Ari Löytynoja

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/186297/publications.pdf

Version: 2024-02-01

36 papers 8,646 citations

331670 21 h-index 35 g-index

52 all docs 52 docs citations

times ranked

52

15484 citing authors

#	Article	IF	CITATIONS
1	Template switching in DNA replication can create and maintain RNA hairpins. Proceedings of the National Academy of Sciences of the United States of America, 2022, 119 , .	7.1	3
2	Genetic population structure constrains local adaptation in sticklebacks. Molecular Ecology, 2021, 30, 1946-1961.	3.9	33
3	Automated improvement of stickleback reference genome assemblies with ⟨scp⟩Lepâ€Anchor⟨/scp⟩ software. Molecular Ecology Resources, 2021, 21, 2166-2176.	4.8	21
4	Phylogeny-Aware Alignment with PRANK and PAGAN. Methods in Molecular Biology, 2021, 2231, 17-37.	0.9	15
5	Evolutionary Sequence Analysis and Visualization with Wasabi. Methods in Molecular Biology, 2021, 2231, 225-240.	0.9	O
6	An inducible genome editing system for plants. Nature Plants, 2020, 6, 766-772.	9.3	77
7	A high-quality assembly of the nine-spined stickleback (Pungitius pungitius) genome. Genome Biology and Evolution, 2019, 11, 3291-3308.	2.5	54
8	Mechanistic insights into the evolution of DUF26-containing proteins in land plants. Communications Biology, 2019, 2, 56.	4.4	75
9	Effects of marker type and filtering criteria on <i>Q</i> _{ST} StS	2.4	12
10	Metabarcoding Gastrointestinal Nematodes in Sympatric Endemic and Nonendemic Species in Ranomafana National Park, Madagascar. International Journal of Primatology, 2018, 39, 49-64.	1.9	6
11	Bracketing phenogenotypic limits of mammalian hybridization. Royal Society Open Science, 2018, 5, 180903.	2.4	24
12	Short template switch events explain mutation clusters in the human genome. Genome Research, 2017, 27, 1039-1049.	5.5	31
13	Wasabi: An Integrated Platform for Evolutionary Sequence Analysis and Data Visualization. Molecular Biology and Evolution, 2016, 33, 1126-1130.	8.9	55
14	A recurrent copy number variation of the NEB triplicate region: only revealed by the targeted nemaline myopathy CGH array. European Journal of Human Genetics, 2016, 24, 574-580.	2.8	32
15	Tracking year-to-year changes in intestinal nematode communities of rufous mouse lemurs (<i>Microcebus rufus</i>). Parasitology, 2015, 142, 1095-1107.	1.5	14
16	Simple chained guide trees give poorer multiple sequence alignments than inferred trees in simulation and phylogenetic benchmarks. Proceedings of the National Academy of Sciences of the United States of America, 2015, 112, E99-100.	7.1	18
17	Analysis of CACTA transposases reveals intron loss as major factor influencing their exon/intron structure in monocotyledonous and eudicotyledonous hosts. Mobile DNA, 2014, 5, 24.	3.6	10
18	\tilde{SA} @ance: reference-based phylogenetic analysis for 18S rRNA studies. BMC Evolutionary Biology, 2014, 14, 235.	3.2	8

#	Article	IF	CITATIONS
19	Cleavage of the Drosophila screw prodomain is critical for a dynamic BMP morphogen gradient in embryogenesis. Developmental Biology, 2014, 389, 149-159.	2.0	26
20	Phylogeny-aware alignment with PRANK. Methods in Molecular Biology, 2014, 1079, 155-170.	0.9	558
21	Accurate extension of multiple sequence alignments using a phylogeny-aware graph algorithm. Bioinformatics, 2012, 28, 1684-1691.	4.1	126
22	Alignment Methods: Strategies, Challenges, Benchmarking, and Comparative Overview. Methods in Molecular Biology, 2012, 855, 203-235.	0.9	17
23	Genome content of uncultivated marine <i>Roseobacters</i> in the surface ocean. Environmental Microbiology, 2012, 14, 41-51.	3.8	42
24	webPRANK: a phylogeny-aware multiple sequence aligner with interactive alignment browser. BMC Bioinformatics, 2010, 11, 579.	2.6	418
25	Uniting Alignments and Trees. Science, 2009, 324, 1528-1529.	12.6	24
26	Phylogeny-Aware Gap Placement Prevents Errors in Sequence Alignment and Evolutionary Analysis. Science, 2008, 320, 1632-1635.	12.6	737
27	Determination and validation of principal gene products. Bioinformatics, 2008, 24, 11-17.	4.1	23
28	A model of evolution and structure for multiple sequence alignment. Philosophical Transactions of the Royal Society B: Biological Sciences, 2008, 363, 3913-3919.	4.0	44
29	Analyses of deep mammalian sequence alignments and constraint predictions for 1% of the human genome. Genome Research, 2007, 17 , $760-774$.	5.5	184
30	Identification and analysis of functional elements in 1% of the human genome by the ENCODE pilot project. Nature, 2007, 447, 799-816.	27.8	4,709
31	MATLIGN: a motif clustering, comparison and matching tool. BMC Bioinformatics, 2007, 8, 189.	2.6	21
32	From The Cover: An algorithm for progressive multiple alignment of sequences with insertions. Proceedings of the National Academy of Sciences of the United States of America, 2005, 102, 10557-10562.	7.1	871
33	A hidden Markov model for progressive multiple alignment. Bioinformatics, 2003, 19, 1505-1513.	4.1	157
34	Matrilinear phylogeography of Atlantic salmon (Salmo salar L.) in Europe and postglacial colonization of the Baltic Sea area. Molecular Ecology, 2001, 10, 89-102.	3.9	145
35	Molecular phylogenetic analyses of the mitochondrial ADP-ATP carriers: The Plantae/Fungi/Metazoa trichotomy revisited. Proceedings of the National Academy of Sciences of the United States of America, 2001, 98, 10202-10207.	7.1	31
36	Thousands of human mutation clusters are explained by short-range template switching. Genome Research, 0, , gr.276478.121.	5 . 5	3

3