Steffen MĶller

List of Publications by Year in descending order

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59 papers 2,775 citations

304743 22 h-index 189892 50 g-index

64 all docs

64
docs citations

times ranked

64

4508 citing authors

#	Article	IF	CITATIONS
1	Scoring functions for drug-effect similarity. Briefings in Bioinformatics, 2021, 22, .	6.5	15
2	Perspectives on automated composition of workflows in the life sciences. F1000Research, 2021, 10, 897.	1.6	7
3	Single-cell analyses of aging, inflammation and senescence. Ageing Research Reviews, 2020, 64, 101156.	10.9	85
4	Suppression of the TGF- \hat{l}^2 pathway by a macrolide antibiotic decreases fibrotic responses by ocular fibroblasts <i>in vitro</i> . Royal Society Open Science, 2020, 7, 200441.	2.4	5
5	Healthspan pathway maps in C. elegans and humans highlight transcription, proliferation/biosynthesis and lipids. Aging, 2020, 12, 12534-12581.	3.1	12
6	Scalable Workflows and Reproducible Data Analysis for Genomics. Methods in Molecular Biology, 2019, 1910, 723-745.	0.9	25
7	Gene-diet interactions associated with complex trait variation in an advanced intercross outbred mouse line. Nature Communications, 2019, 10, 4097.	12.8	35
8	Cloud-Based High Throughput Virtual Screening in Novel Drug Discovery. Lecture Notes in Computer Science, 2019, , 250-278.	1.3	12
9	Health and Aging: Unifying Concepts, Scores, Biomarkers and Pathways. , 2019, 10, 883.		56
10	A workflow for the integrative transcriptomic description of molecular pathology and the suggestion of normalizing compounds, exemplified by Parkinson's disease. Scientific Reports, 2018, 8, 7937.	3.3	5
11	FocusHeuristics – expression-data-driven network optimization and disease gene prediction. Scientific Reports, 2017, 7, 42638.	3. 3	17
12	Robust Cross-Platform Workflows: How Technical and Scientific Communities Collaborate to Develop, Test and Share Best Practices for Data Analysis. Data Science and Engineering, 2017, 2, 232-244.	6.4	19
13	Male-specific association between MT-ND4 11719 A/G polymorphism and ulcerative colitis: a mitochondria-wide genetic association study. BMC Gastroenterology, 2016, 16, 118.	2.0	17
14	Dissecting genetics of cutaneous miRNA in a mouse model of an autoimmune blistering disease. BMC Genomics, 2016, 17, 112.	2.8	8
15	Mitochondrial gene polymorphisms alter hepatic cellular energy metabolism and aggravate diet-induced non-alcoholic steatohepatitis. Molecular Metabolism, 2016, 5, 283-295.	6.5	45
16	Tools and data services registry: a community effort to document bioinformatics resources. Nucleic Acids Research, 2016, 44, D38-D47.	14.5	113
17	Liver steatosis is a risk factor for hepatotoxicity in patients with inflammatory bowel disease under immunosuppressive treatment. European Journal of Gastroenterology and Hepatology, 2015, 27, 698-704.	1.6	28
18	The retinoidâ€related orphan receptor alpha is essential for the endâ€stage effector phase of experimental epidermolysis bullosa acquisita. Journal of Pathology, 2015, 237, 111-122.	4.5	23

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19	Quantitative Trait Locus Analysis Implicates CD4+/CD44high Memory T Cells in the Pathogenesis of Murine Autoimmune Pancreatitis. PLoS ONE, 2015, 10, e0136298.	2.5	7
20	Hardy-Weinberg equilibrium revisited for inferences on genotypes featuring allele and copy-number variations. Scientific Reports, 2015, 5, 9066.	3.3	3
21	Allelic and copy-number variations of Fcl^3Rs affect granulocyte function and susceptibility for autoimmune blistering diseases. Journal of Autoimmunity, 2015, 61, 36-44.	6.5	32
22	A Family with Atypical Hailey Hailey Disease- Is There More to the Underlying Genetics than ATP2C1?. PLoS ONE, 2015, 10, e0121253.	2.5	8
23	ptRNApred: computational identification and classification of post-transcriptional RNA. Nucleic Acids Research, 2014, 42, e167-e167.	14.5	6
24	Nerve Conduction Velocity Is Regulated by the Inositol Polyphosphate-4-Phosphatase II Gene. American Journal of Pathology, 2014, 184, 2420-2429.	3.8	8
25	Community-driven development for computational biology at Sprints, Hackathons and Codefests. BMC Bioinformatics, 2014, 15, S7.	2.6	42
26	Genome-wide mapping of gene–microbiota interactions in susceptibility to autoimmune skin blistering. Nature Communications, 2013, 4, 2462.	12.8	87
27	Genetic control of psoriasis is relatively distinct from that of metabolic syndrome and coronary artery disease. Experimental Dermatology, 2013, 22, 552-553.	2.9	37
28	Combining genetic mapping with genome-wide expression in experimental autoimmune encephalomyelitis highlights a gene network enriched for T cell functions and candidate genes regulating autoimmunity. Human Molecular Genetics, 2013, 22, 4952-4966.	2.9	11
29	Genetic Control of Spontaneous Arthritis in a Four-Way Advanced Intercross Line. PLoS ONE, 2013, 8, e75611.	2.5	2
30	Identification of Quantitative Trait Loci in Experimental Epidermolysis Bullosa Acquisita. Journal of Investigative Dermatology, 2012, 132, 1409-1415.	0.7	35
31	Bioinformatics tools and database resources for systems genetics analysis in mice-a short review and an evaluation of future needs. Briefings in Bioinformatics, 2012, 13, 135-142.	6.5	11
32	xQTL workbench: a scalable web environment for multi-level QTL analysis. Bioinformatics, 2012, 28, 1042-1044.	4.1	16
33	High-throughput Molecular Docking Now in Reach for a Wider Biochemical Community. , 2012, , .		2
34	Scalable Computing for Evolutionary Genomics. Methods in Molecular Biology, 2012, 856, 529-545.	0.9	5
35	AFP-Pred: A random forest approach for predicting antifreeze proteins from sequence-derived properties. Journal of Theoretical Biology, 2011, 270, 56-62.	1.7	226
36	Community-driven computational biology with Debian Linux. BMC Bioinformatics, 2010, 11, S5.	2.6	42

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37	Time course transcriptomics of IFNB1b drug therapy in multiple sclerosis. Autoimmunity, 2010, 43, 172-178.	2.6	34
38	SPRED: A machine learning approach for the identification of classical and non-classical secretory proteins in mammalian genomes. Biochemical and Biophysical Research Communications, 2010, 391, 1306-1311.	2.1	33
39	SNPtoGO: characterizing SNPs by enriched GO terms. Bioinformatics, 2008, 24, 146-148.	4.1	24
40	Integrating ARC grid middleware with Taverna workflows. Bioinformatics, 2008, 24, 1221-1222.	4.1	14
41	Multiple Sclerosis Therapy Monitoring Based on Gene Expression. Current Pharmaceutical Design, 2006, 12, 3761-79.	1.9	7
42	Comparative Genomics for the Investigation of Autoimmune Diseases. Current Pharmaceutical Design, 2006, 12, 3707-22.	1.9	1
43	Definition of a 1.06-Mb Region Linked to Neuroinflammation in Humans, Rats and Mice. Genetics, 2006, 173, 1539-1545.	2.9	20
44	Fine Mapping of Collagen-Induced Arthritis Quantitative Trait Loci in an Advanced Intercross Line. Journal of Immunology, 2006, 177, 7042-7049.	0.8	23
45	Gene expression profiling of arthritis using a QTL chip reveals a complex gene regulation of the Cia5 region in mice. Genes and Immunity, 2005, 6, 575-583.	4.1	16
46	Cytoskeletal Rearrangements in Synovial Fibroblasts as a Novel Pathophysiological Determinant of Modeled Rheumatoid Arthritis. PLoS Genetics, 2005, 1, e48.	3.5	49
47	In silico fine-mapping: narrowing disease-associated loci by intergenomics. Bioinformatics, 2005, 21, 1737-1738.	4.1	2
48	Collection of soluble variants of membrane proteins for transcriptomics and proteomics. In Silico Biology, 2005, 5, 295-311.	0.9	2
49	Non-linear conversion between genetic and physical chromosomal distances. Bioinformatics, 2004, 20, 1966-1967.	4.1	8
50	Intergenomic consensus in multifactorial inheritance loci: the case of multiple sclerosis. Genes and Immunity, 2004, 5, 615-620.	4.1	20
51	Selecting SNPs for association studies based on population frequencies: a novel interactive tool and its application to polygenic diseases. In Silico Biology, 2004, 4, 417-27.	0.9	3
52	Expressionview: visualization of quantitative trait loci and gene-expression data in Ensembl. Genome Biology, 2003, 4, R77.	9.6	21
53	The KOX zinc finger genes: Genome wide mapping of 368 ZNF PAC clones with zinc finger gene clusters predominantly in 23 chromosomal loci are confirmed by human sequences annotated in EnsEMBL. Cytogenetic and Genome Research, 2002, 98, 147-153.	1.1	22
54	Evaluation of methods for the prediction of membrane spanning regions. Bioinformatics, 2002, 18, 218-218.	4.1	12

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55	Consistent integration of non-reliable heterogeneous information resources applied to the annotation of transmembrane proteins. Computers & Chemistry, 2001, 26, 41-49.	1.2	11
56	Evaluation of methods for the prediction of membrane spanning regions. Bioinformatics, 2001, 17, 646-653.	4.1	1,050
57	A collection of well characterised integral membrane proteins. Bioinformatics, 2000, 16, 1159-1160.	4.1	92
58	A novel method for automatic functional annotation of proteins. Bioinformatics, 1999, 15, 228-233.	4.1	84
59	EDITtoTrEMBL: a distributed approach to high-quality automated protein sequence annotation. Bioinformatics, 1999, 15, 219-227.	4.1	38