Martin Hofmann-Apitius

List of Publications by Year in descending order

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118 papers 2,986 citations

28 h-index 223800 46 g-index

154 all docs

154 docs citations

154 times ranked

4141 citing authors

#	Article	IF	CITATIONS
1	Development of a benchmark corpus to support the automatic extraction of drug-related adverse effects from medical case reports. Journal of Biomedical Informatics, 2012, 45, 885-892.	4.3	251
2	A comprehensive, cell specific microRNA catalogue of human peripheral blood. Nucleic Acids Research, 2017, 45, 9290-9301.	14.5	159
3	Hyaluronan-oligosaccharide-induced transcription of metalloproteases. Journal of Cell Science, 2004, 117, 359-367.	2.0	149
4	Detection of IUPAC and IUPAC-like chemical names. Bioinformatics, 2008, 24, i268-i276.	4.1	102
5	The Impact of Pathway Database Choice on Statistical Enrichment Analysis and Predictive Modeling. Frontiers in Genetics, 2019, 10, 1203.	2.3	78
6	ADO: A disease ontology representing the domain knowledge specific to Alzheimer's disease. Alzheimer's and Dementia, 2014, 10, 238-246.	0.8	77
7	COVID-19 Knowledge Graph: a computable, multi-modal, cause-and-effect knowledge model of COVID-19 pathophysiology. Bioinformatics, 2021, 37, 1332-1334.	4.1	70
8	Towards a 21st-century roadmap for biomedical research and drug discovery: consensus report and recommendations. Drug Discovery Today, 2017, 22, 327-339.	6.4	64
9	Using Multi-Scale Genetic, Neuroimaging and Clinical Data for Predicting Alzheimer's Disease and Reconstruction of Relevant Biological Mechanisms. Scientific Reports, 2018, 8, 11173.	3.3	59
10	Grid-enabled Virtual Screening Against Malaria. Journal of Grid Computing, 2008, 6, 29-43.	3.9	56
11	Comorbidity Analysis between Alzheimer's Disease and Type 2 Diabetes Mellitus (T2DM) Based on Shared Pathways and the Role of T2DM Drugs. Journal of Alzheimer's Disease, 2017, 60, 721-731.	2.6	55
12	Bioinformatics Mining and Modeling Methods for the Identification of Disease Mechanisms in Neurodegenerative Disorders. International Journal of Molecular Sciences, 2015, 16, 29179-29206.	4.1	47
13	Design of New Plasmepsin Inhibitors:  A Virtual High Throughput Screening Approach on the EGEE Grid. Journal of Chemical Information and Modeling, 2007, 47, 1818-1828.	5.4	46
14	From integrative disease modeling to predictive, preventive, personalized and participatory (P4) medicine. EPMA Journal, 2013, 4, 23.	6.1	46
15	Towards the taxonomy of human disease. Nature Reviews Drug Discovery, 2015, 14, 75-76.	46.4	46
16	Computable causeâ€andâ€effect models of healthy and Alzheimer's disease states and their mechanistic differential analysis. Alzheimer's and Dementia, 2015, 11, 1329-1339.	0.8	46
17	Challenges and opportunities for oncology biomarker discovery. Drug Discovery Today, 2013, 18, 614-624.	6.4	44
18	Contribution of syndecans to cellular uptake and fibrillation of \hat{l}_{\pm} -synuclein and tau. Scientific Reports, 2019, 9, 16543.	3.3	44

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19	ComPath: an ecosystem for exploring, analyzing, and curating mappings across pathway databases. Npj Systems Biology and Applications, 2018, 4, 3.	3.0	43
20	PathMe: merging and exploring mechanistic pathway knowledge. BMC Bioinformatics, 2019, 20, 243.	2.6	42
21	Multimodal mechanistic signatures for neurodegenerative diseases (NeuroMMSig): a web server for mechanism enrichment. Bioinformatics, 2017, 33, 3679-3681.	4.1	39
22	â€~HypothesisFinder:' A Strategy for the Detection of Speculative Statements in Scientific Text. PLoS Computational Biology, 2013, 9, e1003117.	3.2	37
23	Mining biomarker information in biomedical literature. BMC Medical Informatics and Decision Making, 2012, 12, 148.	3.0	36
24	Multi-omic approach decodes paradoxes of the triple-negative breast cancer: lessons for predictive, preventive and personalised medicine. Amino Acids, 2018, 50, 383-395.	2.7	34
25	Differences in cohort study data affect external validation of artificial intelligence models for predictive diagnostics of dementia - lessons for translation into clinical practice. EPMA Journal, 2020, 11, 367-376.	6.1	34
26	Identification of new drug classification terms in textual resources. Bioinformatics, 2007, 23, i264-i272.	4.1	33
27	Challenges in the association of human single nucleotide polymorphism mentions with unique database identifiers. BMC Bioinformatics, 2011, 12, S4.	2.6	33
28	Construction of biological networks from unstructured information based on a semi-automated curation workflow. Database: the Journal of Biological Databases and Curation, 2015, 2015, bav057.	3.0	33
29	Contribution of syndecans to cellular internalization and fibrillation of amyloid-β(1–42). Scientific Reports, 2019, 9, 1393.	3.3	33
30	OSIRISv1.2: A named entity recognition system for sequence variants of genes in biomedical literature. BMC Bioinformatics, 2008, 9, 84.	2.6	31
31	Text mining for systems biology. Drug Discovery Today, 2014, 19, 140-144.	6.4	31
32	WISDOM-II: Screening against multiple targets implicated in malaria using computational grid infrastructures. Malaria Journal, 2009, 8, 88.	2.3	29
33	PDON: Parkinson's disease ontology for representation and modeling of the Parkinson's disease knowledge domain. Theoretical Biology and Medical Modelling, 2015, 12, 20.	2.1	29
34	Alternative to Consensus ScoringA New Approach Toward the Qualitative Combination of Docking Algorithms. Journal of Chemical Information and Modeling, 2007, 47, 1036-1044.	5.4	28
35	Virtual screening on large scale grids. Parallel Computing, 2007, 33, 289-301.	2.1	27
36	The COVID-19 Ontology. Bioinformatics, 2021, 36, 5703-5705.	4.1	27

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37	Knowledge Retrieval from PubMed Abstracts and Electronic Medical Records with the Multiple Sclerosis Ontology. PLoS ONE, 2015, 10, e0116718.	2.5	26
38	Big data and data repurposing - using existing data to answer new questions in vascular dementia research. BMC Neurology, 2017, 17, 72.	1.8	24
39	Analytical Strategy to Prioritize Alzheimer's Disease Candidate Genes in Gene Regulatory Networks Using Public Expression Data. Journal of Alzheimer's Disease, 2017, 59, 1237-1254.	2.6	23
40	Detecting miRNA Mentions and Relations in Biomedical Literature. F1000Research, 2014, 3, 205.	1.6	22
41	Concept-Based Semi-Automatic Classification of Drugs. Journal of Chemical Information and Modeling, 2009, 49, 1986-1992.	5.4	21
42	Challenges in mining the literature for chemical information. RSC Advances, 2013, 3, 16194.	3.6	20
43	Re-curation and rational enrichment of knowledge graphs in Biological Expression Language. Database: the Journal of Biological Databases and Curation, 2019, 2019, .	3.0	20
44	Bridging Scales in Alzheimer's Disease: Biological Framework for Brain Simulation With The Virtual Brain. Frontiers in Neuroinformatics, 2021, 15, 630172.	2.5	20
45	Knowledge environments representing molecular entities for the virtual physiological human. Philosophical Transactions Series A, Mathematical, Physical, and Engineering Sciences, 2008, 366, 3091-3110.	3.4	19
46	Detecting miRNA Mentions and Relations in Biomedical Literature. F1000Research, 2014, 3, 205.	1.6	19
47	Integration and mining of malaria molecular, functional and pharmacological data: how far are we from a chemogenomic knowledge space?. Malaria Journal, 2006, 5, 110.	2.3	18
48	Of Mice and Men: Comparative Analysis of Neuro-Inflammatory Mechanisms in Human and Mouse Using Cause-and-Effect Models. Journal of Alzheimer's Disease, 2017, 59, 1045-1055.	2.6	18
49	ANMerge: A Comprehensive and Accessible Alzheimer's Disease Patient-Level Dataset. Journal of Alzheimer's Disease, 2021, 79, 423-431.	2.6	18
50	Drug2ways: Reasoning over causal paths in biological networks for drug discovery. PLoS Computational Biology, 2020, 16, e1008464.	3.2	18
51	BEL Commons: an environment for exploration and analysis of networks encoded in Biological Expression Language. Database: the Journal of Biological Databases and Curation, 2018, 2018, .	3.0	17
52	Detecting miRNA Mentions and Relations in Biomedical Literature. F1000Research, 0, 3, 205.	1.6	17
53	On the influence of several factors on pathway enrichment analysis. Briefings in Bioinformatics, 2022, 23, .	6.5	17
54	A new optimization phase for scientific workflow management systems. Future Generation Computer Systems, 2014, 36, 352-362.	7. 5	16

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55	A Computational Approach for Mapping Heme Biology in the Context of Hemolytic Disorders. Frontiers in Bioengineering and Biotechnology, 2020, 8, 74.	4.1	16
56	Linking COVID-19 and Heme-Driven Pathophysiologies: A Combined Computational–Experimental Approach. Biomolecules, 2021, 11, 644.	4.0	16
57	Reconstruction of Chemical Molecules from Images. , 2007, 2007, 4609-12.		15
58	IDENTIFYING GENE-SPECIFIC VARIATIONS IN BIOMEDICAL TEXT. Journal of Bioinformatics and Computational Biology, 2007, 05, 1277-1296.	0.8	15
59	Boosting translational research on Alzheimer's disease in Europe: The Innovative Medicine Initiative AD research platform. Alzheimer's and Dementia, 2015, 11, 1121-1122.	0.8	15
60	Data science in neurodegenerative disease: its capabilities, limitations, and perspectives. Current Opinion in Neurology, 2020, 33, 249-254.	3.6	15
61	Towards a Pathway Inventory of the Human Brain for Modeling Disease Mechanisms Underlying Neurodegeneration. Journal of Alzheimer's Disease, 2016, 52, 1343-1360.	2.6	14
62	Using Drugs as Molecular Probes: AÂComputational Chemical Biology Approach in Neurodegenerative Diseases. Journal of Alzheimer's Disease, 2017, 56, 677-686.	2.6	14
63	Virtual connectomic datasets in Alzheimer's Disease and aging using whole-brain network dynamics modelling. ENeuro, 2021, 8, ENEURO.0475-20.2021.	1.9	14
64	NeuroRDF: semantic integration of highly curated data to prioritize biomarker candidates in Alzheimer's disease. Journal of Biomedical Semantics, 2016, 7, 45.	1.6	13
65	Novel approaches to develop community-built biological network models for potential drug discovery. Expert Opinion on Drug Discovery, 2017, 12, 1-9.	5.0	13
66	Challenges of Integrative Disease Modeling in Alzheimer's Disease. Frontiers in Molecular Biosciences, 2019, 6, 158.	3.5	13
67	A hybrid approach unveils drug repurposing candidates targeting an Alzheimer pathophysiology mechanism. Patterns, 2022, 3, 100433.	5.9	13
68	In Silico Drug Discovery Approaches on Grid Computing Infrastructures. Current Clinical Pharmacology, 2010, 5, 37-46.	0.6	12
69	Variational Autoencoder Modular Bayesian Networks for Simulation of Heterogeneous Clinical Study Data. Frontiers in Big Data, 2020, 3, 16.	2.9	12
70	A method for the rational selection of drug repurposing candidates from multimodal knowledge harmonization. Scientific Reports, 2021, 11, 11049.	3.3	12
71	Computational Modelling Approaches on Epigenetic Factors in Neurodegenerative and Autoimmune Diseases and Their Mechanistic Analysis. Journal of Immunology Research, 2015, 2015, 1-10.	2.2	11
72	Is dementia research ready for big data approaches?. BMC Medicine, 2015, 13, 145.	5 . 5	11

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73	Knowledgeâ€driven computational modeling in Alzheimer's disease research: Current state and future trends. Alzheimer's and Dementia, 2017, 13, 1292-1302.	0.8	11
74	Neuroimaging Feature Terminology: A Controlled Terminology for the Annotation of Brain Imaging Features. Journal of Alzheimer's Disease, 2017, 59, 1153-1169.	2.6	11
75	Clustering of Alzheimer's and Parkinson's disease based on genetic burden of shared molecular mechanisms. Scientific Reports, 2020, 10, 19097.	3.3	11
76	PS4DR: a multimodal workflow for identification and prioritization of drugs based on pathway signatures. BMC Bioinformatics, 2020, 21, 231.	2.6	11
77	STonKGs: a sophisticated transformer trained on biomedical text and knowledge graphs. Bioinformatics, 2022, 38, 1648-1656.	4.1	11
78	@neuLink: a service-oriented application for biomedical knowledge discovery. Studies in Health Technology and Informatics, 2008, 138, 165-72.	0.3	11
79	Reasoning over genetic variance information in cause-and-effect models of neurodegenerative diseases. Briefings in Bioinformatics, 2016, 17, 505-516.	6.5	10
80	Data-Driven Modeling of Knowledge Assemblies in Understanding Comorbidity Between Type 2 Diabetes Mellitus and Alzheimer's Disease. Journal of Alzheimer's Disease, 2020, 78, 87-95.	2.6	10
81	A new optimization phase for scientific workflow management systems. , 2012, , .		9
82	Exploring novel mechanistic insights in Alzheimer's disease by assessing reliability of protein interactions. Scientific Reports, 2015, 5, 13634.	3.3	9
83	GuiltyTargets: Prioritization of Novel Therapeutic Targets With Network Representation Learning. IEEE/ACM Transactions on Computational Biology and Bioinformatics, 2022, 19, 491-500.	3.0	9
84	Pathway Based Analysis of Genes and Interactions Influencing Porcine Testis Samples from Boars with Divergent Androstenone Content in Back Fat. PLoS ONE, 2014, 9, e91077.	2.5	9
85	Grid-Added Value to Address Malaria. IEEE Transactions on Information Technology in Biomedicine, 2008, 12, 173-181.	3.2	8
86	<i>NeuroTransDB</i> : highly curated and structured transcriptomic metadata for neurodegenerative diseases. Database: the Journal of Biological Databases and Curation, 2015, 2015, bav099.	3.0	8
87	Identification of histone modifications in biomedical text for supporting epigenomic research. BMC Bioinformatics, 2009, 10, S28.	2.6	7
88	HuPSON: the human physiology simulation ontology. Journal of Biomedical Semantics, 2013, 4, 35.	1.6	7
89	CSEO – the Cigarette Smoke Exposure Ontology. Journal of Biomedical Semantics, 2014, 5, 31.	1.6	7
90	A systematic approach for identifying shared mechanisms in epilepsy and its comorbidities. Database: the Journal of Biological Databases and Curation, 2018, 2018, .	3.0	7

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91	Realistic simulation of virtual multi-scale, multi-modal patient trajectories using Bayesian networks and sparse auto-encoders. Scientific Reports, 2020, 10, 10971.	3.3	7
92	Identifying multimodal signatures underlying the somatic comorbidity of psychosis: the COMMITMENT roadmap. Molecular Psychiatry, 2021, 26, 722-724.	7.9	7
93	A UNICORE Plugin for HPC-Enabled Scientific Workflows in Taverna 2.2. , 2011, , .		6
94	PLIO: an ontology for formal description of protein–ligand interactions. Bioinformatics, 2011, 27, 1684-1690.	4.1	6
95	Biomarker-guided translation of brain imaging into disease pathway models. Scientific Reports, 2013, 3, 3375.	3.3	6
96	Linking hypothetical knowledge patterns to disease molecular signatures for biomarker discovery in Alzheimer's disease. Genome Medicine, 2014, 6, 97.	8.2	6
97	Systems approach for the selection of micro-RNAs as therapeutic biomarkers of anti-EGFR monoclonal antibody treatment in colorectal cancer. Scientific Reports, 2015, 5, 8013.	3.3	6
98	CLEP: a hybrid data- and knowledge-driven framework for generating patient representations. Bioinformatics, 2021, 37, 3311-3318.	4.1	6
99	Linking hypothetical knowledge patterns to disease molecular signatures for biomarker discovery in Alzheimer's disease. Genome Medicine, 2014, 6, 97.	8.2	6
100	Text mining in livestock animal science: Introducing the potential of text mining to animal sciences 1. Journal of Animal Science, 2012, 90, 3666-3676.	0.5	5
101	Identification of gene co-expression clusters in liver tissues from multiple porcine populations with high and low backfat androstenone phenotype. BMC Genetics, 2015, 16, 21.	2.7	5
102	SVM Based Descriptor Selection and Classification of Neurodegenerative Disease Drugs for Pharmacological Modeling. Molecular Informatics, 2013, 32, 241-249.	2.5	4
103	Using predictive machine learning models for drug response simulation by calibrating patient-specific pathway signatures. Npj Systems Biology and Applications, 2021, 7, 40.	3.0	4
104	Converting Alzheimer's Disease Map into a Heavyweight Ontology: A Formal Network to Integrate Data. Lecture Notes in Computer Science, 2019, , 207-215.	1.3	4
105	Deep Learning-based detection of psychiatric attributes from German mental health records. International Journal of Medical Informatics, 2022, 161, 104724.	3.3	4
106	Integrative data semantics through a model-enabled data stewardship. Bioinformatics, 2022, 38, 3850-3852.	4.1	4
107	DockFlow: Achieving interoperability of protein docking tools across heterogeneous Grid middleware. International Journal of Ad Hoc and Ubiquitous Computing, 2010, 6, 235.	0.5	3
108	Quantifying mechanisms in neurodegenerative diseases (NDDs) using candidate mechanism perturbation amplitude (CMPA) algorithm. BMC Bioinformatics, 2019, 20, 494.	2.6	3

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109	DecoPath: a web application for decoding pathway enrichment analysis. NAR Genomics and Bioinformatics, 2021, 3, Iqab087.	3.2	3
110	Towards a global investigation of transcriptomic signatures through co-expression networks and pathway knowledge for the identification of disease mechanisms. Nucleic Acids Research, 2021, 49, 7939-7953.	14.5	3
111	MultiPaths: a Python framework for analyzing multi-layer biological networks using diffusion algorithms. Bioinformatics, 2021, 37, 137-139.	4.1	3
112	Elucidating gene expression patterns across multiple biological contexts through a large-scale investigation of transcriptomic datasets. BMC Bioinformatics, 2022, 23, .	2.6	3
113	An Improved Weightedâ€Residue Profile Based Method of Using Protein–Ligand Interaction Information in Increasing Hits Selection from Virtual Screening: A Study on Virtual Screening of Human GPCR A2A Receptor Antagonists. Molecular Informatics, 2010, 29, 781-791.	2.5	2
114	A network model of genomic hormone interactions underlying dementia and its translational validation through serendipitous off-target effect. Journal of Translational Medicine, 2013, 11, 177.	4.4	2
115	A Systems Biology Approach for Hypothesizing the Effect of Genetic Variants on Neuroimaging Features in Alzheimer's Disease. Journal of Alzheimer's Disease, 2021, 80, 831-840.	2.6	2
116	A robust framework for rapid deployment of a virtual screening laboratory. Studies in Health Technology and Informatics, 2009, 147, 212-21.	0.3	2
117	Comparison and aggregation of event sequences across ten cohorts to describe the consensus biomarker evolution in Alzheimer's disease. Alzheimer's Research and Therapy, 2022, 14, 55.	6.2	2
118	OUP accepted manuscript. Database: the Journal of Biological Databases and Curation, 2021, 2021, .	3.0	0