Dong Xu

List of Publications by Year in descending order

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		117453	102304
157	5,419	34	66
papers	citations	h-index	g-index
159	159	159	7652
all docs	docs citations	times ranked	citing authors

#	Article	IF	CITATIONS
1	Assessing deep learning methods in <i>cis</i> regulatory motif finding based on genomic sequencing data. Briefings in Bioinformatics, 2022, 23, .	3.2	9
2	RNALocate v2.0: an updated resource for RNA subcellular localization with increased coverage and annotation. Nucleic Acids Research, 2022, 50, D333-D339.	6.5	54
3	Coxiella burnetii Virulent Phase I and Avirulent Phase II Variants Differentially Manipulate Autophagy Pathway in Neutrophils. Infection and Immunity, 2022, 90, IAI0053421.	1.0	4
4	Deep learning shapes single-cell data analysis. Nature Reviews Molecular Cell Biology, 2022, 23, 303-304.	16.1	40
5	Large-Scale Integrative Analysis of Soybean Transcriptome Using an Unsupervised Autoencoder Model. Frontiers in Plant Science, 2022, 13, 831204.	1.7	1
6	Neural relational inference to learn long-range allosteric interactions in proteins from molecular dynamics simulations. Nature Communications, 2022, 13, 1661.	5.8	38
7	A Multi-Level Iterative Bi-Clustering Method for Discovering miRNA Co-regulation Network of Abiotic Stress Tolerance in Soybeans. Frontiers in Plant Science, 2022, 13, 860791.	1.7	2
8	High-Resolution Translatome Analysis Reveals Cortical Cell Programs During Early Soybean Nodulation. Frontiers in Plant Science, 2022, 13, 820348.	1.7	4
9	Discovering trends and hotspots of biosafety and biosecurity research via machine learning. Briefings in Bioinformatics, 2022, 23, .	3.2	1
10	Al-Aided Design of Novel Targeted Covalent Inhibitors against SARS-CoV-2. Biomolecules, 2022, 12, 746.	1.0	43
		1.8	_
11	Zinc is a master-regulator of sperm function associated with binding, motility, and metabolic modulation during porcine sperm capacitation. Communications Biology, 2022, 5, .	2.0	10
11	Zinc is a master-regulator of sperm function associated with binding, motility, and metabolic modulation during porcine sperm capacitation. Communications Biology, 2022, 5, . Diagnosis of thyroid neoplasm using support vector machine algorithms based on platelet RNA-seq. Endocrine, 2021, 72, 758-783.		
	modulation during porcine sperm capacitation. Communications Biology, 2022, 5, . Diagnosis of thyroid neoplasm using support vector machine algorithms based on platelet RNA-seq.	2.0	10
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12	modulation during porcine sperm capacitation. Communications Biology, 2022, 5, . Diagnosis of thyroid neoplasm using support vector machine algorithms based on platelet RNA-seq. Endocrine, 2021, 72, 758-783. DM3Loc: multi-label mRNA subcellular localization prediction and analysis based on multi-head self-attention mechanism. Nucleic Acids Research, 2021, 49, e46-e46. New genotypes of Helicobacter Pylori VacA d-region identified from global strains. BMC Molecular	2.0 1.1 6.5	10 14 95
12 13 14	Diagnosis of thyroid neoplasm using support vector machine algorithms based on platelet RNA-seq. Endocrine, 2021, 72, 758-783. DM3Loc: multi-label mRNA subcellular localization prediction and analysis based on multi-head self-attention mechanism. Nucleic Acids Research, 2021, 49, e46-e46. New genotypes of Helicobacter Pylori VacA d-region identified from global strains. BMC Molecular and Cell Biology, 2021, 22, 4. Domain-specific Topic Model for Knowledge Discovery in Computational and Data-Intensive Scientific	2.0 1.1 6.5	10 14 95 11
12 13 14	modulation during porcine sperm capacitation. Communications Biology, 2022, 5, . Diagnosis of thyroid neoplasm using support vector machine algorithms based on platelet RNA-seq. Endocrine, 2021, 72, 758-783. DM3Loc: multi-label mRNA subcellular localization prediction and analysis based on multi-head self-attention mechanism. Nucleic Acids Research, 2021, 49, e46-e46. New genotypes of Helicobacter Pylori VacA d-region identified from global strains. BMC Molecular and Cell Biology, 2021, 22, 4. Domain-specific Topic Model for Knowledge Discovery in Computational and Data-Intensive Scientific Communities. IEEE Transactions on Knowledge and Data Engineering, 2021, , 1-1. DeePaN: deep patient graph convolutional network integrating clinico-genomic evidence to stratify	2.0 1.1 6.5 1.0	10 14 95 11 5

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19	G2PDeep: a web-based deep-learning framework for quantitative phenotype prediction and discovery of genomic markers. Nucleic Acids Research, 2021, 49, W228-W236.	6.5	12
20	Variation in Gene Expression between Two Sorghum bicolor Lines Differing in Innate Immunity Response. Plants, 2021, 10, 1536.	1.6	3
21	Editorial: Computational Resources for Understanding Biomacromolecular Covalent Modifications. Frontiers in Cell and Developmental Biology, 2021, 9, 728127.	1.8	0
22	Genetic mosaicism, intrafamilial phenotypic heterogeneity, and molecular defects of a novel missense SLC6A1 mutation associated with epilepsy and ADHD. Experimental Neurology, 2021, 342, 113723.	2.0	19
23	Deep learning analysis and age prediction from shoeprints. Forensic Science International, 2021, 327, 110987.	1.3	16
24	Comparison of different parathyroid autograft project after total parathyroidectomy in patients with secondary hyperparathyroidism. American Journal of Otolaryngology - Head and Neck Medicine and Surgery, 2021, 42, 103085.	0.6	0
25	Computational methods for protein localization prediction. Computational and Structural Biotechnology Journal, 2021, 19, 5834-5844.	1.9	12
26	Overlapping Community Detection Based on Membership Degree Propagation. Entropy, 2021, 23, 15.	1.1	5
27	Evolutionary Dynamics of Indels in SARS-CoV-2 Spike Glycoprotein. Evolutionary Bioinformatics, 2021, 17, 117693432110646.	0.6	5
28	MUfoldQA_G: High-accuracy protein model QA via retraining and transformation. Computational and Structural Biotechnology Journal, 2021, 19, 6282-6290.	1.9	3
29	Digital tongue image analyses for health assessment. Medical Review, 2021, 1, 172-198.	0.3	11
30	Acupuncture and Tuina Knowledge Graph for Ancient Literature of Traditional Chinese Medicine. , 2021, , .		0
31	IMPRes-Pro: A high dimensional multiomics integration method for in silico hypothesis generation. Methods, 2020, 173, 16-23.	1.9	9
32	In-Situ Metabolomic Analysis of <i>Setaria viridis</i> Roots Colonized by Beneficial Endophytic Bacteria. Molecular Plant-Microbe Interactions, 2020, 33, 272-283.	1.4	23
33	A deep dense inception network for protein betaâ€turn prediction. Proteins: Structure, Function and Bioinformatics, 2020, 88, 143-151.	1.5	9
34	Fuzzified Image Enhancement for Deep Learning in Iris Recognition. IEEE Transactions on Fuzzy Systems, 2020, 28, 92-99.	6.5	55
35	How oncogenic mutations activate human MAP kinase 1 (MEK1): a molecular dynamics simulation study. Journal of Biomolecular Structure and Dynamics, 2020, 38, 3942-3958.	2.0	7
36	Inductive inference of gene regulatory network using supervised and semi-supervised graph neural networks. Computational and Structural Biotechnology Journal, 2020, 18, 3335-3343.	1.9	31

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37	Impact of frailty on inpatient outcomes in thyroid cancer surgery: 10-year results from the U.S. national inpatient sample. Journal of Otolaryngology - Head and Neck Surgery, 2020, 49, 51.	0.9	13
38	Proteomic and Bioinformatic Profiling of Transporters in Higher Plant Mitochondria. Biomolecules, 2020, 10, 1190.	1.8	10
39	Single-Cell Techniques and Deep Learning in Predicting Drug Response. Trends in Pharmacological Sciences, 2020, 41, 1050-1065.	4.0	27
40	Endoplasmic reticulum retention and degradation of a mutation in SLC6A1 associated with epilepsy and autism. Molecular Brain, 2020, 13, 76.	1.3	30
41	Detecting Cancer Survival Related Gene Markers Based on Rectified Factor Network. Frontiers in Bioengineering and Biotechnology, 2020, 8, 349.	2.0	2
42	A self-attention based message passing neural network for predicting molecular lipophilicity and aqueous solubility. Journal of Cheminformatics, 2020, 12, 15.	2.8	73
43	Leveraging Systematic Functional Analysis to Benchmark an <i>In Silico</i> Framework Distinguishes Driver from Passenger MEK Mutants in Cancer. Cancer Research, 2020, 80, 4233-4243.	0.4	18
44	Overcoming Wnt–β-catenin dependent anticancer therapy resistance in leukaemia stem cells. Nature Cell Biology, 2020, 22, 689-700.	4.6	89
45	Artificial Intelligence-Driven Structurization of Diagnostic Information in Free-Text Pathology Reports. Journal of Pathology Informatics, 2020, 11, 4.	0.8	7
46	A Fast Projection-Based Algorithm for Clustering Big Data. Interdisciplinary Sciences, Computational Life Sciences, 2019, 11, 360-366.	2.2	6
47	PSICA: a fast and accurate web service for protein model quality analysis. Nucleic Acids Research, 2019, 47, W443-W450.	6.5	12
48	A rectified factor network based biclustering method for detecting cancer-related coding genes and miRNAs, and their interactions. Methods, 2019, 166, 22-30.	1.9	4
49	A missense mutation in SLC6A1 associated with Lennox-Gastaut syndrome impairs GABA transporter 1 protein trafficking and function. Experimental Neurology, 2019, 320, 112973.	2.0	37
50	Identification of new loci for salt tolerance in soybean by high-resolution genome-wide association mapping. BMC Genomics, 2019, 20, 318.	1.2	46
51	Extracting Molecular Entities and Their Interactions from Pathway Figures Based on Deep Learning. , 2019, , .		1
52	Trends in Alzheimer's Disease Research Based upon Machine Learning Analysis of PubMed Abstracts. International Journal of Biological Sciences, 2019, 15, 2065-2074.	2.6	15
53	ezCADD: A Rapid 2D/3D Visualization-Enabled Web Modeling Environment for Democratizing Computer-Aided Drug Design. Journal of Chemical Information and Modeling, 2019, 59, 18-24.	2.5	47
54	Use of a Bioinformaticsâ€Based Toxicity Scoring System to Assess Serotonin Burden and Predict Populationâ€Level Adverse Drug Events from Concomitant Serotonergic Drug Therapy. Pharmacotherapy, 2019, 39, 171-181.	1.2	2

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55	Characterization of Select Wild Soybean Accessions in the USDA Germplasm Collection for Seed Composition and Agronomic Traits. Crop Science, 2019, 59, 233-251.	0.8	29
56	Capsule network for protein post-translational modification site prediction. Bioinformatics, 2019, 35, 2386-2394.	1.8	92
57	Plant-derived phosphocholine facilitates cellular uptake of anti-pulmonary fibrotic HJT-sRNA-m7. Science China Life Sciences, 2019, 62, 309-320.	2.3	43
58	MUFOLDâ€S: New deep inceptionâ€insideâ€inception networks for protein secondary structure prediction. Proteins: Structure, Function and Bioinformatics, 2018, 86, 592-598.	1.5	134
59	G2S: a web-service for annotating genomic variants on 3D protein structures. Bioinformatics, 2018, 34, 1949-1950.	1.8	10
60	Molecular insights into the improved clinical performance of PEGylated interferon therapeutics: a molecular dynamics perspective. RSC Advances, 2018, 8, 2315-2322.	1.7	23
61	Comparative Effectiveness of High-Dose Versus Standard-Dose Influenza Vaccines in Preventing Postinfluenza Deaths. Journal of Infectious Diseases, 2018, 218, 336-337.	1.9	4
62	Mapping and confirmation of loci for salt tolerance in a novel soybean germplasm, Fiskeby III. Theoretical and Applied Genetics, 2018, 131, 513-524.	1.8	42
63	Domain-specific Topic Model for Knowledge Discovery through Conversational Agents in Data Intensive Scientific Communities. , 2018, , .		9
64	Integrating Gene Expression Data and Pathway Knowledge for In Silico Hypothesis Generation with IMPRes. , $2018, , .$		1
65	Large-scale prediction of protein ubiquitination sites using a multimodal deep architecture. BMC Systems Biology, 2018, 12, 109.	3.0	29
66	Improving Protein Gamma-Turn Prediction Using Inception Capsule Networks. Scientific Reports, 2018, 8, 15741.	1.6	21
67	Implications of Offâ€Target Serotoninergic Drug Activity: An Analysis of Serotonin Syndrome Reports Using a Systematic Bioinformatics Approach. Pharmacotherapy, 2018, 38, 888-898.	1.2	12
68	An analysis and evaluation of the WeFold collaborative for protein structure prediction and its pipelines in CASP11 and CASP12. Scientific Reports, 2018, 8, 9939.	1.6	19
69	Why Is a High Temperature Needed by Thermus thermophilus Argonaute During mRNA Silencing: A Theoretical Study. Frontiers in Chemistry, 2018, 6, 223.	1.8	9
70	DeepDom: Predicting protein domain boundary from sequence alone using stacked bidirectional LSTM. , $2018, , .$		8
71	iPhosâ€PseEvo: Identifying Human Phosphorylated Proteins by Incorporating Evolutionary Information into General PseAAC via Grey System Theory. Molecular Informatics, 2017, 36, 1600010.	1.4	94
72	Experimental identification and computational characterization of a novel extracellular metalloproteinase produced by Clostridium sordellii. RSC Advances, 2017, 7, 13928-13938.	1.7	4

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73	Understanding the Phosphorylation Mechanism by Using Quantum Chemical Calculations and Molecular Dynamics Simulations. Journal of Physical Chemistry B, 2017, 121, 3565-3573.	1.2	12
74	The Evolution of Soybean Knowledge Base (SoyKB). Methods in Molecular Biology, 2017, 1533, 149-159.	0.4	25
75	BFDCA: A Comprehensive Tool of Using Bayes Factor for Differential Co-Expression Analysis. Journal of Molecular Biology, 2017, 429, 446-453.	2.0	10
76	Understanding the differences of the ligand binding/unbinding pathways between phosphorylated and non-phosphorylated ARH1 using molecular dynamics simulations. Scientific Reports, 2017, 7, 12439.	1.6	21
77	Assessing and predicting drug-induced anticholinergic risks: an integrated computational approach. Therapeutic Advances in Drug Safety, 2017, 8, 361-370.	1.0	21
78	Tongue Image Analysis and Its Mobile App Development for Health Diagnosis. Advances in Experimental Medicine and Biology, 2017, 1005, 99-121.	0.8	17
79	MSBIS: A Multi-Step Biomedical Informatics Screening Approach for Identifying Medications that Mitigate the Risks of Metoclopramide-Induced Tardive Dyskinesia. EBioMedicine, 2017, 26, 132-137.	2.7	7
80	An aggressive car-following model in the view of driving style. Canadian Journal of Civil Engineering, 2017, 44, 775-782.	0.7	4
81	OMPcontact: An Outer Membrane Protein Inter-Barrel Residue Contact Prediction Method. Journal of Computational Biology, 2017, 24, 217-228.	0.8	6
82	A multimodal deep architecture for large-scale protein ubiquitylation site prediction. , 2017, , .		9
83	Tongue Images Classification Based on Constrained High Dispersal Network. Evidence-based Complementary and Alternative Medicine, 2017, 2017, 1-12.	0.5	19
84	BioJava-ModFinder: identification of protein modifications in 3D structures from the Protein Data Bank. Bioinformatics, 2017, 33, 2047-2049.	1.8	8
85	Protein Loop Modeling Using Deep Generative Adversarial Network. , 2017, , .		16
86	Bioinformatics Analysis of Protein Phosphorylation in Plant Systems Biology Using P3DB. Methods in Molecular Biology, 2017, 1558, 127-138.	0.4	12
87	Computational Analysis of the Ligand Binding Site of the Extracellular ATP Receptor, DORN1. PLoS ONE, 2016, 11, e0161894.	1.1	18
88	Identification and Comparative Analysis of Differential Gene Expression in Soybean Leaf Tissue under Drought and Flooding Stress Revealed by RNA-Seq. Frontiers in Plant Science, 2016, 7, 1044.	1.7	116
89	Classification of tongue images based on doublet and color space dictionary. , 2016, , .		2
90	Genome-wide transcriptome analysis of soybean primary root under varying water-deficit conditions. BMC Genomics, 2016, 17, 57.	1.2	99

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91	Improving degree-based variable ordering heuristics for solving constraint satisfaction problems. Journal of Heuristics, 2016, 22, 125-145.	1.1	7
92	Imbalanced multi-label learning for identifying antimicrobial peptides and their functional types. Bioinformatics, 2016, 32, 3745-3752.	1.8	92
93	Differential protein structural disturbances and suppression of assembly partners produced by nonsense GABRG2 epilepsy mutations: implications for disease phenotypic heterogeneity. Scientific Reports, 2016, 6, 35294.	1.6	12
94	Predicting DNA Methylation State of CpG Dinucleotide Using Genome Topological Features and Deep Networks. Scientific Reports, 2016, 6, 19598.	1.6	75
95	Exploring Human Diseases and Biological Mechanisms by Protein Structure Prediction and Modeling. Advances in Experimental Medicine and Biology, 2016, 939, 39-61.	0.8	4
96	FALCON@home: a high-throughput protein structure prediction server based on remote homologue recognition. Bioinformatics, 2016, 32, 462-464.	1.8	34
97	Selective refinement and selection of near-native models in protein structure prediction. Proteins: Structure, Function and Bioinformatics, 2015, 83, 1823-1835.	1.5	3
98	The human epilepsy mutation GABRG2(Q390X) causes chronic subunit accumulation and neurodegeneration. Nature Neuroscience, 2015, 18, 988-996.	7.1	82
99	Soybean (Glycine max) SWEET gene family: insights through comparative genomics, transcriptome profiling and whole genome re-sequence analysis. BMC Genomics, 2015, 16, 520.	1.2	173
100	Identification of quantitative trait loci underlying resistance to southern root-knot and reniform nematodes in soybean accession PI 567516C. Molecular Breeding, 2015, 35, 131.	1.0	34
101	Soybean transcription factor ORFeome associated with drought resistance: a valuable resource to accelerate research on abiotic stress resistance. BMC Genomics, 2015, 16, 596.	1.2	17
102	Effect of lipo-chitooligosaccharide on early growth of C ₄ grass seedlings. Journal of Experimental Botany, 2015, 66, 5727-5738.	2.4	39
103	Polymorphism of the glucosyltransferase gene (ycjM) in Escherichia coli and its use for tracking human fecal pollution in water. Science of the Total Environment, 2015, 537, 260-267.	3.9	6
104	Comparative analysis of the drought-responsive transcriptome in soybean lines contrasting for canopy wilting. Plant Science, 2015, 240, 65-78.	1.7	53
105	Transcriptional responses of Arabidopsis thaliana to chewing and sucking insect herbivores. Frontiers in Plant Science, 2014, 5, 565.	1.7	61
106	A new Hidden Markov Model for protein Quality Assessment using compatibility between protein sequence and structure. Tsinghua Science and Technology, 2014, 19, 559-567.	4.1	4
107	Diverse and divergent protein post-translational modifications in two growth stages of a natural microbial community. Nature Communications, 2014, 5, 4405.	5.8	51
108	DL-PRO: A novel deep learning method for protein model quality assessment. , 2014, 2014, 2071-2078.		38

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109	Genotypic variation of gene expression during the soybean innate immunity response. Plant Genetic Resources: Characterisation and Utilisation, 2014, 12, S27-S30.	0.4	11
110	Disrupting KATP channels diminishes the estrogen-mediated protection in female mutant mice during ischemia-reperfusion. Clinical Proteomics, 2014, 11, 19.	1.1	19
111	xMDFF: molecular dynamics flexible fitting of low-resolution X-ray structures. Acta Crystallographica Section D: Biological Crystallography, 2014, 70, 2344-2355.	2.5	50
112	Flowerâ€visiting insects and their potential impact on transgene flow in rice. Journal of Applied Ecology, 2014, 51, 1357-1365.	1.9	27
113	Differentiating enteric Escherichia coli from environmental bacteria through the putative glucosyltransferase gene (ycjM). Water Research, 2014, 61, 224-231.	5.3	14
114	BridgeDb app: unifying identifier mapping services for Cytoscape. F1000Research, 2014, 3, 148.	0.8	11
115	Major Soybean Maturity Gene Haplotypes Revealed by SNPViz Analysis of 72 Sequenced Soybean Genomes. PLoS ONE, 2014, 9, e94150.	1.1	89
116	NEW MDS AND CLUSTERING BASED ALGORITHMS FOR PROTEIN MODEL QUALITY ASSESSMENT AND SELECTION. International Journal on Artificial Intelligence Tools, 2013, 22, 1360006.	0.7	3
117	Automated Tongue Feature Extraction for ZHENG Classification in Traditional Chinese Medicine. Evidence-based Complementary and Alternative Medicine, 2012, 2012, 1-14.	0.5	64
118	Protein Databases on the Internet. Current Protocols in Molecular Biology, 2012, 97, Unit 19.4	2.9	6
119	Human encoded miRNAs that regulate the inflenenza virus genome. , 2012, , .		2
120	ZHENG classification in Traditional Chinese Medicine based on modified specular-free tongue images. , 2012, , .		7
121	Protein Databases on the Internet. Current Protocols in Protein Science, 2012, 70, Unit2.6.	2.8	11
122	Features for automated tongue image shape classification. , 2012, , .		8
123	Computational Challenges in Characterization of Bacteria and Bacteria-Host Interactions Based on Genomic Data. Journal of Computer Science and Technology, 2012, 27, 225-239.	0.9	9
124	Routine Microsecond Molecular Dynamics Simulations with AMBER on GPUs. 1. Generalized Born. Journal of Chemical Theory and Computation, 2012, 8, 1542-1555.	2.3	1,633
125	Protein structural model selection based on protein-dependent scoring function. Statistics and Its Interface, 2012, 5, 109-115.	0.2	7
126	An automatic tongue detection and segmentation framework for computer-aided tongue image analysis. , $2011, , .$		2

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127	PRIMEGENSw3: A Web-Based Tool for High-Throughput Primer and Probe Design. , 2011, , .		2
128	A Hybrid Consensus and Clustering Method for Protein Structure Selection. , 2011, , .		0
129	CORRELATION BETWEEN POSTTRANSLATIONAL MODIFICATION AND INTRINSIC DISORDER IN PROTEIN. , 2011, , .		16
130	A new clustering-based method for protein structure selection. , 2008, , .		4
131	COMBINING SEQUENCE AND STRUCTURAL PROFILES FOR PROTEIN SOLVENT ACCESSIBILITY PREDICTION. , 2008, , .		5
132	Visual Event Recognition in News Video using Kernel Methods with Multi-Level Temporal Alignment. , 2007, , .		33
133	FACE RECOGNITION — A GENERALIZED MARGINAL FISHER ANALYSIS APPROACH. International Journal of Image and Graphics, 2007, 07, 583-591.	1.2	1
134	Mapping Genes to Pathways Using Ontological Fuzzy Rule Systems. IEEE International Conference on Fuzzy Systems, 2007, , .	0.0	0
135	Rank-One Projections With Adaptive Margins for Face Recognition. IEEE Transactions on Systems, Man, and Cybernetics, 2007, 37, 1226-1236.	5.5	44
136	Element Rearrangement for Tensor-Based Subspace Learning. , 2007, , .		16
137	Human Gait Recognition With Matrix Representation. IEEE Transactions on Circuits and Systems for Video Technology, 2006, 16, 896-903.	5.6	114
138	Bioinformatics and Fuzzy Logic. , 2006, , .		6
139	Survey sequencing of soybean elucidates the genome structure, composition and identifies novel repeats. Functional Plant Biology, 2006, 33, 765.	1.1	10
140	A Fast Algorithm for Low-Resolution Protein Structure Prediction. Annual International Conference of the IEEE Engineering in Medicine and Biology Society, 2006, , .	0.5	1
141	Design and Implementation of Probability-Based Scoring Function for Peptide Mass Fingerprinting Protein Identification. Annual International Conference of the IEEE Engineering in Medicine and Biology Society, 2006, , .	0.5	0
142	Intrinsic Terminator Prediction and Its Application in Synechococcus sp. WH8102. Journal of Computer Science and Technology, 2005, 20, 465-482.	0.9	11
143	PROFILES AND FUZZY K-NEAREST NEIGHBOR ALGORITHM FOR PROTEIN SECONDARY STRUCTURE PREDICTION. , 2005, , .		12
144	Strategies for Inhibition of MDR1 Gene Expression. Molecular Pharmacology, 2004, 66, 268-275.	1.0	56

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145	GENOME-SCALE PROTEIN FUNCTION PREDICTION IN YEAST SACCHAROMYCES CEREVISIAE THROUGH INTEGRATING MULTIPLE SOURCES OF HIGH-THROUGHPUT DATA. , 2004, , .		9
146	P53-Dependent Cell-Killing by Selective Repression of Thymidine Kinase and Reduced Prodrug Activation. Molecular Pharmacology, 2003, 64, 289-297.	1.0	3
147	Characterization of Protein Structure and Function at Genome Scale with a Computational Prediction Pipeline., 2003, 25, 269-293.		3
148	Selective Inhibition of P-glycoprotein Expression in Multidrug-Resistant Tumor Cells by a Designed Transcriptional Regulator. Journal of Pharmacology and Experimental Therapeutics, 2002, 302, 963-971.	1.3	51
149	IDENTIFICATION OF REGULATORY BINDING SITES USING MINIMUM SPANNING TREES., 2002, , .		4
150	Studies of the Mechanism of Phenol Hydroxylase: Mutants Tyr289Phe, Asp54Asn, and Arg281Metâ€. Biochemistry, 2001, 40, 12369-12378.	1.2	39
151	Protein threading using PROSPECT: Design and evaluation. Proteins: Structure, Function and Bioinformatics, 2000, 40, 343-354.	1.5	2
152	The essential tryptophan residues of pig kidney aminoacylase. IUBMB Life, 1997, 43, 1277-1283.	1.5	1
153	An efficient branch-and-bound algorithm for the assignment of protein backbone NMR peaks. , 0, , .		11
154	Rank-one Projections with Adaptive Margins for Face Recognition., 0,,.		2
155	Bioinformatics Analyses of Arabidopsis thaliana Tiling Array Expression Data., 0,, 57-70.		O
156	Predicting Functional Modules Using Microarray and Protein Interaction Data., 0,, 307-329.		0
157	GUV-Net for high fidelity shoeprint generation. Complex & Intelligent Systems, 0, , 1.	4.0	3