, 40, 1279-1281.	9.4	119
42	Association study between genetic variants at the VAMP2 and VAMP3 loci and bipolar affective disorder. Psychiatric Genetics, 2008, 18, 199-203.	0.6	10
43	Transmission ratio distortion and maternal effects confound the analysis of modulators of cystic fibrosis disease severity on 19q13. European Journal of Human Genetics, 2007, 15, 774-778.	1.4	15
44	Identification of probable genotyping errors by consideration of haplotypes. European Journal of Human Genetics, 2006, 14, 450-458.	1.4	15
45	The TNF $\hat{l}\pm$ receptor TNFRSF1A and genes encoding the amiloride-sensitive sodium channel ENaC as modulators in cystic fibrosis. Human Genetics, 2006, 119, 331-343.	1.8	43
46	Detection of Parent-of-Origin Effects in Nuclear Families Using Haplotype Analysis. Human Heredity, 2006, 62, 64-76.	0.4	15
47	Haplotype interaction analysis of unlinked regions. Genetic Epidemiology, 2005, 29, 313-322.	0.6	43
48	Impact of Missing Genotype Data on Monte-Carlo Simulation Based Haplotype Analysis. Human Heredity, 2005, 59, 185-189.	0.4	6
49	Genetic Variation in the Human Androgen Receptor Gene Is the Major Determinant of Common Early-Onset Androgenetic Alopecia. American Journal of Human Genetics, 2005, 77, 140-148.	2.6	198
50	Maximum-likelihood estimation of haplotype frequencies in nuclear families. Genetic Epidemiology, 2004, 27, 21-32.	0.6	191
51	Impact of Genotyping Errors on Type I Error Rate of the Haplotype-Sharing Transmission/Disequilibrium Test (HS-TDT). American Journal of Human Genetics, 2004, 74, 589-591.	2.6	33
52	A Powerful Strategy to Account for Multiple Testing in the Context of Haplotype Analysis. American Journal of Human Genetics, 2004, 75, 561-570.	2.6	153
53	The DTNBP1 (Dysbindin) Gene Contributes to Schizophrenia, Depending on Family History of the Disease. American Journal of Human Genetics, 2003, 73, 1438-1443.	2.6	180
54	Family-Based Association Analysis with Tightly Linked Markers. Human Heredity, 2003, 56, 2-9.	0.4	55

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55	Efficiency of Haplotype Frequency Estimation when Nuclear Familiy Information Is Included. Human Heredity, 2002, 54, 45-53.	0.4	29