Marcin Imielinski

List of Publications by Year in descending order

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Version: 2024-02-01

61 30 papers cit

74

all docs

30,634 42 citations h-index

74

docs citations

h-index g-index

74 50355
times ranked citing authors

138484

58

#	Article	IF	CITATIONS
1	System-wide transcriptome damage and tissue identity loss in COVID-19 patients. Cell Reports Medicine, 2022, 3, 100522.	6.5	24
2	Abstract P2-06-04: Pathognomonic long molecule footprints of backup repair pathways in homologous recombination deficient cancers. Cancer Research, 2022, 82, P2-06-04-P2-06-04.	0.9	0
3	SETD2 Haploinsufficiency Enhances Germinal Center–Associated AICDA Somatic Hypermutation to Drive B-cell Lymphomagenesis. Cancer Discovery, 2022, 12, 1782-1803.	9.4	14
4	Somatic whole genome dynamics of precancer in Barrett's esophagus reveals features associated with disease progression. Nature Communications, 2022, 13, 2300.	12.8	13
5	Identifying synergistic high-order 3D chromatin conformations from genome-scale nanopore concatemer sequencing. Nature Biotechnology, 2022, 40, 1488-1499.	17.5	46
6	Recurrent somatic mutations as predictors of immunotherapy response. Nature Communications, 2022, 13, .	12.8	12
7	Histone H1 loss drives lymphoma by disrupting 3D chromatin architecture. Nature, 2021, 589, 299-305.	27.8	155
8	Whole-genome characterization of lung adenocarcinomas lacking alterations in the RTK/RAS/RAF pathway. Cell Reports, 2021, 34, 108707.	6.4	16
9	Editorial overview: The most difficult of years in cancer research. Current Opinion in Genetics and Development, 2021, 66, iii-iv.	3.3	O
10	A Genomic-Pathologic Annotated Risk Model to Predict Recurrence in Early-Stage Lung Adenocarcinoma. JAMA Surgery, 2021, 156, e205601.	4.3	52
11	Shotgun transcriptome, spatial omics, and isothermal profiling of SARS-CoV-2 infection reveals unique host responses, viral diversification, and drug interactions. Nature Communications, 2021, 12, 1660.	12.8	132
12	Integrated mutational landscape analysis of uterine leiomyosarcomas. Proceedings of the National Academy of Sciences of the United States of America, 2021, 118, .	7.1	48
13	Characterizing genetic intra-tumor heterogeneity across 2,658 human cancer genomes. Cell, 2021, 184, 2239-2254.e39.	28.9	260
14	Structural variant evolution after telomere crisis. Nature Communications, 2021, 12, 2093.	12.8	16
15	Discovery of Candidate DNA Methylation Cancer Driver Genes. Cancer Discovery, 2021, 11, 2266-2281.	9.4	42
16	Molecular Evolution of Classical Hodgkin Lymphoma Revealed Though Whole Genome Sequencing of Hodgkin and Reed-Sternberg Cells. Blood, 2021, 138, 805-805.	1.4	1
17	Distinct Classes of Complex Structural Variation Uncovered across Thousands of Cancer Genome Graphs. Cell, 2020, 183, 197-210.e32.	28.9	141
18	Impact of Lineage Plasticity to and from a Neuroendocrine Phenotype on Progression and Response in Prostate and Lung Cancers. Molecular Cell, 2020, 80, 562-577.	9.7	54

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19	The evolutionary history of 2,658 cancers. Nature, 2020, 578, 122-128.	27.8	690
20	Patterns of somatic structural variation in human cancer genomes. Nature, 2020, 578, 112-121.	27.8	560
21	Analyses of non-coding somatic drivers in 2,658Âcancer whole genomes. Nature, 2020, 578, 102-111.	27.8	424
22	Robust Discovery of Candidate DNA Methylation Cancer Drivers. Blood, 2020, 136, 33-34.	1.4	0
23	Next-generation characterization of the Cancer Cell Line Encyclopedia. Nature, 2019, 569, 503-508.	27.8	2,149
24	Deep Convolutional Neural Networks Enable Discrimination of Heterogeneous Digital Pathology Images. EBioMedicine, 2018, 27, 317-328.	6.1	240
25	SvABA: genome-wide detection of structural variants and indels by local assembly. Genome Research, 2018, 28, 581-591.	5.5	288
26	The chromatin accessibility landscape of primary human cancers. Science, 2018, 362, .	12.6	781
27	Fusion oncogenes—genetic musical chairs. Science, 2018, 361, 848-849.	12.6	4
28	Insertions and Deletions Target Lineage-Defining Genes in Human Cancers. Cell, 2017, 168, 460-472.e14.	28.9	106
29	Clinical hallmarks in whole cancer genomes. Nature Reviews Clinical Oncology, 2017, 14, 265-266.	27.6	1
30	Modeling cancer rearrangement landscapes. Current Opinion in Systems Biology, 2017, 1, 54-61.	2.6	17
31	The cancer precision medicine knowledge base for structured clinical-grade mutations and interpretations. Journal of the American Medical Informatics Association: JAMIA, 2017, 24, 513-519.	4.4	88
32	Distinct patterns of somatic genome alterations in lung adenocarcinomas and squamous cell carcinomas. Nature Genetics, 2016, 48, 607-616.	21.4	933
33	High-throughput Phenotyping of Lung Cancer Somatic Mutations. Cancer Cell, 2016, 30, 214-228.	16.8	171
34	Identification of focally amplified lineage-specific super-enhancers in human epithelial cancers. Nature Genetics, 2016, 48, 176-182.	21.4	283
35	A Pan-Cancer Analysis of Transcriptome Changes Associated with Somatic Mutations in U2AF1 Reveals Commonly Altered Splicing Events. PLoS ONE, 2014, 9, e87361.	2.5	168
36	Genetic modifiers of EGFR dependence in non-small cell lung cancer. Proceedings of the National Academy of Sciences of the United States of America, 2014, 111, 18661-18666.	7.1	46

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37	Oncogenic and sorafenib-sensitive ARAF mutations in lung adenocarcinoma. Journal of Clinical Investigation, 2014, 124, 1582-1586.	8.2	101
38	Signatures of mutational processes in human cancer. Nature, 2013, 500, 415-421.	27.8	8,060
39	Mutational heterogeneity in cancer and the search for new cancer-associated genes. Nature, 2013, 499, 214-218.	27.8	4,761
40	Functional analysis of receptor tyrosine kinase mutations in lung cancer identifies oncogenic extracellular domain mutations of <i>ERBB2</i> . Proceedings of the National Academy of Sciences of the United States of America, 2012, 109, 14476-14481.	7.1	246
41	Mapping the Hallmarks of Lung Adenocarcinoma with Massively Parallel Sequencing. Cell, 2012, 150, 1107-1120.	28.9	1,591
42	A Landscape of Driver Mutations in Melanoma. Cell, 2012, 150, 251-263.	28.9	2,247
43	A Genome-Wide Meta-Analysis of Six Type 1 Diabetes Cohorts Identifies Multiple Associated Loci. PLoS Genetics, 2011, 7, e1002293.	3.5	297
44	Duplication of the SLIT3 Locus on 5q35.1 Predisposes to Major Depressive Disorder. PLoS ONE, 2010, 5, e15463.	2.5	63
45	Comparative genetic analysis of inflammatory bowel disease and type 1 diabetes implicates multiple loci with opposite effects. Human Molecular Genetics, 2010, 19, 2059-2067.	2.9	157
46	Strong synaptic transmission impact by copy number variations in schizophrenia. Proceedings of the National Academy of Sciences of the United States of America, 2010, 107, 10584-10589.	7.1	212
47	Breaking new ground in inflammatory bowel disease genetics: genome-wide association studies and beyond. Pharmacogenomics, 2010, 11, 663-665.	1.3	6
48	Variants of <i>DENND1B </i> Associated with Asthma in Children. New England Journal of Medicine, 2010, 362, 36-44.	27.0	306
49	Follow-Up Analysis of Genome-Wide Association Data Identifies Novel Loci for Type 1 Diabetes. Diabetes, 2009, 58, 290-295.	0.6	136
50	High-resolution mapping and analysis of copy number variations in the human genome: A data resource for clinical and research applications. Genome Research, 2009, 19, 1682-1690.	5.5	313
51	Genome-Wide Analyses of Exonic Copy Number Variants in a Family-Based Study Point to Novel Autism Susceptibility Genes. PLoS Genetics, 2009, 5, e1000536.	3.5	374
52	Autism genome-wide copy number variation reveals ubiquitin and neuronal genes. Nature, 2009, 459, 569-573.	27.8	1,270
53	Common genetic variants on $5p14.1$ associate with autism spectrum disorders. Nature, 2009, 459, 528-533.	27.8	912
54	Common variants at five new loci associated with early-onset inflammatory bowel disease. Nature Genetics, 2009, 41, 1335-1340.	21.4	459

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55	Diverse Genome-wide Association Studies Associate the IL12/IL23 Pathway with Crohn Disease. American Journal of Human Genetics, 2009, 84, 399-405.	6.2	246
56	17q12-21 variants interact with smoke exposure as a risk factor for pediatric asthma but are equally associated with early-onset versus late-onset asthma in North Americans of European ancestry. Journal of Allergy and Clinical Immunology, 2009, 124, 605-607.	2.9	68
57	Loci on 20q13 and 21q22 are associated with pediatric-onset inflammatory bowel disease. Nature Genetics, 2008, 40, 1211-1215.	21.4	310
58	ORMDL3 variants associated with asthma susceptibility in North Americans of European ancestry. Journal of Allergy and Clinical Immunology, 2008, 122, 1225-1227.	2.9	89
59	Association Analysis of the FTO Gene with Obesity in Children of Caucasian and African Ancestry Reveals a Common Tagging SNP. PLoS ONE, 2008, 3, e1746.	2.5	176
60	Association of the BANK1 R61H variant with systemic lupus erythematosus in Americans of European and African ancestry. The Application of Clinical Genetics, 2008, Volume 2, 1-5.	3.0	8
61	Systematic Analysis of Conservation Relations in Escherichia coli Genome-Scale Metabolic Network Reveals Novel Growth Media. Biophysical Journal, 2006, 90, 2659-2672.	0.5	43