Andrey A Yurchenko

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	Frequency and Genomic Aspects of Intrinsic Resistance to Vismodegib in Locally Advanced Basal Cell Carcinoma. Clinical Cancer Research, 2022, 28, 1422-1432.	3.2	6
2	Increased risk of internal tumors in DNA repair-deficient xeroderma pigmentosum patients: analysis of four international cohorts. Orphanet Journal of Rare Diseases, 2022, 17, 104.	1.2	17
3	Resequencing the Yaroslavl cattle genomes reveals signatures of selection and a rare haplotype on <scp>BTA28</scp> likely to be related to breed phenotypes. Animal Genetics, 2022, 53, 680-684.	0.6	4
4	Genomic profiling of lateâ€onset basal cell carcinomas from two brothers with nevoid basal cell carcinoma syndrome. Journal of the European Academy of Dermatology and Venereology, 2021, 35, 396-402.	1.3	2
5	Resequencing and signatures of selection scan in two Siberian native sheep breeds point to candidate genetic variants for adaptation and economically important traits. Animal Genetics, 2021, 52, 126-131.	0.6	6
6	A Novel Case of Gorlin Syndrome Mosaicism Involving an SMO Gene Mutation: Clinical, Histological and Molecular Analysis of Basaloid Tumours. Acta Dermato-Venereologica, 2021, 101, adv00434.	0.6	2
7	The new Internal Transcribed Spacer 2 diagnostic tool clarifies the taxonomic position and geographic distribution of the North American malaria vector Anopheles punctipennis. Malaria Journal, 2021, 20, 141.	0.8	2
8	Demographic History, Adaptation, and NRAP Convergent Evolution at Amino Acid Residue 100 in the World Northernmost Cattle from Siberia. Molecular Biology and Evolution, 2021, 38, 3093-3110.	3.5	27
9	RAS mutations drive proliferative chronic myelomonocytic leukemia via a KMT2A-PLK1 axis. Nature Communications, 2021, 12, 2901.	5.8	44
10	Genotyping and Whole-Genome Resequencing of Welsh Sheep Breeds Reveal Candidate Genes and Variants for Adaptation to Local Environment and Socioeconomic Traits. Frontiers in Genetics, 2021, 12, 612492.	1.1	8
11	DNA polymerase zeta contributes to heterochromatin replication to prevent genome instability. EMBO Journal, 2021, 40, e104543.	3.5	12
12	Whole-Genome Resequencing Points to Candidate DNA Loci Affecting Body Temperature under Cold Stress in Siberian Cattle Populations. Life, 2021, 11, 959.	1.1	8
13	The functional genetic architecture of egg-laying and live-bearing reproduction in common lizards. Nature Ecology and Evolution, 2021, 5, 1546-1556.	3.4	19
14	Genomic differentiation and intercontinental population structure of mosquito vectors Culex pipiens pipiens and Culex pipiens molestus. Scientific Reports, 2020, 10, 7504.	1.6	24
15	XPC deficiency increases risk of hematologic malignancies through mutator phenotype and characteristic mutational signature. Nature Communications, 2020, 11, 5834.	5.8	17
16	Chromosome-Level Assembly of the Common Lizard (Zootoca vivipara) Genome. Genome Biology and Evolution, 2020, 12, 1953-1960.	1.1	13
17	Chromosome and Genome Divergence between the Cryptic Eurasian Malaria Vector-Species Anopheles messeae and Anopheles daciae. Genes, 2020, 11, 165.	1.0	17
18	The Iberian legacy into a young genetic xeroderma pigmentosum cluster in central Brazil. Mutation Research - Genetic Toxicology and Environmental Mutagenesis, 2020, 852, 503164.	0.9	2

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19	Parallelism in eco-morphology and gene expression despite variable evolutionary and genomic backgrounds in a Holarctic fish. PLoS Genetics, 2020, 16, e1008658.	1.5	73
20	First description of ultramutated endometrial cancer caused by germline loss-of-function and somatic exonuclease domain mutations in POLE gene. Genetics and Molecular Biology, 2020, 43, e20200100.	0.6	2
21	A rare splice site mutation in the gene encoding glucokinase/hexokinase 4 in a patient with MODY type 2. Vavilovskii Zhurnal Genetiki I Selektsii, 2020, 24, 299-305.	0.4	1
22	Association of ABCB9 and COL22A1 Gene Polymorphism with Human Predisposition to Severe Forms of Tick-Borne Encephalitis. Russian Journal of Genetics, 2019, 55, 368-377.	0.2	4
23	Exome-wide search and functional annotation of genes associated in patients with severe tick-borne encephalitis in a Russian population. BMC Medical Genomics, 2019, 12, 61.	0.7	9
24	High-density genotyping reveals signatures of selection related to acclimation and economically important traits in 15 local sheep breeds from Russia. BMC Genomics, 2019, 20, 294.	1.2	57
25	Whole Genome Sequencing and Re-sequencing of the Sable Antelope (<i>Hippotragus niger</i>): A Resource for Monitoring Diversity in <i>ex Situ</i> and <i>in Situ</i> Populations. G3: Genes, Genomes, Genetics, 2019, 9, 1785-1793.	0.8	18
26	Exome-wide survey of the Siberian Caucasian population. BMC Medical Genetics, 2019, 20, 51.	2.1	2
27	Genome-wide association study and scan for signatures of selection point to candidate genes for body temperature maintenance under the cold stress in Siberian cattle populations. BMC Genetics, 2019, 20, 26.	2.7	25
28	Population genetic structure and phylogeography of sterlet (<i>Acipenser ruthenus</i> ,) Tj ETQq0 0 0 rgBT /Ove and Analysis, 2019, 30, 156-164.	erlock 107 0.7	Tf 50 387 Td (5
29	Transcriptome of the bivalve Limecola balthica L. from Western Pacific: A new resource for studies of European populations. Marine Genomics, 2018, 40, 58-63.	0.4	4
30	Robust forensic matching of confiscated horns to individual poached African rhinoceros. Current Biology, 2018, 28, R13-R14.	1.8	27
31	A matrix metalloproteinase 9 (MMP9) gene single nucleotide polymorphism is associated with predisposition to tick-borne encephalitis virus-induced severe central nervous system disease. Ticks and Tick-borne Diseases, 2018, 9, 763-767.	1.1	24
32	Genome-wide genotyping uncovers genetic profiles and history of the Russian cattle breeds. Heredity, 2018, 120, 125-137.	1.2	61
33	B Chromosomes of the Asian Seabass (Lates calcarifer) Contribute to Genome Variations at the Level of Individuals and Populations. Genes, 2018, 9, 464.	1.0	11
34	Cloning and Functional Analysis of SaCLCc1, a Gene Belonging to the Chloride Channel Family (CLC), from the Halophyte Suaeda altissima (L.) Pall Doklady Biochemistry and Biophysics, 2018, 481, 186-189.	0.3	8
35	Scans for signatures of selection in Russian cattle breed genomes reveal new candidate genes for environmental adaptation and acclimation. Scientific Reports, 2018, 8, 12984.	1.6	85
36	Whole-Genome Analysis of <i>Mycobacterium tuberculosis</i> from Patients with Tuberculous Spondylitis, Russia. Emerging Infectious Diseases, 2018, 24, 579-583.	2.0	9

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37	De novo transcriptome assembly, annotation and comparison of four ecological and evolutionary model salmonid fish species. BMC Genomics, 2018, 19, 32.	1.2	63
38	Analytical "bake-off―of whole genome sequencing quality for the Genome Russia project using a small cohort for autoimmune hepatitis. PLoS ONE, 2018, 13, e0200423.	1.1	7
39	In silico Analyses of Transcriptomes of the Marine Green Microalga Dunaliella tertiolecta: Identification of Sequences Encoding P-type ATPases. Molecular Biology, 2018, 52, 520-531.	0.4	8
40	Chromosomal-Level Assembly of the Asian Seabass Genome Using Long Sequence Reads and Multi-layered Scaffolding. PLoS Genetics, 2016, 12, e1005954.	1.5	105
41	Pangolin genomes and the evolution of mammalian scales and immunity. Genome Research, 2016, 26, 1312-1322.	2.4	95
42	Comparison of carnivore, omnivore, and herbivore mammalian genomes with a new leopard assembly. Genome Biology, 2016, 17, 211.	3.8	101
43	The Population Origins and Expansion of Feral Cats in Australia. Journal of Heredity, 2016, 107, 104-114.	1.0	21
44	Genomic legacy of the African cheetah, Acinonyx jubatus. Genome Biology, 2015, 16, 277.	3.8	167
45	Genome-wide Evidence Reveals that African and Eurasian Golden Jackals Are Distinct Species. Current Biology, 2015, 25, 2158-2165.	1.8	156
46	Eco-geographic units, population hierarchy, and a two-level conservation strategy with reference to a critically endangered salmonid, Sakhalin taimen Parahucho perryi. Conservation Genetics, 2015, 16, 431-441.	0.8	14
47	A panel of microsatellite loci for population studies of Sakhalin taimen Parahucho perryi (Brevoort). Russian Journal of Genetics, 2012, 48, 831-837.	0.2	2