## Bo Liao

## List of Publications by Year in descending order

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257357 377752 1,401 63 24 34 citations h-index g-index papers 63 1363 63 63 citing authors all docs docs citations times ranked

#	Article	IF	Citations
1	Local-View-Assisted Discriminative Band Selection With Hypergraph Autolearning for Hyperspectral Image Classification. IEEE Transactions on Geoscience and Remote Sensing, 2020, 58, 2042-2055.	2.7	15
2	Scalable One-Pass Self-Representation Learning for Hyperspectral Band Selection. IEEE Transactions on Geoscience and Remote Sensing, 2019, 57, 4360-4374.	2.7	32
3	Prediction of Protein Subcellular Localization Based on Fusion of Multi-view Features. Molecules, 2019, 24, 919.	1.7	21
4	Improved Pre-miRNAs Identification Through Mutual Information of Pre-miRNA Sequences and Structures. Frontiers in Genetics, 2019, 10, 119.	1.1	28
5	Global Similarity Method Based on a Two-tier Random Walk for the Prediction of microRNA–Disease Association. Scientific Reports, 2018, 8, 6481.	1.6	31
6	The Advances and Challenges of Deep Learning Application in Biological Big Data Processing. Current Bioinformatics, 2018, 13, 352-359.	0.7	91
7	New 3D graphical representation for RNA structure analysis and its application in the pre-miRNA identification of plants. RSC Advances, 2018, 8, 30833-30841.	1.7	6
8	A novel information diffusion method based on network consistency for identifying disease related microRNAs. RSC Advances, 2018, 8, 36675-36690.	1.7	14
9	Predicting Influenza Antigenicity by Matrix Completion With Antigen and Antiserum Similarity. Frontiers in Microbiology, 2018, 9, 2500.	1.5	16
10	Computational Tools for Investigating Pathogen, Pathogen-Host Interaction, and Infectious Disease. Canadian Journal of Infectious Diseases and Medical Microbiology, 2018, 2018, 1-2.	0.7	0
11	Gene function prediction based on combining gene ontology hierarchy with multi-instance multi-label learning. RSC Advances, 2018, 8, 28503-28509.	1.7	5
12	BPLLDA: Predicting IncRNA-Disease Associations Based on Simple Paths With Limited Lengths in a Heterogeneous Network. Frontiers in Genetics, 2018, 9, 411.	1.1	52
13	Fisher Discrimination Regularized Robust Coding Based on a Local Center for Tumor Classification. Scientific Reports, 2018, 8, 9152.	1.6	2
14	Semi-Supervised Maximum Discriminative Local Margin for Gene Selection. Scientific Reports, 2018, 8, 8619.	1.6	10
15	Matrix-Based Margin-Maximization Band Selection With Data-Driven Diversity for Hyperspectral Image Classification. IEEE Transactions on Geoscience and Remote Sensing, 2018, 56, 7294-7309.	2.7	9
16	Predicting Drug-Target Interactions with Neighbor Interaction Information and Discriminative Low-rank Representation. Current Protein and Peptide Science, 2018, 19, 455-467.	0.7	3
17	New multilocus linkage disequilibrium measure for tag SNP selection. Journal of Bioinformatics and Computational Biology, 2017, 15, 1750001.	0.3	6
18	Maxdenominator Reweighted Sparse Representation for Tumor Classification. Scientific Reports, 2017, 7, 46030.	1.6	9

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19	Predicting influenza antigenicity from Hemagglutintin sequence data based on a joint random forest method. Scientific Reports, 2017, 7, 1545.	1.6	40
20	Matrix completion with side information and its applications in predicting the antigenicity of influenza viruses. Bioinformatics, 2017, 33, 3195-3201.	1.8	33
21	Subcellular localization prediction of apoptosis proteins based on evolutionary information and support vector machine. Artificial Intelligence in Medicine, 2017, 78, 41-46.	3.8	29
22	Global network random walk for predicting potential human lncRNA-disease associations. Scientific Reports, 2017, 7, 12442.	1.6	89
23	Improving gene regulatory network structure using redundancy reduction in the MRNET algorithm. RSC Advances, 2017, 7, 23222-23233.	1.7	11
24	A novel information fusion strategy based on a regularized framework for identifying disease-related microRNAs. RSC Advances, 2017, 7, 44447-44455.	1.7	10
25	Network-based collaborative filtering recommendation model for inferring novel disease-related miRNAs. RSC Advances, 2017, 7, 44961-44971.	1.7	10
26	Screening drug-target interactions with positive-unlabeled learning. Scientific Reports, 2017, 7, 8087.	1.6	35
27	Improved low-rank matrix recovery method for predicting miRNA-disease association. Scientific Reports, 2017, 7, 6007.	1.6	21
28	Seeksv: an accurate tool for somatic structural variation and virus integration detection. Bioinformatics, 2017, 33, 184-191.	1.8	64
29	An Improved Binary Differential Evolution Algorithm to Infer Tumor Phylogenetic Trees. BioMed Research International, 2017, 2017, 1-13.	0.9	4
30	Gene Regulatory Network Inferences Using a Maximum-Relevance and Maximum-Significance Strategy. PLoS ONE, 2016, 11, e0166115.	1.1	7
31	Network Consistency Projection for Human miRNA-Disease Associations Inference. Scientific Reports, 2016, 6, 36054.	1.6	89
32	Uncover miRNA-Disease Association by Exploiting Global Network Similarity. PLoS ONE, 2016, 11, e0166509.	1.1	16
33	A Hierarchical Clustering Method of Selecting Kernel SNP to Unify Informative SNP and Tag SNP. IEEE/ACM Transactions on Computational Biology and Bioinformatics, 2015, 12, 113-122.	1.9	10
34	A new technique for generating pathogenic barcodes in breast cancer susceptibility analysis. Journal of Theoretical Biology, 2015, 366, 84-90.	0.8	5
35	A co-expression modules based gene selection for cancer recognition. Journal of Theoretical Biology, 2014, 362, 75-82.	0.8	13
36	Gene Selection Using Locality Sensitive Laplacian Score. IEEE/ACM Transactions on Computational Biology and Bioinformatics, 2014, 11, 1146-1156.	1.9	47

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37	Using Multi-Instance Hierarchical Clustering Learning System to Predict Yeast Gene Function. PLoS ONE, 2014, 9, e90962.	1.1	6
38	Learning a Weighted Meta-Sample Based Parameter Free Sparse Representation Classification for Microarray Data. PLoS ONE, 2014, 9, e104314.	1.1	13
39	A new graphical coding of DNA sequence and its similarity calculation. Physica A: Statistical Mechanics and Its Applications, 2013, 392, 4663-4667.	1.2	30
40	Subcellular localization prediction of apoptosis proteins based on the data mining for amino acid index database. , 2013, , .		1
41	Informative SNPs Selection Based on Two-Locus and Multilocus Linkage Disequilibrium: Criteria of Max-Correlation and Min-Redundancy. IEEE/ACM Transactions on Computational Biology and Bioinformatics, 2013, 10, 688-695.	1.9	12
42	Accelerating processing speed in pathway research based on GPU., 2013,,.		0
43	Incorporating Secondary Structural Features into Sequence Information for Predicting Protein Structural Class. Protein and Peptide Letters, 2013, 20, 1079-1087.	0.4	2
44	Incorporating Secondary Features into the General form of Chou's PseAAC for Predicting Protein Structural Class. Protein and Peptide Letters, 2012, 19, 1133-1138.	0.4	36
45	A Novel Method to Select Informative SNPs and Their Application in Genetic Association Studies. IEEE/ACM Transactions on Computational Biology and Bioinformatics, 2012, 9, 1529-1534.	1.9	9
46	Multiple ant colony algorithm method for selecting tag SNPs. Journal of Biomedical Informatics, 2012, 45, 931-937.	2.5	14
47	A novel feature selection method based on CFS in cancer recognition. , 2012, , .		20
48	Predicting Apoptosis Protein Subcellular Location with PseAAC by Incorporating Tripeptide Composition. Protein and Peptide Letters, 2011, 18, 1086-1092.	0.4	30
49	RNA secondary structure alignment based on an extended binary coding method. International Journal of Quantum Chemistry, 2011, 111, 978-982.	1.0	1
50	A novel graphical representation of protein sequences and its application. Journal of Computational Chemistry, 2011, 32, 2539-2544.	1.5	25
51	A Novel method for similarity analysis and protein sub-cellular localization prediction. Bioinformatics, 2010, 26, 2678-2683.	1.8	43
52	A binary coding method of RNA secondary structure and its application. Journal of Computational Chemistry, 2009, 30, 2205-2212.	1.5	7
53	A 2D graphical representation of DNA sequence based on dual nucleotides and its application. International Journal of Quantum Chemistry, 2009, 109, 948-958.	1.0	26
54	A group of 3D graphical representation of DNA sequences based on dual nucleotides. International Journal of Quantum Chemistry, 2008, 108, 1485-1490.	1.0	34

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55	Novel method for analyzing proteome. International Journal of Quantum Chemistry, 2007, 107, 1295-1300.	1.0	8
56	On the Similarity of DNA Primary Sequences Based on 5-D Representation. Journal of Mathematical Chemistry, 2007, 42, 47-57.	0.7	39
57	On a four-dimensional representation of RNA secondary structures. Journal of Mathematical Chemistry, 2007, 42, 1015-1022.	0.7	10
58	RNA secondary structure 2D graphical representation without degeneracy. International Journal of Quantum Chemistry, 2006, 106, 1749-1755.	1.0	21
59	A graphical method to construct a phylogenetic tree. International Journal of Quantum Chemistry, 2006, 106, 1998-2005.	1.0	10
60	Coronavirus phylogeny based on triplets of nucleic acids bases. Chemical Physics Letters, 2006, 421, 313-318.	1.2	24
61	RNA Secondary Structure 3D Graphical Representation Without Degeneracy. Journal of Mathematical Chemistry, 2006, 39, 629-636.	0.7	10
62	Coronavirus phylogeny based on 2D graphical representation of DNA sequence. Journal of Computational Chemistry, 2006, 27, 1196-1202.	1.5	45
63	Graphical approach to analyzing DNA sequences. Journal of Computational Chemistry, 2005, 26, 1519-1523.	1.5	42