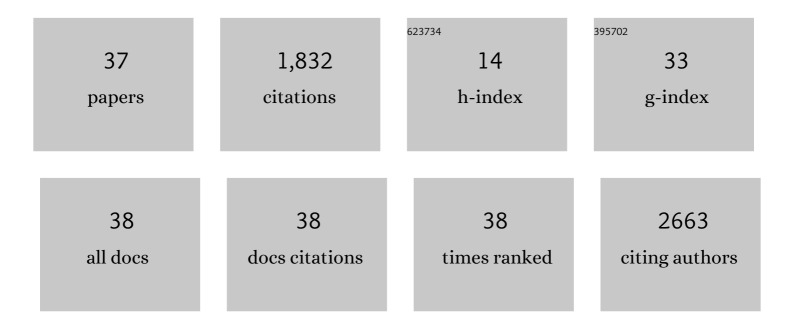
Lisa S Parker

List of Publications by Year in descending order

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LIGA S DADKED

#	Article	IF	CITATIONS
1	The Ethics of Tobacco Harm Reduction: An Analysis of E-Cigarette Availability From the Perspectives of Utilitarianism, Bioethics, and Public Health Ethics. Nicotine and Tobacco Research, 2021, 23, 3-8.	2.6	13
2	Practices and Attitudes toward Returning Genomic Research Results to Low-Resource Research Participants. Public Health Genomics, 2021, 24, 241-252.	1.0	6
3	Ethical Practices and Legal Challenges in Mental Health Research. Asian Bioethics Review, 2020, 12, 87-102.	1.3	3
4	Normative and conceptual ELSI research: what it is, and why it's important. Genetics in Medicine, 2019, 21, 505-509.	2.4	19
5	Research Use of Ecological Momentary Assessment for Adverse Event Monitoring Following Amyloid-β Results Disclosure. Journal of Alzheimer's Disease, 2019, 71, 1071-1079.	2.6	7
6	Jenny Reardon, The Postgenomic Condition: Ethics, Justice, and Knowledge after the Genome. Chicago: University of Chicago Press, 2017, ISBN: 9780226510453 (PB). Hypatia Reviews Online, 2019, 2019, .	0.0	1
7	â€~Recurrent losers unite': Online forums, evidence-based activism, and pregnancy loss. Social Science and Medicine, 2018, 216, 74-80.	3.8	4
8	A qualitative study of Institutional Ethics Committees: Members' understanding of research guidelines, privacy, and challenges to privacy protection. Indian Journal of Medical Ethics, 2018, III, 315-320.	0.4	6
9	Caring for Patients or Organs: New Therapies Raise New Dilemmas in the Emergency Department. American Journal of Bioethics, 2017, 17, 6-16.	0.9	16
10	The Precision Medicine Initiative's All of Us Research Program: an agenda for research on its ethical, legal, and social issues. Genetics in Medicine, 2017, 19, 743-750.	2.4	222
11	EPR and uDCDD: A Response to Commentaries. American Journal of Bioethics, 2017, 17, W1-W3.	0.9	0
12	Advancing Pharmacogenomics Education in the Core PharmD Curriculum through Student Personal Genomic Testing. American Journal of Pharmaceutical Education, 2016, 80, 3.	2.1	73
13	Answers to Questions that Weren't Asked: The Ethical Complexities of Identifying Misattributed Paternity During the Transplant Evaluation Process. International Library of Ethics, Law, and the New Medicine, 2016, , 43-59.	0.5	0
14	Psychometric Properties of a Decisional Capacity Screening Tool for Individuals Contemplating Participation in Alzheimer's Disease Research. Journal of Alzheimer's Disease, 2015, 46, 1-9.	2.6	25
15	Prenatal wholeâ€exome sequencing: parental attitudes. Prenatal Diagnosis, 2015, 35, 1030-1036.	2.3	56
16	False Positives, Outliers, and Other Considerations in Decisional Capacity Screening for Research Consent. Journal of Alzheimer's Disease, 2015, 46, 15-16.	2.6	1
17	Sickle Cell Trait Screening of Collegiate Athletes: Ethical Reasons for Program Reform. Journal of Genetic Counseling, 2015, 24, 873-877.	1.6	11
18	Ethical Considerations Regarding Classroom Use of Personal Genomic Information. Journal of Microbiology and Biology Education, 2014, 15, 191-196.	1.0	6

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#	Article	IF	CITATIONS
19	Managing Incidental Genomic Findings in Clinical Trials: Fulfillment of the Principle of Justice. PLoS Medicine, 2014, 11, e1001584.	8.4	14
20	Subtle Psychosocial Sequelae of Genetic Test Results. Current Genetic Medicine Reports, 2014, 2, 242-249.	1.9	5
21	Incidental findings in the use of DNA to identify human remains: An ethical assessment. Forensic Science International: Genetics, 2013, 7, 221-229.	3.1	28
22	Laboratory informatics based evaluation of methylene tetrahydrofolate reductase C677T genetic test overutilization. Journal of Pathology Informatics, 2013, 4, 33.	1.7	3
23	DNA Identification After Conflict or Disaster. Science, 2013, 341, 1178-1179.	12.6	11
24	Ethical Dimensions of Disparities in Depression Research and Treatment in the Pharmacogenomic Era. Journal of Law, Medicine and Ethics, 2012, 40, 886-903.	0.9	7
25	In Sport and Social Justice, Is Genetic Enhancement a Game Changer?. Health Care Analysis, 2012, 20, 328-346.	2.2	8
26	Managing incidental findings and research results in genomic research involving biobanks and archived data sets. Genetics in Medicine, 2012, 14, 361-384.	2.4	418
27	Comparative effectiveness research: a threat to patient autonomy?. Health Progress, 2011, 92, 64-71.	0.0	0
28	Developing a Community-Academic Partnership to Improve Recognition and Treatment of Depression in Underserved African American and White Elders. American Journal of Geriatric Psychiatry, 2009, 17, 953-964.	1.2	25
29	Managing Incidental Findings in Human Subjects Research: Analysis and Recommendations. Journal of Law, Medicine and Ethics, 2008, 36, 219-248.	0.9	594
30	The Future of Incidental Findings: Should They be Viewed as Benefits?. Journal of Law, Medicine and Ethics, 2008, 36, 341-351.	0.9	50
31	Changing interpretations, stable genes: responsibilities of patients, professionals, and policy makers in the clinical interpretation of complex genetic information. Genetics in Medicine, 2008, 10, 778-783.	2.4	35
32	Disclosure Issues in Neuroscience Research. Accountability in Research, 2008, 15, 226-241.	2.4	2
33	Best Laid Plans for Offering Results Go Awry. American Journal of Bioethics, 2006, 6, 22-23.	0.9	26
34	INFORMATION(AL) MATTERS: BIOETHICS AND THE BOUNDARIES OF THE PUBLIC AND THE PRIVATE. Social Philosophy and Policy, 2002, 19, 83-112.	0.2	6
35	Ethical issues in bipolar disorders pedigree research: privacy concerns, informed consent, and grounds for waiver ¹ . Bipolar Disorders, 2002, 4, 1-16.	1.9	22

#	Article	IF	CITATIONS
37	Preventive Ethics in the Management of Ophthalmic Genetic Disorders. Seminars in Ophthalmology, 1995, 10, 331-343.	1.6	1