## List of Publications by Year in descending order

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394421 395702 1,972 34 19 33 citations h-index g-index papers 35 35 35 3089 all docs docs citations times ranked citing authors

#	Article	IF	CITATIONS
1	Widespread Whole Genome Duplications Contribute to Genome Complexity and Species Diversity in Angiosperms. Molecular Plant, 2018, 11, 414-428.	8.3	251
2	RNA-seq analyses of multiple meristems of soybean: novel and alternative transcripts, evolutionary and functional implications. BMC Plant Biology, 2014, 14, 169.	3.6	229
3	Saccharina genomes provide novel insight into kelp biology. Nature Communications, 2015, 6, 6986.	12.8	222
4	Multiple Polyploidization Events across Asteraceae with Two Nested Events in the Early History Revealed by Nuclear Phylogenomics. Molecular Biology and Evolution, 2016, 33, 2820-2835.	8.9	149
5	The <scp>DYT</scp> 1â€interacting proteins b <scp>HLH</scp> 010, b <scp>HLH</scp> 089 and b <scp>HLH</scp> 091 are redundantly required for <scp>A</scp> rabidopsis anther development and transcriptome. Plant Journal, 2015, 83, 976-990.	5.7	136
6	Analysis of <i>Arabidopsis</i> genome-wide variations before and after meiosis and meiotic recombination by resequencing Landsberg <i>erecta</i> and all four products of a single meiosis. Genome Research, 2012, 22, 508-518.	5.5	125
7	Nuclear phylotranscriptomics and phylogenomics support numerous polyploidization events and hypotheses for the evolution of rhizobial nitrogen-fixing symbiosis in Fabaceae. Molecular Plant, 2021, 14, 748-773.	8.3	86
8	Asterid Phylogenomics/Phylotranscriptomics Uncover Morphological Evolutionary Histories and Support Phylogenetic Placement for Numerous Whole-Genome Duplications. Molecular Biology and Evolution, 2020, 37, 3188-3210.	8.9	82
9	A well-resolved fern nuclear phylogeny reveals the evolution history of numerous transcription factor families. Molecular Phylogenetics and Evolution, 2018, 127, 961-977.	2.7	80
10	inGAP-sv: a novel scheme to identify and visualize structural variation from paired end mapping data. Nucleic Acids Research, 2011, 39, W567-W575.	14.5	74
11	Characterization of meiotic crossovers and gene conversion by whole-genome sequencing in Saccharomyces cerevisiae. BMC Genomics, 2009, 10, 475.	2.8	62
12	inGAP: an integrated next-generation genome analysis pipeline. Bioinformatics, 2010, 26, 127-129.	4.1	58
13	Detection of genomic variations and DNA polymorphisms and impact on analysis of meiotic recombination and genetic mapping. Proceedings of the National Academy of Sciences of the United States of America, 2014, 111, 10007-10012.	7.1	55
14	H3K4me2 functions as a repressive epigenetic mark in plants. Epigenetics and Chromatin, 2019, 12, 40.	3.9	51
15	Alternative splicing during Arabidopsis flower development results in constitutive and stage-regulated isoforms. Frontiers in Genetics, 2014, 5, 25.	2.3	45
16	Recurrent genome duplication events likely contributed to both the ancient and recent rise of ferns. Journal of Integrative Plant Biology, 2020, 62, 433-455.	8.5	43
17	The genome and transcriptome of Trichormus sp. NMC-1: insights into adaptation to extreme environments on the Qinghai-Tibet Plateau. Scientific Reports, 2016, 6, 29404.	3.3	33
18	Association of the molecular regulation of ear leaf senescence/stress response and photosynthesis/metabolism with heterosis at the reproductive stage in maize. Scientific Reports, 2016, 6, 29843.	3.3	23

#	Article	IF	CITATIONS
19	Genomic analysis of field pennycress (Thlaspi arvense) provides insights into mechanisms of adaptation to high elevation. BMC Biology, 2021, 19, 143.	3.8	23
20	Modulation of evening complex activity enables north-to-south adaptation of soybean. Science China Life Sciences, 2021, 64, 179-195.	4.9	22
21	Changing Gly311 to an acidic amino acid in the MATE family protein DTX6 enhances Arabidopsis resistance to the dihydropyridine herbicides. Molecular Plant, 2021, 14, 2115-2125.	8.3	22
22	Analyses of functional conservation and divergence revealÂrequirement of bHLH010/089/091 for pollen development at elevated temperature in Arabidopsis. Journal of Genetics and Genomics, 2020, 47, 477-492.	3.9	13
23	inGAP-Family: Accurate Detection of Meiotic Recombination Loci and Causal Mutations by Filtering Out Artificial Variants due to Genome Complexities. Genomics, Proteomics and Bioinformatics, 2022, 20, 524-535.	6.9	13
24	Effects of parental genetic divergence on gene expression patterns in interspecific hybrids of Camellia. BMC Genomics, 2019, 20, 828.	2.8	11
25	The gut microbiome and microbial metabolites in acute myocardial infarction. Journal of Genetics and Genomics, 2022, 49, 569-578.	3.9	11
26	PhyloMCL: Accurate clustering of hierarchical orthogroups guided by phylogenetic relationship and inference of polyploidy events. Methods in Ecology and Evolution, 2020, 11, 943-954.	5.2	9
27	Polyphyly in 16S rRNA-based LVTree Versus Monophyly in Whole-genome-based CVTree. Genomics, Proteomics and Bioinformatics, 2018, 16, 310-319.	6.9	8
28	Comparative Genomic and Transcriptomic Analysis Reveals Specific Features of Gene Regulation in Kluyveromyces marxianus. Frontiers in Microbiology, 2021, 12, 598060.	3.5	8
29	Evolution of the leucineâ€rich repeat receptorâ€like protein kinase gene family: Ancestral copy number and functional divergence of <i>BAM1</i> and <i>BAM2</i> in Brassicaceae. Journal of Systematics and Evolution, 2016, 54, 204-218.	3.1	7
30	Amino acid based i>de Bruijn i>graph algorithm for identifying complete coding genes from metagenomic and metatranscriptomic short reads. Nucleic Acids Research, 2019, 47, e30-e30.	14.5	7
31	Reply to Zwaenepoel etÂal.: Meeting the Challenges of Detecting Polyploidy Events from Transcriptomic Data. Molecular Plant, 2019, 12, 137-140.	8.3	5
32	Cell free bacterial DNAs in human plasma provide fingerprints for immune-related diseases. Medicine in Microecology, 2020, 5, 100022.	1.6	3
33	Parental folate deficiency induces birth defects in mice accompanied with increased de novo mutations. Cell Discovery, 2022, 8, 18.	6.7	3
34	Histone demethylase IBM1-mediated meiocyte gene expression ensures meiotic chromosome synapsis and recombination. PLoS Genetics, 2022, 18, e1010041.	3.5	1