Jane S Green

List of Publications by Year in descending order

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Version: 2024-02-01



#	Article	IF	CITATIONS
1	A dominant <i>RAD51C</i> pathogenic splicing variant predisposes to breast and ovarian cancer in the Newfoundland population due to founder effect. Molecular Genetics & Genomic Medicine, 2020, 8, e1070.	1.2	6
2	The genetic architecture of Stargardt macular dystrophy (STGD1): a longitudinal 40-year study in a genetic isolate. European Journal of Human Genetics, 2020, 28, 925-937.	2.8	10
3	A novel pathogenic missense ADAMTS17 variant that impairs secretion causes Weill-Marchesani Syndrome with variably dysmorphic hand features. Scientific Reports, 2020, 10, 10827.	3.3	13
4	The long-term survival characteristics of a cohort of colorectal cancer patients and baseline variables associated with survival outcomes with or without time-varying effects. BMC Medicine, 2019, 17, 150.	5.5	32
5	Novel Usher syndrome pathogenic variants identified in cases with hearing and vision loss. BMC Medical Genetics, 2019, 20, 68.	2.1	10
6	Screening of BMPR1a for pathogenic mutations in familial colorectal cancer type X families from Newfoundland. Familial Cancer, 2018, 17, 205-208.	1.9	2
7	Impact of colonoscopic screening in Familial Colorectal Cancer Type X. Molecular Genetics & Genomic Medicine, 2018, 6, 1021-1030.	1.2	10
8	Evidence for <i>GALNT12</i> as a moderate penetrance gene for colorectal cancer. Human Mutation, 2018, 39, 1092-1101.	2.5	20
9	Germline <scp>INDEL</scp> s and <scp>CNV</scp> s in a cohort of colorectal cancer patients: their characteristics, associations with relapseâ€free survival time, and potential timeâ€varying effects on the risk of relapse. Cancer Medicine, 2017, 6, 1220-1232.	2.8	14
10	Novel 25 kb Deletion of MERTK Causes Retinitis Pigmentosa With Severe Progression. , 2017, 58, 1736.		17
11	Structured assessment and followup for patients with hereditary kidney tumour syndromes. Canadian Urological Association Journal, 2016, 10, 214.	0.6	12
12	No associations of a set of SNPs in the Vascular Endothelial Growth Factor (VEGF) and Matrix Metalloproteinase (MMP) genes with survival of colorectal cancer patients. Cancer Medicine, 2016, 5, 2221-2231.	2.8	6
13	Genetic structure of the Newfoundland and Labrador population: founder effects modulate variability. European Journal of Human Genetics, 2016, 24, 1063-1070.	2.8	22
14	MC-100â€Review of a large family with schwannomatosis identifies an expanded phenotype. Journal of Medical Genetics, 2015, 52, A1.1-A1.	3.2	0
15	A Survival Association Study of 102 Polymorphisms Previously Associated with Survival Outcomes in Colorectal Cancer. BioMed Research International, 2015, 2015, 1-9.	1.9	5
16	Community engagement with genetics: public perceptions and expectations about genetics research. Health Expectations, 2015, 18, 1413-1425.	2.6	30
17	A genome wide association study on Newfoundland colorectal cancer patients' survival outcomes. Biomarker Research, 2015, 3, 6	6.8	17
18	Public Attitudes About Genetic Testing in the Newborn Period. JOGNN - Journal of Obstetric, Gynecologic, and Neonatal Nursing, 2012, 41, 191-200.	0.5	28

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19	Impact of Gender and Parent of Origin on the Phenotypic Expression of Hereditary Nonpolyposis Colorectal Cancer in a Large Newfoundland Kindred With a Common MSH2 Mutation. Diseases of the Colon and Rectum, 2002, 45, 1223-1232.	1.3	48
20	Mutations in MKKS cause Bardet-Biedl syndrome. Nature Genetics, 2000, 26, 15-16.	21.4	256
21	Mutations in MKKS cause obesity, retinal dystrophy and renal malformations associated with Bardet-Biedl syndrome. Nature Genetics, 2000, 26, 67-70.	21.4	311
22	Canadian Bardet-Biedl syndrome family reduces the critical region of BBS3 (3p) and presents with a variable phenotype. American Journal of Medical Genetics Part A, 1998, 78, 461-467.	2.4	49
23	Common ancestral mutations in theMEN1 gene is likely responsible for the prolactinoma variant of MEN1 (MEN1Burin) in four kindreds from Newfoundland. Human Mutation, 1998, 11, 264-269.	2.5	120
24	Common ancestral mutations in the MEN1 gene is likely responsible for the prolactinoma variant of MEN1 (MEN1Burin) in four kindreds from Newfoundland. Human Mutation, 1998, 11, 264-269.	2.5	8
25	Mutations of a mutS homolog in hereditary nonpolyposis colorectal cancer. Cell, 1993, 75, 1215-1225.	28.9	2,195