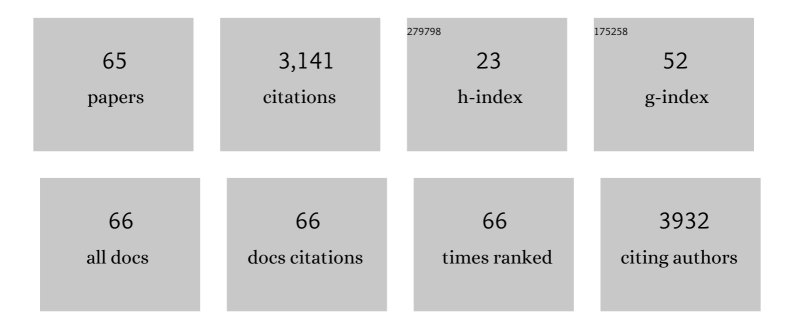
## Nanibaaâ€<sup>™</sup> Garrison

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/5376540/publications.pdf Version: 2024-02-01



#	Article	IF	CITATIONS
1	Return of Genomic Results to Research Participants: The Floor, the Ceiling, and the Choices In Between. American Journal of Human Genetics, 2014, 94, 818-826.	6.2	342
2	A framework for enhancing ethical genomic research with Indigenous communities. Nature Communications, 2018, 9, 2957.	12.8	274
3	Genomic Justice for Native Americans. Science Technology and Human Values, 2013, 38, 201-223.	3.1	213
4	The Human Pangenome Project: a global resource to map genomic diversity. Nature, 2022, 604, 437-446.	27.8	192
5	A systematic literature review of individuals' perspectives on broad consent and data sharing in the United States. Genetics in Medicine, 2016, 18, 663-671.	2.4	181
6	Public Attitudes toward Consent and Data Sharing in Biobank Research: A Large Multi-site Experimental Survey in the US. American Journal of Human Genetics, 2017, 100, 414-427.	6.2	172
7	Human Germline Genome Editing. American Journal of Human Genetics, 2017, 101, 167-176.	6.2	168
8	Rights, interests and expectations: Indigenous perspectives on unrestricted access to genomic data. Nature Reviews Genetics, 2020, 21, 377-384.	16.3	141
9	Genomic Research Through an Indigenous Lens: Understanding the Expectations. Annual Review of Genomics and Human Genetics, 2019, 20, 495-517.	6.2	129
10	Predicting Phenotype from Genotype: Normal Pigmentation*. Journal of Forensic Sciences, 2010, 55, 315-322.	1.6	110
11	Advancing the ethics of paleogenomics. Science, 2018, 360, 384-385.	12.6	110
12	COVID-19 Vaccine Decision-making Factors in Racial and Ethnic Minority Communities in Los Angeles, California. JAMA Network Open, 2021, 4, e2127582.	5.9	108
13	Genome-Wide Association Studies of Quantitatively Measured Skin, Hair, and Eye Pigmentation in Four European Populations. PLoS ONE, 2012, 7, e48294.	2.5	94
14	The Responsibility to Recontact Research Participants after Reinterpretation of Genetic and Genomic Research Results. American Journal of Human Genetics, 2019, 104, 578-595.	6.2	91
15	Family Ties: The Use of DNA Offender Databases to Catch Offenders' Kin. Journal of Law, Medicine and Ethics, 2006, 34, 248-262.	0.9	60
16	Fostering Responsible Research on Ancient DNA. American Journal of Human Genetics, 2020, 107, 183-195.	6.2	57
17	A 122.5-Kilobase Deletion of the P Gene Underlies the High Prevalence of Oculocutaneous Albinism Type 2 in the Navajo Population. American Journal of Human Genetics, 2003, 72, 62-72.	6.2	53
18	Genomic Contextualism: Shifting the Rhetoric of Genetic Exceptionalism. American Journal of Bioethics, 2019, 19, 51-63.	0.9	48

#	Article	IF	CITATIONS
19	Return of results in the genomic medicine projects of the eMERGE network. Frontiers in Genetics, 2014, 5, 50.	2.3	40
20	Creating a data resource: what will it take to build a medical information commons?. Genome Medicine, 2017, 9, 84.	8.2	36
21	Rationales for expanding minority physician representation in the workforce: a scoping review. Medical Education, 2018, 52, 925-935.	2.1	31
22	Access and Management: Indigenous Perspectives on Genomic Data Sharing. Ethnicity and Disease, 2019, 29, 659-668.	2.3	31
23	Awareness and Acceptable Practices: IRB and Researcher Reflections on the Havasupai Lawsuit. American Journal of Bioethics Primary Research, 2013, 4, 55-63.	1.5	29
24	Pathways from autism spectrum disorder diagnosis to genetic testing. Genetics in Medicine, 2018, 20, 737-744.	2.4	29
25	Generations of genomes: advances in paleogenomics technology and engagement for Indigenous people of the Americas. Current Opinion in Genetics and Development, 2020, 62, 91-96.	3.3	27
26	Parents' attitudes toward consent and data sharing in biobanks: A multisite experimental survey. AJOB Empirical Bioethics, 2018, 9, 128-142.	1.6	25
27	Chaco Canyon Dig Unearths Ethical Concerns. Human Biology, 2017, 89, 177.	0.2	25
28	Using Indigenous Standards to Implement the CARE Principles: Setting Expectations through Tribal Research Codes. Frontiers in Genetics, 2022, 13, 823309.	2.3	22
29	Importance of Participant-Centricity and Trust for a Sustainable Medical Information Commons. Journal of Law, Medicine and Ethics, 2019, 47, 12-20.	0.9	20
30	Trust, Precision Medicine Research, and Equitable Participation of Underserved Populations. American Journal of Bioethics, 2018, 18, 34-36.	0.9	19
31	When Participants in Genomic Research Grow Up: Contact and Consent atÂthe Age of Majority. Journal of Pediatrics, 2016, 168, 226-231.e1.	1.8	17
32	Attitudes of Members of Genetics Professional Societies Toward Human Gene Editing. CRISPR Journal, 2019, 2, 331-339.	2.9	15
33	A systematic literature review of Native American and Pacific Islanders' perspectives on health data privacy in the United States. Journal of the American Medical Informatics Association: JAMIA, 2020, 27, 1987-1998.	4.4	15
34	Impact of Psychiatric Information on Potential Jurors in Evaluating High-Functioning Autism Spectrum Disorder (hfASD). Journal of Mental Health Research in Intellectual Disabilities, 2015, 8, 140-167.	2.0	14
35	Stress across generations: A qualitative study of stress, coping, and caregiving among Mexican immigrant mothers Ethnicity and Health, 2019, 24, 378-394.	2.5	13
36	Ethical Guidance in Human Paleogenomics: New and Ongoing Perspectives. Annual Review of Genomics and Human Genetics, 2022, 23, 627-652.	6.2	13

Nanibaa' A Garrison

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37	ls there a way to reduce the inequity in variant interpretation on the basis of ancestry?. American Journal of Human Genetics, 2022, 109, 981-988.	6.2	13
38	Customers or research participants?: Guidance for research practices in commercialization of personal genomics. Genetics in Medicine, 2012, 14, 833-835.	2.4	12
39	Relationships Matter: Ethical Considerations for Returning Results to Family Members of Deceased Subjects. American Journal of Bioethics, 2013, 13, 66-67.	0.9	12
40	Agreed: The Harm Principle Cannot Replace the Best Interest Standard … but the Best Interest Standard Cannot Replace The Harm Principle Either. American Journal of Bioethics, 2018, 18, 38-40.	0.9	12
41	Carnitine palmitoyltransferase 1A P479L and infant death: policy implications of emerging data. Genetics in Medicine, 2017, 19, 851-857.	2.4	11
42	Complicated legacies: The human genome at 20. Science, 2021, 371, 564-569.	12.6	11
43	Considering "Respect for Sovereignty―Beyond the Belmont Report and the Common Rule: Ethical and Legal Implications for American Indian and Alaska Native Peoples. American Journal of Bioethics, 2021, 21, 27-30.	0.9	11
44	Weaving the Strands of Life ( <em>liná BitÅ,'ool</em> ): History of Genetic Research Involving Navajo People. Human Biology, 2019, 91, 189.	0.2	11
45	Ancient-DNA researchers write their own rules. Nature, 2021, 600, 37-37.	27.8	11
46	Forensic familial searching: scientific and social implications. Nature Reviews Genetics, 2013, 14, 445-445.	16.3	10
47	Entwined Processes: Rescripting Consent and Strengthening Governance in Genomics Research with Indigenous Communities. Journal of Law, Medicine and Ethics, 2020, 48, 218-220.	0.9	8
48	Perspectives on Genetic Research: Results From a Survey of Navajo Community Members. Frontiers in Genetics, 2021, 12, 734529.	2.3	8
49	"The System Doesn't Let Us inâ€â€"A Call for Inclusive COVID-19 Vaccine Outreach Rooted in Los Angeles Latinos' Experience of Pandemic Hardships and Inequities. International Journal of Environmental Research and Public Health, 2022, 19, 5785.	5 2.6	8
50	Meningococccal meningitis and complement component 6 deficiency associated with oculocutaneous albinism. European Journal of Pediatrics, 2005, 164, 177-179.	2.7	7
51	Naming Indigenous Concerns, Framing Considerations for Stored Biospecimens. American Journal of Bioethics, 2015, 15, 73-75.	0.9	7
52	Genetic Ancestry Testing with Tribes: Ethics, Identity & Health Implications. Daedalus, 2018, 147, 60-69.	1.8	5
53	Understanding as an Ethical Aspiration in an Era of Digital Technology-Based Communication: An Analysis of Informed Consent Functions. American Journal of Bioethics, 2019, 19, 34-36.	0.9	4
54	The Instrumental Role of Hospital Ethics Committees in Policy Work. American Journal of Bioethics, 2012, 12, 1-2.	0.9	3

Nanibaa' A Garrison

#	Article	IF	CITATIONS
55	Genomics in the clinic: ethical and policy challenges in clinical next-generation sequencing programs at early adopter USA institutions. Personalized Medicine, 2015, 12, 269-282.	1.5	3
56	Considerations for Returning Research Results to Culturally Diverse Participants and Families of Decedents. Journal of Law, Medicine and Ethics, 2015, 43, 569-575.	0.9	3
57	An Ethical Case for Dual-Role Consent: Increasing Research Diversity as a Matter of Respect and Justice. American Journal of Bioethics, 2019, 19, 44-46.	0.9	2
58	Direct-to-Consumer Genomics Companies Should Provide Guidance to Their Customers on (Not) Sharing Personal Genomic Information. American Journal of Bioethics, 2014, 14, 55-57.	0.9	1
59	Genetic Ancestry Testing. , 2015, , 898-904.		1
60	Beyond the Recommendation: Discerning Achievable Goals in Clinical Ethics Consultation. American Journal of Bioethics, 2015, 15, 42-44.	0.9	1
61	New Words and Old Stories: Indigenous Teachings in Health Care and Bioethics. American Journal of Bioethics, 2016, 16, 50-52.	0.9	1
62	Response to Patryn and Zagaja. Genetics in Medicine, 2016, 18, 751-751.	2.4	0
63	Response to Koeller et al Genetics in Medicine, 2017, 19, 1380-1380.	2.4	0
64	Analogies in Genomics Policymaking: Debates and Drawbacks. American Journal of Human Genetics, 2020, 107, 797-801.	6.2	0
65	Informed Consent and the Ethics of IRB Research. , 2020, , 103-108.		О