## Xiangfeng Wang

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/2567013/publications.pdf

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94381 6,384 71 37 citations h-index papers

72 g-index 76 76 76 10392 docs citations times ranked citing authors all docs

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#	Article	IF	CITATIONS
1	Genome optimization via virtual simulation to accelerate maize hybrid breeding. Briefings in Bioinformatics, 2022, 23, .	3.2	3
2	Resequencing 250 Soybean Accessions: New Insights into Genes Associated with Agronomic Traits and Genetic Networks. Genomics, Proteomics and Bioinformatics, 2022, 20, 29-41.	3.0	21
3	Correlation of vacuole morphology with stomatal lineage development by whole-cell electron tomography. Plant Physiology, 2022, 188, 2085-2100.	2.3	11
4	A dirigent family protein confers variation of Casparian strip thickness and salt tolerance in maize. Nature Communications, 2022, 13, 2222.	5.8	55
5	DatabaseÂResources of the National Genomics Data Center, China National Center for Bioinformation in 2021. Nucleic Acids Research, 2021, 49, D18-D28.	6.5	168
6	The microtubule-associated protein WDL4 modulates auxin distribution to promote apical hook opening in Arabidopsis. Plant Cell, 2021, 33, 1927-1944.	3.1	13
7	Metabolomicsâ€driven gene mining and genetic improvement of tolerance to saltâ€induced osmotic stress in maize. New Phytologist, 2021, 230, 2355-2370.	3.5	46
8	The E3 ligase MREL57 modulates microtubule stability and stomatal closure in response to ABA. Nature Communications, 2021, 12, 2181.	5.8	27
9	The genome of the warm-season turfgrass African bermudagrass ( <i>Cynodon transvaalensis</i> ). Horticulture Research, 2021, 8, 93.	2.9	19
10	Forecasting rice latitude adaptation through a daylength-sensing-based environment adaptation simulator. Nature Food, 2021, 2, 348-362.	6.2	16
11	The genetic mechanism of heterosis utilization in maize improvement. Genome Biology, 2021, 22, 148.	3.8	69
12	Genome-Wide Analysis of the HSP20 Gene Family and Expression Patterns of HSP20 Genes in Response to Abiotic Stresses in Cynodon transvaalensis. Frontiers in Genetics, 2021, 12, 732812.	1.1	11
13	LightGBM: accelerated genomically designed crop breeding through ensemble learning. Genome Biology, 2021, 22, 271.	3.8	76
14	Database Resources of the National Genomics Data Center in 2020. Nucleic Acids Research, 2020, 48, D24-D33.	6.5	165
15	The hybrid protein interactome contributes to rice heterosis as epistatic effects. Plant Journal, 2020, 102, 116-128.	2.8	10
16	Genome optimization for improvement of maize breeding. Theoretical and Applied Genetics, 2020, 133, 1491-1502.	1.8	26
17	The roles of endomembrane trafficking in plant abiotic stress responses. Journal of Integrative Plant Biology, 2020, 62, 55-69.	4.1	57
18	Genome Size Evolution Mediated by Gypsy Retrotransposons in Brassicaceae. Genomics, Proteomics and Bioinformatics, 2020, 18, 321-332.	3.0	21

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19	A QTL atlas for grain yield and its component traits in maize ( <i>Zea mays</i> ). Plant Breeding, 2020, 139, 562-574.	1.0	18
20	SR4R: An Integrative SNP Resource for Genomic Breeding and Population Research in Rice. Genomics, Proteomics and Bioinformatics, 2020, 18, 173-185.	3.0	18
21	IbBBX24 Promotes the Jasmonic Acid Pathway and Enhances Fusarium Wilt Resistance in Sweet Potato. Plant Cell, 2020, 32, 1102-1123.	3.1	65
22	Divergent selection and genetic introgression shape the genome landscape of heterosis in hybrid rice. Proceedings of the National Academy of Sciences of the United States of America, 2020, 117, 4623-4631.	3.3	46
23	AP3M harbors actin filament binding activity that is crucial for vacuole morphology and stomatal closure in <i>Arabidopsis</i> . Proceedings of the National Academy of Sciences of the United States of America, 2019, 116, 18132-18141.	3.3	17
24	Understanding the functions and mechanisms of plant cytoskeleton in response to environmental signals. Current Opinion in Plant Biology, 2019, 52, 86-96.	3.5	42
25	Redox-Mediated Endocytosis of a Receptor-Like Kinase during Distal Stem Cell Differentiation Depends on Its Tumor Necrosis Factor Receptor Domain. Plant Physiology, 2019, 181, 1075-1095.	2.3	11
26	<scp>BES</scp> 1 hinders <scp>ABSCISIC ACID INSENSITIVE</scp> 5 and promotes seed germination in <i>Arabidopsis</i> . New Phytologist, 2019, 221, 908-918.	3.5	86
27	Ethylene Signaling Modulates Cortical Microtubule Reassembly in Response to Salt Stress. Plant Physiology, 2018, 176, 2071-2081.	2.3	49
28	CGmapTools improves the precision of heterozygous SNV calls and supports allele-specific methylation detection and visualization in bisulfite-sequencing data. Bioinformatics, 2018, 34, 381-387.	1.8	152
29	Powdery Mildews Are Characterized by Contracted Carbohydrate Metabolism and Diverse Effectors to Adapt to Obligate Biotrophic Lifestyle. Frontiers in Microbiology, 2018, 9, 3160.	1.5	45
30	Machine Learning Models for Genetic Risk Assessment of Infants with Non-syndromic Orofacial Cleft. Genomics, Proteomics and Bioinformatics, 2018, 16, 354-364.	3.0	31
31	Opaque-2 Regulates a Complex Gene Network Associated with Cell Differentiation and Storage Functions of Maize Endosperm. Plant Cell, 2018, 30, 2425-2446.	3.1	83
32	VAMP711 Is Required for Abscisic Acid-Mediated Inhibition of Plasma Membrane H <sup>+</sup> -ATPase Activity. Plant Physiology, 2018, 178, 1332-1343.	2.3	47
33	Transcript profile analysis reveals important roles of jasmonic acid signalling pathway in the response of sweet potato to salt stress. Scientific Reports, 2017, 7, 40819.	1.6	62
34	Isoform Evolution in Primates through Independent Combination of Alternative RNA Processing Events. Molecular Biology and Evolution, 2017, 34, 2453-2468.	3.5	37
35	Transcriptome Association Identifies Regulators of Wheat Spike Architecture. Plant Physiology, 2017, 175, 746-757.	2.3	94
36	Single-Molecule Sequencing Assists Genome Assembly Improvement and Structural Variation Inference. Molecular Plant, 2016, 9, 1085-1087.	3.9	2

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37	Temporal small RNA transcriptome profiling unraveled partitioned miRNA expression in developing maize endosperms between reciprocal crosses. Frontiers in Plant Science, 2015, 6, 744.	1.7	14
38	RNA Sequencing of Laser-Capture Microdissected Compartments of the Maize Kernel Identifies Regulatory Modules Associated with Endosperm Cell Differentiation. Plant Cell, 2015, 27, 513-531.	3.1	206
39	Temporal patterns of gene expression in developing maize endosperm identified through transcriptome sequencing. Proceedings of the National Academy of Sciences of the United States of America, 2014, 111, 7582-7587.	3.3	146
40	Complete genome of Hainan papaya ringspot virus using small RNA deep sequencing. Virus Genes, 2014, 48, 502-508.	0.7	5
41	RNA-Seq analysis of laser-capture microdissected cells of the developing central starchy endosperm of maize. Genomics Data, 2014, 2, 242-245.	1.3	13
42	Machine Learning–Based Differential Network Analysis: A Study of Stress-Responsive Transcriptomes in <i>Arabidopsis</i> . Plant Cell, 2014, 26, 520-537.	3.1	104
43	Machine learning for Big Data analytics in plants. Trends in Plant Science, 2014, 19, 798-808.	4.3	220
44	Dynamic parent-of-origin effects on small interfering RNA expression in the developing maize endosperm. BMC Plant Biology, 2014, 14, 192.	1.6	21
45	A Computational Workflow to Identify Allele-specific Expression and Epigenetic Modification in Maize. Genomics, Proteomics and Bioinformatics, 2013, 11, 247-252.	3.0	10
46	Organ Evolution in Angiosperms Driven by Correlated Divergences of Gene Sequences and Expression Patterns Â. Plant Cell, 2013, 25, 71-82.	3.1	39
47	CrusView: A Java-Based Visualization Platform for Comparative Genomics Analyses in Brassicaceae Species. Plant Physiology, 2013, 163, 354-362.	2.3	33
48	The Reference Genome of the Halophytic Plant Eutrema salsugineum. Frontiers in Plant Science, 2013, 4, 46.	1.7	198
49	Dynamic Expression of Imprinted Genes Associates with Maternally Controlled Nutrient Allocation during Maize Endosperm Development. Plant Cell, 2013, 25, 3212-3227.	3.1	97
50	Application of the Gini Correlation Coefficient to Infer Regulatory Relationships in Transcriptome Analysis Â. Plant Physiology, 2012, 160, 192-203.	2.3	65
51	KGBassembler: a karyotype-based genome assembler for Brassicaceae species. Bioinformatics, 2012, 28, 3141-3143.	1.8	12
52	Transcriptome Comparison of Susceptible and Resistant Wheat in Response to Powdery Mildew Infection. Genomics, Proteomics and Bioinformatics, 2012, 10, 94-106.	3.0	90
53	Computational dissection of Arabidopsis smRNAome leads to discovery of novel microRNAs and short interfering RNAs associated with transcription start sites. Genomics, 2011, 97, 235-243.	1.3	8
54	Systematic Curation of miRBase Annotation Using Integrated Small RNA High-Throughput Sequencing Data for C. elegans and Drosophila. Frontiers in Genetics, 2011, 2, 25.	1.1	52

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55	Genomeâ€wide mapping of the <i>HY5</i> å€mediated genenetworks in Arabidopsis that involve both transcriptional and postâ€transcriptional regulation. Plant Journal, 2011, 65, 346-358.	2.8	274
56	Genome-wide profiling of histone H3 lysine 9 acetylation and dimethylation in Arabidopsis reveals correlation between multiple histone marks and gene expression. Plant Molecular Biology, 2010, 72, 585-595.	2.0	167
57	Development of a versatile, target-oriented tiling microarray assay for measuring allele-specific gene expression. Genomics, 2010, 96, 308-315.	1.3	4
58	Global Epigenetic and Transcriptional Trends among Two Rice Subspecies and Their Reciprocal Hybrids. Plant Cell, 2010, 22, 17-33.	3.1	514
59	Genome-Wide and Organ-Specific Landscapes of Epigenetic Modifications and Their Relationships to mRNA and Small RNA Transcriptomes in Maize. Plant Cell, 2009, 21, 1053-1069.	3.1	291
60	Transcriptionally active gene fragments derived from potentially fast-evolving donor genes in the rice genome. Bioinformatics, 2009, 25, 1215-1218.	1.8	3
61	Characterization of the genome expression trends in the heading-stage panicle of six rice lineages. Genomics, 2009, 93, 169-178.	1.3	11
62	Transcriptional analysis of highly syntenic regions between Medicago truncatula and Glycine max using tiling microarrays. Genome Biology, 2008, 9, R57.	13.9	13
63	High-Resolution Mapping of Epigenetic Modifications of the Rice Genome Uncovers Interplay between DNA Methylation, Histone Methylation, and Gene Expression. Plant Cell, 2008, 20, 259-276.	3.1	281
64	Global Identification and Characterization of Transcriptionally Active Regions in the Rice Genome. PLoS ONE, 2007, 2, e294.	1.1	82
65	Global genome expression analysis of rice in response to drought and high-salinity stresses in shoot, flag leaf, and panicle. Plant Molecular Biology, 2007, 63, 591-608.	2.0	275
66	Genome-wide transcription analyses in rice using tiling microarrays. Nature Genetics, 2006, 38, 124-129.	9.4	154
67	NMPP: a user-customized NimbleGen microarray data processing pipeline. Bioinformatics, 2006, 22, 2955-2957.	1.8	29
68	A Pilot Study of Transcription Unit Analysis in Rice Using Oligonucleotide Tiling-path Microarray. Plant Molecular Biology, 2005, 59, 137-149.	2.0	17
69	A Tiling Microarray Expression Analysis of Rice Chromosome 4 Suggests a Chromosome-Level Regulation of Transcription. Plant Cell, 2005, 17, 1641-1657.	3.1	56
70	A microarray analysis of the rice transcriptome and its comparison to Arabidopsis. Genome Research, 2005, 15, 1274-1283.	2.4	112
71	Tiling microarray analysis of rice chromosome 10 to identify the transcriptome and relate its expression to chromosomal architecture. Genome Biology, 2005, 6, R52.	13.9	41