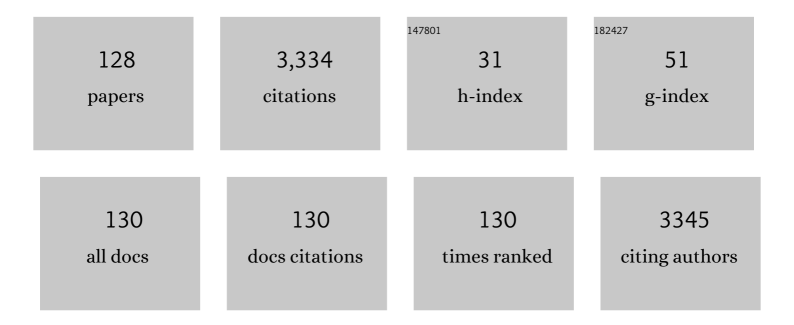
List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	Defining Neuromarketing: Practices and Professional Challenges. Harvard Review of Psychiatry, 2010, 18, 230-237.	2.1	150
2	Attitudes and Practices Among Internists Concerning Genetic Testing. Journal of Genetic Counseling, 2013, 22, 90-100.	1.6	147
3	MDMA (â€~ecstasy') use, and its association with high risk behaviors, mental health, and other factors among gay/bisexual men in New York City. Drug and Alcohol Dependence, 2002, 66, 115-125.	3.2	138
4	MDMA ("Ecstasyâ€) Abuse and High-Risk Sexual Behaviors Among 169 Gay and Bisexual Men. American Journal of Psychiatry, 2000, 157, 1162-1164.	7.2	130
5	Patterns of Communication Between Gay and Lesbian Patients and Their Health Care Providers. Journal of Homosexuality, 2002, 42, 65-75.	2.0	118
6	Researchers' views on return of incidental genomic research results: qualitative and quantitative findings. Genetics in Medicine, 2013, 15, 888-895.	2.4	103
7	The Natural Incubation Period of Kuru and the Episodes of Transmission in Three Clusters of Patients. Neuroepidemiology, 1984, 3, 3-20.	2.3	96
8	Problems in comprehension of informed consent in rural and peri-urban Mali, West Africa. Clinical Trials, 2006, 3, 306-313.	1.6	91
9	Use of Genetic Tests among Neurologists and Psychiatrists: Knowledge, Attitudes, Behaviors, and Needs for Training. Journal of Genetic Counseling, 2014, 23, 156-163.	1.6	87
10	Voluntariness of Consent to Research: A Conceptual Model. Hastings Center Report, 2009, 39, 30-39.	1.0	83
11	Models of Consent to Return of Incidental Findings in Genomic Research. Hastings Center Report, 2014, 44, 22-32.	1.0	72
12	Return of Secondary Genomic Findings vs Patient Autonomy. JAMA - Journal of the American Medical Association, 2013, 310, 369.	7.4	68
13	Decision-Making About Reproductive Choices Among Individuals At-Risk for Huntington's Disease. Journal of Genetic Counseling, 2007, 16, 347-362.	1.6	63
14	Improving Education on Doctor???Patient Relationships and Communication: Lessons from Doctors Who Become Patients. Academic Medicine, 2006, 81, 447-453.	1.6	62
15	Recruiting egg donors online: an analysis of inÂvitro fertilization clinic and agency websites' adherence to American Society for Reproductive Medicine guidelines. Fertility and Sterility, 2012, 98, 995-1000.	1.0	61
16	New Challenges for Electronic Health Records. JAMA - Journal of the American Medical Association, 2015, 313, 29.	7.4	59
17	Informed consent for return of incidental findings in genomic research. Genetics in Medicine, 2014, 16, 367-373.	2.4	58
18	Views of Discrimination among Individuals Confronting Genetic Disease. Journal of Genetic Counseling, 2010, 19, 68-83.	1.6	57

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19	Misunderstandings Concerning Genetics Among Patients Confronting Genetic Disease. Journal of Genetic Counseling, 2010, 19, 430-446.	1.6	57
20	Institutional Review Board Community Members. Academic Medicine, 2012, 87, 975-981.	1.6	56
21	How IRBs view and make decisions about coercion and undue influence: Table 1. Journal of Medical Ethics, 2013, 39, 224-229.	1.8	50
22	Disclosures of Huntington disease risk within families: Patterns of decisionâ€making and implications. American Journal of Medical Genetics, Part A, 2007, 143A, 1835-1849.	1.2	47
23	The Ethics Police?: IRBs' Views Concerning Their Power. PLoS ONE, 2011, 6, e28773.	2.5	43
24	The process of deciding about prophylactic surgery for breast and ovarian cancer: Patient questions, uncertainties, and communication. American Journal of Medical Genetics, Part A, 2010, 152A, 52-66.	1.2	41
25	How local IRBs view central IRBs in the US. BMC Medical Ethics, 2011, 12, 13.	2.4	39
26	"Am I my genes?― Questions of identity among individuals confronting genetic disease. Genetics in Medicine, 2009, 11, 880-889.	2.4	38
27	US IRBS CONFRONTING RESEARCH IN THE DEVELOPING WORLD. Developing World Bioethics, 2012, 12, 63-73.	0.9	36
28	"Patient-timeâ€, "doctor-timeâ€, and "institution-timeâ€, Perceptions and definitions of time among doctors who become patients. Patient Education and Counseling, 2007, 66, 147-155.	2.2	34
29	Payment of egg donors in stem cell research in the USA. Reproductive BioMedicine Online, 2009, 18, 603-608.	2.4	34
30	The roles of family members, health care workers, and others in decision-making processes about genetic testing among individuals at risk for Huntington disease. Genetics in Medicine, 2007, 9, 358-371.	2.4	33
31	Self-Disclosure of HIV Status to Sexual Partners: A Qualitative Study of Issues Faced by Gay Men. Journal of the Gay and Lesbian Medical Association, 1999, 3, 39-49.	0.6	31
32	The Myth of Community Differences as the Cause of Variations Among IRBs. American Journal of Bioethics Primary Research, 2011, 2, 24-33.	1.5	31
33	How old is too old? Challenges faced by clinicians concerning age cutoffs for patients undergoing in vitro fertilization. Fertility and Sterility, 2016, 106, 216-224.	1.0	31
34	Deciding how many embryos to transfer: ongoing challenges and dilemmas. Reproductive Biomedicine and Society Online, 2016, 3, 1-15.	1.8	30
35	Processes and factors involved in decisions regarding return of incidental genomic findings in research. Genetics in Medicine, 2014, 16, 311-317.	2.4	28
36	How much is a child worth? Providers' and patients' views and responses concerning ethical and policy challenges in paying for ART. PLoS ONE, 2017, 12, e0171939.	2.5	28

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37	Challenges and changes in spirituality among doctors who become patients. Social Science and Medicine, 2005, 61, 2396-2406.	3.8	27
38	Views and Experiences of IRBs concerning Research Integrity. Journal of Law, Medicine and Ethics, 2011, 39, 513-528.	0.9	27
39	"Members of the Same Clubâ€: Challenges and Decisions Faced by US IRBs in Identifying and Managing Conflicts of Interest. PLoS ONE, 2011, 6, e22796.	2.5	27
40	Impact of Receiving Secondary Results from Genomic Research: A 12â€Month Longitudinal Study. Journal of Genetic Counseling, 2018, 27, 709-722.	1.6	26
41	How IRBs View and Make Decisions about Social Risks. Journal of Empirical Research on Human Research Ethics, 2013, 8, 58-65.	1.3	25
42	VIEWS OF THE PROCESS AND CONTENT OF ETHICAL REVIEWS OF HIV VACCINE TRIALS AMONG MEMBERS OF US INSTITUTIONAL REVIEW BOARDS AND SOUTH AFRICAN RESEARCH ETHICS COMMITTEES. Developing World Bioethics, 2008, 8, 207-218.	0.9	23
43	Anticipating issues related to increasing preimplantation genetic diagnosis use: a research agenda. Reproductive BioMedicine Online, 2008, 17, 33-42.	2.4	23
44	Advocating for longitudinal follow-up of the health and welfare of egg donors. Fertility and Sterility, 2014, 102, 662-666.	1.0	23
45	Exclusion of Genetic Information From the Medical Record. JAMA - Journal of the American Medical Association, 2010, 304, 1120.	7.4	21
46	"In Sickness and in Health� Disclosures of Genetic Risks in Dating. Journal of Genetic Counseling, 2011, 20, 98-112.	1.6	21
47	Views of internists towards uses of PGD. Reproductive BioMedicine Online, 2013, 26, 142-147.	2.4	21
48	How IRBs View and Make Decisions about Consent Forms. Journal of Empirical Research on Human Research Ethics, 2013, 8, 8-19.	1.3	21
49	Impediments to communication and relationships between infertility care providers and patients. BMC Women's Health, 2018, 18, 84.	2.0	21
50	Voluntariness of consent to research: a preliminary empirical investigation. IRB: Ethics & Human Research, 2009, 31, 10-4.	0.8	21
51	Naming names: Perceptions of name-based HIV reporting, partner notification, and criminalization of non-disclosure among persons living with HIV. Sexuality Research and Social Policy, 2004, 1, 38-57.	2.3	19
52	Challenges, Dilemmas and Factors Involved in PGD Decisionâ€Making: Providers' and Patients' Views, Experiences and Decisions. Journal of Genetic Counseling, 2018, 27, 909-919.	1.6	19
53	From anonymity to "open doors†IRB responses to tensions with researchers. BMC Research Notes, 2012, 5, 347.	1.4	18
54	How good does the science have to be in proposals submitted to Institutional Review Boards? An Interview Study of Institutional Review Board personnel. Clinical Trials, 2013, 10, 761-766.	1.6	18

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55	Psychiatrists' Views of the Genetic Bases of Mental Disorders and Behavioral Traits and Their Use of Genetic Tests. Journal of Nervous and Mental Disease, 2014, 202, 530-538.	1.0	18
56	How Agencies Market Egg Donation on the Internet: A Qualitative Study. Journal of Law, Medicine and Ethics, 2015, 43, 610-618.	0.9	18
57	Buying and selling human eggs: infertility providers' ethical and other concerns regarding egg donor agencies. BMC Medical Ethics, 2016, 17, 71.	2.4	18
58	Reducing the number of fetuses in a pregnancy: providers' and patients' views of challenges. Human Reproduction, 2016, 31, 2570-2576.	0.9	18
59	Impact of patient education videos on genetic counseling outcomes after exome sequencing. Patient Education and Counseling, 2020, 103, 127-135.	2.2	18
60	Ethical issues concerning disclosures of HIV diagnoses to perinatally infected children and adolescents. Journal of Clinical Ethics, 2008, 19, 31-42.	0.3	18
61	Disclosures of illness by doctors to their patients: A qualitative study of doctors with HIV and other serious disorders. Patient Education and Counseling, 2006, 64, 277-284.	2.2	17
62	Preimplantation genetic diagnosis on in vitro fertilization clinic websites: presentations of risks, benefits and other information. Fertility and Sterility, 2009, 92, 1276-1283.	1.0	17
63	Association of Researcher Characteristics with Views on Return of Incidental Findings from Genomic Research. Journal of Genetic Counseling, 2015, 24, 833-841.	1.6	17
64	Reliance agreements and single IRB review of multisite research: Concerns of IRB members and staff. AJOB Empirical Bioethics, 2018, 9, 164-172.	1.6	17
65	HIV and the Law: Integrating Law, Policy, and Social Epidemiology. Journal of Law, Medicine and Ethics, 2002, 30, 533-547.	0.9	16
66	"Post-Residency Disease" and the Medical Self: Identity, Work, and Health Care Among Doctors Who Become Patients. Perspectives in Biology and Medicine, 2006, 49, 542-552.	0.5	16
67	Unconventional combinations of prospective parents: ethical challenges faced by IVF providers. BMC Medical Ethics, 2017, 18, 18.	2.4	16
68	Views and approaches toward risks and benefits among doctors who become patients. Patient Education and Counseling, 2006, 64, 61-68.	2.2	15
69	Designing Babies. , 2019, , .		15
70	The Reporting of Monetary Compensation in Research Articles. Journal of Empirical Research on Human Research Ethics, 2007, 2, 61-67.	1.3	14
71	Controversies concerning mitochondrial replacement therapy. Fertility and Sterility, 2015, 103, 344-346.	1.0	13
72	Pain during embryo transfer is independently associated with clinical pregnancy in fresh/frozen assisted reproductive technology cycles. Journal of Obstetrics and Gynaecology Research, 2016, 42, 684-693.	1.3	13

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73	From "Male Bonding Rituals―to "Suicide Tuesday― Journal of Homosexuality, 2006, 51, 7-32.	2.0	12
74	The Impact of Social Contexts in Testing for Alpha-1 Antitrypsin Deficiency: The Roles of Physicians and Others. Genetic Testing and Molecular Biomarkers, 2009, 13, 269-276.	0.7	12
75	Struggles in Defining and Addressing Requests for "Family Balancingâ€: Ethical Issues Faced by Providers and Patients. Journal of Law, Medicine and Ethics, 2016, 44, 616-629.	0.9	12
76	Local Knowledge and Single IRBs for Multisite Studies: <i>Challenges and Solutions</i> . Ethics & Human Research, 2019, 41, 22-31.	0.9	12
77	To Protect Human Subjects, Review What Was Done, Not Proposed. Science, 2012, 335, 1576-1577.	12.6	11
78	Should Life Insurers Have Access to Genetic Test Results?. JAMA - Journal of the American Medical Association, 2014, 312, 1855.	7.4	11
79	Linguistic and Cultural Challenges in Communication and Translation in US-Sponsored HIV Prevention Research in Emerging Economies. PLoS ONE, 2015, 10, e0133394.	2.5	11
80	Disclosures of funding sources and conflicts of interest in published HIV/AIDS research conducted in developing countries. Journal of Medical Ethics, 2010, 36, 505-510.	1.8	10
81	Evolving Challenges and Research-Needs Concerning Ebola. American Journal of Public Health, 2015, 105, 1513-1515.	2.7	10
82	Evaluation of the cost and effectiveness of diverse recruitment methods for a genetic screening study. Genetics in Medicine, 2019, 21, 2371-2380.	2.4	10
83	Views of preimplantation genetic diagnosis among psychiatrists and neurologists. Journal of reproductive medicine, The, 2014, 59, 385-92.	0.2	10
84	Disclosure of information to potential subjects on research recruitment web sites. IRB: Ethics & Human Research, 2008, 30, 15-20.	0.8	10
85	Contexts, Anyone?: The Need for Contextualization in the Debate About the Moral Status of Embryos. American Journal of Bioethics, 2005, 5, 56-58.	0.9	9
86	The Use of Eggs and Embryos in Stem Cell Research. Seminars in Reproductive Medicine, 2010, 28, 336-344.	1.1	9
87	Kamakahi vs ASRM and the future of compensation for human eggs. American Journal of Obstetrics and Gynecology, 2015, 213, 186-187.e1.	1.3	9
88	THE REPORTING OF IRB REVIEW IN JOURNAL ARTICLES PRESENTING HIV RESEARCH CONDUCTED IN THE DEVELOPING WORLD. Developing World Bioethics, 2011, 11, 161-169.	0.9	8
89	HIV/AIDS Research Conducted in the Developing World and Sponsored by the Developed World: Reporting of Research Ethics Committee Review in Two Countries. Journal of Empirical Research on Human Research Ethics, 2011, 6, 83-91.	1.3	8
90	Infertility providers' and patients' views and experiences concerning doctor shopping in the USA. Human Fertility, 2019, 22, 238-245.	1.7	8

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91	Legal Immunity for Physicians During the COVID-19 Pandemic. Chest, 2020, 158, 1343-1345.	0.8	8
92	Hospital chaplains' communication with patients: Characteristics, functions and potential benefits. Patient Education and Counseling, 2022, 105, 2905-2912.	2.2	8
93	How US institutional review boards decide when researchers need to translate studies. Journal of Medical Ethics, 2014, 40, 193-197.	1.8	7
94	Creating and selling embryos for "donation― ethical challenges. American Journal of Obstetrics and Gynecology, 2015, 212, 167-170.e1.	1.3	7
95	"Will they be good enough parents?†Ethical dilemmas, views, and decisions among assisted reproductive technology (ART) providers. AJOB Empirical Bioethics, 2017, 8, 253-265.	1.6	7
96	How Infertility Patients and Providers View and Confront Religious and Spiritual Issues. Journal of Religion and Health, 2018, 57, 223-239.	1.7	7
97	Gatekeepers for infertility treatment? Views of ART providers concerning referrals by non-ART providers. Reproductive Biomedicine and Society Online, 2018, 5, 17-30.	1.8	7
98	Local IRBs vs. Federal Agencies: Shifting Dynamics, Systems, and Relationships. Journal of Empirical Research on Human Research Ethics, 2012, 7, 50-62.	1.3	6
99	Needs to Prepare for "Post-COVID-19 Syndrome― American Journal of Bioethics, 2020, 20, 4-6.	0.9	6
100	Understanding Ethical Challenges in Medical Education Research. Academic Medicine, 2022, 97, 18-21.	1.6	6
101	Egg donation brokers: an analysis of agency versus in vitro fertilization clinic websites. Journal of reproductive medicine, The, 2014, 59, 534-41.	0.2	6
102	Qualifying Confidentiality: Historical and Empirical Issues and Facts. American Journal of Bioethics, 2006, 6, 26-27.	0.9	5
103	Doctor, Will You Pray for Me? Responding to Patients' Religious and Spiritual Concerns. Academic Medicine, 2021, 96, 349-354.	1.6	5
104	How hospital chaplains develop and use rituals to address medical staff distress. SSM Qualitative Research in Health, 2022, 2, 100087.	1.5	5
105	The Need for Vigilance in the Marketing of Genomic Tests in Psychiatry. Journal of Nervous and Mental Disease, 2015, 203, 809-810.	1.0	4
106	Reducing the Single IRB Burden: Streamlining Electronic IRB Systems. AJOB Empirical Bioethics, 2021, 12, 33-40.	1.6	4
107	Consenting for Molecular Diagnostics. Clinical Chemistry, 2015, 61, 139-141.	3.2	3
108	How Single Institutional Review Boards Manage Their Own Conflicts of Interest. Academic Medicine, 2019, 94, 1554-1560.	1.6	3

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109	Typologies and Meanings of Prayer Among Patients. Journal of Religion and Health, 2021, , 1.	1.7	3
110	Predictive testing and clinical trials in Huntington's disease: An ethical analysis. Movement Disorders, 2018, 33, 243-247.	3.9	3
111	How IRB leaders view and approach challenges raised by industry-funded research. IRB: Ethics & Human Research, 2013, 35, 9-17.	0.8	3
112	Sexual orientation and associated characteristics among north american academic psychiatrists. Journal of Sex Research, 1998, 35, 282-287.	2.5	2
113	Additional Implications of a National Survey on Ethics Consultation in United States Hospitals. American Journal of Bioethics, 2007, 7, 47-48.	0.9	2
114	Views of IRBs Concerning Their Local Ecologies: Perceptions of Relationships, Systems, and Tensions Between IRBs and Their Institutions. American Journal of Bioethics Primary Research, 2013, 4, 31-43.	1.5	2
115	Reviewing <scp>HIV</scp> â€Related Research in Emerging Economies: The Role of Government Reviewing Agencies. Developing World Bioethics, 2016, 16, 4-14.	0.9	2
116	Ethical and Psychosocial Considerations in Informing HIV-Exposed Uninfected Children That They Were Exposed to HIV and Antiretroviral Medications In Utero. American Journal of Public Health, 2016, 106, 1390-1396.	2.7	2
117	Kuru fieldwork in 1981 … and beyond. Philosophical Transactions of the Royal Society B: Biological Sciences, 2008, 363, 3646-3647.	4.0	1
118	Electronic Health Records and Adolescent Privacy—Reply. JAMA - Journal of the American Medical Association, 2015, 313, 1373.	7.4	1
119	Bringing science and advocacy together to address health needs of people who inject drugs. Journal of Medical Ethics, 2018, 44, 165-166.	1.8	1
120	Henrietta Lacks' family's lawsuits: ethical questions and solutions. Trends in Biotechnology, 2022, 40, 769-772.	9.3	1
121	Antoni Van Leeuwenhoek, FRS on Vermeer: a figment of the imagination. FASEB Journal, 2006, 20, 591-594.	0.5	0
122	Vermeer and Leeuwenhoek, Figments of the Imagination?. FASEB Journal, 2012, 26, 2238-2238.	0.5	0
123	Bioethics of Translating Limited Evidence into Clinical Practice: Case Study of the Cerebroplacental Ratio. Journal of Obstetrics and Gynaecology Canada, 2020, 42, 1154-1157.	0.7	0
124	How Artistic Representation Can Inform Current Debates About Chimeras. Journal of Medical Humanities, 2021, 42, 337-343.	0.7	0
125	Preparing for the Next Generation of Ethical Challenges Concerning Heritable Human Genome Editing. American Journal of Bioethics, 2021, 21, 1-4.	0.9	0
126	Experiment on identical siblings separated at birth: ethical implications for researchers, universities, and archives today. Journal of Medical Ethics, 2021, 47, 657-661.	1.8	0

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127	Roles of genetics and blood type in clinical responses to COVID-19: ethical and policy concerns. Journal of Medical Ethics, 2021, 47, 149-151.	1.8	0
128	In Reply to Jain and Kassam:. Academic Medicine, 2022, 97, 8-8.	1.6	0