

Salmon.alevin.quant Documentation

Description: Quantification and analysis of 3' tagged-end single-cell sequencing data using Salmon's Alevin function. See: <u>The Alevin Documentation</u> for detailed usage guidelines.

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Contact: Algorithm and scientific questions: https://github.com/COMBINE-lab/salmon/issues.

Module specific issues: genepattern.org/help

Summary: Quantify gene expression from single-cell sequencing experiments using the Alevin module of the pseudo-alignment based method "Salmon".

Source Publication: Srivastava A, Malik L, Smith T, Sudbery I, Patro R. Alevin efficiently estimates accurate gene abundances from dscRNA-seq data. Genome Biol. 2019 Mar 27;20(1):65. doi: 10.1186/s13059-019-1670-y. PMID: 30917859; PMCID: PMC6437997.

Basic Parameters:

Name	Description
Barcodes	Cellular barcodes+UMI file(s), the FASTQ file containing CB+UMI raw sequences. Alevin also supports parsing of data from multiple files as long as the order is the same as in the "Reads" parameter. Barcodes and reads should share the same base file name in order to ensure order matching, barcodes should have the suffix _R1 (SampleID_R1.fastq.gz). File should be gzipped.
Reads	The FASTQ file containing raw read-sequences. Alevin also supports parsing of data from multiple files as long as the order is the same as in the "Barcodes" parameter. Barcodes and reads should share the same base file name in order to ensure order matching, barcodes should have the suffix _R2 (SampleID_R1.fastq.gz) File should be gzipped.
Transcriptome Index	The indexed transcriptome output from the salmon.index module (or comparable pipeline). File must be .tar.gz
Chemistry Library Type	The chemistry used by the single-cell sequencing platform. Drop-Seq, 10x Chromium v2, 10x Chromium v3, CITE-Seq, CEL-Seq, CEL-Seq2, and Quartz-Seq2 are currently supported. The orientation of the Barcodes and Reads. The
Library Type	Alevin authors recommend using ISR (default) for both Drop-seq and Chromium chemistry.
ТgМар	A two column tsv (tab-separated) file with no header containing the transcript to gene map file. The first column lists each transcript present in the reference, the second column lists the corresponding gene. Alternatively, a .GTF file can be supplied to



	automatically create a transcript to gene map.
Output Basename	The base name to use for naming the alevin results
	file (default: alevin.output)

Advanced Parameters: See: <u>The Alevin Documentation</u> for details on additional parameters offered in this module.

Output Files:

Name	Description
<alevin.output>.tar.gz</alevin.output>	Gzipped files containing the alevin quantification
	results.

Module Language: Shell script

Source Repository: https://github.com/genepattern/salmon.alevin.quant/releases/tag/v1

Docker image: combinelab/salmon:1.4.0

Version	Comment
1	Initial release.