# Package 'BUSpaRse'

November 20, 2024

Type Package

Title kallisto | bustools R utilities

**Version** 1.20.0 **Date** 2024-07-31

**Description** The kallisto | bustools pipeline is a fast and modular set of tools to convert single cell RNA-seq reads in fastq files into gene count or transcript compatibility counts (TCC) matrices for downstream analysis. Central to this pipeline is the barcode, UMI, and set (BUS) file format. This package serves the following purposes: First, this package allows users to manipulate BUS format files as data frames in R and then convert them into gene count or TCC matrices. Furthermore, since R and Rcpp code is easier to handle than pure C++ code, users are encouraged to tweak the source code of this package to experiment with new uses of BUS format and different ways to convert the BUS file into gene count matrix. Second, this package can conveniently generate files required to generate gene count matrices for spliced and unspliced transcripts for RNA velocity. Here biotypes can be filtered and scaffolds and haplotypes can be removed, and the filtered transcriptome can be extracted and written to disk. Third, this package implements utility functions to get transcripts and associated genes required to convert BUS files to gene count matrices, to write the transcript to gene information in the format required by bustools, and to read output of bustools into R as sparses matrices.

BugReports https://github.com/BUStools/BUSpaRse/issues

URL https://github.com/BUStools/BUSpaRse

Imports AnnotationDbi, AnnotationFilter, biomaRt, BiocGenerics, Biostrings, BSgenome, dplyr, ensembldb, GenomeInfoDb, GenomicFeatures, GenomicRanges, ggplot2, IRanges, magrittr, Matrix, methods, plyranges, Rcpp, S4Vectors, stats, stringr, tibble, tidyr, utils, zeallot

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2 Contents

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# **Contents**

.get_velocity_files
annots_from_fa_df
annot_circular
cellranger_biotypes
check_char1
check_genome
check_gff
check_tag_present
check_tx
dl_transcriptome
EC2gene
ensembl_gene_biotypes
$ensembl\_gff\_mcols$
ensembl_gtf_mcols
ensembl_tx_biotypes
get_intron_flanks
get_knee_df
get_velocity_files
make_sparse_matrix
match_style
read_count_output
read_velocity_output
refseq_gff_mcols
save_tr2g_bustools
sort_tr2g
species2dataset

```
standardize tags......
52
Index
```

### Description

Generate RNA velocity files for GRanges

### Usage

```
.get_velocity_files(
 gr,
 L,
 Genome,
 Transcriptome = NULL,
 out_path = ".",
 style = c("genome", "Ensembl", "UCSC", "NCBI", "other"),
  isoform_action = c("separate", "collapse"),
 exon_option = c("full", "junction"),
  transcript_id = "transcript_id",
 gene_id = "gene_id",
  transcript_version = "transcript_version",
 gene_version = "gene_version",
 version_sep = ".",
 transcript_biotype_col = "transcript_biotype",
 gene_biotype_col = "gene_biotype",
 transcript_biotype_use = "all",
 gene_biotype_use = "all",
 chrs_only = TRUE
  save_filtered_gtf = FALSE,
 compress_fa = FALSE,
 width = 80L
)
```

### **Arguments**

gr

A GRanges object for gene annotation.

L

Length of the biological read. For instance, 10xv1: 98 nt, 10xv2: 98 nt, 10xv3: 91 nt, Drop-seq: 50 nt. If in doubt check read length in a fastq file for biological reads with the bash commands: If the fastq file is gzipped, then do zcat your\_file.fastq.gz | head on Linux. If on Mac, then zcat < your\_file.fastq.gz | head. Then you will see lines with nucleotide bases. Copy one of those lines and determine its length with str\_length in R or echo -n <the sequence> | wc -c in bash. Which file corresponds to biological reads depends on the particular technology.

Genome

Either a BSgenome or a XStringSet object of genomic sequences, where the intronic sequences will be extracted from. Use genomeStyles to check which styles are supported for your organism of interest; supported styles can be interconverted. If the style in your genome or annotation is not supported, then the style of chromosome names in the genome and annotation should be manually set to be consistent.

Transcriptome

A XStringSet, a path to a fasta file (can be gzipped) of the transcriptome which contains sequences of spliced transcripts, or NULL. The transcriptome here will be concatenated with the intronic sequences to give one fasta file. When NULL, the transriptome sequences will be extracted from the genome given the gene annotation, so it will be guaranteed that transcript IDs in the transcriptome and in the annotation match. Otherwise, the type of transcript ID in the transcriptome must match that in the gene annotation supplied via argument X.

out\_path

Directory to save the outputs written to disk. If this directory does not exist, then it will be created. Defaults to the current working directory.

style

Formatting of chromosome names. Use genomeStyles to check which styles are supported for your organism of interest and what those styles look like. This can also be a style supported for your organism different from the style used by the annotation and the genome. Then this style will be used for both the annotation and the genome. Can take the following values:

genome If style of the annnotation is different from that of the genome, then the style of the genome will be used.

**other** Custom style, need to manually ensure that the style in annotation matches that of the genome.

**Ensembl** Or UCSC or NCBI, whichever is supported by your species of interest.

isoform\_action Character, indicating action to take with different transcripts of the same gene. Must be one of the following:

> **collapse** First, the union of all exons of different transcripts of a gene will be taken. Then the introns will be inferred from this union. Only the flanked intronic sequences are affected; isoforms will always be taken into account for spliced sequences or exon-exon junctions.

**separate** Introns from different transcripts will be kept separate.

exon\_option

Character, indicating how exonic sequences should be included in the kallisto index. Must be one of the following:

full The full cDNA sequences, which include the full exonic sequences, will be used. This is the default.

**junction** Only the exon-exon junctions, with L-1 bases on each side of the junctions, will be used.

transcript\_id

Character vector of length 1. Tag in attribute field corresponding to transcript IDs. This argument must be supplied and cannot be NA or NULL. Will throw error if tag indicated in this argument does not exist.

gene\_id

Character vector of length 1. Tag in attribute field corresponding to gene IDs. This argument must be supplied and cannot be NA or NULL. Note that this is different from gene symbols, which do not have to be unique. This can be Ensembl or Entrez IDs. However, if the gene symbols are in fact unique for each gene, you may supply the tag for human readable gene symbols to this argument. Will throw error if tag indicated in this argument does not exist. This is typically "gene\_id" for annotations from Ensembl and "gene" for refseq.

### transcript\_version

Character vector of length 1. Tag in attribute field corresponding to *transcript* version number. If your GTF file does not include transcript version numbers, or if you do not wish to include the version number, then use NULL for this argument. To decide whether to include transcript version number, check whether version numbers are included in the transcripts.txt in the kallisto output directory. If that file includes version numbers, then transcript version numbers must be included here as well. If that file does not include version numbers, then transcript version numbers must not be included here.

gene\_version

Character vector of length 1. Tag in attribute field corresponding to *gene* version number. If your GTF file does not include gene version numbers, or if you do not wish to include the version number, then use NULL for this argument. Unlike transcript version number, it's up to you whether to include gene version number.

version\_sep

Character to separate bewteen the main ID and the version number. Defaults to ".", as in Ensembl.

# transcript\_biotype\_col

Character vector of length 1. Tag in attribute field corresponding to *transcript* biotype.

#### gene\_biotype\_col

Character vector of length 1. Tag in attribute field corresponding to *gene* biotype.

#### transcript\_biotype\_use

Character, can be "all" or a vector of *transcript* biotypes to be used. Transcript biotypes aren't entirely the same as gene biotypes. For instance, in Ensembl annotation, retained\_intron is a transcript biotype, but not a gene biotype. If "cellranger", then a warning will be given. See data("ensembl\_tx\_biotypes") for all available transcript biotypes from Ensembl.

### gene\_biotype\_use

Character, can be "all", "cellranger", or a vector of *gene* biotypes to be used. If "cellranger", then the biotypes used by Cell Ranger's reference are used. See data("cellranger\_biotypes") for gene biotypes the Cell Ranger reference uses. See data("ensembl\_gene\_biotypes") for all available gene biotypes from Ensembl. Note that gene biotypes and transcript biotypes are not always the same.

chrs\_only

Logical, whether to include chromosomes only, for GTF and GFF files can contain annotations for scaffolds, which are not incorporated into chromosomes. This will also exclude haplotypes. Defaults to TRUE. Only applicable to species found in genomeStyles().

6 annots\_from\_fa\_df

save\_filtered\_gtf

Logical. If filtering type, biotypes, and/or chromosomes, whether to save the

filtered GRanges as a GTF file.

compress\_fa Logical, whether to compress the output fasta file. If TRUE, then the fasta file

will be gzipped.

width Maximum number of letters per line of sequence in the output fasta file. Must

be an integer.

#### Value

```
See get_velocity_files
```

annots\_from\_fa\_df

Get genome annotation from Ensembl FASTA file

# **Description**

Ensembl FASTA files for RNA contain much of the information contained in GTF files, such as chromosome, genome assembly version, coordinates, strand, gene ID, gene symbol, and gene description. Given such a FASTA file, this function can extract all the genome annotation information and return a data frame or a GRanges object.

### Usage

```
annots_from_fa_df(file)
annots_from_fa_GRanges(file)
```

# Arguments

file

Path to the FASTA file to be read. The file can remain gzipped.

# Value

```
annots_from_fa_df returns a data frame and annots_from_fa_GRanges returns GRanges. A data frame with genome annotations.
```

#### **Examples**

```
fn <- system.file("testdata/fasta_test.fasta", package = "BUSpaRse")\\ annots\_from\_fa\_df(fn)\\ annots\_from\_fa\_GRanges(fn)
```

annot\_circular 7

ć	annot_circular	Transfer information about circular chromosomes between genome and annotation

### **Description**

Internal use, called after calling subset\_annot.

### Usage

```
annot_circular(Genome, annot)
```

### **Arguments**

Genome Either a BSgenome or a XStringSet object of genomic sequences, where the

intronic sequences will be extracted from. Use genomeStyles to check which styles are supported for your organism of interest; supported styles can be interconverted. If the style in your genome or annotation is not supported, then the style of chromosome names in the genome and annotation should be manually

set to be consistent.

annot Genome annotation, an object of a class with a seqlevels method, such as

GRanges, TxDb, and EnsDb.

#### Value

If neither genome nor annotation indicates which chromosome is circular, then the input will be returned unchanged. If only one of genome and annotation has such information, then it will be transferred to the one that does not. If both do have such information, the information from the genome will be transferred to the annotation if they're different.

cellranger\_biotypes Cell Ranger gene biotypes

# Description

In the GRCh38 Cell Ranger reference package, an Ensembl human GTF file is filtered by gene biotypes. This vector includes all gene biotypes included by this Cell Ranger reference package.

#### Usage

```
cellranger_biotypes
```

# **Format**

A character vector with all Cell Ranger reference package gene biotypes.

### **Source**

https://support.10xgenomics.com/single-cell-gene-expression/software/pipelines/latest/advanced/references

8 check\_genome

#### See Also

ensembl\_gene\_biotypes ensembl\_tx\_biotypes

check\_char1

Check that an object is a character vector of length 1

# Description

Just in case the user passes something with length more than 1 and messes up everything thanks to vectorization.

### Usage

```
check_char1(x)
```

### **Arguments**

Х

Named vector of arguments to be checked.

### Value

Error if x is not a character vector with length 1.

check\_genome

Check for chromosomes in genome but not annotation

### **Description**

Check for chromosomes in genome but not annotation

### Usage

```
check_genome(chrs_use, Genome)
```

# Arguments

chrs\_use

Character vector of names of chromosomes present in both the annotation and

the genome.

Genome

Either a BSgenome or a XStringSet object of genomic sequences, where the intronic sequences will be extracted from. Use genomeStyles to check which styles are supported for your organism of interest; supported styles can be interconverted. If the style in your genome or annotation is not supported, then the style of chromosome names in the genome and annotation should be manually

set to be consistent.

### Value

Nothing. Will emit message if the genome contains chromosomes absent from the annotation.

check\_gff 9

check_gff	Check inputs to tr2g_gtf and tr2g_gtf3
_0	1 0-50 0-50

### **Description**

This function validates inputs to tr2g\_gtf and tr2g\_gtf3 and throws error early if some inputs are wrong.

### Usage

```
check_gff(format, file, transcript_id, gene_id)
```

### **Arguments**

format	Whether it's gtf or gff3.
file	Path to a GTF file to be read. The file can remain gzipped. Use getGTF from the biomartr package to download GTF files from Ensembl, and use getGFF from biomartr to download GFF3 files from Ensembl and RefSeq.
transcript_id	Character vector of length 1. Tag in attribute field corresponding to transcript IDs. This argument must be supplied and cannot be NA or NULL. Will throw error

if tag indicated in this argument does not exist.

gene\_id

Character vector of length 1. Tag in attribute field corresponding to gene IDs. This argument must be supplied and cannot be NA or NULL. Note that this is different from gene symbols, which do not have to be unique. This can be

Ensembl or Entrez IDs. However, if the gene symbols are in fact unique for each gene, you may supply the tag for human readable gene symbols to this argument. Will throw error if tag indicated in this argument does not exist. This is typically "gene\_id" for annotations from Ensembl and "gene" for refseq.

#### Value

Nothing, will throw error if there's a problem.

check_tag_present Check that a tag is present in attribute field of GTF/GFF
---

# **Description**

The attribute field of GTF/GFF files are very complicated and is very inconsistent between sources. This function is to make sure that transcript and gene IDs can be extracted properly.

# Usage

```
check_tag_present(tags_use, tags, error = TRUE)
```

10 dl\_transcriptome

### **Arguments**

tags\_use The tags to be checked.

tags The tags present in attribute field.

error Whether to throw an error for absent tags. If FALSE, then a warning will be

given.

### Value

Error or warning if tag is absent.

check\_tx

Check if transcript ID in transcriptome and annotation match

# Description

This function throws an error if transcript IDs in transcriptome and annotation do not overlap. If they do overlap, this function will give a message about transcript IDs that do not agree in the transcriptome and the annotation

### Usage

```
check_tx(tx_annot, tx)
```

### **Arguments**

tx\_annot Character vector of transcript IDs from the annotation.

tx Character vector of transcript IDs from the transcriptome.

### Value

Character vector of the overlapping transcript IDs.

# Description

This function downloads the cDNA fasta file from specific version of Ensembl. It can also filter the fasta file by gene and transcript biotype and remove scaffolds and haplotypes.

dl\_transcriptome 11

#### Usage

```
dl_transcriptome(
   species,
   out_path = ".",
   type = c("vertebrate", "metazoa", "plant", "fungus", "protist"),
   transcript_biotype_use = "all",
   gene_biotype_use = "all",
   chrs_only = TRUE,
   ensembl_version = NULL,
   verbose = TRUE,
   ...
)
```

#### **Arguments**

species

Character vector of length 1, Latin name of the species of interest.

out\_path

Directory to save the outputs written to disk. If this directory does not exist, then it will be created. Defaults to the current working directory.

type

Character, must be one of "vertebrate", "metazoa", "plant", "fungus" and "protist". Passing "vertebrate" will use the default www.ensembl.org host. Gene annotation of some common invertebrate model organisms, such as *Drosophila melanogaster*, are available on www.ensembl.org so for these invertebrate model organisms, "vertebrate" can be used for this argument. Passing values other than "vertebrate" will use other Ensembl hosts. For animals absent from www.ensembl.org, try "metazoa".

transcript\_biotype\_use

Character, can be "all" or a vector of *transcript* biotypes to be used. Transcript biotypes aren't entirely the same as gene biotypes. For instance, in Ensembl annotation, retained\_intron is a transcript biotype, but not a gene biotype. If "cellranger", then a warning will be given. See data("ensembl\_tx\_biotypes") for all available transcript biotypes from Ensembl.

gene\_biotype\_use

Character, can be "all", "cellranger", or a vector of *gene* biotypes to be used. If "cellranger", then the biotypes used by Cell Ranger's reference are used. See data("cellranger\_biotypes") for gene biotypes the Cell Ranger reference uses. See data("ensembl\_gene\_biotypes") for all available gene biotypes from Ensembl. Note that gene biotypes and transcript biotypes are not always the same

chrs\_only

Logical, whether to include chromosomes only, for GTF and GFF files can contain annotations for scaffolds, which are not incorporated into chromosomes. This will also exclude haplotypes. Defaults to TRUE. Only applicable to species found in genomeStyles().

ensembl\_version

Integer version number of Ensembl (e.g. 94 for the October 2018 release). This argument defaults to NULL, which will use the current release of Ensembl. Use listEnsemblArchives to see the version number corresponding to the Ensembl release of a particular date. The version specified here must match the version of Ensembl where the transcriptome used to build the kallisto index was downloaded. This only works for vertebrates and the most common invertebrate model organisms like *Drosophila melanogaster* and *C. elegans* (i.e.

12 EC2gene

www.ensembl.org and its mirrors), not the other Ensembl sites for plants, protists, fungi, and metazoa.

verbose Whether to display progress.

... Other arguments passed to tr2g\_fasta.

#### Value

Invisibly returns the path to the fasta file. The following files are written to disk, in the out\_path directory:

species.genome.cdna.all.fa.gz The cDNA fasta file from Ensembl, from the specified version.

**cdna\_filtered.fa** The filtered cDNA fasta file, only keeping the desired biotypes and without scaffolds and haplotypes (if chrs\_only = TRUE). This file will not be written if all gene and transcript biotypes are used and scaffolds and haplotypes are not removed.

tr2g.tsv The transcript to gene file, without headers so can be directly used for bustools.

### **Examples**

EC2gene

Map EC Index to Genes Compatible with the EC

# Description

In the output file output.bus, equivalence classes (EC) are denoted by an index, which is related to the set of transcripts the EC is compatible to in the output file matrix.ec. This function further relates the set of transcripts to the set of genes the EC is compatible to. This function first reads in matrix.ec, and then translates the transcripts into genes.

### Usage

```
EC2gene(tr2g, kallisto_out_path, verbose = TRUE)
```

# Arguments

tr2g A Data frame with columns gene and transcript, in the same order as in the transcriptome index for kallisto.

kallisto\_out\_path

Path to the kallisto bus output directory.

verbose Logical, whether to display progress.

#### **Details**

The data frame passed to tr2g can be generated from function transcript2gene in this package for any organism that has gene and transcript ID on Ensembl, or from the tr2g\_\* family of function. You no longer need to use this function before running make\_sparse\_matrix; the purpose of this function is to query which genes equivalence classes map to.

Calling this function is unnessary when working with gene count matrices. However, this function is useful for finding genes the ECs map to in TCC matrices, such as when finding species-specific ECs in mixed species datasets and identifying ECs mapped to known marker genes of cell types.

#### Value

A data frame with 3 columns:

**EC\_ind** Index of the EC as appearing in the matrix.ec file.

EC A list column each element of which is a numeric vector of the transcripts in the EC corresponding to the EC index. To learn more about list columns, see the relevant section in the R for Data Science book.

gene A list column each element of which is a character vector of genes the EC maps to.

#### See Also

```
transcript2gene
```

# **Examples**

```
# Load toy example for testing
toy_path <- system.file("testdata", package = "BUSpaRse")
load(paste(toy_path, "toy_example.RData", sep = "/"))
EC2gene(tr2g_toy, toy_path, verbose = FALSE)</pre>
```

ensembl\_gene\_biotypes Gene biotypes from Ensembl

### **Description**

These vectors are here to make it easy to look up which biotypes are available for Ensembl without having to parse GTF and fasta files every time. See this page from Ensembl for what the biotypes mean.

# Usage

```
ensembl_gene_biotypes
```

### Format

A character vector with all Ensembl gene biotypes.

### Source

This vector is all the unique gene biotypes in the Ensembl version 99 human GTF file.

14 ensembl\_gtf\_mcols

### See Also

ensembl\_tx\_biotypes cellranger\_biotypes

ensembl\_gff\_mcols

These are the column names of the mcols when the Ensembl GTF file is read into R as a GRanges, including gene\_id, transcript\_id, biotype, description, and so on, and the mandatory tags like ID, Name, and Parent.

### **Description**

These are the column names of the mcols when the Ensembl GTF file is read into R as a GRanges, including gene\_id, transcript\_id, biotype, description, and so on, and the mandatory tags like ID, Name, and Parent.

# Usage

ensembl\_gff\_mcols

#### **Format**

A character vector

### **Source**

Ensembl version 99 human GFF3 file

ensembl\_gtf\_mcols

Tags in the attributes field of Ensembl GTF files

### **Description**

These are the column names of the mcols when the Ensembl GTF file is read into R as a GRanges, including gene\_id, transcript\_id, gene\_biotype, transcript\_biotype, description, and so on

# Usage

```
ensembl_gtf_mcols
```

# Format

A character vector

# Source

Ensembl version 99 human GTF file

ensembl\_tx\_biotypes 15

ensembl\_tx\_biotypes Tra

Transcript biotypes from Ensembl

### **Description**

These vectors are here to make it easy to look up which biotypes are available for Ensembl without having to parse GTF and fasta files every time. See this page from Ensembl for what the biotypes mean.

### Usage

```
ensembl_tx_biotypes
```

### **Format**

A character vector with all Ensembl transcript biotypes.

#### **Source**

This vector is all the unique transcript biotypes in the Ensembl version 99 human GTF file.

#### See Also

ensembl\_gene\_biotypes cellranger\_biotypes

get\_intron\_flanks

Get flanked intronic ranges

### **Description**

Get flanked intronic ranges

### Usage

```
get_intron_flanks(grl, L, get_junctions)
```

### **Arguments**

grl A CompressedGRangesList for exonic ranges, each element for one transcript.

L Read length.

get\_junctions Logical, whether to also return exon-exon junctions.

#### Value

If get\_junctions is FALSE, then a GRanges object with ranges for flanked intronic regions. If get\_junctions is TRUE, then in addition to the flanked intronic ranges, a CompressedGRangesList with exon-exon junction ranges and ranges for transcripts without introns.

16 get\_knee\_df

get	knee	dt

Plot the transposed knee plot and inflection point

### **Description**

Plot a transposed knee plot, showing the inflection point and the number of remaining cells after inflection point filtering. It's transposed since it's more generalizable to multi-modal data.

### Usage

```
get_knee_df(mat)
get_inflection(df, lower = 100)
knee_plot(df, inflection)
```

### **Arguments**

mat Gene count matrix, a dgCMatrix.

df The data frame from get\_knee\_df.

lower Minimum total UMI counts for barcode for it to be considered when calculating

the inflection point; this helps to avoid the noisy part of the curve for barcodes

with very few counts.

inflection Output of get\_inflection.

### Value

get\_knee\_df returns a tibble with two columns: total for total UMI counts for each barcode, and rank for rank of the total counts, with number 1 for the barcode with the most counts.

get\_inflection returns a numeric(1) for the total UMI count at the inflection point.

knee\_plot returns a ggplot2 object.

#### Note

Code in part adapted from barcodeRanks from DropetUtils.

# **Examples**

```
# Download dataset already in BUS format
library(TENxBUSData)
TENxBUSData(".", dataset = "hgmm100")
tr2g <- transcript2gene(c("Homo sapiens", "Mus musculus"),
    type = "vertebrate",
    ensembl_version = 99, kallisto_out_path = "./out_hgmm100")
m <- make_sparse_matrix("./out_hgmm100/output.sorted.txt",
    tr2g = tr2g, est_ncells = 1e5,
    est_ngenes = nrow(tr2g), TCC = FALSE)
df <- get_knee_df(m)
infl <- get_inflection(df)
knee_plot(df, infl)
# Clean up files from the example
unlink("out_hgmm100")</pre>
```

get\_velocity\_files

Get files required for RNA velocity with bustools

### **Description**

Computation of RNA velocity requires the number of unspliced transcripts, which can be quantified with reads containing intronic sequences. This function extracts intronic sequences flanked by L-1 bases of exonic sequences where L is the biological read length of the single cell technology of interest. The flanking exonic sequences are included for reads partially mapping to an intron and an exon.

### Usage

```
get_velocity_files(
  Χ,
  L,
  Genome,
  Transcriptome = NULL,
  out_path = ".",
  style = c("genome", "Ensembl", "UCSC", "NCBI", "other"),
  isoform_action = c("separate", "collapse"),
  exon_option = c("full", "junction"),
  compress_fa = FALSE,
  width = 80L,
)
## S4 method for signature 'GRanges'
get_velocity_files(
  Χ,
  L,
  Genome,
  Transcriptome = NULL,
  out_path = ".",
  style = c("genome", "Ensembl", "UCSC", "NCBI", "other"),
  isoform_action = c("separate", "collapse"),
  exon_option = c("full", "junction"),
  compress_fa = FALSE,
  width = 80L,
  transcript_id = "transcript_id",
  gene_id = "gene_id",
  transcript_version = "transcript_version",
  gene_version = "gene_version",
  version_sep = ".",
  transcript_biotype_col = "transcript_biotype",
  gene_biotype_col = "gene_biotype",
  transcript_biotype_use = "all",
  gene_biotype_use = "all",
  chrs_only = TRUE,
  save_filtered_gtf = FALSE
```

```
## S4 method for signature 'character'
get_velocity_files(
  Χ,
  L,
  Genome,
  Transcriptome = NULL,
  out_path = ".",
  style = c("genome", "Ensembl", "UCSC", "NCBI", "other"),
isoform_action = c("separate", "collapse"),
  exon_option = c("full", "junction"),
  compress_fa = FALSE,
  width = 80L,
  is_circular = NULL,
  transcript_id = "transcript_id",
  gene_id = "gene_id",
  transcript_version = "transcript_version",
  gene_version = "gene_version",
  version_sep = ".",
  transcript_biotype_col = "transcript_biotype",
  gene_biotype_col = "gene_biotype",
  transcript_biotype_use = "all",
  gene_biotype_use = "all",
  chrs_only = TRUE,
  save_filtered_gtf = FALSE
)
## S4 method for signature 'TxDb'
get_velocity_files(
  Χ,
  L,
  Genome,
  Transcriptome,
  out_path,
  style = c("genome", "Ensembl", "UCSC", "NCBI", "other"),
  isoform_action = c("separate", "collapse"),
  exon_option = c("full", "junction"),
  compress_fa = FALSE,
  width = 80L,
  chrs_only = TRUE
)
## S4 method for signature 'EnsDb'
get_velocity_files(
  Χ,
  L,
  Genome,
  Transcriptome,
  out_path,
  style = c("genome", "Ensembl", "UCSC", "NCBI", "other"),
  isoform_action = c("separate", "collapse"),
  exon_option = c("full", "junction"),
```

```
compress_fa = FALSE,
width = 80L,
use_transcript_version = TRUE,
use_gene_version = TRUE,
transcript_biotype_col = "TXBIOTYPE",
gene_biotype_col = "GENEBIOTYPE",
transcript_biotype_use = "all",
gene_biotype_use = "all",
chrs_only = TRUE
```

#### Arguments

Χ

Gene annotation with transcript and exon information. It can be a path to a GTF file with annotation of exon coordinates of transcripts, preferably from Ensembl. In the metadata, the following fields are required: type (e.g. whether the range of interest is a gene or transcript or exon or CDS), gene ID, and transcript ID. These fields need not to have standard names, as long as their names are specified in arguments of this function. It can also be a TxDb object, such as from the Bioconductor package TxDb.Hsapiens.UCSC.hg38.knownGene. It can also be a EnsDb object.

L

Length of the biological read. For instance, 10xv1: 98 nt, 10xv2: 98 nt, 10xv3: 91 nt, Drop-seq: 50 nt. If in doubt check read length in a fastq file for biological reads with the bash commands: If the fastq file is gzipped, then do zcat your\_file.fastq.gz | head on Linux. If on Mac, then zcat < your\_file.fastq.gz | head. Then you will see lines with nucleotide bases. Copy one of those lines and determine its length with str\_length in R or echo -n <the sequence> | wc -c in bash. Which file corresponds to biological reads depends on the particular technology.

Genome

Either a BSgenome or a XStringSet object of genomic sequences, where the intronic sequences will be extracted from. Use genomeStyles to check which styles are supported for your organism of interest; supported styles can be interconverted. If the style in your genome or annotation is not supported, then the style of chromosome names in the genome and annotation should be manually set to be consistent.

Transcriptome

A XStringSet, a path to a fasta file (can be gzipped) of the transcriptome which contains sequences of spliced transcripts, or NULL. The transcriptome here will be concatenated with the intronic sequences to give one fasta file. When NULL, the transriptome sequences will be extracted from the genome given the gene annotation, so it will be guaranteed that transcript IDs in the transcriptome and in the annotation match. Otherwise, the type of transcript ID in the transcriptome must match that in the gene annotation supplied via argument X.

out\_path

Directory to save the outputs written to disk. If this directory does not exist, then it will be created. Defaults to the current working directory.

style

Formatting of chromosome names. Use genomeStyles to check which styles are supported for your organism of interest and what those styles look like. This can also be a style supported for your organism different from the style used by the annotation and the genome. Then this style will be used for both the annotation and the genome. Can take the following values:

**genome** If style of the annnotation is different from that of the genome, then the style of the genome will be used.

> **other** Custom style, need to manually ensure that the style in annotation matches that of the genome.

**Ensembl** Or UCSC or NCBI, whichever is supported by your species of interest.

isoform\_action Character, indicating action to take with different transcripts of the same gene. Must be one of the following:

> collapse First, the union of all exons of different transcripts of a gene will be taken. Then the introns will be inferred from this union. Only the flanked intronic sequences are affected; isoforms will always be taken into account for spliced sequences or exon-exon junctions.

**separate** Introns from different transcripts will be kept separate.

exon\_option

Character, indicating how exonic sequences should be included in the kallisto index. Must be one of the following:

full The full cDNA sequences, which include the full exonic sequences, will be used. This is the default.

**junction** Only the exon-exon junctions, with L-1 bases on each side of the junctions, will be used.

compress\_fa

Logical, whether to compress the output fasta file. If TRUE, then the fasta file will be gzipped.

width

Maximum number of letters per line of sequence in the output fasta file. Must be an integer.

Extra arguments for methods.

transcript\_id

Character vector of length 1. Tag in attribute field corresponding to transcript IDs. This argument must be supplied and cannot be NA or NULL. Will throw error if tag indicated in this argument does not exist.

gene\_id

Character vector of length 1. Tag in attribute field corresponding to gene IDs. This argument must be supplied and cannot be NA or NULL. Note that this is different from gene symbols, which do not have to be unique. This can be Ensembl or Entrez IDs. However, if the gene symbols are in fact unique for each gene, you may supply the tag for human readable gene symbols to this argument. Will throw error if tag indicated in this argument does not exist. This is typically "gene\_id" for annotations from Ensembl and "gene" for refseq.

### transcript\_version

Character vector of length 1. Tag in attribute field corresponding to transcript version number. If your GTF file does not include transcript version numbers, or if you do not wish to include the version number, then use NULL for this argument. To decide whether to include transcript version number, check whether version numbers are included in the transcripts.txt in the kallisto output directory. If that file includes version numbers, then transscript version numbers must be included here as well. If that file does not include version numbers, then transcript version numbers must not be included here.

gene\_version

Character vector of length 1. Tag in attribute field corresponding to gene version number. If your GTF file does not include gene version numbers, or if you do not wish to include the version number, then use NULL for this argument. Unlike transcript version number, it's up to you whether to include gene version number.

version\_sep

Character to separate bewteen the main ID and the version number. Defaults to ".", as in Ensembl.

transcript\_biotype\_col

Character vector of length 1. Tag in attribute field corresponding to *transcript* biotype.

gene\_biotype\_col

Character vector of length 1. Tag in attribute field corresponding to *gene* biotype.

transcript\_biotype\_use

Character, can be "all" or a vector of *transcript* biotypes to be used. Transcript biotypes aren't entirely the same as gene biotypes. For instance, in Ensembl annotation, retained\_intron is a transcript biotype, but not a gene biotype. If "cellranger", then a warning will be given. See data("ensembl\_tx\_biotypes") for all available transcript biotypes from Ensembl.

gene\_biotype\_use

Character, can be "all", "cellranger", or a vector of *gene* biotypes to be used. If "cellranger", then the biotypes used by Cell Ranger's reference are used. See data("cellranger\_biotypes") for gene biotypes the Cell Ranger reference uses. See data("ensembl\_gene\_biotypes") for all available gene biotypes from Ensembl. Note that gene biotypes and transcript biotypes are not always the same.

chrs\_only

Logical, whether to include chromosomes only, for GTF and GFF files can contain annotations for scaffolds, which are not incorporated into chromosomes. This will also exclude haplotypes. Defaults to TRUE. Only applicable to species found in genomeStyles().

save\_filtered\_gtf

Logical. If filtering type, biotypes, and/or chromosomes, whether to save the filtered GRanges as a GTF file.

is\_circular

Logical vector of the same length as the number of sequences in the annotation and with the same names as the sequences, indicating whether the sequence is circular. If NULL, then all sequences will be assumed to be linear.

use\_transcript\_version

Logical, whether to include version number in the Ensembl transcript ID.

use\_gene\_version

Logical, whether to include version number in the Ensembl gene ID. Unlike transcript version number, it's up to you whether to include gene version number.

# Value

The following files will be written to disk in the directory out\_path:

- **cDNA\_introns.fa** A fasta file containing both the spliced transcripts and the flanked intronic sequences. The intronic sequences are flanked by L-1 nt of exonic sequences to capture reads from nascent transcript partially mapping to exons. If the exon is shorter than 2\*(L-1) nt, then the entire exon will be included in the intronic sequence. This will be used to build the kallisto index.
- **cDNA\_tx\_to\_capture.txt** A text file of transcript IDs of spliced transcripts. If exon\_option == "junction", then IDs of the exon-exon junctions. These IDs will have the pattern "transcript ID"-Jx, where x is a number differentiating between different junctions of the same transcript. Here x will always be ordered from 5' to 3' as on the plus strand.
- **introns\_tx\_to\_capture.txt** A text file of IDs of introns. The names will have the pattern "transcript ID"-Ix, where x is a number differentiating between introns of the same transcript. If all transcripts of the same gene are collapsed before inferring intronic sequences, gene ID will

22 make\_sparse\_matrix

be used in place of transcript ID. Here x will always be ordered from 5' to 3' as on the plus strand.

**tr2g.txt** A text file with two columns matching transcripts and introns to genes. The first column is transcript or intron ID, and the second column is the corresponding gene ID. The part for transcripts are generated from the gene annotation supplied.

Nothing is returned into the R session.

### **Examples**

make\_sparse\_matrix

Convert the Output of kallisto bus into Gene by Gell Matrix

### Description

This function takes the output file of kallisto bus, after being sorted and converted into text with bustools. See vignettes on the website of this package for a tutorial. The bustools output has 4 columns: barcode, UMI, equivalence class, and counts. This function converts that file into a sparse matrix that can be used in downstream analyses.

#### Usage

```
make_sparse_matrix(
  bus_path,
  tr2g,
  est_ncells,
  est_ngenes,
  whitelist = NULL,
  gene_count = TRUE,
  TCC = TRUE,
  single_gene = TRUE,
  verbose = TRUE,
  progress_unit = 5e+06
)
```

#### **Arguments**

bus\_path

Path to the sorted text bus output file.

tr2g

A Data frame with columns gene and transcript, in the same order as in the transcriptome index for kallisto. This argument can be missing or is ignored if only the TCC matrix, not the gene count matrix, is made.

make\_sparse\_matrix 23

est\_ncells Estimated number of cells; providing this argument will speed up computation

as it minimizes memory reallocation as vectors grow.

est\_ngenes Estimated number of genes or equivalence classes.

whitelist A character vector with valid cell barcodes. This is an optional argument, that

defaults to NULL. When it is NULL, all cell barcodes present that have some UMI assignable to genes or ECs will be included in the sparse matrix whether they are known to be valid or not. Barcodes with only UMIs that are not assignable

to genes or ECs will still be excluded.

gene\_count Logical, whether the gene count matrix should be returned.

TCC Logical, whether the TCC matrix should be returned.

single\_gene Logical, whether to use single gene mode. In single gene mode, only UMIs that

can be uniquely mapped to one gene are kept. Without single gene mode, UMIs

mapped to multiple genes will be evenly distributed to those genes.

verbose Whether to display progress.

progress\_unit How many iteration to print one progress update when reading in the kallisto

bus file.

### **Details**

This function can generate both the gene count matrix and the transcript compatibility count (TCC) matrix. The TCC matrix has barcodes in the columns and equivalence classes in the rows. See Ntranos et al. 2016 for more information about the RCC matrix.

For 10x data sets, you can find a barcode whitelist file that comes with CellRanger installation. You don't need to run CellRanger to get that. An example path to get the whitelist file is cellranger-2.1.0/cellranger-cs. for v2 chemistry.

### Value

If both gene count and TCC matrices are returned, then this function returns a list with two matrices, each with genes/equivalence classes in the rows and barcodes in the columns. If only one of gene count and TCC matrices is returned, then a dgCMatrix with genes/equivalence classes in the rows and barcodes in the columns. These matrices are unfiltered. Please filter the empty droplets before downstream analysis.

# See Also

EC2gene

### **Examples**

```
# Load toy example for testing
toy_path <- system.file("testdata", package = "BUSpaRse")
load(paste(toy_path, "toy_example.RData", sep = "/"))
out_fn <- paste0(toy_path, "/output.sorted.txt")
# With whitelist
m <- make_sparse_matrix(out_fn, tr2g_toy, 10, 3, whitelist = whitelist,
    gene_count = TRUE, TCC = FALSE, single_gene = TRUE,
    verbose = FALSE)</pre>
```

24 read\_count\_output

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Match chromosome naming styles of annotation and genome

### **Description**

Internal use. This function matches chromosome naming styles. It will also give the genome and the annotation the same genome slot. This function assumes that the annotation and the genome refer to the same version of genome. If more than one style, then the first element will be used.

#### Usage

```
match_style(Genome, annot, style)
```

#### **Arguments**

Genome Either a BSgenome or a XStringSet object of genomic sequences, where the

intronic sequences will be extracted from. Use genomeStyles to check which styles are supported for your organism of interest; supported styles can be interconverted. If the style in your genome or annotation is not supported, then the style of chromosome names in the genome and annotation should be manually

set to be consistent.

annot Genome annotation, an object of a class with a seqlevels method, such as

GRanges, TxDb, and EnsDb.

style Formatting of chromosome names. Use genomeStyles to check which styles

are supported for your organism of interest and what those styles look like. This can also be a style supported for your organism different from the style used by the annotation and the genome. Then this style will be used for both the

annotation and the genome. Can take the following values:

**genome** If style of the annnotation is different from that of the genome, then the style of the genome will be used.

**other** Custom style, need to manually ensure that the style in annotation matches that of the genome.

Ensembl Or UCSC or NCBI, whichever is supported by your species of interest.

### Value

A list of two. The first element is the genome with the proper style, and the second element is the annotation with the proper style.

 $read\_count\_output$ 

Read matrix along with barcode and gene names

### **Description**

This function takes in a directory and name and reads the mtx file, genes, and barcodes from the output of bustools to return a sparse matrix with column names and row names.

read\_velocity\_output 25

#### Usage

```
read_count_output(dir, name, tcc = FALSE)
```

#### **Arguments**

dir Directory with the bustools count outputs.

name The files in the output directory should be "name".mtx, "name".genes.txt, and

"name".barcodes.txt.

tcc Logical, whether the matrix of interest is a TCC matrix. Defaults to FALSE.

#### Value

A dgCMatrix with barcodes as column names and genes as row names.

### **Examples**

```
# Internal toy data used for unit testing
toy_path <- system.file("testdata", package = "BUSpaRse")
m <- read_count_output(toy_path, name = "genes", tcc = FALSE)</pre>
```

read\_velocity\_output Read intronic and exonic matrices into R

### **Description**

Read intronic and exonic matrices into R

### Usage

```
read_velocity_output(spliced_dir, unspliced_dir, spliced_name, unspliced_name)
```

### **Arguments**

spliced\_dir Directory with mtx file for UMI counts of spliced transcripts.

unspliced\_dir Directory with mtx file for UMI counts of unspliced transcripts.

spliced\_name The files in the splicedd directory should be <spliced\_name>.mtx, <spliced\_name>.genes.txt,

 $and <\!\!spliced\_name\!\!>\!.barcodes.txt.$ 

unspliced\_name The files in the unsplicedd directory should be <unspliced\_name>.mtx, <un-

 $spliced\_name{>}.genes.txt, and <\!unspliced\_name{>}.barcodes.txt.$ 

#### Value

A list of two dgCMatrix with barcodes as column names and genes as row names. The elements of the list will be spliced and unspliced.

# Examples

```
# Internal toy data used for unit testing
toy_path <- system.file("testdata", package = "BUSpaRse")
m <- read_velocity_output(toy_path, toy_path,
    spliced_name = "genes",
    unspliced_name = "genes")</pre>
```

26 save\_tr2g\_bustools

refseq\_gff\_mcols

Tags in the attributes field of RefSeq GFF files

### **Description**

These are the column names of the mcols when the Ensembl GTF file is read into R as a GRanges, including gene, transcript\_id, gene\_biotype, description, and so on, and the mandatory tags like ID, Name, and Parent.

#### Usage

```
refseq_gff_mcols
```

#### **Format**

A character vector

#### Source

Ensembl version 99 human GTF file

save\_tr2g\_bustools

Save transcript to gene file for use in bustools

### **Description**

This function saves the transcript to gene data frame generated by this package in whatever means in a format required by bustools. In order to use bustools to generate the gene count or TCC matrix, a file that maps transcripts to genes is required. This should be a tsv file with 2 columns: the first column for transcript ID and the second for gene ID. The order of transcripts in this file must be the same as the order in the kallisto index, and this ordering can be ensured by the function sort\_tr2g. There must also be no headers. All columns other than transcript and gene will be discarded. To save a file with those columns, directly save the transcript to gene data frame with function like write.table, readr::write\_delim.

### Usage

```
save_tr2g_bustools(tr2g, file_save = "./tr2g.tsv")
```

### **Arguments**

tr2g The data frame output from the tr2g\_\* family of functions.

file\_save File name of the file to be saved. The directory in which the file is to be saved

must exist.

### Value

Nothing is returned into the R session. A tsv file of the format required by bustools with the name and directory specified will be written to disk.

sort\_tr2g

#### Note

This function has been superseded by the new version of tr2g\_\* functions that can extract transcriptome for only the biotypes specified and with only the standard chromosomes. The new version of tr2g\_\* functions also sorts the transcriptome so the tr2g and the transcriptome have transcripts in the same order, and write the tr2g.tsv file in the bustools format.

### **Examples**

```
toy_path <- system.file("testdata", package = "BUSpaRse")
file_use <- paste(toy_path, "gtf_test.gtf", sep = "/")
tr2g <- tr2g_gtf(file = file_use, get_transcriptome = FALSE,
    write_tr2g = FALSE, save_filtered_gtf = FALSE)
save_tr2g_bustools(tr2g, file_save = "./tr2g.tsv")
# Clean up files from the example
file.remove("tr2g.tsv")</pre>
```

sort\_tr2g

Sort transcripts to the same order as in kallisto index

### **Description**

This function takes the data frame output from the tr2g\_\* family of functions in this package as the input, and sorts it so the transcripts are in the same order as in the kallisto index used to generate the bus file. Sorting is vital to obtain the correct sparse matrix from the bus file as equivalence class notations are based on the index of transcripts in the kallisto index.

### Usage

```
sort_tr2g(tr2g, file, kallisto_out_path)
```

### **Arguments**

tr2g The data frame output from the tr2g\_\* family of functions.

file Character vector of length 1, path to a tsv file with transcript IDs and the corre-

sponding gene IDs, in the format required for bustools, or written by save\_tr2g\_bustools.

kallisto\_out\_path

Character vector of length 1, path to the directory for the outputs of kallisto bus.

#### Details

Since the attribute field of GTF and GFF3 files varies across sources, output from tr2g\_gtf and tr2g\_gff3 may need further clean up. You may also supply gene and transcript IDs from other sources. This function should be used after the clean up, when the transcript IDs in the cleaned up data frame have the same format as those in transcript

### Value

A data frame with columns transcript and gene and the other columns present in tr2g or the data frame in file, with the transcript IDs sorted to be in the same order as in the kallisto index.

28 species2dataset

#### Note

This function has been superseded by the new version of tr2g\_\* functions that can extract transcriptome for only the biotypes specified and with only the standard chromosomes. The new version of tr2g \* functions also sorts the transcriptome so the tr2g and the transcriptome have transcripts in the same order.

#### See Also

```
Other functions to retrieve transcript and gene info: tr2g_EnsDb(), tr2g_TxDb(), tr2g_ensembl(),
tr2g_fasta(), tr2g_gff3(), tr2g_gtf(), transcript2gene()
```

### **Examples**

```
toy_path <- system.file("testdata", package = "BUSpaRse")</pre>
file_use <- paste(toy_path, "gtf_test.gtf", sep = "/")</pre>
tr2g <- tr2g_gtf(file = file_use, get_transcriptome = FALSE,</pre>
  write_tr2g = FALSE, save_filtered_gtf = FALSE, transcript_version = NULL)
tr2g <- sort_tr2g(tr2g, kallisto_out_path = toy_path)</pre>
```

species2dataset

Convert Latin species name to dataset name

### **Description**

This function converts Latin species name to a dataset name in biomart to query gene and transcript ID.

### Usage

```
species2dataset(
  species,
  type = c("vertebrate", "metazoa", "plant", "fungus", "protist")
)
```

### **Arguments**

species

A character vector of Latin names of species present in this scRNA-seq dataset.

This is used to retrieve Ensembl information from biomart.

type

A character vector indicating the type of each species. Each element must be one of "vertebrate", "metazoa", "plant", "fungus", and "protist". If length is 1, then this type will be used for all species specified here. Can be missing if

fasta\_file is specified.

#### Value

The appropriate dataset name for biomart.

### **Examples**

```
species2dataset(species = "Homo sapiens")
```

standardize\_tags 29

standardize_tags	Standardize GRanges field names	
------------------	---------------------------------	--

### **Description**

To avoid introducing rlang as another dependency for tidyeval. This function will also convert exon numbers to integer.

### Usage

```
standardize_tags(gr, gene_id, transcript_id)
```

# **Arguments**

gr A GRanges object.

gene\_id Name of the metadata field for gene ID.

transcript\_id Name of the metadata field for transcript ID.

### Value

A GRanges object with standardized names: gene ID as gene\_id, and transcript ID as transcript\_id.

### **Description**

Exclude chromosomes present in the annotation but absent from the genome and add information about circular chromosomes.

# Usage

```
subset_annot(Genome, annot)
## S4 method for signature 'DNAStringSet'
subset_annot(Genome, annot)
## S4 method for signature 'BSgenome'
subset_annot(Genome, annot)
```

### **Arguments**

Genome Either a BSgenome or a XStringSet object of genomic sequences, where the

intronic sequences will be extracted from. Use <code>genomeStyles</code> to check which styles are supported for your organism of interest; supported styles can be interconverted. If the style in your genome or annotation is not supported, then the style of chromosome names in the genome and annotation should be manually

set to be consistent.

annot Either a GRanges object or a TxDb object for gene annotation.

30 tr2g\_EnsDb

#### Value

A subsetted genome annotation of the same type of the input genome annotation.

### **Examples**

```
library(BSgenome.Hsapiens.UCSC.hg38)
library(EnsDb.Hsapiens.v86)
library(GenomeInfoDb)
gn <- BSgenome.Hsapiens.UCSC.hg38
seqlevelsStyle(gn) <- "Ensembl"
subset_annot(gn, EnsDb.Hsapiens.v86)</pre>
```

sub\_annot

Remove chromosomes in anotation absent from genome

### **Description**

Remove chromosomes in anotation absent from genome

### Usage

```
sub_annot(chrs_use, annot)
```

### **Arguments**

chrs\_use Character vector of names of chromosomes present in both the annotation and

the genome.

annot Either a GRanges object or a TxDb object for gene annotation.

## Value

A subsetted genome annotation of the same type of the input genome annotation.

tr2g\_EnsDb Get transcript and gene info from EnsDb objects

# Description

Bioconductor provides Ensembl genome annotation in AnnotationHub; older versions of Ensembl annotation can be obtained from packages like EnsDb.Hsapiens.v86. This is an alternative to querying Ensembl with biomart; Ensembl's server seems to be less stable than that of Bioconductor. However, more information and species are available on Ensembl biomart than on AnnotationHub.

tr2g\_EnsDb 31

#### Usage

```
tr2g_EnsDb(
  ensdb,
  Genome = NULL,
 get_transcriptome = TRUE,
 out_path = ".",
 write_tr2g = TRUE,
 other_attrs = NULL,
  use_gene_name = TRUE,
  use_transcript_version = TRUE,
  use_gene_version = TRUE,
  transcript_biotype_col = "TXBIOTYPE",
  gene_biotype_col = "GENEBIOTYPE",
  transcript_biotype_use = "all",
  gene_biotype_use = "all",
  chrs_only = TRUE,
  compress_fa = FALSE,
  overwrite = FALSE
)
```

### **Arguments**

ensdb

Ann EnsDb object, such as from AnnotationHub or EnsDb. Hsapiens. v86.

Genome

Either a BSgenome or a XStringSet object of genomic sequences, where the intronic sequences will be extracted from. Use genomeStyles to check which styles are supported for your organism of interest; supported styles can be interconverted. If the style in your genome or annotation is not supported, then the style of chromosome names in the genome and annotation should be manually set to be consistent.

get\_transcriptome

Logical, whether to extract transcriptome from genome with the GTF file. If filtering biotypes or chromosomes, the filtered GRanges will be used to extract transcriptome.

out\_path

Directory to save the outputs written to disk. If this directory does not exist, then it will be created. Defaults to the current working directory.

write\_tr2g

Logical, whether to write tr2g to disk. If TRUE, then a file tr2g.tsv will be written into out\_path.

other\_attrs

Character vector. Other attributes to get from the EnsDb object, such as gene symbol and position on the genome. Use columns to see which attributes are available.

use\_gene\_name

Logical, whether to get gene names.

use\_transcript\_version

Logical, whether to include version number in the Ensembl transcript ID. To decide whether to include transcript version number, check whether version numbers are included in the transcripts.txt in the kallisto output directory. If that file includes version numbers, then transcript version numbers must be included here as well. If that file does not include version numbers, then transcript version numbers must not be included here.

use\_gene\_version

Logical, whether to include version number in the Ensembl gene ID. Unlike transcript version number, it's up to you whether to include gene version number.

32 tr2g\_EnsDb

transcript\_biotype\_col

Character vector of length 1. Tag in attribute field corresponding to *transcript* biotype.

gene\_biotype\_col

Character vector of length 1. Tag in attribute field corresponding to *gene* biotype.

transcript\_biotype\_use

Character, can be "all" or a vector of *transcript* biotypes to be used. Transcript biotypes aren't entirely the same as gene biotypes. For instance, in Ensembl annotation, retained\_intron is a transcript biotype, but not a gene biotype. If "cellranger", then a warning will be given. See data("ensembl\_tx\_biotypes") for all available transcript biotypes from Ensembl.

gene\_biotype\_use

Character, can be "all", "cellranger", or a vector of *gene* biotypes to be used. If "cellranger", then the biotypes used by Cell Ranger's reference are used. See data("cellranger\_biotypes") for gene biotypes the Cell Ranger reference uses. See data("ensembl\_gene\_biotypes") for all available gene biotypes from Ensembl. Note that gene biotypes and transcript biotypes are not always the same.

chrs\_only

Logical, whether to include chromosomes only, for GTF and GFF files can contain annotations for scaffolds, which are not incorporated into chromosomes. This will also exclude haplotypes. Defaults to TRUE. Only applicable to species found in genomeStyles().

compress\_fa

Logical, whether to compress the output fasta file. If TRUE, then the fasta file will be gzipped.

overwrite

Logical, whether to overwrite if files with names of outputs written to disk already exist.

#### Value

A data frame with at least 2 columns: gene for gene ID, transcript for transcript ID, and optionally gene\_name for gene names. If other\_attrs has been specified, then those will also be columns in the data frame returned.

# See Also

```
ensembl_gene_biotypes ensembl_tx_biotypes cellranger_biotypes
```

```
Other functions to retrieve transcript and gene info: sort_tr2g(), tr2g_TxDb(), tr2g_ensembl(), tr2g_fasta(), tr2g_gff3(), tr2g_gtf(), transcript2gene()
```

#### **Examples**

```
library(EnsDb.Hsapiens.v86)
tr2g_EnsDb(EnsDb.Hsapiens.v86, get_transcriptome = FALSE, write_tr2g = FALSE,
use_transcript_version = FALSE,
use_gene_version = FALSE)
```

 $tr2g\_ensembl$  33

tr2g_ensembl	Get transcript and gene info from Ensembl	

### **Description**

This function queries Ensembl biomart to convert transcript IDs to gene IDs.

# Usage

```
tr2g_ensembl(
  species,
  type = c("vertebrate", "metazoa", "plant", "fungus", "protist"),
  out_path = ".",
  write_tr2g = TRUE,
  other_attrs = NULL,
  use_gene_name = TRUE,
  use_transcript_version = TRUE,
  use_gene_version = TRUE,
  transcript_biotype_col = "transcript_biotype",
  gene_biotype_col = "gene_biotype",
  transcript_biotype_use = "all",
  gene_biotype_use = "all",
  chrs_only = TRUE,
  ensembl_version = NULL,
  overwrite = FALSE,
  verbose = TRUE,
)
```

# Arguments

species	Character vector of length 1, Latin name of the species of interest.
type	Character, must be one of "vertebrate", "metazoa", "plant", "fungus" and "protist". Passing "vertebrate" will use the default www.ensembl.org host. Gene annotation of some common invertebrate model organisms, such as <i>Drosophila melanogaster</i> , are available on www.ensembl.org so for these invertebrate model organisms, "vertebrate" can be used for this argument. Passing values other than "vertebrate" will use other Ensembl hosts. For animals absent from www.ensembl.org, try "metazoa".
out_path	Directory to save the outputs written to disk. If this directory does not exist, then it will be created. Defaults to the current working directory.
write_tr2g	Logical, whether to write tr2g to disk. If TRUE, then a file tr2g.tsv will be written into out_path.
other_attrs	Character vector. Other attributes to get from Ensembl, such as gene symbol and position on the genome. Use listAttributes to see which attributes are available.
use_gene_name	Logical, whether to get gene names.

34 tr2g\_ensembl

### use\_transcript\_version

Logical, whether to include version number in the Ensembl transcript ID. To decide whether to include transcript version number, check whether version numbers are included in the transcripts.txt in the kallisto output directory. If that file includes version numbers, then transcript version numbers must be included here as well. If that file does not include version numbers, then transcript version numbers must not be included here.

#### use\_gene\_version

Logical, whether to include version number in the Ensembl gene ID. Unlike transcript version number, it's up to you whether to include gene version number.

#### transcript\_biotype\_col

Character vector of length 1. Tag in attribute field corresponding to *transcript* biotype.

#### gene\_biotype\_col

Character vector of length 1. Tag in attribute field corresponding to *gene* biotype.

### transcript\_biotype\_use

Character, can be "all" or a vector of *transcript* biotypes to be used. Transcript biotypes aren't entirely the same as gene biotypes. For instance, in Ensembl annotation, retained\_intron is a transcript biotype, but not a gene biotype. If "cellranger", then a warning will be given. See data("ensembl\_tx\_biotypes") for all available transcript biotypes from Ensembl.

#### gene\_biotype\_use

Character, can be "all", "cellranger", or a vector of *gene* biotypes to be used. If "cellranger", then the biotypes used by Cell Ranger's reference are used. See data("cellranger\_biotypes") for gene biotypes the Cell Ranger reference uses. See data("ensembl\_gene\_biotypes") for all available gene biotypes from Ensembl. Note that gene biotypes and transcript biotypes are not always the same.

### chrs\_only

Logical, whether to include chromosomes only, for GTF and GFF files can contain annotations for scaffolds, which are not incorporated into chromosomes. This will also exclude haplotypes. Defaults to TRUE. Only applicable to species found in genomeStyles().

### ensembl\_version

Integer version number of Ensembl (e.g. 94 for the October 2018 release). This argument defaults to NULL, which will use the current release of Ensembl. Use listEnsemblArchives to see the version number corresponding to the Ensembl release of a particular date. The version specified here must match the version of Ensembl where the transcriptome used to build the kallisto index was downloaded. This only works for vertebrates and the most common invertebrate model organisms like *Drosophila melanogaster* and *C. elegans* (i.e. www.ensembl.org and its mirrors), not the other Ensembl sites for plants, protists, fungi, and metazoa.

#### overwrite

Logical, whether to overwrite if files with names of outputs written to disk already exist.

### verbose

Whether to display progress.

. . .

Othe arguments to be passed to useMart, such as mirror. Note that setting mirrors other than the default, e.g. uswest, does not work for archived versions.

 $tr2g_fasta$  35

#### Value

A data frame with at least 2 columns: gene for gene ID, transcript for transcript ID, and optionally gene\_name for gene names. If other\_attrs has been specified, then those will also be columns in the data frame returned.

#### See Also

```
dl_transcriptome
```

```
Other functions to retrieve transcript and gene info: sort_tr2g(), tr2g_EnsDb(), tr2g_TxDb(), tr2g_fasta(), tr2g_gff3(), tr2g_gtf(), transcript2gene()
```

### **Examples**

```
tr2g <- tr2g_ensembl(species = "Danio rerio",
other_attrs = "description", write_tr2g = FALSE)
# This will use plants.ensembl.org as host instead of www.ensembl.org
tr2g <- tr2g_ensembl(species = "Arabidopsis thaliana", type = "plant",
    write_tr2g = FALSE)</pre>
```

tr2g\_fasta

Get transcript and gene info from names in FASTA files

### **Description**

FASTA files, such as those for cDNA and ncRNA from Ensembl, might have genome annotation information in the name of each sequence entry. This function extracts the transcript and gene IDs from such FASTA files.

### Usage

```
tr2g_fasta(
   file,
   out_path = ".",
   write_tr2g = TRUE,
   use_gene_name = TRUE,
   use_transcript_version = TRUE,
   use_gene_version = TRUE,
   transcript_biotype_use = "all",
   gene_biotype_use = "all",
   chrs_only = TRUE,
   save_filtered = TRUE,
   compress_fa = FALSE,
   overwrite = FALSE
)
```

### **Arguments**

file Path to the FASTA file to be read. The file can remain gzipped.

out\_path Directory to save the outputs written to disk. If this directory does not exist, then it will be created. Defaults to the current working directory.

 $tr2g_fasta$ 

write\_tr2g Logical, whether to write tr2g to disk. If TRUE, then a file tr2g.tsv will be written into out\_path.

use\_gene\_name Logical, whether to get gene names.

use\_transcript\_version

Logical, whether to include version number in the Ensembl transcript ID. To decide whether to include transcript version number, check whether version numbers are included in the transcripts.txt in the kallisto output directory. If that file includes version numbers, then transcript version numbers must be included here as well. If that file does not include version numbers, then transcript version numbers must not be included here.

use\_gene\_version

Logical, whether to include version number in the Ensembl gene ID. Unlike transcript version number, it's up to you whether to include gene version number.

transcript\_biotype\_use

Character, can be "all" or a vector of *transcript* biotypes to be used. Transcript biotypes aren't entirely the same as gene biotypes. For instance, in Ensembl annotation, retained\_intron is a transcript biotype, but not a gene biotype. If "cellranger", then a warning will be given. See data("ensembl\_tx\_biotypes") for all available transcript biotypes from Ensembl.

gene\_biotype\_use

Character, can be "all", "cellranger", or a vector of *gene* biotypes to be used. If "cellranger", then the biotypes used by Cell Ranger's reference are used. See data("cellranger\_biotypes") for gene biotypes the Cell Ranger reference uses. See data("ensembl\_gene\_biotypes") for all available gene biotypes from Ensembl. Note that gene biotypes and transcript biotypes are not always the same.

the same

chrs\_only Logical, whether to include chromosomes only, for GTF and GFF files can contain annotations for scaffolds, which are not incorporated into chromosomes. This will also exclude haplotypes. Defaults to TRUE. Only applicable to species

found in genomeStyles().

If TRUE, the file will be tx\_filtered.fa in out\_path.

compress\_fa Logical, whether to compress the output fasta file. If TRUE, then the fasta file

will be gzipped.

overwrite Logical, whether to overwrite if files with names of outputs written to disk al-

ready exist.

# Details

At present, this function only works with FASTA files from Ensembl, and uses regex to extract vertebrate Ensembl IDs. Sequence names should be formatted as follows:

```
ENST00000632684.1 cdna chromosome:GRCh38:7:142786213:142786224:1
gene:ENSG00000282431.1 gene_biotype:TR_D_gene transcript_biotype:TR_D_gene
gene_symbol:TRBD1 description:T cell receptor beta diversity 1
[Source:HGNC Symbol;Acc:HGNC:12158]
```

If your FASTA file sequence names are formatted differently, then you must extract the transcript and gene IDs by some other means. The Bioconductor package Biostrings is recommended; after reading the FASTA file into R, the sequence names can be accessed by the names function.

 $tr2g\_gff3$  37

While normally, you should call sort\_tr2g to sort the transcript IDs from the output of the tr2g\_\* family of functions, If the FASTA file supplied here is the same as the one used to build the kallisto index, then the transcript IDs in the output of this function are in the same order as in the kallisto index, so you can skip sort\_tr2g and proceed directly to EC2gene with the output of this function.

#### Value

A data frame with at least 2 columns: gene for gene ID, transcript for transcript ID, and optionally gene\_name for gene names.

#### See Also

```
ensembl_gene_biotypes ensembl_tx_biotypes cellranger_biotypes

Other functions to retrieve transcript and gene info: sort_tr2g(), tr2g_EnsDb(), tr2g_TxDb(), tr2g_ensembl(), tr2g_gff3(), tr2g_gtf(), transcript2gene()
```

# Examples

```
toy_path <- system.file("testdata", package = "BUSpaRse")
file_use <- paste(toy_path, "fasta_test.fasta", sep = "/")
tr2g <- tr2g_fasta(file = file_use, save_filtered = FALSE, write_tr2g = FALSE)</pre>
```

tr2g\_gff3

Get transcript and gene info from GFF3 file

## **Description**

This function reads a GFF3 file and extracts the transcript ID and corresponding gene ID. This function assumes that the GFF3 file is properly formatted. See <a href="http://gmod.org/wiki/GFF3">http://gmod.org/wiki/GFF3</a> for a detailed description of proper GFF3 format. Note that GTF files have a somewhat different and simpler format in the attribute field, which this function does not support. See <a href="http://mblab.wustl.edu/GTF2.html">http://mblab.wustl.edu/GTF2.html</a> for a detailed description of proper GTF format. To extract transcript and gene information from GTF files, see the function  $tr2g_gtf$  in this package. Some files bearing the .gff3 are in fact more like the GTF format. If this is so, then change the extension to .gtf and use the function  $tr2g_gtf$  in this package instead.

```
tr2g_gff3(
    file,
    Genome = NULL,
    get_transcriptome = TRUE,
    out_path = ".",
    write_tr2g = TRUE,
    transcript_id = "transcript_id",
    gene_id = "gene_id",
    gene_name = "Name",
    transcript_version = "version",
    gene_version = "version",
    version_sep = ".",
    transcript_biotype_col = "biotype",
```

 $tr2g_gff3$ 

```
gene_biotype_col = "biotype",
  transcript_biotype_use = "all",
  gene_biotype_use = "all",
  chrs_only = TRUE,
  compress_fa = FALSE,
  save_filtered_gff = TRUE,
  overwrite = FALSE,
  source = c("ensembl", "refseq")
)
```

## **Arguments**

file

Path to a GTF file to be read. The file can remain gzipped. Use getGTF from the biomartr package to download GTF files from Ensembl, and use getGFF from biomartr to download GFF3 files from Ensembl and RefSeq.

Genome

Either a BSgenome or a XStringSet object of genomic sequences, where the intronic sequences will be extracted from. Use genomeStyles to check which styles are supported for your organism of interest; supported styles can be interconverted. If the style in your genome or annotation is not supported, then the style of chromosome names in the genome and annotation should be manually set to be consistent.

get\_transcriptome

Logical, whether to extract transcriptome from genome with the GTF file. If filtering biotypes or chromosomes, the filtered GRanges will be used to extract transcriptome.

out\_path

Directory to save the outputs written to disk. If this directory does not exist, then it will be created. Defaults to the current working directory.

write\_tr2g

Logical, whether to write tr2g to disk. If TRUE, then a file tr2g.tsv will be written into out\_path.

transcript\_id

Character vector of length 1. Tag in attribute field corresponding to transcript IDs. This argument must be supplied and cannot be NA or NULL. Will throw error if tag indicated in this argument does not exist.

gene\_id

Character vector of length 1. Tag in attribute field corresponding to gene IDs. This argument must be supplied and cannot be NA or NULL. Note that this is different from gene symbols, which do not have to be unique. This can be Ensembl or Entrez IDs. However, if the gene symbols are in fact unique for each gene, you may supply the tag for human readable gene symbols to this argument. Will throw error if tag indicated in this argument does not exist. This is typically "gene\_id" for annotations from Ensembl and "gene" for refseq.

gene\_name

Character vector of length 1. Tag in attribute field corresponding to gene symbols. This argument can be NA or NULL if you are fine with non-human readable gene IDs and do not wish to extract human readable gene symbols.

transcript\_version

Character vector of length 1. Tag in attribute field corresponding to *transcript* version number. If your GTF file does not include transcript version numbers, or if you do not wish to include the version number, then use NULL for this argument. To decide whether to include transcript version number, check whether version numbers are included in the transcripts.txt in the kallisto output directory. If that file includes version numbers, then transcript version numbers must be included here as well. If that file does not include version numbers, then transcript version numbers must not be included here.

tr2g\_gff3 39

gene\_version

Character vector of length 1. Tag in attribute field corresponding to *gene* version number. If your GTF file does not include gene version numbers, or if you do not wish to include the version number, then use NULL for this argument. Unlike transcript version number, it's up to you whether to include gene version number.

version\_sep

Character to separate bewteen the main ID and the version number. Defaults to ".", as in Ensembl.

transcript\_biotype\_col

Character vector of length 1. Tag in attribute field corresponding to *transcript* biotype.

gene\_biotype\_col

Character vector of length 1. Tag in attribute field corresponding to *gene* biotype.

transcript\_biotype\_use

Character, can be "all" or a vector of *transcript* biotypes to be used. Transcript biotypes aren't entirely the same as gene biotypes. For instance, in Ensembl annotation, retained\_intron is a transcript biotype, but not a gene biotype. If "cellranger", then a warning will be given. See data("ensembl\_tx\_biotypes") for all available transcript biotypes from Ensembl.

gene\_biotype\_use

Character, can be "all", "cellranger", or a vector of *gene* biotypes to be used. If "cellranger", then the biotypes used by Cell Ranger's reference are used. See data("cellranger\_biotypes") for gene biotypes the Cell Ranger reference uses. See data("ensembl\_gene\_biotypes") for all available gene biotypes from Ensembl. Note that gene biotypes and transcript biotypes are not always the same.

chrs\_only

Logical, whether to include chromosomes only, for GTF and GFF files can contain annotations for scaffolds, which are not incorporated into chromosomes. This will also exclude haplotypes. Defaults to TRUE. Only applicable to species found in genomeStyles().

compress\_fa

Logical, whether to compress the output fasta file. If TRUE, then the fasta file will be gzipped.

save\_filtered\_gff

Logical. If filtering type, biotypes, and/or chromosomes, whether to save the filtered GRanges as a GFF3 file.

overwrite

Logical, whether to overwrite if files with names of outputs written to disk already exist.

source

Name of the database where this GFF3 file was downloaded. Must be either "ensembl" or "refseq".

#### **Details**

Transcript and gene versions may not be present in all GTF files, so these arguments are optional. This function has arguments for transcript and gene version numbers because Ensembl IDs have version numbers. For Ensembl IDs, we recommend including the version number, since a change in version number signals a change in the entity referred to by the ID after reannotation. If a version is used, then it will be appended to the ID, separated by version\_sep.

The transcript and gene IDs are The attribute field (the last field) of GTF files can be complicated and inconsistent across different sources. Please check the attribute tags in your GTF file and consider the arguments of this function carefully. The defaults are set according to Ensembl GTF

40 tr2g\_GRanges

files; defaults may not work for files from other sources. Due to the general lack of standards for the attribute field, you may need to further clean up the output of this function.

#### Value

A data frame at least 2 columns: gene for gene ID, transcript for transcript ID, and optionally, gene\_name for gene names.

## Note

The defaults here are for Ensembl GFF3 files. To see all attribute tags for Ensembl and RefSeq GFF3 files, see data("ensembl\_gff\_mcols") and data("refseq\_gff\_mcols").

#### See Also

 $ensembl\_gene\_biotypes\ ensembl\_tx\_biotypes\ cellranger\_biotypes\ ensembl\_gtf\_mcols\ ensembl\_gff\_mcols\ refseq\_gff\_mcols$ 

```
Other functions to retrieve transcript and gene info: sort_tr2g(), tr2g_EnsDb(), tr2g_TxDb(), tr2g_ensembl(), tr2g_fasta(), tr2g_gtf(), transcript2gene()
```

## **Examples**

```
toy_path <- system.file("testdata", package = "BUSpaRse")
file_use <- paste(toy_path, "gff3_test.gff3", sep = "/")
# Default
tr2g <- tr2g_gff3(file = file_use, write_tr2g = FALSE,
get_transcriptome = FALSE, save_filtered_gff = FALSE)
# Excluding version numbers
tr2g <- tr2g_gff3(file = file_use, transcript_version = NULL,
    gene_version = NULL, write_tr2g = FALSE, get_transcriptome = FALSE,
    save_filtered_gff = FALSE)</pre>
```

tr2g\_GRanges

Get transcript and gene info from GRanges

# Description

Internal use, for GRanges from GTF files

```
tr2g_GRanges(
   gr,
   Genome = NULL,
   get_transcriptome = TRUE,
   out_path = ".",
   write_tr2g = TRUE,
   transcript_id = "transcript_id",
   gene_id = "gene_id",
   gene_name = "gene_name",
   transcript_version = "transcript_version",
   gene_version = "gene_version",
```

tr2g\_GRanges 41

```
version_sep = ".",
  transcript_biotype_col = "transcript_biotype",
  gene_biotype_col = "gene_biotype",
  transcript_biotype_use = "all",
  gene_biotype_use = "all",
  chrs_only = TRUE,
  compress_fa = FALSE,
  save_filtered_gtf = TRUE,
  overwrite = FALSE
)
```

#### **Arguments**

gr

A GRanges object. The metadata columns should be atomic vectors, not lists.

Genome

Either a BSgenome or a XStringSet object of genomic sequences, where the intronic sequences will be extracted from. Use genomeStyles to check which styles are supported for your organism of interest; supported styles can be interconverted. If the style in your genome or annotation is not supported, then the style of chromosome names in the genome and annotation should be manually set to be consistent.

get\_transcriptome

Logical, whether to extract transcriptome from genome with the GTF file. If filtering biotypes or chromosomes, the filtered GRanges will be used to extract transcriptome.

out\_path

Directory to save the outputs written to disk. If this directory does not exist, then it will be created. Defaults to the current working directory.

write\_tr2g

Logical, whether to write tr2g to disk. If TRUE, then a file tr2g.tsv will be written into out\_path.

transcript\_id

Character vector of length 1. Tag in attribute field corresponding to transcript IDs. This argument must be supplied and cannot be NA or NULL. Will throw error if tag indicated in this argument does not exist.

gene\_id

Character vector of length 1. Tag in attribute field corresponding to gene IDs. This argument must be supplied and cannot be NA or NULL. Note that this is different from gene symbols, which do not have to be unique. This can be Ensembl or Entrez IDs. However, if the gene symbols are in fact unique for each gene, you may supply the tag for human readable gene symbols to this argument. Will throw error if tag indicated in this argument does not exist. This is typically "gene\_id" for annotations from Ensembl and "gene" for refseq.

gene\_name

Character vector of length 1. Tag in attribute field corresponding to gene symbols. This argument can be NA or NULL if you are fine with non-human readable gene IDs and do not wish to extract human readable gene symbols.

transcript\_version

Character vector of length 1. Tag in attribute field corresponding to *transcript* version number. If your GTF file does not include transcript version numbers, or if you do not wish to include the version number, then use NULL for this argument. To decide whether to include transcript version number, check whether version numbers are included in the transcripts.txt in the kallisto output directory. If that file includes version numbers, then transcript version numbers must be included here as well. If that file does not include version numbers, then transcript version numbers must not be included here.

42 tr2g\_gtf

gene\_version

Character vector of length 1. Tag in attribute field corresponding to *gene* version number. If your GTF file does not include gene version numbers, or if you do not wish to include the version number, then use NULL for this argument. Unlike transcript version number, it's up to you whether to include gene version number.

version\_sep

Character to separate bewteen the main ID and the version number. Defaults to ".", as in Ensembl.

transcript\_biotype\_col

Character vector of length 1. Tag in attribute field corresponding to *transcript* biotype.

gene\_biotype\_col

Character vector of length 1. Tag in attribute field corresponding to *gene* biotype.

transcript\_biotype\_use

Character, can be "all" or a vector of *transcript* biotypes to be used. Transcript biotypes aren't entirely the same as gene biotypes. For instance, in Ensembl annotation, retained\_intron is a transcript biotype, but not a gene biotype. If "cellranger", then a warning will be given. See data("ensembl\_tx\_biotypes") for all available transcript biotypes from Ensembl.

gene\_biotype\_use

Character, can be "all", "cellranger", or a vector of *gene* biotypes to be used. If "cellranger", then the biotypes used by Cell Ranger's reference are used. See data("cellranger\_biotypes") for gene biotypes the Cell Ranger reference uses. See data("ensembl\_gene\_biotypes") for all available gene biotypes from Ensembl. Note that gene biotypes and transcript biotypes are not always the same.

chrs\_only

Logical, whether to include chromosomes only, for GTF and GFF files can contain annotations for scaffolds, which are not incorporated into chromosomes. This will also exclude haplotypes. Defaults to TRUE. Only applicable to species found in genomeStyles().

 $compress\_fa$ 

Logical, whether to compress the output fasta file. If TRUE, then the fasta file will be gzipped.

save\_filtered\_gtf

Logical. If filtering type, biotypes, and/or chromosomes, whether to save the filtered GRanges as a GTF file.

overwrite

Logical, whether to overwrite if files with names of outputs written to disk already exist.

## Value

A data frame at least 2 columns: gene for gene ID, transcript for transcript ID, and optionally, gene\_name for gene names.

 $tr2g\_gtf$  43

#### **Description**

This function reads a GTF file and extracts the transcript ID and corresponding gene ID. This function assumes that the GTF file is properly formatted. See <a href="http://mblab.wustl.edu/GTF2">http://mblab.wustl.edu/GTF2</a>. <a href="http://mblab.wustl.edu/GTF2">httml</a> for a detailed description of proper GTF format. Note that GFF3 files have a somewhat different and more complicated format in the attribute field, which this function does not support. See <a href="http://gmod.org/wiki/GFF3">http://gmod.org/wiki/GFF3</a> for a detailed description of proper GFF3 format. To extract transcript and gene information from GFF3 files, see the function <a href="https://creativecommons.org/representation-r

## Usage

```
tr2g_gtf(
  file,
 Genome = NULL,
  get_transcriptome = TRUE,
  out_path = ".",
  write_tr2g = TRUE,
  transcript_id = "transcript_id",
  gene_id = "gene_id",
  gene_name = "gene_name",
  transcript_version = "transcript_version",
  gene_version = "gene_version",
  version_sep = ".",
  transcript_biotype_col = "transcript_biotype",
  gene_biotype_col = "gene_biotype",
  transcript_biotype_use = "all",
  gene_biotype_use = "all",
  chrs_only = TRUE,
  compress_fa = FALSE,
  save_filtered_gtf = TRUE,
  overwrite = FALSE
)
```

## **Arguments**

file

Path to a GTF file to be read. The file can remain gzipped. Use getGTF from the biomartr package to download GTF files from Ensembl, and use getGFF from biomartr to download GFF3 files from Ensembl and RefSeq.

Genome

Either a BSgenome or a XStringSet object of genomic sequences, where the intronic sequences will be extracted from. Use genomeStyles to check which styles are supported for your organism of interest; supported styles can be interconverted. If the style in your genome or annotation is not supported, then the style of chromosome names in the genome and annotation should be manually set to be consistent.

get\_transcriptome

Logical, whether to extract transcriptome from genome with the GTF file. If filtering biotypes or chromosomes, the filtered GRanges will be used to extract transcriptome.

out\_path

Directory to save the outputs written to disk. If this directory does not exist, then it will be created. Defaults to the current working directory.

write\_tr2g

Logical, whether to write tr2g to disk. If TRUE, then a file tr2g.tsv will be written into out\_path.

44 tr2g\_gtf

transcript\_id Character vector of length 1. Tag in attribute field corresponding to transcript IDs. This argument must be supplied and cannot be NA or NULL. Will throw error

if tag indicated in this argument does not exist.

Character vector of length 1. Tag in attribute field corresponding to gene IDs. This argument must be supplied and cannot be NA or NULL. Note that this is different from gene symbols, which do not have to be unique. This can be Ensembl or Entrez IDs. However, if the gene symbols are in fact unique for each gene, you may supply the tag for human readable gene symbols to this argument. Will throw error if tag indicated in this argument does not exist. This

is typically "gene\_id" for annotations from Ensembl and "gene" for refseq.

Character vector of length 1. Tag in attribute field corresponding to gene gene\_name symbols. This argument can be NA or NULL if you are fine with non-human readable gene IDs and do not wish to extract human readable gene symbols.

Character vector of length 1. Tag in attribute field corresponding to transcript version number. If your GTF file does not include transcript version numbers, or if you do not wish to include the version number, then use NULL for this argument. To decide whether to include transcript version number, check whether

> directory. If that file includes version numbers, then transscript version numbers must be included here as well. If that file does not include version numbers, then

> version numbers are included in the transcripts.txt in the kallisto output

transcript version numbers must not be included here.

Character vector of length 1. Tag in attribute field corresponding to gene version number. If your GTF file does not include gene version numbers, or if you do not wish to include the version number, then use NULL for this argument. Unlike transcript version number, it's up to you whether to include gene version number.

version\_sep Character to separate bewteen the main ID and the version number. Defaults to ".", as in Ensembl.

transcript\_biotype\_col

Character vector of length 1. Tag in attribute field corresponding to transcript biotype.

gene\_biotype\_col

Character vector of length 1. Tag in attribute field corresponding to gene biotype.

transcript\_biotype\_use

Character, can be "all" or a vector of transcript biotypes to be used. Transcript biotypes aren't entirely the same as gene biotypes. For instance, in Ensembl annotation, retained\_intron is a transcript biotype, but not a gene biotype. If "cellranger", then a warning will be given. See data("ensembl\_tx\_biotypes") for all available transcript biotypes from Ensembl.

Character, can be "all", "cellranger", or a vector of gene biotypes to be used. If "cellranger", then the biotypes used by Cell Ranger's reference are used. See data("cellranger\_biotypes") for gene biotypes the Cell Ranger reference uses. See data("ensembl\_gene\_biotypes") for all available gene biotypes from Ensembl. Note that gene biotypes and transcript biotypes are not always the same.

Logical, whether to include chromosomes only, for GTF and GFF files can contain annotations for scaffolds, which are not incorporated into chromosomes.

gene\_id

transcript\_version

gene\_version

gene\_biotype\_use

chrs\_only

 $tr2g\_gtf$  45

This will also exclude haplotypes. Defaults to TRUE. Only applicable to species found in genomeStyles().

compress\_fa Logical, whether to compress the output fasta file. If TRUE, then the fasta file will be gzipped.

save\_filtered\_gtf

Logical. If filtering type, biotypes, and/or chromosomes, whether to save the filtered GRanges as a GTF file.

overwrite Logical, whether to overwrite if files with names of outputs written to disk already exist.

#### **Details**

Transcript and gene versions may not be present in all GTF files, so these arguments are optional. This function has arguments for transcript and gene version numbers because Ensembl IDs have version numbers. For Ensembl IDs, we recommend including the version number, since a change in version number signals a change in the entity referred to by the ID after reannotation. If a version is used, then it will be appended to the ID, separated by version\_sep.

The transcript and gene IDs are The attribute field (the last field) of GTF files can be complicated and inconsistent across different sources. Please check the attribute tags in your GTF file and consider the arguments of this function carefully. The defaults are set according to Ensembl GTF files; defaults may not work for files from other sources. Due to the general lack of standards for the attribute field, you may need to further clean up the output of this function.

## Value

A data frame at least 2 columns: gene for gene ID, transcript for transcript ID, and optionally, gene\_name for gene names.

## See Also

```
ensembl_gene_biotypes ensembl_tx_biotypes cellranger_biotypes

Other functions to retrieve transcript and gene info: sort_tr2g(), tr2g_EnsDb(), tr2g_ensembl(), tr2g_fasta(), tr2g_gff3(), transcript2gene()
```

## **Examples**

```
toy_path <- system.file("testdata", package = "BUSpaRse")
file_use <- paste(toy_path, "gtf_test.gtf", sep = "/")
# Default
tr2g <- tr2g_gtf(file = file_use, get_transcriptome = FALSE,
    write_tr2g = FALSE, save_filtered_gtf = FALSE)
# Excluding version numbers
tr2g <- tr2g_gtf(file = file_use, transcript_version = NULL,
    gene_version = NULL, get_transcriptome = FALSE,
    write_tr2g = FALSE, save_filtered_gtf = FALSE)</pre>
```

 $46 tr2g_TxDb$ 

tr2g\_junction

tr2g for exon-exon junctions

## **Description**

```
tr2g for exon-exon junctions
```

## Usage

```
tr2g_junction(tr2g_cdna, junction_names)
```

## **Arguments**

```
tr2g_cdna The original tr2g_cdna.
junction_names Names of junctions internally generated.
```

## Value

A tr2g data frame where "transcripts" are the exon-exon junctions and genes are the corresponding genes.

tr2g\_TxDb

Get transcript and gene info from TxDb objects

## Description

The genome and gene annotations of some species can be conveniently obtained from Bioconductor packages. This is more convenient than downloading GTF files from Ensembl and reading it into R. In these packages, the gene annotation is stored in a TxDb object, which has standardized names for gene IDs, transcript IDs, exon IDs, and so on, which are stored in the metadata fields in GTF and GFF3 files, which are not standardized. This function extracts transcript and corresponding gene information from gene annotation stored in a TxDb object.

```
tr2g_TxDb(
   txdb,
   Genome = NULL,
   get_transcriptome = TRUE,
   out_path = ".",
   write_tr2g = TRUE,
   chrs_only = TRUE,
   compress_fa = FALSE,
   overwrite = FALSE
)
```

 $tr2g\_TxDb$  47

## **Arguments**

txdb A TxDb object with gene annotation.

Genome Either a BSgenome or a XStringSet object of genomic sequences, where the

intronic sequences will be extracted from. Use <code>genomeStyles</code> to check which styles are supported for your organism of interest; supported styles can be interconverted. If the style in your genome or annotation is not supported, then the style of chromosome names in the genome and annotation should be manually

set to be consistent.

get\_transcriptome

Logical, whether to extract transcriptome from genome with the GTF file. If filtering biotypes or chromosomes, the filtered GRanges will be used to extract

transcriptome.

out\_path Directory to save the outputs written to disk. If this directory does not exist, then

it will be created. Defaults to the current working directory.

write\_tr2g Logical, whether to write tr2g to disk. If TRUE, then a file tr2g.tsv will be

written into out\_path.

chrs\_only Logical, whether to include chromosomes only, for GTF and GFF files can con-

tain annotations for scaffolds, which are not incorporated into chromosomes. This will also exclude haplotypes. Defaults to TRUE. Only applicable to species

found in genomeStyles().

compress\_fa Logical, whether to compress the output fasta file. If TRUE, then the fasta file

will be gzipped.

overwrite Logical, whether to overwrite if files with names of outputs written to disk al-

ready exist.

## Value

A data frame with 3 columns: gene for gene ID, transcript for transcript ID, and tx\_id for internal transcript IDs used to avoid duplicate transcript names. For TxDb packages from Bioconductor, gene ID is Entrez ID, while transcript IDs are Ensembl IDs with version numbers for TxDb.Hsapiens.UCSC.hg38.knownGene. In some cases, the transcript ID have duplicates, and this is resolved by adding numbers to make the IDs unique.

A data frame with 3 columns: gene for gene ID, transcript for transcript ID, and gene\_name for gene names. If other\_attrs has been specified, then those will also be columns in the data frame returned.

## See Also

```
Other functions to retrieve transcript and gene info: sort_tr2g(), tr2g_EnsDb(), tr2g_ensembl(), tr2g_fasta(), tr2g_gff3(), tr2g_gtf(), transcript2gene()

Other functions to retrieve transcript and gene info: sort_tr2g(), tr2g_EnsDb(), tr2g_ensembl(),
```

## **Examples**

```
library(TxDb.Hsapiens.UCSC.hg38.knownGene)
library(BSgenome.Hsapiens.UCSC.hg38)
tr2g_TxDb(TxDb.Hsapiens.UCSC.hg38.knownGene, BSgenome.Hsapiens.UCSC.hg38)
# Clean up
file.remove("transcriptome.fa", "tr2g.tsv")
```

tr2g\_fasta(), tr2g\_gff3(), tr2g\_gtf(), transcript2gene()

48 transcript2gene

transcript2gene

Map Ensembl transcript ID to gene ID

# Description

This function is a shortcut to get the correctly sorted data frame with transcript IDs and the corresponding gene IDs from Ensembl biomart or Ensembl transcriptome FASTA files. For biomart query, it calls tr2g\_ensembl and then sort\_tr2g. For FASTA files, it calls tr2g\_fasta and then sort\_tr2g. Unlike in tr2g\_ensembl and tr2g\_fasta, multiple species can be supplied if cells from different species were sequenced together. This function should only be used if the kallisto inidex was built with transcriptomes from Ensembl. Also, if querying biomart, please make sure to set ensembl\_version to match the version where the transcriptomes were downloaded.

#### Usage

```
transcript2gene(
  species,
  fasta_file,
  kallisto_out_path,
  type = "vertebrate",
   ...
)
```

#### **Arguments**

species A character vector of Latin names of species present in this scRNA-seq dataset.

This is used to retrieve Ensembl information from biomart.

fasta\_file Character vector of paths to the transcriptome FASTA files used to build the

kallisto index. Exactly one of species and fasta\_file can be missing.

kallisto\_out\_path

Path to the kallisto bus output directory.

type A character vector indicating the type of each species. Each element must be

one of "vertebrate", "metazoa", "plant", "fungus", and "protist". If length is 1, then this type will be used for all species specified here. Can be missing if

fasta\_file is specified.

... Other arguments passed to tr2g\_ensembl such as other\_attrs, ensembl\_version,

and arguments passed to useMart. If fasta\_files is supplied instead of species, then this will be extra argumennts to tr2g\_fasta, such as use\_transcript\_version

and use\_gene\_version.

## Value

A data frame with two columns: gene and transcript, with Ensembl gene and transcript IDs (with version number), in the same order as in the transcriptome index used in kallisto.

## Note

This function has been superseded by the new version of tr2g\_\* functions that can extract transcriptome for only the biotypes specified and with only the standard chromosomes. The new version of tr2g\_\* functions also sorts the transcriptome so the tr2g and the transcriptome have transcripts in the same order.

#### See Also

```
Other functions to retrieve transcript and gene info: sort_tr2g(), tr2g_EnsDb(), tr2g_TxDb(), tr2g_ensembl(), tr2g_fasta(), tr2g_gff3(), tr2g_gff()
```

## **Examples**

```
# Download dataset already in BUS format
library(TENxBUSData)
TENxBUSData(".", dataset = "hgmm100")
tr2g <- transcript2gene(c("Homo sapiens", "Mus musculus"),
   type = "vertebrate", save_filtered = FALSE,
   ensembl_version = 99, kallisto_out_path = "./out_hgmm100")
# Clean up files from the example
unlink("out_hgmm100")</pre>
```

validate\_velocity\_input

Validate input to get\_velocity\_files

## **Description**

Validate input to get\_velocity\_files

#### Usage

```
validate_velocity_input(
   L,
   Genome,
   Transcriptome,
   out_path,
   compress_fa,
   width,
   exon_option
)
```

## Arguments

L

Length of the biological read. For instance, 10xv1: 98 nt, 10xv2: 98 nt, 10xv3: 91 nt, Drop-seq: 50 nt. If in doubt check read length in a fastq file for biological reads with the bash commands: If the fastq file is gzipped, then do zcat your\_file.fastq.gz | head on Linux. If on Mac, then zcat < your\_file.fastq.gz | head. Then you will see lines with nucleotide bases. Copy one of those lines and determine its length with str\_length in R or echo -n < the sequence > | wc -c in bash. Which file corresponds to biological reads depends on the particular technology.

Genome

Either a BSgenome or a XStringSet object of genomic sequences, where the intronic sequences will be extracted from. Use genomeStyles to check which styles are supported for your organism of interest; supported styles can be interconverted. If the style in your genome or annotation is not supported, then the style of chromosome names in the genome and annotation should be manually set to be consistent.

Transcriptome	A XStringSet, a path to a fasta file (can be gzipped) of the transcriptome which contains sequences of spliced transcripts, or NULL. The transcriptome here will be concatenated with the intronic sequences to give one fasta file. When NULL, the transcriptome sequences will be extracted from the genome given the gene annotation, so it will be guaranteed that transcript IDs in the transcriptome and in the annotation match. Otherwise, the type of transcript ID in the transcriptome must match that in the gene annotation supplied via argument X.
out_path	Directory to save the outputs written to disk. If this directory does not exist, then it will be created. Defaults to the current working directory.
compress_fa	Logical, whether to compress the output fasta file. If TRUE, then the fasta file will be gzipped.
width	Maximum number of letters per line of sequence in the output fasta file. Must be an integer.
exon_option	Character, indicating how exonic sequences should be included in the kallisto index. Must be one of the following:
	<b>full</b> The full cDNA sequences, which include the full exonic sequences, will be used. This is the default.

## Value

50

Will throw error if validation fails. Returns a named list whose first element is the normalized path to output directory, and whose second element is the normalized path to the transcriptome file if specified.

junction Only the exon-exon junctions, with L-1 bases on each side of the junc-

write\_velocity\_output Write the files for RNA velocity to disk

tions, will be used.

# Description

Write the files for RNA velocity to disk, in the specified output directory.

```
write_velocity_output(
  out_path,
  introns,
  Genome,
  Transcriptome,
  isoform_action,
  exon_option,
  tr2g_cdna,
  compress_fa,
  width
)
```

## **Arguments**

out\_path Directory to save the outputs written to disk. If this directory does not exist, then

it will be created.

introns Intronic ranges plus flanking region, returned by get\_intron\_flanks.

Genome Either a BSgenome or a XStringSet object of genomic sequences, where the

intronic sequences will be extracted from. Use <code>genomeStyles</code> to check which styles are supported for your organism of interest; supported styles can be interconverted. If the style in your genome or annotation is not supported, then the style of chromosome names in the genome and annotation should be manually

set to be consistent.

Transcriptome A XStringSet, a path to a fasta file (can be gzipped) of the transcriptome which

contains sequences of spliced transcripts, or NULL. The transcriptome here will be concatenated with the intronic sequences to give one fasta file. When NULL, the transriptome sequences will be extracted from the genome given the gene annotation, so it will be guaranteed that transcript IDs in the transcriptome and in the annotation match. Otherwise, the type of transcript ID in the transcriptome

must match that in the gene annotation supplied via argument X.

isoform\_action Character, indicating action to take with different transcripts of the same gene.

Must be one of the following:

**collapse** First, the union of all exons of different transcripts of a gene will be taken. Then the introns will be inferred from this union. Only the flanked intronic sequences are affected; isoforms will always be taken into account for spliced sequences or exon-exon junctions.

separate Introns from different transcripts will be kept separate.

exon\_option Character, indicating how exonic sequences should be included in the kallisto

index. Must be one of the following:

**full** The full cDNA sequences, which include the full exonic sequences, will be used. This is the default.

**junction** Only the exon-exon junctions, with L-1 bases on each side of the junctions, will be used.

tr2g\_cdna A data frame with columns transcript and gene that maps transcripts to genes

for spliced transcripts.

compress\_fa Logical, whether to compress the output fasta file. If TRUE, then the fasta file

will be gzipped.

width Maximum number of letters per line of sequence in the output fasta file. Must

be an integer.

## Value

Nothing into the R session. The files are written to disk.

# Index

* datasets	genomeStyles, 4, 7, 8, 19, 24, 29, 31, 38, 41,	
cellranger_biotypes, 7	43, 47, 49, 51	
<pre>ensembl_gene_biotypes, 13</pre>	get_inflection, 16	
<pre>ensembl_gff_mcols, 14</pre>	<pre>get_inflection(get_knee_df), 16</pre>	
<pre>ensembl_gtf_mcols, 14</pre>	get_intron_flanks, 15, 51	
<pre>ensembl_tx_biotypes, 15</pre>	get_knee_df, <i>16</i> , 16	
<pre>refseq_gff_mcols, 26</pre>	get_velocity_files, 6, 17	
* functions to retrieve transcript and gene	<pre>get_velocity_files,character-method</pre>	
info	(get_velocity_files), 17	
sort_tr2g, 27	<pre>get_velocity_files,EnsDb-method</pre>	
tr2g_EnsDb, 30	(get_velocity_files), 17	
tr2g_ensembl, 33	<pre>get_velocity_files,GRanges-method</pre>	
tr2g_fasta, 35	(get_velocity_files), 17	
tr2g_gff3, 37	<pre>get_velocity_files,TxDb-method</pre>	
tr2g_gtf, 42	(get_velocity_files), 17	
tr2g_TxDb,46	GRanges, 41	
transcript2gene, 48	1 ( ) 16	
<pre>.get_velocity_files, 3</pre>	knee_plot(get_knee_df), 16	
	listAttributes, 33	
annot_circular, 7	listEnsemblArchives, 11, 34	
annots_from_fa_df, 6	113 CENSEMBLAI CHIVES, 11, 34	
annots_from_fa_GRanges	<pre>make_sparse_matrix, 22</pre>	
(annots_from_fa_df), 6	match_style, 24	
	<b>-</b> ,	
BSgenome, 4, 7, 8, 19, 24, 29, 31, 38, 41, 43,	read_count_output, 24	
47, 49, 51	<pre>read_velocity_output, 25</pre>	
	refseq_gff_mcols, 26	
cellranger_biotypes, 7		
check_char1, 8	save_tr2g_bustools, 26, 27	
check_genome, 8	seqlevels, 7, 24	
check_gff, 9	sort_tr2g, 26, 27, 32, 35, 37, 40, 45, 47–49	
check_tag_present, 9	species2dataset, 28	
check_tx, 10	standardize_tags, 29	
columns, 31	str_length, 4, 19, 49	
	sub_annot, 30	
dl_transcriptome, 10	subset_annot, 7, 29	
_	subset_annot,BSgenome-method	
EC2gene, 12, 23, 37	(subset_annot), 29	
EnsDb, 19	subset_annot, DNAStringSet-method	
ensembl_gene_biotypes, 13	(subset_annot), 29	
ensembl_gff_mcols, 14	tr2g_EnsDb, 28, 30, 35, 37, 40, 45, 47, 49	
ensembl_gtf_mcols, 14	tr2g_ensembl, 28, 32, 33, 37, 40, 45, 47–49	
ensembl_tx_biotypes, 15	tr2g_fasta, 28, 32, 35, 35, 40, 45, 47–49	

INDEX 53

```
tr2g_gff3, 27, 28, 32, 35, 37, 37, 43, 45, 47, 49

tr2g_GRanges, 40

tr2g_gtf, 27, 28, 32, 35, 37, 40, 42, 47, 49

tr2g_junction, 46

tr2g_TxDb, 28, 32, 35, 37, 40, 45, 46, 49

transcript2gene, 13, 28, 32, 35, 37, 40, 45, 47, 48

TxDb, 19, 46, 47

useMart, 34, 48

validate_velocity_input, 49

write.table, 26

write_velocity_output, 50

XStringSet, 4, 7, 8, 19, 24, 29, 31, 38, 41, 43, 47, 49–51
```