

## **PreprocessVelocityTranscriptome Documentation**

**Description:** Extract transcript and intron sequences from the genome sequence using the <u>eisaR</u> package in order to quantify both intronic (unspliced) and exonic (spliced) sequences.

Author: Anthony S. Castanza genepattern.org/help

**Summary**: In order to build a transcriptome index for single-cell RNA velocity quantification, intronic (unprocessed) and exonic (processed) RNA sequences must be extracted from the genome. This module prepares the files necessary to produce velocity-compatible input files for the salmon index module.

## Parameters:

Name	Description
GTF	A GTF file containing the genomic ranges to extract
	features from for quantification.
Genome FASTA	A FASTA file of the genomic sequence
	corresponding to the organism's GTF file.
Intron Extraction	Consider transcripts separately ("separate") when
	extracting intronic regions, or collapsed to gene level
	("collapse").
Insert Size	Length of the insert being sequenced by the single-
	cell platform not including any adapters or barcodes.

## **Output Files:**

Output riles.		
Name	Description	
<gtf.basename>.annotation.velocity.</gtf.basename>	A gzipped GTF file containing the intronic and	
<pre><intron.flank.length>bp_flank.gtf.gz</intron.flank.length></pre>	exonic genomic ranges extracted.	
	Input for the salmon.index module.	
<gtf.basename>.annotation.velocity.</gtf.basename>	A gzipped FASTA file of the genomic sequence	
<pre><intron.flank.length>bp_flank.fa.gz</intron.flank.length></pre>	corresponding to intronic and exonic genomic	
	ranges extracted.	
	Input for the salmon.index module.	
<gtf.basename>.annotation.velocity.</gtf.basename>	A tab delimited file containing the list of spliced	
<pre><intron.flank.length>bp_flank.features.tsv</intron.flank.length></pre>	gene ids in column 1, the unspliced gene ids in	
	column 2, and gene names (symbols) in column 3.	
<gtf.basename>.annotation.velocity.</gtf.basename>	A two-column file containing the mappings of	
<pre><intron.flank.length>bp_flank.tgMap.tsv</intron.flank.length></pre>	transcript level features to gene level features Input	
	for the salmon.alevin.quant module.	
<gtf.basename>.annotation.velocity.</gtf.basename>	A list of the gene ids for mitochondrial genes.	
<pre><intron.flank.length>bp_flank.mtGenes.txt</intron.flank.length></pre>		
<gtf.basename>.annotation.velocity.</gtf.basename>	A list of the gene ids with the biotype "rRNA"	
<pre><intron.flank.length>bp_flank.rrnaGenes.txt</intron.flank.length></pre>	(ribosomal RNA genes).	

Module Language: R 4.0.5

Source Repository: https://github.com/genepattern/PreprocessVelocityTranscriptome/releases/tag/v1

**Docker image:** genepattern/prepvelocitytxome:1.0

Version	Comment
1	Initial release.