Sandra Soo-Jin Lee

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/3502869/publications.pdf

Version: 2024-02-01

62 papers

1,744 citations

331670 21 h-index 302126 39 g-index

64 all docs

64 docs citations

times ranked

64

1844 citing authors

#	Article	IF	CITATIONS
1	Genetic Research and Health Disparities. JAMA - Journal of the American Medical Association, 2004, 291, 2985.	7.4	227
2	Beyond Consent: Building Trusting Relationships With Diverse Populations in Precision Medicine Research. American Journal of Bioethics, 2018, 18, 3-20.	0.9	152
3	Patient Perspectives on the Learning Health System: The Importance of Trust and Shared Decision Making. American Journal of Bioethics, 2015, 15, 4-17.	0.9	94
4	Attitudes Toward Risk and Informed Consent for Research on Medical Practices. Annals of Internal Medicine, 2015, 162, 690-696.	3.9	87
5	Racializing Drug Design: Implications of Pharmacogenomics for Health Disparities. American Journal of Public Health, 2005, 95, 2133-2138.	2.7	82
6	The ethics of characterizing difference: guiding principles on using racial categories in human genetics. Genome Biology, 2008, 9, 404.	9.6	75
7	Reflections on the Cost of "Low-Cost" Whole Genome Sequencing: Framing the Health Policy Debate. PLoS Biology, 2013, 11, e1001699.	5.6	67
8	Research 2.0: Social Networking and Direct-To-Consumer (DTC) Genomics. American Journal of Bioethics, 2009, 9, 35-44.	0.9	62
9	The Illusive Gold Standard in Genetic Ancestry Testing. Science, 2009, 325, 38-39.	12.6	62
10	Personal genome testing in medical education: student experiences with genotyping in the classroom. Genome Medicine, 2013, 5, 24.	8.2	53
11	Enacting the molecular imperative: How gene-environment interaction research links bodies and environments in the post-genomic age. Social Science and Medicine, 2016, 155, 51-60.	3.8	52
12	A randomized study of multimedia informational aids for research on medical practices: Implications for informed consent. Clinical Trials, 2017, 14, 94-102.	1.6	48
13	Views of Genetics Health Professionals on the Return of Genomic Results. Journal of Genetic Counseling, 2014, 23, 531-538.	1.6	43
14	Race, Distributive Justice and the Promise of Pharmacogenomics. Molecular Diagnosis and Therapy, 2003, 3, 385-392.	3.3	37
15	A comparison of institutional review board professionals' and patients' views on consent for research on medical practices. Clinical Trials, 2016, 13, 555-565.	1.6	36
16	Obligations of the "Gift― Reciprocity and Responsibility in Precision Medicine. American Journal of Bioethics, 2021, 21, 57-66.	0.9	34
17	Adrift in the gray zone: IRB perspectives on research in the learning health system. AJOB Empirical Bioethics, 2016, 7, 125-134.	1.6	29
18	Racing Forward: The Genomics and Personalized Medicine Act. Science, 2009, 323, 342-342.	12.6	27

#	Article	IF	CITATIONS
19	Ethics of inclusion: Cultivate trust in precision medicine. Science, 2019, 364, 941-942.	12.6	27
20	Anticipating uncertainty and irrevocable decisions: provider perspectives on implementing whole-genome sequencing in critically ill children with heart disease. Genetics in Medicine, 2018, 20, 1455-1461.	2.4	26
21	Race, Risk, and Recreation in Personal Genomics: The Limits of Play. Medical Anthropology Quarterly, 2013, 27, 550-569.	1.4	25
22	Homogeneity and heterogeneity as situational properties: Producing – and moving beyond? – race in post-genomic science. Social Studies of Science, 2014, 44, 579-599.	2.5	25
23	Health equality, race and pharmacogenomics. British Journal of Clinical Pharmacology, 2022, 88, 27-33.	2.4	24
24	Accounting for Complexity. Science Technology and Human Values, 2016, 41, 194-218.	3.1	21
25	Dys-appearing Tongues and Bodily Memories: The Aging of First-Generation Resident Koreans in Japan. Ethos, 2000, 28, 198-223.	0.2	20
26	Biobanks of a †racial kind': mining for difference in the new genetics. Patterns of Prejudice, 2006, 40, 443-460.	0.6	20
27	Cancer Health Assessments Reaching Many (CHARM): A clinical trial assessing a multimodal cancer genetics services delivery program and its impact on diverse populations. Contemporary Clinical Trials, 2021, 106, 106432.	1.8	19
28	Race and Ancestry in the Age of Inclusion: Technique and Meaning in Post-Genomic Science. Journal of Health and Social Behavior, 2014, 55, 504-518.	4.8	18
29	American DNA. Current Anthropology, 2013, 54, S77-S86.	1.6	17
30	Protecting Posted Genes: Social Networking and the Limits of GINA. American Journal of Bioethics, 2014, 14, 32-44.	0.9	17
31	Is precision medicine relevant in the age of COVID-19?. Genetics in Medicine, 2021, 23, 999-1000.	2.4	17
32	Lessons learned about harmonizing survey measures for the CSER consortium. Journal of Clinical and Translational Science, 2020, 4, 537-546.	0.6	16
33	Introduction to the article collection †Translation in healthcare: ethical, legal, and social implications'. BMC Medical Ethics, 2016, 17, 74.	2.4	14
34	Translating genomic testing results for pediatric critical care: Opportunities for genetic counselors. Journal of Genetic Counseling, 2020, 29, 78-87.	1.6	13
35	Family secrets: Experiences and outcomes of participating in direct-to-consumer genetic relative-finder services. American Journal of Human Genetics, 2022, 109, 486-497.	6.2	12
36	Integrating stakeholder feedback in translational genomics research: an ethnographic analysis of a study protocol's evolution. Genetics in Medicine, 2020, 22, 1094-1101.	2.4	11

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37	Participant Reactions to a Literacy-Focused, Web-Based Informed Consent Approach for a Genomic Implementation Study. AJOB Empirical Bioethics, 2021, 12, 1-11.	1.6	11
38	Strategies for recruiting representative samples of Asian Americans for a population-based case–control study. Journal of Epidemiology and Community Health, 2016, 70, 974-982.	3.7	10
39	Consuming DNA: The Good Citizen in the Age of Precision Medicine. Annual Review of Anthropology, 2017, 46, 33-48.	1.5	9
40	Institutional culture is the key to team science. Nature Biotechnology, 2017, 35, 1212-1214.	17.5	9
41	Studying "Friends― The Ethics of Using Social Media as Research Platforms. American Journal of Bioethics, 2017, 17, 1-2.	0.9	8
42	Beyond inclusion: Enacting team equity in precision medicine research. PLoS ONE, 2022, 17, e0263750.	2.5	8
43	Attitudes towards Social Networking and Sharing Behaviors among Consumers of Direct-to-Consumer Personal Genomics. Journal of Personalized Medicine, 2013, 3, 275-287.	2.5	7
44	The Biobank as Political Artifact. Annals of the American Academy of Political and Social Science, 2015, 661, 143-159.	1.6	7
45	personalized medicine and pharmacogenomics: ethical and social challenges. Personalized Medicine, 2005, 2, 29-35.	1.5	6
46	Race and the science of difference in the age of genomics. , 0, , 26-39.		6
47	Trustworthiness in Untrustworthy Times: Response to Open Peer Commentaries on <i>Beyond Consent</i> . American Journal of Bioethics, 2018, 18, W6-W8.	0.9	6
48	The Ethics of Consent in a Shifting Genomic Ecosystem. Annual Review of Biomedical Data Science, 2021, 4, 145-164.	6.5	6
49	Laboratory-related outcomes from integrating an accessible delivery model for hereditary cancer risk assessment and genetic testing in populations with barriers to access. Genetics in Medicine, 2022, 24, 1196-1205.	2.4	6
50	Strategies of inclusion: The tradeoffs of pursuing "baked in―diversity through place-based recruitment. Social Science and Medicine, 2022, 306, 115132.	3.8	6
51	The time is ripe for an ethics of entrepreneurship. Nature Biotechnology, 2014, 32, 316-318.	17.5	5
52	Lessons Learned From the U.S. Public Health Service Syphilis Study at Tuskegee: Incorporating a Discourse on Relationships Into the Ethics of Research Participation Among Asian Americans. Ethics and Behavior, 2012, 22, 489-492.	1.8	4
53	Assessing the Pedagogical Goals of Self-Testing in Evaluating the Consultation Needs of Different Student Populations. American Journal of Bioethics, 2012, 12, 41-43.	0.9	4
54	Excavating the Personal Genome: The Good Biocitizen in the Age of Precision Health. Hastings Center Report, 2020, 50, S54-S61.	1.0	4

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55	Parental Attitudes Toward Clinical Genomic Sequencing in Children With Critical Cardiac Disease. Pediatric Critical Care Medicine, 2021, 22, e419-e426.	0.5	4
56	Challenges and potential solutions to health disparities in genomic medicine. Cell, 2022, 185, 2007-2010.	28.9	3
57	Response to Open Peer Commentaries: Distinguishing the "Gift―from "Donation―as a Path toward Reciprocity and Relational Ethics. American Journal of Bioethics, 2021, 21, W1-W3.	0.9	2
58	Response to Open Peer Commentaries on "Research 2.0: Social Networking and Direct-to-Consumer Personal Genomicsâ€. American Journal of Bioethics, 2009, 9, W1-W3.	0.9	1
59	Ethical Considerations for Pharmacogenomics: Privacy and Confidentiality., 0,, 66-73.		1
60	The Role of Patient Perspectives in Clinical Research Ethics and Policy: Response to Open Peer Commentaries on "Patient Perspectives on the Learning Health System― American Journal of Bioethics, 2016, 16, W7-W9.	0.9	1
61	Feasibility of a Traceback Approach for Using Pathology Specimens to Facilitate Genetic Testing in the Genetic Risk Analysis in Ovarian Cancer (GRACE) Study Protocol. Journal of Personalized Medicine, 2021, 11, 1194.	2.5	1
62	Direct-to-Consumer Personal Genetic Testing. , 2015, , 440-445.		0