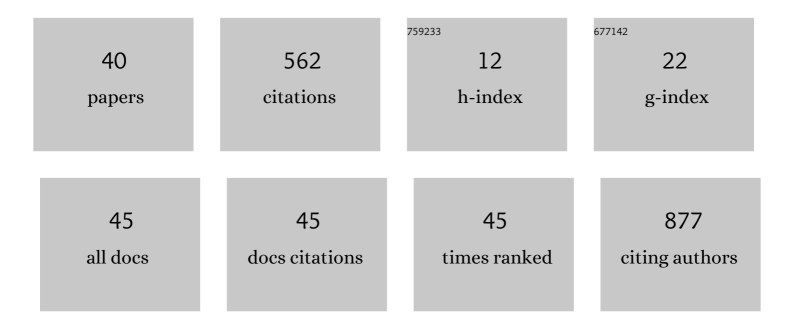
Cinzia Colombo

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/784663/publications.pdf Version: 2024-02-01



#	Article	IF	CITATIONS
1	ldentifying unanswered questions and setting the agenda for future systematic research in Multiple Sclerosis. A worldwide, multi-stakeholder Priority Setting project. Multiple Sclerosis and Related Disorders, 2022, 60, 103688.	2.0	3
2	Critical observations on and suggested ways forward for healthcare communication during COVID-19: pEACH position paper. Patient Education and Counseling, 2021, 104, 217-222.	2.2	41
3	The CORBEL matrix on informed consent in clinical studies: a multidisciplinary approach of Research Infrastructures Building Enduring Life-science Services. BMC Medical Ethics, 2021, 22, 95.	2.4	2
4	The IN-DEEP project "INtegrating and Deriving Evidence, Experiences, Preferences― a web information model on magnetic resonance imaging for people with multiple sclerosis. Journal of Neurology, 2020, 267, 2421-2431.	3.6	1
5	A dynamic web-based decision aid to improve informed choice in organised breast cancer screening. A pragmatic randomised trial in Italy. British Journal of Cancer, 2020, 123, 714-721.	6.4	22
6	Industry funding of patient and health consumer organisations: systematic review with meta-analysis. BMJ, The, 2020, 368, l6925.	6.0	44
7	Sharing individual participant data from clinical studies: a cross-sectional online survey among Italian patient and citizen groups. BMJ Open, 2019, 9, e024863.	1.9	13
8	Choosing Wisely Italy: online survey on opinions and behaviors of 1006 people and 355 volunteers of healthcare advocacy associations. European Journal of Public Health, 2019, 29, 396-401.	0.3	2
9	HIV priorities by Italian AIDS advocacy groups: information on prevention (still) comes first. An on-line survey. Epidemiologia E Prevenzione, 2019, 43, 270-274.	1.1	2
10	Deciding on cystic fibrosis carrier screening: three citizens' juries and an online survey. European Journal of Public Health, 2018, 28, 973-977.	0.3	6
11	Producing an evidenceâ€based treatment information website in partnership with people affected by multiple sclerosis. Health Science Reports, 2018, 1, e24.	1.5	7
12	Personalised informed choice on evidence and controversy on mammography screening: study protocol for a randomized controlled trial. BMC Cancer, 2017, 17, 429.	2.6	10
13	Patient-reported outcomes: nothing without engagement. Acta Oncológica, 2016, 55, 1494-1495.	1.8	1
14	Development and assessment of a website presenting evidence-based information for people with multiple sclerosis: the IN-DEEP project. BMC Neurology, 2016, 16, 30.	1.8	15
15	Citizens' juries could help to guide screening policy. BMJ, The, 2016, 353, i3520.	6.0	0
16	Online health information seeking: how people with multiple sclerosis find, assess and integrate treatment information to manage their health. Health Expectations, 2016, 19, 727-737.	2.6	68
17	A European multi-language initiative to make the general population aware of independent clinical research: the European Communication on Research Awareness Need project. Trials, 2016, 17, 19.	1.6	11
18	Involving a Citizens' Jury in Decisions on Individual Screening for Prostate Cancer. PLoS ONE, 2016, 11, e0143176.	2.5	10

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#	Article	IF	CITATIONS
19	Transparency of funding of patient groups is mandatory but is not enough. BMJ, The, 2014, 349, g6301-g6301.	6.0	О
20	Web Search Behavior and Information Needs of People With Multiple Sclerosis: Focus Group Study and Analysis of Online Postings. Interactive Journal of Medical Research, 2014, 3, e12.	1.4	51
21	Informing Women on Menopause and Hormone Therapy: Know the Menopause a Multidisciplinary Project Involving Local Healthcare System. PLoS ONE, 2013, 8, e85121.	2.5	12
22	Does a consumer training work? a follow-up survey of the PartecipaSalute training programs. Health Research Policy and Systems, 2012, 10, 27.	2.8	6
23	Patient Organizations' Funding from Pharmaceutical Companies: Is Disclosure Clear, Complete and Accessible to the Public? An Italian Survey. PLoS ONE, 2012, 7, e34974.	2.5	33
24	Patient empowerment as a component of health system reforms: rights, benefits and vested interests. Internal and Emergency Medicine, 2012, 7, 183-187.	2.0	23
25	A new informed consent form model for cancer patients: Preliminary results of a prospective study by the Italian Association of Medical Oncology (AIOM). Patient Education and Counseling, 2012, 87, 243-249.	2.2	12
26	Presenting evidence-based health information for people with multiple sclerosis: the IN-DEEP project protocol. BMC Medical Informatics and Decision Making, 2012, 12, 20.	3.0	19
27	Family Burden after Severe Brain Injury. Patient, 2011, 4, 55-65.	2.7	12
28	CIGNOweb.it. Tumori, 2011, 97, 133-135.	1.1	3
29	Role of Hormone Therapy in the Management of Menopause. Obstetrics and Gynecology, 2010, 116, 442.	2.4	2
30	Introduction. Journal of Ambulatory Care Management, 2010, 33, 178-181.	1.1	1
31	Fostering a Strategic Alliance Between Patients' Associations and Health Care Professionals. Journal of Ambulatory Care Management, 2010, 33, 223-230.	1.1	7
32	International Health Consumers in The Cochrane Collaboration. Journal of Ambulatory Care Management, 2010, 33, 182-189.	1.1	12
33	Press coverage of hormone replacement therapy and menopause. European Journal of Obstetrics, Gynecology and Reproductive Biology, 2010, 153, 56-61.	1.1	8
34	Citizens' juries in health care. BMJ: British Medical Journal, 2010, 341, c5141-c5141.	2.3	2
35	Informing women about hormone replacement therapy: the consensus conference statement. BMC Women's Health, 2009, 9, 14.	2.0	17
36	Menopause: Knowledge, attitude and practice among Italian women. Maturitas, 2009, 63, 246-252.	2.4	40

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#	Article	IF	CITATIONS
37	Informing Women About Hormone Replacement Therapy. Cancer Journal (Sudbury, Mass), 2009, 15, 344.	2.0	1
38	PartecipaSalute, an Italian project to involve lay people, patients? associations and scientific-medical representatives in the health debate. Health Expectations, 2007, 10, 194-204.	2.6	30
39	Oncologists' opinions about research ethics committees in Italy: an update, 2004. European Journal of Cancer Prevention, 2006, 15, 91-94.	1.3	5
40	Consumers and health providers working in partnership for the promotion of person-centred health services: a co-produced qualitative evidence synthesis. The Cochrane Library, 0, , .	2.8	8