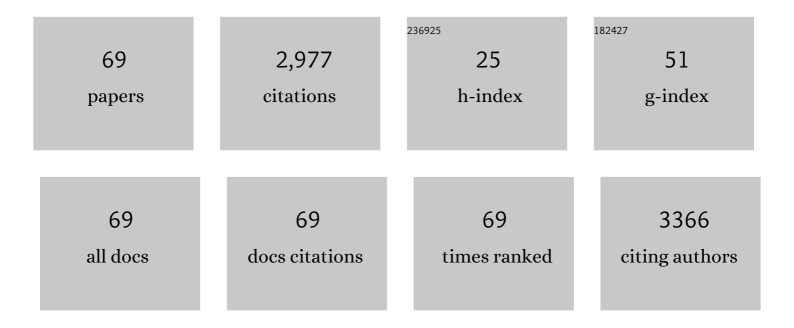
Yunpeng Zhang

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	LncACTdb 3.0: an updated database of experimentally supported ceRNA interactions and personalized networks contributing to precision medicine. Nucleic Acids Research, 2022, 50, D183-D189.	14.5	45
2	Shedding light on the hidden human proteome expands immunopeptidome in cancer. Briefings in Bioinformatics, 2022, 23, .	6.5	4
3	Identifying individualized risk subpathways reveals pan-cancer molecular classification based on multi-omics data. Computational and Structural Biotechnology Journal, 2022, 20, 838-849.	4.1	2
4	Applicability of Anticancer Drugs for the Triple-Negative Breast Cancer Based on Homologous Recombination Repair Deficiency. Frontiers in Cell and Developmental Biology, 2022, 10, 845950.	3.7	4
5	Revealing the contribution of somatic gene mutations to shaping tumor immune microenvironment. Briefings in Bioinformatics, 2022, 23, .	6.5	6
6	GABC : A comprehensive resource and Genome Atlas for Breast Cancer. International Journal of Cancer, 2021, 148, 988-994.	5.1	1
7	Lnc2Cancer 3.0: an updated resource for experimentally supported IncRNA/circRNA cancer associations and web tools based on RNA-seq and scRNA-seq data. Nucleic Acids Research, 2021, 49, D1251-D1258.	14.5	162
8	LnCeCell: a comprehensive database of predicted lncRNA-associated ceRNA networks at single-cell resolution. Nucleic Acids Research, 2021, 49, D125-D133.	14.5	42
9	LincSNP 3.0: an updated database for linking functional variants to human long non-coding RNAs, circular RNAs and their regulatory elements. Nucleic Acids Research, 2021, 49, D1244-D1250.	14.5	28
10	Systematic analysis of enhancer regulatory circuit perturbation driven by copy number variations in malignant glioma. Theranostics, 2021, 11, 3060-3073.	10.0	6
11	Alternative splicing perturbation landscape identifies RNA binding proteins as potential therapeutic targets in cancer. Molecular Therapy - Nucleic Acids, 2021, 24, 792-806.	5.1	15
12	Dissecting immune cell stat regulation network reveals biomarkers to predict ICB therapy responders in melanoma. Journal of Translational Medicine, 2021, 19, 296.	4.4	4
13	Combined homologous recombination repair deficiency and immune activation analysis for predicting intensified responses of anthracycline, cyclophosphamide and taxane chemotherapy in triple-negative breast cancer. BMC Medicine, 2021, 19, 190.	5.5	11
14	Dynamic regulatory networks of T cell trajectory dissect transcriptional control of T cell state transition. Molecular Therapy - Nucleic Acids, 2021, 26, 1115-1129.	5.1	11
15	Single-Cell Transcriptomic Analysis Reveals a Tumor-Reactive T Cell Signature Associated With Clinical Outcome and Immunotherapy Response In Melanoma. Frontiers in Immunology, 2021, 12, 758288.	4.8	13
16	Identifying and characterizing lincRNA genomic clusters reveals its cooperative functions in human cancer. Journal of Translational Medicine, 2021, 19, 509.	4.4	5
17	LnCeVar: a comprehensive database of genomic variations that disturb ceRNA network regulation. Nucleic Acids Research, 2020, 48, D111-D117.	14.5	59
18	LncTarD: a manually-curated database of experimentally-supported functional lncRNA–target regulations in human diseases. Nucleic Acids Research, 2020, 48, D118-D126.	14.5	77

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19	A comprehensive overview of oncogenic pathways in human cancer. Briefings in Bioinformatics, 2020, 21, 957-969.	6.5	29
20	Identifying functions and prognostic biomarkers of network motifs marked by diverse chromatin states in human cell lines. Oncogene, 2020, 39, 677-689.	5.9	11
21	RNAactDrug: a comprehensive database of RNAs associated with drug sensitivity from multi-omics data. Briefings in Bioinformatics, 2020, 21, 2167-2174.	6.5	32
22	Identification and comprehensive characterization of IncRNAs with copy number variations and their driving transcriptional perturbed subpathways reveal functional significance for cancer. Briefings in Bioinformatics, 2020, 21, 2153-2166.	6.5	12
23	Pan-cancer characterization of immune-related lncRNAs identifies potential oncogenic biomarkers. Nature Communications, 2020, 11, 1000.	12.8	293
24	Identifying subpathway signatures for individualized anticancer drug response by integrating multi-omics data. Journal of Translational Medicine, 2019, 17, 255.	4.4	11
25	Prioritization of candidate cancer drugs based on a drug functional similarity network constructed by integrating pathway activities and drug activities. Molecular Oncology, 2019, 13, 2259-2277.	4.6	27
26	Comprehensive Characterization of Somatic Mutations Impacting IncRNA Expression for Pan-Cancer. Molecular Therapy - Nucleic Acids, 2019, 18, 66-79.	5.1	27
27	Molecular characterization and clinical relevance of m6A regulators across 33 cancer types. Molecular Cancer, 2019, 18, 137.	19.2	286
28	Identification of Cancer Dysfunctional Subpathways by Integrating DNA Methylation, Copy Number Variation, and Gene-Expression Data. Frontiers in Genetics, 2019, 10, 441.	2.3	27
29	Gain-of-Function Mutations: An Emerging Advantage for Cancer Biology. Trends in Biochemical Sciences, 2019, 44, 659-674.	7.5	38
30	Systematic identification of lincRNAâ€based prognostic biomarkers by integrating lincRNA expression and copy number variation in lung adenocarcinoma. International Journal of Cancer, 2019, 144, 1723-1734.	5.1	85
31	LncACTdb 2.0: an updated database of experimentally supported ceRNA interactions curated from low- and high-throughput experiments. Nucleic Acids Research, 2019, 47, D121-D127.	14.5	97
32	MetSigDis: a manually curated resource for the metabolic signatures of diseases. Briefings in Bioinformatics, 2019, 20, 203-209.	6.5	106
33	System level characterization of small molecule drugs and their affected long noncoding RNAs. Aging, 2019, 11, 12428-12451.	3.1	8
34	LncMAP: Pan-cancer atlas of long noncoding RNA-mediated transcriptional network perturbations. Nucleic Acids Research, 2018, 46, 1113-1123.	14.5	115
35	MSDD: a manually curated database of experimentally supported associations among miRNAs, SNPs and human diseases. Nucleic Acids Research, 2018, 46, D181-D185.	14.5	42
36	DiseaseEnhancer: a resource of human disease-associated enhancer catalog. Nucleic Acids Research, 2018, 46, D78-D84.	14.5	69

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37	SEECancer: a resource for somatic events in evolution of cancer genome. Nucleic Acids Research, 2018, 46, D1018-D1026.	14.5	9
38	Global view of a drug-sensitivity gene network. Oncotarget, 2018, 9, 3254-3266.	1.8	3
39	Genome-Wide Mapping of SNPs in Non-coding RNAs. Advances in Experimental Medicine and Biology, 2018, 1094, 39-48.	1.6	0
40	Computational Inferring of Risk Subpathways Mediated by Dysfunctional Non-coding RNAs. Advances in Experimental Medicine and Biology, 2018, 1094, 87-95.	1.6	0
41	Inference of patientâ€specific subpathway activities reveals a functional signature associated with the prognosis of patients with breast cancer. Journal of Cellular and Molecular Medicine, 2018, 22, 4304-4316.	3.6	28
42	FACER: comprehensive molecular and functional characterization of epigenetic chromatin regulators. Nucleic Acids Research, 2018, 46, 10019-10033.	14.5	66
43	LncRNAs2Pathways: Identifying the pathways influenced by a set of IncRNAs of interest based on a global network propagation method. Scientific Reports, 2017, 7, 46566.	3.3	24
44	Dynamic Organization of IncRNA and Circular RNA Regulators Collectively Controlled Cardiac Differentiation in Humans. EBioMedicine, 2017, 24, 137-146.	6.1	73
45	RNA Function Prediction. Methods in Molecular Biology, 2017, 1654, 17-28.	0.9	17
46	The LncRNA Connectivity Map: Using LncRNA Signatures to Connect Small Molecules, LncRNAs, and Diseases. Scientific Reports, 2017, 7, 6655.	3.3	21
47	BioM2MetDisease: a manually curated database for associations between microRNAs, metabolites, small molecules and metabolic diseases. Database: the Journal of Biological Databases and Curation, 2017, 2017, .	3.0	5
48	LncSubpathway: a novel approach for identifying dysfunctional subpathways associated with risk IncRNAs by integrating IncRNA and mRNA expression profiles and pathway topologies. Oncotarget, 2017, 8, 15453-15469.	1.8	9
49	Integrating gene and IncRNA expression to infer subpathway activity for tumor analyses. Oncotarget, 2017, 8, 111433-111443.	1.8	2
50	Comprehensive characterization of IncRNA-mRNA related ceRNA network across 12 major cancers. Oncotarget, 2016, 7, 64148-64167.	1.8	171
51	Subpathway-LNCE: Identify dysfunctional subpathways competitively regulated by IncRNAs through integrating IncRNA-mRNA expression profile and pathway topologies. Oncotarget, 2016, 7, 69857-69870.	1.8	18
52	Identification of a IncRNA involved functional module for esophageal cancer subtypes. Molecular BioSystems, 2016, 12, 3312-3323.	2.9	10
53	MiRSEA: Discovering the pathways regulated by dysfunctional MicroRNAs. Oncotarget, 2016, 7, 55012-55025.	1.8	10
54	Dissecting dysfunctional crosstalk pathways regulated by miRNAs during glioma progression. Oncotarget, 2016, 7, 25769-25782.	1.8	7

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#	Article	IF	CITATIONS
55	ESEA: Discovering the Dysregulated Pathways based on Edge Set Enrichment Analysis. Scientific Reports, 2015, 5, 13044.	3.3	31
56	Global Prioritization of Disease Candidate Metabolites Based on a Multi-omics Composite Network. Scientific Reports, 2015, 5, 17201.	3.3	43
57	Genome-wide DNA methylome reveals the dysfunction of intronic microRNAs in major psychosis. BMC Medical Genomics, 2015, 8, 62.	1.5	27
58	miRSponge: a manually curated database for experimentally supported miRNA sponges and ceRNAs. Database: the Journal of Biological Databases and Curation, 2015, 2015, bav098.	3.0	112
59	Identification of Subtype Specific miRNA-mRNA Functional Regulatory Modules in Matched miRNA-mRNA Expression Data: Multiple Myeloma as a Case. BioMed Research International, 2015, 2015, 1-15.	1.9	5
60	Genome-wide DNA methylome analysis reveals epigenetically dysregulated non-coding RNAs in human breast cancer. Scientific Reports, 2015, 5, 8790.	3.3	54
61	Identification of IncRNA-associated competing triplets reveals global patterns and prognostic markers for cancer. Nucleic Acids Research, 2015, 43, 3478-3489.	14.5	219
62	Subpathway-GMir: identifying miRNA-mediated metabolic subpathways by integrating condition-specific genes, microRNAs, and pathway topologies. Oncotarget, 2015, 6, 39151-39164.	1.8	33
63	The DNA Methylome and Transcriptome of Different Brain Regions in Schizophrenia and Bipolar Disorder. PLoS ONE, 2014, 9, e95875.	2.5	75
64	MPINet: Metabolite Pathway Identification via Coupling of Global Metabolite Network Structure and Metabolomic Profile. BioMed Research International, 2014, 2014, 1-14.	1.9	12
65	The detection of risk pathways, regulated by miRNAs, via the integration of sample-matched miRNA-mRNA profiles and pathway structure. Journal of Biomedical Informatics, 2014, 49, 187-197.	4.3	9
66	Identification of miRNA-Mediated Core Gene Module for Glioma Patient Prediction by Integrating High-Throughput miRNA, mRNA Expression and Pathway Structure. PLoS ONE, 2014, 9, e96908.	2.5	26
67	Prioritizing Candidate Disease Metabolites Based on Global Functional Relationships between Metabolites in the Context of Metabolic Pathways. PLoS ONE, 2014, 9, e104934.	2.5	23
68	Network Analysis Reveals Functional Cross-links between Disease and Inflammation Genes. Scientific Reports, 2013, 3, 3426.	3.3	22
69	Systematic Identification of Core Transcription Factors Mediating Dysregulated Links Bridging Inflammatory Bowel Diseases and Colorectal Cancer. PLoS ONE, 2013, 8, e83495.	2.5	3