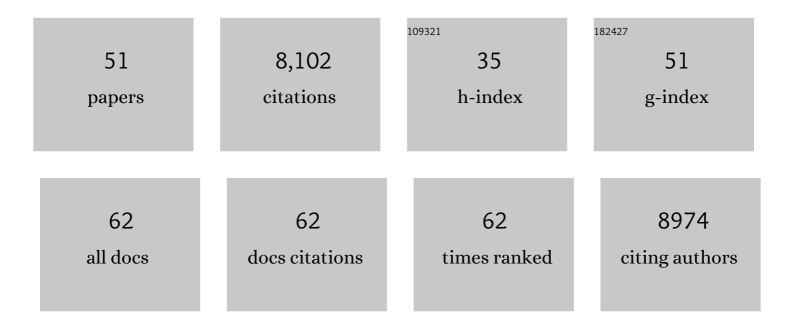
Kevin R Thornton

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	The Drosophila melanogaster Genetic Reference Panel. Nature, 2012, 482, 173-178.	27.8	1,756
2	The origin of new genes: glimpses from the young and old. Nature Reviews Genetics, 2003, 4, 865-875.	16.3	775
3	Comparative genome sequencing of Drosophila pseudoobscura: Chromosomal, gene, and cis-element evolution. Genome Research, 2005, 15, 1-18.	5.5	453
4	A New Approach for Using Genome Scans to Detect Recent Positive Selection in the Human Genome. PLoS Biology, 2007, 5, e171.	5.6	413
5	Genome-wide analysis of a long-term evolution experiment with Drosophila. Nature, 2010, 467, 587-590.	27.8	410
6	Retroposed New Genes Out of the X in Drosophila. Genome Research, 2002, 12, 1854-1859.	5.5	399
7	libsequence: a C++ class library for evolutionary genetic analysis. Bioinformatics, 2003, 19, 2325-2327.	4.1	307
8	Multilocus patterns of nucleotide variability and the demographic and selection history of Drosophila melanogaster populations. Genome Research, 2005, 15, 790-799.	5.5	247
9	Approximate Bayesian Inference Reveals Evidence for a Recent, Severe Bottleneck in a Netherlands Population of Drosophila melanogaster. Genetics, 2006, 172, 1607-1619.	2.9	239
10	Genome sequencing reveals complex speciation in the <i>Drosophila simulans</i> clade. Genome Research, 2012, 22, 1499-1511.	5.5	220
11	EXTENSIVE INTROGRESSION OF MITOCHONDRIAL DNA RELATIVE TO NUCLEAR GENES IN THE DROSOPHILA YAKUBA SPECIES GROUP. Evolution; International Journal of Organic Evolution, 2006, 60, 292-302.	2.3	187
12	Controlling the False-Positive Rate in Multilocus Genome Scans for Selection. Genetics, 2007, 175, 737-750.	2.9	163
13	A second-generation assembly of the <i>Drosophila simulans</i> genome provides new insights into patterns of lineage-specific divergence. Genome Research, 2013, 23, 89-98.	5.5	157
14	Efficient ancestry and mutation simulation with msprime 1.0. Genetics, 2022, 220, .	2.9	133
15	Efficient pedigree recording for fast population genetics simulation. PLoS Computational Biology, 2018, 14, e1006581.	3.2	130
16	Progress and prospects in mapping recent selection in the genome. Heredity, 2007, 98, 340-348.	2.6	121
17	Abundance and Distribution of Transposable Elements in Two Drosophila QTL Mapping Resources. Molecular Biology and Evolution, 2013, 30, 2311-2327.	8.9	118
18	Characterizing natural variation using next-generation sequencing technologies. Trends in Genetics, 2009. 25. 463-471.	6.7	116

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#	Article	IF	CITATIONS
19	Gene Duplication and Evolution. Science, 2001, 293, 1551a-1551.	12.6	103
20	Retroposed New Genes Out of the X in <i>Drosophila</i> . Genome Research, 2002, 12, 1854-1859.	5.5	99
21	An Approximate Bayesian Estimator Suggests Strong, Recurrent Selective Sweeps in Drosophila. PLoS Genetics, 2008, 4, e1000198.	3.5	94
22	Evidence of directional and stabilizing selection in contemporary humans. Proceedings of the National Academy of Sciences of the United States of America, 2018, 115, 151-156.	7.1	90
23	Landscape of Standing Variation for Tandem Duplications in Drosophila yakuba and Drosophila simulans. Molecular Biology and Evolution, 2014, 31, 1750-1766.	8.9	89
24	The Power to Detect Quantitative Trait Loci Using Resequenced, Experimentally Evolved Populations of Diploid, Sexual Organisms. Molecular Biology and Evolution, 2014, 31, 1040-1055.	8.9	88
25	On the Utility of Linkage Disequilibrium as a Statistic for Identifying Targets of Positive Selection in Nonequilibrium Populations. Genetics, 2007, 176, 2371-2379.	2.9	84
26	Rapid Divergence of Gene Duplicates on the Drosophila melanogaster X Chromosome. Molecular Biology and Evolution, 2002, 19, 918-925.	8.9	83
27	Nucleotide Variation Along the Drosophila melanogaster Fourth Chromosome. Science, 2002, 295, 134-137.	12.6	81
28	A C++ Template Library for Efficient Forward-Time Population Genetic Simulation of Large Populations. Genetics, 2014, 198, 157-166.	2.9	78
29	Properties and Modeling of GWAS when Complex Disease Risk Is Due to Non-Complementing, Deleterious Mutations in Genes of Large Effect. PLoS Genetics, 2013, 9, e1003258.	3.5	72
30	Extensive introgression of mitochondrial DNA relative to nuclear genes in the Drosophila yakuba species group. Evolution; International Journal of Organic Evolution, 2006, 60, 292-302.	2.3	69
31	X chromosomes and autosomes evolve at similar rates in <i>Drosophila</i> : No evidence for faster-X protein evolution. Genome Research, 2006, 16, 498-504.	5.5	67
32	Tandem duplications lead to novel expression patterns through exon shuffling in Drosophila yakuba. PLoS Genetics, 2017, 13, e1006795.	3.5	53
33	Efficiently Summarizing Relationships in Large Samples: A General Duality Between Statistics of Genealogies and Genomes. Genetics, 2020, 215, 779-797.	2.9	51
34	Polygenic Adaptation to an Environmental Shift: Temporal Dynamics of Variation Under Gaussian Stabilizing Selection and Additive Effects on a Single Trait. Genetics, 2019, 213, 1513-1530.	2.9	49
35	Identification by full-coverage array CGH of human DNA copy number increases relative to chimpanzee and gorilla. Genome Research, 2006, 16, 173-181.	5.5	48
36	Genetic architecture and selective sweeps after polygenic adaptation to distant trait optima. PLoS Genetics, 2018, 14, e1007794.	3.5	48

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37	Excess of Amino Acid Substitutions Relative to Polymorphism Between X-Linked Duplications in Drosophila melanogaster. Molecular Biology and Evolution, 2004, 22, 273-284.	8.9	46
38	Validation of Rearrangement Break Points Identified by Paired-End Sequencing in Natural Populations of Drosophila melanogaster. Genome Biology and Evolution, 2010, 2, 83-101.	2.5	46
39	Gene Expression Variation in <i>Drosophila melanogaster</i> Due to Rare Transposable Element Insertion Alleles of Large Effect. Genetics, 2015, 199, 85-93.	2.9	45
40	Recombination and the Properties of Tajima's D in the Context of Approximate-Likelihood Calculation. Genetics, 2005, 171, 2143-2148.	2.9	35
41	Automating approximate Bayesian computation by local linear regression. BMC Genetics, 2009, 10, 35.	2.7	33
42	A Model of Compound Heterozygous, Loss-of-Function Alleles Is Broadly Consistent with Observations from Complex-Disease GWAS Datasets. PLoS Genetics, 2017, 13, e1006573.	3.5	31
43	The Neutral Coalescent Process for Recent Gene Duplications and Copy-Number Variants. Genetics, 2007, 177, 987-1000.	2.9	26
44	Nucleotide Variation and Recombination Along the Fourth Chromosome in Drosophila simulans. Genetics, 2004, 166, 1783-1794.	2.9	25
45	Tandem Duplications and the Limits of Natural Selection in Drosophila yakuba and Drosophila simulans. PLoS ONE, 2015, 10, e0132184.	2.5	25
46	Revised Annotations, Sex-Biased Expression, and Lineage-Specific Genes in the <i>Drosophila melanogaster</i> Group. G3: Genes, Genomes, Genetics, 2014, 4, 2345-2351.	1.8	17
47	Inferring Selection in Partially Sequenced Regions. Molecular Biology and Evolution, 2008, 25, 438-446.	8.9	13
48	Nucleotide Variation and Recombination Along the Fourth Chromosome in <i>Drosophila simulans</i> . Genetics, 2004, 166, 1783-1794.	2.9	13
49	Experiments with the Site Frequency Spectrum. Bulletin of Mathematical Biology, 2011, 73, 829-872.	1.9	11
50	Efficient Software for Multi-marker, Region-Based Analysis of GWAS Data. G3: Genes, Genomes, Genetics, 2016, 6, 1023-1030.	1.8	6
51	EXTENSIVE INTROGRESSION OF MITOCHONDRIAL DNA RELATIVE TO NUCLEAR GENES IN THE DROSOPHILA YAKUBA SPECIES GROUP. Evolution; International Journal of Organic Evolution, 2006, 60, 292.	2.3	4