Eva C Winkler

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/1177711/publications.pdf

Version: 2024-02-01

43 papers

1,183 citations

567281 15 h-index 32 g-index

59 all docs 59 docs citations

59 times ranked

1766 citing authors

#	Article	IF	CITATIONS
1	When do we need to care about the caregiver? Supportive care needs, anxiety, and depression among informal caregivers of patients with cancer and cancer survivors. Cancer, 2015, 121, 1513-1519.	4.1	295
2	Patient Involvement in Decisions to Limit Treatment: The Crucial Role of Agreement Between Physician and Patient. Journal of Clinical Oncology, 2009, 27, 2225-2230.	1.6	102
3	GA4GH: International policies and standards for data sharing across genomic research and healthcare. Cell Genomics, 2021, 1, 100029.	6.5	94
4	What Keeps Oncologists From Addressing Palliative Care Early on With Incurable Cancer Patients? An Active Stance Seems Key. Oncologist, 2015, 20, 56-61.	3.7	84
5	Stakeholders' perspectives on biobank-based genomic research: systematic review of the literature. European Journal of Human Genetics, 2015, 23, 1607-1614.	2.8	61
6	High prevalence of moral distress reported by oncologists and oncology nurses in endâ€ofâ€ife decision making. Psycho-Oncology, 2018, 27, 2733-2739.	2.3	58
7	The second patient? Family members of cancer patients and their role in end-of-life decision making. BMC Palliative Care, 2018, 17, 29.	1.8	39
8	The patient-level effect of the cost of Cancer care – financial burden in German Cancer patients. BMC Cancer, 2020, 20, 529.	2.6	30
9	Evaluating a patient's request for life-prolonging treatment: an ethical framework. Journal of Medical Ethics, 2012, 38, 647-651.	1.8	27
10	Ethical assessment of life-prolonging treatment. Lancet Oncology, The, 2011, 12, 720-722.	10.7	26
11	Digitalizing Health Services by Implementing a Personal Electronic Health Record in Germany: Qualitative Analysis of Fundamental Prerequisites From the Perspective of Selected Experts. Journal of Medical Internet Research, 2020, 22, e15102.	4.3	25
12	Applying systems biology to biomedical research and health care: a prÃ@cising definition of systems medicine. BMC Health Services Research, 2017, 17, 761.	2.2	24
13	Do patients and research subjects have a right to receive their genomic raw data? An ethical and legal analysis. BMC Medical Ethics, 2020, 21, 7.	2.4	24
14	A Personal Electronic Health Record: Study Protocol of a Feasibility Study on Implementation in a Real-World Health Care Setting. JMIR Research Protocols, 2017, 6, e33.	1.0	22
15	Complexity of care and strategies of self-management in patients with colorectal cancer. Patient Preference and Adherence, 2017, Volume 11, 731-742.	1.8	21
16	So rare we need to hunt for them: reframing the ethical debate on incidental findings. Genome Medicine, 2015, 7, 83.	8.2	19
17	Patient-Reported Financial Distress in Cancer: A Systematic Review of Risk Factors in Universal Healthcare Systems. Cancers, 2021, 13, 5015.	3.7	18
18	Cancer Patients' Preferences for Quantity or Quality of Life: German Translation and Validation of the Quality and Quantity Questionnaire. Oncology Research and Treatment, 2014, 37, 472-478.	1.2	16

#	Article	IF	CITATIONS
19	Secondary Use of Clinical Data in Data-Gathering, Non-Interventional Research or Learning Activities: Definition, Types, and a Framework for Risk Assessment. Journal of Medical Internet Research, 2021, 23, e26631.	4.3	16
20	Utilizing a Prototype Patient-Controlled Electronic Health Record in Germany: Qualitative Analysis of User-Reported Perceptions and Perspectives. JMIR Formative Research, 2018, 2, e10411.	1.4	16
21	"Rather one more chemo than one less…― Oncologists and Oncology Nurses' Reasons for Aggressive Treatment of Young Adults with Advanced Cancer. Oncologist, 2018, 23, 256-262.	3.7	15
22	The COVID-19 Pandemic and Cancer Patients in Germany: Impact on Treatment, Follow-Up Care and Psychological Burden. Frontiers in Public Health, 2021, 9, 788598.	2.7	14
23	"Often Relatives are the Key […]―–Family Involvement in Treatment Decision Making in Patients with Advanced Cancer Near the End of Life. Oncologist, 2021, 26, e831-e837.	3.7	12
24	Patients' Willingness to Provide Their Clinical Data for Research Purposes and Acceptance of Different Consent Models: Findings From a Representative Survey of Patients With Cancer. Journal of Medical Internet Research, 2022, 24, e37665.	4.3	12
25	Patients' Preferences in Non-Curable Cancer Disease. Oncology Research and Treatment, 2019, 42, 31-34.	1.2	11
26	Assessing patient–caregiver communication in cancer—a psychometric validation of the Cancer Communication Assessment Tool (CCAT-PF) in a German sample. Supportive Care in Cancer, 2014, 22, 2473-2478.	2.2	9
27	Caregivers' role in using a personal electronic health record: a qualitative study of cancer patients and caregivers in Germany. BMC Medical Informatics and Decision Making, 2020, 20, 158.	3.0	9
28	Sekundänutzung klinischer Daten aus der Patientenversorgung fÃ⅓r Forschungszwecke – Eine qualitative Interviewstudie zu Nutzen- und Risikopotenzialen aus Sicht von Expertinnen und Experten fÃ⅓r den deutschen Forschungskontext. , 2022, , 185-210.		8
29	Late decisions about treatment limitation in patients with cancer: empirical analysis of end-of-life practices in a haematology and oncology unit at a German university hospital. ESMO Open, 2020, 5, e000950.	4.5	6
30	Trust and responsibility in molecular tumour boards. Bioethics, 2018, 32, 464-472.	1.4	5
31	Cancer Patients' Preferences for either Quality of Life or a Longer Life Determine Their Willingness to Talk about Forgoing Cancer-Specific Treatment. Oncology Research and Treatment, 2019, 42, 458-465.	1.2	5
32	Researchers' Duty to Share Pre-publication Data: From the Prima Facie Duty to Practice. Law, Governance and Technology Series, 2016, , 309-337.	0.4	5
33	Do Researchers in Empirical Ethics Studies Have a Duty to Act Upon their Findings? Case Study in End-of-Life Decision Making. Journal of Empirical Research on Human Research Ethics, 2019, 14, 438-440.	1.3	3
34	Evaluating requests for physicianâ€assisted suicide. A survey among German oncologists. Cancer Medicine, 2023, 12, 1813-1820.	2.8	3
35	Development and Evaluation of an Ethical Guideline for Decisions to Limit Life-Prolonging Treatment in Advanced Cancer: Protocol for a Monocentric Mixed-Method Interventional Study. JMIR Research Protocols, 2018, 7, e157.	1.0	2
36	Sekundänutzung klinischer Daten in datensammelnden, nicht-interventionellen Forschungs- oder LernaktivitÃæn – Begriff, Studientypen und ethische Herausforderungen. , 2022, , 71-98.		2

#	Article	IF	CITATIONS
37	Is Dupras and Bunnik's Framework for Assessing Privacy Risks in Multi-Omic Research and Databases Still Too Exceptionalist?. American Journal of Bioethics, 2021, 21, 80-82.	0.9	2
38	Ethische Herausforderungen der Genomsequenzierung in der translationalen Forschung und Antworten aus dem EURAT-Projekt. Laboratoriums Medizin, 2014, 38, .	0.6	1
39	Ethical challenges of whole genome sequencing in translational research and answers by the EURAT-project. Laboratoriums Medizin, 2015, 38, .	0.6	1
40	Between Minimal and Greater Than Minimal Risk: How Research Participants and Oncologists Assess Data-Sharing and the Risk of Re-identification in Genomic Research. Philosophy and Technology, 2019, 32, 39-55.	4.3	1
41	Nipping Diseases in the Bud? Ethical and Social Considerations of the Concept of †Disease Interceptionâ€. Public Health Ethics, 2021, 14, 100-108.	1.0	0
42	Development and Evaluation of a Decision Aid to Support Patients' Participatory Decision-Making for Tumor-Specific and Palliative Therapy for Advanced Cancer: Protocol for a Pre-Post Study. JMIR Research Protocols, 2021, 10, e24954.	1.0	0
43	Onkologische Ethik. , 2013, , 479-495.		O