## **Resource Summary Report**

Generated by FDI Lab - SciCrunch.org on May 15, 2025

# **StemID**

RRID:SCR\_017242

Type: Tool

### **Proper Citation**

StemID (RRID:SCR\_017242)

#### Resource Information

URL: https://github.com/dgrun/StemID

**Proper Citation:** StemID (RRID:SCR\_017242)

**Description:** Algorithm for derivation of cell lineage trees based on RaceID2 results and predicts multipotent cell identites. StemID2 is algorithm for identification of lineage trees based on RaceID3 analysis. Used for better understanding of differentiation dynamics in variety of systems. Written in R computing language.

Synonyms: StemID2

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

software resource, algorithm resource

**Defining Citation: PMID:27345837** 

Keywords: derivation, cell, linage, tree, predict, multipotent, identity, dynamic

Funding: European Research Council;

Nederlandse Organisatie voor Wetenschappelijk Onderzoek;

DON Foundation;

**Dutch Diabetes Research Foundation** 

Availability: Free, Available for download, Freely available

Resource Name: StemID

Resource ID: SCR 017242

Alternate URLs: https://github.com/dgrun/RaceID3 StemID2,

https://github.com/dgrun/RaceID3\_StemID2\_package

**Record Creation Time:** 20220129T080334+0000

**Record Last Update:** 20250514T061806+0000

## **Ratings and Alerts**

No rating or validation information has been found for StemID.

No alerts have been found for StemID.

### **Data and Source Information**

Source: SciCrunch Registry

## **Usage and Citation Metrics**

We found 10 mentions in open access literature.

Listed below are recent publications. The full list is available at FDI Lab - SciCrunch.org.

Henon C, et al. (2024) Single-cell multiomics profiling reveals heterogeneous transcriptional programs and microenvironment in DSRCTs. Cell reports. Medicine, 5(6), 101582.

Wang X, et al. (2024) MarsGT: Multi-omics analysis for rare population inference using single-cell graph transformer. Nature communications, 15(1), 338.

Wang S, et al. (2024) scSID: A lightweight algorithm for identifying rare cell types by capturing differential expression from single-cell sequencing data. Computational and structural biotechnology journal, 23, 589.

Wang X, et al. (2023) MarsGT: Multi-omics analysis for rare population inference using single-cell graph transformer. bioRxiv: the preprint server for biology.

Pérez-Mojica JE, et al. (2023) Single-embryo RNA sequencing for continuous and sexspecific gene expression analysis on Drosophila. STAR protocols, 4(3), 102535.

Zhang F, et al. (2022) FitDevo: accurate inference of single-cell developmental potential using sample-specific gene weight. Briefings in bioinformatics, 23(5).

Zhang L, et al. (2022) Digital Cell Atlas of Mouse Uterus: From Regenerative Stage to Maturational Stage. Frontiers in genetics, 13, 847646.

Molenaar B, et al. (2021) Single-cell transcriptomics following ischemic injury identifies a role for B2M in cardiac repair. Communications biology, 4(1), 146.

Ouadah Y, et al. (2019) Rare Pulmonary Neuroendocrine Cells Are Stem Cells Regulated by Rb, p53, and Notch. Cell, 179(2), 403.

Honkoop H, et al. (2019) Single-cell analysis uncovers that metabolic reprogramming by ErbB2 signaling is essential for cardiomyocyte proliferation in the regenerating heart. eLife, 8.