Julia Abelson

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/3787687/publications.pdf

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96 papers

5,491 citations

33 h-index 72 g-index

98 all docs 98 docs citations 98 times ranked 5051 citing authors

#	Article	IF	CITATIONS
1	How Can Research Organizations More Effectively Transfer Research Knowledge to Decision Makers?. Milbank Quarterly, 2003, 81, 221-248.	4.4	860
2	Deliberations about deliberative methods: issues in the design and evaluation of public participation processes. Social Science and Medicine, 2003, 57, 239-251.	3.8	787
3	Public participation in health care priority setting: A scoping review. Health Policy, 2009, 91, 219-228.	3.0	283
4	Bringing â€~the public' into health technology assessment and coverage policy decisions: From principles to practice. Health Policy, 2007, 82, 37-50.	3.0	203
5	Does deliberation make a difference? Results from a citizens panel study of health goals priority setting. Health Policy, 2003, 66, 95-106.	3.0	198
6	â€The public is too subjective': public involvement at different levels of health-care decision making. Social Science and Medicine, 2002, 54, 1825-1837.	3.8	179
7	Deliberative dialogues as a mechanism for knowledge translation and exchange in health systems decision-making. Social Science and Medicine, 2012, 75, 1938-1945.	3.8	164
8	Patient and public engagement in research and health system decision making: A systematic review of evaluation tools. Health Expectations, 2018, 21, 1075-1084.	2.6	153
9	What is important to continuity in home care?. Social Science and Medicine, 2004, 58, 177-192.	3.8	125
10	Evaluating patient and public involvement in research. BMJ: British Medical Journal, 2018, 363, k5147.	2.3	116
11	Supporting quality public and patient engagement in health system organizations: development and usability testing of the <scp>P</scp> ublic and <scp>P</scp> atient <scp>E</scp> ngagement <scp>E</scp> valuation <scp>T</scp> ool. Health Expectations, 2016, 19, 817-827.	2.6	114
12	"lt all dependsâ€. Conceptualizing public involvement in the context of health technology assessment agencies. Social Science and Medicine, 2010, 70, 1518-1526.	3.8	101
13	Primary Care In Canada: So Much Innovation, So Little Change. Health Affairs, 2001, 20, 116-131.	5.2	100
14	PUBLIC AND PATIENT INVOLVEMENT IN HEALTH TECHNOLOGY ASSESSMENT: A FRAMEWORK FOR ACTION. International Journal of Technology Assessment in Health Care, 2016, 32, 256-264.	0.5	98
15	How Contexts and Issues Influence the Use of Policyâ€Relevant Research Syntheses: A Critical Interpretive Synthesis. Milbank Quarterly, 2013, 91, 604-648.	4.4	92
16	Examining the role of context in the implementation of a deliberative public participation experiment: Results from a Canadian comparative study. Social Science and Medicine, 2007, 64, 2115-2128.	3.8	88
17	What Is Public Deliberation?. Hastings Center Report, 2012, 42, 14-16.	1.0	87
18	Alternatives to Project-specific Consent for Access to Personal Information for Health Research: What Is the Opinion of the Canadian Public?. Journal of the American Medical Informatics Association: JAMIA, 2007, 14, 706-712.	4.4	78

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19	Why and When Should We Use Public Deliberation?. Hastings Center Report, 2012, 42, 17-20.	1.0	75
20	Eliciting ethical and social values in health technology assessment: A participatory approach. Social Science and Medicine, 2011, 73, 135-144.	3.8	72
21	Why do policies change? Institutions, interests, ideas and networks in three cases of policy reform. Health Policy and Planning, 2016, 31, 1200-1211.	2.7	72
22	What does it mean to trust a health system?. Health Policy, 2009, 91, 63-70.	3.0	70
23	Understanding the role of contextual influences on local health-care decision making: case study results from Ontario, Canada. Social Science and Medicine, 2001, 53, 777-793.	3.8	67
24	Are we expecting too much from print media? An analysis of newspaper coverage of the 2002 Canadian healthcare reform debate. Social Science and Medicine, 2006, 63, 89-102.	3.8	64
25	Questioning patient engagement: research scientists' perceptions of the challenges of patient engagement in a cardiovascular research network. Patient Preference and Adherence, 2017, Volume 11, 1573-1583.	1.8	63
26	Participation in health care priority-setting through the eyes of the participants. Journal of Health Services Research and Policy, 2002, 7, 222-229.	1.7	62
27	Enhancing Citizen Engagement in Cancer Screening Through Deliberative Democracy. Journal of the National Cancer Institute, 2013, 105, 380-386.	6.3	56
28	The policy analysis of â€~values talk': lessons from Canadian health reform. Health Policy, 2004, 67, 15-24.	3.0	53
29	Conceptualizing the use of public involvement in health policy decision-making. Social Science and Medicine, 2015, 138, 14-21.	3.8	48
30	Will it make a difference if I show up and share? A citizens' perspective on improving public involvement processes for health system decision-making. Journal of Health Services Research and Policy, 2004, 9, 205-212.	1.7	45
31	Is Reasonable Access What We Want? Implications of, and Challenges to, Current Canadian Policy on Equity in Health Care. International Journal of Health Services, 1993, 23, 629-653.	2.5	38
32	Obtaining public input for healthâ€systems decisionâ€making: Past experiences and future prospects. Canadian Public Administration, 2002, 45, 70-97.	0.9	38
33	Citizens' perspectives on personalized medicine: a qualitative public deliberation study. European Journal of Human Genetics, 2013, 21, 1197-1201.	2.8	38
34	ASSESSING THE IMPACTS OF CITIZEN DELIBERATIONS ON THE HEALTH TECHNOLOGY PROCESS. International Journal of Technology Assessment in Health Care, 2013, 29, 282-289.	0.5	35
35	Moving cautiously: Public involvement and the health technology assessment community. International Journal of Technology Assessment in Health Care, 2011, 27, 43-49.	0.5	34
36	Managing under managed community care: the experiences of clients, providers and managers in Ontario's competitive home care sector. Health Policy, 2004, 68, 359-372.	3.0	33

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37	Media Hyping and the "Herceptin Access Story": An Analysis of Canadian and UK Newspaper Coverage. Healthcare Policy, 2009, 4, e113-28.	0.6	31
38	Supporting the evaluation of public and patient engagement in health system organizations: Results from an implementation research study. Health Expectations, 2019, 22, 1132-1143.	2.6	30
39	Knowledge into action?. Health Policy, 2007, 80, 158-171.	3.0	29
40	The integration of citizens into a science/policy network in genetics: governance arrangements and asymmetry in expertise. Health Expectations, 2011, 14, 261-271.	2.6	28
41	Mental health service delivery in Ontario, Canada: how do policy legacies shape prospects for reform?. Health Economics, Policy and Law, 2007, 2, 363-389.	1.8	27
42	Enabling the participation of marginalized populations: case studies from a health service organization in Ontario, Canada. Health Promotion International, 2017, 32, dav118.	1.8	25
43	From rhetoric to reality: including patient voices in supportive cancer care planning. Health Expectations, 2005, 8, 195-209.	2.6	24
44	Evaluating Patient, Family and Public Engagement in Health Services Improvement and System Redesign. Healthcare Quarterly, 2018, 21, 31-37.	0.7	24
45	The visible politics of the privatization debate in Quebec. Healthcare Policy, 2012, 8, 67-79.	0.6	22
46	Applying priority-setting frameworks: A review of public and vulnerable populations' participation in health-system priority setting. Health Policy, 2020, 124, 133-142.	3.0	21
47	Experiences of Novice Physiotherapists Adapting to Their Role in Acute Care Hospitals. Physiotherapy Canada Physiotherapie Canada, 2005, 57, 145.	0.6	19
48	Development of a framework for effective community engagement in Ontario, Canada. Health Policy, 2011, 101, 59-69.	3.0	18
49	Addressing the affordability of cancer drugs: using deliberative public engagement to inform health policy. Health Research Policy and Systems, 2019, 17, 17.	2.8	18
50	Patient engagement in health technology assessment: what constitutes †meaningful†and how we might get there. Journal of Health Services Research and Policy, 2018, 23, 69-71.	1.7	17
51	Measuring Children's Mental Health in Ontario: Policy Issues and Prospects for Change. Journal of the Canadian Academy of Child and Adolescent Psychiatry, 2018, 27, 88-98.	0.6	17
52	Who is in and who is out? A qualitative analysis of stakeholder participation in priority setting for health in three districts in Uganda. Health Policy and Planning, 2019, 34, 358-369.	2.7	16
53	Joining the conversation: newspaper journalists' views on working with researchers. Healthcare Policy, 2005, 1, 123-39.	0.6	16
54	The Future of Public Deliberation on Health Issues. Hastings Center Report, 2012, 42, 27-29.	1.0	15

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55	Priority setting and equity in COVID-19 pandemic plans: a comparative analysis of 18 African countries. Health Policy and Planning, 2022, 37, 297-309.	2.7	15
56	Developing a Canadian evaluation framework for patient and public engagement in research: study protocol. Research Involvement and Engagement, 2021, 7, 10.	2.9	14
57	Canadians Confront Health Care Reform. Health Affairs, 2004, 23, 186-193.	5.2	13
58	Ethical Challenges Related to Patient Involvement in Health Technology Assessment. International Journal of Technology Assessment in Health Care, 2019, 35, 253-256.	0.5	13
59	In Search of Informed Input: A Systematic Approach to Involving the Public in Community Decision Making. Healthcare Management Forum, 1996, 9, 48-52.	1.4	12
60	Uncertain times: A survey of Canadian women's perspectives toward mammography screening. Preventive Medicine, 2018, 112, 209-215.	3.4	12
61	Members of Minority and Underserved Communities Set Priorities for Health Research. Milbank Quarterly, 2018, 96, 675-705.	4.4	12
62	The value of frameworks as knowledge translation mechanisms to guide community participation practice in Ontario CHCs. Social Science and Medicine, 2015, 142, 223-231.	3.8	11
63	Does moral reasoning influence public values for health care priority setting?: A population-based randomized stated preference survey. Health Policy, 2020, 124, 647-658.	3.0	11
64	An institutional ethnographic analysis of public and patient engagement activities at a national health technology assessment agency. International Journal of Technology Assessment in Health Care, 2021, 37, e37.	0.5	11
65	The impact of COVID‶9 on patient engagement in the health system: Results from a Panâ€Canadian survey of patient, family and caregiver partners. Health Expectations, 2022, 25, 744-753.	2.6	11
66	Designing and Implementing Deliberative Processes for Health Technology Assessment: A Good Practices Report of a Joint HTAi/ISPOR Task Force. Value in Health, 2022, 25, 869-886.	0.3	11
67	Evaluating community deliberations about health research priorities. Health Expectations, 2019, 22, 772-784.	2.6	10
68	â€The problem is small enough, the problem is big enough': a qualitative study of health technology assessment and public policy on drug funding decisions for children. International Journal for Equity in Health, 2020, 19, 45.	3.5	10
69	Primary Care Performance Measurement and Reporting at a Regional Level: Could a Matrix Approach Provide Actionable Information for Policy Makers and Clinicians?. Healthcare Policy, 2016, 12, 33-51.	0.6	10
70	Fielding standardized patients in primary care settings: lessons from a study using unannounced standardized patients to assess preventive care practices. International Journal for Quality in Health Care, 1998, 10, 199-206.	1.8	9
71	Framing Incremental Expansions to Public Health Insurance Systems: The Case of Canadian Pharmacare. Journal of Health Politics, Policy and Law, 2014, 39, 295-330.	1.9	9
72	†I just want to be able to make a choice': Results from citizen deliberations about mammography screening in Ontario, Canada. Health Policy, 2018, 122, 1364-1371.	3.0	9

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73	Uncomfortable trade-offs: Canadian policy makers' perspectives on setting objectives for their health systems. Health Policy, 2017, 121, 9-16.	3.0	8
74	Designing and Implementing Deliberative Processes for Health Technology Assessment: A Good Practices Report of a Joint HTAi/ISPOR Task Force. International Journal of Technology Assessment in Health Care, 2022, 38, .	0.5	8
75	Ethics in Health Technology Assessment: Understanding Health Technologies as Policies. Healthcare Management Forum, 2013, 26, 72-76.	1.4	7
76	Evidence-informed policymaking and policy innovation in a low-income country: does policy network structure matter?. Evidence and Policy, 2018, 14, 381-401.	1.0	7
77	Do assisted living facilities that offer a dementia care program differ from those that do not? A population-level cross-sectional study in Ontario, Canada. BMC Geriatrics, 2021, 21, 463.	2.7	7
78	Association Between Dementia Care Programs in Assisted Living Facilities and Transitions to Nursing Homes in Ontario, Canada: AÂPopulation-Based Cohort Study. Journal of the American Medical Directors Association, 2021, 22, 2115-2120.e6.	2.5	7
79	Understanding the Role of Values in Health Policy Decision-Making From the Perspective of Policy-Makers and Stakeholders: A Multiple-Case Embedded Study in Chile and Colombia. International Journal of Health Policy and Management, 2020, 9, 185-197.	0.9	7
80	The Moral Foundations of Child Health and Social Policies: A Critical Interpretive Synthesis. Children, 2021, 8, 43.	1.5	6
81	Citizen perspectives on the use of publicly reported primary care performance information: Results from citizenâ€patient dialogues in three Canadian provinces. Health Expectations, 2019, 22, 974-982.	2.6	5
82	Une expérience de consultation publique délibérative dans Charlevoix. Recherches Sociographiques, 2004, 45, 77-104.	0.1	4
83	A Fresh Approach to Reform? A Policy Analysis of the Development and Implementation of Ontario's Mental Health and Addictions Strategy. Healthcare Policy, 2019, 14, 29-42.	0.6	4
84	Identifying approaches for synthesizing and summarizing information to support informed citizen deliberations in health policy: a scoping review. Journal of Health Services Research and Policy, 2020, 25, 59-66.	1.7	4
85	Priority measures for publicly reporting primary care performance: Results of public engagement through deliberative dialogues in 3 Canadian provinces. Health Expectations, 2020, 23, 1213-1223.	2.6	4
86	Alternative funding and delivery models. Journal of Ambulatory Care Management, 1993, 16, 19-29.	1.1	3
87	Opportunities and Challenges in the Use of Public Deliberation to Inform Public Health Policies. American Journal of Bioethics, 2009, 9, 24-25.	0.9	3
88	Exploring the conundrum of the new knowledge production regime: an ethnographic case study on the governance and outcomes of a science/policy network in genetics. Science and Public Policy, 2010, 37, 737-750.	2.4	3
89	Supporting women at average risk to make informed decisions about mammography when there is no "right―answer: a qualitative citizen deliberation study. CMAJ Open, 2019, 7, E730-E737.	2.4	2
90	Policy Agenda-Setting and Causal Stories: Examining How Organized Interests redefined the Problem of Refugee Health Policy in Canada. Healthcare Policy, 2020, 15, 116-131.	0.6	2

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91	Ethical and Social Values for Paediatric Health Technology Assessment and Drug Policy. International Journal of Health Policy and Management, 2020, , .	0.9	2
92	Barriers to Equitable Public Participation in Health-System Priority Setting Within the Context of Decentralization: The Case of Vulnerable Women in a Ugandan District. International Journal of Health Policy and Management, 2020, , .	0.9	2
93	Health technology assessment processes: a North-South comparison of the evaluation andÂrecommendation of health technologies in Canada and Chile. International Journal of Health Governance, 2022, 27, 312-328.	1.2	2
94	Engaging the Public Voice in Health Care Decision-Making., 2008, , 123-138.		1
95	Ethical Challenges at the Intersection of Policy and Practice in Humanitarian Contexts. Prehospital and Disaster Medicine, 2017, 32, S48-S49.	1.3	O
96	Diverse Discussion in Public Deliberation on Cancer Drug Funding. Journal of Deliberative Democracy, 2022, 18, .	0.6	0