Helen Parkinson

List of Publications by Citations

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The third column is the impact factor (IF) of the journal, and the fourth column is the number of citations of the article.

16,667 48 129 121 h-index g-index citations papers 20,837 5.88 14.1 132 avg, IF L-index ext. papers ext. citations

#	Paper	IF	Citations
121	Minimum information about a microarray experiment (MIAME)-toward standards for microarray data. <i>Nature Genetics</i> , 2001 , 29, 365-71	36.3	3326
120	The NHGRI GWAS Catalog, a curated resource of SNP-trait associations. <i>Nucleic Acids Research</i> , 2014 , 42, D1001-6	20.1	2124
119	The NHGRI-EBI GWAS Catalog of published genome-wide association studies, targeted arrays and summary statistics 2019. <i>Nucleic Acids Research</i> , 2019 , 47, D1005-D1012	20.1	1422
118	The new NHGRI-EBI Catalog of published genome-wide association studies (GWAS Catalog). <i>Nucleic Acids Research</i> , 2017 , 45, D896-D901	20.1	1321
117	ArrayExpressa public repository for microarray gene expression data at the EBI. <i>Nucleic Acids Research</i> , 2003 , 31, 68-71	20.1	637
116	ArrayExpress updatesimplifying data submissions. <i>Nucleic Acids Research</i> , 2015 , 43, D1113-6	20.1	550
115	Disease Ontology 2015 update: an expanded and updated database of human diseases for linking biomedical knowledge through disease data. <i>Nucleic Acids Research</i> , 2015 , 43, D1071-8	20.1	403
114	ArrayExpress updatefrom an archive of functional genomics experiments to the atlas of gene expression. <i>Nucleic Acids Research</i> , 2009 , 37, D868-72	20.1	346
113	Modeling sample variables with an Experimental Factor Ontology. <i>Bioinformatics</i> , 2010 , 26, 1112-8	7.2	302
112	ArrayExpress updatetrends in database growth and links to data analysis tools. <i>Nucleic Acids Research</i> , 2013 , 41, D987-90	20.1	286
111	Natural variation in a Drosophila clock gene and temperature compensation. <i>Science</i> , 1997 , 278, 2117-2	2033.3	278
110	ArrayExpress updatean archive of microarray and high-throughput sequencing-based functional genomics experiments. <i>Nucleic Acids Research</i> , 2011 , 39, D1002-4	20.1	273
109	A global map of human gene expression. <i>Nature Biotechnology</i> , 2010 , 28, 322-4	44.5	271
108	Expression Atlas updatea database of gene and transcript expression from microarray- and sequencing-based functional genomics experiments. <i>Nucleic Acids Research</i> , 2014 , 42, D926-32	20.1	247
107	Open Targets: a platform for therapeutic target identification and validation. <i>Nucleic Acids Research</i> , 2017 , 45, D985-D994	20.1	241
106	The International Mouse Phenotyping Consortium Web Portal, a unified point of access for knockout mice and related phenotyping data. <i>Nucleic Acids Research</i> , 2014 , 42, D802-9	20.1	189
105	Modeling biomedical experimental processes with OBI. <i>Journal of Biomedical Semantics</i> , 2010 , 1 Suppl 1, S7	2.2	187

(2010-2010)

104	Gene expression atlas at the European bioinformatics institute. Nucleic Acids Research, 2010, 38, D690-	820.1	167
103	The EBI RDF platform: linked open data for the life sciences. <i>Bioinformatics</i> , 2014 , 30, 1338-9	7.2	160
102	The MGED Ontology: a resource for semantics-based description of microarray experiments. <i>Bioinformatics</i> , 2006 , 22, 866-73	7.2	152
101	The Human Phenotype Ontology: Semantic Unification of Common and Rare Disease. <i>American Journal of Human Genetics</i> , 2015 , 97, 111-24	11	147
100	Disease model discovery from 3,328 gene knockouts by The International Mouse Phenotyping Consortium. <i>Nature Genetics</i> , 2017 , 49, 1231-1238	36.3	145
99	Finding our way through phenotypes. <i>PLoS Biology</i> , 2015 , 13, e1002033	9.7	144
98	The Ontology for Biomedical Investigations. <i>PLoS ONE</i> , 2016 , 11, e0154556	3.7	143
97	Gene Expression Atlas updatea value-added database of microarray and sequencing-based functional genomics experiments. <i>Nucleic Acids Research</i> , 2012 , 40, D1077-81	20.1	133
96	Prevalence of sexual dimorphism in mammalian phenotypic traits. <i>Nature Communications</i> , 2017 , 8, 154	17 5 7.4	130
95	Standards for microarray data. <i>Science</i> , 2002 , 298, 539	33.3	120
94	Gramene 2016: comparative plant genomics and pathway resources. <i>Nucleic Acids Research</i> , 2016 , 44, D1133-40	20.1	102
93	Minimum information specification for in situ hybridization and immunohistochemistry experiments (MISFISHIE). <i>Nature Biotechnology</i> , 2008 , 26, 305-12	44.5	97
92	A standardized framework for representation of ancestry data in genomics studies, with application to the NHGRI-EBI GWAS Catalog. <i>Genome Biology</i> , 2018 , 19, 21	18.3	87
91	Submission of microarray data to public repositories. <i>PLoS Biology</i> , 2004 , 2, E317	9.7	87
90	Large scale comparison of global gene expression patterns in human and mouse. <i>Genome Biology</i> , 2010 , 11, R124	18.3	85
89	Tools and data services registry: a community effort to document bioinformatics resources. <i>Nucleic Acids Research</i> , 2016 , 44, D38-47	20.1	81
88	The EMBL nucleotide sequence database. <i>Nucleic Acids Research</i> , 2001 , 29, 17-21	20.1	77
87	The MOLGENIS toolkit: rapid prototyping of biosoftware at the push of a button. <i>BMC Bioinformatics</i> , 2010 , 11 Suppl 12, S12	3.6	76

86	CLO: The cell line ontology. <i>Journal of Biomedical Semantics</i> , 2014 , 5, 37	2.2	70
85	Improving reporting standards for polygenic scores in risk prediction studies. <i>Nature</i> , 2021 , 591, 211-2	19 ₅ 0.4	70
84	Importing ArrayExpress datasets into R/Bioconductor. <i>Bioinformatics</i> , 2009 , 25, 2092-4	7.2	68
83	PDX-MI: Minimal Information for Patient-Derived Tumor Xenograft Models. <i>Cancer Research</i> , 2017 , 77, e62-e66	10.1	65
82	The human-induced pluripotent stem cell initiative-data resources for cellular genetics. <i>Nucleic Acids Research</i> , 2017 , 45, D691-D697	20.1	63
81	Identifiers for the 21st century: How to design, provision, and reuse persistent identifiers to maximize utility and impact of life science data. <i>PLoS Biology</i> , 2017 , 15, e2001414	9.7	63
80	ArrayExpress: a public database of gene expression data at EBI. <i>Comptes Rendus - Biologies</i> , 2003 , 326, 1075-8	1.4	58
79	The Polygenic Score Catalog as an open database for reproducibility and systematic evaluation. <i>Nature Genetics</i> , 2021 , 53, 420-425	36.3	58
78	Applying the ARRIVE Guidelines to an In Vivo Database. <i>PLoS Biology</i> , 2015 , 13, e1002151	9.7	56
77	LifeTime and improving European healthcare through cell-based interceptive medicine. <i>Nature</i> , 2020 , 587, 377-386	50.4	56
76	Data standards for Omics data: the basis of data sharing and reuse. <i>Methods in Molecular Biology</i> , 2011 , 719, 31-69	1.4	52
75	Open Targets Genetics: systematic identification of trait-associated genes using large-scale genetics and functional genomics. <i>Nucleic Acids Research</i> , 2021 , 49, D1311-D1320	20.1	49
74	The BioSample Database (BioSD) at the European Bioinformatics Institute. <i>Nucleic Acids Research</i> , 2012 , 40, D64-70	20.1	47
73	Rapid establishment of the European Bank for induced Pluripotent Stem Cells (EBiSC) - the Hot Start experience. <i>Stem Cell Research</i> , 2017 , 20, 105-114	1.6	45
72	Development of FuGO: an ontology for functional genomics investigations. <i>OMICS A Journal of Integrative Biology</i> , 2006 , 10, 199-204	3.8	45
71	PDX Finder: A portal for patient-derived tumor xenograft model discovery. <i>Nucleic Acids Research</i> , 2019 , 47, D1073-D1079	20.1	42
70	Mouse Phenotype Database Integration Consortium: integration [corrected] of mouse phenome data resources. <i>Mammalian Genome</i> , 2007 , 18, 157-63	3.2	41
69	A Standard Nomenclature for Referencing and Authentication of Pluripotent Stem Cells. <i>Stem Cell Reports</i> , 2018 , 10, 1-6	8	39

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68	The Software Ontology (SWO): a resource for reproducibility in biomedical data analysis, curation and digital preservation. <i>Journal of Biomedical Semantics</i> , 2014 , 5, 25	2.2	39	
67	The European Bioinformatics Instituteß data resources. <i>Nucleic Acids Research</i> , 2003 , 31, 43-50	20.1	38	
66	Leveraging European infrastructures to access 1 million human genomes by 2022. <i>Nature Reviews Genetics</i> , 2019 , 20, 693-701	30.1	36	
65	The MGED ontology: a framework for describing functional genomics experiments. <i>Comparative and Functional Genomics</i> , 2003 , 4, 127-32		36	
64	The ArrayExpress gene expression database: a software engineering and implementation perspective. <i>Bioinformatics</i> , 2005 , 21, 1495-501	7.2	33	
63	Data storage and analysis in ArrayExpress. <i>Methods in Enzymology</i> , 2006 , 411, 370-86	1.7	33	
62	eQTL Catalogue: a compendium of uniformly processed human gene expression and splicing QTLs		33	
61	Accessing data from the International Mouse Phenotyping Consortium: state of the art and future plans. <i>Mammalian Genome</i> , 2012 , 23, 641-52	3.2	32	
60	Updates to BioSamples database at European Bioinformatics Institute. <i>Nucleic Acids Research</i> , 2014 , 42, D50-2	20.1	30	
59	A compendium of uniformly processed human gene expression and splicing quantitative trait loci. <i>Nature Genetics</i> , 2021 , 53, 1290-1299	36.3	28	
58	The underlying principles of scientific publication. <i>Bioinformatics</i> , 2002 , 18, 1409	7.2	27	
57	BioSamples database: an updated sample metadata hub. <i>Nucleic Acids Research</i> , 2019 , 47, D1172-D117	820.1	27	
56	Linking rare and common disease: mapping clinical disease-phenotypes to ontologies in therapeutic target validation. <i>Journal of Biomedical Semantics</i> , 2016 , 7, 8	2.2	26	
55	Toward richer metadata for microbial sequences: replacing strain-level NCBI taxonomy taxids with BioProject, BioSample and Assembly records. <i>Standards in Genomic Sciences</i> , 2014 , 9, 1275-7		26	
54	Meeting Report from the Second "Minimum Information for Biological and Biomedical Investigations" (MIBBI) workshop. <i>Standards in Genomic Sciences</i> , 2010 , 3, 259-66		26	
53	Human and mouse essentiality screens as a resource for disease gene discovery. <i>Nature Communications</i> , 2020 , 11, 655	17.4	25	
52	Identification of Cancer Related Genes Using a Comprehensive Map of Human Gene Expression. <i>PLoS ONE</i> , 2016 , 11, e0157484	3.7	24	
51	OntoCATsimple ontology search and integration in Java, R and REST/JavaScript. <i>BMC Bioinformatics</i> , 2011 , 12, 218	3.6	23	

50	A mouse informatics platform for phenotypic and translational discovery. <i>Mammalian Genome</i> , 2015 , 26, 413-21	3.2	20
49	ArrayExpress service for reviewers/editors of DNA microarray papers. <i>Nature Biotechnology</i> , 2006 , 24, 1321-2	44.5	20
48	Identification of genes required for eye development by high-throughput screening of mouse knockouts. <i>Communications Biology</i> , 2018 , 1, 236	6.7	20
47	Annotation of environmental OMICS data: application to the transcriptomics domain. <i>OMICS A Journal of Integrative Biology</i> , 2006 , 10, 172-8	3.8	18
46	The cellular microscopy phenotype ontology. <i>Journal of Biomedical Semantics</i> , 2016 , 7, 28	2.2	17
45	Harmonising phenomics information for a better interoperability in the rare disease field. <i>European Journal of Medical Genetics</i> , 2018 , 61, 706-714	2.6	16
44	Observ-OM and Observ-TAB: Universal syntax solutions for the integration, search, and exchange of phenotype and genotype information. <i>Human Mutation</i> , 2012 , 33, 867-73	4.7	16
43	REMBI: Recommended Metadata for Biological Images-enabling reuse of microscopy data in biology. <i>Nature Methods</i> , 2021 , 18, 1418-1422	21.6	16
42	Standards for microarray data: an open letter. Environmental Health Perspectives, 2004, 112, A666-7	8.4	13
41	PhenoImageShare: an image annotation and query infrastructure. <i>Journal of Biomedical Semantics</i> , 2016 , 7, 35	2.2	12
40	Annotarea tool for annotating high-throughput biomedical investigations and resulting data. <i>Bioinformatics</i> , 2010 , 26, 2470-1	7.2	12
39	MAGETabulator, a suite of tools to support the microarray data format MAGE-TAB. <i>Bioinformatics</i> , 2009 , 25, 279-80	7.2	12
38	Data storage and analysis in ArrayExpress and Expression Profiler. <i>Current Protocols in Bioinformatics</i> , 2008 , Chapter 7, Unit 7.13	24.2	12
37	The European Variation Archive: a FAIR resource of genomic variation for all species. <i>Nucleic Acids Research</i> , 2021 ,	20.1	10
36	Plant-based microarray data at the European Bioinformatics Institute. Introducing AtMIAMExpress, a submission tool for Arabidopsis gene expression data to ArrayExpress. <i>Plant Physiology</i> , 2005 , 139, 632-6	6.6	9
35	Contributions of the EMERALD project to assessing and improving microarray data quality. <i>BioTechniques</i> , 2011 , 50, 27-31	2.5	9
34	The SOFG Anatomy Entry List (SAEL): an annotation tool for functional genomics data. <i>Comparative and Functional Genomics</i> , 2004 , 5, 521-7		8
33	A guide to microarray experiments-an open letter to the scientific journals. <i>Lancet, The</i> , 2002 , 360, 101	940	8

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32	Mouse mutant phenotyping at scale reveals novel genes controlling bone mineral density. <i>PLoS Genetics</i> , 2020 , 16, e1009190	6	8
31	Precision medicine: Look to the mice. <i>Science</i> , 2015 , 349, 390	33.3	7
30	Reporting phenotypes in mouse models when considering body size as a potential confounder. <i>Journal of Biomedical Semantics</i> , 2016 , 7, 2	2.2	7
29	Gene Ontology Curation of Neuroinflammation Biology Improves the Interpretation of Alzheimerß Disease Gene Expression Data. <i>Journal of Alzheimerts Disease</i> , 2020 , 75, 1417-1435	4.3	6
28	Webulous and the Webulous Google Add-Ona web service and application for ontology building from templates. <i>Journal of Biomedical Semantics</i> , 2016 , 7, 17	2.2	6
27	Using ontologies to annotate microarray experiments. <i>Methods in Enzymology</i> , 2006 , 411, 325-39	1.7	6
26	Wrestling with SUMO and bio-ontologies. <i>Nature Biotechnology</i> , 2006 , 24, 21-2; author reply 23	44.5	6
25	Improving the Gene Ontology Resource to Facilitate More Informative Analysis and Interpretation of Alzheimerß Disease Data. <i>Genes</i> , 2018 , 9,	4.2	6
24	Anatomy ontologies and potential users: bridging the gap. <i>Journal of Biomedical Semantics</i> , 2011 , 2 Suppl 4, S3	2.2	5
23	The European Genome-phenome Archive in 2021. Nucleic Acids Research, 2021,	20.1	5
22	OpenStats: A robust and scalable software package for reproducible analysis of high-throughput phenotypic data. <i>PLoS ONE</i> , 2020 , 15, e0242933	3.7	5
21	Soft windowing application to improve analysis of high-throughput phenotyping data. <i>Bioinformatics</i> , 2020 , 36, 1492-1500	7.2	5
20	Standards and ontologies for functional genomics 2. Comparative and Functional Genomics, 2004, 5, 618	-22	4
19	Workshop proceedings: GWAS summary statistics standards and sharing. <i>Cell Genomics</i> , 2021 , 1, 100004		4
18	Improving reporting standards for polygenic scores in risk prediction studies		4
17	Desiderata for the development of next-generation electronic health record phenotype libraries. <i>GigaScience</i> , 2021 , 10,	7.6	4
16	Comparison, alignment, and synchronization of cell line information between CLO and EFO. <i>BMC Bioinformatics</i> , 2017 , 18, 557	3.6	3
15	Sequencing-based genome-wide association studies reporting standards. <i>Cell Genomics</i> , 2021 , 1, 10000	5-1000	005

14	Standards and ontologies for functional genomics: towards unified ontologies for biology and biomedicine. <i>Comparative and Functional Genomics</i> , 2003 , 4, 116-20		2
13	Pedro Ontology Services: A Framework for Rapid Ontology Markup. <i>Lecture Notes in Computer Science</i> , 2005 , 578-591	0.9	2
12	PDX Finder: A Portal for Patient-Derived tumor Xenograft Model Discovery		2
11	A standardized framework for representation of ancestry data in genomics studies, with application to the NHGRI-EBI GWAS Catalog		2
10	Extensive identification of genes involved in congenital and structural heart disorders and cardiomyopathy 2022 , 1, 157-173		2
9	MageCometweb application for harmonizing existing large-scale experiment descriptions. <i>Bioinformatics</i> , 2012 , 28, 1402-3	7.2	1
8	NCRI informatics initiative. <i>Nature Biotechnology</i> , 2005 , 23, 1212	44.5	1
7	OpenStats: A Robust and Scalable Software Package for Reproducible Analysis of High-Throughput Phenotypic Data		1
6	Analyzing the heterogeneity of rule-based EHR phenotyping algorithms in CALIBER and the UK Biobank	(1
5	Identifiers for the 21st century: How to design, provision, and reuse persistent identifiers to maximize utility and impact of life science data		1
4	The EurOPDX Data Portal: an open platform for patient-derived cancer xenograft data sharing and visualization <i>BMC Genomics</i> , 2022 , 23, 156	4.5	1
3	Identifying genetic determinants of inflammatory pain in mice using a large-scale gene-targeted screen <i>Pain</i> , 2022 , 163, 1139-1157	8	O
2	Pleiotropy data resource as a primer for investigating co-morbidities/multi-morbidities and their role in disease. <i>Mammalian Genome</i> , 2021 , 1	3.2	О
1	Standards for Functional Genomics 2009 , 293-329		