## Verena Heinrich

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/2286754/publications.pdf

Version: 2024-02-01

21 papers 3,431 citations

471509 17 h-index 752698 20 g-index

24 all docs

24 docs citations

times ranked

24

5185 citing authors

#	Article	IF	CITATIONS
1	Non-coding deletions identify Maenli lncRNA as a limb-specific En1 regulator. Nature, 2021, 592, 93-98.	27.8	53
2	Position effects at the FGF8 locus are associated with femoral hypoplasia. American Journal of Human Genetics, 2021, 108, 1725-1734.	6.2	4
3	The mole genome reveals regulatory rearrangements associated with adaptive intersexuality. Science, 2020, 370, 208-214.	12.6	41
4	Enhancer hijacking determines extrachromosomal circular MYCN amplicon architecture in neuroblastoma. Nature Communications, 2020, 11, 5823.	12.8	104
5	Hi-C Identifies Complex Genomic Rearrangements and TAD-Shuffling in Developmental Diseases. American Journal of Human Genetics, 2020, 106, 872-884.	6.2	85
6	CRUP: a comprehensive framework to predict condition-specific regulatory units. Genome Biology, 2019, 20, 227.	8.8	26
7	Preformed chromatin topology assists transcriptional robustness of <i>Shh</i> during limb development. Proceedings of the National Academy of Sciences of the United States of America, 2019, 116, 12390-12399.	7.1	131
8	Serial genomic inversions induce tissue-specific architectural stripes, gene misexpression and congenital malformations. Nature Cell Biology, 2019, 21, 305-310.	10.3	107
9	Dynamic 3D chromatin architecture contributes to enhancer specificity and limb morphogenesis. Nature Genetics, 2018, 50, 1463-1473.	21.4	147
10	Characterization of hundreds of regulatory landscapes in developing limbs reveals two regimes of chromatin folding. Genome Research, 2017, 27, 223-233.	5.5	123
11	A likelihood ratio-based method to predict exact pedigrees for complex families from next-generation sequencing data. Bioinformatics, 2017, 33, 72-78.	4.1	8
12	Composition and dosage of a multipartite enhancer cluster control developmental expression of Ihh (Indian hedgehog). Nature Genetics, 2017, 49, 1539-1545.	21.4	107
13	Formation of new chromatin domains determines pathogenicity of genomic duplications. Nature, 2016, 538, 265-269.	27.8	582
14	Somatic neurofibromatosis type 1 (NF1) inactivation events in cutaneous neurofibromas of a single NF1 patient. European Journal of Human Genetics, 2015, 23, 870-873.	2.8	20
15	Disruptions of Topological Chromatin Domains Cause Pathogenic Rewiring of Gene-Enhancer Interactions. Cell, 2015, 161, 1012-1025.	28.9	1,725
16	Strategies to improve the performance of rare variant association studies by optimizing the selection of controls. Bioinformatics, 2015, 31, btv457.	4.1	0
17	Screening for single nucleotide variants, small indels and exon deletions with a nextâ€generation sequencing based gene panel approach for ⟨scp⟩U⟨/scp⟩ sher syndrome. Molecular Genetics & amp; Genomic Medicine, 2014, 2, 393-401.	1.2	22
18	Homozygous and Compound-Heterozygous Mutations in TGDS Cause Catel-Manzke Syndrome. American Journal of Human Genetics, 2014, 95, 763-770.	6.2	37

#	Article	IF	CITATIONS
19	Estimating exome genotyping accuracy by comparing to data from large scale sequencing projects. Genome Medicine, 2013, 5, 69.	8.2	23
20	Filtering for Compound Heterozygous Sequence Variants in Non-Consanguineous Pedigrees. PLoS ONE, 2013, 8, e70151.	2.5	41
21	The allele distribution in next-generation sequencing data sets is accurately described as the result of a stochastic branching process. Nucleic Acids Research, 2012, 40, 2426-2431.	14.5	40