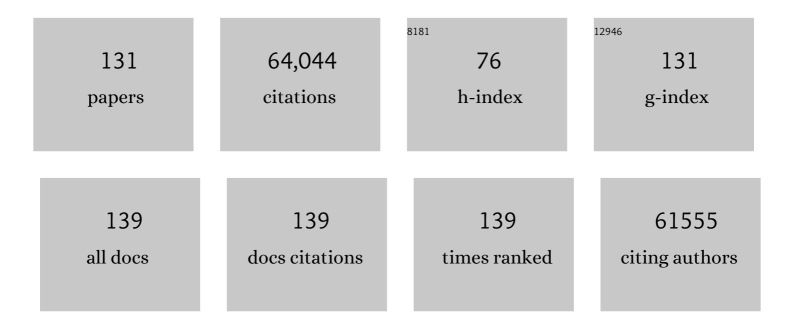
Jane Grimwood

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	Novel metabolic interactions and environmental conditions mediate the boreal peatmoss-cyanobacteria mutualism. ISME Journal, 2022, 16, 1074-1085.	9.8	25
2	Multiplex knockout of trichome-regulating MYB duplicates in hybrid poplar using a single gRNA. Plant Physiology, 2022, 189, 516-526.	4.8	18
3	The Common Bean V Gene Encodes Flavonoid 3′5′ Hydroxylase: A Major Mutational Target for Flavonoid Diversity in Angiosperms. Frontiers in Plant Science, 2022, 13, 869582.	3.6	7
4	eP144: Long-read genome sequencing secondary processing pipelines provide variant call accuracy that exceeds current clinical standards for short-read genome sequencing. Genetics in Medicine, 2022, 24, S89.	2.4	0
5	Habitatâ€adapted microbial communities mediate <i>Sphagnum</i> peatmoss resilience to warming. New Phytologist, 2022, 234, 2111-2125.	7.3	18
6	Phylogenomics of the genus Glycine sheds light on polyploid evolution and life-strategy transition. Nature Plants, 2022, 8, 233-244.	9.3	26
7	A generalist–specialist trade-off between switchgrass cytotypes impacts climate adaptation and geographic range. Proceedings of the National Academy of Sciences of the United States of America, 2022, 119, e2118879119.	7.1	5
8	The recent evolutionary rescue of a staple crop depended on over half a century of global germplasm exchange. Science Advances, 2022, 8, eabj4633.	10.3	9
9	Genomic mechanisms of climate adaptation in polyploid bioenergy switchgrass. Nature, 2021, 590, 438-444.	27.8	144
10	Widespread polycistronic gene expression in green algae. Proceedings of the National Academy of Sciences of the United States of America, 2021, 118, .	7.1	30
11	Gene disruption by structural mutations drives selection in US rice breeding over the last century. PLoS Genetics, 2021, 17, e1009389.	3.5	6
12	Long-read sequence assembly: a technical evaluation in barley. Plant Cell, 2021, 33, 1888-1906.	6.6	180
13	LCM and RNA-seq analyses revealed roles of cell cycle and translational regulation and homoeolog expression bias in cotton fiber cell initiation. BMC Genomics, 2021, 22, 309.	2.8	7
14	The tepary bean genome provides insight into evolution and domestication under heat stress. Nature Communications, 2021, 12, 2638.	12.8	43
15	Pests, diseases, and aridity have shaped the genome of Corymbia citriodora. Communications Biology, 2021, 4, 537.	4.4	21
16	Gene-rich UV sex chromosomes harbor conserved regulators of sexual development. Science Advances, 2021, 7, .	10.3	53
17	Genomic variation within the maize stiffâ€stalk heterotic germplasm pool. Plant Genome, 2021, 14, e20114.	2.8	14
18	Four chromosome scale genomes and a pan-genome annotation to accelerate pecan tree breeding. Nature Communications, 2021, 12, 4125.	12.8	49

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19	Comparative Transcriptomics of Non-Embryogenic and Embryogenic Callus in Semi-Recalcitrant and Non-Recalcitrant Upland Cotton Lines. Plants, 2021, 10, 1775.	3.5	10
20	Legacy genetics of <i>Arachis cardenasii</i> in the peanut crop shows the profound benefits of international seed exchange. Proceedings of the National Academy of Sciences of the United States of America, 2021, 118, .	7.1	18
21	A draft genome provides hypotheses on drought tolerance in a keystone plant species in Western North America threatened by climate change. Ecology and Evolution, 2021, 11, 15417-15429.	1.9	6
22	PEATmoss (<i>Physcomitrella</i> Expression Atlas Tool): a unified gene expression atlas for the model plant <i>Physcomitrella patens</i> . Plant Journal, 2020, 102, 165-177.	5.7	74
23	A genome assembly and the somatic genetic and epigenetic mutation rate in a wild long-lived perennial Populus trichocarpa. Genome Biology, 2020, 21, 259.	8.8	68
24	A genome resource for green millet Setaria viridis enables discovery of agronomically valuable loci. Nature Biotechnology, 2020, 38, 1203-1210.	17.5	103
25	The barley pan-genome reveals the hidden legacy of mutation breeding. Nature, 2020, 588, 284-289.	27.8	314
26	Genome biology of the paleotetraploid perennial biomass crop Miscanthus. Nature Communications, 2020, 11, 5442.	12.8	67
27	Chloroplast genome sequences of Carya illinoinensis from two distinct geographic populations. Tree Genetics and Genomes, 2020, 16, 1.	1.6	9
28	Genome Sequence of the Chestnut Blight Fungus <i>Cryphonectria parasitica</i> EP155: A Fundamental Resource for an Archetypical Invasive Plant Pathogen. Phytopathology, 2020, 110, 1180-1188.	2.2	34
29	Genomic diversifications of five Cossypium allopolyploid species and their impact on cotton improvement. Nature Genetics, 2020, 52, 525-533.	21.4	249
30	Construction and comparison of three referenceâ€quality genome assemblies for soybean. Plant Journal, 2019, 100, 1066-1082.	5.7	113
31	Genome mapping of quantitative trait loci (QTL) controlling domestication traits of intermediate wheatgrass (Thinopyrum intermedium). Theoretical and Applied Genetics, 2019, 132, 2325-2351.	3.6	30
32	The genome sequence of segmental allotetraploid peanut Arachis hypogaea. Nature Genetics, 2019, 51, 877-884.	21.4	439
33	A chromosome-scale genome assembly and dense genetic map for Xenopus tropicalis. Developmental Biology, 2019, 452, 8-20.	2.0	43
34	Genome sequence of the model rice variety KitaakeX. BMC Genomics, 2019, 20, 905.	2.8	59
35	The <i>Physcomitrella patens</i> gene atlas project: largeâ€scale <scp>RNA</scp> â€seq based expression data. Plant Journal, 2018, 95, 168-182.	5.7	115
36	Exploring the loblolly pine (Pinus taeda L.) genome by BAC sequencing and Cot analysis. Gene, 2018, 663, 165-177.	2.2	13

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37	The <i>Physcomitrella patens</i> chromosomeâ€scale assembly reveals moss genome structure and evolution. Plant Journal, 2018, 93, 515-533.	5.7	406
38	The <i>Sorghum bicolor</i> reference genome: improved assembly, gene annotations, a transcriptome atlas, and signatures of genome organization. Plant Journal, 2018, 93, 338-354.	5.7	431
39	Hardwood Tree Genomics: Unlocking Woody Plant Biology. Frontiers in Plant Science, 2018, 9, 1799.	3.6	50
40	The genomic landscape of molecular responses to natural drought stress in Panicum hallii. Nature Communications, 2018, 9, 5213.	12.8	101
41	The Aquilegia genome provides insight into adaptive radiation and reveals an extraordinarily polymorphic chromosome with a unique history. ELife, 2018, 7, .	6.0	120
42	Divergent cytosine DNA methylation patterns in single ell, soybean root hairs. New Phytologist, 2017, 214, 808-819.	7.3	75
43	Insights into Land Plant Evolution Garnered from the Marchantia polymorpha Genome. Cell, 2017, 171, 287-304.e15.	28.9	973
44	Insights into the red algae and eukaryotic evolution from the genome of <i>Porphyra umbilicalis</i> (Bangiophyceae, Rhodophyta). Proceedings of the National Academy of Sciences of the United States of America, 2017, 114, E6361-E6370.	7.1	233
45	Sub genome anchored physical frameworks of the allotetraploid Upland cotton (Gossypium hirsutum) Tj ETQq1 1 7, 15274.	0.784314 3.3	rgBT /Over 23
46	Genome organization of the vg1 and nodal3 gene clusters in the allotetraploid frog Xenopus laevis. Developmental Biology, 2017, 426, 236-244.	2.0	4
47	Transposable Elements versus the Fungal Genome: Impact on Whole-Genome Architecture and Transcriptional Profiles. PLoS Genetics, 2016, 12, e1006108.	3.5	177
48	Expansion of Signal Transduction Pathways in Fungi by Extensive Genome Duplication. Current Biology, 2016, 26, 1577-1584.	3.9	175
49	Sequencing wild and cultivated cassava and related species reveals extensive interspecific hybridization and genetic diversity. Nature Biotechnology, 2016, 34, 562-570.	17.5	340
50	On the origin and evolutionary consequences of gene body DNA methylation. Proceedings of the National Academy of Sciences of the United States of America, 2016, 113, 9111-9116.	7.1	260
51	Empty Niches after Extinctions Increase Population Sizes of Modern Corals. Current Biology, 2016, 26, 3190-3194.	3.9	79
52	Genome evolution in the allotetraploid frog Xenopus laevis. Nature, 2016, 538, 336-343.	27.8	849
53	Genetic analysis of Physcomitrella patens identifies ABSCISIC ACID NON-RESPONSIVE (ANR), a regulator of ABA responses unique to basal land plants and required for desiccation tolerance. Plant Cell, 2016, 28, tpc.00091.2016.	6.6	98
54	The genome of the seagrass Zostera marina reveals angiosperm adaptation to the sea. Nature, 2016, 530, 331-335.	27.8	460

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55	Comprehensive analysis of Hox gene expression in the amphipod crustacean Parhyale hawaiensis. Developmental Biology, 2016, 409, 297-309.	2.0	51
56	Evolving New Skeletal Traits by cis -Regulatory Changes in Bone Morphogenetic Proteins. Cell, 2016, 164, 45-56.	28.9	132
57	Targeted Switchgrass BAC Library Screening and Sequence Analysis Identifies Predicted Biomass and Stress Response-Related Genes. Bioenergy Research, 2016, 9, 109-122.	3.9	10
58	Genome Analysis of Planctomycetes Inhabiting Blades of the Red Alga Porphyra umbilicalis. PLoS ONE, 2016, 11, e0151883.	2.5	39
59	A Comparative Epigenomic Analysis of Polyploidy-Derived Genes in Soybean and Common Bean. Plant Physiology, 2015, 168, 1433-1447.	4.8	88
60	Sequencing of 15Â622 geneâ€bearing BAC s clarifies the geneâ€dense regions of the barley genome. Plant Journal, 2015, 84, 216-227.	5.7	36
61	Complete Plastome Sequences from <i>Glycine syndetika</i> and Six Additional Perennial Wild Relatives of Soybean. G3: Genes, Genomes, Genetics, 2014, 4, 2023-2033.	1.8	26
62	Sequencing of diverse mandarin, pummelo and orange genomes reveals complex history of admixture during citrus domestication. Nature Biotechnology, 2014, 32, 656-662.	17.5	572
63	The genome of Eucalyptus grandis. Nature, 2014, 510, 356-362.	27.8	725
64	The Chlamydomonas genome project: a decade on. Trends in Plant Science, 2014, 19, 672-680.	8.8	145
65	A reference genome for common bean and genome-wide analysis of dual domestications. Nature Genetics, 2014, 46, 707-713.	21.4	1,159
66	Insights into bilaterian evolution from three spiralian genomes. Nature, 2013, 493, 526-531.	27.8	564
67	The high-quality draft genome of peach (Prunus persica) identifies unique patterns of genetic diversity, domestication and genome evolution. Nature Genetics, 2013, 45, 487-494.	21.4	1,031
68	The Capsella rubella genome and the genomic consequences of rapid mating system evolution. Nature Genetics, 2013, 45, 831-835.	21.4	374
69	The Reference Genome of the Halophytic Plant Eutrema salsugineum. Frontiers in Plant Science, 2013, 4, 46.	3.6	198
70	Comparative Genome Structure, Secondary Metabolite, and Effector Coding Capacity across Cochliobolus Pathogens. PLoS Genetics, 2013, 9, e1003233.	3.5	232
71	Development of an integrated transcript sequence database and a gene expression atlas for gene discovery and analysis in switchgrass (<i>Panicum virgatum</i> L.). Plant Journal, 2013, 74, 160-173.	5.7	70
72	Genome sequence of the button mushroom <i>Agaricus bisporus</i> reveals mechanisms governing adaptation to a humic-rich ecological niche. Proceedings of the National Academy of Sciences of the United States of America, 2012, 109, 17501-17506.	7.1	359

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73	Algal genomes reveal evolutionary mosaicism and the fate of nucleomorphs. Nature, 2012, 492, 59-65.	27.8	377
74	Insight into tradeâ€off between wood decay and parasitism from the genome of a fungal forest pathogen. New Phytologist, 2012, 194, 1001-1013.	7.3	210
75	Repeated polyploidization of Gossypium genomes and the evolution of spinnable cotton fibres. Nature, 2012, 492, 423-427.	27.8	1,204
76	A Genome-Wide Survey of Switchgrass Genome Structure and Organization. PLoS ONE, 2012, 7, e33892.	2.5	50
77	The genomic basis of adaptive evolution in threespine sticklebacks. Nature, 2012, 484, 55-61.	27.8	1,600
78	Reference genome sequence of the model plant Setaria. Nature Biotechnology, 2012, 30, 555-561.	17.5	864
79	A Genome-wide SNP Genotyping Array Reveals Patterns of Global and Repeated Species-Pair Divergence in Sticklebacks. Current Biology, 2012, 22, 83-90.	3.9	212
80	Comparative genomics of citric-acid-producing <i>Aspergillus niger</i> ATCC 1015 versus enzyme-producing CBS 513.88. Genome Research, 2011, 21, 885-897.	5.5	329
81	Comparative genomic analysis of the thermophilic biomass-degrading fungi Myceliophthora thermophila and Thielavia terrestris. Nature Biotechnology, 2011, 29, 922-927.	17.5	428
82	The Arabidopsis lyrata genome sequence and the basis of rapid genome size change. Nature Genetics, 2011, 43, 476-481.	21.4	814
83	Finished Genome of the Fungal Wheat Pathogen Mycosphaerella graminicola Reveals Dispensome Structure, Chromosome Plasticity, and Stealth Pathogenesis. PLoS Genetics, 2011, 7, e1002070.	3.5	532
84	The GC-Rich Mitochondrial and Plastid Genomes of the Green Alga Coccomyxa Give Insight into the Evolution of Organelle DNA Nucleotide Landscape. PLoS ONE, 2011, 6, e23624.	2.5	53
85	Hydractinia Allodeterminant alr1 Resides in an Immunoglobulin Superfamily-like Gene Complex. Current Biology, 2010, 20, 1122-1127.	3.9	61
86	Genome sequence of the palaeopolyploid soybean. Nature, 2010, 463, 178-183.	27.8	3,854
87	Complete HOX cluster characterization of the coelacanth provides further evidence for slow evolution of its genome. Proceedings of the National Academy of Sciences of the United States of America, 2010, 107, 3622-3627.	7.1	65
88	Adaptive Evolution of Pelvic Reduction in Sticklebacks by Recurrent Deletion of a <i>Pitx1</i> Enhancer. Science, 2010, 327, 302-305.	12.6	901
89	The Genome of the Western Clawed Frog <i>Xenopus tropicalis</i> . Science, 2010, 328, 633-636.	12.6	708
90	Evolution of an Expanded Sex-Determining Locus in <i>Volvox</i> . Science, 2010, 328, 351-354.	12.6	159

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91	Genomic Analysis of Organismal Complexity in the Multicellular Green Alga <i>Volvox carteri</i> . Science, 2010, 329, 223-226.	12.6	536
92	The completion of the Mammalian Gene Collection (MGC). Genome Research, 2009, 19, 2324-2333.	5.5	125
93	The Genome of Nectria haematococca: Contribution of Supernumerary Chromosomes to Gene Expansion. PLoS Genetics, 2009, 5, e1000618.	3.5	402
94	A Hypervariable Invertebrate Allodeterminant. Current Biology, 2009, 19, 583-589.	3.9	95
95	The Sorghum bicolor genome and the diversification of grasses. Nature, 2009, 457, 551-556.	27.8	2,642
96	A role for a neo-sex chromosome in stickleback speciation. Nature, 2009, 461, 1079-1083.	27.8	327
97	Green Evolution and Dynamic Adaptations Revealed by Genomes of the Marine Picoeukaryotes <i>Micromonas</i> . Science, 2009, 324, 268-272.	12.6	591
98	The Trichoplax genome and the nature of placozoans. Nature, 2008, 454, 955-960.	27.8	801
99	The Phaeodactylum genome reveals the evolutionary history of diatom genomes. Nature, 2008, 456, 239-244.	27.8	1,458
100	The tiny eukaryote Ostreococcus provides genomic insights into the paradox of plankton speciation. Proceedings of the National Academy of Sciences of the United States of America, 2007, 104, 7705-7710.	7.1	563
101	Genome sequence of the lignocellulose-bioconverting and xylose-fermenting yeast Pichia stipitis. Nature Biotechnology, 2007, 25, 319-326.	17.5	449
102	A physical map of the highly heterozygous Populus genome: integration with the genome sequence and genetic map and analysis of haplotype variation. Plant Journal, 2007, 50, 1063-1078.	5.7	70
103	The <i>Chlamydomonas</i> Genome Reveals the Evolution of Key Animal and Plant Functions. Science, 2007, 318, 245-250.	12.6	2,354
104	Sequencing and analysis of 10,967 full-length cDNA clones from Xenopus laevis and Xenopus tropicalis reveals post-tetraploidization transcriptome remodeling. Genome Research, 2006, 16, 796-803.	5.5	73
105	Toward a Reference Sequence of the Soybean Genome: A Multiagency Effort. Crop Science, 2006, 46, S-55.	1.8	37
106	Widespread Parallel Evolution in Sticklebacks by Repeated Fixation of Ectodysplasin Alleles. Science, 2005, 307, 1928-1933.	12.6	1,299
107	Coelacanth genome sequence reveals the evolutionary history of vertebrate genes. Genome Research, 2004, 14, 2397-2405.	5.5	70
108	Closing the Gaps on Human Chromosome 19 Revealed Genes With a High Density of Repetitive Tandemly Arrayed Elements. Genome Research, 2004, 14, 239-246.	5.5	47

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109	The Status, Quality, and Expansion of the NIH Full-Length cDNA Project: The Mammalian Gene Collection (MGC). Genome Research, 2004, 14, 2121-2127.	5.5	486
110	Quality assessment of the human genome sequence. Nature, 2004, 429, 365-368.	27.8	172
111	The DNA sequence and biology of human chromosome 19. Nature, 2004, 428, 529-535.	27.8	298
112	The DNA sequence and comparative analysis of human chromosome 5. Nature, 2004, 431, 268-274.	27.8	102
113	The sequence and analysis of duplication-rich human chromosome 16. Nature, 2004, 432, 988-994.	27.8	156
114	Fowl sequence. Nature, 2004, 432, 679-680.	27.8	24
115	The Master Sex-Determination Locus in Threespine Sticklebacks Is on a Nascent Y Chromosome. Current Biology, 2004, 14, 1416-1424.	3.9	367
116	Gene Conversion and the Evolution of Protocadherin Gene Cluster Diversity. Genome Research, 2004, 14, 354-366.	5.5	100
117	New Genomic Tools for Molecular Studies of Evolutionary Change in Threespine Sticklebacks. Behaviour, 2004, 141, 1331-1344.	0.8	64
118	Longitudinal serum concentrations of placental growth factor: Evidence for abnormal placental angiogenesis in pathologic pregnancies. American Journal of Obstetrics and Gynecology, 2003, 188, 177-182.	1.3	373
119	Six is seventh. Nature, 2003, 425, 775-776.	27.8	8
120	Extensive Linkage Disequilibrium, a Common 16.7-Kilobase Deletion, and Evidence of Balancing Selection in the Human Protocadherin α Cluster. American Journal of Human Genetics, 2003, 72, 621-635.	6.2	51
121	Coding potential of laboratory and clinical strains of human cytomegalovirus. Proceedings of the National Academy of Sciences of the United States of America, 2003, 100, 14976-14981.	7.1	450
122	Generation and initial analysis of more than 15,000 full-length human and mouse cDNA sequences. Proceedings of the National Academy of Sciences of the United States of America, 2002, 99, 16899-16903.	7.1	1,610
123	Comparative DNA Sequence Analysis of Mouse and Human Protocadherin Gene Clusters. Genome Research, 2001, 11, 389-404.	5.5	224
124	Initial sequencing and analysis of the human genome. Nature, 2001, 409, 860-921.	27.8	21,074
125	Expression of Chlamydia pneumoniaePolymorphic Membrane Protein Family Genes. Infection and Immunity, 2001, 69, 2383-2389.	2.2	66
126	Attachment of Toxoplasma gondii to a Specific Membrane Fraction of CHO Cells. Infection and Immunity, 2000, 68, 7198-7201.	2.2	5

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127	Computational Analysis of the Polymorphic Membrane Protein Superfamily of <i>Chlamydia trachomatis</i> and <i>Chlamydia pneumoniae</i> . Microbial & Comparative Genomics, 1999, 4, 187-201.	0.4	95
128	Comparative genomes of Chlamydia pneumoniae and C. trachomatis. Nature Genetics, 1999, 21, 385-389.	21.4	636
129	Toxoplasma gondii: the role of parasite surface and secreted proteins in host cell invasion. International Journal for Parasitology, 1996, 26, 169-173.	3.1	104
130	Toxoplasma gondii: The role of a 30-kDa surface protein in host cell invasion. Experimental Parasitology, 1992, 74, 106-111.	1.2	98
131	Kinetics of the growth and variation in infectivity of <i>Toxoplasma gondii</i> in mice. Annals of Tropical Medicine and Parasitology, 1991, 85, 659-661.	1.6	8