## Andrew S Allen

List of Publications by Year in descending order

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ANDREW SALLEN

#	Article	IF	CITATIONS
1	Ancestry adjustment improves genome-wide estimates of regional intolerance. Genetics, 2022, , .	1.2	2
2	Casual Genetic Variants in Stillbirth. Obstetrical and Gynecological Survey, 2021, 76, 79-81.	0.2	0
3	Correcting signal biases and detecting regulatory elements in STARR-seq data. Genome Research, 2021, 31, 877-889.	2.4	11
4	Diverse genetic causes of polymicrogyria with epilepsy. Epilepsia, 2021, 62, 973-983.	2.6	12
5	Sub-genic intolerance, ClinVar, and the epilepsies: A whole-exome sequencing study of 29,165 individuals. American Journal of Human Genetics, 2021, 108, 965-982.	2.6	35
6	Coding and noncoding variants in EBF3 are involved in HADDS and simplex autism. Human Genomics, 2021, 15, 44.	1.4	16
7	Focused goodness of fit tests for gene set analyses. Briefings in Bioinformatics, 2021, , .	3.2	Ο
8	Bayesian estimation of genetic regulatory effects in high-throughput reporter assays. Bioinformatics, 2020, 36, 331-338.	1.8	0
9	Hypnotics with novel modes of action. British Journal of Clinical Pharmacology, 2020, 86, 244-249.	1.1	25
10	Causal Genetic Variants in Stillbirth. New England Journal of Medicine, 2020, 383, 1107-1116.	13.9	67
11	Incorporating external information to improve sparse signal detection in rareâ€variant geneâ€setâ€based analyses. Genetic Epidemiology, 2020, 44, 330-338.	0.6	5
12	Rare-variant collapsing analyses for complex traits: guidelines and applications. Nature Reviews Genetics, 2019, 20, 747-759.	7.7	147
13	Evaluating Chromatin Accessibility Differences Across Multiple Primate Species Using a Joint Modeling Approach. Genome Biology and Evolution, 2019, 11, 3035-3053.	1.1	12
14	Improved Pathogenic Variant Localization via a Hierarchical Model of Sub-regional Intolerance. American Journal of Human Genetics, 2019, 104, 299-309.	2.6	29
15	Exome-Based Rare-Variant Analyses in CKD. Journal of the American Society of Nephrology: JASN, 2019, 30, 1109-1122.	3.0	40
16	A new approach for rare variation collapsing on functional protein domains implicates specific genic regions in ALS. Genome Research, 2019, 29, 809-818.	2.4	21
17	Genome-wide association study (GWAS) of human host factors influencing viral severity of herpes simplex virus type 2 (HSV-2). Genes and Immunity, 2019, 20, 112-120.	2.2	17
18	Facilitating the Calculation of the Efficient Score Using Symbolic Computing. American Statistician, 2018, 72, 199-205.	0.9	0

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19	Loss of tubulin deglutamylase <scp>CCP</scp> 1 causes infantileâ€onset neurodegeneration. EMBO Journal, 2018, 37, .	3.5	86
20	Human genome-wide measurement of drug-responsive regulatory activity. Nature Communications, 2018, 9, 5317.	5.8	34
21	meaRtools: An R package for the analysis of neuronal networks recorded on microelectrode arrays. PLoS Computational Biology, 2018, 14, e1006506.	1.5	22
22	<tt>bcSeq</tt> : an R package for fast sequence mapping in high-throughput shRNA and CRISPR screens. Bioinformatics, 2018, 34, 3581-3583.	1.8	4
23	De novo and inherited private variants in MAP1B in periventricular nodular heterotopia. PLoS Genetics, 2018, 14, e1007281.	1.5	40
24	High-throughput interpretation of gene structure changes in human and nonhuman resequencing data, using ACE. Bioinformatics, 2017, 33, 1437-1446.	1.8	2
25	An Exome Sequencing Study to Assess the Role of Rare Genetic Variation in Pulmonary Fibrosis. American Journal of Respiratory and Critical Care Medicine, 2017, 196, 82-93.	2.5	185
26	Quantifying the Impact of Non-coding Variants on Transcription Factor-DNA Binding. Lecture Notes in Computer Science, 2017, 10229, 336-352.	1.0	16
27	Leveraging population information in familyâ€based rare variant association analyses of quantitative traits. Genetic Epidemiology, 2017, 41, 98-107.	0.6	3
28	Transversions have larger regulatory effects than transitions. BMC Genomics, 2017, 18, 394.	1.2	83
29	A case-control collapsing analysis identifies epilepsy genes implicated in trio sequencing studies focused on de novo mutations. PLoS Genetics, 2017, 13, e1007104.	1.5	25
30	Mapping eQTL by leveraging multiple tissues and DNA methylation. BMC Bioinformatics, 2017, 18, 455.	1.2	5
31	Orion: Detecting regions of the human non-coding genome that are intolerant to variation using population genetics. PLoS ONE, 2017, 12, e0181604.	1.1	31
32	Robust analysis of secondary phenotypes in case ontrol genetic association studies. Statistics in Medicine, 2016, 35, 4226-4237.	0.8	8
33	Exploiting expression patterns across multiple tissues to map expression quantitative trait loci. BMC Bioinformatics, 2016, 17, 257.	1.2	12
34	The intolerance to functional genetic variation of protein domains predicts the localization of pathogenic mutations within genes. Genome Biology, 2016, 17, 9.	3.8	118
35	Polymorphisms in HLA Class II Genes Are Associated With Susceptibility to <i>Staphylococcus aureus</i> Infection in a White Population. Journal of Infectious Diseases, 2016, 213, 816-823.	1.9	44
36	Lifting EMMeT to OWL Getting the Most from SKOS. Lecture Notes in Computer Science, 2016, , 69-80.	1.0	3

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37	Testing the Effect of Rare Compoundâ€Heterozygous and Recessive Mutations in Case–Parent Sequencing Studies. Genetic Epidemiology, 2015, 39, 166-172.	0.6	5
38	The Intolerance of Regulatory Sequence to Genetic Variation Predicts Gene Dosage Sensitivity. PLoS Genetics, 2015, 11, e1005492.	1.5	123
39	Exome sequencing in amyotrophic lateral sclerosis identifies risk genes and pathways. Science, 2015, 347, 1436-1441.	6.0	823
40	Incorporating Functional Information in Tests of Excess De Novo Mutational Load. American Journal of Human Genetics, 2015, 97, 272-283.	2.6	29
41	The Genetics of Neuropsychiatric Diseases: Looking In and Beyond the Exome. Annual Review of Neuroscience, 2015, 38, 47-68.	5.0	27
42	Testing for risk and protective trends in genetic analyses of HIV acquisition. Biostatistics, 2015, 16, 268-280.	0.9	0
43	Utilizing Population Controls in Rare-Variant Case-Parent Association Tests. American Journal of Human Genetics, 2014, 94, 845-853.	2.6	15
44	De novo mutations in epileptic encephalopathies. Nature, 2013, 501, 217-221.	13.7	1,351
45	Sequencing studies in human genetics: design and interpretation. Nature Reviews Genetics, 2013, 14, 460-470.	7.7	236
46	Genic Intolerance to Functional Variation and the Interpretation of Personal Genomes. PLoS Genetics, 2013, 9, e1003709.	1.5	844
47	Stratification core Matching Improves Correction for Confounding by Population Stratification in Caseâ€Control Association Studies. Genetic Epidemiology, 2012, 36, 195-205.	0.6	21
48	Control for Confounding in Case-Control Studies Using the Stratification Score, a Retrospective Balancing Score. American Journal of Epidemiology, 2011, 173, 752-760.	1.6	21
49	Scoreâ€based adjustment for confounding by population stratification in genetic association studies. Genetic Epidemiology, 2010, 34, 383-385.	0.6	7
50	SNPs in CAST are associated with Parkinson disease: A confirmation study. American Journal of Medical Genetics Part B: Neuropsychiatric Genetics, 2010, 153B, 973-979.	1.1	6
51	Fast and Robust Association Tests for Untyped SNPs in Case-Control Studies. Human Heredity, 2010, 70, 167-176.	0.4	3
52	Effect of population stratification on the identification of significant single-nucleotide polymorphisms in genome-wide association studies. BMC Proceedings, 2009, 3, S13.	1.8	12
53	Genome-wide association analysis of rheumatoid arthritis data via haplotype sharing. BMC Proceedings, 2009, 3, S30.	1.8	9
54	A novel haplotypeâ€sharing approach for genomeâ€wide caseâ€control association studies implicates the calpastatin gene in Parkinson's disease. Genetic Epidemiology, 2009, 33, 657-667.	0.6	29

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55	A Regressionâ€based Association Test for Caseâ€control Studies that Uses Inferred Ancestral Haplotype Similarity. Annals of Human Genetics, 2009, 73, 520-526.	0.3	6
56	Robust estimation and testing of haplotype effects in case ontrol studies. Genetic Epidemiology, 2008, 32, 29-40.	0.6	8
57	Response to Lee etÂal American Journal of Human Genetics, 2008, 82, 526-528.	2.6	6
58	Invited Keynote Talk: Haplotype Sharing for Genome-Wide Case-Control Association Studies. , 2008, , 183-183.		0
59	Statistical Models for Haplotype Sharing in Case-Parent Trio Data. Human Heredity, 2007, 64, 35-44.	0.4	19
60	A Simple and Improved Correction for Population Stratification in Case-Control Studies. American Journal of Human Genetics, 2007, 80, 921-930.	2.6	150
61	Association mapping via a class of haplotype-sharing statistics. BMC Proceedings, 2007, 1, S123.	1.8	4
62	Inference on haplotype/disease association using parent-affected-child data: the projection conditional on parental haplotypes method. Genetic Epidemiology, 2007, 31, 211-223.	0.6	21
63	Summary of contributions to GAW15 Group 13: candidate gene association studies. Genetic Epidemiology, 2007, 31, S110-S117.	0.6	2
64	Robust testing of haplotype/disease association. BMC Genetics, 2005, 6, S69.	2.7	5
65	Locally-efficient robust estimation of haplotype-disease association in family-based studies. Biometrika, 2005, 92, 559-571.	1.3	21
66	Informative Missingness in Genetic Association Studies: Case-Parent Designs. American Journal of Human Genetics, 2003, 72, 671-680.	2.6	67
67	Joint Models for Toxicology Studies with Doseâ€Dependent Number of Implantations. Risk Analysis, 2002, 22, 1165-1173.	1.5	8