## Joanne Ngeow

List of Publications by Year in descending order

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#	Article	lF	CITATIONS
1	Cancer Risks Associated With <i>BRCA1</i> and <i>BRCA2</i> Pathogenic Variants. Journal of Clinical Oncology, 2022, 40, 1529-1541.	1.6	90
2	Polygenic risk modeling for prediction of epithelial ovarian cancer risk. European Journal of Human Genetics, 2022, 30, 349-362.	2.8	23
3	Studying Kidney Diseases Using Organoid Models. Frontiers in Cell and Developmental Biology, 2022, 10, 845401.	3.7	9
4	Male breast cancer: a Singapore perspective. ANZ Journal of Surgery, 2022, , .	0.7	0
5	Overlap of high-risk individuals predicted by family history, and genetic and non-genetic breast cancer risk prediction models: implications for risk stratification. BMC Medicine, 2022, 20, 150.	5.5	9
6	Association between Breast Cancer Polygenic Risk Score and Chemotherapy-Induced Febrile Neutropenia: Null Results. Cancers, 2022, 14, 2714.	3.7	2
7	Oh GxE! The Complexity of Body Mass Index and Colon Cancer Risk. Journal of the National Cancer Institute, 2021, 113, 5-6.	6.3	3
8	Investigation into the origins of an ancient BRCA1 founder mutation identified among Chinese families in Singapore. International Journal of Cancer, 2021, 148, 637-645.	5.1	5
9	Maternal and fetal outcomes in phaeochromocytoma and pregnancy: a multicentre retrospective cohort study and systematic review of literature. Lancet Diabetes and Endocrinology,the, 2021, 9, 13-21.	11.4	37
10	Molecular connections between circadian rhythm and genome maintenance pathways. Endocrine-Related Cancer, 2021, 28, R55-R66.	3.1	13
11	Cost-effectiveness of olaparib maintenance therapy when used with and without restriction by <i>BRCA1/2</i> mutation status for platinum-sensitive relapsed ovarian cancer. Expert Review of Pharmacoeconomics and Outcomes Research, 2021, 21, 441-448.	1.4	7
12	Impact of Variant Reclassification in Cancer Predisposition Genes on Clinical Care. JCO Precision Oncology, 2021, 5, 577-584.	3.0	18
13	CDKN2A germline alterations and the relevance of genotype-phenotype associations in cancer predisposition. Hereditary Cancer in Clinical Practice, 2021, 19, 21.	1.5	36
14	Clinical implementation of an oncologyâ€specific family health history risk assessment tool. Hereditary Cancer in Clinical Practice, 2021, 19, 20.	1.5	2
15	DNA methylation and breast cancer-associated variants. Breast Cancer Research and Treatment, 2021, 188, 713-727.	2.5	7
16	Evaluation of treatment effects in patients with endometrial cancer and <i>POLE</i> mutations: An individual patient data metaâ€analysis. Cancer, 2021, 127, 2409-2422.	4.1	62
17	Management of individuals with germline variants in PALB2: a clinical practice resource of the American College of Medical Genetics and Genomics (ACMG). Genetics in Medicine, 2021, 23, 1416-1423.	2.4	34
18	Understanding patients' views and willingness toward the use of telehealth in a cancer genetics service in Asia. Journal of Genetic Counseling, 2021, 30, 1658-1670.	1.6	5

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19	Telomere biology disorders. Npj Genomic Medicine, 2021, 6, 36.	3.8	29
20	The predictive ability of the 313 variant–based polygenic risk score for contralateral breast cancer risk prediction in women of European ancestry with a heterozygous BRCA1 or BRCA2 pathogenic variant. Genetics in Medicine, 2021, 23, 1726-1737.	2.4	16
21	Homologous Recombination Deficiency: Cancer Predispositions and Treatment Implications. Oncologist, 2021, 26, e1526-e1537.	3.7	53
22	The Skin in Cowden Syndrome. Frontiers in Medicine, 2021, 8, 658842.	2.6	7
23	How practice setting affects family physicians' views on genetic screening: a qualitative study. BMC Family Practice, 2021, 22, 141.	2.9	2
24	Variation in the risk of colorectal cancer in families with Lynch syndrome: a retrospective cohort study. Lancet Oncology, The, 2021, 22, 1014-1022.	10.7	58
25	A Multiplex Thyroid-Specific Assay for Quantification of Circulating Thyroid Cell-Free RNA in Plasma of Thyroid Cancer Patients. Frontiers in Genetics, 2021, 12, 721832.	2.3	2
26	Spectrum of Germline Mutations Within Fanconi Anemia–Associated Genes Across Populations of Varying Ancestry. JNCI Cancer Spectrum, 2021, 5, .	2.9	3
27	Identifying ataxiaâ€ŧelangiectasia in cancer patients: Novel insights from an interesting case and review of literature. Clinical Case Reports (discontinued), 2021, 9, 995-1009.	0.5	6
28	Experience and Perceptions of a Family Health History Risk Assessment Tool among Multi-Ethnic Asian Breast Cancer Patients. Journal of Personalized Medicine, 2021, 11, 1046.	2.5	0
29	Germline breast cancer susceptibility genes, tumor characteristics, and survival. Genome Medicine, 2021, 13, 185.	8.2	3
30	Evaluating empowerment in genetic counseling using patientâ€reported outcomes. Clinical Genetics, 2020, 97, 246-256.	2.0	19
31	Understanding the Psychological Impact of COVID-19 Pandemic on Patients With Cancer, Their Caregivers, and Health Care Workers in Singapore. JCO Global Oncology, 2020, 6, 1494-1509.	1.8	95
32	European polygenic risk score for prediction of breast cancer shows similar performance in Asian women. Nature Communications, 2020, 11, 3833.	12.8	88
33	An in-depth exploration of the post-test informational needs of BRCA1 and BRCA2 pathogenic variant carriers in Asia. Hereditary Cancer in Clinical Practice, 2020, 18, 22.	1.5	8
34	Multiple neoplasia in a patient with Gitelman syndrome harboring germline monoallelic MUTYH mutation. Npj Genomic Medicine, 2020, 5, 39.	3.8	3
35	Early-onset breast cancer in a woman with a germline mobile element insertion resulting in BRCA2 disruption: a case report. Human Genome Variation, 2020, 7, 24.	0.7	3
36	The management of BRCA1 and BRCA2 carriers in Singapore. Chinese Clinical Oncology, 2020, 9, 62-62.	1.2	6

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37	Functional characterisation guides classification of novel BAP1 germline variants. Npj Genomic Medicine, 2020, 5, 50.	3.8	3
38	Predictive Testing for Tumor Predisposition Syndromes in Pediatric Relatives: An Asian Experience. Frontiers in Pediatrics, 2020, 8, 568528.	1.9	1
39	Emerging functions of Fanconi anemia genes in replication fork protection pathways. Human Molecular Genetics, 2020, 29, R158-R164.	2.9	17
40	Factors shaping atâ€risk individuals' decisions to undergo genetic testing for cancer in Asia. Health and Social Care in the Community, 2020, 28, 1569-1577.	1.6	7
41	Biallelic NF1 inactivation in high grade serous ovarian cancers from patients with neurofibromatosis type 1. Familial Cancer, 2020, 19, 353-358.	1.9	3
42	Use of telephone intake for family history taking at a cancer genetics service in Asia. Journal of Genetic Counseling, 2020, 29, 1192-1199.	1.6	3
43	Characterization of the Cancer Spectrum in Men With Germline <i>BRCA1</i> and <i>BRCA2</i> Pathogenic Variants. JAMA Oncology, 2020, 6, 1218.	7.1	48
44	Missense PALB2 germline variant disrupts nuclear localization of PALB2 in a patient with breast cancer. Familial Cancer, 2020, 19, 123-131.	1.9	3
45	PTEN in Hereditary and Sporadic Cancer. Cold Spring Harbor Perspectives in Medicine, 2020, 10, a036087.	6.2	28
46	Comparison of Pheochromocytoma-Specific Morbidity and Mortality Among Adults With Bilateral Pheochromocytomas Undergoing Total Adrenalectomy vs Cortical-Sparing Adrenalectomy. JAMA Network Open, 2019, 2, e198898.	5.9	80
47	Functional analysis of clinical BARD1 germline variants. Journal of Physical Education and Sports Management, 2019, 5, a004093.	1.2	6
48	IL13RA2 Is Differentially Regulated in Papillary Thyroid Carcinoma vs Follicular Thyroid Carcinoma. Journal of Clinical Endocrinology and Metabolism, 2019, 104, 5573-5584.	3.6	14
49	Impact of free cancer predisposition cascade genetic testing on uptake in Singapore. Npj Genomic Medicine, 2019, 4, 22.	3.8	26
50	PRL3-zumab as an immunotherapy to inhibit tumors expressing PRL3 oncoprotein. Nature Communications, 2019, 10, 2484.	12.8	30
51	Advances in Sarcoma Genomics and Therapeutic Management. , 2019, , 609-621.		Ο
52	The Global State of the Genetic Counseling Profession. European Journal of Human Genetics, 2019, 27, 183-197.	2.8	215
53	PTEN-opathies: from biological insights to evidence-based precision medicine. Journal of Clinical Investigation, 2019, 129, 452-464.	8.2	128
54	SAT-317 The Importance of Accurate Genetic Diagnosis Highlighted in a Case of Recurrent Bilateral Pheochromocytomas. Journal of the Endocrine Society, 2019, 3, .	0.2	0

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55	Abstract 2780: IL13RA2 is differentially regulated in papillary thyroid carcinoma versus follicular thyroid carcinoma. , 2019, , .		0
56	Delayed diagnosis of Shwachman diamond syndrome with short telomeres and a review of cases in Asia. Leukemia Research Reports, 2018, 9, 54-57.	0.4	6
57	Mutational spectrum in a worldwide study of 29,700 families with <i>BRCA1</i> or <i>BRCA2</i> mutations. Human Mutation, 2018, 39, 593-620.	2.5	224
58	Risk management adherence following genetic testing for hereditary cancer syndromes: a Singaporean experience. Familial Cancer, 2018, 17, 621-626.	1.9	6
59	Factors influencing the decision to share cancer genetic results among family members: An inâ€depth interview study of women in an Asian setting. Psycho-Oncology, 2018, 27, 998-1004.	2.3	22
60	Evaluation of the relative effectiveness of the 2017 updated Manchester scoring system for predicting BRCA1/2 mutations in a Southeast Asian country. Journal of Medical Genetics, 2018, 55, 344-350.	3.2	5
61	Proffered Papers and Posters Presented at the Seventh International Symposium on Hereditary Breast and Ovarian Cancer—BrcA: From the Personal to the Population. Current Oncology, 2018, 25, 224-262.	2.2	2
62	Clinical relevance of screening checklists for detecting cancer predisposition syndromes in Asian childhood tumours. Npj Genomic Medicine, 2018, 3, 30.	3.8	13
63	Germline Pathogenic Variants in Homologous Recombination and DNA Repair Genes in an Asian Cohort of Young-Onset Colorectal Cancer. JNCI Cancer Spectrum, 2018, 2, pky054.	2.9	21
64	Whole exome sequencing identifies recessive germline mutations in FAM160A1 in familial NK/T cell lymphoma. Blood Cancer Journal, 2018, 8, 111.	6.2	5
65	A delayed diagnosis of Pallister-Hall syndrome in an adult male following the incidental detection of a hypothalamic hamartoma. Human Genome Variation, 2018, 5, 31.	0.7	3
66	The influence of Malay cultural beliefs on breast cancer screening and genetic testing: A focus group study. Psycho-Oncology, 2018, 27, 2855-2861.	2.3	30
67	Geneâ€specific criteria for <i>PTEN</i> variant curation: Recommendations from the ClinGen PTEN Expert Panel. Human Mutation, 2018, 39, 1581-1592.	2.5	123
68	Hamartomatous Polyposis Syndromes. , 2018, , 165-183.		1
69	65 YEARS OF THE DOUBLE HELIX: Genetics informs precision practice in the diagnosis and management of pheochromocytoma. Endocrine-Related Cancer, 2018, 25, T201-T219.	3.1	52
70	Impact of Appointment Waiting Time on Attendance Rates at a Clinical Cancer Genetics Service. Journal of Genetic Counseling, 2018, 27, 1473-1481.	1.6	15
71	65 YEARS OF THE DOUBLE HELIX: The advancements of gene editing and potential application to hereditary cancer. Endocrine-Related Cancer, 2018, 25, T141-T158.	3.1	3
72	Predictors of next-generation sequencing panel selection using a shared decision-making approach. Npj Genomic Medicine, 2018, 3, 11.	3.8	9

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73	Preventive medicine of von Hippel–Lindau disease-associated pancreatic neuroendocrine tumors. Endocrine-Related Cancer, 2018, 25, 783-793.	3.1	42
74	Inherited Thyroid Cancer. , 2018, , 163-171.		3
75	Breast cancer risk and clinical implications for germline PTEN mutation carriers. Breast Cancer Research and Treatment, 2017, 165, 1-8.	2.5	78
76	Clinical Implications for Germline PTEN Spectrum Disorders. Endocrinology and Metabolism Clinics of North America, 2017, 46, 503-517.	3.2	39
77	Clinical Characterization of the Pheochromocytoma and Paraganglioma Susceptibility Genes <i>SDHA</i> , <i>TMEM127</i> , <i>MAX</i> , and <i>SDHAF2</i> for Gene-Informed Prevention. JAMA Oncology, 2017, 3, 1204.	7.1	149
78	An International Collaborative Standardizing a Comprehensive Patient-Centered Outcomes Measurement Set for Colorectal Cancer. JAMA Oncology, 2017, 3, 686.	7.1	94
79	Germline Mutations in Cancer Predisposition Genes are Frequent in Sporadic Sarcomas. Scientific Reports, 2017, 7, 10660.	3.3	52
80	Impact of subsidies on cancer genetic testing uptake in Singapore. Journal of Medical Genetics, 2017, 54, 254-259.	3.2	24
81	Clinical management of pheochromocytoma and paraganglioma in Singapore: missed opportunities for genetic testing. Molecular Genetics & amp; Genomic Medicine, 2017, 5, 602-607.	1.2	7
82	Germline mutation contribution to chromosomal instability. Endocrine-Related Cancer, 2017, 24, T33-T46.	3.1	13
83	Immune dysregulation in patients with PTEN hamartoma tumor syndrome: Analysis of FOXP3 regulatory TÂcells. Journal of Allergy and Clinical Immunology, 2017, 139, 607-620.e15.	2.9	77
84	Sensing of dangerous DNA. Mechanisms of Ageing and Development, 2017, 165, 33-46.	4.6	33
85	Reply to G. Le Flahec et al. Journal of Clinical Oncology, 2017, 35, 377-377.	1.6	Ο
86	Abstract 4285: Impact of subsidies on cancer genetic testing uptake in Singapore. , 2017, , .		0
87	Cancer immunotherapy: unique perspectives for endocrine-related cancers. Endocrine-Related Cancer, 2017, 24, E13-E14.	3.1	Ο
88	Genetic predisposition resulting in sinusoidal obstruction syndrome in a patient with resected sigmoid cancer on adjuvant oxaliplatin. BMJ Case Reports, 2016, 2016, bcr2015212978.	0.5	2
89	Cost Effectiveness of Universal Hepatitis B Virus Screening in Patients Beginning Chemotherapy for Sarcomas or GI Stromal Tumors. Journal of Global Oncology, 2016, 2, 186-199.	0.5	4
90	Germline hemizygous deletion of CDKN2A–CDKN2B locus in a patient presenting with Li–Fraumeni syndrome. Npj Genomic Medicine, 2016, 1, 16015.	3.8	9

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91	Tissue Microbiome Profiling Identifies an Enrichment of Specific Enteric Bacteria in Opisthorchis viverrini Associated Cholangiocarcinoma. EBioMedicine, 2016, 8, 195-202.	6.1	94
92	The DNA Structure-Specific Endonuclease MUS81 Mediates DNA Sensor STINC-Dependent Host Rejection of Prostate Cancer Cells. Immunity, 2016, 44, 1177-1189.	14.3	162
93	Using Quality Improvement Methods and Time-Driven Activity-Based Costing to Improve Value-Based Cancer Care Delivery at a Cancer Genetics Clinic. Journal of Oncology Practice, 2016, 12, e320-e331.	2.5	24
94	Sarcomas Associated With Genetic Cancer Predisposition Syndromes: A Review. Oncologist, 2016, 21, 1002-1013.	3.7	78
95	Precision medicine in heritable cancer: when somatic tumour testing and germline mutations meet. Npj Genomic Medicine, 2016, 1, 15006.	3.8	41
96	Inherited breast cancer predisposition in Asians: multigene panel testing outcomes from Singapore. Npj Genomic Medicine, 2016, 1, 15003.	3.8	44
97	Familial non-medullary thyroid cancer: unraveling the genetic maze. Endocrine-Related Cancer, 2016, 23, R577-R595.	3.1	97
98	Clinical and molecular characteristics of East Asian patients with von Hippel–Lindau syndrome. Chinese Journal of Cancer, 2016, 35, 79.	4.9	6
99	Mismatch Repair Deficiency in Colorectal Cancers: Is Somatic Genomic Testing the Grab-Bag for All Answers?. Journal of Clinical Oncology, 2016, 34, 2085-2087.	1.6	5
100	Evaluation of the methods to identify patients who may benefit from PARP inhibitor use. Endocrine-Related Cancer, 2016, 23, R267-R285.	3.1	28
101	Mutation spectrum of POLE and POLD1 mutations in South East Asian women presenting with grade 3 endometrioid endometrial carcinomas. Gynecologic Oncology, 2016, 141, 113-120.	1.4	34
102	<i>HABP2</i> in Familial Non-medullary Thyroid Cancer: Will the Real Mutation Please Stand Up?. Journal of the National Cancer Institute, 2016, 108, djw013.	6.3	14
103	Rectal Cancer: Age Matters in the Affairs of Stage. Journal of the National Cancer Institute, 2016, 108, djv325.	6.3	3
104	Germline PTEN Mutation Analysis for PTEN Hamartoma Tumor Syndrome. Methods in Molecular Biology, 2016, 1388, 63-73.	0.9	9
105	Abstract 1793: POLE gene mutations confer good progression free survival in Asian women with FIGO grade 3 endometrioid endometrial carcinoma. , 2016, , .		0
106	A Case of Two Young Brothers with Natural-Killer/T-Cell Lymphoma. Blood, 2016, 128, 5293-5293.	1.4	0
107	Wholeâ€exome sequencing of breast cancer, malignant peripheral nerve sheath tumor and neurofibroma from a patient with neurofibromatosis type 1. Cancer Medicine, 2015, 4, 1871-1878.	2.8	30
108	Hereditary Syndromes Manifesting as Endometrial Carcinoma: How Can Pathological Features Aid Risk Assessment?. BioMed Research International, 2015, 2015, 1-17.	1.9	18

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109	PTEN hamartoma tumor syndrome: Clinical risk assessment and management protocol. Methods, 2015, 77-78, 11-19.	3.8	51
110	Cowden syndrome-associated germline SDHD variants alter PTEN nuclear translocation through SRC-induced PTEN oxidation. Human Molecular Genetics, 2015, 24, 142-153.	2.9	25
111	Germline and somatic SDHx alterations in apparently sporadic differentiated thyroid cancer. Endocrine-Related Cancer, 2015, 22, 121-130.	3.1	30
112	New Genetic and Genomic Approaches After the Genome-wide Association Study Era—Back to the Future. Gastroenterology, 2015, 149, 1138-1141.	1.3	6
113	Exome Sequencing Reveals Germline SMAD9 Mutation That Reduces Phosphatase and Tensin Homolog Expression and Is Associated With Hamartomatous Polyposis and Gastrointestinal Ganglioneuromas. Gastroenterology, 2015, 149, 886-889.e5.	1.3	24
114	Detecting Germline <i>PTEN</i> Mutations Among At-Risk Patients With Cancer: An Age- and Sex-Specific Cost-Effectiveness Analysis. Journal of Clinical Oncology, 2015, 33, 2537-2544.	1.6	16
115	Germline Heterozygous Variants in SEC23B Are Associated with Cowden Syndrome and Enriched in Apparently Sporadic Thyroid Cancer. American Journal of Human Genetics, 2015, 97, 661-676.	6.2	76
116	Germline <scp><i>PTEN</i></scp> , <scp><i>SDHBâ€Ð</i></scp> , and <scp><i>KLLN</i></scp> alterations in endometrial cancer patients with Cowden and Cowdenâ€like syndromes: An international, multicenter, prospective study. Cancer, 2015, 121, 688-696.	4.1	46
117	The Singapore Liver Cancer Recurrence (SLICER) Score for Relapse Prediction in Patients with Surgically Resected Hepatocellular Carcinoma. PLoS ONE, 2015, 10, e0118658.	2.5	46
118	Hereditary diffuse gastric cancer: What the clinician should know. World Journal of Gastrointestinal Oncology, 2015, 7, 153.	2.0	25
119	Abstract 3468: Impact of histology on serum thyroglobulin as a biomarker for nonmedullary thyroid cancer recurrence. , 2015, , .		0
120	Abstract 3732: Detecting germline PTEN mutations among at-risk cancer patients: An age and gender-specific cost effectiveness analysis. , 2015, , .		0
121	Germline Alterations in <i>RASAL1</i> in Cowden Syndrome Patients Presenting with Follicular Thyroid Cancer and in Individuals with Apparently Sporadic Epithelial Thyroid Cancer. Journal of Clinical Endocrinology and Metabolism, 2014, 99, E1316-E1321.	3.6	20
122	Whole-genome sequencing: not yet making the clinical grade. Personalized Medicine, 2014, 11, 471-475.	1.5	0
123	BluepRINT for Moderate-to-Low Penetrance Cancer Susceptibility Genes Needed: Breast Cancer and Beyond. Cancer Discovery, 2014, 4, 762-763.	9.4	1
124	Increased Prevalence of Eosinophilic Gastrointestinal Disorders in Pediatric <i>PTEN</i> Hamartoma Tumor Syndromes. Journal of Pediatric Gastroenterology and Nutrition, 2014, 58, 553-560.	1.8	41
125	Potential genetic anticipation in hereditary leiomyomatosis-renal cell cancer (HLRCC). Familial Cancer, 2014, 13, 281-289.	1.9	20
126	Second Malignant Neoplasms in Patients With Cowden Syndrome With Underlying Germline <i>PTEN</i> Mutations. Journal of Clinical Oncology, 2014, 32, 1818-1824.	1.6	105

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127	<i>TERT</i> and <i>BRAF</i> in Thyroid Cancer: Teaming Up for Trouble. Journal of Clinical Oncology, 2014, 32, 2683-2684.	1.6	17
128	Prevalence of Germline PTEN, BMPR1A, SMAD4, STK11, and ENG Mutations in Patients With Moderate-Load Colorectal Polyps. Gastroenterology, 2013, 144, 1402-1409.e5.	1.3	61
129	RASAL1 in Thyroid Cancer: Wisdom From an Old Foe. Journal of the National Cancer Institute, 2013, 105, 1597-1599.	6.3	9
130	Population-Based Universal Screening for Lynch Syndrome: Ready, Set… How?. Journal of Clinical Oncology, 2013, 31, 2527-2529.	1.6	12
131	Research Highlights: Highlights from the latest articles in personalized medicine. Personalized Medicine, 2013, 10, 231-233.	1.5	Ο
132	Into the eye of the storm: breast cancer's somatic mutation landscape points to DNA damage and repair. Translational Cancer Research, 2013, 2, 59-61.	1.0	1
133	Utility of PTEN Protein Dosage in Predicting for Underlying Germline <i>PTEN</i> Mutations among Patients Presenting with Thyroid Cancer and Cowden-Like Phenotypes. Journal of Clinical Endocrinology and Metabolism, 2012, 97, E2320-E2327.	3.6	17
134	<i>PTEN</i> mutations: help spot thyroid cancer before it occurs. Expert Review of Endocrinology and Metabolism, 2012, 7, 251-254.	2.4	1
135	GATA2 negatively regulates PTEN by preventing nuclear translocation of androgen receptor and by androgen-independent suppression of PTEN transcription in breast cancer. Human Molecular Genetics, 2012, 21, 569-576.	2.9	52
136	Lifetime Cancer Risks in Individuals with Germline <i>PTEN</i> Mutations. Clinical Cancer Research, 2012, 18, 400-407.	7.0	738
137	Thyroid cancer genetics: how close are we to personalizing clinical management?. Personalized Medicine, 2012, 9, 355-358.	1.5	1
138	PTEN Lipid Phosphatase Activity and Proper Subcellular Localization Are Necessary and Sufficient for Down-Regulating AKT Phosphorylation in the Nucleus in Cowden Syndrome. Journal of Clinical Endocrinology and Metabolism, 2012, 97, E2179-E2187.	3.6	18
139	Autoimmunity, Intestinal Lymphoid Hyperplasia, and Defects in Mucosal B-Cell Homeostasis in Patients With PTEN Hamartoma Tumor Syndrome. Gastroenterology, 2012, 142, 1093-1096.e6.	1.3	61
140	Use of PTEN protein dosage to predict for underlying germ-line PTEN mutations among patients presenting with thyroid cancer and Cowden-like phenotypes Journal of Clinical Oncology, 2012, 30, 1508-1508.	1.6	0
141	Evil lurks in the heart of man: cardiac paraganglioma presenting as recurrent dyspnoea and chronic cough. BMJ Case Reports, 2011, 2011, bcr1120115170-bcr1120115170.	0.5	5
142	Targeted therapies in the treatment of gastric cancer. Asia-Pacific Journal of Clinical Oncology, 2011, 7, 224-235.	1.1	12
143	Nuclear expression of MATK is a novel marker of type II enteropathy-associated T-cell lymphoma. Leukemia, 2011, 25, 555-557.	7.2	59
144	Prognostic impact of bleomycin-induced pneumonitis on the outcome of Hodgkin's lymphoma. Annals of Hematology, 2011, 90, 67-72.	1.8	22

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145	Docetaxel is effective in heavily pretreated patients with disseminated nasopharyngeal carcinoma. Annals of Oncology, 2011, 22, 718-722.	1.2	64
146	Incidence and Clinical Characteristics of Thyroid Cancer in Prospective Series of Individuals with Cowden and Cowden-Like Syndrome Characterized by Germline PTEN, SDH, or KLLN Alterations. Journal of Clinical Endocrinology and Metabolism, 2011, 96, E2063-E2071.	3.6	172
147	The Role of Pemetrexed Combined with Gemcitabine for Non-Small-Cell Lung Cancer. Current Drug Targets, 2010, 11, 61-66.	2.1	5
148	Impact of comorbidities on clinical outcomes in non-small cell lung cancer patients who are elderly and/or have poor performance status. Critical Reviews in Oncology/Hematology, 2010, 76, 53-60.	4.4	27
149	Recurring Mouth Ulcer and Skin Rash in a Man With Abnormal Chest Radiograph. Chest, 2010, 137, 983-988.	0.8	0
150	Second hematologic malignancies after ABVD: Two case reports and a retrospective study of 183 Hodgkin lymphoma patients. Acta Oncológica, 2010, 49, 257-259.	1.8	3
151	Abstract 2167: Unbiased genomic approaches identify 2 major subclasses of Gastric Cancer with prognostic and predictive value superior to Lauren's and also reveals subtype-specific treatment opportunities. , 2010, , .		0
152	A multimodality approach to reversible paraneoplastic encephalitis associated with ovarian teratomas. Acta Oncológica, 2009, 48, 1079-1082.	1.8	1
153	High SUV uptake on FDG–PET/CT predicts for an aggressive B-cell lymphoma in a prospective study of primary FDG–PET/CT staging in lymphoma. Annals of Oncology, 2009, 20, 1543-1547.	1.2	131
154	Comparative analysis of extraâ€nodal NK/Tâ€cell lymphoma and peripheral Tâ€cell lymphoma: significant differences in clinical characteristics and prognosis. European Journal of Haematology, 2008, 80, 55-60.	2.2	57
155	Incomplete Immunity and Missed Vaccination Opportunities in East African Immigrants Settling in Australia. Journal of Immigrant and Minority Health, 2008, 10, 263-268.	1.6	24
156	Prognostic factors in patients with diffuse large B cell lymphoma: Before and after the introduction of rituximab. Leukemia and Lymphoma, 2008, 49, 462-469.	1.3	95
157	Capecitabine-induced oromandibular dystonia: A case report and literature review. Acta Oncológica, 2008, 47, 1161-1165.	1.8	11
158	Health services utilisation and barriers for settlers from the Horn of Africa. Australian and New Zealand Journal of Public Health, 2007, 31, 333-335.	1.8	20
159	Undiagnosed and Potentially Lethal Parasite Infections Among Immigrants and Refugees in Australia. Journal of Travel Medicine, 2006, 13, 233-239.	3.0	68
160	Vitamin D deficiency is common and unrecognized among recently arrived adult immigrants from The Horn of Africa. Internal Medicine Journal, 2003, 33, 47-51.	0.8	60
161	Oncology workload in a tertiary hospital during the COVID-19 pandemic. Proceedings of Singapore Healthcare, 0, , 201010582110511.	0.6	1