Wenyi Wang

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The third column is the impact factor (IF) of the journal, and the fourth column is the number of citations of the article.

61 6,197 24 75 g-index

75 q-index

75 ext. papers ext. citations avg, IF

61 75 avg, IF

12.6 L-index

#	Paper	IF	Citations
61	Comprehensive molecular characterization of urothelial bladder carcinoma. <i>Nature</i> , 2014 , 507, 315-22	50.4	1963
60	The Molecular Taxonomy of Primary Prostate Cancer. <i>Cell</i> , 2015 , 163, 1011-25	56.2	1713
59	The somatic genomic landscape of chromophobe renal cell carcinoma. <i>Cancer Cell</i> , 2014 , 26, 319-330	24.3	521
58	The evolutionary history of 2,658 cancers. <i>Nature</i> , 2020 , 578, 122-128	50.4	307
57	Prediction of germline mutations and cancer risk in the Lynch syndrome. <i>JAMA - Journal of the American Medical Association</i> , 2006 , 296, 1479-87	27.4	271
56	PancPRO: risk assessment for individuals with a family history of pancreatic cancer. <i>Journal of Clinical Oncology</i> , 2007 , 25, 1417-22	2.2	150
55	MuSE: accounting for tumor heterogeneity using a sample-specific error model improves sensitivity and specificity in mutation calling from sequencing data. <i>Genome Biology</i> , 2016 , 17, 178	18.3	120
54	Inference of combinatorial regulation in yeast transcriptional networks: a case study of sporulation. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , 2005 , 102, 1998-2003	11.5	87
53	Immuno-genomic landscape of osteosarcoma. <i>Nature Communications</i> , 2020 , 11, 1008	17.4	77
52	DeMix: deconvolution for mixed cancer transcriptomes using raw measured data. <i>Bioinformatics</i> , 2013 , 29, 1865-71	7.2	72
51	A systematic approach to reconstructing transcription networks in Saccharomycescerevisiae. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , 2002 , 99, 16893-8	11.5	71
50	An ensemble approach to accurately detect somatic mutations using SomaticSeq. <i>Genome Biology</i> , 2015 , 16, 197	18.3	64
49	Characterizing genetic intra-tumor heterogeneity across 2,658 human cancer genomes. <i>Cell</i> , 2021 , 184, 2239-2254.e39	56.2	57
48	BayesMendel: an R environment for Mendelian risk prediction. <i>Statistical Applications in Genetics and Molecular Biology</i> , 2004 , 3, Article21	1.2	56
47	Siah1 interacts with the scaffold protein POSH to promote JNK activation and apoptosis. <i>Journal of Biological Chemistry</i> , 2006 , 281, 303-12	5.4	53
46	Mammalian Sug1 and c-Fos in the nuclear 26S proteasome. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , 1996 , 93, 8236-40	11.5	53
45	High-quality DNA sequence capture of 524 disease candidate genes. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , 2011 , 108, 6549-54	11.5	47

(2014-2013)

44	Rare variant detection using family-based sequencing analysis. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , 2013 , 110, 3985-90	11.5	42
43	Neutral tumor evolution?. <i>Nature Genetics</i> , 2018 , 50, 1630-1633	36.3	38
42	Transcriptome Deconvolution of Heterogeneous Tumor Samples with Immune Infiltration. <i>IScience</i> , 2018 , 9, 451-460	6.1	36
41	Estimating CDKN2A carrier probability and personalizing cancer risk assessments in hereditary melanoma using MelaPRO. <i>Cancer Research</i> , 2010 , 70, 552-9	10.1	31
40	Identification of rare DNA variants in mitochondrial disorders with improved array-based sequencing. <i>Nucleic Acids Research</i> , 2011 , 39, 44-58	20.1	26
39	Accurate RNA Sequencing From Formalin-Fixed Cancer Tissue To Represent High-Quality Transcriptome From Frozen Tissue. <i>JCO Precision Oncology</i> , 2018 , 2018,	3.6	26
38	Characterizing genetic intra-tumor heterogeneity across 2,658 human cancer genomes		25
37	Bayesian variable selection for binary outcomes in high-dimensional genomic studies using non-local priors. <i>Bioinformatics</i> , 2016 , 32, 1338-45	7.2	21
36	Next-Generation Molecular Testing of Newborn Dried Blood Spots for Cystic Fibrosis. <i>Journal of Molecular Diagnostics</i> , 2016 , 18, 267-82	5.1	21
35	RNA-seq mixology: designing realistic control experiments to compare protocols and analysis methods. <i>Nucleic Acids Research</i> , 2017 , 45, e30	20.1	19
34	Identification of germline DICER1 mutations and loss of heterozygosity in familial Wilms tumour. <i>Journal of Medical Genetics</i> , 2016 , 53, 385-8	5.8	19
33	Slow down to stay alive: HER4 protects against cellular stress and confers chemoresistance in neuroblastoma. <i>Cancer</i> , 2012 , 118, 5140-54	6.4	18
32	Evaluation of school-based smoking-cessation interventions for self-described adolescent smokers. <i>Pediatrics</i> , 2009 , 124, e187-94	7.4	18
31	Cardelino: computational integration of somatic clonal substructure and single-cell transcriptomes. <i>Nature Methods</i> , 2020 , 17, 414-421	21.6	17
30	Estimating genome-wide copy number using allele-specific mixture models. <i>Journal of Computational Biology</i> , 2008 , 15, 857-66	1.7	17
29	Multiplex target capture with double-stranded DNA probes. <i>Genome Medicine</i> , 2013 , 5, 50	14.4	15
28	Cardelino: Integrating whole exomes and single-cell transcriptomes to reveal phenotypic impact of somatic variants		12
27	FamSeq: a variant calling program for family-based sequencing data using graphics processing units. <i>PLoS Computational Biology</i> , 2014 , 10, e1003880	5	10

26	Penetrance of Different Cancer Types in Families with Li-Fraumeni Syndrome: A Validation Study Using Multicenter Cohorts. <i>Cancer Research</i> , 2020 , 80, 354-360	10.1	9
25	Single-cell ATAC and RNA sequencing reveal pre-existing and persistent cells associated with prostate cancer relapse. <i>Nature Communications</i> , 2021 , 12, 5307	17.4	9
24	BAYESIAN VARIABLE SELECTION FOR SURVIVAL DATA USING INVERSE MOMENT PRIORS. <i>Annals of Applied Statistics</i> , 2020 , 14, 809-828	2.1	8
23	Estimating Mutation Carrier Probability in Families with Li-Fraumeni Syndrome Using LFSPRO. Cancer Epidemiology Biomarkers and Prevention, 2017, 26, 837-844	4	7
22	Integration of transcriptional and mutational data simplifies the stratification of peripheral T-cell lymphoma. <i>American Journal of Hematology</i> , 2019 , 94, 628-634	7.1	7
21	A DNA resequencing array for genes involved in Parkinson's disease. <i>Parkinsonism and Related Disorders</i> , 2012 , 18, 386-90	3.6	7
20	Systematic Assessment of Tumor Purity and Its Clinical Implications. <i>JCO Precision Oncology</i> , 2020 , 4,	3.6	7
19	A molecular inversion probe assay for detecting alternative splicing. <i>BMC Genomics</i> , 2010 , 11, 712	4.5	6
18	Abstract 3000: Pervasive intra-tumour heterogeneity and subclonal selection across cancer types 2018 ,		6
17	HepatoScore-14: Measures of Biological Heterogeneity Significantly Improve Prediction of Hepatocellular Carcinoma Risk. <i>Hepatology</i> , 2021 , 73, 2278-2292	11.2	5
16	Integrating multi-platform genomic data using hierarchical Bayesian relevance vector machines. <i>Eurasip Journal on Bioinformatics and Systems Biology</i> , 2013 , 2013, 9		4
15	A recessive Mendelian model to predict carrier probabilities of DFNB1 for nonsyndromic deafness. <i>Human Mutation</i> , 2006 , 27, 1135-42	4.7	4
14	Penetrance Estimates Over Time to First and Second Primary Cancer Diagnosis in Families with Li-Fraumeni Syndrome: A Single Institution Perspective. <i>Cancer Research</i> , 2020 , 80, 347-353	10.1	3
13	Bayesian Semiparametric Estimation of Cancer-specific Age-at-onset Penetrance with Application to Li-Fraumeni Syndrome. <i>Journal of the American Statistical Association</i> , 2019 , 114, 541-552	2.8	3
12	Bayesian latent-class mixed-effect hybrid models for dyadic longitudinal data with non-ignorable dropouts. <i>Biometrics</i> , 2013 , 69, 914-24	1.8	2
11	Tumor cell total mRNA expression shapes the molecular and clinical phenotype of cancer		2
10	Transcriptome Deconvolution of Heterogeneous Tumor Samples with Immune Infiltration		2
9	Bayesian analysis of longitudinal dyadic data with informative missing data using a dyadic shared-parameter model. <i>Statistical Methods in Medical Research</i> , 2019 , 28, 70-83	2.3	2

LIST OF PUBLICATIONS

8	SRMA: an R package for resequencing array data analysis. <i>Bioinformatics</i> , 2012 , 28, 1928-30	7.2	1
7	Bayesian Edge Regression in Undirected Graphical Models to Characterize Interpatient Heterogeneity in Cancer. <i>Journal of the American Statistical Association</i> ,1-31	2.8	1
6	Estimating Genome-Wide Copy Number Using Allele Specific Mixture Models. <i>Lecture Notes in Computer Science</i> , 2007 , 137-150	0.9	1
5	A pedigree-based prediction model identifies carriers of deleterious de novo mutations in families with Li-Fraumeni syndrome. <i>Genome Research</i> , 2020 , 30, 1170-1180	9.7	1
4	Bayesian estimation of a semiparametric recurrent event model with applications to the penetrance estimation of multiple primary cancers in Li-Fraumeni syndrome. <i>Biostatistics</i> , 2020 , 21, 467	-482	1
3	Single-cell ATAC and RNA sequencing reveal pre-existing and persistent subpopulations of cells associated with relapse of prostate cancer		1
2	DNA Variant Calling in Targeted Sequencing Data54-76		
1	MuSE: A Novel Approach to Mutation Calling with Sample-Specific Error Modeling. <i>Methods in Molecular Biology</i> , 2022 , 21-27	1.4	