## Anna Middleton

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/8666863/publications.pdf

Version: 2024-02-01

69 papers 2,974 citations

236925 25 h-index 50 g-index

71 all docs

71 docs citations

times ranked

71

4584 citing authors

| #  | Article   | IF   | Citations |
|----|---|------|-----------|
| 1  | Genetic diagnosis of developmental disorders in the DDD study: a scalable analysis of genome-wide research data. Lancet, The, 2015, 385, 1305-1314.   | 13.7 | 651       |
| 2  | The Global State of the Genetic Counseling Profession. European Journal of Human Genetics, 2019, 27, 183-197.   | 2.8  | 215       |
| 3  | Human Germline Genome Editing. American Journal of Human Genetics, 2017, 101, 167-176.  | 6.2  | 168       |
| 4  | Attitudes of nearly 7000 health professionals, genomic researchers and publics toward the return of incidental results from sequencing research. European Journal of Human Genetics, 2016, 24, 21-29. | 2.8  | 161       |
| 5  | Attitudes of Deaf Adults toward Genetic Testing for Hereditary Deafness. American Journal of Human Genetics, 1998, 63, 1175-1180.   | 6.2  | 153       |
| 6  | Genetic counseling globally: Where are we now?. American Journal of Medical Genetics, Part C: Seminars in Medical Genetics, 2018, 178, 98-107.  | 1.6  | 109       |
| 7  | GA4GH: International policies and standards for data sharing across genomic research and healthcare. Cell Genomics, 2021, 1, 100029.  | 6.5  | 94        |
| 8  | Genetic counselling in the era of genomic medicine. British Medical Bulletin, 2018, 126, 27-36.   | 6.9  | 85        |
| 9  | Prenatal Diagnosis for Inherited Deafnessâ€"What is the Potential Demand?. Journal of Genetic Counseling, 2001, 10, 121-131.  | 1.6  | 82        |
| 10 | Global Public Perceptions of Genomic Data Sharing: What Shapes the Willingness to Donate DNA and Health Data?. American Journal of Human Genetics, 2020, 107, 743-752.                                | 6.2  | 76        |
| 11 | Trust in genomic data sharing among members of the general public in the UK, USA, Canada and Australia. Human Genetics, 2019, 138, 1237-1246.   | 3.8  | 69        |
| 12 | The Global Landscape of Nursing and Genomics. Journal of Nursing Scholarship, 2018, 50, 249-256.  | 2.4  | 59        |
| 13 | Attitudes of publics who are unwilling to donate DNA data for research. European Journal of Medical Genetics, 2019, 62, 316-323.  | 1.3  | 53        |
| 14 | Policy challenges of clinical genome sequencing. BMJ, The, 2013, 347, f6845-f6845.  | 6.0  | 50        |
| 15 | APPLaUD: access for patients and participants to individual level uninterpreted genomic data. Human Genomics, 2018, 12, 7.  | 2.9  | 45        |
| 16 | Direct-to-consumer genetic testing: where and how does genetic counseling fit?. Personalized Medicine, 2017, 14, 249-257.   | 1.5  | 44        |
| 17 | Preferences for communication in clinic from deaf people: a crossâ€sectional study. Journal of Evaluation in Clinical Practice, 2010, 16, 811-817.  | 1.8  | 41        |
| 18 | Demonstrating trustworthiness when collecting and sharing genomic data: public views across 22 countries. Genome Medicine, 2021, 13, 92.  | 8.2  | 39        |

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|----|---|------|-----------|
| 19 | Potential research participants support the return of raw sequence data. Journal of Medical Genetics, 2015, 52, 571-574.  | 3.2  | 38        |
| 20 | Towards equitable and trustworthy genomics research. EBioMedicine, 2022, 76, 103879.  | 6.1  | 34        |
| 21 | The role of genetic counsellors in genomic healthcare in the United Kingdom: a statement by the Association of Genetic Nurses and Counsellors. European Journal of Human Genetics, 2017, 25, 659-661. | 2.8  | 32        |
| 22 | Increasing nursing capacity in genomics: Overview of existing global genomics resources. Nurse Education Today, 2018, 69, 53-59.  | 3.3  | 32        |
| 23 | Genomic variant sharing: a position statement. Wellcome Open Research, 2019, 4, 22.   | 1.8  | 31        |
| 24 | Society and personal genome data. Human Molecular Genetics, 2018, 27, R8-R13.   | 2.9  | 29        |
| 25 | Members of the public in the USA, UK, Canada and Australia expressing genetic exceptionalism say they are more willing to donate genomic data. European Journal of Human Genetics, 2020, 28, 424-434. | 2.8  | 29        |
| 26 | Genetic counselors and Genomic Counseling in the United Kingdom. Molecular Genetics & Samp; Genomic Medicine, 2015, 3, 79-83.   | 1.2  | 28        |
| 27 | Willingness to donate genomic and other medical data: results from Germany. European Journal of Human Genetics, 2020, 28, 1000-1009.  | 2.8  | 28        |
| 28 | Communicating in a healthcare setting with people who have hearing loss. BMJ: British Medical Journal, 2010, 341, c4672-c4672.  | 2.3  | 26        |
| 29 | †Your DNA, Your Say': global survey gathering attitudes toward genomics: design, delivery and methods. Personalized Medicine, 2018, 15, 311-318.  | 1.5  | 26        |
| 30 | Position statement on opportunistic genomic screening from the Association of Genetic Nurses and Counsellors (UK and Ireland). European Journal of Human Genetics, 2014, 22, 955-956.                 | 2.8  | 25        |
| 31 | A Roadmap for Global Acceleration of Genomics Integration Across Nursing. Journal of Nursing Scholarship, 2020, 52, 329-338.  | 2.4  | 24        |
| 32 | Returning genome sequences to research participants: Policy and practice. Wellcome Open Research, 2017, 2, 15.  | 1.8  | 24        |
| 33 | Stakeholders in psychiatry and their attitudes toward receiving pertinent and incident findings in genomic research. American Journal of Medical Genetics, Part A, 2017, 173, 2649-2658.              | 1.2  | 20        |
| 34 | No expectation to share incidental findings in genomic research. Lancet, The, 2015, 385, 1289-1290.   | 13.7 | 19        |
| 35 | Whose Deaf Genes Are They Anyway?: The Deaf Community's Challenge to Legislation on Embryo Selection. Sign Language Studies, 2010, 10, 155-169.   | 0.3  | 18        |
| 36 | Report from the UK and Eire Association of Genetic Nurses and Counsellors (AGNC) Supervision Working Group on Genetic Counselling Supervision. Journal of Genetic Counseling, 2007, 16, 127-142.      | 1.6  | 17        |

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|----|--|------|-----------|
| 37 | Reflections on the Experience of Counseling Supervision by a Team of Genetic Counselors from the UK. Journal of Genetic Counseling, 2007, 16, 143-155.   | 1.6  | 17        |
| 38 | Empirical research on the ethics of genomic research. American Journal of Medical Genetics, Part A, 2013, 161, 2099-2101.  | 1.2  | 17        |
| 39 | Australians' perspectives on support around use of personal genomic testing: Findings from the Genioz study. European Journal of Medical Genetics, 2019, 62, 290-299.                              | 1.3  | 17        |
| 40 | Genetics in the 21st Century: Implications for patients, consumers and citizens. F1000Research, 2017, 6, 2020.   | 1.6  | 15        |
| 41 | Views, Knowledge, and Beliefs about Genetics and Genetic Counseling among Deaf People. Sign Language Studies, 2010, 10, 170-196.   | 0.3  | 14        |
| 42 | Australians' views on personal genomic testing: focus group findings from the Genioz study. European Journal of Human Genetics, 2018, 26, 1101-1112.   | 2.8  | 14        |
| 43 | Australians' views and experience of personal genomic testing: survey findings from the Genioz study. European Journal of Human Genetics, 2019, 27, 711-720.                                       | 2.8  | 14        |
| 44 | A Maturity Matrix for Nurse Leaders to Facilitate and Benchmark Progress in Genomic Healthcare Policy, Infrastructure, Education, and Delivery. Journal of Nursing Scholarship, 2020, 52, 583-592. | 2.4  | 14        |
| 45 | A roadmap for restoring trust in Big Data. Lancet Oncology, The, 2018, 19, 1014-1015.  | 10.7 | 13        |
| 46 | Genetics in the 21st Century: Implications for patients, consumers and citizens. F1000Research, 2017, 6, 2020.   | 1.6  | 13        |
| 47 | Point of View: An evolution from genetic counselling to genomic counselling. European Journal of Medical Genetics, 2019, 62, 288-289.  | 1.3  | 12        |
| 48 | Popular culture and genetics; friend, foe or something more complex?. European Journal of Medical Genetics, 2019, 62, 368-375.   | 1.3  | 12        |
| 49 | A pilot study of inhaled dry-powder mannitol during cystic fibrosis-related pulmonary exacerbation.<br>European Respiratory Journal, 2015, 45, 541-544.  | 6.7  | 11        |
| 50 | Your DNA, Your Say. New Bioethics, 2017, 23, 74-80.  | 1.1  | 11        |
| 51 | Socialising the genome. Lancet, The, 2017, 389, 1603-1604.   | 13.7 | 10        |
| 52 | The preferences of potential stakeholders in psychiatric genomic research regarding consent procedures and information delivery. European Psychiatry, 2019, 55, 29-35.                             | 0.2  | 10        |
| 53 | Communication about DTC Testing: Commentary on a †Family Experience of Personal Genomics'. Journal of Genetic Counseling, 2012, 21, 392-398.   | 1.6  | 9         |
| 54 | Should doctors have a legal duty to warn relatives of their genetic risks? Lancet, The, 2019, 394, 2133-2135.  | 13.7 | 9         |

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|----|--|-----|-----------|
| 55 | From Expectations to Experiences: Consumer Autonomy and Choice in Personal Genomic Testing. AJOB Empirical Bioethics, 2020, 11, 63-76.   | 1.6 | 9         |
| 56 | Return of genomic results does not motivate intentÂtoÂparticipate in research for all: Perspectives across 22 countries. Genetics in Medicine, 2022, 24, 1120-1129.  | 2.4 | 8         |
| 57 | Providing a Transcultural Genetic Counseling Service in the UK. Journal of Genetic Counseling, 2007, 16, 567-582.  | 1.6 | 7         |
| 58 | Genomic variant sharing: a position statement. Wellcome Open Research, 0, 4, 22.   | 1.8 | 7         |
| 59 | Tailoring genetic information and services to clients' culture, knowledge and language level. Nursing Standard (Royal College of Nursing (Great Britain): 1987), 2005, 20, 52-56.  | 0.1 | 6         |
| 60 | Engaged genomic science produces better and fairer outcomes: an engagement framework for engaging and involving participants, patients and publics in genomics research and healthcare implementation. Wellcome Open Research, 2021, 6, 311. | 1.8 | 6         |
| 61 | Clause 14(4)(9) of embryo bill should be amended or deleted. BMJ: British Medical Journal, 2008, 336, 976.1-976.   | 2.3 | 5         |
| 62 | Professional duties are now considered legal duties of care within genomic medicine. European Journal of Human Genetics, 2020, 28, 1301-1304.  | 2.8 | 5         |
| 63 | Attitudes of Costa Rican individuals towards donation of personal genetic data for research. Personalized Medicine, 2021, 18, 141-152.   | 1.5 | 4         |
| 64 | The expectations and realities of nutrigenomic testing in australia: A qualitative study. Health Expectations, 2021, 24, 670-686.  | 2.6 | 3         |
| 65 | Editorial on Supervision. Journal of Genetic Counseling, 2007, 16, 123-125.  | 1.6 | 2         |
| 66 | Public trust and genomic medicine in Canada and the UK. Wellcome Open Research, 2021, 6, 124.  | 1.8 | 2         |
| 67 | World congress on genetic counselling. European Journal of Medical Genetics, 2019, 62, 287.  | 1.3 | 1         |
| 68 | Public trust and genomic medicine in Canada and the UK. Wellcome Open Research, 2021, 6, 124.  | 1.8 | 1         |
| 69 | A public backlash towards genomics is a risk all of us working in genomics must share. Lancet<br>Regional Health - Europe, The, 2022, 15, 100347.  | 5.6 | 1         |