## Hans Ehrencrona

List of Publications by Year in descending order

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71061 54882 7,525 88 41 84 citations h-index g-index papers 90 90 90 11481 docs citations times ranked citing authors all docs

#	Article	IF	CITATIONS
1	Extended genetic diagnostics for children with profound sensorineural hearing loss by implementing massive parallel sequencing. Diagnostic outcome, family experience and clinical implementation. International Journal of Pediatric Otorhinolaryngology, 2022, 159, 111218.	0.4	5
2	A search for modifying genetic factors in CHEK2:c.1100delC breast cancer patients. Scientific Reports, 2021, 11, 14763.	1.6	3
3	Genetic testing in women with early-onset breast cancer: a Traceback pilot study. Breast Cancer Research and Treatment, 2021, 190, 307-315.	1.1	1
4	FINAL Analysis of a PAN European STOP Tyrosine Kinase Inhibitor Trial in Chronic Myeloid Leukemia : The EURO-SKI Study. Blood, 2021, 138, 633-633.	0.6	10
5	Association of Genomic Domains in <i>BRCA1</i> and <i>BRCA2</i> with Prostate Cancer Risk and Aggressiveness. Cancer Research, 2020, 80, 624-638.	0.4	39
6	Cancer Risks Associated With Germline <i>PALB2</i> Pathogenic Variants: An International Study of 524 Families. Journal of Clinical Oncology, 2020, 38, 674-685.	0.8	270
7	Comprehensive molecular comparison of BRCA1 hypermethylated and BRCA1 mutated triple negative breast cancers. Nature Communications, 2020, 11, 3747.	5.8	53
8	Public support for healthcare-mediated disclosure of hereditary cancer risk information: Results from a population-based survey in Sweden. Hereditary Cancer in Clinical Practice, 2020, 18, 18.	0.6	11
9	Variations in the Referral Pattern for Genetic Counseling of Patients with Early-Onset Breast Cancer: A Population-Based Study in Southern Sweden. Public Health Genomics, 2020, 23, 100-109.	0.6	1
10	Ovarian and Breast Cancer Risks Associated With Pathogenic Variants in <i>RAD51C</i> and <i>RAD51D</i> Journal of the National Cancer Institute, 2020, 112, 1242-1250.	3.0	106
11	The Spectrum of FANCM Protein Truncating Variants in European Breast Cancer Cases. Cancers, 2020, 12, 292.	1.7	11
12	Large scale multifactorial likelihood quantitative analysis of <i>BRCA1</i> and <i>BRCA2</i> variants: An ENIGMA resource to support clinical variant classification. Human Mutation, 2019, 40, 1557-1578.	1.1	102
13	Whole-genome sequencing of triple-negative breast cancers in a population-based clinical study. Nature Medicine, 2019, 25, 1526-1533.	15.2	218
14	Hereditary colorectal cancer diagnostics in southern Sweden: retrospective evaluation and future considerations with emphasis on Lynch syndrome. Journal of Community Genetics, 2019, 10, 259-266.	0.5	2
15	Clinicians' use of breast cancer risk assessment tools according to their perceived importance of breast cancer risk factors: an international survey. Journal of Community Genetics, 2019, 10, 61-71.	0.5	7
16	Mutational spectrum in a worldwide study of 29,700 families with <i>BRCA1</i> brcai>BRCA2mutations. Human Mutation, 2018, 39, 593-620.	1.1	224
17	Discontinuation of tyrosine kinase inhibitor therapy in chronic myeloid leukaemia (EURO-SKI): a prespecified interim analysis of a prospective, multicentre, non-randomised, trial. Lancet Oncology, The, 2018, 19, 747-757.	5.1	444
18	The <i>BRCA1</i> c. 5096G> A p.Arg1699Gln (R1699Q) intermediate risk variant: breast and ovarian cancer risk estimation and recommendations for clinical management from the ENIGMA consortium. Journal of Medical Genetics, 2018, 55, 15-20.	1.5	50

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19	Use of the BOADICEA Web Application in clinical practice: appraisals by clinicians from various countries. Familial Cancer, 2018, 17, 31-41.	0.9	9
20	Individuals with FANCM biallelic mutations do not develop Fanconi anemia, but show risk for breast cancer, chemotherapy toxicity and may display chromosome fragility. Genetics in Medicine, 2018, 20, 452-457.	1.1	59
21	Identification of ten variants associated with risk of estrogen-receptor-negative breast cancer. Nature Genetics, 2017, 49, 1767-1778.	9.4	289
22	Increased proportion of mature NK cells is associated with successful imatinib discontinuation in chronic myeloid leukemia. Leukemia, 2017, 31, 1108-1116.	3.3	201
23	Evaluation of Polygenic Risk Scores for Breast and Ovarian Cancer Risk Prediction in BRCA1 and BRCA2 Mutation Carriers. Journal of the National Cancer Institute, 2017, 109, .	3.0	242
24	Identification of four novel susceptibility loci for oestrogen receptor negative breast cancer. Nature Communications, 2016, 7, 11375.	5.8	93
25	Functional mechanisms underlying pleiotropic risk alleles at the 19p13.1 breast–ovarian cancer susceptibility locus. Nature Communications, 2016, 7, 12675.	5.8	78
26	Breast cancer risk variants at 6q25 display different phenotype associations and regulate ESR1, RMND1 and CCDC170. Nature Genetics, 2016, 48, 374-386.	9.4	125
27	No clinical utility of KRAS variant rs61764370 for ovarian or breast cancer. Gynecologic Oncology, 2016, 141, 386-401.	0.6	18
28	Cessation of Tyrosine Kinase Inhibitors Treatment in Chronic Myeloid Leukemia Patients with Deep Molecular Response: Results of the Euro-Ski Trial. Blood, 2016, 128, 787-787.	0.6	43
29	No Differences in Molecular Relapse-Free Survival after Stopping Imatinib Treatment of Chronic Myeloid Leukemia Between Patients with Prior 4.5 Log Reduction (MR4.5) but Detectable and Patients with Undetectable Disease in the EURO-SKI Trial. Blood, 2016, 128, 789-789.	0.6	9
30	An original phylogenetic approach identified mitochondrial haplogroup T1a1 as inversely associated with breast cancer risk in BRCA2 mutation carriers. Breast Cancer Research, 2015, 17, 61.	2.2	26
31	Laboratory recommendations for scoring deep molecular responses following treatment for chronic myeloid leukemia. Leukemia, 2015, 29, 999-1003.	3.3	280
32	<i>FANCM</i> c.5791C>T nonsense mutation (rs144567652) induces exon skipping, affects DNA repair activity and is a familial breast cancer risk factor. Human Molecular Genetics, 2015, 24, 5345-5355.	1.4	91
33	Association of Type and Location of <i>BRCA1 </i> BRCA2 Mutations With Risk of Breast and Ovarian Cancer. JAMA - Journal of the American Medical Association, 2015, 313, 1347.	3.8	390
34	Candidate Genetic Modifiers for Breast and Ovarian Cancer Risk in <i>BRCA1</i> Mutation Carriers. Cancer Epidemiology Biomarkers and Prevention, 2015, 24, 308-316.	1.1	22
35	Dasatinib induces fast and deep responses in newly diagnosed chronic myeloid leukaemia patients in chronic phase: clinical results from a randomised phaseâ€2 study ( <scp>N</scp> ord <scp>CML</scp> 006). European Journal of Haematology, 2015, 94, 243-250.	1.1	61
36	Mature, Adaptive-like CD56DIM NK Cells in Chronic Myeloid Leukemia Patients in Treatment Free Remission. Blood, 2015, 126, 343-343.	0.6	3

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37	RNA-seq identifies clinically relevant fusion genes in leukemia including a novel MEF2D/CSF1R fusion responsive to imatinib. Leukemia, 2014, 28, 977-979.	3.3	49
38	DNA Glycosylases Involved in Base Excision Repair May Be Associated with Cancer Risk in BRCA1 and BRCA2 Mutation Carriers. PLoS Genetics, 2014, 10, e1004256.	1.5	47
39	Functional implications of the p.Cys680Arg mutation in the MLH1 mismatch repair protein. Molecular Genetics & Canada Gen	0.6	1
40	Associations of common breast cancer susceptibility alleles with risk of breast cancer subtypes in BRCA1 and BRCA2 mutation carriers. Breast Cancer Research, 2014, 16, 3416.	2.2	57
41	Interim Analysis of a Pan European Stop Tyrosine Kinase Inhibitor Trial in Chronic Myeloid Leukemia : The EURO-SKI study. Blood, 2014, 124, 151-151.	0.6	38
42	Early Disease Relapse after Tyrosine Kinase Inhibitor Treatment Discontinuation in CML Is Related Both to Low Number and Impaired Function of NK-Cells. Blood, 2014, 124, 812-812.	0.6	33
43	Potential of Digital PCR in CML Calibration. Blood, 2014, 124, 1817-1817.	0.6	0
44	Impact of malignant stem cell burden on therapy outcome in newly diagnosed chronic myeloid leukemia patients. Leukemia, 2013, 27, 1520-1526.	3.3	60
45	Whole-genome-amplified DNA as a source for mutational analysis underestimates the frequency of mutations in pediatric acute myeloid leukemia. Leukemia, 2013, 27, 510-512.	3.3	1
46	Analysis of Mice Lacking the Heparin-Binding Splice Isoform of Platelet-Derived Growth Factor A. Molecular and Cellular Biology, 2013, 33, 4030-4040.	1.1	8
47	Dasatinib Treatment Induces Fast and Deep Responses In Newly Diagnosed Chronic Myeloid Leukemia (CML) Patients In Chronic Phase: Clinical Results From a Randomized Phase 2 Study (NordCML006). Blood, 2013, 122, 4032-4032.	0.6	4
48	Distinct transcriptional control in major immunogenetic subsets of chronic lymphocytic leukemia exhibiting subset-biased global DNA methylation profiles. Epigenetics, 2012, 7, 1435-1442.	1.3	37
49	Common Variants at the 19p13.1 and <i>ZNF365</i> Loci Are Associated with ER Subtypes of Breast Cancer and Ovarian Cancer Risk in <i>BRCA1</i> and <i>BRCA2</i> Mutation Carriers. Cancer Epidemiology Biomarkers and Prevention, 2012, 21, 645-657.	1.1	47
50	Response: high ERG gene expression is an unfavorable prognostic marker in pediatric acute myeloid leukemia. Blood, 2012, 119, 1087-1088.	0.6	7
51	Effects of BRCA2 cis-regulation in normal breast and cancer risk amongst BRCA2 mutation carriers. Breast Cancer Research, 2012, 14, R63.	2.2	22
52	Evaluation of chromosome 6p22 as a breast cancer risk modifier locus in a follow-up study of BRCA2 mutation carriers. Breast Cancer Research and Treatment, 2012, 136, 295-302.	1.1	4
53	Breast Cancer Risk and 6q22.33: Combined Results from Breast Cancer Association Consortium and Consortium of Investigators on Modifiers of BRCA1/2. PLoS ONE, 2012, 7, e35706.	1.1	11
54	Association of PHB 1630 C>T and MTHFR 677 C>T polymorphisms with breast and ovarian cancer risk in BRCA1/2 mutation carriers: results from a multicenter study. British Journal of Cancer, 2012, 106, 2016-2024.	2.9	27

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55	Ovarian cancer susceptibility alleles and risk of ovarian cancer in <i>BRCA1</i> and <i>BRCA2</i> mutation carriers. Human Mutation, 2012, 33, 690-702.	1.1	34
56	Mantle cell lymphoma displays a homogenous methylation profile: A comparative analysis with chronic lymphocytic leukemia. American Journal of Hematology, 2012, 87, 361-367.	2.0	13
57	Exploring the link between MORF4L1 and risk of breast cancer. Breast Cancer Research, 2011, 13, R40.	2.2	23
58	Common breast cancer susceptibility alleles are associated with tumour subtypes in BRCA1 and BRCA2 mutation carriers: results from the Consortium of Investigators of Modifiers of BRCA1/2. Breast Cancer Research, 2011, 13, R110.	2.2	71
59	Bilateral Prophylactic Mastectomy in Swedish Women at High Risk of Breast Cancer. Annals of Surgery, 2011, 253, 1147-1154.	2.1	54
60	Combination of pegylated IFN- $\hat{l}$ ±2b with imatinib increases molecular response rates in patients with low- or intermediate-risk chronic myeloid leukemia. Blood, 2011, 118, 3228-3235.	0.6	174
61	The frequency and prognostic impact of dic(9;20)(p13.2;q11.2) in childhood B-cell precursor acute lymphoblastic leukemia: results from the NOPHO ALL-2000 trial. Leukemia, 2011, 25, 622-628.	3.3	25
62	Evaluation of the XRCC1 gene as a phenotypic modifier in BRCA1/2 mutation carriers. Results from the consortium of investigators of modifiers of BRCA1/BRCA2. British Journal of Cancer, 2011, 104, 1356-1361.	2.9	7
63	Presence of FLT3-ITD and high BAALC expression are independent prognostic markers in childhood acute myeloid leukemia. Blood, 2011, 118, 5905-5913.	0.6	83
64	Haplotype structure in Ashkenazi Jewish BRCA1 and BRCA2 mutation carriers. Human Genetics, 2011, 130, 685-699.	1.8	18
65	Highâ€resolution genomic screening in mantle cell lymphomaâ€"specific changes correlate with genomic complexity, the proliferation signature and survival. Genes Chromosomes and Cancer, 2011, 50, 113-121.	1.5	29
66	Common alleles at 6q25.1 and 1p11.2 are associated with breast cancer risk for BRCA1 and BRCA2 mutation carriers. Human Molecular Genetics, 2011, 20, 3304-3321.	1.4	68
67	Common Genetic Variation at BARD1 Is Not Associated with Breast Cancer Risk in BRCA1 or BRCA2 Mutation Carriers. Cancer Epidemiology Biomarkers and Prevention, 2011, 20, 1032-1038.	1.1	16
68	Common variants of the BRCA1 wild-type allele modify the risk of breast cancer in BRCA1 mutation carriers. Human Molecular Genetics, 2011, 20, 4732-4747.	1.4	32
69	Impact of TP53 mutation and 17p deletion in mantle cell lymphoma. Leukemia, 2011, 25, 1904-1908.	3.3	88
70	Interplay between BRCA1 and RHAMM Regulates Epithelial Apicobasal Polarization and May Influence Risk of Breast Cancer. PLoS Biology, 2011, 9, e1001199.	2.6	91
71	A locus on 19p13 modifies risk of breast cancer in BRCA1 mutation carriers and is associated with hormone receptor–negative breast cancer in the general population. Nature Genetics, 2010, 42, 885-892.	9.4	309
72	Common Breast Cancer Susceptibility Alleles and the Risk of Breast Cancer for <i>BRCA1</i> and <i>BRCA2</i> Mutation Carriers: Implications for Risk Prediction. Cancer Research, 2010, 70, 9742-9754.	0.4	169

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73	How to Handle Genetic Information: A Comparison of Attitudes among Patients and the General Population. Public Health Genomics, 2010, 13, 396-405.	0.6	7
74	Association of the Variants < i > CASP8 < /i > D302H and < i > CASP10 < /i > V410I with Breast and Ovarian Cancer Risk in < i > BRCA1 < /i > and < i > BRCA2 < /i > Mutation Carriers. Cancer Epidemiology Biomarkers and Prevention, 2010, 19, 2859-2868.	1.1	37
75	Establishment of the first World Health Organization International Genetic Reference Panel for quantitation of BCR-ABL mRNA. Blood, 2010, 116, e111-e117.	0.6	141
76	Stability of Conversion Factors for BCR-ABL Monitoring -– Implications for the Frequency of Validation Rounds. Blood, 2010, 116, 893-893.	0.6	16
77	Genome-Wide Array-Based Methylation Profiling Reveals Preferential Methylation of Homeobox Transcription Factor Genes In Mantle Cell Lymphoma and Pro-Apoptotic Genes In Chronic Lymphocytic Leukemia. Blood, 2010, 116, 536-536.	0.6	O
78	Abstract P6-10-03: Bilateral Prophylactic Mastectomy in Swedish Women at High Risk of Breast Cancer $\hat{a} \in$ " A National Survey. , 2010, , .		1
79	Common variants in LSP1, 2q35 and 8q24 and breast cancer risk for BRCA1 and BRCA2 mutation carriers. Human Molecular Genetics, 2009, 18, 4442-4456.	1.4	99
80	Clinical and cytogenetic features of a populationâ€based consecutive series of 285 pediatric Tâ€cell acute lymphoblastic leukemias: Rare Tâ€cell receptor gene rearrangements are associated with poor outcome. Genes Chromosomes and Cancer, 2009, 48, 795-805.	1.5	33
81	Evaluation of a candidate breast cancer associated SNP in ERCC4 as a risk modifier in BRCA1 and BRCA2 mutation carriers. Results from the Consortium of Investigators of Modifiers of BRCA1/BRCA2 (CIMBA). British Journal of Cancer, 2009, 101, 2048-2054.	2.9	15
82	Comparison of imatinib 400 mg and 800 mg daily in the front-line treatment of high-risk, Philadelphia-positive chronic myeloid leukemia: a European LeukemiaNet Study. Blood, 2009, 113, 4497-4504.	0.6	173
83	A Randomized Phase II Study Comparing Imatinib and the Combination of Imatinib and Pegylated Interferon Alpha-2b in Newly Diagnosed Non-High Risk Chronic Myeloid Leukemia (CML) Patients in Complete Hematological Remission After Imatinib Induction Therapy Blood, 2009, 114, 3280-3280.	0.6	2
84	Uâ€2973, a novel Bâ€cell line established from a patient with a mature Bâ€cell leukemia displaying concurrent t(14;18) and <i>MYC</i> translocation to a nonâ€ <i>IG</i> gene partner. European Journal of Haematology, 2008, 81, 218-225.	1.1	6
85	A specific requirement for PDGF-C in palate formation and PDGFR-α signaling. Nature Genetics, 2004, 36, 1111-1116.	9.4	199
86	PDGF-C is a new protease-activated ligand for the PDGF α-receptor. Nature Cell Biology, 2000, 2, 302-309.	4.6	548
87	PDGFB Regulates the Development of the Labyrinthine Layer of the Mouse Fetal Placenta. Developmental Biology, 1999, 212, 124-136.	0.9	108
88	PDGF-A Signaling Is a Critical Event in Lung Alveolar Myofibroblast Development and Alveogenesis. Cell, 1996, 85, 863-873.	13.5	787