

Package ‘txcutr’

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Title Transcriptome CUTteR

Description Various mRNA sequencing library preparation methods generate sequencing reads specifically from the transcript ends. Analyses that focus on quantification of isoform usage from such data can be aided by using truncated versions of transcriptome annotations, both at the alignment or pseudo-alignment stage, as well as in downstream analysis. This package implements some convenience methods for readily generating such truncated annotations and their corresponding sequences.

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Encoding UTF-8

Depends R (>= 4.1.0)

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biocViews Alignment, Annotation, RNASeq, Sequencing, Transcriptomics

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<code>.clipTranscript</code>	<i>Clip Transcript to Given Length</i>
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Description

Internal function for operating on individual GRanges, where ranges represent exons in a transcript. This is designed to be used in an `*apply` function over a GRangesList object.

Usage

```
.clipTranscript(gr, maxTxLength)
```

Arguments

<code>gr</code>	a GRanges object
<code>maxTxLength</code>	a positive integer

Value

the clipped GRanges object

.fillReduce *Convert GRanges to Single Range*

Description

Convert GRanges to Single Range

Usage

```
.fillReduce(gr, validate = TRUE)
```

Arguments

gr	a GRanges with ranges to be merged.
validate	logical determining whether entries should be checked for compatible seqnames and strands.

Details

The validation assumes seqnames and strand are Rle objects.

Value

GRanges with single interval

.mutateEach, CompressedGRangesList-method
Efficient Metadata Columns Mutation

Description

Efficient Metadata Columns Mutation

Usage

```
## S4 method for signature 'CompressedGRangesList'  
.mutateEach(gr1, ...)
```

Arguments

gr1	a CompressedGRangesList
...	named list of vectors to insert as metadata columns on each element GRanges. Each vector length must match the length of the GRangesList.

Value

a CompressedGRangesList with all element GRanges updated with supplied metadata columns

<code>.propagateMap</code>	<i>Propagate Transcript Merge Map</i>
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Description

Propagate Transcript Merge Map

Usage

```
.propagateMap(df, MAXITERS = 1000)
```

Arguments

<code>df</code>	a <code>data.frame</code> with columns <code>tx_in</code> and <code>tx_out</code>
<code>MAXITERS</code>	a numeric controlling the maximum number of iterations

Value

a converged `data.frame`, such that, `tx_out` is not present in any `tx_in`

<code>exportFASTA</code>	<i>Export Transcriptome as FASTA</i>
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Description

Export Transcriptome as FASTA

Usage

```
exportFASTA(txdb, genome, file, ...)
```

Arguments

<code>txdb</code>	a <code>TxDb</code> object representing a transcriptome annotation
<code>genome</code>	a <code>BSgenome</code> object from which to extract sequences
<code>file</code>	a string for output FASTA file. File names ending in <code>".gz"</code> will automatically use gzip compression.
<code>...</code>	additional arguments to pass through to writeXStringSet

Value

The `txdb` argument is invisibly returned.

Examples

```
library(TxDb.Scerevisiae.UCSC.sacCer3.sgdGene)
library(BSgenome.Scerevisiae.UCSC.sacCer3)

## load annotation and genome
txdb <- TxDb.Scerevisiae.UCSC.sacCer3.sgdGene
sacCer3 <- BSgenome.Scerevisiae.UCSC.sacCer3

## restrict to 'chrI' transcripts (makes for briefer example runtime)
seqlevels(txdb) <- c("chrI")

## last 500 nts per tx
txdb_w500 <- truncateTxome(txdb)

## export uncompressed
outfile <- tempfile("sacCer3.sgdGene.w500", fileext=".fa")
exportFASTA(txdb_w500, sacCer3, outfile)

## export compressed
outfile <- tempfile("sacCer3.sgdGene.w500", fileext=".fa.gz")
exportFASTA(txdb_w500, sacCer3, outfile)
```

exportGTF

Export GTF

Description

Exports a TxDb annotation to a GTF file

Usage

```
exportGTF(txdb, file, source = "txcutr")
```

Arguments

txdb	transcriptome to be output
file	a string or connection to output GTF file. Automatically recognizes strings ending with ".gz" for zipped output.
source	a string to go in the source column

Value

The txdb argument is invisibly returned.

Examples

```
library(Txdb.Scerevisiae.UCSC.sacCer3.sgdGene)

## load annotation
txdb <- Txdb.Scerevisiae.UCSC.sacCer3.sgdGene

## restrict to 'chrI' transcripts
seqlevels(txdb) <- c("chrI")

## last 500 nts per tx
txdb_w500 <- truncateTxome(txdb)

## export uncompressed
outfile <- tempfile("sacCer3.sgdGene.w500", fileext=".gtf")
exportGTF(txdb_w500, outfile)

## export compressed
outfile <- tempfile("sacCer3.sgdGene.w500", fileext=".gtf.gz")
exportGTF(txdb_w500, outfile)
```

exportMergeTable	<i>Export Merge Table for Transcriptome</i>
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Description

Export Merge Table for Transcriptome

Usage

```
exportMergeTable(txdb, file, minDistance = 200L)
```

Arguments

txdb	a TxDb object representing a transcriptome annotation
file	a string or connection to output TSV file. Automatically recognizes strings ending with ".gz" for zipped output.
minDistance	the minimum separation to regard overlapping transcripts as unique.

Value

The txdb argument is invisibly returned.

Examples

```
library(TxDB.Scerevisiae.UCSC.sacCer3.sgdGene)

## load annotation
txdb <- TxDb.Scerevisiae.UCSC.sacCer3.sgdGene

## restrict to 'chrI' transcripts (makes for briefer example runtime)
seqlevels(txdb) <- c("chrI")

## last 500 nts per tx
txdb_w500 <- truncateTxome(txdb)

## export plain format
outfile <- tempfile("sacCer3.sgdGene.w500", fileext=".tsv")
exportMergeTable(txdb_w500, outfile)

## export compressed format
outfile <- tempfile("sacCer3.sgdGene.w500", fileext=".tsv.gz")
exportMergeTable(txdb_w500, outfile)
```

generateMergeTable	<i>Generate Merge Table</i>
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Description

Generate Merge Table

Usage

```
generateMergeTable(txdb, minDistance = 200)

## S4 method for signature 'TxDb'
generateMergeTable(txdb, minDistance = 200L)
```

Arguments

txdb	an object representing a transcriptome
minDistance	the minimum separation to regard overlapping transcripts as unique

Value

a data.frame with three columns - tx_in the input transcript - tx_out the transcript merged into - gene_out the gene merged into

a data.frame with three columns - tx_in the input transcript - tx_out the transcript merged into - gene_out the gene merged into

Examples

```

library(TxDb.Scerevisiae.UCSC.sacCer3.sgdGene)

## load annotation
txdb <- TxDb.Scerevisiae.UCSC.sacCer3.sgdGene

## restrict to 'chrI' transcripts
seqlevels(txdb) <- c("chrI")

## last 500 nts per tx
txdb_w500 <- truncateTxome(txdb)
txdb_w500

## last 100 nts per tx
txdb_w100 <- truncateTxome(txdb, maxTxLength=100)
txdb_w100

```

truncateTxome

Truncate Transcriptome

Description

Truncate Transcriptome

Usage

```

truncateTxome(txdb, maxTxLength = 500, ...)

## S4 method for signature 'TxDb'
truncateTxome(txdb, maxTxLength = 500, BPPARAM = bpparam())

```

Arguments

txdb	a TxDb object
maxTxLength	the maximum length of transcripts
...	additional arguments
BPPARAM	A BiocParallelParam object specifying whether and how the method should be parallelized.

Value

a TxDb object
a TxDb object

Examples

```
library(TxDb.Scerevisiae.UCSC.sacCer3.sgdGene)

## load annotation
txdb <- TxDb.Scerevisiae.UCSC.sacCer3.sgdGene

## restrict to 'chrI' transcripts
seqlevels(txdb) <- c("chrI")

## last 500 nts per tx
txdb_w500 <- truncateTxome(txdb)
txdb_w500

## last 100 nts per tx
txdb_w100 <- truncateTxome(txdb, maxTxLength=100)
txdb_w100
```

txdbToGRangesList *Convert TxDb object to GRangesList*

Description

Convert TxDb object to GRangesList

Usage

```
txdbToGRangesList(
  txdb,
  geneCols = c("gene_id"),
  transcriptCols = c("gene_id", "tx_name"),
  exonCols = c("gene_id", "tx_name", "exon_id", "exon_rank")
)
```

Arguments

txdb a TxDb object

geneCols names of columns to include in the genes ranges

transcriptCols names of columns to include in the transcripts ranges

exonCols names of columns to include in the exons ranges

Value

a GRangesList object with entries c(genes, transcripts, exons)

Examples

```
library(TxDb.Scerevisiae.UCSC.sacCer3.sgdGene)

## load annotation
txdb <- TxDb.Scerevisiae.UCSC.sacCer3.sgdGene

grl <- txdbToGRangesList(txdb)
grl
```

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