Takashi Gojobori

List of Publications by Year in descending order

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312 papers 18,200 citations

20036 63 h-index 20625 120 g-index

328 all docs 328 docs citations

times ranked

328

23850 citing authors

#	Article	IF	CITATIONS
1	An alternative, zeaxanthin epoxidase-independent abscisic acid biosynthetic pathway in plants. Molecular Plant, 2022, 15, 151-166.	3.9	55
2	Seawater desalination based drinking water: Microbial characterization during distribution with and without residual chlorine. Water Research, 2022, 210, 117975.	5.3	13
3	An open chat with…Takashi Gojobori. FEBS Open Bio, 2022, 12, 338-339.	1.0	O
4	From shallow to deep: some lessons learned from application of machine learning for recognition of functional genomic elements in human genome. Human Genomics, 2022, 16, 7.	1.4	3
5	Affinity2Vec: drug-target binding affinity prediction through representation learning, graph mining, and machine learning. Scientific Reports, 2022, 12, 4751.	1.6	23
6	A cautionary signal from the Red Sea on the impact of increased dust activity on marine microbiota. BMC Genomics, 2022, 23, 277.	1.2	8
7	DeepSVP: integration of genotype and phenotype for structural variant prioritization using deep learning. Bioinformatics, 2022, 38, 1677-1684.	1.8	7
8	Role of C-Reactive Protein in Diabetic Inflammation. Mediators of Inflammation, 2022, 2022, 1-15.	1.4	22
9	Identification of lipolytic enzymes using high-throughput single-cell screening and sorting of a metagenomic library. New Biotechnology, 2022, 70, 102-108.	2.4	6
10	Population structure of indigenous inhabitants of Arabia. PLoS Genetics, 2021, 17, e1009210.	1.5	14
11	MetaCancer: A deep learning-based pan-cancer metastasis prediction model developed using multi-omics data. Computational and Structural Biotechnology Journal, 2021, 19, 4404-4411.	1.9	19
12	IBDDB: a manually curated and text-mining-enhanced database of genes involved in inflammatory bowel disease. Database: the Journal of Biological Databases and Curation, 2021, 2021, .	1.4	8
13	Characterization of microbiologically influenced corrosion by comprehensive metagenomic analysis of an inland oil field. Gene, 2021, 774, 145425.	1.0	10
14	Biomedical computing in the Arab world. Communications of the ACM, 2021, 64, 108-113.	3.3	0
15	Significant variants of type 2 diabetes in the Arabian Region through an Integration of exome databases. PLoS ONE, 2021, 16, e0249226.	1.1	0
16	Rise and fall of the global conversation and shifting sentiments during the COVID-19 pandemic. Humanities and Social Sciences Communications, $2021, 8, .$	1.3	12
17	Leptin and Obesity: Role and Clinical Implication. Frontiers in Endocrinology, 2021, 12, 585887.	1.5	363
18	A single neuron subset governs a single coactive neuron circuit in Hydra vulgaris, representing a possible ancestral feature of neural evolution. Scientific Reports, 2021, 11, 10828.	1.6	5

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19	Transmission dynamics of SARS-CoV-2 on the Diamond Princess uncovered using viral genome sequence analysis. Gene, 2021, 779, 145496.	1.0	7
20	CovMT: an interactive SARS-CoV-2 mutation tracker, with a focus on critical variants. Lancet Infectious Diseases, The, 2021, 21, 602.	4.6	57
21	Evolution of memory systemâ€related genes. FEBS Open Bio, 2021, 11, 3201-3210.	1.0	4
22	Development of a time-series shotgun metagenomics database for monitoring microbial communities at the Pacific coast of Japan. Scientific Reports, 2021, 11, 12222.	1.6	6
23	DES-Tcell is a knowledgebase for exploring immunology-related literature. Scientific Reports, 2021, 11, 14344.	1.6	1
24	Detection of SARS-CoV-2 variants requires urgent global coordination. International Journal of Infectious Diseases, 2021, 109, 50-53.	1.5	4
25	Seasonal and annual changes in the microbial communities of Ofunato Bay, Japan, based on metagenomics. Scientific Reports, 2021, 11, 17277.	1.6	4
26	Genome sequencing and identification of cellulase genes in Bacillus paralicheniformis strains from the Red Sea. BMC Microbiology, 2021, 21, 254.	1.3	10
27	DTi2Vec: Drug–target interaction prediction using network embedding and ensemble learning. Journal of Cheminformatics, 2021, 13, 71.	2.8	27
28	Re-infection with a different SARS-CoV-2 clade and prolonged viral shedding in a hematopoietic stem cell transplantation patient. International Journal of Infectious Diseases, 2021, 110, 267-271.	1.5	9
29	Angiotensin–Converting Enzyme (ACE) 1 Gene Polymorphism and Phenotypic Expression of COVID-19 Symptoms. Genes, 2021, 12, 1572.	1.0	28
30	Machine learning and deep learning methods that use omics data for metastasis prediction. Computational and Structural Biotechnology Journal, 2021, 19, 5008-5018.	1.9	69
31	Metagenomic analysis provides functional insights into seasonal change of a non-cyanobacterial prokaryotic community in temperate coastal waters. PLoS ONE, 2021, 16, e0257862.	1.1	5
32	Levothyroxine Treatment and the Risk of Cardiac Arrhythmias – Focus on the Patient Submitted to Thyroid Surgery. Frontiers in Endocrinology, 2021, 12, 758043.	1.5	9
33	Predicting Bone Metastasis Using Gene Expression-Based Machine Learning Models. Frontiers in Genetics, 2021, 12, 771092.	1.1	10
34	Integration of Droplet Microfluidic Tools for Single-cell Functional Metagenomics: An Engineering Head Start. Genomics, Proteomics and Bioinformatics, 2021, 19, 504-518.	3.0	4
35	What is the right sequencing approach? Solo VS extended family analysis in consanguineous populations. BMC Medical Genomics, 2020, 13, 103.	0.7	10
36	PATH ^{cre8} : A Tool That Facilitates the Searching for Heterologous Biosynthetic Routes. ACS Synthetic Biology, 2020, 9, 3217-3227.	1.9	7

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37	Functional Pangenome Analysis Shows Key Features of E Protein Are Preserved in SARS and SARS-CoV-2. Frontiers in Cellular and Infection Microbiology, 2020, 10, 405.	1.8	40
38	Sequencing effort dictates gene discovery in marine microbial metagenomes. Environmental Microbiology, 2020, 22, 4589-4603.	1.8	13
39	Atherosclerosis Linked to Aberrant Amino Acid Metabolism and Immunosuppressive Amino Acid Catabolizing Enzymes. Frontiers in Immunology, 2020, 11, 551758.	2.2	44
40	Splice2Deep: An ensemble of deep convolutional neural networks for improved splice site prediction in genomic DNA. Gene: X, 2020, 763, 100035.	2.3	33
41	Behavioral and brain-transcriptomic synchronization between the two opponents of a fighting pair of the fish Betta splendens. PLoS Genetics, 2020, 16, e1008831.	1.5	22
42	DTiGEMS+: drug–target interaction prediction using graph embedding, graph mining, and similarity-based techniques. Journal of Cheminformatics, 2020, 12, 44.	2.8	62
43	SARS-CoV-2 infections and COVID-19 mortalities strongly correlate with ACE1 I/D genotype. Gene, 2020, 758, 144944.	1.0	116
44	DES-ROD: Exploring Literature to Develop New Links between RNA Oxidation and Human Diseases. Oxidative Medicine and Cellular Longevity, 2020, 2020, 1-13.	1.9	3
45	Marine Metagenomic Sequence Counts of Reads Assigned to Taxa Consistently Proportionate to Read Counts Obtained for per g of Seawater Sample. , 2019, , 183-188.		1
46	Monitoring of the toxic dinoflagellate Alexandrium catenella in Osaka Bay, Japan using a massively parallel sequencing (MPS)-based technique. Harmful Algae, 2019, 89, 101660.	2.2	24
47	Construction of complete Tupaia belangeri transcriptome database by whole-genome and comprehensive RNA sequencing. Scientific Reports, 2019, 9, 12372.	1.6	16
48	Mining biosynthetic gene clusters in Virgibacillus genomes. BMC Genomics, 2019, 20, 696.	1.2	7
49	Regionalized nervous system in Hydra and the mechanism of its development. Gene Expression Patterns, 2019, 31, 42-59.	0.3	29
50	A Preliminary Metagenome Analysis Based on a Combination of Protein Domains. Proteomes, 2019, 7, 19.	1.7	0
51	Finding Nemo's Genes: A chromosomeâ€scale reference assembly of the genome of the orange clownfish <i>Amphiprion percula</i> . Molecular Ecology Resources, 2019, 19, 570-585.	2.2	55
52	The X Files: "The Mystery of X Chromosome Instability in Alzheimer's Disease― Frontiers in Genetics, 2019, 10, 1368.	1.1	25
53	Metagenomic Methods: From Seawater to the Database. , 2019, , 3-16.		1
54	In Memory of Vladimir B. Bajic (1952–2019). Genomics, Proteomics and Bioinformatics, 2019, 17, 473-474.	3.0	1

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55	Metagenome-based diversity analyses suggest a strong locality signal for bacterial communities associated with oyster aquaculture farms in Ofunato Bay. Gene, 2018, 665, 149-154.	1.0	14
56	Seasonal changes in the communities of photosynthetic picoeukaryotes in Ofunato Bay as revealed by shotgun metagenomic sequencing. Gene, 2018, 665, 127-132.	1.0	6
57	Seasonal changes in the abundance of bacterial genes related to dimethylsulfoniopropionate catabolism in seawater from Ofunato Bay revealed by metagenomic analysis. Gene, 2018, 665, 174-184.	1.0	17
58	Taxonomic profiles in metagenomic analyses of free-living microbial communities in the Ofunato Bay. Gene, 2018, 665, 192-200.	1.0	15
59	Basin-scale seasonal changes in marine free-living bacterioplankton community in the Ofunato Bay. Gene, 2018, 665, 185-191.	1.0	10
60	Whole-Genome Sequencing of 84 Japanese Eels Reveals Evidence against Panmixia and Support for Sympatric Speciation. Genes, 2018, 9, 474.	1.0	13
61	Global Ramifications of Dust and Sandstorm Microbiota. Genome Biology and Evolution, 2018, 10, 1970-1987.	1.1	44
62	Gene-by-gene or localized dosage compensation on the neo-X chromosome in Drosophila miranda. Genome Biology and Evolution, 2018, 10, 1875-1881.	1.1	8
63	In silico exploration of Red Sea Bacillus genomes for natural product biosynthetic gene clusters. BMC Genomics, 2018, 19, 382.	1.2	17
64	Functional interrogation of Plasmodium genus metabolism identifies species- and stage-specific differences in nutrient essentiality and drug targeting. PLoS Computational Biology, 2018, 14, e1005895.	1.5	24
65	FARNA: knowledgebase of inferred functions of non-coding RNA transcripts. Nucleic Acids Research, 2017, 45, gkw973.	6. 5	30
66	The genome of Chenopodium quinoa. Nature, 2017, 542, 307-312.	13.7	569
67	In silico screening for candidate chassis strains of free fatty acid-producing cyanobacteria. BMC Genomics, 2017, 18, 33.	1.2	11
68	Comparative genomic analysis of translation initiation mechanisms for genes lacking the Shine–Dalgarno sequence in prokaryotes. Nucleic Acids Research, 2017, 45, 3922-3931.	6.5	43
69	Building a bio-based industry in the Middle East through harnessing the potential of the Red Sea biodiversity. Applied Microbiology and Biotechnology, 2017, 101, 4837-4851.	1.7	10
70	Viva Europa, a Land of Excellence in Research and Innovation for Health and Wellbeing. Progress in Preventive Medicine (New York, N Y), 2017, 2, e006.	0.7	6
71	The Emerging Facets of Non-Cancerous Warburg Effect. Frontiers in Endocrinology, 2017, 8, 279.	1.5	59
72	Low Concentrations of Vitamin C Reduce the Synthesis of Extracellular Polymers and Destabilize Bacterial Biofilms. Frontiers in Microbiology, 2017, 8, 2599.	1.5	66

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73	Accelerated pseudogenization on the neo-X chromosome in Drosophila miranda. Nature Communications, 2016, 7, 13659.	5.8	15
74	V-GAP: Viral genome assembly pipeline. Gene, 2016, 576, 676-680.	1.0	9
75	Editorial. Gene, 2016, 576, 592.	1.0	O
76	Two-step evolution of endosymbiosis between hydra and algae. Molecular Phylogenetics and Evolution, 2016, 103, 19-25.	1.2	9
77	VaProS: a database-integration approach for protein/genome information retrieval. Journal of Structural and Functional Genomics, 2016, 17, 69-81.	1.2	9
78	Structural adaption of extremophile proteins to the environments with special reference to hydrophobic networks. Ecological Genetics and Genomics, 2016, 1, 1-5.	0.3	7
79	Selection pressure on human STR loci and its relevance in repeat expansion disease. Molecular Genetics and Genomics, 2016, 291, 1851-1869.	1.0	15
80	Evolutionary Transitions of MicroRNA-Target Pairs. Genome Biology and Evolution, 2016, 8, 1621-1633.	1.1	23
81	Different Endosymbiotic Interactions in Two Hydra Species Reflect the Evolutionary History of Endosymbiosis. Genome Biology and Evolution, 2016, 8, 2155-2163.	1.1	24
82	DESM: portal for microbial knowledge exploration systems. Nucleic Acids Research, 2016, 44, D624-D633.	6.5	12
83	Databases of the marine metagenomics. Gene, 2016, 576, 724-728.	1.0	20
84	Speciation of two gobioid species, Pterogobius elapoides and Pterogobius zonoleucus revealed by multi-locus nuclear and mitochondrial DNA analyses. Gene, 2016, 576, 593-602.	1.0	8
85	Single-cell technologies in environmental omics. Gene, 2016, 576, 701-707.	1.0	20
86	Metagenomic studies of the Red Sea. Gene, 2016, 576, 717-723.	1.0	36
87	Influences of diurnal sampling bias on fixed-point monitoring of plankton biodiversity determined using a massively parallel sequencing-based technique. Gene, 2016, 576, 667-675.	1.0	11
88	Comparative miRNAs analysis of Two contrasting broccoli inbred lines with divergent head-forming capacity under temperature stress. BMC Genomics, 2015, 16, 1026.	1.2	22
89	Function and Evolutionary Origin of Unicellular Camera-Type Eye Structure. PLoS ONE, 2015, 10, e0118415.	1.1	31
90	Challenges and Opportunities of Airborne Metagenomics. Genome Biology and Evolution, 2015, 7, 1216-1226.	1.1	97

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91	Eye-like ocelloids are built from different endosymbiotically acquired components. Nature, 2015, 523, 204-207.	13.7	74
92	Ecological genomics in Xanthomonas: the nature of genetic adaptation with homologous recombination and host shifts. BMC Genomics, 2015, 16, 188.	1.2	23
93	Marine Metagenome as A Resource for Novel Enzymes. Genomics, Proteomics and Bioinformatics, 2015, 13, 290-295.	3.0	41
94	Marine metagenomics as a source for bioprospecting. Marine Genomics, 2015, 24, 21-30.	0.4	51
95	Coevolution of Axon Guidance Molecule Slit and Its Receptor Robo. PLoS ONE, 2014, 9, e94970.	1.1	7
96	Tissue- and Stage-Dependent Dosage Compensation on the Neo-X Chromosome in Drosophila pseudoobscura. Molecular Biology and Evolution, 2014, 31, 614-624.	3.5	36
97	Adaptive divergence with gene flow in incipient speciation of <i><scp>M</scp>iscanthus floridulus</i> /i>/i>sinensis/i> complex (<scp>P</scp> oaceae). Plant Journal, 2014, 80, 834-847.	2.8	17
98	Evolutionarily conserved transcription factor Apontic controls the G1/S progression by inducing <i>cyclin E</i> during eye development. Proceedings of the National Academy of Sciences of the United States of America, 2014, 111, 9497-9502.	3.3	18
99	Origins of Japanese flowering cherry (Prunus subgenus Cerasus) cultivars revealed using nuclear SSR markers. Tree Genetics and Genomes, 2014, 10, 477-487.	0.6	48
100	Innexin gap junctions in nerve cells coordinate spontaneous contractile behavior in Hydra polyps. Scientific Reports, 2014, 4, 3573.	1.6	43
101	Dopamine Receptor Genes and Evolutionary Differentiation in the Domestication of Fighting Cocks and Long-Crowing Chickens. PLoS ONE, 2014, 9, e101778.	1.1	21
102	Evolutionary changes of multiple visual pigment genes in the complete genome of Pacific bluefin tuna. Proceedings of the National Academy of Sciences of the United States of America, 2013, 110, 11061-11066.	3.3	106
103	Computational analysis and functional expression of ancestral copepod luciferase. Gene, 2013, 528, 201-205.	1.0	20
104	Dynamic Evolution of Endogenous Retrovirus-Derived Genes Expressed in Bovine Conceptuses during the Period of Placentation. Genome Biology and Evolution, 2013, 5, 296-306.	1.1	30
105	H-InvDB in 2013: an omics study platform for human functional gene and transcript discovery. Nucleic Acids Research, 2013, 41, D915-D919.	6.5	22
106	Comparison of Gene Expression Profile of Epiretinal Membranes Obtained from Eyes with Proliferative Vitreoretinopathy to That of Secondary Epiretinal Membranes. PLoS ONE, 2013, 8, e54191.	1.1	30
107	The First Symbiont-Free Genome Sequence of Marine Red Alga, Susabi-nori (Pyropia yezoensis). PLoS ONE, 2013, 8, e57122.	1.1	150
108	The DNA Data Bank of Japan launches a new resource, the DDBJ Omics Archive of functional genomics experiments. Nucleic Acids Research, 2012, 40, D38-D42.	6.5	43

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109	Evolution of Bioluminescence in Marine Planktonic Copepods. Molecular Biology and Evolution, 2012, 29, 1669-1681.	3.5	48
110	A New Database (GCD) on Genome Composition for Eukaryote and Prokaryote Genome Sequences and Their Initial Analyses. Genome Biology and Evolution, 2012, 4, 501-512.	1.1	16
111	Comparative Genome Analysis of Three Eukaryotic Parasites with Differing Abilities To Transform Leukocytes Reveals Key Mediators of <i>Theileria</i> Induced Leukocyte Transformation. MBio, 2012, 3, e00204-12.	1.8	64
112	Infectious Endogenous Retroviruses in Cats and Emergence of Recombinant Viruses. Journal of Virology, 2012, 86, 8634-8644.	1.5	68
113	TP Atlas: integration and dissemination of advances in Targeted Proteins Research Program (TPRP)â€"structural biology project phase II in Japan. Journal of Structural and Functional Genomics, 2012, 13, 145-154.	1.2	2
114	Multiple Plastids Collected by the Dinoflagellate Dinophysis mitra through Kleptoplastidy. Applied and Environmental Microbiology, 2012, 78, 813-821.	1.4	56
115	A prioritization analysis of disease association by data-mining of functional annotation of human genes. Genomics, 2012, 99, 1-9.	1.3	11
116	The Temporal Sequence of the Mammalian Neocortical Neurogenetic Program Drives Mediolateral Pattern in the Chick Pallium. Developmental Cell, 2012, 22, 863-870.	3.1	81
117	Genome and Transcriptome Analysis of the Food-Yeast Candida utilis. PLoS ONE, 2012, 7, e37226.	1.1	28
118	Prediction of Protein-Destabilizing Polymorphisms by Manual Curation with Protein Structure. PLoS ONE, 2012, 7, e50445.	1.1	4
119	Gene dosage imbalance of human chromosome 21 in mouse embryonic stem cells differentiating to neurons. Gene, 2011, 481, 93-101.	1.0	6
120	Evolutionary Patterns of Recently Emerged Animal Duplogs. Genome Biology and Evolution, 2011, 3, 1119-1135.	1.1	18
121	Evolutionary history of dog rabies in Brazil. Journal of General Virology, 2011, 92, 85-90.	1.3	13
122	Welcome to Genome Biology and Evolution. Genome Biology and Evolution, 2010, 1, 1-1.	1.1	1
123	The transcript repeat element: the human Alu sequence as a component of gene networks influencing cancer. Functional and Integrative Genomics, 2010, 10, 307-319.	1.4	28
124	Low genetic diversities of rabies virus populations within different hosts in Brazil. Infection, Genetics and Evolution, 2010, 10, 278-283.	1.0	5
125	Endogenous non-retroviral RNA virus elements in mammalian genomes. Nature, 2010, 463, 84-87.	13.7	404
126	The dynamic genome of Hydra. Nature, 2010, 464, 592-596.	13.7	743

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127	Dynamic evolution of translation initiation mechanisms in prokaryotes. Proceedings of the National Academy of Sciences of the United States of America, 2010, 107, 6382-6387.	3.3	137
128	Evolutionary Pattern of Gene Homogenization between Primate-Specific Paralogs after Human and Macaque Speciation Using the 4-2-4 Method. Molecular Biology and Evolution, 2010, 27, 2152-2171.	3.5	19
129	DDBJ launches a new archive database with analytical tools for next-generation sequence data. Nucleic Acids Research, 2010, 38, D33-D38.	6.5	86
130	Nematogalectin, a nematocyst protein with GlyXY and galectin domains, demonstrates nematocyte-specific alternative splicing in <i>Hydra</i> Proceedings of the National Academy of Sciences of the United States of America, 2010, 107, 18539-18544.	3.3	69
131	A Non-sulfated Chondroitin Stabilizes Membrane Tubulation in Cnidarian Organelles. Journal of Biological Chemistry, 2010, 285, 25613-25623.	1.6	35
132	Gene expression profile of fibrovascular membranes from patients with proliferative diabetic retinopathy. British Journal of Ophthalmology, 2010, 94, 795-801.	2.1	19
133	Long-range neural and gap junction protein-mediated cues control polarity during planarian regeneration. Developmental Biology, 2010, 339, 188-199.	0.9	176
134	Genome Network Project: An Integrated Genomic Platform. Nature Precedings, 2009, , .	0.1	0
135	DDBJ Activities: Contribution to the Research in Information Biology. Nature Precedings, 2009, , .	0.1	0
136	Methods for Incorporating the Hypermutability of CpG Dinucleotides in Detecting Natural Selection Operating at the Amino Acid Sequence Level. Molecular Biology and Evolution, 2009, 26, 2275-2284.	3.5	18
137	VarySysDB: a human genetic polymorphism database based on all H-InvDB transcripts. Nucleic Acids Research, 2009, 37, D810-D815.	6.5	14
138	Midline governs axon pathfinding by coordinating expression of two major guidance systems. Genes and Development, 2009, 23, 1165-1170.	2.7	32
139	The transcriptional network that controls growth arrest and differentiation in a human myeloid leukemia cell line. Nature Genetics, 2009, 41, 553-562.	9.4	408
140	An evolutionary origin and selection process of goldfish. Gene, 2009, 430, 5-11.	1.0	75
141	The evolutionary study of small RNA-directed gene silencing pathways by investigating RNase III enzymes. Gene, 2009, 435, 1-8.	1.0	5
142	Mapping Human Genetic Diversity in Asia. Science, 2009, 326, 1541-1545.	6.0	557
143	Detecting Linkage between a Trait and a Marker in a Random Mating Population without Pedigree Record. PLoS ONE, 2009, 4, e4956.	1.1	0
144	The Promoter Signatures in Rice LEA Genes Can Be Used to Build a Co-expressing LEA Gene Network. Rice, 2008, 1, 177-187.	1.7	14

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145	The future of biocuration. Nature, 2008, 455, 47-50.	13.7	648
146	<i>Brachyury</i> â€downstream gene sets in a chordate, <i>Ciona intestinalis</i> : integrating notochord specification, morphogenesis and chordate evolution. Evolution & Development, 2008, 10, 37-51.	1.1	53
147	Evolution of complex structures: minicollagens shape the cnidarian nematocyst. Trends in Genetics, 2008, 24, 431-438.	2.9	117
148	Eukaryotic nuclear structure explains the evolutionary rate difference of ribosome export factors. Gene, 2008, 421, 7-13.	1.0	1
149	The origin of nucleus: Rebuild from the prokaryotic ancestors of ribosome export factors. Gene, 2008, 423, 149-152.	1.0	7
150	The evolutionary relationship between gene duplication and alternative splicing. Gene, 2008, 427, 19-31.	1.0	34
151	Evolution of Pacific Ocean and the Sea of Japan populations of the gobiid species, Pterogobius elapoides and Pterogobius zonoleucus, based on molecular and morphological analyses. Gene, 2008, 427, 7-18.	1.0	38
152	Cilium Evolution: Identification of a Novel Protein, Nematocilin, in the Mechanosensory Cilium of Hydra Nematocytes. Molecular Biology and Evolution, 2008, 25, 2009-2017.	3.5	27
153	Low conservation and species-specific evolution of alternative splicing in humans and mice: comparative genomics analysis using well-annotated full-length cDNAs. Nucleic Acids Research, 2008, 36, 6386-6395.	6.5	27
154	Distribution and Effects of Nonsense Polymorphisms in Human Genes. PLoS ONE, 2008, 3, e3393.	1.1	35
155	Transcriptional Interferences in cis Natural Antisense Transcripts of Humans and Mice. Genetics, 2007, 176, 1299-1306.	1.2	124
156	Compensatory Change of Interacting Amino Acids in the Coevolution of Transcriptional Coactivator MBF1 and TATA-Box–Binding Protein. Molecular Biology and Evolution, 2007, 24, 1458-1463.	3.5	20
157	Gene Cluster Analysis Method Identifies Horizontally Transferred Genes with High Reliability and Indicates that They Provide the Main Mechanism of Operon Gain in 8 Species of Î ³ -Proteobacteria. Molecular Biology and Evolution, 2007, 24, 805-813.	3.5	28
158	Curated genome annotation of Oryza sativa ssp. japonica and comparative genome analysis with Arabidopsis thaliana. Genome Research, 2007, 17, 175-183.	2.4	218
159	The evolutionary emergence of cell type-specific genes inferred from the gene expression analysis of <i>Hydra</i> . Proceedings of the National Academy of Sciences of the United States of America, 2007, 104, 14735-14740.	3.3	83
160	Increasing genetic diversity of hepatitis C virus in haemophiliacs with human immunodeficiency virus coinfection. Journal of General Virology, 2007, 88, 2513-2519.	1.3	9
161	Genetic Stability of Human T Lymphotropic Virus Type I despite Antiviral Pressures by CTLs. Journal of Immunology, 2007, 178, 5966-5972.	0.4	24
162	Diversity of preferred nucleotide sequences around the translation initiation codon in eukaryote genomes. Nucleic Acids Research, 2007, 36, 861-871.	6.5	181

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163	The H-Invitational Database (H-InvDB), a comprehensive annotation resource for human genes and transcripts. Nucleic Acids Research, 2007, 36, D793-D799.	6.5	57
164	Frequent emergence and functional resurrection of processed pseudogenes in the human and mouse genomes. Gene, 2007, 389, 196-203.	1.0	44
165	Mapping of chimpanzee full-length cDNAs onto the human genome unveils large potential divergence of the transcriptome. Gene, 2007, 399, 1-10.	1.0	13
166	Evolutionary origin of sex-related genes in the mouse brain. Gene, 2007, 406, 108-112.	1.0	1
167	Comparative genome analyses of nervous system-specific genes. Gene, 2006, 365, 130-136.	1.0	22
168	Evolution of metabolic networks by gain and loss of enzymatic reaction in eukaryotes. Gene, 2006, 365, 88-94.	1.0	18
169	magp4 gene may contribute to the diversification of cichlid morphs and their speciation. Gene, 2006, 373, 126-133.	1.0	23
170	Alternative splicing in human transcriptome: Functional and structural influence on proteins. Gene, 2006, 380, 63-71.	1.0	60
171	Differential evolutionary rates of duplicated genes in protein interaction network. Gene, 2006, 385, 57-63.	1.0	23
172	Radical amino acid change versus positive selection in the evolution of viral envelope proteins. Gene, 2006, 385, 83-88.	1.0	23
173	Degeneration after sexual differentiation in hydra and its relevance to the evolution of aging. Gene, 2006, 385, 64-70.	1.0	76
174	Cancer-related mutations in BRCA1-BRCT cause long-range structural changes in protein-protein binding sites: A molecular dynamics study. Proteins: Structure, Function and Bioinformatics, 2006, 66, 69-86.	1.5	20
175	Effects of U0126 and fibroblast growth factor on gene expression profile in Ciona intestinalis embryos as revealed by microarray analysis. Development Growth and Differentiation, 2006, 48, 391-400.	0.6	9
176	Exploration and Grading of Possible Genes from 183 Bacterial Strains by a Common Protocol to Identification of New Genes: Gene Trek in Prokaryote Space (GTPS). DNA Research, 2006, 13, 245-254.	1.5	18
177	Large-scale identification and characterization of alternative splicing variants of human gene transcripts using 56 419 completely sequenced and manually annotated full-length cDNAs. Nucleic Acids Research, 2006, 34, 3917-3928.	6.5	46
178	Rate of Evolution in Brain-Expressed Genes in Humans and Other Primates. PLoS Biology, 2006, 5, e13.	2.6	93
179	Rapid Evolution of Major Histocompatibility Complex Class I Genes in Primates Generates New Disease Alleles in Humans via Hitchhiking Diversity. Genetics, 2006, 173, 1555-1570.	1.2	100
180	The Evolutionary Rate of a Protein Is Influenced by Features of the Interacting Partners. Molecular Biology and Evolution, 2006, 23, 784-789.	3.5	32

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181	Comparative study of flux redistribution of metabolic pathway in glutamate production by two coryneform bacteria. Metabolic Engineering, 2005, 7, 59-69.	3.6	59
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