Robert Cook-Deegan

List of Publications by Year in descending order

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125 papers 4,225 citations

147801 31 h-index 60 g-index

133 all docs

133
docs citations

times ranked

133

4437 citing authors

#	Article	IF	Citations
1	Neurocognitive enhancement: what can we do and what should we do?. Nature Reviews Neuroscience, 2004, 5, 421-425.	10.2	546
2	Disclosure of <i> APOE </i> Genotype for Risk of Alzheimer's Disease. New England Journal of Medicine, 2009, 361, 245-254.	27.0	490
3	Broad Consent for Research With Biological Samples: Workshop Conclusions. American Journal of Bioethics, 2015, 15, 34-42.	0.9	221
4	The Human Pangenome Project: a global resource to map genomic diversity. Nature, 2022, 604, 437-446.	27.8	192
5	BRCA Challenge: BRCA Exchange as a global resource for variants in BRCA1 and BRCA2. PLoS Genetics, 2018, 14, e1007752.	3.5	148
6	Genetic Testing For Alzheimer's Disease And Its Impact On Insurance Purchasing Behavior. Health Affairs, 2005, 24, 483-490.	5.2	127
7	Commercial landscape of noninvasive prenatal testing in the United States. Prenatal Diagnosis, 2013, 33, 521-531.	2.3	115
8	Fertility patients' views about frozen embryo disposition: results of a multi-institutional U.S. survey. Fertility and Sterility, 2010, 93, 499-509.	1.0	113
9	Evidence and anecdotes: an analysis of human gene patenting controversies. Nature Biotechnology, 2006, 24, 1091-1094.	17.5	101
10	Patents in Genomics and Human Genetics. Annual Review of Genomics and Human Genetics, 2010, 11, 383-425.	6.2	88
11	Factors that affect infertility patients' decisions about disposition of frozen embryos. Fertility and Sterility, 2006, 85, 1623-1630.	1.0	87
12	Is Bayh-Dole Good for Developing Countries? Lessons from the US Experience. PLoS Biology, 2008, 6, e262.	5.6	83
13	Variants of uncertain significance in BRCA: a harbinger of ethical and policy issues to come?. Genome Medicine, 2014, 6, 121.	8.2	74
14	The next controversy in genetic testing: clinical data as trade secrets?. European Journal of Human Genetics, 2013, 21, 585-588.	2.8	73
15	The licensing of DNA patents by US academic institutions: an empirical survey. Nature Biotechnology, 2006, 24, 31-39.	17.5	72
16	Reflections on the Cost of "Low-Cost" Whole Genome Sequencing: Framing the Health Policy Debate. PLoS Biology, 2013, 11, e1001699.	5.6	67
17	Gene Patenting â€" The Supreme Court Finally Speaks. New England Journal of Medicine, 2013, 369, 869-875.	27.0	58
18	Columbia University's Axel Patents: Technology Transfer and Implications for the Bayhâ€Dole Act. Milbank Quarterly, 2009, 87, 683-715.	4.4	54

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19	Patents, Secrecy, and DNA. Science, 2001, 293, 217-217.	12.6	48
20	Non-reporting and inconsistent reporting of race and ethnicity in articles that claim associations among genotype, outcome, and race or ethnicity. Journal of Medical Ethics, 2006, 32, 724-728.	1.8	44
21	Intellectual property, technology transfer and manufacture of low-cost HPV vaccines in India. Nature Biotechnology, 2010, 28, 671-678.	17.5	42
22	The Bermuda Triangle: The Pragmatics, Policies, and Principles for Data Sharing in the History of the Human Genome Project. Journal of the History of Biology, 2018, 51, 693-805.	0.5	42
23	The effects of business practices, licensing, and intellectual property on development and dissemination of the polymerase chain reaction: case study. Journal of Biomedical Discovery and Collaboration, 2006, 1, 7.	2.0	40
24	Harm, hype and evidence: ELSI research and policy guidance. Genome Medicine, 2013, 5, 21.	8.2	39
25	Impact of gene patents and licensing practices on access to genetic testing for inherited susceptibility to cancer: Comparing breast and ovarian cancers with colon cancers. Genetics in Medicine, 2010, 12, S15-S38.	2.4	37
26	Incorporating ethnicity into genetic risk assessment for Alzheimer disease: the REVEAL study experience. Genetics in Medicine, 2008, 10, 207-214.	2.4	36
27	Impact of gene patents and licensing practices on access to genetic testing for cystic fibrosis. Genetics in Medicine, 2010, 12, S194-S211.	2.4	36
28	Creating a data resource: what will it take to build a medical information commons?. Genome Medicine, 2017, 9, 84.	8.2	36
29	A randomized controlled trial of disclosing genetic risk information for Alzheimer disease via telephone. Genetics in Medicine, 2018, 20, 132-141.	2.4	36
30	The alta summit, December 1984. Genomics, 1989, 5, 661-663.	2.9	35
31	Disclosing Pleiotropic Effects During Genetic Risk Assessment for Alzheimer Disease. Annals of Internal Medicine, 2016, 164, 155.	3.9	34
32	Beyond Our Borders? Public Resistance to Global Genomic Data Sharing. PLoS Biology, 2016, 14, e2000206.	5.6	33
33	Balancing openness with Indigenous data sovereignty: An opportunity to leave no one behind in the journey to sequence all of life. Proceedings of the National Academy of Sciences of the United States of America, 2022, 119, .	7.1	33
34	Direct-to-consumer genetic tests: beyond medical regulation?. Genome Medicine, 2009, 1, 17.	8.2	32
35	The science commons in health research: structure, function, and value. Journal of Technology Transfer, 2007, 32, 133-156.	4.3	31
36	Barriers to clinical adoption of next generation sequencing: Perspectives of a policy Delphi panel. Applied & Translational Genomics, 2016, 10, 19-24.	2.1	30

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37	Patents and Innovation in Cancer Therapeutics: Lessons from CellPro. Milbank Quarterly, 2002, 80, 637-676.	4.4	29
38	After Myriad: Genetic Testing in the Wake of Recent Supreme Court Decisions about Gene Patents. Current Genetic Medicine Reports, 2014, 2, 223-241.	1.9	29
39	Ethical and Practical Issues Associated with Aggregating Databases. PLoS Medicine, 2008, 5, e190.	8.4	28
40	A randomized noninferiority trial of condensed protocols for genetic risk disclosure of Alzheimer's disease. Alzheimer's and Dementia, 2015, 11, 1222-1230.	0.8	28
41	Fostering reproducibility in industry-academia research. Science, 2017, 357, 759-761.	12.6	28
42	The dangers of diagnostic monopolies. Nature, 2009, 458, 405-406.	27.8	27
43	DNA patents and diagnostics: not a pretty picture. Nature Biotechnology, 2010, 28, 784-791.	17.5	27
44	Moving beyond Bermuda: sharing data to build a medical information commons. Genome Research, 2017, 27, 897-901.	5.5	27
45	Impact of gene patents and licensing practices on access to genetic testing and carrier screening for Tay-Sachs and Canavan disease. Genetics in Medicine, 2010, 12, S5-S14.	2.4	23
46	Sharing Data to Build a Medical Information Commons: From Bermuda to the Global Alliance. Annual Review of Genomics and Human Genetics, 2017, 18, 389-415.	6.2	22
47	Public variant databases: liability?. Genetics in Medicine, 2017, 19, 838-841.	2.4	21
48	The Human Genome Project after a decade: policy issues. Nature Genetics, 1998, 20, 333-335.	21.4	20
49	Spinocerebellar ataxia: Patient and health professional perspectives on whether and how patents affect access to clinical genetic testing. Genetics in Medicine, 2010, 12, S83-S110.	2.4	20
50	Sharing data under the 21st Century Cures Act. Genetics in Medicine, 2017, 19, 1289-1294.	2.4	20
51	Characterizing the Biomedical Data-Sharing Landscape. Journal of Law, Medicine and Ethics, 2019, 47, 21-30.	0.9	18
52	The science commons in life science research: structure, function, and value of access to genetic diversity. International Social Science Journal, 2006, 58, 299-317.	1.6	17
53	Gene patents and personalized medicine - what lies ahead?. Genome Medicine, 2009, 1, 92.	8.2	17
54	Impact of gene patents and licensing practices on access to genetic testing for long QT syndrome. Genetics in Medicine, 2010, 12, S111-S154.	2.4	17

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55	Constructing narratives of heroism and villainy: case study of Myriad's BRACAnalysis® compared to Genentech's Herceptin®. Genome Medicine, 2013, 5, 8.	8.2	17
56	Universities: The Fallen Angels of Bayh-Dole?. Daedalus, 2018, 147, 76-89.	1.8	17
57	MYRIAD AFTER THE PROPRIETARY DATA DILEMMA. North Carolina Journal of Law & Technology, 2014, 15, 597-637.	2.0	17
58	Privacy, families, and human subject protections: Some lessons from pedigree research. Journal of Continuing Education in the Health Professions, 2001, 21, 224-237.	1.3	16
59	Metastasizing patent claims on BRCA1. Genomics, 2010, 95, 312-314.	2.9	16
60	Perspectives on Genetic and Genomic Technologies in an Academic Medical Center: The Duke Experience. Journal of Personalized Medicine, 2015, 5, 67-82.	2.5	15
61	Barriers to clinical adoption of next-generation sequencing: a policy Delphi panel's solutions. Personalized Medicine, 2017, 14, 339-354.	1.5	15
62	Reactions to the National Academies/Royal Society Report on <i>Heritable Human Genome Editing</i> CRISPR Journal, 2020, 3, 332-349.	2.9	15
63	Perceptions of Personalized Medicine in an Academic Health System: Educational Findings. Journal of Contemporary Medical Education, 2015, 3, 14.	0.2	15
64	Law and Science Collide Over Human Gene Patents. Science, 2012, 338, 745-747.	12.6	14
65	Gene patents and licensing: Case studies prepared for the Secretary's Advisory Committee on Genetics, Health, and Society. Genetics in Medicine, 2010, 12, S1-S2.	2.4	13
66	Distributing the future: The weak justifications for keeping human genomic databases secret and the challenges and opportunities in reverse engineering them. Applied & Translational Genomics, 2014, 3, 124-127.	2.1	13
67	Genomic Data-Sharing Practices. Journal of Law, Medicine and Ethics, 2019, 47, 31-40.	0.9	12
68	Patents and Misplaced Angst: Lessons for Translational Stem Cell Research from Genomics. Cell Stem Cell, 2013, 12, 508-512.	11.1	11
69	The emergence of commercial genomics: analysis of the rise of a biotechnology subsector during the Human Genome Project, 1990 to 2004. Genome Medicine, 2013, 5, 83.	8.2	11
70	<i>AMP</i> v. <i>Myriad</i> : A Surgical Strike on Blockbuster Business Models. Science Translational Medicine, 2013, 5, 192ed9.	12.4	11
71	Introduction: Sharing Data in a Medical Information Commons. Journal of Law, Medicine and Ethics, 2019, 47, 7-11.	0.9	11
72	Complicated legacies: The human genome at 20. Science, 2021, 371, 564-569.	12.6	11

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73	SCIENCE AND SECURITY: Practical Experiences in Dual-Use Review. Science, 2007, 316, 1432-1433.	12.6	10
74	Impact of gene patents and licensing practices on access to genetic testing for Alzheimer disease. Genetics in Medicine, 2010, 12, S71-S82.	2.4	10
75	Moving Beyond "lsolated―Gene Patents. Science, 2013, 341, 137-138.	12.6	10
76	In support of mitochondrial replacement therapy. Nature Medicine, 2019, 25, 870-871.	30.7	10
77	A survey of U.S. public perspectives on facial recognition technology and facial imaging data practices in health and research contexts. PLoS ONE, 2021, 16, e0257923.	2.5	10
78	Impact of gene patents and licensing practices on access to genetic testing for hereditary hemochromatosis. Genetics in Medicine, 2010, 12, S155-S170.	2.4	9
79	A collaboratively derived international research agenda on legislative science advice. Palgrave Communications, 2019, 5, .	4.7	9
80	Racing for academic glory and patents: Lessons from CRISPR. Science, 2017, 358, 874-876.	12.6	8
81	Ethical, legal, and social issues in the Earth BioGenome Project. Proceedings of the National Academy of Sciences of the United States of America, 2022, 119, e2115859119.	7.1	8
82	The mouse that trolled: the long and tortuous history of a gene mutation patent that became an expensive impediment to Alzheimer's research. Journal of Law and the Biosciences, 2015, 2, 213-262.	1.6	7
83	Open science precision medicine in Canada: Points to consider. Facets, 2019, 4, 1-19.	2.4	7
84	Boosting Health Services Research. Science, 2011, 333, 1384-1385.	12.6	6
85	Patents and Genome-Wide DNA Sequence Analysis: Is it Safe to Go into the Human Genome?. Journal of Law, Medicine and Ethics, 2014, 42, 42-50.	0.9	6
86	Keeping score, strengthening policy and fighting bad actors over access to research tools. Nature Biotechnology, 2015, 33, 143-147.	17.5	5
87	Developing context-specific next-generation sequencing policy. Nature Biotechnology, 2016, 34, 466-470.	17.5	5
88	The continuing saga of patents and nonâ€invasive prenatal testing. Prenatal Diagnosis, 2019, 39, 441-447.	2.3	5
89	BRCA1/2 Variant Data-Sharing Practices. Journal of Law, Medicine and Ethics, 2019, 47, 88-96.	0.9	4
90	Report of the X chromosome workshop. Genomics, 1990, 7, 647-654.	2.9	3

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91	The Genesis of The Human Genome Project. , 1991, 1, 1-75.		3
92	Gene Patents: The Shadow of Uncertainty. Science, 2011, 331, 873-874.	12.6	3
93	Are Human Genes Patentable?. Annals of Internal Medicine, 2013, 159, 298-9.	3.9	3
94	What's the Use? <i>Disparate Purposes of U.S. Federal Bioethics Commissions </i> Report, 2017, 47, S14-S16.	1.0	3
95	Challenges to Building a Gene Variant Commons to Assess Hereditary Cancer Risk: Results of a Modified Policy Delphi Panel Deliberation. Journal of Personalized Medicine, 2021, 11, 646.	2.5	3
96	U.S. Adult Perspectives on Facial Images, DNA, and Other Biometrics. IEEE Transactions on Technology and Society, 2022, 3, 9-15.	3.2	3
97	P <scp>rivate</scp> P <scp>arts</scp> . The Sciences, 1994, 34, 18-23.	0.1	2
98	PUBLIC HEALTH: Genomics and Medicine at a Crossroads in Chernobyl. Science, 2006, 314, 62-63.	12.6	2
99	Rules for Growth: Promoting Innovation and Growth Through Legal Reform. SSRN Electronic Journal, 2011, , .	0.4	2
100	<i>Sequenom v. Ariosa</i> â€" The Death of a Genetic Testing Patent. New England Journal of Medicine, 2016, 375, 2418-2419.	27.0	2
101	CRISPR Patents: Aspiring to Coherent Patent Policy. American Journal of Bioethics, 2018, 18, 51-54.	0.9	2
102	15. Does NIH need a DARPA?. , 2019, , 453-460.		2
103	Participant-Partners in Genetic Research: An Exome Study with Families of Children with Unexplained Medical Conditions. Journal of Participatory Medicine, 2018, 10, e2.	1.3	2
104	The researcher as a congressional director. Trends in Biochemical Sciences, 1982, 7, 434.	7.5	1
105	Social and Ethical Implications of Advances in Human Genetics. Southern Medical Journal, 1990, 83, 879-882.	0.7	1
106	The large DNA insert cloning workshop. Genomics, 1990, 7, 654-660.	2.9	1
107	Alzheimer Testing at Silver Years. Cambridge Quarterly of Healthcare Ethics, 1998, 7, 294-307.	0.8	1
108	Reply to Wrong fixes for gene patents. Nature Biotechnology, 2010, 28, 1243-1243.	17.5	1

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109	The mouse that trolled (again). Journal of Law and the Biosciences, 2016, 3, 185-191.	1.6	1
110	Community crystal gazing. Nature Biotechnology, 2016, 34, 276-283.	17.5	1
111	National Partnership for Maternal Safety: Consensus Bundle on Venous Thromboembolism. Obstetrics and Gynecology, 2019, 134, 1115-1117.	2.4	1
112	Governing Heritable Human Genome Editing: A Textual History and a Proposal for the Future. CRISPR Journal, 2021, 4, 469-476.	2.9	1
113	Public—Private Interactions in Genomic Medicine: Research and Development. , 2009, , 434-444.		1
114	Is Bayh-Dole Good for Developing Countries? Lessons from the U.S. Experience. , 2014, , 201-215.		1
115	Cystic Fibrosis Patents: A Case Study of Successful Licensing. LES Nouvelles, 2013, , 21-30.	0.5	1
116	Mapping the Human Genome Biotechnology and the Human Genome: Innovations and Impact Avril D. Woodhead Benjamin J. Barnhart. BioScience, 1989, 39, 402-403.	4.9	0
117	Gene quests and the social ethics of research. Amyloid: the International Journal of Experimental and Clinical Investigation: the Official Journal of the International Society of Amyloidosis, 1994, 1, 283-285.	3.0	0
118	Commentary on "distinguishing genetic from nongenetic medical tests: Some implications for antidiscrimination legislation―(J. S. Alper and J. Beckwith). Science and Engineering Ethics, 1998, 4, 151-154.	2.9	0
119	DNA sequence patents are not in the grave yet. Nature Biotechnology, 2009, 27, 122-122.	17.5	0
120	LeRoy Walters's Legacy of Bioethics in Genetics and Biotechnology Policy. Kennedy Institute of Ethics Journal, 2019, 29, 51-66.	0.5	0
121	How Bioethics Can Inform Policy Decisions About Genetic Enhancement. Philosophy and Medicine, 2008, , 161-198.	0.3	0
122	Dr. Varmus Goes to Washington. American Scientist, 2009, 97, 152.	0.1	0
123	Genomics and Patents. , 2013, , 464-473.		0
124	Ethics of Global Health Photography: A Focus on Being More Human. Health and Human Rights, 2019, 21, 49-62.	1.3	0
125	Biomedical Research Policy and Innovation (1940s–Present). , 0, , 181-196.		0