## Zohar Yakhini

List of Publications by Year in descending order

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53794 22166 15,736 123 45 113 citations h-index g-index papers 141 141 141 23949 docs citations times ranked citing authors all docs

#	Article	IF	CITATIONS
1	Electroporation-based proteome sampling ex vivo enables the detection of brain melanoma protein signatures in a location proximate to visible tumor margins. PLoS ONE, 2022, 17, e0265866.	2.5	5
2	SOLQC: Synthetic Oligo Library Quality Control tool. Bioinformatics, 2021, 37, 720-722.	4.1	14
3	miRNA normalization enables joint analysis of several datasets to increase sensitivity and to reveal novel miRNAs differentially expressed in breast cancer. PLoS Computational Biology, 2021, 17, e1008608.	3.2	1
4	Overcoming the design, build, test bottleneck for synthesis of nonrepetitive protein-RNA cassettes. Nature Communications, 2021, 12, 1576.	12.8	8
5	CRISPECTOR provides accurate estimation of genome editing translocation and off-target activity from comparative NGS data. Nature Communications, 2021, 12, 3042.	12.8	23
6	On the stability of log-rank test under labeling errors. Bioinformatics, 2021, 37, 4451-4459.	4.1	0
7	Assessing heterogeneity in spatial data using the HTA index with applications to spatial transcriptomics and imaging. Bioinformatics, 2021, 37, 3796-3804.	4.1	6
8	A broad analysis of splicing regulation in yeast using a large library of synthetic introns. PLoS Genetics, 2021, 17, e1009805.	3.5	16
9	Efficient gene expression signature for a breast cancer immuno-subtype. PLoS ONE, 2021, 16, e0245215.	2.5	2
10	Sequence Reconstruction Under Stutter Noise in Enzymatic DNA Synthesis., 2021,,.		4
11	Spatial transcriptomics inferred from pathology whole-slide images links tumor heterogeneity to survival in breast and lung cancer. Scientific Reports, 2020, 10, 18802.	3.3	78
12	Increasing CRISPR Efficiency and Measuring Its Specificity in HSPCs Using a Clinically Relevant System. Molecular Therapy - Methods and Clinical Development, 2020, 17, 1097-1107.	4.1	46
13	IoT or NoT: Identifying IoT Devices in a Short Time Scale. , 2020, , .		18
14	Distributed flux balance analysis simulations of serial biomass fermentation by two organisms. PLoS ONE, 2020, 15, e0227363.	2.5	10
15	Molecular harvesting with electroporation for tissue profiling. Scientific Reports, 2019, 9, 15750.	3.3	9
16	Data storage in DNA with fewer synthesis cycles using composite DNA letters. Nature Biotechnology, 2019, 37, 1229-1236.	17.5	110
17	The Functional 3D Organization of Unicellular Genomes. Scientific Reports, 2019, 9, 12734.	3.3	O
18	Predicting Methylation from Sequence and Gene Expression Using Deep Learning with Attention. Lecture Notes in Computer Science, 2019, , 179-190.	1.3	4

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19	Synthetic 5′ UTRs Can Either Up- or Downregulate Expression upon RNA-Binding Protein Binding. Cell Systems, 2019, 9, 93-106.e8.	6.2	11
20	Reciprocal Reprogramming of Cancer Cells and Associated Mesenchymal Stem Cells in Gastric Cancer. Stem Cells, 2019, 37, 176-189.	3.2	18
21	Sparse NIR optimization method (SNIRO) to quantify analyte composition with visible (VIS)/near infrared (NIR) spectroscopy (350â€nm-2500â€nm). Analytica Chimica Acta, 2019, 1051, 32-40.	5.4	4
22	Design and Analysis of Offshore Macroalgae Biorefineries. Methods in Molecular Biology, 2018, 1980, 9-33.	0.9	2
23	Macroalgae Biorefinery from Kappaphycus alvarezii: Conversion Modeling and Performance Prediction for India and Philippines as Examples. Bioenergy Research, 2018, 11, 22-32.	3.9	42
24	An <i>in Vivo</i> Binding Assay for RNA-Binding Proteins Based on Repression of a Reporter Gene. ACS Synthetic Biology, 2018, 7, 2765-2774.	3.8	16
25	Genome-wide analysis of fitness data and its application to improve metabolic models. BMC Bioinformatics, 2018, 19, 368.	2.6	3
26	SMARTIV: combined sequence and structure de-novo motif discovery for in-vivo RNA binding data. Nucleic Acids Research, 2018, 46, W221-W228.	14.5	7
27	Systematic Investigation of Transcription Factor Activity in the Context of Chromatin Using Massively Parallel Binding and Expression Assays. Molecular Cell, 2017, 65, 604-617.e6.	9.7	48
28	A combined sequence and structure based method for discovering enriched motifs in RNA from in vivo binding data. Methods, 2017, 118-119, 73-81.	3.8	13
29	A Synthetic Oligo Library and Sequencing Approach Reveals an Insulation Mechanism Encoded within Bacterial $\parallel f \parallel $	6.4	23
30	Net primary productivity, biofuel production and CO 2 emissions reduction potential of Ulva sp. (Chlorophyta) biomass in a coastal area of the Eastern Mediterranean. Energy Conversion and Management, 2017, 148, 1497-1507.	9.2	40
31	Prediction of Scar Size in Rats Six Months after Burns Based on Early Post-injury Polarization-Sensitive Optical Frequency Domain Imaging. Frontiers in Physiology, 2017, 8, 967.	2.8	6
32	Mutual enrichment in aggregated ranked lists with applications to gene expression regulation. Bioinformatics, 2016, 33, btw727.	4.1	0
33	<scp>LIMT</scp> is a novel metastasis inhibiting lnc <scp>RNA</scp> suppressed by <scp>EGF</scp> and downregulated in aggressive breast cancer. EMBO Molecular Medicine, 2016, 8, 1052-1064.	6.9	77
34	Thermochemical hydrolysis of macroalgae Ulva for biorefinery: Taguchi robust design method. Scientific Reports, 2016, 6, 27761.	3.3	22
35	Serum Nâ€glycan analysis in breast cancer patients – Relation to tumour biology and clinical outcome. Molecular Oncology, 2016, 10, 59-72.	4.6	34
36	Extending partial haplotypes to full genome haplotypes using chromosome conformation capture data. Bioinformatics, 2016, 32, i559-i566.	4.1	17

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37	Mutual enrichment in aggregated ranked lists with applications to gene expression regulation. Bioinformatics, 2016, 32, i464-i472.	4.1	5
38	Systematic discovery of cap-independent translation sequences in human and viral genomes. Science, 2016, 351, .	12.6	258
39	BioLEGO â€" a web-based application for biorefinery design and evaluation of serial biomass fermentation. Technology, 2015, 03, 89-98.	1.4	10
40	An Efficient Minimum Free Energy Structure-Based Search Method for Riboswitch Identification Based on Inverse RNA Folding. PLoS ONE, 2015, 10, e0134262.	2.5	11
41	Systematic Dissection of the Sequence Determinants of Gene 3' End Mediated Expression Control. PLoS Genetics, 2015, 11, e1005147.	3.5	70
42	Unraveling determinants of transcription factor binding outside the core binding site. Genome Research, 2015, 25, 1018-1029.	5.5	146
43	ENViz: a Cytoscape App for integrated statistical analysis and visualization of sample-matched data with multiple data types. Bioinformatics, 2015, 31, 1683-1685.	4.1	12
44	RNAPattMatch: a web server for RNA sequence/structure motif detection based on pattern matching with flexible gaps. Nucleic Acids Research, 2015, 43, W507-W512.	14.5	11
45	Optimizing analytical depth and cost efficiency of IEF-LC/MS proteomics. , 2014, , .		0
46	Proposed design of distributed macroalgal biorefineries: thermodynamics, bioconversion technology, and sustainability implications for developing economies. Biofuels, Bioproducts and Biorefining, 2014, 8, 67-82.	3.7	49
47	Molecular Rules Governing <i>De Novo</i> Methylation in Cancer. Cancer Research, 2014, 74, 1475-1483.	0.9	55
48	Deregulation of cancer-related miRNAs is a common event in both benign and malignant human breast tumors. Carcinogenesis, 2014, 35, 76-85.	2.8	119
49	Probing the effect of promoters on noise in gene expression using thousands of designed sequences. Genome Research, 2014, 24, 1698-1706.	5.5	118
50	Peer group normalization and urine to blood context in steroid metabolomics: The case of CAH and obesity. Steroids, 2014, 88, 83-89.	1.8	15
51	Mutual enrichment in ranked lists and the statistical assessment of position weight matrix motifs. Algorithms for Molecular Biology, 2014, 9, 11.	1.2	6
52	Association of N-Glycosylation with Breast Carcinoma and Systemic Features Using High-Resolution Quantitative UPLC. Journal of Proteome Research, 2014, 13, 2314-2327.	3.7	123
53	Spatial localization of co-regulated genes exceeds genomic gene clustering in the Saccharomyces cerevisiae genome. Nucleic Acids Research, 2013, 41, 2191-2201.	14.5	50
54	miRNA target enrichment analysis reveals directly active miRNAs in health and disease. Nucleic Acids Research, 2013, 41, e45-e45.	14.5	51

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55	DRIMust: a web server for discovering rank imbalanced motifs using suffix trees. Nucleic Acids Research, 2013, 41, W174-W179.	14.5	53
56	Mutual Enrichment in Ranked Lists and the Statistical Assessment of Position Weight Matrix Motifs. Lecture Notes in Computer Science, 2013, , 273-286.	1.3	1
57	Identifying In-Trans Process Associated Genes in Breast Cancer by Integrated Analysis of Copy Number and Expression Data. PLoS ONE, 2013, 8, e53014.	2.5	54
58	Efficient motif search in ranked lists and applications to variable gap motifs. Nucleic Acids Research, 2012, 40, 5832-5847.	14.5	22
59	Dotted interval graphs. ACM Transactions on Algorithms, 2012, 8, 1-21.	1.0	2
60	Inferring gene regulatory logic from high-throughput measurements of thousands of systematically designed promoters. Nature Biotechnology, 2012, 30, 521-530.	17.5	439
61	Global Methylation Patterns in Idiopathic Pulmonary Fibrosis. PLoS ONE, 2012, 7, e33770.	2.5	169
62	Systematic Determination of Replication Activity Type Highlights Interconnections between Replication, Chromatin Structure and Nuclear Localization. PLoS ONE, 2012, 7, e48986.	2.5	13
63	Ischemia caused by time to freezing induces systematic microRNA and mRNA responses in cancer tissue. Molecular Oncology, 2011, 5, 564-576.	4.6	29
64	Divergent RNA binding specificity of yeast Puf2p. Rna, 2011, 17, 1479-1488.	3.5	25
65	miRNA-mRNA Integrated Analysis Reveals Roles for miRNAs in Primary Breast Tumors. PLoS ONE, 2011, 6, e16915.	2.5	278
66	A structural-based statistical approach suggests a cooperative activity of PUM1 and miR-410 in human 3'-untranslated regions. Silence: A Journal of RNA Regulation, 2010, 1, 17.	8.1	20
67	Comparative Analysis of DNA Replication Timing Reveals Conserved Large-Scale Chromosomal Architecture. PLoS Genetics, 2010, 6, e1001011.	3.5	158
68	EGF Decreases the Abundance of MicroRNAs That Restrain Oncogenic Transcription Factors. Science Signaling, 2010, 3, ra43.	3.6	100
69	A Novel Translocation Breakpoint within the BPTF Gene Is Associated with a Pre-Malignant Phenotype. PLoS ONE, 2010, 5, e9657.	2.5	53
70	Novel Rank-Based Statistical Methods Reveal MicroRNAs with Differential Expression in Multiple Cancer Types. PLoS ONE, 2009, 4, e8003.	2.5	150
71	It Takes Two to Tango: Genotyping and Phenotyping in Genome-Wide Association Studies. Biological Theory, 2009, 4, 294-301.	1.5	1
72	GOrilla: a tool for discovery and visualization of enriched GO terms in ranked gene lists. BMC Bioinformatics, 2009, 10, 48.	2.6	3,032

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73	Developmental programming of CpG island methylation profiles in the human genome. Nature Structural and Molecular Biology, 2009, 16, 564-571.	8.2	345
74	The Fine-Scale and Complex Architecture of Human Copy-Number Variation. American Journal of Human Genetics, 2008, 82, 685-695.	6.2	315
75	Global organization of replication time zones of the mouse genome. Genome Research, 2008, 18, 1562-1570.	5.5	148
76	Clinically driven semi-supervised class discovery in gene expression data. Bioinformatics, 2008, 24, i90-i97.	4.1	15
77	<i>GSTP1</i> Promoter Haplotypes Affect DNA Methylation Levels and Promoter Activity in Breast Carcinomas. Cancer Research, 2008, 68, 5562-5571.	0.9	44
78	Small Deletion Variants Have Stable Breakpoints Commonly Associated with Alu Elements. PLoS ONE, 2008, 3, e3104.	2.5	52
79	Framework for Identifying Common Aberrations in DNA Copy Number Data. , 2007, , 122-136.		8
80	Array CGH analysis of copy number variation identifies 1284 new genes variant in healthy white males: implications for association studies of complex diseases. Human Molecular Genetics, 2007, 16, 2783-2794.	2.9	200
81	Discovering Motifs in Ranked Lists of DNA Sequences. PLoS Computational Biology, 2007, 3, e39.	3.2	633
82	Similarities and differences of gene expression in yeast stress conditions. Bioinformatics, 2007, 23, e184-e190.	4.1	24
83	A supervised approach for identifying discriminating genotype patterns and its application to breast cancer data. Bioinformatics, 2007, 23, e91-e98.	4.1	20
84	Optimization of probe coverage for high-resolution oligonucleotide aCGH. Bioinformatics, 2007, 23, e77-e83.	4.1	13
85	Gene expression and the concept of the phenotype. Studies in History and Philosophy of Science Part C:Studies in History and Philosophy of Biological and Biomedical Sciences, 2007, 38, 238-254.	1.3	32
86	Polycomb-mediated methylation on Lys27 of histone H3 pre-marks genes for de novo methylation in cancer. Nature Genetics, 2007, 39, 232-236.	21.4	1,062
87	Semi-supervised class discovery using quantitative phenotypes – CVD as a case study. BMC Bioinformatics, 2007, 8, S6.	2.6	5
88	Efficient Calculation of Interval Scores for DNA Copy Number Data Analysis. Journal of Computational Biology, 2006, 13, 215-228.	1.6	132
89	Molecular Signatures Determining Coronary Artery and Saphenous Vein Smooth Muscle Cell Phenotypes. Arteriosclerosis, Thrombosis, and Vascular Biology, 2006, 26, 1058-1065.	2.4	61
90	A High-Throughput Approach for Associating MicroRNAs with Their Activity Conditions. Journal of Computational Biology, 2006, 13, 245-266.	1.6	11

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91	Differences in Vascular Bed Disease Susceptibility Reflect Differences in Gene Expression Response to Atherogenic Stimuli. Circulation Research, 2006, 98, 200-208.	4.5	71
92	ANALYSIS OF SNP-EXPRESSION ASSOCIATION MATRICES. Journal of Bioinformatics and Computational Biology, 2006, 04, 259-274.	0.8	4
93	Genetic variation in putative regulatory loci controlling gene expression in breast cancer. Proceedings of the National Academy of Sciences of the United States of America, 2006, 103, 7735-7740.	7.1	32
94	Pathway analysis of coronary atherosclerosis. Physiological Genomics, 2005, 23, 103-118.	2.3	144
95	Designing optimally multiplexed SNP genotyping assays. Journal of Computer and System Sciences, 2005, 70, 399-417.	1.2	1
96	Using Expression Data to Discover RNA and DNA Regulatory Sequence Motifs. Lecture Notes in Computer Science, 2005, , 65-78.	1.3	9
97	Multiplexing Schemes for Generic SNP Genotyping Assays. Journal of Computational Biology, 2005, 12, 514-533.	1.6	8
98	Exploratory Visualization of Array-Based Comparative Genomic Hybridization. Information Visualization, 2005, 4, 176-190.	1.9	6
99	Marek's disease virus Meq transforms chicken cells via the v-Jun transcriptional cascade: A converging transforming pathway for avian oncoviruses. Proceedings of the National Academy of Sciences of the United States of America, 2005, 102, 14831-14836.	7.1	92
100	Analysis of SNP-expression association matrices. , 2005, , 135-43.		2
101	Finding Approximate Tandem Repeats in Genomic Sequences. Journal of Computational Biology, 2005, 12, 928-942.	1.6	82
102	Joint Analysis of DNA Copy Numbers and Gene Expression Levels. Lecture Notes in Computer Science, 2004, , 135-146.	1.3	20
103	Finding approximate tandem repeats in genomic sequences. , 2004, , .		19
104	Towards Optimally Multiplexed Applications of Universal Arrays. Journal of Computational Biology, 2004, 11, 476-492.	1.6	4
105	Comparative genomic hybridization using oligonucleotide microarrays and total genomic DNA. Proceedings of the National Academy of Sciences of the United States of America, 2004, 101, 17765-17770.	7.1	336
106	Discovering Local Structure in Gene Expression Data: The Order-Preserving Submatrix Problem. Journal of Computational Biology, 2003, 10, 373-384.	1.6	391
107	Novel Role for the Potent Endogenous Inotrope Apelin in Human Cardiac Dysfunction. Circulation, 2003, 108, 1432-1439.	1.6	311
108	Molecular classification of familial non- <i>BRCA1/BRCA2</i> breast cancer. Proceedings of the National Academy of Sciences of the United States of America, 2003, 100, 2532-2537.	7.1	182

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109	Identification of endothelial cell genes by combined database mining and microarray analysis. Physiological Genomics, 2003, 13, 249-262.	2.3	107
110	Designing Optimally Multiplexed SNP Genotyping Assays. Lecture Notes in Computer Science, 2003, , 320-338.	1.3	5
111	Towards optimally multiplexed applications of universal DNA tag systems. , 2003, , .		7
112	Gene expression analysis reveals matrilysin as a key regulator of pulmonary fibrosis in mice and humans. Proceedings of the National Academy of Sciences of the United States of America, 2002, 99, 6292-6297.	7.1	576
113	Transcriptional Profiling of Non-small Cell Lung Cancer Using Oligonucleotide Microarrays. Chest, 2002, 121, 44S.	0.8	3
114	METHODS FOR ANALYSIS AND VISUALIZATION OF SNP GENOTYPE DATA FOR COMPLEX DISEASES. , 2002, , 548-61.		9
115	Use of oligonucleotide microarrays to analyze gene expression patterns in pulmonary fibrosis reveals distinct patterns of gene expression in mice and humans. Chest, 2002, 121, 31S-32S.	0.8	5
116	Gene-Expression Profiles in Hereditary Breast Cancer. New England Journal of Medicine, 2001, 344, 539-548.	27.0	1,669
117	Analysis of Expression Patterns: The Scope of the Problem, the Problem of Scope. Disease Markers, 2001, 17, 59-65.	1.3	8
118	Class discovery in gene expression data. , 2001, , .		73
119	Universal DNA tag systems. , 2000, , .		13
120	Tissue Classification with Gene Expression Profiles. Journal of Computational Biology, 2000, 7, 559-583.	1.6	623
121	Universal DNA Tag Systems: A Combinatorial Design Scheme. Journal of Computational Biology, 2000, 7, 503-519.	1.6	52
122	Clustering gene expression patterns. , 1999, , .		61
123	Clustering Gene Expression Patterns. Journal of Computational Biology, 1999, 6, 281-297.	1.6	959