Michael Q Zhang

List of Publications by Year in descending order

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71102 24258 110 17,120 115 41 citations h-index g-index papers 151 151 151 32814 docs citations times ranked citing authors all docs

#	Article	IF	CITATIONS
1	Integrative analysis of 111 reference human epigenomes. Nature, 2015, 518, 317-330.	27.8	5,653
2	Combinatorial patterns of histone acetylations and methylations in the human genome. Nature Genetics, 2008, 40, 897-903.	21.4	2,034
3	Analysis of the Vertebrate Insulator Protein CTCF-Binding Sites in the Human Genome. Cell, 2007, 128, 1231-1245.	28.9	910
4	CRISPR Inversion of CTCF Sites Alters Genome Topology and Enhancer/Promoter Function. Cell, 2015, 162, 900-910.	28.9	846
5	Networkâ€based global inference of human disease genes. Molecular Systems Biology, 2008, 4, 189.	7.2	583
6	NONCODE 2016: an informative and valuable data source of long non-coding RNAs. Nucleic Acids Research, 2016, 44, D203-D208.	14.5	574
7	NONCODEV5: a comprehensive annotation database for long non-coding RNAs. Nucleic Acids Research, 2018, 46, D308-D314.	14.5	434
8	Allelic reprogramming of 3D chromatin architecture during early mammalian development. Nature, 2017, 547, 232-235.	27.8	406
9	An integrative genomics approach identifies Hypoxia Inducible Factor-1 (HIF-1)-target genes that form the core response to hypoxia. Nucleic Acids Research, 2009, 37, 4587-4602.	14.5	400
10	HITS-CLIP and Integrative Modeling Define the Rbfox Splicing-Regulatory Network Linked to Brain Development and Autism. Cell Reports, 2014, 6, 1139-1152.	6.4	326
11	Recurrently deregulated lncRNAs in hepatocellular carcinoma. Nature Communications, 2017, 8, 14421.	12.8	279
12	Genome-wide map of regulatory interactions in the human genome. Genome Research, 2014, 24, 1905-1917.	5 . 5	259
13	In Situ Capture of Chromatin Interactions by Biotinylated dCas9. Cell, 2017, 170, 1028-1043.e19.	28.9	236
14	Pervasive Chromatin-RNA Binding Protein Interactions Enable RNA-Based Regulation of Transcription. Cell, 2019, 178, 107-121.e18.	28.9	224
15	Integrative analysis of haplotype-resolved epigenomes across human tissues. Nature, 2015, 518, 350-354.	27.8	201
16	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.	4.1	152
17	Co-inhibitory Molecule B7 Superfamily Member 1 Expressed by Tumor-Infiltrating Myeloid Cells Induces Dysfunction of Anti-tumor CD8+ T Cells. Immunity, 2018, 48, 773-786.e5.	14.3	150
18	Histone Deacetylases Positively Regulate Transcription through the Elongation Machinery. Cell Reports, 2015, 13, 1444-1455.	6.4	138

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19	Fast dimension reduction and integrative clustering of multi-omics data using low-rank approximation: application to cancer molecular classification. BMC Genomics, 2015, 16, 1022.	2.8	124
20	Reconstructing cell cycle pseudo time-series via single-cell transcriptome data. Nature Communications, 2017, 8, 22.	12.8	121
21	Model-guided quantitative analysis of microRNA-mediated regulation on competing endogenous RNAs using a synthetic gene circuit. Proceedings of the National Academy of Sciences of the United States of America, 2015, 112, 3158-3163.	7.1	117
22	Activity-dependent FUS dysregulation disrupts synaptic homeostasis. Proceedings of the National Academy of Sciences of the United States of America, 2014, 111, E4769-78.	7.1	116
23	DNA motifs in human and mouse proximal promoters predict tissue-specific expression. Proceedings of the National Academy of Sciences of the United States of America, 2006, 103, 6275-6280.	7.1	114
24	Identifying tissue-selective transcription factor binding sites in vertebrate promoters. Proceedings of the National Academy of Sciences of the United States of America, 2005, 102, 1560-1565.	7.1	113
25	ChIA-PET2: a versatile and flexible pipeline for ChIA-PET data analysis. Nucleic Acids Research, 2017, 45, e4-e4.	14.5	104
26	Epigenome-Wide Association of Liver Methylation Patterns and Complex Metabolic Traits in Mice. Cell Metabolism, 2015, 21, 905-917.	16.2	98
27	Similarity of position frequency matrices for transcription factor binding sites. Bioinformatics, 2005, 21, 307-313.	4.1	97
28	PAF1 regulation of promoter-proximal pause release via enhancer activation. Science, 2017, 357, 1294-1298.	12.6	95
29	Super-resolution dipole orientation mapping via polarization demodulation. Light: Science and Applications, 2016, 5, e16166-e16166.	16.6	93
30	Chd5 orchestrates chromatin remodelling during sperm development. Nature Communications, 2014, 5, 3812.	12.8	82
31	SEAM is a spatial single nuclear metabolomics method for dissecting tissue microenvironment. Nature Methods, 2021, 18, 1223-1232.	19.0	78
32	FIND: difFerential chromatin INteractions Detection using a spatial Poisson process. Genome Research, 2018, 28, 412-422.	5 . 5	69
33	Statistical significance of cis-regulatory modules. BMC Bioinformatics, 2007, 8, 19.	2.6	68
34	Design and bioinformatics analysis of genome-wide CLIP experiments. Nucleic Acids Research, 2015, 43, 5263-5274.	14.5	65
35	A common set of distinct features that characterize noncoding RNAs across multiple species. Nucleic Acids Research, 2015, 43, 104-114.	14.5	63
36	BL-Hi-C is an efficient and sensitive approach for capturing structural and regulatory chromatin interactions. Nature Communications, 2017, 8, 1622.	12.8	60

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37	Inhibiting the integrated stress response pathway prevents aberrant chondrocyte differentiation thereby alleviating chondrodysplasia. ELife, 2018, 7, .	6.0	59
38	Synergistic co-regulation and competition by a SOX9-GLI-FOXA phasic transcriptional network coordinate chondrocyte differentiation transitions. PLoS Genetics, 2018, 14, e1007346.	3.5	56
39	Network embedding-based representation learning for single cell RNA-seq data. Nucleic Acids Research, 2017, 45, e166-e166.	14.5	54
40	Tissueâ€specific regulatory elements in mammalian promoters. Molecular Systems Biology, 2007, 3, 73.	7.2	52
41	HiCDB: a sensitive and robust method for detecting contact domain boundaries. Nucleic Acids Research, 2018, 46, 11239-11250.	14.5	52
42	SuperCT: a supervised-learning framework for enhanced characterization of single-cell transcriptomic profiles. Nucleic Acids Research, 2019, 47, e48-e48.	14.5	52
43	MyoD is a 3D genome structure organizer for muscle cell identity. Nature Communications, 2022, 13, 205.	12.8	50
44	Engineering EMT using 3D micro-scaffold to promote hepatic functions for drug hepatotoxicity evaluation. Biomaterials, 2016, 91, 11-22.	11.4	45
45	Integrating Hi-C and FISH data for modeling of the 3D organization of chromosomes. Nature Communications, 2019, 10, 2049.	12.8	45
46	Emergent Lévy behavior in single-cell stochastic gene expression. Physical Review E, 2017, 96, 040402.	2.1	40
47	Distinct and Predictive Histone Lysine Acetylation Patterns at Promoters, Enhancers, and Gene Bodies. G3: Genes, Genomes, Genetics, 2014, 4, 2051-2063.	1.8	39
48	Quantitative combination of natural anti-oxidants prevents metabolic syndrome by reducing oxidative stress. Redox Biology, 2015, 6, 206-217.	9.0	39
49	The landscape of RNA polymerase II–associated chromatin interactions in prostate cancer. Journal of Clinical Investigation, 2020, 130, 3987-4005.	8.2	37
50	The loss of heterochromatin is associated with multiscale three-dimensional genome reorganization and aberrant transcription during cellular senescence. Genome Research, 2021, 31, 1121-1135.	5.5	36
51	MICC: an R package for identifying chromatin interactions from ChIA-PET data. Bioinformatics, 2015, 31, 3832-3834.	4.1	34
52	MTGIpick allows robust identification of genomic islands from a single genome. Briefings in Bioinformatics, 2016, 19, bbw118.	6.5	34
53	Global transcriptional activity dynamics reveal functional enhancer RNAs. Genome Research, 2018, 28, 1799-1811.	5.5	34
54	Differential connectivity of splicing activators and repressors to the human spliceosome. Genome Biology, 2015, 16, 119.	9.6	33

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55	<i>De novo</i> deciphering three-dimensional chromatin interaction and topological domains by wavelet transformation of epigenetic profiles. Nucleic Acids Research, 2016, 44, e106-e106.	14.5	33
56	Single-cell stochastic gene expression kinetics with coupled positive-plus-negative feedback. Physical Review E, 2019, 100, 052406.	2.1	33
57	Sox2 and Klf4 as the Functional Core in Pluripotency Induction without Exogenous Oct4. Cell Reports, 2019, 29, 1986-2000.e8.	6.4	32
58	Alterations of specific chromatin conformation affect ATRA-induced leukemia cell differentiation. Cell Death and Disease, 2018, 9, 200.	6.3	29
59	2SigFinder: the combined use of small-scale and large-scale statistical testing for genomic island detection from a single genome. BMC Bioinformatics, 2020, 21, 159.	2.6	29
60	Super-resolution imaging of a 2.5 kb non-repetitive DNA in situ in the nuclear genome using molecular beacon probes. ELife, $2017, 6, .$	6.0	29
61	Hsa-miR-1246, hsa-miR-320a and hsa-miR-196b-5p inhibitors can reduce the cytotoxicity of Ebola virus glycoprotein in vitro. Science China Life Sciences, 2014, 57, 959-972.	4.9	28
62	Multiplexed capture of spatial configuration and temporal dynamics of locus-specific 3D chromatin by biotinylated dCas9. Genome Biology, 2020, 21, 59.	8.8	27
63	FastDMA: An Infinium HumanMethylation450 Beadchip Analyzer. PLoS ONE, 2013, 8, e74275.	2.5	25
64	Relaxation rates of gene expression kinetics reveal the feedback signs of autoregulatory gene networks. Journal of Chemical Physics, 2018, 148, .	3.0	24
65	Developing novel methods to image and visualize 3D genomes. Cell Biology and Toxicology, 2018, 34, 367-380.	5.3	24
66	3D genome alterations associated with dysregulated HOXA13 expression in high-risk T-lineage acute lymphoblastic leukemia. Nature Communications, 2021, 12, 3708.	12.8	24
67	Quantifying the phase separation property of chromatin-associated proteins under physiological conditions using an anti-1,6-hexanediol index. Genome Biology, 2021, 22, 229.	8.8	24
68	Population dynamics of cancer cells with cell state conversions. Quantitative Biology, 2013, 1, 201-208.	0.5	22
69	Stochastic fluctuations can reveal the feedback signs of gene regulatory networks at the single-molecule level. Scientific Reports, 2017, 7, 16037.	3.3	22
70	CTCF functions as an insulator for somatic genes and a chromatin remodeler for pluripotency genes during reprogramming. Cell Reports, 2022, 39, 110626.	6.4	22
71	Web3DMol: interactive protein structure visualization based on WebGL. Nucleic Acids Research, 2017, 45, W523-W527.	14.5	21
72	Genome-wide analysis of the response to nitric oxide in uropathogenic Escherichia coli CFT073. Microbial Genomics, 2015, 1, e000031.	2.0	21

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73	3CPET: finding co-factor complexes from ChIA-PET data using a hierarchical Dirichlet process. Genome Biology, 2015, 16, 288.	8.8	20
74	ChIP-Array 2: integrating multiple omics data to construct gene regulatory networks. Nucleic Acids Research, 2015, 43, W264-W269.	14.5	19
75	Analysis of <i>C. elegans</i> muscle transcriptome using trans-splicing-based RNA tagging (SRT). Nucleic Acids Research, 2016, 44, gkw734.	14.5	19
76	HPVMD-C: a disease-based mutation database of human papillomavirus in China. Database: the Journal of Biological Databases and Curation, 2022, 2022, .	3.0	19
77	Nucleosome eviction and multiple co-factor binding predict estrogen-receptor-alpha-associated long-range interactions. Nucleic Acids Research, 2014, 42, 6935-6944.	14.5	17
78	Regulatory RNA binding proteins contribute to the transcriptome-wide splicing alterations in human cellular senescence. Aging, 2018, 10, 1489-1505.	3.1	17
79	Dynamic Alternative Splicing During Mouse Preimplantation Embryo Development. Frontiers in Bioengineering and Biotechnology, 2020, 8, 35.	4.1	17
80	Resolving the genetic heterogeneity of prelingual hearing loss within one family: Performance comparison and application of two targeted next generation sequencing approaches. Journal of Human Genetics, 2014, 59, 599-607.	2.3	16
81	FAT10 Is Critical in Influenza A Virus Replication by Inhibiting Type I IFN. Journal of Immunology, 2016, 197, 824-833.	0.8	16
82	HCSGD: An integrated database of human cellular senescence genes. Journal of Genetics and Genomics, 2017, 44, 227-234.	3.9	15
83	Molecular basis for histone H3 "K4me3-K9me3/2―methylation pattern readout by Spindlin1. Journal of Biological Chemistry, 2020, 295, 16877-16887.	3.4	15
84	CAPTURE: <i>In Situ</i> Analysis of Chromatin Composition of Endogenous Genomic Loci by Biotinylated dCas9. Current Protocols in Molecular Biology, 2018, 123, e64.	2.9	14
85	Computational prediction of novel components of lung transcriptional networks. Bioinformatics, 2007, 23, 21-29.	4.1	13
86	Deciphering hierarchical organization of topologically associated domains through change-point testing. BMC Bioinformatics, 2021, 22, 183.	2.6	12
87	Single-cell alternative polyadenylation analysis delineates GABAergic neuron types. BMC Biology, 2021, 19, 144.	3.8	12
88	Integrative molecular analysis of metastatic hepatocellular carcinoma. BMC Medical Genomics, 2019, 12, 164.	1.5	11
89	Characterizing microRNA-mediated modulation of gene expression noise and its effect on synthetic gene circuits. Cell Reports, 2021, 36, 109573.	6.4	11
90	Exome Sequencing Identifies a Novel Frameshift Mutation of <i>MYO6 </i> as the Cause of Autosomal Dominant Nonsyndromic Hearing Loss in a Chinese Family. Annals of Human Genetics, 2014, 78, 410-423.	0.8	10

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91	MarkovHC: Markov hierarchical clustering for the topological structure of high-dimensional single-cell omics data with transition pathway and critical point detection. Nucleic Acids Research, 2022, 50, 46-56.	14.5	9
92	Mammalian non-CG methylations are conserved and cell-type specific and may have been involved in the evolution of transposon elements. Scientific Reports, 2016, 6, 32207.	3.3	8
93	A large-scale CRISPR screen and identification of essential genes in cellular senescence bypass. Aging, 2019, 11, 4011-4031.	3.1	8
94	Tn5-FISH, a novel cytogenetic method to image chromatin interactions with sub-kilobase resolution. Journal of Genetics and Genomics, 2020, 47, 727-734.	3.9	8
95	Assembly and Validation of Versatile Transcription Activator-Like Effector Libraries. Scientific Reports, 2014, 4, 4857.	3.3	7
96	Reconfigurable hybrid interface for molecular marker diagnostics and in-situ reporting. Biosensors and Bioelectronics, 2015, 74, 744-750.	10.1	7
97	ScaffComb: A Phenotypeâ€Based Framework for Drug Combination Virtual Screening in Largeâ€Scale Chemical Datasets. Advanced Science, 2021, 8, e2102092.	11.2	7
98	Small C-terminal Domain Phosphatase 3 Dephosphorylates the Linker Sites of Receptor-regulated Smads (R-Smads) to Ensure Transforming Growth Factor \hat{l}^2 (TGF \hat{l}^2)-mediated Germ Layer Induction in Xenopus Embryos. Journal of Biological Chemistry, 2015, 290, 17239-17249.	3.4	6
99	ModuleRole: A Tool for Modulization, Role Determination and Visualization in Protein-Protein Interaction Networks. PLoS ONE, 2014, 9, e94608.	2.5	5
100	Genome wide mapping of Foxo1 binding-sites in murine T lymphocytes. Genomics Data, 2014, 2, 280-281.	1.3	5
101	Advances in computational ChIAâ€PET data analysis. Quantitative Biology, 2016, 4, 217-225.	0.5	5
102	DAGM: A novel modelling framework to assess the risk of HER2-negative breast cancer based on germline rare coding mutations. EBioMedicine, 2021, 69, 103446.	6.1	4
103	Integration of single cell data by disentangled representation learning. Nucleic Acids Research, 2022, 50, e8-e8.	14.5	4
104	Bivalent-Like Chromatin Markers Are Predictive for Transcription Start Site Distribution in Human. PLoS ONE, 2012, 7, e38112.	2.5	3
105	Highly diversified core promoters in the human genome and their effects on gene expression and disease predisposition. BMC Genomics, 2020, 21, 842.	2.8	3
106	SCT Promoter Methylation Is a Highly Discriminative Biomarker for Lung and Many Other Cancers. IEEE Life Sciences Letters, 2015, 1, 30-33.	1.2	2
107	Single‑cell RNA sequencing of t(8;21) acute myeloid leukemia for risk prediction. Oncology Reports, 2020, 43, 1278-1288.	2.6	2
108	A data-driven method to learn a jump diffusion process from aggregate biological gene expression data. Journal of Theoretical Biology, 2022, 532, 110923.	1.7	2

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109	Role of emergent palliative radiation therapy in the care of patients with cancer Journal of Clinical Oncology, 2016, 34, 201-201.	1.6	2
110	Rathke's cleft cyst with xanthogranulomatous change: A case report and review of the literature. , 2020, 11, 246.		2
111	Model-based analysis of chromatin interactions from dCas9-Based CAPTURE-3C-seq. PLoS ONE, 2020, 15, e0236666.	2.5	1
112	A personal journey on cracking the genomic codes. Quantitative Biology, 2021, 9, 8-22.	0.5	1
113	Super-resolution fluorescence dipole orientation microscopy., 2016,,.		0
114	DE MERVLs are Enriched Around Two-Cell-Specific Genes During Zygotic Genome Activation in Mouse., 2018,,.		0
115	Computational modeling and analysis of the morphogenetic domain signaling networks regulating C. elegans embryogenesis. Computational and Structural Biotechnology Journal, 2022, 20, 3653-3666.	4.1	0