

Salmon.Indexer Documentation

Description: Indexing a transcriptome in order to perform quantification with Salmon and Alevin. See: The Salmon User Guide 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: In order to quantify gene expression using Salmon or Alevin, the transcriptome must first be indexed. This module performs this indexing step.

Source Publication: Patro R, Duggal G, Love MI, Irizarry RA, Kingsford C. Salmon provides fast and bias-aware quantification of transcript expression. Nat Methods. 2017 Apr;14(4):417-419. doi: 10.1038/nmeth.4197. Epub 2017 Mar 6. PMID: 28263959; PMCID: PMC5600148.

Parameters:

Name	Description
GTF gz	A GTF file containing the genomic features to be
	indexed. Must be gzipped.
Transcriptome fa gz	A FASTA file of the transcript sequences
	corresponding to the organism's GTF file.
	Optional: if a Fasta file is not provided the module
	will attempt to use gffread to extract the sequences
	from the Genomic fasta file. Must be gzipped.
Genome fa gz	A FASTA file of the genomic sequences
	corresponding to the organism's genome assembly
	used for the transcript annotation. Must be gzipped.
kmer	An odd numbered integer. Used to index the
	transcriptome, used as the minimum acceptable
	length of a valid match. (Default: 31, for reads
	<70bp, consider adjusting this to a smaller value.)
Index Mode	Options for generating a decoy-aware transcriptome
	index
	Build a full decoy index (full), or a partial decoy index
	(partial). See parameter description for details.
Use Gencode	Gencode uses " " characters in their FASTA record
	IDs, "true" allows salmon.index to be aware of these
	characters when processing gene IDs from records

Output Files:

Name	Description
[index_name]_index.tar.gx	A gzipped file containing the salmon index for downstream quantification. Name will be based on either the GTF or Transcriptome file name depending on indexing mode, and contain the kmer value utilized.
<gtf.basename>.fa.gz</gtf.basename>	If a transcriptome fasta was not provided, this output is generated from the provided GTF file and genome fasta file using GFFread.



Module Language: Shell script

Source Repository: https://github.com/genepattern/Salmon.Indexer/releases/tag/v1
Docker image: combinelab/salmon:1.5.2

Version	Comment
1	Initial release.