Lars Arvestad

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	The Mitogenome of Norway Spruce and a Reappraisal of Mitochondrial Recombination in Plants. Genome Biology and Evolution, 2020, 12, 3586-3598.	2.5	35
2	Diversity and evolution of chitin synthases in oomycetes (Straminipila: Oomycota). Molecular Phylogenetics and Evolution, 2019, 139, 106558.	2.7	14
3	RecPhyloXML: a format for reconciled gene trees. Bioinformatics, 2018, 34, 3646-3652.	4.1	22
4	alv: a console-based viewer for molecular sequence alignments. Journal of Open Source Software, 2018, 3, 955.	4.6	3
5	VMCMC: a graphical and statistical analysis tool for Markov chain Monte Carlo traces. BMC Bioinformatics, 2017, 18, 97.	2.6	20
6	Structural Variation Detection with Read Pair Information: An Improved Null Hypothesis Reduces Bias. Journal of Computational Biology, 2017, 24, 581-589.	1.6	3
7	GenFamClust: an accurate, synteny-aware and reliable homology inference algorithm. BMC Evolutionary Biology, 2016, 16, 120.	3.2	14
8	Probabilistic inference of lateral gene transfer events. BMC Bioinformatics, 2016, 17, 431.	2.6	5
9	Assembly scaffolding with PE-contaminated mate-pair libraries. Bioinformatics, 2016, 32, 1925-1932.	4.1	42
10	Structural Variation Detection with Read Pair Information—An Improved Null-Hypothesis Reduces Bias. Lecture Notes in Computer Science, 2016, , 176-188.	1.3	0
11	Gene-pseudogene evolution: a probabilistic approach. BMC Genomics, 2015, 16, S12.	2.8	6
12	BESST - Efficient scaffolding of large fragmented assemblies. BMC Bioinformatics, 2014, 15, 281.	2.6	157
13	A Bayesian Method for Analyzing Lateral Gene Transfer. Systematic Biology, 2014, 63, 409-420.	5.6	73
14	Forensic Informativity of ~3000Âbp of Coding Sequence of Domestic Dog mt <scp>DNA</scp> . Journal of Forensic Sciences, 2014, 59, 898-908.	1.6	4
15	GAM-NGS: genomic assemblies merger for next generation sequencing. BMC Bioinformatics, 2013, 14, S6.	2.6	76
16	GenPhyloData: realistic simulation of gene family evolution. BMC Bioinformatics, 2013, 14, 209.	2.6	39
17	Quantitative synteny scoring improves homology inference and partitioning of gene families. BMC Bioinformatics, 2013, 14, S12.	2.6	32
18	Fastphylo: Fast tools for phylogenetics. BMC Bioinformatics, 2013, 14, 334.	2.6	18

LARS ARVESTAD

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19	The Norway spruce genome sequence and conifer genome evolution. Nature, 2013, 497, 579-584.	27.8	1,303
20	Excap: Maximization of Haplotypic Diversity of Linked Markers. PLoS ONE, 2013, 8, e79012.	2.5	0
21	DLRS: gene tree evolution in light of a species tree. Bioinformatics, 2012, 28, 2994-2995.	4.1	36
22	Improved gap size estimation for scaffolding algorithms. Bioinformatics, 2012, 28, 2215-2222.	4.1	23
23	Evolution and human tissue expression of the <i>Cres/Testatin</i> subgroup genes, a reproductive tissue specific subgroup of the type 2 cystatins. Evolution & Development, 2010, 12, 329-342.	2.0	6
24	Classification of DNA sequences using Bloom filters. Bioinformatics, 2010, 26, 1595-1600.	4.1	57
25	The gene evolution model and computing its associated probabilities. Journal of the ACM, 2009, 56, 1-44.	2.2	66
26	Simultaneous Bayesian gene tree reconstruction and reconciliation analysis. Proceedings of the National Academy of Sciences of the United States of America, 2009, 106, 5714-5719.	7.1	140
27	Identification of the cellulose synthase genes from the Oomycete Saprolegnia monoica and effect of cellulose synthesis inhibitors on gene expression and enzyme activity. Fungal Genetics and Biology, 2009, 46, 759-767.	2.1	27
28	MAP20, a Microtubule-Associated Protein in the Secondary Cell Walls of Hybrid Aspen, Is a Target of the Cellulose Synthesis Inhibitor 2,6-Dichlorobenzonitrile A. Plant Physiology, 2008, 148, 1283-1294.	4.8	76
29	Evolution of a domain conserved in microtubule-associated proteins of eukaryotes. Advances and Applications in Bioinformatics and Chemistry, 2008, 1, 51.	2.6	6
30	Evolution after gene duplication: models, mechanisms, sequences, systems, and organisms. Journal of Experimental Zoology Part B: Molecular and Developmental Evolution, 2007, 308B, 58-73.	1.3	148
31	primetv: a viewer for reconciled trees. BMC Bioinformatics, 2007, 8, 148.	2.6	20
32	Efficient Methods for Estimating Amino Acid Replacement Rates. Journal of Molecular Evolution, 2006, 62, 663-673.	1.8	13
33	Genome-Wide Survey for Biologically Functional Pseudogenes. PLoS Computational Biology, 2006, 2, e46.	3.2	75
34	Expressed sequence tags from the midgut and an epithelial cell line of Chironomus tentans: annotation, bioinformatic classification of unknown transcripts and analysis of expression levels. Insect Molecular Biology, 2005, 14, 689-695.	2.0	9
35	The genome sequence of black cottonwood (Populus trichocarpa) reveals 18 conserved cellulose synthase (CesA) genes. Planta, 2005, 221, 739-746.	3.2	121
36	Assessment of Protein Distance Measures and Tree-Building Methods for Phylogenetic Tree Reconstruction. Molecular Biology and Evolution, 2005, 22, 2257-2264.	8.9	16

LARS ARVESTAD

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37	ESTs from brain and testis of White Leghorn and red junglefowl: annotation, bioinformatic classification of unknown transcripts and analysis of expression levels. Cytogenetic and Genome Research, 2005, 111, 79-87.	1.1	21
38	Gene tree reconstruction and orthology analysis based on an integrated model for duplications and sequence evolution. , 2004, , .		82
39	Bayesian gene/species tree reconciliation and orthology analysis using MCMC. Bioinformatics, 2003, 19, i7-i15.	4.1	137
40	mtDNA Tandem Repeats in Domestic Dogs and Wolves: Mutation Mechanism Studied by Analysis of the Sequence of Imperfect Repeats. Molecular Biology and Evolution, 2000, 17, 474-488.	8.9	42
41	A Novel Method for Forensic DNA Investigations: Repeat- Type Sequence Analysis of Tandemly Repeated mtDNA in Domestic Dogs. Journal of Forensic Sciences, 2000, 45, 990-999.	1.6	21
42	Estimation of Reversible Substitution Matrices from Multiple Pairs of Sequences. Journal of Molecular Evolution, 1997, 45, 696-703.	1.8	18
43	Aligning coding DNA in the presence of frame-shift errors. Lecture Notes in Computer Science, 1997, , 180-190.	1.3	4