

# Resource Summary Report

Generated by SPARC SAWG on Sep 18, 2026

## Eddy Lab Software

RRID:SCR\_001458

Type: Tool

### Proper Citation

Eddy Lab Software (RRID:SCR\_001458)

### Resource Information

**URL:** <http://eddylab.org/software.html>

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**Description:** Software library containing tools for statistical manipulations of data. Tools include profile hidden Markov models for biological sequence analysis, RNA structure analysis, and a prototype noncoding RNA genefinder.

**Synonyms:** Eddy Lab: Software, Eddy Lab - Software

**Resource Type:** software resource

**Keywords:** software repository, statistics, data, statistical analysis, statistical manipulation, markov model, bio.tools

**Availability:** Free, Available for download, Freely available

**Resource Name:** Eddy Lab Software

**Resource ID:** SCR\_001458

**Alternate IDs:** biotools:pknots, nif-0000-08778

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

**Old URLs:** <http://selab.janelia.org/software.html>

**Record Creation Time:** 20220129T080207+0000

**Record Last Update:** 20260912T195526+0000

### Ratings and Alerts

No rating or validation information has been found for Eddy Lab Software.

No alerts have been found for Eddy Lab Software.

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## Data and Source Information

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

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## Usage and Citation Metrics

We found 23 mentions in open access literature.

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

Yamagami R, et al. (2025) tRNA pseudouridine synthase D (TruD) from *Thermus thermophilus* modifies U13 in tRNA<sup>Asp</sup>, tRNA<sup>Glu</sup>, and tRNA<sup>Gln</sup> and U35 in tRNA<sup>Tyr</sup>. RNA (New York, N.Y.), 31(6), 850.

Liu G, et al. (2024) Identification of Hammerhead-variant ribozyme sequences in SARS-CoV-2. Nucleic acids research, 52(6), 3262.

Shen L, et al. (2024) Genomic basis of environmental adaptation in the widespread poly-extremophilic *Exiguobacterium* group. The ISME journal, 18(1).

Zhang Z, et al. (2024) Minimal twister sister-like self-cleaving ribozymes in the human genome revealed by deep mutational scanning. eLife, 12.

Déjosez M, et al. (2023) Bat pluripotent stem cells reveal unusual entanglement between host and viruses. Cell, 186(5), 957.

Shimwell C, et al. (2023) A first molecular characterization of the scorpion telson microbiota of *Hadrurus arizonensis* and *Smeringurus mesaensis*. PloS one, 18(1), e0277303.

Weinberg CE, et al. (2019) Novel ribozymes: discovery, catalytic mechanisms, and the quest to understand biological function. Nucleic acids research, 47(18), 9480.

Boyd JA, et al. (2018) GraftM: a tool for scalable, phylogenetically informed classification of genes within metagenomes. Nucleic acids research, 46(10), e59.

Královicová J, et al. (2018) PUF60-activated exons uncover altered 3' splice-site selection by germline missense mutations in a single RRM. Nucleic acids research, 46(12), 6166.

Dudek NK, et al. (2017) Novel Microbial Diversity and Functional Potential in the Marine Mammal Oral Microbiome. Current biology : CB, 27(24), 3752.

Wang Y, et al. (2015) Genome-wide comparative analysis reveals similar types of NBS genes in hybrid *Citrus sinensis* genome and original *Citrus clementine* genome and provides new insights into non-TIR NBS genes. PloS one, 10(3), e0121893.

Rees MA, et al. (2015) Changes in protein abundance are observed in bacterial isolates from a natural host. *Frontiers in cellular and infection microbiology*, 5, 71.

Sangwan N, et al. (2014) Reconstructing an ancestral genotype of two hexachlorocyclohexane-degrading *Sphingobium* species using metagenomic sequence data. *The ISME journal*, 8(2), 398.

Gillett CP, et al. (2014) Bulk de novo mitogenome assembly from pooled total DNA elucidates the phylogeny of weevils (Coleoptera: Curculionoidea). *Molecular biology and evolution*, 31(8), 2223.

Tang YT, et al. (2014) Genome of the human hookworm *Necator americanus*. *Nature genetics*, 46(3), 261.

Sahl JW, et al. (2013) Evolution of a pathogen: a comparative genomics analysis identifies a genetic pathway to pathogenesis in *Acinetobacter*. *PloS one*, 8(1), e54287.

Sridhar J, et al. (2013) Computational small RNA prediction in bacteria. *Bioinformatics and biology insights*, 7, 83.

Mariotti M, et al. (2013) SECISearch3 and Sebastian: new tools for prediction of SECIS elements and selenoproteins. *Nucleic acids research*, 41(15), e149.

Mitreva M, et al. (2011) The draft genome of the parasitic nematode *Trichinella spiralis*. *Nature genetics*, 43(3), 228.

Sievers F, et al. (2011) Fast, scalable generation of high-quality protein multiple sequence alignments using Clustal Omega. *Molecular systems biology*, 7, 539.