Zhengqing Ouyang

List of Publications by Year in descending order

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44 papers 3,772 citations

394421 19 h-index 345221 36 g-index

44 all docs

44 docs citations

44 times ranked

8046 citing authors

#	Article	IF	CITATIONS
1	Differential RNA methylation using multivariate statistical methods. Briefings in Bioinformatics, 2022, 23, .	6.5	3
2	Direct DNA crosslinking with CAP-C uncovers transcription-dependent chromatin organization at high resolution. Nature Biotechnology, 2021, 39, 225-235.	17.5	37
3	Targeting eIF4A-Dependent Translation of KRAS Signaling Molecules. Cancer Research, 2021, 81, 2002-2014.	0.9	17
4	Multimodal single-cell analysis of cutaneous T-cell lymphoma reveals distinct subclonal tissue-dependent signatures. Blood, 2021, 138, 1456-1464.	1.4	39
5	LISA2: Learning Complex Single-Cell Trajectory and Expression Trends. Frontiers in Genetics, 2021, 12, 681206.	2.3	1
6	A new graph-based clustering method with application to single-cell RNA-seq data from human pancreatic islets. NAR Genomics and Bioinformatics, 2021, 3, Iqaa087.	3.2	2
7	Low SATB1 Expression Promotes IL-5 and IL-9 Expression in Sézary Syndrome. Journal of Investigative Dermatology, 2020, 140, 713-716.	0.7	5
8	Identification and analysis of RNA structural disruptions induced by single nucleotide variants using Riprap and RiboSNitchDB. NAR Genomics and Bioinformatics, 2020, 2, Iqaa057.	3.2	10
9	Largeâ€scale analysis of the positionâ€dependent binding and regulation of human RNA binding proteins. Quantitative Biology, 2020, 8, 119-129.	0.5	O
10	Gene selection for optimal prediction of cell position in tissues from single-cell transcriptomics data. Life Science Alliance, 2020, 3, e202000867.	2.8	20
11	Meta-analysis of peptides to detect protein significance. Statistics and Its Interface, 2020, 13, 465-474.	0.3	O
12	PATH: An interactive web platform for analysis of time-course high-dimensional genomic data. International Journal of Computational Biology and Drug Design, 2020, 13, 529.	0.3	0
13	PATH: An interactive web platform for analysis of time-course high-dimensional genomic data. International Journal of Computational Biology and Drug Design, 2020, 13, 529.	0.3	0
14	Single-cell transcriptomics–based MacSpectrum reveals macrophage activation signatures in diseases. JCI Insight, 2019, 4, .	5.0	86
15	PRAS: Predicting functional targets of RNA binding proteins based on CLIP-seq peaks. PLoS Computational Biology, 2019, 15, e1007227.	3.2	8
16	c-MYC regulates mRNA translation efficiency and start-site selection in lymphoma. Journal of Experimental Medicine, 2019, 216, 1509-1524.	8.5	32
17	Multiplexed detection of proteins, transcriptomes, clonotypes and CRISPR perturbations in single cells. Nature Methods, 2019, 16, 409-412.	19.0	364
18	Dissecting Pathway Disturbances Using Network Topology and Multi-platform Genomics Data. Statistics in Biosciences, 2018, 10, 86-106.	1.2	9

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19	Joint Principal Trend Analysis for Longitudinal High-dimensional Data. Biometrics, 2018, 74, 430-438.	1.4	4
20	DNA methylomes of bovine gametes and in vivo produced preimplantation embryos. Biology of Reproduction, 2018, 99, 949-959.	2.7	59
21	LISA: Accurate reconstruction of cell trajectory and pseudo-time for massive single cell RNA-seq data. , 2018, , .		2
22	A statistical framework for data integration through graphical models with application to cancer genomics. Annals of Applied Statistics, 2017, 11, 161-184.	1.1	19
23	HSA: integrating multi-track Hi-C data for genome-scale reconstruction of 3D chromatin structure. Genome Biology, 2016, 17, 40.	8.8	60
24	Genome-Wide Probing of RNA Structures In Vitro Using Nucleases and Deep Sequencing. Methods in Molecular Biology, 2016, 1361, 141-160.	0.9	4
25	Joint modeling of RNase footprint sequencing profiles for genome-wide inference of RNA structure. Nucleic Acids Research, 2015, 43, 9187-9197.	14.5	7
26	Landscape and variation of RNA secondary structure across the human transcriptome. Nature, 2014, 505, 706-709.	27.8	519
27	SeqFold: Genome-scale reconstruction of RNA secondary structure integrating high-throughput sequencing data. Genome Research, 2013, 23, 377-387.	5.5	102
28	Genome-wide mapping of RNA structure using nuclease digestion and high-throughput sequencing. Nature Protocols, 2013, 8, 849-869.	12.0	85
29	STAT3 Targets Suggest Mechanisms of Aggressive Tumorigenesis in Diffuse Large B-Cell Lymphoma. G3: Genes, Genomes, Genetics, 2013, 3, 2173-2185.	1.8	34
30	Correlation of Global MicroRNA Expression With Basal Cell Carcinoma Subtype. G3: Genes, Genomes, Genetics, 2012, 2, 279-286.	1.8	36
31	Genome-wide Measurement of RNA Folding Energies. Molecular Cell, 2012, 48, 169-181.	9.7	192
32	Architecture of the human regulatory network derived from ENCODE data. Nature, 2012, 489, 91-100.	27.8	1,384
33	A New FACS Approach Isolates hESC Derived Endoderm Using Transcription Factors. PLoS ONE, 2011, 6, e17536.	2.5	23
34	Hedgehog pathway-regulated gene networks in cerebellum development and tumorigenesis. Proceedings of the National Academy of Sciences of the United States of America, 2010, 107, 9736-9741.	7.1	109
35	Noncoding RNA Landmarks of Pluripotency and Reprogramming. Cell Stem Cell, 2010, 7, 649-650.	11.1	2
36	ChIP-Seq of transcription factors predicts absolute and differential gene expression in embryonic stem cells. Proceedings of the National Academy of Sciences of the United States of America, 2009, 106, 21521-21526.	7.1	312

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37	Assessing the Conservation of Mammalian Gene Expression Using High-Density Exon Arrays. Molecular Biology and Evolution, 2007, 24, 1283-1285.	8.9	49
38	Exon arrays provide accurate assessments of gene expression. Genome Biology, 2007, 8, R82.	9.6	103
39	Hierarchical structure analysis describing abnormal base composition of genomes. Physical Review E, 2005, 72, 041915.	2.1	2
40	Scaling and Hierarchical Structures in DNA Sequences. Physical Review Letters, 2004, 93, 078103.	7.8	15
41	MULTIVARIATE ENTROPY DISTANCE METHOD FOR PROKARYOTIC GENE IDENTIFICATION. Journal of Bioinformatics and Computational Biology, 2004, 02, 353-373.	0.8	17
42	Unsupervised gene selection for predicting cell spatial positions in the Drosophila embryo. F1000Research, 0, 9, 124.	1.6	0
43	Structural inference of timeâ€varying mixed graphical models. Stat, 0, , e414.	0.4	O
44	An unsupervised learning method for reconstructing cell spatial organization withÂapplication to the DREAM Single Cell Transcriptomics Challenge. F1000Research, 0, 9, 124.	1.6	0