## **Resource Summary Report**

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# <u>GATK</u>

RRID:SCR\_001876 Type: Tool

**Proper Citation** 

GATK (RRID:SCR\_001876)

#### **Resource Information**

URL: https://software.broadinstitute.org/gatk/

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**Description:** A software package to analyze next-generation resequencing data. The toolkit offers a wide variety of tools, with a primary focus on variant discovery and genotyping as well as strong emphasis on data quality assurance. Its robust architecture, powerful processing engine and high-performance computing features make it capable of taking on projects of any size. This software library makes writing efficient analysis tools using next-generation sequencing data very easy, and second it's a suite of tools for working with human medical resequencing projects such as 1000 Genomes and The Cancer Genome Atlas. These tools include things like a depth of coverage analyzers, a quality score recalibrator, a SNP/indel caller and a local realigner. (entry from Genetic Analysis Software)

Abbreviations: GATK

Synonyms: Genome Analysis ToolKit

**Resource Type:** software resource, data analysis software, software application, software library, data processing software, software toolkit

Defining Citation: PMID:21478889

Keywords: gene, genetic, genomic, next-generation resequencing, bio.tools

Funding:

Resource Name: GATK

Resource ID: SCR\_001876

Alternate IDs: nlx\_154324, OMICS\_00286, biotools:gatk

Alternate URLs: http://www.broadinstitute.org/gsa/wiki/index.php/The\_Genome\_Analysis\_Toolkit, https://bio.tools/gatk

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### **Ratings and Alerts**

No rating or validation information has been found for GATK.

No alerts have been found for GATK.

### Data and Source Information

Source: SciCrunch Registry

#### **Usage and Citation Metrics**

We found 14330 mentions in open access literature.

Listed below are recent publications. The full list is available at <u>NIF</u>.

Liu JN, et al. (2025) Pan-genome analyses of 11 Fraxinus species provide insights into salt adaptation in ash trees. Plant communications, 6(1), 101137.

Alfayyadh MM, et al. (2025) PathVar: A Customisable NGS Variant Calling Algorithm Implicates Novel Candidate Genes and Pathways in Hemiplegic Migraine. Clinical genetics, 107(2), 157.

Heimer G, et al. (2025) Biallelic PIGM Coding Variant Causes Intractable Epilepsy and Intellectual Disability Without Thrombotic Events. Clinical genetics, 107(2), 179.

Shi W, et al. (2025) VDGE: a data repository of variation database for gene-edited animals across multiple species. Nucleic acids research, 53(D1), D1250.

Xu X, et al. (2025) Redefining the accumulated temperature index for accurate prediction of rice flowering time in diverse environments. Plant biotechnology journal, 23(1), 302.

Van Haute L, et al. (2025) Pathogenic PDE12 variants impair mitochondrial RNA processing causing neonatal mitochondrial disease. EMBO molecular medicine, 17(1), 193.

McGuire E, et al. (2025) A case of penicillin-resistant group B Streptococcus isolated from a patient in the UK. The Journal of antimicrobial chemotherapy, 80(2), 399.

Wang J, et al. (2025) Research note: A low-density SNP genotyping panel for Chinese native chickens. Poultry science, 104(1), 104609.

Rostamzadeh Mahdabi E, et al. (2025) Comparative Analysis of Runs of Homozygosity Islands in Indigenous and Commercial Chickens Revealed Candidate Loci for Disease Resistance and Production Traits. Veterinary medicine and science, 11(1), e70074.

Tejedor JR, et al. (2025) Integration of multi-omics layers empowers precision diagnosis through unveiling pathogenic mechanisms on maple syrup urine disease. Journal of inherited metabolic disease, 48(1), e12829.

Assis BA, et al. (2025) Genomic signatures of adaptation in native lizards exposed to humanintroduced fire ants. Nature communications, 16(1), 89.

Yi Y, et al. (2025) Mitochondrial-cytochrome c oxidase II promotes glutaminolysis to sustain tumor cell survival upon glucose deprivation. Nature communications, 16(1), 212.

Aizpurua-Iraola J, et al. (2025) A reduction in effective population size has not relaxed purifying selection in the human population of Eivissa (Balearic Islands). Scientific reports, 15(1), 660.

Sasa N, et al. (2025) Blood DNA virome associates with autoimmune diseases and COVID-19. Nature genetics, 57(1), 65.

Mapendano CK, et al. (2025) Longer survival with precision medicine in late-stage cancer patients. ESMO open, 10(1), 104089.

Bayam E, et al. (2025) Bi-allelic variants in WDR47 cause a complex neurodevelopmental syndrome. EMBO molecular medicine, 17(1), 129.

Peillard-Fiorente F, et al. (2025) Point mutations in functionally diverse genes are associated with increased natural DNA transformation in multidrug resistant Streptococcus pneumoniae. Nucleic acids research, 53(1).

Yang L, et al. (2025) A novel de novo GABRA2 gene missense variant causing developmental epileptic encephalopathy in a Chinese patient. Annals of clinical and translational neurology, 12(1), 137.

Lee H, et al. (2025) A Korean Patient With Leber Congenital Amaurosis and a Homozygous RPE65 Variant Originating From a Paternal Uniparental Isodisomy. Molecular genetics & genomic medicine, 13(1), e70060.

Wang M, et al. (2025) YHSeqY3000 panel captures all founding lineages in the Chinese