## **Resource Summary Report**

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# **Centrifuge Classifier**

RRID:SCR\_016665 Type: Tool

#### **Proper Citation**

Centrifuge Classifier (RRID:SCR\_016665)

#### **Resource Information**

URL: http://www.ccb.jhu.edu/software/centrifuge/

Proper Citation: Centrifuge Classifier (RRID:SCR\_016665)

**Description:** Software for rapid and sensitive classification of metagenomic sequences. Used for the classification of DNA sequences from microbial samples and analysis of large metagenomics data sets on conventional desktop computers.

**Resource Type:** data analysis software, sequence analysis software, software resource, data processing software, software application

Defining Citation: DOI:10.1101/gr.210641.116

**Keywords:** classification, large, metagenomic, sequence, DNA, microbial, sample, analysis, data, desktop, computer, bio.tools

Funding: U. S. Army Research Office W911NF1410490; NSF ABI1356078; NHGRI R01 HG006677; NIGMS R01 GM083873

Availability: Free, Available for download, Freely available

Resource Name: Centrifuge Classifier

Resource ID: SCR\_016665

Alternate IDs: biotools:centrifuge, OMICS\_12217

Alternate URLs: https://github.com/infphilo/centrifuge, https://bio.tools/centrifuge,

https://sources.debian.org/src/centrifuge/

License: GNU General Public License v3.0

**Record Creation Time:** 20220129T080331+0000

Record Last Update: 20250430T060101+0000

## **Ratings and Alerts**

No rating or validation information has been found for Centrifuge Classifier.

No alerts have been found for Centrifuge Classifier.

### Data and Source Information

Source: <u>SciCrunch Registry</u>

## **Usage and Citation Metrics**

We found 7 mentions in open access literature.

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

Leiendecker L, et al. (2023) Human Papillomavirus 42 Drives Digital Papillary Adenocarcinoma and Elicits a Germ Cell-like Program Conserved in HPV-Positive Cancers. Cancer discovery, 13(1), 70.

Lötter A, et al. (2022) Haplogenome assembly reveals structural variation in Eucalyptus interspecific hybrids. GigaScience, 12.

Pham GM, et al. (2020) Construction of a chromosome-scale long-read reference genome assembly for potato. GigaScience, 9(9).

Nikolaisen NK, et al. (2020) Employing MIC Data for Mink Pathogens to Propose Tentative Epidemiological Cut-Off Values: A Step Toward Rationalizing Antimicrobial Use in Mink. Frontiers in veterinary science, 7, 544594.

Rousseau-Gueutin M, et al. (2020) Long-read assembly of the Brassica napus reference genome Darmor-bzh. GigaScience, 9(12).

Watts GS, et al. (2019) Identification and quantitation of clinically relevant microbes in patient samples: Comparison of three k-mer based classifiers for speed, accuracy, and sensitivity. PLoS computational biology, 15(11), e1006863.

Nelson MT, et al. (2019) Human and Extracellular DNA Depletion for Metagenomic Analysis of Complex Clinical Infection Samples Yields Optimized Viable Microbiome Profiles. Cell reports, 26(8), 2227.