Carolyn Petersen

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/3418322/publications.pdf

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36	752	14	25
papers	citations	h-index	g-index
37	37	37	869
all docs	docs citations	times ranked	citing authors

#	Article	IF	CITATIONS
1	Defining AMIA's artificial intelligence principles. Journal of the American Medical Informatics Association: JAMIA, 2022, 29, 585-591.	4.4	36
2	Selecting venues for AMIA events and conferences: guiding ethical principles. Journal of the American Medical Informatics Association: JAMIA, 2022, 29, 1319-1322.	4.4	5
3	Use of patient-generated health data for shared decision-making in the clinical environment: ready for prime time. MHealth, 2021, 7, 39-39.	1.6	4
4	Recommendations for the safe, effective use of adaptive CDS in the US healthcare system: an AMIA position paper. Journal of the American Medical Informatics Association: JAMIA, 2021, 28, 677-684.	4.4	46
5	Defining participatory health informatics – a scoping review. Informatics for Health and Social Care, 2021, 46, 234-243.	2.6	16
6	Digital health & Digita	1.3	16
7	A Policy Framework to Support Shared Decision-Making through the Use of Person-Generated Health Data. ACI Open, 2021, 05, e104-e115.	0.5	1
8	Citizen science to further precision medicine: from vision to implementation. JAMIA Open, 2020, 3, 2-8.	2.0	24
9	Special Section on Ethics in Health Informatics. Yearbook of Medical Informatics, 2020, 29, 077-080.	1.0	4
10	Ethical Considerations for Participatory Health through Social Media: Healthcare Workforce and Policy Maker Perspectives. Yearbook of Medical Informatics, 2020, 29, 071-076.	1.0	7
11	From Commercialization to Accountability: Responsible Health Data Collection, Use, and Disclosure for the 21st Century. Applied Clinical Informatics, 2020, 11, 366-373.	1.7	7
12	Evidence-Based Health Informatics as the Foundation for the COVID-19 Response: A Joint Call for Action. Methods of Information in Medicine, 2020, 59, 183-192.	1.2	8
13	Health 'Big Data' Value, Benefit, and Control: The Patient eHealth Equity Gap. Studies in Health Technology and Informatics, 2020, 270, 1123-1127.	0.3	4
14	Beyond TAM and UTAUT: Future directions for HIT implementation research. Journal of Biomedical Informatics, 2019, 100, 103315.	4.3	100
15	User-focused data sharing agreements: a foundation for the genomic future. JAMIA Open, 2019, 2, 402-406.	2.0	5
16	Patients' access to health records. BMJ: British Medical Journal, 2019, 367, l5725.	2.3	28
17	Content Strategy: Writing for Health Consumers on the Web. , 2019, , 275-285.		2
18	Patients as Agents in Behavioral Health Research and Service Provision: Recommendations to Support the Learning Health System. Applied Clinical Informatics, 2019, 10, 841-848.	1.7	15

#	Article	IF	CITATIONS
19	Advance Directives and Code Status Information Exchange: A Consensus Proposal for a Minimum Set of Attributes. Cambridge Quarterly of Healthcare Ethics, 2019, 28, 178-185.	0.8	10
20	Managing Privacy and Data Sharing Through the Use of Health Care Information Fiduciaries. Studies in Health Technology and Informatics, 2019, 265, 157-162.	0.3	2
21	Through Patients' Eyes: Regulation, Technology, Privacy, and the Future. Yearbook of Medical Informatics, 2018, 27, 010-015.	1.0	11
22	Balancing Health Information Exchange and Privacy Governance from a Patient-Centred Connected Health and Telehealth Perspective. Yearbook of Medical Informatics, 2018, 27, 048-054.	1.0	25
23	Patient informaticians: Turning patient voice into patient action. JAMIA Open, 2018, 1, 130-135.	2.0	14
24	Social Media in Health Care: Time for Transparent Privacy Policies and Consent for Data Use and Disclosure. Applied Clinical Informatics, 2018, 09, 856-859.	1.7	7
25	AMIA's code of professional and ethical conduct 2018. Journal of the American Medical Informatics Association: JAMIA, 2018, 25, 1579-1582.	4.4	15
26	Samantha Adams Festschrift: Adamsian Discourseâ€"The Patient, and Everything Else. Applied Clinical Informatics, 2018, 09, 500-502.	1.7	0
27	Balancing Between Privacy and Patient Needs for Health Information in the Age of Participatory Health and Social Media: A Scoping Review. Yearbook of Medical Informatics, 2018, 27, 029-036.	1.0	30
28	The Future of Patient Engagement in the Governance of Shared Data. EGEMS (Washington, DC), 2017, 4, 6.	2.0	9
29	Precision medicine: opportunities, possibilities, and challenges for patients and providers. Journal of the American Medical Informatics Association: JAMIA, 2016, 23, 787-790.	4.4	64
30	Patient-generated health data: a pathway to enhanced long-term cancer survivorship. Journal of the American Medical Informatics Association: JAMIA, 2016, 23, 456-461.	4.4	23
31	Legal and Regulatory Considerations Associated with Use of Patient-Generated Health Data from Social Media and Mobile Health (mHealth) Devices. Applied Clinical Informatics, 2015, 06, 16-26.	1.7	55
32	mHealth: Don't Forget All the Stakeholders in the Business Case. Medicine 2 0, 2015, 4, e4.	2.4	38
33	Sorrell v. IMS Health: issues and opportunities for informaticians. Journal of the American Medical Informatics Association: JAMIA, 2013, 20, 35-37.	4.4	7
34	AMIA's Code of Professional and Ethical Conduct. Journal of the American Medical Informatics Association: JAMIA, 2013, 20, 141-143.	4.4	25
35	A Code of Professional Ethical Conduct for the American Medical Informatics Association: An AMIA Board of Directors Approved White Paper. Journal of the American Medical Informatics Association: JAMIA, 2007, 14, 391-393.	4.4	11
36	Health Information on the Internet. JAMA - Journal of the American Medical Association, 2002, 287, 2713.	7.4	78

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