## Chunlin Xiao

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/1774126/publications.pdf

Version: 2024-02-01

21 papers

15,676 citations

12 h-index

759233

18 g-index

28 all docs

28 docs citations

28 times ranked

15624 citing authors

#	Article	IF	CITATIONS
1	The Sequence of the Human Genome. Science, 2001, 291, 1304-1351.	12.6	12,623
2	The complete sequence of a human genome. Science, 2022, 376, 44-53.	12.6	1,222
3	Extensive sequencing of seven human genomes to characterize benchmark reference materials. Scientific Data, 2016, 3, 160025.	5.3	575
4	An open resource for accurately benchmarking small variant and reference calls. Nature Biotechnology, 2019, 37, 561-566.	17.5	277
5	A robust benchmark for detection of germline large deletions and insertions. Nature Biotechnology, 2020, 38, 1347-1355.	17.5	233
6	A complete reference genome improves analysis of human genetic variation. Science, 2022, 376, eabl3533.	12.6	144
7	Curated variation benchmarks for challenging medically relevant autosomal genes. Nature Biotechnology, 2022, 40, 672-680.	17.5	90
8	Benchmarking challenging small variants with linked and long reads. Cell Genomics, 2022, 2, 100128.	6.5	77
9	Toward best practice in cancer mutation detection with whole-genome and whole-exome sequencing. Nature Biotechnology, 2021, 39, 1141-1150.	17.5	66
10	A diploid assembly-based benchmark for variants in the major histocompatibility complex. Nature Communications, 2020, 11, 4794.	12.8	56
11	Establishing community reference samples, data and call sets for benchmarking cancer mutation detection using whole-genome sequencing. Nature Biotechnology, 2021, 39, 1151-1160.	17.5	39
12	Cross-oncopanel study reveals high sensitivity and accuracy with overall analytical performance depending on genomic regions. Genome Biology, 2021, 22, 109.	8.8	20
13	Assessing reproducibility of inherited variants detected with short-read whole genome sequencing. Genome Biology, 2022, 23, 2.	8.8	18
14	Whole genome and exome sequencing reference datasets from a multi-center and cross-platform benchmark study. Scientific Data, 2021, 8, 296.	5.3	15
15	High-coverage, long-read sequencing of Han Chinese trio reference samples. Scientific Data, 2019, 6, 91.	5.3	13
16	Achieving robust somatic mutation detection with deep learning models derived from reference data sets of a cancer sample. Genome Biology, 2022, 23, 12.	8.8	11
17	A crowdsourced set of curated structural variants for the human genome. PLoS Computational Biology, 2020, 16, e1007933.	3.2	6
18	A crowdsourced set of curated structural variants for the human genome. , 2020, 16, e1007933.		0

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#	Article	IF	CITATIONS
19	A crowdsourced set of curated structural variants for the human genome. , 2020, 16, e1007933.		O
20	A crowdsourced set of curated structural variants for the human genome. , 2020, 16, e1007933.		O
21	A crowdsourced set of curated structural variants for the human genome. , 2020, 16, e1007933.		O