Haiyi Lou

List of Publications by Year in descending order

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HAIVILOU

#	Article	IF	CITATIONS
1	Improved NGS variant calling tool for the <i>PRSS1–PRSS2</i> locus. Gut, 2023, 72, 210-212.	12.1	3
2	Haplotype-resolved de novo assembly of a Tujia genome suggests the necessity for high-quality population-specific genome references. Cell Systems, 2022, 13, 321-333.e6.	6.2	10
3	Refining models of archaic admixture in Eurasia with ArchaicSeeker 2.0. Nature Communications, 2021, 12, 6232.	12.8	19
4	<i>De novo</i> assembly of a Tibetan genome and identification of novel structural variants associated with high-altitude adaptation. National Science Review, 2020, 7, 391-402.	9.5	28
5	Prioritizing natural-selection signals from the deep-sequencing genomic data suggests multi-variant adaptation in Tibetan highlanders. National Science Review, 2019, 6, 1201-1222.	9.5	30
6	Analysis of five deep-sequenced trio-genomes of the Peninsular Malaysia Orang Asli and North Borneo populations. BMC Genomics, 2019, 20, 842.	2.8	3
7	Genome-wide comparison of allele-specific gene expression between African and European populations. Human Molecular Genetics, 2018, 27, 1067-1077.	2.9	15
8	A missense point mutation in COL10A1 identified with whole-genome deep sequencing in a 7-generation Pakistan dwarf family. Heredity, 2018, 120, 83-89.	2.6	6
9	A metagenomic approach to dissect the genetic composition of enterotypes in Han Chinese and two Muslim groups. Systematic and Applied Microbiology, 2018, 41, 1-12.	2.8	24
10	Genome-wide variants of Eurasian facial shape differentiation and a prospective model of DNA based face prediction. Journal of Genetics and Genomics, 2018, 45, 419-432.	3.9	38
11	Genetic History of Xinjiang's Uyghurs Suggests Bronze Age Multiple-Way Contacts in Eurasia. Molecular Biology and Evolution, 2017, 34, 2572-2582.	8.9	63
12	CNVbase: Batch identification of novel and rare copy number variations based on multi-ethnic population data. Journal of Genetics and Genomics, 2017, 44, 367-370.	3.9	1
13	Assessing genome-wide copy number variation in the Han Chinese population. Journal of Medical Genetics, 2017, 54, 685-692.	3.2	7
14	Differentiated demographic histories and local adaptations between Sherpas and Tibetans. Genome Biology, 2017, 18, 115.	8.8	67
15	Genome-wide scans reveal variants at EDAR predominantly affecting hair straightness in Han Chinese and Uyghur populations. Human Genetics, 2016, 135, 1279-1286.	3.8	27
16	Ancestral Origins and Genetic History of Tibetan Highlanders. American Journal of Human Genetics, 2016, 99, 580-594.	6.2	208
17	Copy number variations and genetic admixtures in three Xinjiang ethnic minority groups. European Journal of Human Genetics, 2015, 23, 536-542.	2.8	22
18	A 3.4-kb Copy-Number Deletion near EPAS1 Is Significantly Enriched in High-Altitude Tibetans but Absent from the Denisovan Sequence. American Journal of Human Genetics, 2015, 97, 54-66.	6.2	69

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19	Quantitating and Dating Recent Gene Flow between European and East Asian Populations. Scientific Reports, 2015, 5, 9500.	3.3	25
20	Genetic architectures of ADME genes in five Eurasian admixed populations and implications for drug safety and efficacy. Journal of Medical Genetics, 2014, 51, 614-622.	3.2	22
21	A panel of ancestry informative markers to estimate and correct potential effects of population stratification in Han Chinese. European Journal of Human Genetics, 2014, 22, 248-253.	2.8	39
22	A Map of Copy Number Variations in Chinese Populations. PLoS ONE, 2011, 6, e27341.	2.5	44
23	A Genome-Wide Search for Signals of High-Altitude Adaptation in Tibetans. Molecular Biology and Evolution, 2011, 28, 1003-1011.	8.9	311
24	Genomic Dissection of Population Substructure of Han Chinese and Its Implication in Association Studies. American Journal of Human Genetics, 2009, 85, 762-774.	6.2	338