## Fiona Cunningham

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/5958482/publications.pdf

Version: 2024-02-01

49 papers

32,718 citations

39 h-index 50 g-index

56 all docs 56
docs citations

56 times ranked 62040 citing authors

#	Article	IF	CITATIONS
1	The Ensembl Variant Effect Predictor. Genome Biology, 2016, 17, 122.	3.8	5,181
2	The NHGRI-EBI GWAS Catalog of published genome-wide association studies, targeted arrays and summary statistics 2019. Nucleic Acids Research, 2019, 47, D1005-D1012.	6.5	3,179
3	Ensembl 2018. Nucleic Acids Research, 2018, 46, D754-D761.	6.5	2,710
4	GENCODE reference annotation for the human and mouse genomes. Nucleic Acids Research, 2019, 47, D766-D773.	6.5	2,350
5	The new NHGRI-EBI Catalog of published genome-wide association studies (GWAS Catalog). Nucleic Acids Research, 2017, 45, D896-D901.	6.5	1,932
6	Deriving the consequences of genomic variants with the Ensembl API and SNP Effect Predictor. Bioinformatics, 2010, 26, 2069-2070.	1.8	1,461
7	Ensembl 2016. Nucleic Acids Research, 2016, 44, D710-D716.	6.5	1,372
8	Ensembl 2021. Nucleic Acids Research, 2021, 49, D884-D891.	6.5	1,231
9	Ensembl 2014. Nucleic Acids Research, 2014, 42, D749-D755.	6.5	1,211
10	Ensembl 2015. Nucleic Acids Research, 2015, 43, D662-D669.	6.5	1,145
11	Ensembl 2022. Nucleic Acids Research, 2022, 50, D988-D995.	6.5	1,103
12	Ensembl 2020. Nucleic Acids Research, 2020, 48, D682-D688.	6.5	1,076
13	Ensembl 2019. Nucleic Acids Research, 2019, 47, D745-D751.	6.5	879
14	Ensembl 2013. Nucleic Acids Research, 2012, 41, D48-D55.	6.5	856
15	Ensembl 2012. Nucleic Acids Research, 2012, 40, D84-D90.	6.5	840
16	GENCODE 2021. Nucleic Acids Research, 2021, 49, D916-D923.	6.5	633
17	Ensembl 2011. Nucleic Acids Research, 2011, 39, D800-D806.	6.5	630
18	WormBase: a multi-species resource for nematode biology and genomics. Nucleic Acids Research, 2004, 32, 411D-417.	6.5	610

#	Article	IF	Citations
19	Ensembl 2017. Nucleic Acids Research, 2017, 45, D635-D642.	6.5	535
20	Modernizing Reference Genome Assemblies. PLoS Biology, 2011, 9, e1001091.	2.6	458
21	Ensembl variation resources. Database: the Journal of Biological Databases and Curation, 2018, 2018, .	1.4	377
22	Integrated allelic, transcriptional, and phenomic dissection of the cardiac effects of titin truncations in health and disease. Science Translational Medicine, 2015, 7, 270ra6.	5.8	375
23	Integrative Annotation of Variants from 1092 Humans: Application to Cancer Genomics. Science, 2013, 342, 1235587.	6.0	341
24	Whole-genome sequencing of patients with rare diseases in a national health system. Nature, 2020, 583, 96-102.	13.7	338
25	Ensembl's 10th year. Nucleic Acids Research, 2010, 38, D557-D562.	6.5	251
26	A joint NCBI and EMBL-EBI transcript set for clinical genomics and research. Nature, 2022, 604, 310-315.	13.7	162
27	A standardized framework for representation of ancestry data in genomics studies, with application to the NHGRI-EBI GWAS Catalog. Genome Biology, 2018, 19, 21.	3.8	159
28	WormBase: a comprehensive data resource for Caenorhabditis biology and genomics. Nucleic Acids Research, 2004, 33, D383-D389.	6.5	155
29	Ensembl variation resources. BMC Genomics, 2010, 11, 293.	1.2	124
30	WormBase: a cross-species database for comparative genomics. Nucleic Acids Research, 2003, 31, 133-137.	6.5	107
31	Locus Reference Genomic sequences: an improved basis for describing human DNA variants. Genome Medicine, 2010, 2, 24.	3.6	100
32	GA4GH: International policies and standards for data sharing across genomic research and healthcare. Cell Genomics, 2021, 1, 100029.	3.0	94
33	A Combined Functional Annotation Score for Non-Synonymous Variants. Human Heredity, 2012, 73, 47-51.	0.4	90
34	Flexible and scalable diagnostic filtering of genomic variants using G2P with Ensembl VEP. Nature Communications, 2019, 10, 2373.	5.8	86
35	A standard variation file format for human genome sequences. Genome Biology, 2010, 11, R88.	13.9	79
36	Locus Reference Genomic: reference sequences for the reporting of clinically relevant sequence variants. Nucleic Acids Research, 2014, 42, D873-D878.	<b>6.</b> 5	73

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37	The Gene Curation Coalition: A global effort to harmonize gene–disease evidence resources. Genetics in Medicine, 2022, 24, 1732-1742.	1.1	56
38	The European Variation Archive: a FAIR resource of genomic variation for all species. Nucleic Acids Research, 2022, 50, D1216-D1220.	6.5	50
39	IQSEC2-related encephalopathy in males and females: a comparative study including 37 novel patients. Genetics in Medicine, 2019, 21, 837-849.	1.1	47
40	Paralogous annotation of disease-causing variants in long QT syndrome genes. Human Mutation, 2012, 33, 1188-1191.	1.1	44
41	A database and API for variation, dense genotyping and resequencing data. BMC Bioinformatics, 2010, 11, 238.	1.2	33
42	Scaling up data curation using deep learning: An application to literature triage in genomic variation resources. PLoS Computational Biology, 2018, 14, e1006390.	1.5	33
43	Annotating and prioritizing genomic variants using the Ensembl Variant Effect Predictor—A tutorial. Human Mutation, 2022, 43, 986-997.	1.1	30
44	Improving the Sequence Ontology terminology for genomic variant annotation. Journal of Biomedical Semantics, 2015, 6, 32.	0.9	23
45	Haplosaurus computes protein haplotypes for use in precision drug design. Nature Communications, 2018, 9, 4128.	5.8	21
46	TranscriptSNPView: a genome-wide catalog of mouse coding variation. Nature Genetics, 2006, 38, 853-853.	9.4	14
47	DECIPHER: Supporting the interpretation and sharing of rare disease phenotypeâ€linked variant data to advance diagnosis and research. Human Mutation, 2022, , .	1.1	10
48	The value of primary transcripts to the clinical and nonâ€clinical genomics community: Survey results and roadmap for improvements. Molecular Genetics & Enomic Medicine, 2021, 9, e1786.	0.6	5
49	Annotation and curation of human genomic variations: an ELIXIR Implementation Study. F1000Research, 0, 9, 1207.	0.8	o