## **Michael Baudis**

## List of Publications by Year in Descending Order

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The third column is the impact factor (IF) of the journal, and the fourth column is the number of citations of the article.

86 56 29 3,273 h-index g-index citations papers 3,996 102 7.3 4.91 L-index avg, IF ext. citations ext. papers

#	Paper	IF	Citations
86	Beacon v2 and Beacon Networks: a "lingua franca" for federated data discovery in biomedical genomics, and beyond <i>Human Mutation</i> , <b>2022</b> ,	4.7	1
85	GA4GH: International policies and standards for data sharing across genomic research and healthcare <i>Cell Genomics</i> , <b>2021</b> , 1, 100029-100029		20
84	The GA4GH Variation Representation Specification: A computational framework for variation representation and federated identification <i>Cell Genomics</i> , <b>2021</b> , 1, 100027-100027		4
83	International federation of genomic medicine databases using GA4GH standards <i>Cell Genomics</i> , <b>2021</b> , 1, 100032-100032		7
82	Signatures of Discriminative Copy Number Aberrations in 31 Cancer Subtypes. <i>Frontiers in Genetics</i> , <b>2021</b> , 12, 654887	4.5	1
81	The Progenetix oncogenomic resource in 2021. <i>Database: the Journal of Biological Databases and Curation</i> , <b>2021</b> , 2021,	5	1
80	Geographic assessment of cancer genome profiling studies. <i>Database: the Journal of Biological Databases and Curation</i> , <b>2020</b> , 2020,	5	1
79	Oncology Informatics: Status Quo and Outlook. <i>Oncology</i> , <b>2020</b> , 98, 329-331	3.6	5
78	Minimum error calibration and normalization for genomic copy number analysis. <i>Genomics</i> , <b>2020</b> , 112, 3331-3341	4.3	1
77	Enabling population assignment from cancer genomes with SNP2pop. Scientific Reports, 2020, 10, 4846	4.9	3
76	A harmonized meta-knowledgebase of clinical interpretations of somatic genomic variants in cancer. <i>Nature Genetics</i> , <b>2020</b> , 52, 448-457	36.3	58
75	The ELIXIR Human Copy Number Variations Community: building bioinformatics infrastructure for research. <i>F1000Research</i> , <b>2020</b> , 9,	3.6	4
74	DNA Copy Number Changes in Diffuse Large B Cell Lymphomas. Frontiers in Oncology, <b>2020</b> , 10, 584095	5.3	4
73	The Ubiquitin Ligase TRIP12 Limits PARP1 Trapping and Constrains PARP Inhibitor Efficiency. <i>Cell Reports</i> , <b>2020</b> , 32, 107985	10.6	22
72	Mountains and Chasms: Surveying the Oncogenomic Publication Landscape. <i>Oncology</i> , <b>2020</b> , 98, 332-34	<b>-3</b> 3.6	3
71	Leveraging European infrastructures to access 1 million human genomes by 2022. <i>Nature Reviews Genetics</i> , <b>2019</b> , 20, 693-701	30.1	36
70	DNA copy number imbalances in primary cutaneous lymphomas. <i>Journal of the European Academy of Dermatology and Venereology</i> , <b>2019</b> , 33, 1062-1075	4.6	3

69	Federated discovery and sharing of genomic data using Beacons. <i>Nature Biotechnology</i> , <b>2019</b> , 37, 220-2	2 <b>24</b> 4.5	42
68	Krppel-Like Factor 10 participates in cervical cancer immunoediting through transcriptional regulation of Pregnancy-Specific Beta-1 Glycoproteins. <i>Scientific Reports</i> , <b>2018</b> , 8, 9445	4.9	5
67	Registered access: authorizing data access. European Journal of Human Genetics, 2018, 26, 1721-1731	5.3	17
66	segment_liftover : a Python tool to convert segments between genome assemblies. <i>F1000Research</i> , <b>2018</b> , 7, 319	3.6	12
65	segment_liftover : a Python tool to convert segments between genome assemblies. <i>F1000Research</i> , <b>2018</b> , 7, 319	3.6	13
64	Integrated Molecular Meta-Analysis of 1,000 Pediatric High-Grade and Diffuse Intrinsic Pontine Glioma. <i>Cancer Cell</i> , <b>2017</b> , 32, 520-537.e5	24.3	423
63	The SIB Swiss Institute of BioinformaticsR esources: focus on curated databases. <i>Nucleic Acids Research</i> , <b>2016</b> , 44, D27-37	20.1	41
62	CNARA: reliability assessment for genomic copy number profiles. <i>BMC Genomics</i> , <b>2016</b> , 17, 799	4.5	2
61	HG-95INTEGRATED MOLECULAR META-ANALYSIS OF 1000 PAEDIATRIC HIGH GRADE GLIOMA AND DIPG. <i>Neuro-Oncology</i> , <b>2016</b> , 18, iii70.3-iii70	1	78
60	PKCland HMGB1 antagonistically control hydrogen peroxide-induced poly-ADP-ribose formation. <i>Nucleic Acids Research</i> , <b>2016</b> , 44, 7630-45	20.1	12
59	arrayMap 2014: an updated cancer genome resource. <i>Nucleic Acids Research</i> , <b>2015</b> , 43, D825-30	20.1	15
58	HG-11 * INTEGRATIVE MOLECULAR META-ANALYSIS OF 700 PEDIATRIC HIGH GRADE GLIOMA AND DIPG DEFINES WIDESPREAD INTER- AND INTRA-TUMORAL HETEROGENEITY. <i>Neuro-Oncology</i> , <b>2015</b> , 17, iii12-iii13	1	1
57	Genomic instability of osteosarcoma cell lines in culture: impact on the prediction of metastasis relevant genes. <i>PLoS ONE</i> , <b>2015</b> , 10, e0125611	3.7	21
56	Chromothripsis-like patterns are recurring but heterogeneously distributed features in a survey of 22,347 cancer genome screens. <i>BMC Genomics</i> , <b>2014</b> , 15, 82	4.5	78
55	Progenetix: 12 years of oncogenomic data curation. <i>Nucleic Acids Research</i> , <b>2014</b> , 42, D1055-62	20.1	28
54	HIGH GRADE GLIOMAS AND DIPG. <i>Neuro-Oncology</i> , <b>2014</b> , 16, i40-i59	1	78
53	Biopsying parapsoriasis: quo vadis? Are morphological stains enough or are ancillary tests needed?. <i>Romanian Journal of Morphology and Embryology</i> , <b>2014</b> , 55, 1085-92	0.6	1
52	High resolution copy number analysis of IRF4 translocation-positive diffuse large B-cell and follicular lymphomas. <i>Genes Chromosomes and Cancer</i> , <b>2013</b> , 52, 150-5	5	19

51	PKC signaling prevents irradiation-induced apoptosis of primary human fibroblasts. <i>Cell Death and Disease</i> , <b>2013</b> , 4, e498	9.8	33
50	SIL1 mutations and clinical spectrum in patients with Marinesco-Sjogren syndrome. <i>Brain</i> , <b>2013</b> , 136, 3634-44	11.2	58
49	Recurrent loss of heterozygosity in 1p36 associated with TNFRSF14 mutations in IRF4 translocation negative pediatric follicular lymphomas. <i>Haematologica</i> , <b>2013</b> , 98, 1237-41	6.6	43
48	DNA copy number alterations in central primitive neuroectodermal tumors and tumors of the pineal region: an international individual patient data meta-analysis. <i>Journal of Neuro-Oncology</i> , <b>2012</b> , 109, 415-23	4.8	12
47	Molecular karyotyping as a relevant diagnostic tool in children with growth retardation with Silver-Russell features. <i>Journal of Pediatrics</i> , <b>2012</b> , 161, 933-42	3.6	32
46	2p21 Deletions in hypotonia-cystinuria syndrome. <i>European Journal of Medical Genetics</i> , <b>2012</b> , 55, 561-3	2.6	13
45	Losses at chromosome 4q are associated with poor survival in operable ductal pancreatic adenocarcinoma. <i>Pancreatology</i> , <b>2012</b> , 12, 16-22	3.8	7
44	Integrative genome-wide expression profiling identifies three distinct molecular subgroups of renal cell carcinoma with different patient outcome. <i>BMC Cancer</i> , <b>2012</b> , 12, 310	4.8	23
43	Improved multiplex ligation-dependent probe amplification analysis identifies a deleterious PMS2 allele generated by recombination with crossover between PMS2 and PMS2CL. <i>Genes Chromosomes and Cancer</i> , <b>2012</b> , 51, 819-31	5	16
42	Array-basierter Nachweis chromosomaler Aberrationen bei malignen Neoplasien. <i>Medizinische Genetik</i> , <b>2012</b> , 24, 114-122	0.5	
41	arrayMap: a reference resource for genomic copy number imbalances in human malignancies. <i>PLoS ONE</i> , <b>2012</b> , 7, e36944	3.7	27
40	Specific genomic regions are differentially affected by copy number alterations across distinct cancer types, in aggregated cytogenetic data. <i>PLoS ONE</i> , <b>2012</b> , 7, e43689	3.7	3
39	Supernumerary Asymmetric Dic(15;15) With Secondary Mosaic Formation in One of Two Developmentally Retarded Twins. <i>International Journal of Human Genetics</i> , <b>2011</b> , 11, 75-82	1	
38	Silver-Russell patients showing a broad range of ICR1 and ICR2 hypomethylation in different tissues. <i>Clinical Genetics</i> , <b>2011</b> , 80, 83-8	4	51
37	CDCOCA: a statistical method to define complexity dependence of co-occuring chromosomal aberrations. <i>BMC Medical Genomics</i> , <b>2011</b> , 4, 21	3.7	6
36	Pathways and Crossroads to Colorectal Cancer <b>2011</b> , 369-394		3
35	Submicroscopic chromosomal imbalances in idiopathic Silver-Russell syndrome (SRS): the SRS phenotype overlaps with the 12q14 microdeletion syndrome. <i>Journal of Medical Genetics</i> , <b>2010</b> , 47, 356	-ēo <sup>8</sup>	38
34	CDCOCA: a statistical method to define complexity dependent co-occurring chromosomal aberrations <b>2010</b> , 11, P23		1

## (2006-2010)

33	Increased expression of cellular retinol-binding protein 1 in laryngeal squamous cell carcinoma.  Journal of Cancer Research and Clinical Oncology, 2010, 136, 931-8	4.9	18
32	MUC1 oncogene amplification correlates with protein overexpression in invasive breast carcinoma cells. <i>Cancer Genetics and Cytogenetics</i> , <b>2010</b> , 201, 102-10		37
31	Identification of a 21q22 duplication in a Silver-Russell syndrome patient further narrows down the Down syndrome critical region. <i>American Journal of Medical Genetics, Part A</i> , <b>2010</b> , 152A, 356-9	2.5	12
30	Chromosome 11p15 duplication in Silver-Russell syndrome due to a maternally inherited translocation t(11;15). <i>American Journal of Medical Genetics, Part A</i> , <b>2010</b> , 152A, 1484-7	2.5	11
29	Inferring progression models for CGH data. <i>Bioinformatics</i> , <b>2009</b> , 25, 2208-15	7.2	9
28	Recurrent loss, but lack of mutations, of the SMARCB1 tumor suppressor gene in T-cell prolymphocytic leukemia with TCL1A-TCRAD juxtaposition. <i>Cancer Genetics and Cytogenetics</i> , <b>2009</b> , 192, 44-7		19
27	Translocations involving 8q24 in Burkitt lymphoma and other malignant lymphomas: a historical review of cytogenetics in the light of todays knowledge. <i>Leukemia</i> , <b>2009</b> , 23, 225-34	10.7	165
26	Quantifying cancer progression with conjunctive Bayesian networks. <i>Bioinformatics</i> , <b>2009</b> , 25, 2809-15	7.2	76
25	Comprehensive characterization of genomic aberrations in gangliogliomas by CGH, array-based CGH and interphase FISH. <i>Brain Pathology</i> , <b>2008</b> , 18, 326-37	6	53
24	Recurrent loss of the Y chromosome and homozygous deletions within the pseudoautosomal region 1: association with male predominance in mantle cell lymphoma. <i>Haematologica</i> , <b>2008</b> , 93, 949-5	0 <sup>6.6</sup>	13
23	Chromosomal changes characterize head and neck cancer with poor prognosis. <i>Journal of Molecular Medicine</i> , <b>2008</b> , 86, 1353-65	5.5	27
22	A 10.7 Mb interstitial deletion of 13q21 without phenotypic effect defines a further non-pathogenic euchromatic variant. <i>American Journal of Medical Genetics, Part A</i> , <b>2008</b> , 146A, 2417-20	2.5	8
21	Translocations Involving 8q24 in Burkitt Lymphoma and Other Malignant Lymphomas: A Historical Review of Cytogenetics in the Light of Todays Knowledge. <i>Blood</i> , <b>2008</b> , 112, 2814-2814	2.2	
20	Combined single nucleotide polymorphism-based genomic mapping and global gene expression profiling identifies novel chromosomal imbalances, mechanisms and candidate genes important in the pathogenesis of T-cell prolymphocytic leukemia with inv(14)(q11q32). <i>Leukemia</i> , <b>2007</b> , 21, 2153-63	10.7	80
19	Genomic imbalances in 5918 malignant epithelial tumors: an explorative meta-analysis of chromosomal CGH data. <i>BMC Cancer</i> , <b>2007</b> , 7, 226	4.8	109
18	Allele-specific loss of heterozygosity in multiple colorectal adenomas: toward an integrated molecular cytogenetic map II. <i>Cancer Genetics and Cytogenetics</i> , <b>2006</b> , 167, 1-14		17
17	Distance-based clustering of CGH data. <i>Bioinformatics</i> , <b>2006</b> , 22, 1971-8	7.2	47
16	Online database and bioinformatics toolbox to support data mining in cancer cytogenetics. <i>BioTechniques</i> , <b>2006</b> , 40, 269-70, 272	2.5	22

15	ABCB1 over-expression and drug-efflux in acute lymphoblastic leukemia cell lines with t(17;19) and E2A-HLF expression. <i>Pediatric Blood and Cancer</i> , <b>2006</b> , 47, 757-64	3	13
14	Genomisches Profiling mittels SCOMP bei duktalen Adenokarzinomen des Pankreas. <i>Langenbecks Archiv Fu r Chirurgie Supplement</i> , <b>2006</b> , 27-28		
13	Genetic losses in breast cancer: toward an integrated molecular cytogenetic map. <i>Cancer Genetics and Cytogenetics</i> , <b>2005</b> , 160, 141-51		12
12	Microarray comparative genomic hybridization detection of chromosomal imbalances in uterine cervix carcinoma. <i>BMC Cancer</i> , <b>2005</b> , 5, 77	4.8	63
11	Unequivocal delineation of clinicogenetic subgroups and development of a new model for improved outcome prediction in neuroblastoma. <i>Journal of Clinical Oncology</i> , <b>2005</b> , 23, 2280-99	2.2	145
10	Randomized study to evaluate the use of high-dose therapy as part of primary treatment for "aggressive" lymphoma. <i>Journal of Clinical Oncology</i> , <b>2002</b> , 20, 4413-9	2.2	118
9	Gain of chromosome arm 9p is characteristic of primary mediastinal B-cell lymphoma (MBL): comprehensive molecular cytogenetic analysis and presentation of a novel MBL cell line. <i>Genes Chromosomes and Cancer</i> , <b>2001</b> , 30, 393-401	5	117
8	Progenetix.net: an online repository for molecular cytogenetic aberration data. <i>Bioinformatics</i> , <b>2001</b> , 17, 1228-9	7.2	155
7	Comparative genomic hybridization for the analysis of leukemias and lymphomas. <i>Methods in Molecular Medicine</i> , <b>2001</b> , 55, 43-64		1
6	t(11;14)-positive mantle cell lymphomas exhibit complex karyotypes and share similarities with B-cell chronic lymphocytic leukemia <b>2000</b> , 27, 285-294		112
5	Analysis of genomic alterations in benign, atypical, and anaplastic meningiomas: toward a genetic model of meningioma progression. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , <b>1997</b> , 94, 14719-24	11.5	324
4	Chromosome imbalances in papillary renal cell carcinoma and first cytogenetic data of familial cases analyzed by comparative genomic hybridization. <i>Cytogenetic and Genome Research</i> , <b>1996</b> , 75, 17-	21 <sup>1.9</sup>	36
3	Identification of genetic imbalances in malignant lymphoma using comparative genomic hybridization. <i>Stem Cells</i> , <b>1995</b> , 13 Suppl 3, 83-7	5.8	14
2	A harmonized meta-knowledgebase of clinical interpretations of cancer genomic variants		5
1	The GA4GH Variation Representation Specification (VRS): a Computational Framework for the Precise Representation and Federated Identification of Molecular Variation		3