List of Publications by Year in descending order

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		38742	25787
212	13,133	50	108
papers	citations	h-index	g-index
221	221	221	20939
all docs	docs citations	times ranked	citing authors

Кемісні Уоснірл

#	Article	IF	CITATIONS
1	Dyserythropoietic anaemia with an intronic GATA1 splicing mutation in patients suspected to have Diamondâ€Blackfan anaemia. EJHaem, 2022, 3, 163-167.	1.0	1
2	Description of longitudinal tumor evolution in a case of multiply relapsed clear cell sarcoma of the kidney. Cancer Reports, 2022, 5, e1458.	1.4	3
3	Novel TENM3–ALK fusion is an alternate mechanism for ALK activation in neuroblastoma. Oncogene, 2022, 41, 2789-2797.	5.9	3
4	The landscape of genetic aberrations in myxofibrosarcoma. International Journal of Cancer, 2022, 151, 565-577.	5.1	13
5	Abstract 6085: Clonal evolution of mammary epithelial cells into breast cancers. Cancer Research, 2022, 82, 6085-6085.	0.9	0
6	Amplified <i>EPOR</i> / <i>JAK2</i> Genes Define a Unique Subtype of Acute Erythroid Leukemia. Blood Cancer Discovery, 2022, 3, 410-427.	5.0	7
7	Frequent mutations in HLA and related genes in extranodal NK/T cell lymphomas. Leukemia and Lymphoma, 2021, 62, 95-103.	1.3	12
8	Acquisition of monosomy 7 and a <i>RUNX1</i> mutation in Pearson syndrome. Pediatric Blood and Cancer, 2021, 68, e28799.	1.5	9
9	Reduced-intensity conditioning is effective for hematopoietic stem cell transplantation in young pediatric patients with Diamond–Blackfan anemia. Bone Marrow Transplantation, 2021, 56, 1013-1020.	2.4	10
10	Association of high-risk neuroblastoma classification based on expression profiles with differentiation and metabolism. PLoS ONE, 2021, 16, e0245526.	2.5	11
11	Frequent genetic alterations in immune checkpoint–related genes in intravascular large B-cell lymphoma. Blood, 2021, 137, 1491-1502.	1.4	49
12	Poor Myocardial Compaction in a Patient with Recessive <i>MYL2</i> Myopathy. International Heart Journal, 2021, 62, 445-447.	1.0	1
13	Clinical significance of RAS pathway alterations in pediatric acute myeloid leukemia. Haematologica, 2021, , .	3.5	10
14	Clonal evolution and clinical implications of genetic abnormalities in blastic transformation of chronic myeloid leukaemia. Nature Communications, 2021, 12, 2833.	12.8	39
15	A Possible Association Between a Nucleotideâ€Binding Domain LRRâ€Containing Protein Family PYDâ€Containing Protein 1 Mutation and an Autoinflammatory Disease Involving Liver Cirrhosis. Hepatology, 2021, 74, 2296-2299.	7.3	6
16	Clonal evidence for the development of neuroblastoma with extensive copyâ€neutral loss of heterozygosity arising in a mature teratoma. Cancer Science, 2021, 112, 2921-2927.	3.9	3
17	Molecular classification and diagnostics of upper urinary tract urothelial carcinoma. Cancer Cell, 2021, 39, 793-809.e8.	16.8	65
18	Optimization of prediction methods for risk assessment of pathogenic germline variants in the Japanese population. Cancer Science, 2021, 112, 3338-3348.	3.9	3

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19	Droplet digital polymerase chain reaction assay for the detection of the minor clone of <i>KIT</i> D816V in paediatric acute myeloid leukaemia especially showing <i>RUNX1â€RUNX1T1</i> transcripts. British Journal of Haematology, 2021, 194, 414-422.	2.5	2
20	Successful treatment of hepatosplenic T-cell lymphoma with fludarabine, high-dose cytarabine and subsequent unrelated umbilical cord blood transplantation. International Journal of Hematology, 2021, , 1.	1.6	0
21	Unbiased Detection of Driver Mutations in Extramammary Paget Disease. Clinical Cancer Research, 2021, 27, 1756-1765.	7.0	24
22	EPOR/JAK/STAT Signaling Pathway As Therapeutic Target of Acute Erythroid Leukemia. Blood, 2021, 138, 610-610.	1.4	2
23	Clonal Evolution Pattern and Prognostic Significance of Clonal Architecture in KMT2A-Rearranged Acute Myeloid Leukemia. Blood, 2021, 138, 2358-2358.	1.4	0
24	A founder variant in the South Asian population leads to a high prevalence of <i>FANCL</i> Fanconi anemia cases in India. Human Mutation, 2020, 41, 122-128.	2.5	10
25	Novel DDX41 variants in Thai patients with myeloid neoplasms. International Journal of Hematology, 2020, 111, 241-246.	1.6	20
26	DNA methylation-based classification reveals difference between pediatric T-cell acute lymphoblastic leukemia and normal thymocytes. Leukemia, 2020, 34, 1163-1168.	7.2	14
27	Single-cell analysis based dissection of clonality in myelofibrosis. Nature Communications, 2020, 11, 73.	12.8	46
28	Frequent mutations that converge on the NFKBIZ pathway in ulcerative colitis. Nature, 2020, 577, 260-265.	27.8	168
29	Fusion partner–specific mutation profiles and KRAS mutations as adverse prognostic factors in MLL-rearranged AML. Blood Advances, 2020, 4, 4623-4631.	5.2	7
30	Genetic and clinical landscape of breast cancers with germline BRCA1/2 variants. Communications Biology, 2020, 3, 578.	4.4	20
31	Comprehensive genetic analysis of pediatric germ cell tumors identifies potential drug targets. Communications Biology, 2020, 3, 544.	4.4	9
32	Integrated multiomics analysis of hepatoblastoma unravels its heterogeneity and provides novel druggable targets. Npj Precision Oncology, 2020, 4, 20.	5.4	30
33	Clinical utility of target captureâ€based panel sequencing in hematological malignancies: A multicenter feasibility study. Cancer Science, 2020, 111, 3367-3378.	3.9	11
34	Landscape of driver mutations and their clinical impacts in pediatric B-cell precursor acute lymphoblastic leukemia. Blood Advances, 2020, 4, 5165-5173.	5.2	33
35	Novel COL4A1 mutations identified in infants with congenital hemolytic anemia in association with brain malformations. Human Genome Variation, 2020, 7, 42.	0.7	2
36	Two Aldehyde Clearance Systems Are Essential to Prevent Lethal Formaldehyde Accumulation in Mice and Humans. Molecular Cell, 2020, 80, 996-1012.e9.	9.7	92

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37	Genomic analysis of multiple myeloma using targeted capture sequencing in the Japanese cohort. British Journal of Haematology, 2020, 191, 755-763.	2.5	0
38	Combined Cohesin–RUNX1 Deficiency Synergistically Perturbs Chromatin Looping and Causes Myelodysplastic Syndromes. Cancer Discovery, 2020, 10, 836-853.	9.4	51
39	Clinical Impacts of Germline <i>DDX41</i> Mutations on Myeloid Neoplasms. Blood, 2020, 136, 38-40.	1.4	7
40	Prognostic Relevance of Genetic Abnormalities in Blastic Transformation of Chronic Myeloid Leukemia. Blood, 2020, 136, 3-4.	1.4	3
41	Hematopoietic stem cell transplantation for progressive combined immunodeficiency and lymphoproliferation in patients with activated phosphatidylinositol-3-OH kinase I´ syndrome type 1. Journal of Allergy and Clinical Immunology, 2019, 143, 266-275.	2.9	49
42	Differential expression of individual transcript variants of PD-1 and PD-L2 genes on Th-1/Th-2 status is guaranteed for prognosis prediction in PCNSL. Scientific Reports, 2019, 9, 10004.	3.3	24
43	Mechanisms of Progression of Myeloid Preleukemia to Transformed Myeloid Leukemia in Children with Down Syndrome. Cancer Cell, 2019, 36, 123-138.e10.	16.8	93
44	Integrated genetic and epigenetic analysis revealed heterogeneity of acute lymphoblastic leukemia in Down syndrome. Cancer Science, 2019, 110, 3358-3367.	3.9	15
45	Defective Epstein–Barr virus in chronic active infection and haematological malignancy. Nature Microbiology, 2019, 4, 404-413.	13.3	152
46	Frequent structural variations involving programmed death ligands in Epstein-Barr virus-associated lymphomas. Leukemia, 2019, 33, 1687-1699.	7.2	98
47	Remission clone in acute myeloid leukemia shows growth advantage after chemotherapy but is distinct from leukemic clone. Experimental Hematology, 2019, 75, 26-30.	0.4	1
48	Molecular heterogeneity in peripheral T-cell lymphoma, not otherwise specified revealed by comprehensive genetic profiling. Leukemia, 2019, 33, 2867-2883.	7.2	148
49	KLF1 mutation E325K induces cell cycle arrest in erythroid cells differentiated from congenital dyserythropoietic anemia patient-specific induced pluripotent stem cells. Experimental Hematology, 2019, 73, 25-37.e8.	0.4	17
50	Pathogenic mutations identified by a multimodality approach in 117 Japanese Fanconi anemia patients. Haematologica, 2019, 104, 1962-1973.	3.5	22
51	Comprehensive analysis of genetic aberrations linked to tumorigenesis in regenerative nodules of liver cirrhosis. Journal of Gastroenterology, 2019, 54, 628-640.	5.1	33
52	Frequent germline mutations of HAVCR2 in sporadic subcutaneous panniculitis-like T-cell lymphoma. Blood Advances, 2019, 3, 588-595.	5.2	73
53	Transcriptome analysis offers a comprehensive illustration of the genetic background of pediatric acute myeloid leukemia. Blood Advances, 2019, 3, 3157-3169.	5.2	51
54	Ring sideroblasts in AML are associated with adverse risk characteristics and have a distinct gene expression pattern. Blood Advances, 2019, 3, 3111-3122.	5.2	6

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55	Invariant patterns of clonal succession determine specific clinical features of myelodysplastic syndromes. Nature Communications, 2019, 10, 5386.	12.8	53
56	Paraneoplastic hypereosinophilic syndrome associated with <i>IL3â€IgH</i> positive acute lymphoblastic leukemia. Pediatric Blood and Cancer, 2019, 66, e27449.	1.5	12
57	Molecular pathogenesis of disease progression in MLL-rearranged AML. Leukemia, 2019, 33, 612-624.	7.2	26
58	Age-related remodelling of oesophageal epithelia by mutated cancer drivers. Nature, 2019, 565, 312-317.	27.8	476
59	Genomic landscape and clonal evolution of acute myeloid leukemia with t(8;21): an international study on 331 patients. Blood, 2019, 133, 1140-1151.	1.4	96
60	Novel neuroblastoma amplified sequence (NBAS) mutations in a Japanese boy with fever-triggered recurrent acute liver failure. Human Genome Variation, 2019, 6, 2.	0.7	26
61	Genetic and transcriptional landscape of plasma cells in POEMS syndrome. Leukemia, 2019, 33, 1723-1735.	7.2	28
62	<scp>NOTCH</scp> 1 pathway activating mutations and clonal evolution in pediatric Tâ€cell acute lymphoblastic leukemia. Cancer Science, 2019, 110, 784-794.	3.9	26
63	Modification of cellular and humoral immunity by somatically reverted T cells in X-linked lymphoproliferative syndrome type 1. Journal of Allergy and Clinical Immunology, 2019, 143, 421-424.e11.	2.9	8
64	Associations of complementation group, ALDH2 genotype, and clonal abnormalities with hematological outcome in Japanese patients with Fanconi anemia. Annals of Hematology, 2019, 98, 271-280.	1.8	19
65	Distinct, Ethnic, Clinical, and Genetic Characteristics of Myelodysplastic Syndromes with Der(1;7). Blood, 2019, 134, 5392-5392.	1.4	2
66	PPM1D and DNMT3A Mutations in Myelodysplasia and Clonal Hematopoiesis. Blood, 2019, 134, 1709-1709.	1.4	2
67	Hematopoietic lineage distribution and evolutionary dynamics of clonal hematopoiesis. Leukemia, 2018, 32, 1908-1919.	7.2	137
68	Dysregulation of Epstein-Barr Virus Infection in Hypomorphic ZAP70 Mutation. Journal of Infectious Diseases, 2018, 218, 825-834.	4.0	22
69	Sudden Intracranial Hemorrhage in a Patient With Atypical Chronic Myeloid Leukemia in Chronic Phase. Journal of Pediatric Hematology/Oncology, 2018, 40, e553-e556.	0.6	3
70	Integrated molecular profiling of juvenile myelomonocytic leukemia. Blood, 2018, 131, 1576-1586.	1.4	78
71	Early detection and evolution of preleukemic clones in therapy-related myeloid neoplasms following autologous SCT. Blood, 2018, 131, 1846-1857.	1.4	35
72	Prognostic relevance of integrated genetic profiling in adult T-cell leukemia/lymphoma. Blood, 2018, 131, 215-225.	1.4	124

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73	Genetic heterogeneity of uncharacterized childhood autoimmune diseases with lymphoproliferation. Pediatric Blood and Cancer, 2018, 65, e26831.	1.5	18
74	Integrated Molecular Characterization of the Lethal Pediatric Cancer Pancreatoblastoma. Cancer Research, 2018, 78, 865-876.	0.9	25
75	Hidden <i>FLT3</i> -D835Y clone in <i>FLT3</i> -ITD-positive acute myeloid leukemia that evolved into very late relapse with T-lymphoblastic leukemia. Leukemia and Lymphoma, 2018, 59, 1490-1493.	1.3	2
76	Physiological Srsf2 P95H expression causes impaired hematopoietic stem cell functions and aberrant RNA splicing in mice. Blood, 2018, 131, 621-635.	1.4	64
77	Clonally related diffuse large B-cell lymphoma and interdigitating dendritic cell sarcoma sharing MYC translocation. Haematologica, 2018, 103, e553-e556.	3.5	14
78	Two siblings with familial neuroblastoma with distinct clinical phenotypes harboring an <i>ALK</i> germline mutation. Genes Chromosomes and Cancer, 2018, 57, 665-669.	2.8	2
79	Germline loss-of-function SAMD9 and SAMD9L alterations in adult myelodysplastic syndromes. Blood, 2018, 132, 2309-2313.	1.4	38
80	Gain-of-function <i>IKBKB</i> mutation causes human combined immune deficiency. Journal of Experimental Medicine, 2018, 215, 2715-2724.	8.5	69
81	Aberrant splicing and defective mRNA production induced by somatic spliceosome mutations in myelodysplasia. Nature Communications, 2018, 9, 3649.	12.8	140
82	Recurrent CCND3 mutations in MLL-rearranged acute myeloid leukemia. Blood Advances, 2018, 2, 2879-2889.	5.2	19
83	Whole-exome analysis to detect congenital hemolytic anemia mimicking congenital dyserythropoietic anemia. International Journal of Hematology, 2018, 108, 306-311.	1.6	8
84	RUNX1 mutations in pediatric acute myeloid leukemia are associated with distinct genetic features and an inferior prognosis. Blood, 2018, 131, 2266-2270.	1.4	15
85	De Novo Mutations Activating Germline TP53 in an Inherited Bone-Marrow-Failure Syndrome. American Journal of Human Genetics, 2018, 103, 440-447.	6.2	33
86	Analysis of Genomic Predispositions to Sporadic Myeloid Neoplasms Mediated By DDX41 in Japan. Blood, 2018, 132, 4371-4371.	1.4	0
87	Clinical utility of next-generation sequencing for inherited bone marrow failure syndromes. Genetics in Medicine, 2017, 19, 796-802.	2.4	66
88	<i>ASXL2</i> mutations are frequently found in pediatric AML patients with t(8;21)/ <i>RUNX1â€RUNX1T1</i> and associated with a better prognosis. Genes Chromosomes and Cancer, 2017, 56, 382-393.	2.8	18
89	Genetic abnormalities in myelodysplasia and secondary acute myeloid leukemia: impact on outcome of stem cell transplantation. Blood, 2017, 129, 2347-2358.	1.4	268
90	Exome sequencing identified <i>RPS15A</i> as a novel causative gene for Diamond-Blackfan anemia. Haematologica, 2017, 102, e93-e96.	3.5	30

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91	Hypoxic adaptation of leukemic cells infiltrating the CNS affords a therapeutic strategy targeting VEGFA. Blood, 2017, 129, 3126-3129.	1.4	23
92	Clonal evolution in myelodysplastic syndromes. Nature Communications, 2017, 8, 15099.	12.8	118
93	Common Variable Immunodeficiency Caused by FANC Mutations. Journal of Clinical Immunology, 2017, 37, 434-444.	3.8	18
94	The E-Id Protein Axis Specifies Adaptive Lymphoid Cell Identity and Suppresses Thymic Innate Lymphoid Cell Development. Immunity, 2017, 46, 818-834.e4.	14.3	73
95	Constitutional abnormalities of <i>IDH1</i> combined with secondary mutations predispose a patient with Maffucci syndrome to acute lymphoblastic leukemia. Pediatric Blood and Cancer, 2017, 64, e26647.	1.5	9
96	BCL6 locus is hypermethylated in angioimmunoblastic T-cell lymphoma. International Journal of Hematology, 2017, 105, 465-469.	1.6	23
97	Dynamics of clonal evolution in myelodysplastic syndromes. Nature Genetics, 2017, 49, 204-212.	21.4	348
98	Abnormal hematopoiesis and autoimmunity in human subjects with germline IKZF1 mutations. Journal of Allergy and Clinical Immunology, 2017, 140, 223-231.	2.9	99
99	Gene expression and risk of leukemic transformation in myelodysplasia. Blood, 2017, 130, 2642-2653.	1.4	64
100	Loss of DNA Damage Response in Neuroblastoma and Utility of a PARP Inhibitor. Journal of the National Cancer Institute, 2017, 109, .	6.3	43
101	Molecular studies reveal MLL-MLLT10/AF10 and ARID5B-MLL gene fusions displaced in a case of infantile acute lymphoblastic leukemia with complex karyotype. Oncology Letters, 2017, 14, 2295-2299.	1.8	6
102	Atypical dyskeratosis congenita diagnosed using wholeâ€exome sequencing. Pediatrics International, 2017, 59, 933-935.	0.5	2
103	Recurrent SPI1 (PU.1) fusions in high-risk pediatric T cell acute lymphoblastic leukemia. Nature Genetics, 2017, 49, 1274-1281.	21.4	100
104	Haploinsufficiency of TNFAIP3 ( A20 ) by germline mutation is involved in autoimmune lymphoproliferative syndrome. Journal of Allergy and Clinical Immunology, 2017, 139, 1914-1922.	2.9	91
105	Metachronous anaplastic sarcoma of the kidney and thyroid follicular carcinoma as manifestations of DICER1 abnormalities. Human Pathology, 2017, 61, 205-209.	2.0	18
106	Diagnostic challenge of Diamond–Blackfan anemia in mothers and children by whole-exome sequencing. International Journal of Hematology, 2017, 105, 515-520.	1.6	18
107	Autonomous feedback loop of RUNX1-p53-CBFB in acute myeloid leukemia cells. Scientific Reports, 2017, 7, 16604.	3.3	29
108	Recurrent genetic defects on chromosome 5q in myeloid neoplasms. Oncotarget, 2017, 8, 6483-6495.	1.8	34

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109	Identification of the genetic and clinical characteristics of neuroblastomas using genome-wide analysis. Oncotarget, 2017, 8, 107513-107529.	1.8	23
110	Frequent NFKBIE deletions are associated with poor outcome in primary mediastinal B-cell lymphoma. Blood, 2016, 128, 2666-2670.	1.4	82
111	Somatic mosaicism in chronic myeloid leukemia in remission. Blood, 2016, 128, 2863-2866.	1.4	13
112	Single cell genotyping of exome sequencing-identified mutations to characterize the clonal composition and evolution of inv(16) AML in a CBL mutated clonal hematopoiesis. Leukemia Research, 2016, 47, 41-46.	0.8	12
113	Aberrant PD-L1 expression through 3′-UTR disruption in multiple cancers. Nature, 2016, 534, 402-406.	27.8	536
114	ATP11C is a major flippase in human erythrocytes and its defect causes congenital hemolytic anemia. Haematologica, 2016, 101, 559-565.	3.5	72
115	The phenotype and clinical course of Japanese Fanconi Anaemia infants is influenced by patient, but not maternal <i>ALDH2</i> genotype. British Journal of Haematology, 2016, 175, 457-461.	2.5	10
116	<i>TERT</i> promoter mutations and chromosome 8p loss are characteristic of nonalcoholic fatty liver diseaseâ€related hepatocellular carcinoma. International Journal of Cancer, 2016, 139, 2512-2518.	5.1	36
117	Genomic analysis of clonal origin of Langerhans cell histiocytosis following acute lymphoblastic leukaemia. British Journal of Haematology, 2016, 175, 169-172.	2.5	12
118	Wholeâ€exome sequencing reveals the spectrum of gene mutations and the clonal evolution patterns in paediatric acute myeloid leukaemia. British Journal of Haematology, 2016, 175, 476-489.	2.5	60
119	Phosphatase and tensin homolog ( PTEN ) mutation can cause activated phosphatidylinositol 3-kinase Î́′ syndrome–like immunodeficiency. Journal of Allergy and Clinical Immunology, 2016, 138, 1672-1680.e10.	2.9	87
120	Variegated RHOA mutations in adult T-cell leukemia/lymphoma. Blood, 2016, 127, 596-604.	1.4	98
121	Mutation allele burden remains unchanged in chronic myelomonocytic leukaemia responding to hypomethylating agents. Nature Communications, 2016, 7, 10767.	12.8	177
122	ALDH2 polymorphism in patients with Diamond-Blackfan anemia in Japan. International Journal of Hematology, 2016, 103, 112-114.	1.6	1
123	PIEZO1 gene mutation in a Japanese family with hereditary high phosphatidylcholine hemolytic anemia and hemochromatosis-induced diabetes mellitus. International Journal of Hematology, 2016, 104, 125-129.	1.6	25
124	Gene Expression Profiles and Methylation Analysis in Down Syndrome Related Acute Lymphoblastic Leukemia. Blood, 2016, 128, 4084-4084.	1.4	0
125	Structural Variations Involving Programmed Death Ligands in B-Cell and T-Cell Lymphomas. Blood, 2016, 128, 4105-4105.	1.4	0
126	Recurrent VAV1 Abnormalities in Angioimmunoblastic T Cell Lymphoma. Blood, 2016, 128, 4104-4104.	1.4	1

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127	Profiling of somatic mutations in acute myeloid leukemia with FLT3-ITD at diagnosis and relapse. Blood, 2015, 126, 2491-2501.	1.4	180
128	Mutations in the Gene Encoding the E2 Conjugating Enzyme UBE2T Cause Fanconi Anemia. American Journal of Human Genetics, 2015, 96, 1001-1007.	6.2	100
129	BRCC3 mutations in myeloid neoplasms. Haematologica, 2015, 100, 1051-7.	3.5	20
130	Integrated genetic and epigenetic analysis defines novel molecular subgroups in rhabdomyosarcoma. Nature Communications, 2015, 6, 7557.	12.8	149
131	Inherited and Somatic Defects in DDX41 in Myeloid Neoplasms. Cancer Cell, 2015, 27, 658-670.	16.8	341
132	Mutational landscape and clonal architecture in grade II and III gliomas. Nature Genetics, 2015, 47, 458-468.	21.4	729
133	Integrated molecular analysis of adult T cell leukemia/lymphoma. Nature Genetics, 2015, 47, 1304-1315.	21.4	659
134	Clinical and genetic features of dyskeratosis congenita, cryptic dyskeratosis congenita, and Hoyeraal-Hreidarsson syndrome in Japan. International Journal of Hematology, 2015, 102, 544-552.	1.6	21
135	GATA2 and secondary mutations in familial myelodysplastic syndromes and pediatric myeloid malignancies. Haematologica, 2015, 100, e398-e401.	3.5	48
136	Late-Onset Combined Immunodeficiency with a Novel IL2RG Mutation and Probable Revertant Somatic Mosaicism. Journal of Clinical Immunology, 2015, 35, 610-614.	3.8	26
137	Genomon ITDetector: a tool for somatic internal tandem duplication detection from cancer genome sequencing data. Bioinformatics, 2015, 31, 116-118.	4.1	58
138	Genetic Background of Idiopathic Bone Marrow Failure Syndromes in Children. Blood, 2015, 126, 3610-3610.	1.4	2
139	Aberrant DNA Methylation Is Associated with a Poor Outcome in Juvenile Myelomonocytic Leukemia. PLoS ONE, 2015, 10, e0145394.	2.5	25
140	Genomic landscape of liposarcoma. Oncotarget, 2015, 6, 42429-42444.	1.8	94
141	Next-Generation Sequencing Reveal Proviral Genome and Transcriptome in Adult T-Cell Leukemia/Lymphoma. Blood, 2015, 126, 3882-3882.	1.4	0
142	Clonal Evolution in Relapsed Pediatric Acute Lymphoblastic Leukemia. Blood, 2015, 126, 1425-1425.	1.4	0
143	Detection of the G17V RHOA Mutation in Angioimmunoblastic T-Cell Lymphoma and Related Lymphomas Using Quantitative Allele-Specific PCR. PLoS ONE, 2014, 9, e109714.	2.5	24
144	Splicing factor mutations and cancer. Wiley Interdisciplinary Reviews RNA, 2014, 5, 445-459.	6.4	126

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145	Biallelic <i>DICER1</i> Mutations in Sporadic Pleuropulmonary Blastoma. Cancer Research, 2014, 74, 2742-2749.	0.9	67
146	Recurrent somatic mutations underlie corticotropin-independent Cushing's syndrome. Science, 2014, 344, 917-920.	12.6	177
147	The nicotinic cholinergic system is affected in rats with delayed carbon monoxide encephalopathy. Neuroscience Letters, 2014, 569, 33-37.	2.1	19
148	Somatic RHOA mutation in angioimmunoblastic T cell lymphoma. Nature Genetics, 2014, 46, 171-175.	21.4	542
149	Acquired Initiating Mutations in Early Hematopoietic Cells of CLL Patients. Cancer Discovery, 2014, 4, 1088-1101.	9.4	213
150	Whole-exome sequence analysis of ataxia telangiectasia-like phenotype. Journal of the Neurological Sciences, 2014, 340, 86-90.	0.6	12
151	DDX41 Is a Tumor Suppressor Gene Associated with Inherited and Acquired Mutations. Blood, 2014, 124, 125-125.	1.4	1
152	Chronological Analysis of Clonal Evolution in Acquired Aplastic Anemia. Blood, 2014, 124, 253-253.	1.4	4
153	Landscape of Genetic Alterations in Adult T-Cell Leukemia/Lymphoma. Blood, 2014, 124, 75-75.	1.4	1
154	In Analogy to AML, MDS Can be Sub-Classified By Ancestral Mutations. Blood, 2014, 124, 823-823.	1.4	4
155	Comprehensive Analysis of Aberrant RNA Splicing in Myelodysplastic Syndromes. Blood, 2014, 124, 826-826.	1.4	6
156	Clinical Features of Patients with ASXL1 and ASXL2 Mutations in Pediatric Acute Myeloid Leukemia. Blood, 2014, 124, 1024-1024.	1.4	0
157	The Clinical and Genetic Features of Dyskeratosis Congenita, Cryptic Dyskeratosis Congenita, and Hoyeraal-Hreidarsson Syndrome in Japan. Blood, 2014, 124, 1608-1608.	1.4	Ο
158	An Adverse Prognostic Effect of Homozygous TET2 Mutational Status on the Relapse Risk of Acute Myeloid Leukemia Patients of Normal Karyotype. Blood, 2014, 124, 1052-1052.	1.4	0
159	Whole Exome and Transcriptome Analyses in Pediatric T-Cell Acute Lymphoblastic Leukemia. Blood, 2014, 124, 3527-3527.	1.4	Ο
160	Novel Biological Effects and Distinct Patterns of Rhoa Mutations in Adult T-Cell Leukemia/Lymphoma and Angioimmunoblastic T Cell Lymphoma. Blood, 2014, 124, 2215-2215.	1.4	0
161	Clinical and Molecular Significance of Peripheral Blood Cell-Free DNA in B-Cell Lymphomas for Detection of Genetic Mutations and Correlation with Disease Status. Blood, 2014, 124, 1658-1658.	1.4	0
162	Impact and Function of Somatic PHF6 Mutations in Myeloid Neoplasms. Blood, 2014, 124, 3581-3581.	1.4	0

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163	Identification of Cell-Type-Specific Mutations in Angioimmunoblastic T-Cell Lymphoma. Blood, 2014, 124, 3025-3025.	1.4	1
164	Diagnostic Efficacy of Whole-Exome Sequencing in 250 Patients with Congenital Bone Marrow Failure. Blood, 2014, 124, 4385-4385.	1.4	0
165	Whole-Exome Sequencing Reveals a Paucity of Somatic Gene Mutations in Aplastic Anemia and Refractory Cytopenia of Childhood. Blood, 2014, 124, 4388-4388.	1.4	Ο
166	The landscape of somatic mutations in Down syndrome–related myeloid disorders. Nature Genetics, 2013, 45, 1293-1299.	21.4	324
167	Deep Sequencing in Cancer Research. Japanese Journal of Clinical Oncology, 2013, 43, 110-115.	1.3	19
168	Integrated molecular analysis of clear-cell renal cell carcinoma. Nature Genetics, 2013, 45, 860-867.	21.4	955
169	Whole Exome Sequencing Reveals Clonal Evolution Pattern and Driver Mutations Of Relapsed Pediatric AML. Blood, 2013, 122, 1410-1410.	1.4	1
170	PRPF8 Defects Cause Missplicing In Myeloid Malignancies. Blood, 2013, 122, 2838-2838.	1.4	1
171	Spliceosomal Gene LUC7L2 Mutation Causes Missplicing and Alteration Of Gene Expression In Myeloid Neoplasms. Blood, 2013, 122, 470-470.	1.4	1
172	Clinical "MUTATOME―Of Myelodysplastic Syndrome; Comparison To Primary Acute Myelogenous Leukemia. Blood, 2013, 122, 518-518.	1.4	2
173	Landscape Of Genetic Lesions In 944 Patients With Myelodysplastic Syndromes. Blood, 2013, 122, 521-521.	1.4	14
174	Somatic G17V Rhoa Mutation Specifies Angioimmunoblastic T-Cell Lymphoma. Blood, 2013, 122, 815-815.	1.4	2
175	Whole Exome Sequencing Detecting Kinesin Family Gene Defects In Myeloid Neoplasm. Blood, 2013, 122, 2762-2762.	1.4	Ο
176	Whole Exome Sequencing Shows a Paucity Of Somatic Gene Mutations In Pediatric Idiopathic Bone Marrow Failure Syndrome. Blood, 2013, 122, 3708-3708.	1.4	0
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