

# Resource Summary Report

Generated by [Kravitz](#) on Sep 10, 2026

## SimSeq

RRID:SCR\_006947

Type: Tool

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### Proper Citation

SimSeq (RRID:SCR\_006947)

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### Resource Information

**URL:** <https://github.com/jstjohn/SimSeq>

**Proper Citation:** SimSeq (RRID:SCR\_006947)

**Description:** An illumina paired-end and mate-pair short read simulator. This project attempts to model as many of the quirks that exist in Illumina data as possible. Some of these quirks include the potential for chimeric reads, and non-biotinylated fragment pull down in mate-pair libraries .

**Abbreviations:** SimSeq

**Resource Type:** simulation software, software application, software resource

**Keywords:** bio.tools

**Availability:** Free

**Resource Name:** SimSeq

**Resource ID:** SCR\_006947

**Alternate IDs:** biotools:simseq, OMICS\_00258

**Alternate URLs:** <https://bio.tools/simseq>

**Record Creation Time:** 20220129T080239+0000

**Record Last Update:** 20260905T132604+0000

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### Ratings and Alerts

No rating or validation information has been found for SimSeq.

No alerts have been found for SimSeq.

## Data and Source Information

**Source:** [SciCrunch Registry](#)

## Usage and Citation Metrics

We found 30 mentions in open access literature.

**Listed below are recent publications.** The full list is available at [Kravitz](#).

Buratin A, et al. (2023) Systematic benchmarking of statistical methods to assess differential expression of circular RNAs. *Briefings in bioinformatics*, 24(1).

Tonkin-Hill G, et al. (2023) Robust analysis of prokaryotic pangenome gene gain and loss rates with Panstripe. *Genome research*, 33(1), 129.

Fleming JF, et al. (2023) nRCFV: a new, dataset-size-independent metric to quantify compositional heterogeneity in nucleotide and amino acid datasets. *BMC bioinformatics*, 24(1), 145.

Carruthers T, et al. (2022) The Implications of Incongruence between Gene Tree and Species Tree Topologies for Divergence Time Estimation. *Systematic biology*, 71(5), 1124.

Venturini C, et al. (2022) Haplotype assignment of longitudinal viral deep sequencing data using covariation of variant frequencies. *Virus evolution*, 8(2), veac093.

Sekizuka T, et al. (2022) Comprehensive Genome and Plasmidome Analysis of Antimicrobial Resistant Bacteria in Wastewater Treatment Plant Effluent of Tokyo. *Antibiotics (Basel, Switzerland)*, 11(10).

Knyazev S, et al. (2021) Accurate assembly of minority viral haplotypes from next-generation sequencing through efficient noise reduction. *Nucleic acids research*, 49(17), e102.

Sekizuka T, et al. (2020) A Genome Epidemiological Study of SARS-CoV-2 Introduction into Japan. *mSphere*, 5(6).

Lammers F, et al. (2019) Retrophylogenomics in rorquals indicate large ancestral population sizes and a rapid radiation. *Mobile DNA*, 10, 5.

Lewitus E, et al. (2019) A non-parametric analytic framework for within-host viral phylogenies and a test for HIV-1 founder multiplicity. *Virus evolution*, 5(2), vez044.

Assefa AT, et al. (2018) Differential gene expression analysis tools exhibit substandard performance for long non-coding RNA-sequencing data. *Genome biology*, 19(1), 96.

Eisfeldt J, et al. (2018) AMYCNE: Confident copy number assessment using whole genome sequencing data. *PloS one*, 13(3), e0189710.

Nakada-Tsukui K, et al. (2018) AIG1 affects in vitro and in vivo virulence in clinical isolates of *Entamoeba histolytica*. PLoS pathogens, 14(3), e1006882.

Baaijens JA, et al. (2017) De novo assembly of viral quasispecies using overlap graphs. Genome research, 27(5), 835.

Klopfstein S, et al. (2017) More on the Best Evolutionary Rate for Phylogenetic Analysis. Systematic biology, 66(5), 769.

Imamura D, et al. (2017) Comparative genome analysis of VSP-II and SNPs reveals heterogenic variation in contemporary strains of *Vibrio cholerae* O1 isolated from cholera patients in Kolkata, India. PLoS neglected tropical diseases, 11(2), e0005386.

Sikdar S, et al. (2017) EAMA: Empirically adjusted meta-analysis for large-scale simultaneous hypothesis testing in genomic experiments. PloS one, 12(10), e0187287.

Zoer M, et al. (2017) Variant profiling of evolving prokaryotic populations. PeerJ, 5, e2997.

Tripathi R, et al. (2017) Unraveling long non-coding RNAs through analysis of high-throughput RNA-sequencing data. Non-coding RNA research, 2(2), 111.

Briskine RV, et al. (2017) Positional bias in variant calls against draft reference assemblies. BMC genomics, 18(1), 263.