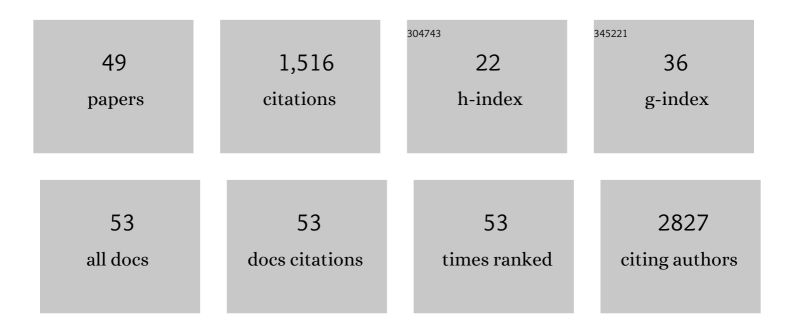
## Adrian Thorogood

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/37740/publications.pdf Version: 2024-02-01



#	Article	IF	CITATIONS
1	Isolated Aerobic Exercise and Weight Loss: A Systematic Review and Meta-Analysis of Randomized Controlled Trials. American Journal of Medicine, 2011, 124, 747-755.	1.5	192
2	BRCA Challenge: BRCA Exchange as a global resource for variants in BRCA1 and BRCA2. PLoS Genetics, 2018, 14, e1007752.	3.5	148
3	Reproducibility and Reuse of Adaptive Immune Receptor Repertoire Data. Frontiers in Immunology, 2017, 8, 1418.	4.8	102
4	GA4GH: International policies and standards for data sharing across genomic research and healthcare. Cell Genomics, 2021, 1, 100029.	6.5	94
5	Global Public Perceptions of Genomic Data Sharing: What Shapes the Willingness to Donate DNA and Health Data?. American Journal of Human Genetics, 2020, 107, 743-752.	6.2	76
6	Return of individual genomic research results: are laws and policies keeping step?. European Journal of Human Genetics, 2019, 27, 535-546.	2.8	73
7	Trust in genomic data sharing among members of the general public in the UK, USA, Canada and Australia. Human Genetics, 2019, 138, 1237-1246.	3.8	69
8	Consent recommendations for research and international data sharing involving persons with dementia. Alzheimer's and Dementia, 2018, 14, 1334-1343.	0.8	68
9	Attitudes of publics who are unwilling to donate DNA data for research. European Journal of Medical Genetics, 2019, 62, 316-323.	1.3	53
10	APPLaUD: access for patients and participants to individual level uninterpreted genomic data. Human Genomics, 2018, 12, 7.	2.9	45
11	An implementation framework for the feedback of individual research results and incidental findings in research. BMC Medical Ethics, 2014, 15, 88.	2.4	39
12	Demonstrating trustworthiness when collecting and sharing genomic data: public views across 22 countries. Genome Medicine, 2021, 13, 92.	8.2	39
13	Registered access: a â€ <sup></sup> Triple-A' approach. European Journal of Human Genetics, 2016, 24, 1676-1680.	2.8	33
14	Consent Processes for Mobile App Mediated Research: Systematic Review. JMIR MHealth and UHealth, 2017, 5, e126.	3.7	33
15	Genomics: data sharing needs an international code of conduct. Nature, 2020, 578, 31-33.	27.8	31
16	Toward better governance of human genomic data. Nature Genetics, 2021, 53, 2-8.	21.4	31
17	The Data Use Ontology to streamline responsible access to human biomedical datasets. Cell Genomics, 2021, 1, 100028.	6.5	31
18	Members of the public in the USA, UK, Canada and Australia expressing genetic exceptionalism say they are more willing to donate genomic data. European Journal of Human Genetics, 2020, 28, 424-434.	2.8	29

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19	Reporting results from whole-genome and whole-exome sequencing in clinical practice: a proposal for Canada?. Journal of Medical Genetics, 2014, 51, 68-70.	3.2	27
20	â€~Your DNA, Your Say': global survey gathering attitudes toward genomics: design, delivery and methods. Personalized Medicine, 2018, 15, 311-318.	1.5	26
21	International Guidelines for Privacy in Genomic Biobanking (or the Unexpected Virtue of Pluralism). Journal of Law, Medicine and Ethics, 2015, 43, 690-702.	0.9	25
22	A P3G generic access agreement for population genomic studies. Nature Biotechnology, 2013, 31, 384-385.	17.5	24
23	International federation of genomic medicine databases using GA4GH standards. Cell Genomics, 2021, 1, 100032.	6.5	22
24	Public variant databases: liability?. Genetics in Medicine, 2017, 19, 838-841.	2.4	21
25	A Tale of Two Capacities: Including Children and Decisionally Vulnerable Adults in Biomedical Research. Frontiers in Genetics, 2019, 10, 289.	2.3	16
26	COVID-19 Research: Navigating the European General Data Protection Regulation. Journal of Medical Internet Research, 2020, 22, e19799.	4.3	16
27	Who should have access to genomic data and how should they be held accountable? Perspectives of Data Access Committee members and experts. European Journal of Human Genetics, 2016, 24, 1671-1675.	2.8	15
28	Population Biobanking and International Collaboration. Pathobiology, 2014, 81, 276-285.	3.8	13
29	The Human Genome Organisation: towards next-generation ethics. Genome Medicine, 2013, 5, 38.	8.2	12
30	Bringing Code to Data: Do Not Forget Governance. Journal of Medical Internet Research, 2020, 22, e18087.	4.3	11
31	Canada: will privacy rules continue to favour open science?. Human Genetics, 2018, 137, 595-602.	3.8	10
32	Genetic database software as medical devices. Human Mutation, 2018, 39, 1702-1712.	2.5	10
33	Consent insufficient for data release. Science, 2019, 364, 445-446.	12.6	9
34	Letter: Relearning the 3 R's? Reinterpretation, recontact, and return of genetic variants. Genetics in Medicine, 2019, 21, 2401-2402.	2.4	9
35	Genomic Sequencing Capacity, Data Retention, and Personal Access to Raw Data in Europe. Frontiers in Genetics, 2020, 11, 303.	2.3	9
36	Essential Requirements for Establishing and Operating Data Trusts. International Journal of Population Data Science, 2020, 5, 1353.	0.1	9

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37	Return of genomic results does not motivate intentÂtoÂparticipate in research for all: Perspectives across 22 countries. Genetics in Medicine, 2022, 24, 1120-1129.	2.4	8
38	Open science precision medicine in Canada: Points to consider. Facets, 2019, 4, 1-19.	2.4	7
39	International Data Sharing and Rare Disease: The Importance of Ethics and Patient Involvement. , 0, , .		5
40	Substitute consent to data sharing: a way forward for international dementia research?. Journal of Law and the Biosciences, 2017, 4, lsw063.	1.6	4
41	Accelerating evidence gathering and approval of precision medicine therapies: the FDA takes aim at rare mutations. Genetics in Medicine, 2019, 21, 542-544.	2.4	4
42	Policy-aware data lakes: a flexible approach to achieve legal interoperability for global research collaborations. Journal of Law and the Biosciences, 2020, 7, Isaa065.	1.6	4
43	Openness, inclusion, and respect in dementia research. Lancet Neurology, The, 2019, 18, 135-136.	10.2	3
44	International Coordination of Research Ethics Review: An Adequacy Model. Philosophies, 2021, 6, 93.	0.7	3
45	Applying GDPR roles and responsibilities to scientific data sharing. International Data Privacy Law, 2022, 12, 207-219.	1.2	3
46	Streamlining ethics review for international health research. Science, 2022, 375, 825-826.	12.6	2
47	Oversight, governance, and policy for making decisions about return of individual genomic findings. , 2020, , 29-41.		1
48	Sharing Bioinformatic Data for Machine Learning: Maximizing Interoperability through License Selection. , 2020, , .		1
49	Point-of-Care Genetic Tests for Infectious Disease: Legal Considerations. Current Pharmacogenomics and Personalized Medicine, 2014, 12, 43-50.	0.2	1