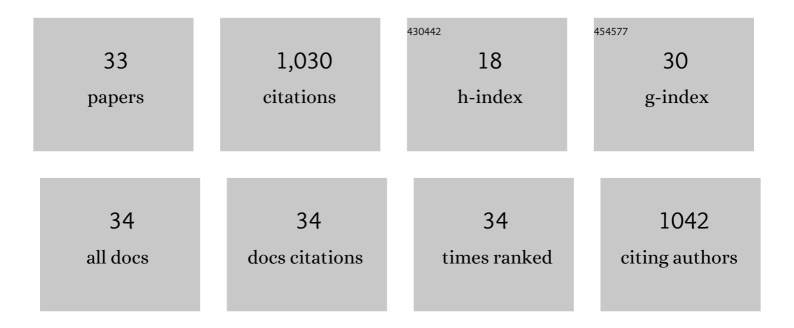
## Susan Bull

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/7051477/publications.pdf Version: 2024-02-01



SUSAN RUU

#	Article	IF	CITATIONS
1	Key criteria for the ethical acceptability of COVID-19 human challenge studies: Report of a WHO Working Group. Vaccine, 2021, 39, 633-640.	1.7	42
2	Vaccine-enhanced disease: case studies and ethical implicationsÂforÂresearch andÂpublic health. Wellcome Open Research, 2021, 6, 154.	0.9	4
3	Equitable data sharing in epidemics and pandemics. BMC Medical Ethics, 2021, 22, 136.	1.0	10
4	SARS-CoV-2 challenge studies: ethics and risk minimisation. Journal of Medical Ethics, 2020, , medethics-2020-106504.	1.0	15
5	"Are we getting the biometric bioethics right?―– the use of biometrics within the healthcare system in Malawi. Global Bioethics, 2020, 31, 67-80.	0.5	3
6	Researcher and study participants' perspectives of consent in clinical studies in four referral hospitals in Vietnam. BMC Medical Ethics, 2020, 21, 4.	1.0	5
7	The ethics of data sharing and biobanking in health research. Wellcome Open Research, 2020, 5, 270.	0.9	7
8	How should assent to research be sought in low income settings? Perspectives from parents and children in Southern Malawi. BMC Medical Ethics, 2019, 20, 32.	1.0	7
9	<i>â€~lt is an entrustment'</i> : Broad consent for genomic research and biobanks in sub‣aharan Africa. Developing World Bioethics, 2019, 19, 9-17.	0.6	42
10	Challenges arising when seeking broad consent for health research data sharing: a qualitative study of perspectives in Thailand. BMC Medical Ethics, 2018, 19, 86.	1.0	18
11	Sharing Individual-Level Health Research Data: Experiences, Challenges and a Research Agenda. Asian Bioethics Review, 2017, 9, 393-400.	0.9	7
12	Trust, Respect, and Reciprocity. Journal of Empirical Research on Human Research Ethics, 2015, 10, 251-263.	0.6	35
13	Sharing Public Health Research Data. Journal of Empirical Research on Human Research Ethics, 2015, 10, 217-224.	0.6	28
14	Views of Ethical Best Practices in Sharing Individual-Level Data From Medical and Public Health Research. Journal of Empirical Research on Human Research Ethics, 2015, 10, 225-238.	0.6	60
15	Research Stakeholders' Views on Benefits and Challenges for Public Health Research Data Sharing in Kenya: The Importance of Trust and Social Relations. PLoS ONE, 2015, 10, e0135545.	1.1	39
16	Sweat, Skepticism, and Uncharted Territory. Journal of Empirical Research on Human Research Ethics, 2015, 10, 239-250.	0.6	37
17	Involving Research Stakeholders in Developing Policy on Sharing Public Health Research Data in Kenya. Journal of Empirical Research on Human Research Ethics, 2015, 10, 264-277.	0.6	46
18	Developing Ethical Practices for Public Health Research Data Sharing in South Africa. Journal of Empirical Research on Human Research Ethics, 2015, 10, 290-301.	0.6	44

Susan Bull

#	Article	IF	CITATIONS
19	Perceived Benefits, Harms, and Views About How to Share Data Responsibly. Journal of Empirical Research on Human Research Ethics, 2015, 10, 278-289.	0.6	45
20	Understandings of genomic research in developing countries: a qualitative study of the views of MalariaGEN participants in Mali. BMC Medical Ethics, 2015, 16, 42.	1.0	23
21	Best Practices for Ethical Sharing of Individual-Level Health Research Data From Low- and Middle-Income Settings. Journal of Empirical Research on Human Research Ethics, 2015, 10, 302-313.	0.6	60
22	Ethical issues in the export, storage and reuse of human biological samples in biomedical research: perspectives of key stakeholders in Ghana and Kenya. BMC Medical Ethics, 2014, 15, 76.	1.0	37
23	Consensus standards for introductory e-learning courses in human participants research ethics. Journal of Medical Ethics, 2014, 40, 426-428.	1.0	6
24	Ethics and Tropical Diseases. , 2014, , 31-39.e1.		0
25	Consent and Community Engagement in Diverse Research Contexts: Reviewing and Developing Research and Practice. Journal of Empirical Research on Human Research Ethics, 2013, 8, 1-18.	0.6	123
26	Tailoring Information Provision and Consent Processes to Research Contexts: The Value of Rapid Assessments. Journal of Empirical Research on Human Research Ethics, 2012, 7, 37-52.	0.6	45
27	Seeking consent to genetic and genomic research in a rural Ghanaian setting: A qualitative study of the MalariaGEN experience. BMC Medical Ethics, 2012, 13, 15.	1.0	97
28	Ensuring Consent to Research is Voluntary: How Far Do We Need to Go?. American Journal of Bioethics, 2011, 11, 27-29.	0.5	22
29	Ethics in collaborative global health research networks. Clinical Ethics, 2009, 4, 165-168.	0.5	17
30	Impact of social stigma on the process of obtaining informed consent for genetic research on podoconiosis: a qualitative study. BMC Medical Ethics, 2009, 10, 13.	1.0	85
31	Engaging publics in biobanking and genetic research governance - a literature review towards informing practice in India. Wellcome Open Research, 0, 6, 5.	0.9	2
32	Engaging publics in biobanking and genetic research governance - a literature review towards informing practice in India. Wellcome Open Research, 0, 6, 5.	0.9	1
33	Respecting values and perspectives in biobanking and genetic research governance: Outcomes of a qualitative study in Bengaluru, India. Wellcome Open Research, 0, 7, 78.	0.9	1