Mahsa Shabani

List of Publications by Year in descending order

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430874 501196 38 982 18 28 citations h-index g-index papers 39 39 39 1164 docs citations times ranked citing authors all docs

#	Article	IF	CITATIONS
1	Rules for processing genetic data for research purposes in view of the new EU General Data Protection Regulation. European Journal of Human Genetics, 2018, 26, 149-156.	2.8	95
2	Attitudes of research participants and the general public towards genomic data sharing: a systematic literature review. Expert Review of Molecular Diagnostics, 2014, 14, 1053-1065.	3.1	82
3	Blockchain-based platforms for genomic data sharing: a de-centralized approach in response to the governance problems?. Journal of the American Medical Informatics Association: JAMIA, 2019, 26, 76-80.	4.4	69
4	From the principles of genomic data sharing to the practices of data access committees. EMBO Molecular Medicine, 2015, 7, 507-509.	6.9	51
5	Reâ€identifiability of genomic data and the <scp>GDPR</scp> . EMBO Reports, 2019, 20, .	4.5	49
6	The challenges of the expanded availability of genomic information: an agenda-setting paper. Journal of Community Genetics, 2018, 9, 103-116.	1.2	45
7	Ethics review for international data-intensive research. Science, 2016, 351, 1399-1400.	12.6	44
8	Controlled Access under Review: Improving the Governance of Genomic Data Access. PLoS Biology, 2015, 13, e1002339.	5.6	42
9	Ethics review of big data research: What should stay and what should be reformed?. BMC Medical Ethics, 2021, 22, 51.	2.4	39
10	DNA Data Marketplace: An Analysis of the Ethical Concerns Regarding the Participation of the Individuals. Frontiers in Genetics, 2019, 10, 1107.	2.3	37
11	Registered access: a â€~Triple-A' approach. European Journal of Human Genetics, 2016, 24, 1676-1680.	2.8	33
12	Registered access: authorizing data access. European Journal of Human Genetics, 2018, 26, 1721-1731.	2.8	33
13	"You want the right amount of oversight― interviews with data access committee members and experts on genomic data access. Genetics in Medicine, 2016, 18, 892-897.	2.4	32
14	Toward better governance of human genomic data. Nature Genetics, 2021, 53, 2-8.	21.4	31
15	Challenges of web-based personal genomic data sharing. Life Sciences, Society and Policy, 2015, 11, 3.	3.2	26
16	Oversight of Genomic Data Sharing: What Roles for Ethics and Data Access Committees?. Biopreservation and Biobanking, 2017, 15, 469-474.	1.0	25
17	Forensic Epigenetic Age Estimation and Beyond: Ethical and Legal Considerations. Trends in Genetics, 2018, 34, 489-491.	6.7	24
18	Will the European Health Data Space change data sharing rules?. Science, 2022, 375, 1357-1359.	12.6	24

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19	Review of policies of companies and databases regarding access to customers' genealogy data for law enforcement purposes. Personalized Medicine, 2020, 17, 141-153.	1.5	23
20	Data Sharing in Biomedical Sciences: A Systematic Review of Incentives. Biopreservation and Biobanking, 2021, 19, 219-227.	1.0	20
21	Raw Genomic Data: Storage, Access, and Sharing. Trends in Genetics, 2018, 34, 8-10.	6.7	18
22	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
23	Factors that influence data sharing through data sharing platforms: A qualitative study on the views and experiences of cohort holders and platform developers. PLoS ONE, 2021, 16, e0254202.	2.5	15
24	Research ethics review for the use of anonymized samples and data: A systematic review of normative documents. Accountability in Research, 2017, 24, 483-496.	2.4	12
25	Variant data sharing by clinical laboratories through public databases: consent, privacy and further contact for research policies. Genetics in Medicine, 2019, 21, 1031-1037.	2.4	12
26	The Data Governance Act and the EU's move towards facilitating data sharing. Molecular Systems Biology, 2021, 17, e10229.	7.2	12
27	Transparency and objectivity in governance of clinical trials data sharing: Current practices and approaches. Clinical Trials, 2019, 16, 547-551.	1.6	11
28	The Impact of the GDPR on the Governance of Biobank Research. Law, Governance and Technology Series, 2021, , 45-60.	0.4	11
29	Credit and Recognition for Contributions to Data-Sharing Platforms Among Cohort Holders and Platform Developers in Europe: Interview Study. Journal of Medical Internet Research, 2022, 24, e25983.	4.3	10
30	Consent insufficient for data release. Science, 2019, 364, 445-446.	12.6	9
31	Genomic Sequencing Capacity, Data Retention, and Personal Access to Raw Data in Europe. Frontiers in Genetics, 2020, 11, 303.	2.3	9
32	Data sharing platforms and the academic evaluation system. EMBO Reports, 2020, 21, e50690.	4.5	9
33	Data sharing platforms: instruments to inform and shape science policy on data sharing?. Scientometrics, 2022, 127, 3007-3019.	3.0	7
34	An agenda-setting paper on data sharing platforms: euCanSHare workshop. Open Research Europe, $0, 1, 80$.	2.0	3
35	Reporting, recording, and communication of COVID-19 cases in workplace: data protection as a moving target. Journal of Law and the Biosciences, 2020, 7, Isaa008.	1.6	2
36	The Use of Samples Originating From Doping Control Procedures for Research Purposes: A Qualitative Study. Journal of Empirical Research on Human Research Ethics, 2019, 14, 254-261.	1.3	1

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37	Editorial: Personal Genomes: Accessing, Sharing, and Interpretation. Frontiers in Genetics, 2021, 12, 687584.	2.3	1
38	An agenda-setting paper on data sharing platforms: euCanSHare workshop. Open Research Europe, 0, 1, 80.	2.0	1