## **Resource Summary Report**

Generated by <u>dkNET</u> on May 18, 2025

# **Ngmlr**

RRID:SCR\_017620 Type: Tool

**Proper Citation** 

Ngmlr (RRID:SCR\_017620)

#### **Resource Information**

URL: https://github.com/philres/ngmlr

Proper Citation: Ngmlr (RRID:SCR\_017620)

**Description:** Software tool as long read mapper designed to align PacBio or Oxford Nanopore reads to reference genome and optimized for structural variation detection.

Abbreviations: NGMLR

Synonyms: coNvex Gap-cost alignMent for Long Reads

**Resource Type:** software application, alignment software, data processing software, image analysis software, software resource

Defining Citation: PMID:29713083

**Keywords:** Long, read, mapper, align, PacBio, Oxford Nanopore, read, reference, genome, structural, variantion, detection, bio.tools

**Funding:** National Science Foundation ; NHGRI R01 HG006677; NHGRI UM1 HG008898

Availability: Free, Available for download, Freely available

Resource Name: Ngmlr

Resource ID: SCR\_017620

Alternate IDs: biotools:ngmlr

Alternate URLs: https://bio.tools/ngmlr

License: MIT License

**Record Creation Time:** 20220129T080336+0000

Record Last Update: 20250517T060328+0000

### **Ratings and Alerts**

No rating or validation information has been found for Ngmlr.

No alerts have been found for Ngmlr.

### Data and Source Information

Source: SciCrunch Registry

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

We found 28 mentions in open access literature.

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

Liang X, et al. (2024) Genome comparisons reveal accessory genes crucial for the evolution of apple Glomerella leaf spot pathogenicity in Colletotrichum fungi. Molecular plant pathology, 25(4), e13454.

Ijaz J, et al. (2024) Haplotype-specific assembly of shattered chromosomes in esophageal adenocarcinomas. Cell genomics, 4(2), 100484.

Low EL, et al. (2024) Chromosome-scale Elaeis guineensis and E. oleifera assemblies: comparative genomics of oil palm and other Arecaceae. G3 (Bethesda, Md.), 14(9).

Helal AA, et al. (2024) Benchmarking long-read aligners and SV callers for structural variation detection in Oxford nanopore sequencing data. Scientific reports, 14(1), 6160.

Wang C, et al. (2024) High-depth whole-genome sequencing identifies structure variants, copy number variants and short tandem repeats associated with Parkinson's disease. NPJ Parkinson's disease, 10(1), 134.

Romagnoli S, et al. (2023) Resolving complex structural variants via nanopore sequencing. Frontiers in genetics, 14, 1213917.

Higuera A, et al. (2023) Draft genomes of Blastocystis subtypes from human samples of Colombia. Parasites & vectors, 16(1), 52.

Sedeek K, et al. (2023) Multi-omics resources for targeted agronomic improvement of pigmented rice. Nature food, 4(5), 366.

Feng L, et al. (2022) The highly continuous reference genome of a leaf-chimeric red pineapple (Ananas comosus var. bracteatus f. tricolor) provides insights into elaboration of leaf color. G3 (Bethesda, Md.), 12(2).

van der Lee M, et al. (2022) Application of long-read sequencing to elucidate complex pharmacogenomic regions: a proof of principle. The pharmacogenomics journal, 22(1), 75.

Du Y, et al. (2022) Dynamic Interplay between Structural Variations and 3D Genome Organization in Pancreatic Cancer. Advanced science (Weinheim, Baden-Wurttemberg, Germany), 9(18), e2200818.

Xiao C, et al. (2022) Personalized genome assembly for accurate cancer somatic mutation discovery using tumor-normal paired reference samples. Genome biology, 23(1), 237.

Walker K, et al. (2022) The third international hackathon for applying insights into large-scale genomic composition to use cases in a wide range of organisms. F1000Research, 11, 530.

Lou H, et al. (2022) Haplotype-resolved de novo assembly of a Tujia genome suggests the necessity for high-quality population-specific genome references. Cell systems, 13(4), 321.

Liu H, et al. (2022) Assessment of two-pool multiplex long-amplicon nanopore sequencing of SARS-CoV-2. Journal of medical virology, 94(1), 327.

Alser M, et al. (2021) Technology dictates algorithms: recent developments in read alignment. Genome biology, 22(1), 249.

Hu T, et al. (2021) Bioinformatics resources for SARS-CoV-2 discovery and surveillance. Briefings in bioinformatics, 22(2), 631.

Chawla HS, et al. (2021) Long-read sequencing reveals widespread intragenic structural variants in a recent allopolyploid crop plant. Plant biotechnology journal, 19(2), 240.

Minoche AE, et al. (2021) ClinSV: clinical grade structural and copy number variant detection from whole genome sequencing data. Genome medicine, 13(1), 32.

Peñaloza C, et al. (2021) A chromosome-level genome assembly for the Pacific oyster Crassostrea gigas. GigaScience, 10(3).