Erik Scott Wright

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/3093119/publications.pdf

Version: 2024-02-01

23 papers 1,076 citations

933447 10 h-index 19 g-index

26 all docs

26 docs citations

times ranked

26

1972 citing authors

#	Article	IF	Citations
1	FindNonCoding: rapid and simple detection of non-coding RNAs in genomes. Bioinformatics, 2022, 38, 841-843.	4.1	O
2	Sample-efficient identification of high-dimensional antibiotic synergy with a normalized diagonal sampling design. PLoS Computational Biology, 2022, 18, e1010311.	3.2	0
3	Ten simple rules for hitting a home run with your elevator pitch. PLoS Computational Biology, 2021, 17, e1008756.	3.2	6
4	Multi-stable bacterial communities exhibit extreme sensitivity to initial conditions. FEMS Microbiology Ecology, 2021, 97, .	2.7	10
5	Accurate annotation of protein coding sequences with IDTAXA. NAR Genomics and Bioinformatics, 2021, 3, lqab080.	3.2	O
6	SubTap, a Versatile 3D Printed Platform for Eavesdropping on Extracellular Interactions. MSystems, 2021, 6, e0090221.	3.8	1
7	Parallel evolution between genomic segments of seasonal human influenza viruses reveals RNA-RNA relationships. ELife, 2021, 10, .	6.0	11
8	Identification of antibiotic pairs that evade concurrent resistance via a retrospective analysis of antimicrobial susceptibility test results. Lancet Microbe, The, 2021, 2, e545-e554.	7.3	26
9	Identification of antibiotic pairs with disjoint resistance: innovative progress made but questions remain $\hat{a} \in \text{``Authors''}$ reply. Lancet Microbe, The, 2021, 2, e653.	7.3	0
10	Discordance Among Antibiotic Prescription Guidelines Reflects a Lack of Clear Best Practices. Open Forum Infectious Diseases, 2021, 8, ofaa571.	0.9	1
11	AssessORF: combining evolutionary conservation and proteomics to assess prokaryotic gene predictions. Bioinformatics, 2020, 36, 1022-1029.	4.1	10
12	RNAconTest: comparing tools for noncoding RNA multiple sequence alignment based on structural consistency. Rna, 2020, 26, 531-540.	3.5	37
13	Stochastic exits from dormancy give rise to heavyâ€tailed distributions of descendants in bacterial populations. Molecular Ecology, 2019, 28, 3915-3928.	3.9	21
14	Capturing the Complex Interplay Between Drugs and the Intestinal Microbiome. Clinical Pharmacology and Therapeutics, 2019, 106, 501-504.	4.7	5
15	Exclusivity offers a sound yet practical species criterion for bacteria despite abundant gene flow. BMC Genomics, 2018, 19, 724.	2.8	14
16	IDTAXA: a novel approach for accurate taxonomic classification of microbiome sequences. Microbiome, 2018, 6, 140.	11.1	336
17	Inhibitory interactions promote frequent bistability among competing bacteria. Nature Communications, 2016, 7, 11274.	12.8	82
18	Quality filtering of Illumina index reads mitigates sample cross-talk. BMC Genomics, 2016, 17, 876.	2.8	103

#	Article	IF	CITATIONS
19	DesignSignatures: a tool for designing primers that yields amplicons with distinct signatures. Bioinformatics, 2016, 32, 1565-1567.	4.1	10
20	DECIPHER: harnessing local sequence context to improve protein multiple sequence alignment. BMC Bioinformatics, 2015, 16, 322.	2.6	271
21	Exploiting extension bias in polymerase chain reaction to improve primer specificity in ensembles of nearly identical <scp>DNA</scp> templates. Environmental Microbiology, 2014, 16, 1354-1365.	3.8	72
22	Automated Design of Probes for rRNA-Targeted Fluorescence <i>In Situ</i> Hybridization Reveals the Advantages of Using Dual Probes for Accurate Identification. Applied and Environmental Microbiology, 2014, 80, 5124-5133.	3.1	39
23	Modeling Formamide Denaturation of Probe-Target Hybrids for Improved Microarray Probe Design in Microbial Diagnostics. PLoS ONE, 2012, 7, e43862.	2.5	16