Jue Ruan

List of Publications by Year in descending order

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ILLE RUAN

#	Article	IF	CITATIONS
1	The First High-quality Reference Genome of Sika Deer Provides Insights into High-tannin Adaptation. Genomics, Proteomics and Bioinformatics, 2023, 21, 203-215.	3.0	9
2	A Chromosome-level Genome Assembly of Wild Castor Provides New Insights into its Adaptive Evolution in Tropical Desert. Genomics, Proteomics and Bioinformatics, 2022, 20, 42-59.	3.0	18
3	En Route to Completion: What Is An Ideal Reference Genome?. Genomics, Proteomics and Bioinformatics, 2022, 20, 1-3.	3.0	0
4	The Cycas genome and the early evolution of seed plants. Nature Plants, 2022, 8, 389-401.	4.7	80
5	A super pan-genomic landscape of rice. Cell Research, 2022, 32, 878-896.	5.7	99
6	Transcriptome Analysis of Ginkgo biloba L. Leaves across Late Developmental Stages Based on RNA-Seq and Co-Expression Network. Forests, 2021, 12, 315.	0.9	1
7	African lungfish genome sheds light on the vertebrate water-to-land transition. Cell, 2021, 184, 1362-1376.e18.	13.5	99
8	The nearly complete genome of Ginkgo biloba illuminates gymnosperm evolution. Nature Plants, 2021, 7, 748-756.	4.7	98
9	High-quality reference genome of Fasciola gigantica: Insights into the genomic signatures of transposon-mediated evolution and specific parasitic adaption in tropical regions. PLoS Neglected Tropical Diseases, 2021, 15, e0009750.	1.3	12
10	Screening and identification of female-specific DNA sequences in octaploid sturgeon using comparative genomics with high-throughput sequencing. Genomics, 2021, 113, 4237-4244.	1.3	6
11	Fast and accurate long-read assembly with wtdbg2. Nature Methods, 2020, 17, 155-158.	9.0	917
12	Population sequencing enhances understanding of tea plant evolution. Nature Communications, 2020, 11, 4447.	5.8	123
13	Genome and single-cell RNA-sequencing of the earthworm Eisenia andrei identifies cellular mechanisms underlying regeneration. Nature Communications, 2020, 11, 2656.	5.8	43
14	From asymmetrical to balanced genomic diversification during rediploidization: Subgenomic evolution in allotetraploid fish. Science Advances, 2020, 6, eaaz7677.	4.7	59
15	Comparison of Long Non-Coding RNA Expression Profiles of Cattle and Buffalo Differing in Muscle Characteristics. Frontiers in Genetics, 2020, 11, 98.	1.1	19
16	A genome-wide survey of copy number variations reveals an asymmetric evolution of duplicated genes in rice. BMC Biology, 2020, 18, 73.	1.7	7
17	Understanding divergent domestication traits from the whole-genome sequencing of swamp- and river-buffalo populations. National Science Review, 2020, 7, 686-701.	4.6	43
18	Structural variation during dog domestication: insights from gray wolf and dhole genomes. National Science Review, 2019, 6, 110-122.	4.6	30

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19	Accurate circular consensus long-read sequencing improves variant detection and assembly of a human genome. Nature Biotechnology, 2019, 37, 1155-1162.	9.4	1,010
20	Sequencing of Chinese castor lines reveals genetic signatures of selection and yield-associated loci. Nature Communications, 2019, 10, 3418.	5.8	26
21	A chromosome-scale genome assembly of cucumber (Cucumis sativus L.). GigaScience, 2019, 8, .	3.3	138
22	LRScaf: improving draft genomes using long noisy reads. BMC Genomics, 2019, 20, 955.	1.2	53
23	Genomic variation in 3,010 diverse accessions of Asian cultivated rice. Nature, 2018, 557, 43-49.	13.7	1,091
24	Novel sequences, structural variations and gene presence variations of Asian cultivated rice. Scientific Data, 2018, 5, 180079.	2.4	14
25	Genome analysis of Taraxacum kok-saghyz Rodin provides new insights into rubber biosynthesis. National Science Review, 2018, 5, 78-87.	4.6	90
26	pBACode: a random-barcode-based high-throughput approach for BAC paired-end sequencing and physical clone mapping. Nucleic Acids Research, 2017, 45, gkw1261.	6.5	9
27	novoBreak: local assembly for breakpoint detection in cancer genomes. Nature Methods, 2017, 14, 65-67.	9.0	93
28	Genomic analysis of snub-nosed monkeys (Rhinopithecus) identifies genes and processes related to high-altitude adaptation. Nature Genetics, 2016, 48, 947-952.	9.4	109
29	DBC2OLC: Efficient Assembly of Large Genomes Using Long Erroneous Reads of the Third Generation Sequencing Technologies. Scientific Reports, 2016, 6, 31900.	1.6	268
30	The Evolution of Small Insertions and Deletions in the Coding Genes of Drosophila melanogaster. Molecular Biology and Evolution, 2013, 30, 2699-2708.	3.5	6
31	Reference-Free Comparative Genomics of 174 Chloroplasts. PLoS ONE, 2012, 7, e48995.	1.1	4
32	Rapid growth of a hepatocellular carcinoma and the driving mutations revealed by cell-population genetic analysis of whole-genome data. Proceedings of the National Academy of Sciences of the United States of America, 2011, 108, 12042-12047.	3.3	117
33	The sequence and de novo assembly of the giant panda genome. Nature, 2010, 463, 311-317.	13.7	1,058
34	De novo assembly of human genomes with massively parallel short read sequencing. Genome Research, 2010, 20, 265-272.	2.4	2,516
35	The genome of the cucumber, Cucumis sativus L. Nature Genetics, 2009, 41, 1275-1281.	9.4	1,317
36	The Sequence Alignment/Map format and SAMtools. Bioinformatics, 2009, 25, 2078-2079.	1.8	49,124

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37	The diploid genome sequence of an Asian individual. Nature, 2008, 456, 60-65.	13.7	834
38	Mapping short DNA sequencing reads and calling variants using mapping quality scores. Genome Research, 2008, 18, 1851-1858.	2.4	2,275
39	A genetic variation map for chicken with 2.8 million single-nucleotide polymorphisms. Nature, 2004, 432, 717-722.	13.7	391
40	SMARTdenovo: a de novo assembler using long noisy reads. GigaByte, 0, 2021, 1-9.	0.0	109