Beverly M Yashar

List of Publications by Year in descending order

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Version: 2024-02-01

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	50	2,016	21		43
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#	Article	lF	CITATIONS
1	A report of the AGCPD task force to evaluate associations between select admissions requirements, demographics, and performance on ABGC certification examination. Journal of Genetic Counseling, 2022, 31, 302-315.	1.6	8
2	Improving genderâ€affirming care in genetic counseling: Using educational tools that amplify transgender and/or gender nonâ€binary community voices. Journal of Genetic Counseling, 2022, 31, 1102-1112.	1.6	8
3	Searching for Answers: Information-Seeking by Young People At-Risk for Huntington's Disease. Journal of Huntington's Disease, 2022, , 1-10.	1.9	O
4	Rewards and challenges of parenting a child with Down syndrome: a qualitative study of fathers' perceptions. Disability and Rehabilitation, 2021, 43, 3562-3573.	1.8	13
5	The introduction of genetic counseling in Ethiopia: Results of a training workshop and lessons learned. PLoS ONE, 2021, 16, e0255278.	2.5	7
6	Graduate training during the COVIDâ€19 pandemic: North American genetic counseling students' challenges, intolerance of uncertainty, and psychological wellâ€being. Journal of Genetic Counseling, 2021, 30, 1325-1335.	1.6	6
7	Improving access to cancer genetic services: perspectives of high-risk clients in a community-based setting. Journal of Community Genetics, 2020, 11, 119-123.	1.2	O
8	Optimizing efficiency and skill utilization: Analysis of genetic counselors' attitudes regarding delegation in a clinical setting. Journal of Genetic Counseling, 2020, 29, 67-77.	1.6	5
9	Development of a decision support tool in pediatric Differences/Disorders of Sex Development. Seminars in Pediatric Surgery, 2019, 28, 150838.	1.1	20
10	Primary care physicians' understanding and utilization of pediatric exome sequencing results. Journal of Genetic Counseling, 2019, 28, 1130-1138.	1.6	6
11	Gender destinies: assigning gender in Disorders of Sex Developmentâ€Intersex clinics. Sociology of Health and Illness, 2019, 41, 1520-1534.	2.1	13
12	Certified Nurse-Midwives' Experiences With Provision of Prenatal Genetic Screening. Journal of Perinatal and Neonatal Nursing, 2019, 33, E3-E14.	0.7	2
13	Evaluating and improving the implementation of a community-based hereditary cancer screening program. Journal of Community Genetics, 2019, 10, 51-60.	1.2	8
14	Genetic testing impacts the utility of prospective familial screening in hypertrophic cardiomyopathy through identification of a nonfamilial subgroup. Genetics in Medicine, 2018, 20, 69-75.	2.4	33
15	"My Plate is Full― Reasons for Declining a Genetic Evaluation of Hearing Loss. Journal of Genetic Counseling, 2018, 27, 597-607.	1.6	6
16	Physician Experiences and Understanding of Genomic Sequencing in Oncology. Journal of Genetic Counseling, 2018, 27, 187-196.	1.6	24
17	Preadoption Genetic Testing: Social workers' decision-making process. Adoption Quarterly, 2018, 21, 141-160.	1.0	1
18	Does Patient-centered Care Change Genital Surgery Decisions? The Strategic Use of Clinical Uncertainty in Disorders of Sex Development Clinics. Journal of Health and Social Behavior, 2018, 59, 520-535.	4.8	46

#	Article	IF	Citations
19	A Qualitative Study of Anticipated Decision Making around Type 2 Diabetes Genetic Testing: the Role of Scientifically Concordant and Discordant Expectations. Journal of Genetic Counseling, 2017, 26, 469-479.	1.6	2
20	Nonâ€invasive prenatal screening for trisomy 21: Consumers' perspectives. American Journal of Medical Genetics, Part A, 2016, 170, 375-385.	1.2	7
21	Parents' Perspectives on Variants of Uncertain Significance from Chromosome Microarray Analysis. Journal of Genetic Counseling, 2016, 25, 101-111.	1.6	42
22	Expanding the genetic counseling workforce: program directors' views on increasing the size of genetic counseling graduate programs. Genetics in Medicine, 2016, 18, 842-849.	2.4	54
23	The Lived Experience of MRKH: Sharing Health Information with Peers. Journal of Pediatric and Adolescent Gynecology, 2016, 29, 154-158.	0.7	43
24	Genetics educational needs in China: physicians' experience and knowledge of genetic testing. Genetics in Medicine, 2015, 17, 757-760.	2.4	11
25	Presented Abstracts from the Thirty Fourth Annual Education Conference of the National Society of Genetic Counselors (Pittsburgh, PA, October 2015). Journal of Genetic Counseling, 2015, 24, 1044-1143.	1.6	1
26	Family Communication in a Population at Risk for Hypertrophic Cardiomyopathy. Journal of Genetic Counseling, 2015, 24, 336-348.	1.6	21
27	Peering into a Chilean Black Box: Parental Storytelling in Pediatric Genetic Counseling. Journal of Genetic Counseling, 2013, 22, 805-816.	1.6	1
28	9p partial monosomy and disorders of sex development: Review and postulation of a pathogenetic mechanism. American Journal of Medical Genetics, Part A, 2013, 161, 1882-1896.	1.2	24
29	Introduction to the Special Issue on Genetic Counseling: A Global Perspective. Journal of Genetic Counseling, 2013, 22, 685-687.	1.6	9
30	Ehlers–Danlos syndrome, hypermobility type: A characterization of the patients' lived experience. American Journal of Medical Genetics, Part A, 2013, 161, 2981-2988.	1.2	64
31	Quality of Life and Autonomy in Emerging Adults with Earlyâ€Onset Neuromuscular Disorders. Journal of Genetic Counseling, 2012, 21, 713-725.	1.6	9
32	Mutations in <i>RPGR </i> and <i>RP2 </i> Account for 15% of Males with Simplex Retinal Degenerative Disease., 2012, 53, 8232.		108
33	Direct-to-consumer genetic testing: An assessment of genetic counselors' knowledge and beliefs. Genetics in Medicine, 2011, 13, 325-332.	2.4	61
34	Perceptions of Licensure: A Survey of Michigan Genetic Counselors. Journal of Genetic Counseling, 2009, 18, 357-365.	1.6	4
35	Molecular Testing for Hereditary Retinal Disease as Part of Clinical Care. JAMA Ophthalmology, 2007, 125, 252.	2.4	37
36	An unusual X-linked retinoschisis phenotype and biochemical characterization of the W112C RS1 mutation. Vision Research, 2006, 46, 3845-3852.	1.4	16

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37	Meta-analysis of genome scans of age-related macular degeneration. Human Molecular Genetics, 2005, 14, 2257-2264.	2.9	224
38	Toll-like receptor 4 variant D299G is associated with susceptibility to age-related macular degeneration. Human Molecular Genetics, 2005, 14, 1449-1455.	2.9	177
39	Association of Apolipoprotein E Alleles with Susceptibility to Age-Related Macular Degeneration in a Large Cohort from a Single Center. Investigative Ophthalmology and Visual Science, 2004, 45, 1306-1310.	3.3	129
40	The Interface Between the Practice of Medical Genetics and Human Genetic Research: What Every Genetic Counselor Needs to Know. Journal of Genetic Counseling, 2004, 13, 351-368.	1.6	10
41	Age-Related Macular Degeneration: A High-Resolution Genome Scan for Susceptibility Loci in a Population Enriched for Late-Stage Disease. American Journal of Human Genetics, 2004, 74, 482-494.	6.2	157
42	Clinical studies of X-linked retinitis pigmentosa in three Swedish families with newly identified mutations in the RP2 and RPGR-ORF15 genes. Ophthalmic Genetics, 2003, 24, 215-223.	1.2	25
43	Retinal Histopathology of an XLRP Carrier with a Mutation in the RPGR Exon ORF15. Experimental Eye Research, 2002, 75, 431-443.	2.6	37
44	A Comprehensive Mutation Analysis of RP2 and RPGR in a North American Cohort of Families with X-Linked Retinitis Pigmentosa. American Journal of Human Genetics, 2002, 70, 1545-1554.	6.2	224
45	DNA-Sequence Patenting: National Society of Genetic Counselors (NSGC) Position Paper. Journal of Genetic Counseling, 2002, 11, 241-243.	1.6	0
46	Microarray analysis of gene expression in the aging human retina. Investigative Ophthalmology and Visual Science, 2002, 43, 2554-60.	3.3	79
47	Evaluation of the ELOVL4 gene in patients with age-related macular degeneration. Ophthalmic Genetics, 2001, 22, 233-239.	1.2	53
48	Searching for Genotype-Phenotype Correlations in X-Linked Juvenile Retinoschisis., 2001,, 45-53.		8
49	Novel mutations in XLRS1 causing retinoschisis, including first evidence of putative leader sequence change. Human Mutation, 1999, 14, 423-427.	2.5	40
50	Dynamics and organization of MAP kinase signal pathways. Molecular Reproduction and Development, 1995, 42, 477-485.	2.0	133