Haoyu Cheng

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/6802277/publications.pdf

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	933410	1199563	
3,637	10	12	
citations	h-index	g-index	
18	18	1848	
docs citations	times ranked	citing authors	
		18 18	

#	Article	IF	CITATIONS
1	Haplotype-resolved de novo assembly using phased assembly graphs with hifiasm. Nature Methods, 2021, 18, 170-175.	19.0	1,675
2	The complete sequence of a human genome. Science, 2022, 376, 44-53.	12.6	1,222
3	Haplotype-resolved assembly of diploid genomes without parental data. Nature Biotechnology, 2022, 40, 1332-1335.	17.5	139
4	Chromosome-scale, haplotype-resolved assembly of human genomes. Nature Biotechnology, 2021, 39, 309-312.	17.5	109
5	Curated variation benchmarks for challenging medically relevant autosomal genes. Nature Biotechnology, 2022, 40, 672-680.	17.5	90
6	Metagenome assembly of high-fidelity long reads with hifiasm-meta. Nature Methods, 2022, 19, 671-674.	19.0	56
7	Real-time mapping of nanopore raw signals. Bioinformatics, 2021, 37, i477-i483.	4.1	41
8	Fast alignment and preprocessing of chromatin profiles with Chromap. Nature Communications, 2021, 12, 6566.	12.8	39
9	BitMapper: an efficient all-mapper based on bit-vector computing. BMC Bioinformatics, 2015, 16, 192.	2.6	26
10	LVMapper: A Large-Variance Clone Detector Using Sequencing Alignment Approach. IEEE Access, 2020, 8, 27986-27997.	4.2	14
11	FMtree: a fast locating algorithm of FM-indexes for genomic data. Bioinformatics, 2018, 34, 416-424.	4.1	7
12	BitMapper2: A GPU-Accelerated All-Mapper Based on the Sparseq-Gram Index. IEEE/ACM Transactions on Computational Biology and Bioinformatics, 2019, 16, 886-897.	3.0	7
13	An Efficient Filtration Method Based on Variable-Length Seeds for Sequence Alignment. Communications in Computer and Information Science, 2017, , 214-223.	0.5	O