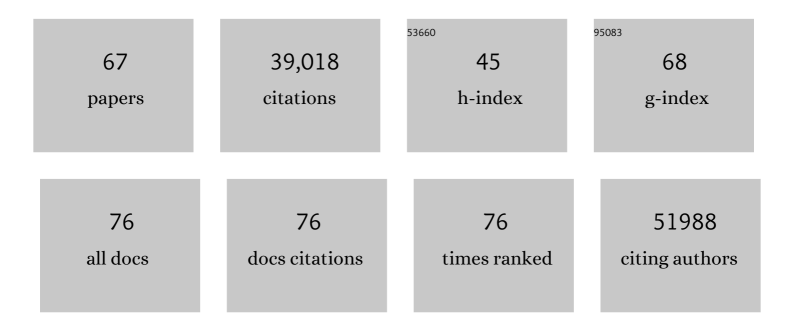
Kim M Rutherford

List of Publications by Year in descending order

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KIM M RUTHEREORD

#	Article	IF	CITATIONS
1	Genome sequence of the human malaria parasite Plasmodium falciparum. Nature, 2002, 419, 498-511.	13.7	3,881
2	The Gene Ontology Resource: 20 years and still GOing strong. Nucleic Acids Research, 2019, 47, D330-D338.	6.5	3,474
3	Complete genome sequence of the model actinomycete Streptomyces coelicolor A3(2). Nature, 2002, 417, 141-147.	13.7	2,940
4	Artemis: sequence visualization and annotation. Bioinformatics, 2000, 16, 944-945.	1.8	2,912
5	Gene Ontology Consortium: going forward. Nucleic Acids Research, 2015, 43, D1049-D1056.	6.5	2,743
6	The Gene Ontology resource: enriching a GOld mine. Nucleic Acids Research, 2021, 49, D325-D334.	6.5	2,416
7	The genome sequence of the food-borne pathogen Campylobacter jejuni reveals hypervariable sequences. Nature, 2000, 403, 665-668.	13.7	1,869
8	Expansion of the Gene Ontology knowledgebase and resources. Nucleic Acids Research, 2017, 45, D331-D338.	6.5	1,838
9	Massive gene decay in the leprosy bacillus. Nature, 2001, 409, 1007-1011.	13.7	1,607
10	ACT: the Artemis comparison tool. Bioinformatics, 2005, 21, 3422-3423.	1.8	1,536
11	The genome sequence of Schizosaccharomyces pombe. Nature, 2002, 415, 871-880.	13.7	1,508
12	Complete genome sequence of a multiple drug resistant Salmonella enterica serovar Typhi CT18. Nature, 2001, 413, 848-852.	13.7	1,192
13	Genome sequence of Yersinia pestis, the causative agent of plague. Nature, 2001, 413, 523-527.	13.7	1,144
14	Identification of Functional Elements and Regulatory Circuits by <i>Drosophila</i> modENCODE. Science, 2010, 330, 1787-1797.	6.0	1,124
15	Integrative Analysis of the <i>Caenorhabditis elegans</i> Genome by the modENCODE Project. Science, 2010, 330, 1775-1787.	6.0	912
16	Complete genomes of two clinical Staphylococcus aureus strains: Evidence for the rapid evolution of virulence and drug resistance. Proceedings of the National Academy of Sciences of the United States of America, 2004, 101, 9786-9791.	3.3	830
17	A Comprehensive Survey of the Plasmodium Life Cycle by Genomic, Transcriptomic, and Proteomic Analyses. Science, 2005, 307, 82-86.	6.0	743
18	Complete DNA sequence of a serogroup A strain of Neisseria meningitidis Z2491. Nature, 2000, 404, 502-506.	13.7	687

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19	Genomic plasticity of the causative agent of melioidosis, Burkholderia pseudomallei. Proceedings of the United States of America, 2004, 101, 14240-14245.	3.3	675
20	Gene Ontology Annotations and Resources. Nucleic Acids Research, 2012, 41, D530-D535.	6.5	456
21	FlyMine: an integrated database for Drosophila and Anopheles genomics. Genome Biology, 2007, 8, R129.	13.9	345
22	A Human-Curated Annotation of the Candida albicans Genome. PLoS Genetics, 2005, 1, e1.	1.5	293
23	PomBase: a comprehensive online resource for fission yeast. Nucleic Acids Research, 2012, 40, D695-D699.	6.5	288
24	The complete genome sequence and analysis of Corynebacterium diphtheriae NCTC13129. Nucleic Acids Research, 2003, 31, 6516-6523.	6.5	285
25	Dense sampling of bird diversity increases power of comparative genomics. Nature, 2020, 587, 252-257.	13.7	251
26	PHI-base: a new interface and further additions for the multi-species pathogen–host interactions database. Nucleic Acids Research, 2017, 45, D604-D610.	6.5	231
27	InterMine: a flexible data warehouse system for the integration and analysis of heterogeneous biological data. Bioinformatics, 2012, 28, 3163-3165.	1.8	229
28	GeneDB: a resource for prokaryotic and eukaryotic organisms. Nucleic Acids Research, 2004, 32, 339D-343.	6.5	199
29	The Gene Ontology: enhancements for 2011. Nucleic Acids Research, 2012, 40, D559-D564.	6.5	191
30	PomBase 2018: user-driven reimplementation of the fission yeast database provides rapid and intuitive access to diverse, interconnected information. Nucleic Acids Research, 2019, 47, D821-D827.	6.5	157
31	Sequence of Plasmodium falciparum chromosomes 1, 3–9 and 13. Nature, 2002, 419, 527-531.	13.7	156
32	RNAcentral: a hub of information for non-coding RNA sequences. Nucleic Acids Research, 2019, 47, D221-D229.	6.5	153
33	PHI-base: the pathogen–host interactions database. Nucleic Acids Research, 2020, 48, D613-D620.	6.5	145
34	Viewing and annotating sequence data with Artemis. Briefings in Bioinformatics, 2003, 4, 124-132.	3.2	136
35	modMine: flexible access to modENCODE data. Nucleic Acids Research, 2012, 40, D1082-D1088.	6.5	126
36	The tuatara genome reveals ancient features of amniote evolution. Nature, 2020, 584, 403-409.	13.7	105

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37	Large-scale transcriptome sequencing reveals novel expression patterns for key sex-related genes in a sex-changing fish. Biology of Sex Differences, 2015, 6, 26.	1.8	100
38	Stress, novel sex genes, and epigenetic reprogramming orchestrate socially controlled sex change. Science Advances, 2019, 5, eaaw7006.	4.7	99
39	PomBase 2015: updates to the fission yeast database. Nucleic Acids Research, 2015, 43, D656-D661.	6.5	95
40	WebACT-an online companion for the Artemis Comparison Tool. Bioinformatics, 2005, 21, 3665-3666.	1.8	84
41	Hidden in plain sight: what remains to be discovered in the eukaryotic proteome?. Open Biology, 2019, 9, 180241.	1.5	80
42	A Re-Annotation of theSaccharomyces cerevisiaeGenome. Comparative and Functional Genomics, 2001, 2, 143-154.	2.0	72
43	Fission stories: using PomBase to understand <i>Schizosaccharomyces pombe</i> biology. Genetics, 2022, 220, .	1.2	60
44	The DNA sequence of chromosome I of an African trypanosome: gene content, chromosome organisation, recombination and polymorphism. Nucleic Acids Research, 2003, 31, 4864-4873.	6.5	56
45	PHI-base in 2022: a multi-species phenotype database for Pathogen–Host Interactions. Nucleic Acids Research, 2022, 50, D837-D847.	6.5	53
46	Histological and transcriptomic effects of 17α-methyltestosterone on zebrafish gonad development. BMC Genomics, 2017, 18, 557.	1.2	52
47	Canto: an online tool for community literature curation. Bioinformatics, 2014, 30, 1791-1792.	1.8	41
48	De novo draft assembly of the Botrylloides leachii genome provides further insight into tunicate evolution. Scientific Reports, 2018, 8, 5518.	1.6	36
49	Uncovering the pathways underlying whole body regeneration in a chordate model, Botrylloides leachi using de novo transcriptome analysis. BMC Genomics, 2016, 17, 114.	1.2	34
50	PomBase: The Scientific Resource for Fission Yeast. Methods in Molecular Biology, 2018, 1757, 49-68.	0.4	32
51	Female Mimicry by Sneaker Males Has a Transcriptomic Signature in Both the Brain and the Gonad in a Sex-Changing Fish. Molecular Biology and Evolution, 2018, 35, 225-241.	3.5	29
52	Analysis of the genome of the New Zealand giant collembolan (Holacanthella duospinosa) sheds light on hexapod evolution. BMC Genomics, 2017, 18, 795.	1.2	28
53	Improving functional annotation for industrial microbes: a case study with Pichia pastoris. Trends in Biotechnology, 2014, 32, 396-399.	4.9	23
54	Identification of sex differences in zebrafish (Danio rerio) brains during early sexual differentiation and masculinization using 17α-methyltestoteroneâ€. Biology of Reproduction, 2018, 99, 446-460.	1.2	21

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55	JaponicusDB: rapid deployment of a model organism database for an emerging model species. Genetics, 2022, 220, .	1.2	21
56	Molecular evolution of <i>Dmrt1</i> accompanies change of sex-determining mechanisms in reptilia. Biology Letters, 2014, 10, 20140809.	1.0	20
57	Community curation in PomBase: enabling fission yeast experts to provide detailed, standardized, sharable annotation from research publications. Database: the Journal of Biological Databases and Curation, 2020, 2020, .	1.4	19
58	Genetic sex assignment in wild populations using genotypingâ€byâ€sequencing data: A statistical threshold approach. Molecular Ecology Resources, 2018, 18, 179-190.	2.2	17
59	Adipose transcriptome analysis provides novel insights into molecular regulation of prolonged fasting in northern elephant seal pups. Physiological Genomics, 2018, 50, 495-503.	1.0	15
60	DNA from mollusc shell: a valuable and underutilised substrate for genetic analyses. PeerJ, 2020, 8, e9420.	0.9	14
61	Molecular structure of sauropsid β-keratins from tuatara (Sphenodon punctatus). Journal of Structural Biology, 2019, 207, 21-28.	1.3	13
62	Male–female relatedness at specific SNP-linkage groups influences cryptic female choice in Chinook salmon (<i>Oncorhynchus tshawytscha</i>). Proceedings of the Royal Society B: Biological Sciences, 2017, 284, 20170853.	1.2	9
63	Reduced representation sequencing detects only subtle regional structure in a heavily exploited and rapidly recolonizing marine mammal species. Ecology and Evolution, 2018, 8, 8736-8749.	0.8	9
64	Term Matrix: a novel Gene Ontology annotation quality control system based on ontology term co-annotation patterns. Open Biology, 2020, 10, 200149.	1.5	7
65	Analysis of 114 kb of DNA sequence from fission yeast chromosome 2 immediately centromere-distal tohis5. Yeast, 2000, 16, 1405-1411.	0.8	6
66	Evolutionary history of the podoplanin gene. Gene Reports, 2018, 13, 28-37.	0.4	3
67	A genomeâ€wide investigation of adaptive signatures in proteinâ€coding genes related to tool behaviour in New Caledonian and Hawaiian crows. Molecular Ecology, 2021, 30, 973-986.	2.0	2