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List of Publications by Year in descending order

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567281 888059 4,526 17 15 17 h-index citations g-index papers 19 19 19 6186 docs citations times ranked citing authors all docs

#	Article	IF	CITATIONS
1	Nuclear organization of active and inactive chromatin domains uncovered by chromosome conformation capture–on-chip (4C). Nature Genetics, 2006, 38, 1348-1354.	21.4	1,219
2	Looping and Interaction between Hypersensitive Sites in the Active \hat{l}^2 -globin Locus. Molecular Cell, 2002, 10, 1453-1465.	9.7	1,205
3	A Single Oncogenic Enhancer Rearrangement Causes Concomitant EVI1 and GATA2 Deregulation in Leukemia. Cell, 2014, 157, 369-381.	28.9	571
4	The second decade of 3C technologies: detailed insights into nuclear organization. Genes and Development, 2016, 30, 1357-1382.	5.9	320
5	Regulation of disease-associated gene expression in the 3D genome. Nature Reviews Molecular Cell Biology, 2016, 17, 771-782.	37.0	294
6	4C Technology: Protocols and Data Analysis. Methods in Enzymology, 2012, 513, 89-112.	1.0	203
7	Determining long-range chromatin interactions for selected genomic sites using 4C-seq technology: From fixation to computation. Methods, 2012, 58, 221-230.	3.8	198
8	Enhancer hubs and loop collisions identified from single-allele topologies. Nature Genetics, 2018, 50, 1151-1160.	21.4	189
9	Epigenomic annotation of gene regulatory alterations during evolution of the primate brain. Nature Neuroscience, 2016, 19, 494-503.	14.8	113
10	Building regulatory landscapes reveals that an enhancer can recruit cohesin to create contact domains, engage CTCF sites and activate distant genes. Nature Structural and Molecular Biology, 2022, 29, 563-574.	8.2	49
11	Hominin-specific regulatory elements selectively emerged in oligodendrocytes and are disrupted in autism patients. Nature Communications, 2020, 11, 301.	12.8	37
12	Identification of context-specific gene regulatory networks with <tt>GEMULA</tt> â€"gene expression modeling using LAsso. Bioinformatics, 2012, 28, 214-221.	4.1	35
13	Interplay between CTCF boundaries and a super enhancer controls cohesin extrusion trajectories and gene expression. Molecular Cell, 2021, 81, 3082-3095.e6.	9.7	29
14	Multi-contact 4C: long-molecule sequencing of complex proximity ligation products to uncover local cooperative and competitive chromatin topologies. Nature Protocols, 2020, 15, 364-397.	12.0	25
15	LLM3D: a log-linear modeling-based method to predict functional gene regulatory interactions from genome-wide expression data. Nucleic Acids Research, 2011, 39, 5313-5327.	14.5	19
16	Recently Evolved Enhancers Emerge with High Interindividual Variability and Less Frequently Associate with Disease. Cell Reports, 2020, 31, 107799.	6.4	7
17	Novel orthogonal methods to uncover the complexity and diversity of nuclear architecture. Current Opinion in Genetics and Development, 2021, 67, 10-17.	3.3	6