## Remco R Bouckaert

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/844741/publications.pdf

Version: 2024-02-01

33 papers 12,687 citations

331670 21 h-index 395702 33 g-index

54 all docs

54 docs citations

54 times ranked 16566 citing authors

#	Article	IF	CITATIONS
1	Multidimensional Phylogenetic Metrics Identify Class I Aminoacyl-tRNA Synthetase Evolutionary Mosaicity and Inter-Modular Coupling. International Journal of Molecular Sciences, 2022, 23, 1520.	4.1	5
2	StarBeast3: Adaptive Parallelized Bayesian Inference under the Multispecies Coalescent. Systematic Biology, 2022, 71, 901-916.	5.6	29
3	An Efficient Coalescent Epoch Model for Bayesian Phylogenetic Inference. Systematic Biology, 2022, 71, 1549-1560.	5.6	19
4	Detecting contact in language trees: a Bayesian phylogenetic model with horizontal transfer. Humanities and Social Sciences Communications, 2022, 9, .	2.9	9
5	Bayesian Inference of Species Trees using Diffusion Models. Systematic Biology, 2021, 70, 145-161.	5.6	21
6	Adaptive dating and fast proposals: Revisiting the phylogenetic relaxed clock model. PLoS Computational Biology, 2021, 17, e1008322.	3.2	60
7	Phylodynamics reveals the role of human travel and contact tracing in controlling the first wave of COVID-19 in four island nations. Virus Evolution, 2021, 7, veab052.	4.9	35
8	Real-Time Genomics for Tracking Severe Acute Respiratory Syndrome Coronavirus 2 Border Incursions after Virus Elimination, New Zealand. Emerging Infectious Diseases, 2021, 27, 2361-2368.	4.3	27
9	Bayesian phylogenetic analysis of linguistic data using BEAST. Journal of Language Evolution, 2021, 6, 119-135.	1.7	9
10	Combinatorial perspectives on Dollo-k characters in phylogenetics. Advances in Applied Mathematics, 2021, 131, 102252.	0.7	1
11	Triangulation supports agricultural spread of the Transeurasian languages. Nature, 2021, 599, 616-621.	27.8	58
12	OBAMA: OBAMA for Bayesian amino-acid model averaging. PeerJ, 2020, 8, e9460.	2.0	11
13	Adaptive Metropolis-coupled MCMC for BEAST 2. PeerJ, 2020, 8, e9473.	2.0	28
14	Phylodynamic Model Adequacy Using Posterior Predictive Simulations. Systematic Biology, 2019, 68, 358-364.	5.6	25
15	Model Selection and Parameter Inference in Phylogenetics Using Nested Sampling. Systematic Biology, 2019, 68, 219-233.	5.6	138
16	Bayesian phylodynamics of avian influenza A virus H9N2 in Asia with time-dependent predictors of migration. PLoS Computational Biology, 2019, 15, e1007189.	3.2	22
17	BEAST 2.5: An advanced software platform for Bayesian evolutionary analysis. PLoS Computational Biology, 2019, 15, e1006650.	3.2	2,484
18	A Bayesian phylogenetic study of the Dravidian language family. Royal Society Open Science, 2018, 5, 171504.	2.4	54

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19	The origin and expansion of Pama–Nyungan languages across Australia. Nature Ecology and Evolution, 2018, 2, 741-749.	7.8	54
20	Bayesian Phylogenetic Estimation of Clade Ages Supports Trans-Atlantic Dispersal of Cichlid Fishes. Systematic Biology, 2017, 66, syw076.	5.6	86
21	StarBEAST2 Brings Faster Species Tree Inference and Accurate Estimates of Substitution Rates. Molecular Biology and Evolution, 2017, 34, 2101-2114.	8.9	371
22	bModelTest: Bayesian phylogenetic site model averaging and model comparison. BMC Evolutionary Biology, 2017, 17, 42.	3.2	609
23	Tracing hepatitis B virus (HBV) genotype B5 (formerly B6) evolutionary history in the circumpolar Arctic through phylogeographic modelling. PeerJ, 2017, 5, e3757.	2.0	29
24	Phylogeography by diffusion on a sphere: whole world phylogeography. PeerJ, 2016, 4, e2406.	2.0	37
25	BEAST 2: A Software Platform for Bayesian Evolutionary Analysis. PLoS Computational Biology, 2014, 10, e1003537.	3.2	5,301
26	Species Delimitation using Genome-Wide SNP Data. Systematic Biology, 2014, 63, 534-542.	5.6	390
27	Looking for trees in the forest: summary tree from posterior samples. BMC Evolutionary Biology, 2013, 13, 221.	3.2	131
28	Evolutionary Rates and Hbv: Issues of Rate Estimation with Bayesian Molecular Methods. Antiviral Therapy, 2013, 18, 497-504.	1.0	68
29	Mapping the Origins and Expansion of the Indo-European Language Family. Science, 2012, 337, 957-960.	12.6	549
30	Inferring Species Trees Directly from Biallelic Genetic Markers: Bypassing Gene Trees in a Full Coalescent Analysis. Molecular Biology and Evolution, 2012, 29, 1917-1932.	8.9	828
31	DensiTree: making sense of sets of phylogenetic trees. Bioinformatics, 2010, 26, 1372-1373.	4.1	565
32	Racing algorithms for conditional independence inference. International Journal of Approximate Reasoning, 2007, 45, 386-401.	3.3	12
33	A modified simulation scheme for inference in Bayesian networks. International Journal of Approximate Reasoning, 1996, 14, 55-80.	3.3	27