Yao-Cheng Lin

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/6888773/publications.pdf

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42 papers 10,812 citations

186265
28
h-index

254184 43 g-index

47 all docs

47 docs citations

times ranked

47

15001 citing authors

#	Article	IF	CITATIONS
1	Toward automatic plant phenotyping: starting from leaf counting. Multimedia Tools and Applications, 2022, 81, 11865-11879.	3.9	3
2	DNA-free CRISPR-Cas9 gene editing of wild tetraploid tomato <i>Solanum peruvianum</i> using protoplast regeneration. Plant Physiology, 2022, 188, 1917-1930.	4.8	39
3	Whole genome resequencing and complementation tests reveal candidate loci contributing to bacterial wilt (Ralstonia sp.) resistance in tomato. Scientific Reports, 2022, 12, 8374.	3.3	5
4	Efficient and Economical Targeted Insertion in Plant Genomes via Protoplast Regeneration. CRISPR Journal, 2021, 4, 752-760.	2.9	9
5	Comparative Genomic Analysis Uncovered Evolution of Pathogenicity Factors, Horizontal Gene Transfer Events, and Heavy Metal Resistance Traits in Citrus Canker Bacterium Xanthomonas citri subsp. citri. Frontiers in Microbiology, 2021, 12, 731711.	3.5	5
6	Automatic Leaf Counting Using Improved YOLOv3., 2020,,.		3
7	Small RNA profiling in Pinus pinaster reveals the transcriptome of developing seeds and highlights differences between zygotic and somatic embryos. Scientific Reports, 2019, 9, 11327.	3.3	28
8	Reference genome and comparative genome analysis for the WHO reference strain for Mycobacterium bovis BCG Danish, the present tuberculosis vaccine. BMC Genomics, 2019, 20, 561.	2.8	16
9	Draft Genome Assembly of the False Spider Mite Brevipalpus yothersi. Microbiology Resource Announcements, 2019, 8, .	0.6	6
10	Functional and evolutionary genomic inferences in <i>Populus</i> through genome and population sequencing of American and European aspen. Proceedings of the National Academy of Sciences of the United States of America, 2018, 115, E10970-E10978.	7.1	84
11	The Apostasia genome and the evolution of orchids. Nature, 2017, 549, 379-383.	27.8	305
12	Complete Genome Sequence of Bacillus cereus C1L, a Plant Growth-Promoting Rhizobacterium from the Rhizosphere of Formosa Lily in Taiwan. Genome Announcements, 2017, 5, .	0.8	2
13	Systems genetics reveals a transcriptional network associated with susceptibility in the maize–grey leaf spot pathosystem. Plant Journal, 2017, 89, 746-763.	5.7	49
14	Draft genome of Cercospora zeina, Fusarium pininemorale, Hawksworthiomyces lignivorus, Huntiella decipiens and Ophiostoma ips. IMA Fungus, 2017, 8, 385-396.	3.8	37
15	The genome of the seagrass Zostera marina reveals angiosperm adaptation to the sea. Nature, 2016, 530, 331-335.	27.8	460
16	Comparative in silico analysis of SSRs in coding regions of high confidence predicted genes in Norway spruce (Picea abies) and Loblolly pine (Pinus taeda). BMC Genetics, 2015, 16, 149.	2.7	3
17	Serendipitous Meta-Transcriptomics: The Fungal Community of Norway Spruce (Picea abies). PLoS ONE, 2015, 10, e0139080.	2.5	27
18	Genome-Wide Analysis Reveals Diverged Patterns of Codon Bias, Gene Expression, and Rates of Sequence Evolution in Picea Gene Families. Genome Biology and Evolution, 2015, 7, 1002-1015.	2.5	63

#	Article	lF	CITATIONS
19	Expanding the repertoire of secretory peptides controlling root development with comparative genome analysis and functional assays. Journal of Experimental Botany, 2015, 66, 5257-5269.	4.8	71
20	The Plant Genome Integrative Explorer Resource: PlantGen <scp>IE</scp> .org. New Phytologist, 2015, 208, 1149-1156.	7.3	282
21	Genome dynamics of the human embryonic kidney 293 lineage in response to cell biology manipulations. Nature Communications, 2014, 5, 4767.	12.8	421
22	miRNA profiling in leaf and cork tissues of Quercus suber reveals novel miRNAs and tissue-specific expression patterns. Tree Genetics and Genomes, 2014, 10, 721-737.	1.6	20
23	Comparative in silicoanalysis of EST-SSRs in angiosperm and gymnosperm tree genera. BMC Plant Biology, 2014, 14, 220.	3.6	29
24	Pan genome of the phytoplankton Emiliania underpins its global distribution. Nature, 2013, 499, 209-213.	27.8	448
25	SILAC-Based Proteome Analysis of <i>Starmerella bombicola</i> Sophorolipid Production. Journal of Proteome Research, 2013, 12, 4376-4392.	3.7	22
26	<i>Candida bombicola</i> as a platform organism for the production of tailorâ€made biomolecules. Biotechnology and Bioengineering, 2013, 110, 2494-2503.	3.3	42
27	The Norway spruce genome sequence and conifer genome evolution. Nature, 2013, 497, 579-584.	27.8	1,303
28	The biosynthetic gene cluster for sophorolipids: a biotechnological interesting biosurfactant produced by <i><scp>S</scp>tarmerella bombicola</i> <.i>Molecular Microbiology, 2013, 88, 501-509.	2.5	94
29	A Comprehensive Analysis of Genes Encoding Small Secreted Proteins Identifies Candidate Effectors in <i>Melampsora larici-populina</i> (Poplar Leaf Rust). Molecular Plant-Microbe Interactions, 2012, 25, 279-293.	2.6	150
30	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
31	GOLVEN Secretory Peptides Regulate Auxin Carrier Turnover during Plant Gravitropic Responses. Developmental Cell, 2012, 22, 678-685.	7.0	182
32	Obligate biotrophy features unraveled by the genomic analysis of rust fungi. Proceedings of the National Academy of Sciences of the United States of America, 2011, 108, 9166-9171.	7.1	640
33	Development and implementation of a highly-multiplexed SNP array for genetic mapping in maritime pine and comparative mapping with loblolly pine. BMC Genomics, 2011, 12, 368.	2.8	66
34	Oil Bodies and Oleosins in <i>Physcomitrella</i> Possess Characteristics Representative of Early Trends in Evolution Â. Plant Physiology, 2009, 150, 1192-1203.	4.8	73
35	Genome sequence of the recombinant protein production host Pichia pastoris. Nature Biotechnology, 2009, 27, 561-566.	17.5	422
36	Open access to sequence: Browsing the Pichia pastoris genome. Microbial Cell Factories, 2009, 8, 53.	4.0	55

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#	Article	IF	CITATIONS
37	The genome of Laccaria bicolor provides insights into mycorrhizal symbiosis. Nature, 2008, 452, 88-92.	27.8	1,003
38	The Rice Annotation Project Database (RAP-DB): 2008 update. Nucleic Acids Research, 2007, 36, D1028-D1033.	14.5	295
39	Curated genome annotation of Oryza sativa ssp. japonica and comparative genome analysis with Arabidopsis thaliana. Genome Research, 2007, 17, 175-183.	5.5	218
40	A rice gene activation/knockout mutant resource for high throughput functional genomics. Plant Molecular Biology, 2007, 63, 351-364.	3.9	197
41	A rice phenomics study—phenotype scoring and seed propagation of a T-DNA insertion-induced rice mutant population. Plant Molecular Biology, 2007, 65, 427-438.	3.9	52
42	The map-based sequence of the rice genome. Nature, 2005, 436, 793-800.	27.8	3,365