# Package 'ORFik'

February 21, 2025

Type Package

```
Title Open Reading Frames in Genomics
Version 1.27.3
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Description
     R package for analysis of transcript and translation features through manipulation of sequence data
     and NGS data like Ribo-Seq, RNA-Seq, TCP-
     Seg and CAGE. It is generalized in the sense that any transcript region
     can be analysed, as the name hints to it was made with investigation of ribosomal patterns over
     Open Reading Frames (ORFs) as it's primary use case.
     ORFik is extremely fast through use of C++, data.table and GenomicRanges.
     Package allows to reassign starts of the transcripts with the use of CAGE-Seq data,
     automatic shifting of RiboSeq reads, finding of Open Reading Frames for
     whole genomes and much more.
biocViews ImmunoOncology, Software, Sequencing, RiboSeq, RNASeq,
     FunctionalGenomics, Coverage, Alignment, DataImport
License MIT + file LICENSE
LazyData TRUE
BugReports https://github.com/Roleren/ORFik/issues
URL https://github.com/Roleren/ORFik
Depends R (>= 4.1.0), IRanges (>= 2.17.1), GenomicRanges (>= 1.35.1),
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```

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# Description

# Main goals:

- 1. Finding Open Reading Frames (very fast) in the genome of interest or on the set of transcripts/sequences.
- 2. Utilities for metaplots of RiboSeq coverage over gene START and STOP codons allowing to spot the shift.
- 3. Shifting functions for the RiboSeq data.
- 4. Finding new Transcription Start Sites with the use of CageSeq data.
- 5. Various measurements of gene identity e.g. FLOSS, coverage, ORFscore, entropy that are recreated based on many scientific publications.
- 6. Utility functions to extend GenomicRanges for faster grouping, splitting, tiling etc.

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#### See Also

Useful links:

- https://github.com/Roleren/ORFik
- Report bugs at https://github.com/Roleren/ORFik/issues

addCdsOnLeaderEnds

Extends leaders downstream

#### **Description**

When finding uORFs, often you want to allow them to end inside the cds.

# Usage

```
addCdsOnLeaderEnds(fiveUTRs, cds, onlyFirstExon = FALSE)
```

#### **Arguments**

fiveUTRs The 5' leader sequences as GRangesList

cds If you want to extend 5' leaders downstream, to catch uorfs going into cds,

include it.

onlyFirstExon logical (F), include whole cds or only first exons.

# **Details**

This is a simple way to do that

#### Value

a GRangesList of cds exons added to ends

#### See Also

Other uorfs: filterUORFs(), removeORFsWithSameStartAsCDS(), removeORFsWithSameStopAsCDS(), removeORFsWithStartInsideCDS(), removeORFsWithinCDS(), uORFSearchSpace()

addNewTSSOnLeaders Add cage max peaks as new transcript start sites for each 5' leader (\*) strands are not supported, since direction must be known.

# Description

Add cage max peaks as new transcript start sites for each 5' leader (\*) strands are not supported, since direction must be known.

#### Usage

```
addNewTSSOnLeaders(fiveUTRs, maxPeakPosition, removeUnused, cageMcol)
```

# **Arguments**

fiveUTRs (GRangesList) The 5' leaders or full transcript sequences

maxPeakPosition

The max peak for each 5' leader found by cage

removeUnused logical (FALSE), if False: (standard is to set them to original annotation), If

TRUE: remove leaders that did not have any cage support.

cageMcol a logical (FALSE), if TRUE, add a meta column to the returned object with the

raw CAGE counts in support for new TSS.

#### Value

a GRanges object of first exons

### **Description**

```
add_pseudo_5utrs_txdb_if_needed
```

# Usage

```
add_pseudo_5utrs_txdb_if_needed(
   txdb,
   pseudo_5UTRS_if_needed = NULL,
   minimum_5UTR_percentage = 30
)
```

#### **Arguments**

```
txdb a TxDb object pseudo_5UTRS_if_needed
```

integer, default NULL. If defined > 0, will add pseudo 5' UTRs of maximum this length if 'minimum\_5UTR\_percentage" (default 30 mRNAs (coding transcripts) do not have a leader. (NULL and 0 are both the ignore command)

minimum\_5UTR\_percentage

numeric, default 30. What minimum percentage of mRNAs most have a 5' UTRs (leaders), to not do the pseudo\_UTR addition. If percentage is higher, addition is ignored, set to 101 to always do it.

#### Value

txdb (new txdb if it was done, old if not)

a lignment Feature Statistics

Create alignment feature statistcs

#### **Description**

Among others how much reads are in mRNA, introns, intergenic, and check of reads from rRNA and other ncRNAs. The better the annotation / gtf used, the more results you get.

### Usage

```
alignmentFeatureStatistics(
   df,
   type = "ofst",
   force = TRUE,
   library.names = bamVarName(df),
   BPPARAM = bpparam()
)
```

# Arguments

df

an ORFik experiment

type

a character(default: "default"), load files in experiment or some precomputed variant, like "ofst" or "pshifted". These are made with ORFik:::convertLibs(), shiftFootprintsByExperiment(), etc. Can also be custom user made folders inside the experiments bam folder. It acts in a recursive manner with priority: If you state "pshifted", but it does not exist, it checks "ofst". If no .ofst files, it uses "default", which always must exists.

Presets are (folder is relative to default lib folder, some types fall back to other formats if folder does not exist):

- "default": load the original files for experiment, usually bam.

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- "ofst": loads ofst files from the ofst folder, relative to lib folder (falls back to default)

- "pshifted": loads ofst, wig or bigwig from pshifted folder (falls back to ofst, then default)
- "cov": Load covRle objects from cov\_RLE folder (fail if not found)
- "covl": Load covRleList objects, from cov\_RLE\_List folder (fail if not found)
- "bed": Load bed files, from bed folder (falls back to default)
- Other formats must be loaded directly with fimport

force

logical, default TRUE If TRUE, reload library files even if matching named variables are found in environment used by experiment (see envExp) A simple way to make sure correct libraries are always loaded. FALSE is faster if data is loaded correctly already.

library.names
BPPARAM

character vector, names of libraries, default: name\_decider(df, naming)

how many cores/threads to use? default: bpparam(). To see number of threads used, do bpparam()\$workers. You can also add a time remaining bar, for a

more detailed pipeline.

#### Value

a data.table of the statistcs

allFeaturesHelper

Calculate the features in computeFeatures function

#### **Description**

Not used directly, calculates all features internally for computeFeatures.

#### Usage

```
allFeaturesHelper(
  grl,
  RFP,
  RNA,
  tx,
  fiveUTRs,
  cds,
  threeUTRs,
  faFile,
  riboStart,
  riboStop,
  sequenceFeatures,
  uorfFeatures,
  grl.is.sorted,
 weight.RFP = 1L,
 weight.RNA = 1L,
  st = NULL
)
```

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# Arguments

grl	a GRangesList object with usually ORFs, but can also be either leaders, cds', 3' utrs, etc. This is the regions you want to score.
RFP	RiboSeq reads as GAlignments, GRanges or GRangesList object
RNA	RnaSeq reads as GAlignments, GRanges or GRangesList object
tx	a GRangesList of transcripts, normally called from: exonsBy(Gtf, by = "tx", use.names = $T$ ) only add this if you are not including Gtf file If you are using CAGE, you do not need to reassign these to the cage peaks, it will do it for you.
fiveUTRs	fiveUTRs as GRangesList, if you used cage-data to extend 5' utrs, remember to input CAGE assigned version and not original!
cds	a GRangesList of coding sequences
threeUTRs	a GRangesList of transcript 3' utrs, normally called from: threeUTRsByTranscript(Gtf, use.names = $T$ )
faFile	a path to fasta indexed genome, an open FaFile, a BSgenome, or path to ORFik experiment with valid genome.
riboStart	usually 26, the start of the floss interval, see ?floss
riboStop	usually 34, the end of the floss interval
sequenceFeature	es
	a logical, default TRUE, include all sequence features, that is: Kozak, fraction-Lengths, distORFCDS, isInFrame, isOverlapping and rankInTx. uorfFeatures = FALSE will remove the 4 last.
uorfFeatures	a logical, default TRUE, include all uORF sequence features, that is: distOR-FCDS, isInFrame, isOverlapping and rankInTx
grl.is.sorted	logical (F), a speed up if you know argument grl is sorted, set this to TRUE.
weight.RFP	a vector (default: 1L). Can also be character name of column in RFP. As in translationalEff(weight = "score") for: GRanges("chr1", 1, "+", score = 5), would mean score column tells that this alignment region was found 5 times.
weight.RNA	Same as weightRFP but for RNA weights. (default: 1L)
st	(NULL), if defined must be: st = startRegion(grl, tx, T, -3, 9)

# Value

a data.table with features

ppendZeroes Append zero values to data.table
--

# Description

For every position in width max.pos - min.pos + 1, append 0 values in data.table. Needed when coveragePerTiling was run on coverage window with drop.zero.dt as TRUE and you need to plot 0 positions after a transformation by coverageScorings.

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#### Usage

```
appendZeroes(dt, max.pos, min.pos = 1L, fractions = unique(dt$fraction))
```

#### **Arguments**

dt a data.table from coverageByTiling that is normalized by coverageScorings.

max.pos integer, max position of dt

min.pos integer, default 1L. Minimum position of dt

fractions default unique(dt\( \)fraction), will repeat each fraction max.pos - min.pos + 1

times.

#### Value

a data.table with appended 0 values

artificial.orfs

Create small artificial orfs from cds

# **Description**

Usefull to see if short ORFs prediction is dependent on length.

Split cds first in two, a start part and stop part. Then say how large the two parts can be and merge them together. It will sample a value in range give.

Parts will be forced to not overlap and can not extend outside original cds

#### Usage

```
artificial.orfs(
  cds,
  start5 = 1,
  end5 = 4,
  start3 = -4,
  end3 = 0,
  bin.if.few = TRUE
)
```

#### **Arguments**

```
a GRangesList of orfs, must have width %% 3 == 0 and length >= 6
start5 integer, default: 1 (start of orf)
end5 integer, default: 4 (max 4 codons from start codon)
start3 integer, default -4 (max 4 codons from stop codon)
end3 integer, default: 0 (end of orf)
```

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bin.if.few

logical, default TRUE, instead of per codon, do per 2, 3, 4 codons if you have few samples compared to lengths wanted, If you have 4 cds' and you want 7 different lengths, which is the standard, it will give you possible nt length: 6-12-18-24 instead of original 6-9-12-15-18-21-24.

If you have more than 30x cds than lengths wanted this is skipped. (for default arguments this is: 7\*30 = 210 cds)

#### **Details**

If artificial cds length is not divisible by 2, like 3 codons, the second codon will always be from the start region etc.

Also If there are many very short original cds, the distribution will be skewed towards more smaller artificial cds.

#### Value

GRangesList of new ORFs (sorted: + strand increasing start, - strand decreasing start)

#### **Examples**

```
txdb <- ORFik.template.experiment()
#cds <- loadRegion(txdb, "cds")
## To get enough CDSs, just replicate them
# cds <- rep(cds, 100)
#artificial.orfs(cds)</pre>
```

assignAnnotations

Overlaps GRanges object with provided annotations.

#### **Description**

It will return same list of GRanges, but with metdata columns: trainscript\_id - id of transcripts that overlap with each ORF gene\_id - id of gene that this transcript belongs to isoform - for coding protein alignment in relation to cds on coresponding transcript, for non-coding transcripts alignment in relation to the transcript.

#### Usage

```
assignAnnotations(ORFs, con)
```

# **Arguments**

ORFs - GRanges or GRangesList object of your ORFs.

con - Path to gtf file with annotations.

#### Value

A GRanges object of your ORFs with metadata columns 'gene', 'transcript', isoform' and 'biotype'.

assignFirstExonsStartSite

Reassign the start positions of the first exons per group in grl

#### Description

Per group in GRangesList, assign the most upstream site.

#### Usage

```
assignFirstExonsStartSite(
  grl,
  newStarts,
  is.circular = all(isCircular(grl) %in% TRUE)
)
```

#### **Arguments**

grl a GRangesList object

newStarts an integer vector of same length as grl, with new start values (absolute coordi-

nates, not relative)

is.circular logical, default FALSE if not any is: all(isCircular(grl) Where grl is the ranges

checked. If TRUE, allow ranges to extend below position 1 on chromosome.

Since circular genomes can have negative coordinates.

#### **Details**

make sure your grl is sorted, since start of "-" strand objects should be the max end in group, use ORFik:::sortPerGroup(grl) to get sorted grl.

#### Value

the same GRangesList with new start sites

#### See Also

Other GRanges: assignLastExonsStopSite(), downstreamFromPerGroup(), downstreamOfPerGroup(), upstreamFromPerGroup(), upstreamOfPerGroup()

assignLastExonsStopSite

Reassign the stop positions of the last exons per group

#### Description

Per group in GRangesList, assign the most downstream site.

#### Usage

```
assignLastExonsStopSite(
  grl,
  newStops,
  is.circular = all(isCircular(grl) %in% TRUE)
)
```

#### **Arguments**

grl a GRangesList object

newStops an integer vector of same length as grl, with new start values (absolute coordi-

nates, not relative)

is.circular logical, default FALSE if not any is: all(isCircular(grl) Where grl is the ranges

checked. If TRUE, allow ranges to extend below position 1 on chromosome.

Since circular genomes can have negative coordinates.

#### **Details**

make sure your grl is sorted, since stop of "-" strand objects should be the min start in group, use ORFik:::sortPerGroup(grl) to get sorted grl.

#### Value

the same GRangesList with new stop sites

#### See Also

```
Other GRanges: assignFirstExonsStartSite(), downstreamFromPerGroup(), downstreamOfPerGroup(), upstreamFromPerGroup(), upstreamOfPerGroup()
```

20 assignTSSByCage

assignTSSByCage	Input a txdb and add a 5' leader for each transcript, that does not have
	one.

# Description

For all cds in txdb, that does not have a 5' leader: Start at 1 base upstream of cds and use CAGE, to assign leader start. All these leaders will be 1 exon based, if you really want exon splicings, you can use exon prediction tools, or run sequencing experiments.

# Usage

```
assignTSSByCage(
  txdb,
  cage,
  extension = 1000,
  filterValue = 1,
  restrictUpstreamToTx = FALSE,
  removeUnused = FALSE,
  preCleanup = TRUE,
  pseudoLength = 1
)
```

# Arguments

txdb	a TxDb file, a path to one of: (.gtf ,.gff, .gff2, .gff2, .db or .sqlite) or an ORFik experiment	
cage	Either a filePath for the CageSeq file as .bed .bam or .wig, with possible compressions (".gzip", ".gz", ".bgz"), or already loaded CageSeq peak data as GRanges or GAlignment. NOTE: If it is a .bam file, it will add a score column by running: convertToOneBasedRanges(cage, method = "5prime", addScoreColumn = TRUE) The score column is then number of replicates of read, if score column is something else, like read length, set the score column to NULL first.	
extension	The maximum number of basses upstream of the TSS to search for CageSeq peak.	
filterValue	The minimum number of reads on cage position, for it to be counted as possible new tss. (represented in score column in CageSeq data) If you already filtered, set it to 0.	
restrictUpstreamToTx		
	a logical (FALSE). If TRUE: restrict leaders to not extend closer than 5 bases from closest upstream leader, set this to TRUE.	
removeUnused	logical (FALSE), if False: (standard is to set them to original annotation), If TRUE: remove leaders that did not have any cage support.	
preCleanup	logical (TRUE), if TRUE, remove all reads in region (-5:-1, 1:5) of all original tss in leaders. This is to keep original TSS if it is only +/- 5 bases from the original.	

asTX

pseudoLength

a numeric, default 1. Add a pseudo length for all the UTRs. Will not extend a leader if it would make it go outside the defined seqlengths of the chromosome (for non circular chromosomes), or extending closer than 50 nucleotides to upstream cds. So this length is not guaranteed for all!

#### **Details**

Given a TxDb object, reassign the start site per transcript using max peaks from CageSeq data. A max peak is defined as new TSS if it is within boundary of 5' leader range, specified by 'extension' in bp. A max peak must also be higher than minimum CageSeq peak cutoff specified in 'filter-Value'. The new TSS will then be the positioned where the cage read (with highest read count in the interval). If no CAGE supports a leader, the width will be set to 1 base.

#### Value

a TxDb obect of reassigned transcripts

#### See Also

```
Other CAGE: reassignTSSbyCage(), reassignTxDbByCage()
```

# **Examples**

```
txdbFile <- system.file("extdata", "hg19_knownGene_sample.sqlite",
    package = "GenomicFeatures")
cagePath <- system.file("extdata", "cage-seq-heart.bed.bgz",
    package = "ORFik")

## Not run:
    assignTSSByCage(txdbFile, cagePath)
    #Minimum 20 cage tags for new TSS
    assignTSSByCage(txdbFile, cagePath, filterValue = 20)
    # Create pseudo leaders for the ones without hits
    assignTSSByCage(txdbFile, cagePath, pseudoLength = 100)
    # Create only pseudo leaders (in example 2 leaders are added)
    assignTSSByCage(txdbFile, cage = NULL, pseudoLength = 100)

## End(Not run)</pre>
```

asTX

Map genomic to transcript coordinates by reference

#### **Description**

Map range coordinates between features in the genome and transcriptome (reference) space.

asTX

# Usage

```
asTX(
  grl,
  reference,
  ignore.strand = FALSE,
  x.is.sorted = TRUE,
  tx.is.sorted = TRUE
)
```

# **Arguments**

grl	a GRangesList of ranges within the reference, grl must either have names matching, or a meta column called 'names' that gives grouping names. i.e. grl named uORF_1_ENST00001, must then have a names meta column with ENST00001.
reference	a GRangesList of ranges that include grl as a subset of ranges. Example: cds is grl and mrna can be reference
ignore.strand	When ignore.strand is TRUE, strand is ignored in overlaps operations (i.e., all strands are considered "+") and the strand in the output is '*'.  When ignore.strand is FALSE (default) strand in the output is taken from the transcripts argument. When transcripts is a GRangesList, all inner list elements of a common list element must have the same strand or an error is thrown.  Mapped position is computed by counting from the transcription start site (TSS) and is not affected by the value of ignore.strand.
x.is.sorted	if x is a GRangesList object, are "-" strand groups pre-sorted in decreasing order within group, default: TRUE $$
tx.is.sorted	if transcripts is a GRangesList object, are "-" strand groups pre-sorted in decreasing order within group, default: TRUE

#### **Details**

Similar to GenomicFeatures' pmapToTranscripts, but in this version the grl ranges are compared to reference ranges with same name, not by index. This gives a large speedup, but also requires all objects must be named.

#### Value

a GRangesList in transcript coordinates

# See Also

```
Other ExtendGenomicRanges: coveragePerTiling(), extendLeaders(), extendTrailers(), reduceKeepAttr(), tile1(), txSeqsFromFa(), windowPerGroup()
```

# **Examples**

```
seqname <- c("tx1", "tx2", "tx3")
seqs <- c("ATGGGTATTTATA", "AAAAA", "ATGGGTAATA")
grIn1 <- GRanges(seqnames = "1",</pre>
```

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```
ranges = IRanges(start = c(21, 10), end = c(23, 19)),
                  strand = "-")
grIn2 <- GRanges(seqnames = "1",</pre>
                  ranges = IRanges(start = c(1), end = c(5)),
                  strand = "-")
grIn3 <- GRanges(seqnames = "1",</pre>
                  ranges = IRanges(start = c(1010), end = c(1019)),
                  strand = "-")
grl <- GRangesList(grIn1, grIn2, grIn3)</pre>
names(grl) <- seqname</pre>
# Find ORFs
test_ranges <- findMapORFs(grl, seqs,</pre>
                  "ATG|TGG|GGG",
                  "TAA|AAT|ATA",
                  longestORF = FALSE,
                  minimumLength = 0)
# Genomic coordinates ORFs
test_ranges
# Transcript coordinate ORFs
asTX(test_ranges, reference = grl)
# seqnames will here be index of transcript it came from
```

bamVarName

Get library variable names from ORFik experiment

# Description

What will each sample be called given the columns of the experiment? A column is included if more than 1 unique element value exist in that column.

#### Usage

```
bamVarName(
   df,
   skip.replicate = length(unique(df$rep)) == 1,
   skip.condition = length(unique(df$condition)) == 1,
   skip.stage = length(unique(df$stage)) == 1,
   skip.fraction = length(unique(df$fraction)) == 1,
   skip.experiment = !df@expInVarName,
   skip.libtype = FALSE,
   fraction_prepend_f = TRUE
)
```

# **Arguments**

```
df an ORFik experiment
skip.replicate a logical (FALSE), don't include replicate in variable name.
```

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```
skip.condition a logical (FALSE), don't include condition in variable name.

skip.stage a logical (FALSE), don't include stage in variable name.

skip.fraction a logical (FALSE), don't include fraction

skip.experiment a logical (FALSE), don't include experiment

skip.libtype a logical (FALSE), don't include libtype

fraction_prepend_f

a logical (TRUE), include "f" in front of fraction, useful for knowing what fraction is.
```

#### Value

variable names of libraries (character vector)

#### See Also

```
Other ORFik_experiment: ORFik.template.experiment(), ORFik.template.experiment.zf(), create.experiment(), experiment-class, filepath(), libraryTypes(), organism, experiment-method, outputLibs(), read.experiment(), save.experiment(), validateExperiments()
```

# **Examples**

```
df <- ORFik.template.experiment()
bamVarName(df)

## without libtype
bamVarName(df, skip.libtype = TRUE)
## Without experiment name
bamVarName(df, skip.experiment = TRUE)</pre>
```

bamVarNamePicker

Get variable name per filepath in experiment

# **Description**

Get variable name per filepath in experiment

#### Usage

```
bamVarNamePicker(
   df,
   skip.replicate = FALSE,
   skip.condition = FALSE,
   skip.stage = FALSE,
   skip.fraction = FALSE,
   skip.experiment = FALSE,
   skip.libtype = FALSE,
   fraction_prepend_f = TRUE
)
```

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#### **Arguments**

```
df an ORFik experiment

skip.replicate a logical (FALSE), don't include replicate in variable name.

skip.condition a logical (FALSE), don't include condition in variable name.

skip.stage a logical (FALSE), don't include stage in variable name.

skip.fraction a logical (FALSE), don't include fraction

skip.experiment a logical (FALSE), don't include experiment

skip.libtype a logical (FALSE), don't include libtype

fraction_prepend_f
 a logical (TRUE), include "f" in front of fraction, useful for knowing what fraction is.
```

# Value

variable name of library (character vector)

Get batch name variants
Get batch hame variants

#### **Description**

Used to standardize nomeclature for experiments. Example: Biological samples (batches) batch will become b1

#### Usage

```
batchNames()
```

#### Value

a data.table with 2 columns, the main name, and all name variants of the main name in second column as a list.

#### See Also

```
Other experiment_naming: cellLineNames(), cellTypeNames(), conditionNames(), fractionNames(), inhibitorNames(), libNames(), mainNames(), repNames(), stageNames(), tissueNames()
```

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bedToGR

Converts bed style data.frame to GRanges

# **Description**

For info on columns, see: https://www.ensembl.org/info/website/upload/bed.html

# Usage

```
bedToGR(x, skip.name = TRUE)
```

# **Arguments**

x A data. frame from imported bed-file, to convert to GRanges skip.name default (TRUE), skip name column (column 4)

#### Value

a GRanges object from bed

#### See Also

```
Other utils: convertToOneBasedRanges(), export.bed12(), export.bigWig(), export.fstwig(), export.wiggle(), fimport(), findFa(), fread.bed(), optimizeReads(), readBam(), readBigWig(), readWig()
```

browseSRA

Open SRA in browser for specific bioproject

#### **Description**

Open SRA in browser for specific bioproject

#### Usage

```
browseSRA(x, browser = getOption("browser"))
```

# **Arguments**

x character, bioproject ID.

browser a non-empty character string giving the name of the program to be used as the

HTML browser. It should be in the PATH, or a full path specified. Alternatively,

an R function to be called to invoke the browser.

Under Windows NULL is also allowed (and is the default), and implies that the

file association mechanism will be used.

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#### Value

invisible(NULL), opens webpage only

#### See Also

```
Other sra: download.SRA(), download.SRA.metadata(), download.ebi(), get_bioproject_candidates(), install.sratoolkit(), rename.SRA.files()
```

# **Examples**

```
#browseSRA("PRJNA336542")

#' # For windows make sure a valid browser is defined:
browser <- getOption("browser")
#browseSRA("PRJNA336542", browser)</pre>
```

cellLineNames

Get cell-line name variants

# **Description**

Used to standardize nomeclature for experiments.

Example: THP1 is main naming, but a variant is THP-1 THP-1 will then be renamed to THP1 (variables in R, can not have - in them)

# Usage

```
cellLineNames(convertToTissue = FALSE)
```

#### **Arguments**

convertToTissue

logical, FALSE. If TRUE, return tissue type. NONE is returned for general non-differentiated cell lines like 3T3.

#### Value

a data.table with 2 columns, the main name, and all name variants of the main name in second column as a list.

### See Also

```
Other experiment_naming: batchNames(), cellTypeNames(), conditionNames(), fractionNames(), inhibitorNames(), libNames(), mainNames(), repNames(), stageNames(), tissueNames()
```

changePointAnalysis

cellTypeNames

Get cell type name variants

# Description

Used to standardize nomeclature for experiments.

Example: 1 is main naming, but a variant is rep1 rep1 will then be renamed to 1

#### Usage

```
cellTypeNames()
```

#### Value

a data.table with 2 columns, the main name, and all name variants of the main name in second column as a list.

#### See Also

```
Other experiment_naming: batchNames(), cellLineNames(), conditionNames(), fractionNames(), inhibitorNames(), libNames(), mainNames(), repNames(), stageNames(), tissueNames()
```

changePointAnalysis

Get the offset for specific RiboSeq read width

# **Description**

Creates sliding windows of transcript normalized counts per position and check which window has most in upstream window vs downstream window. Pick the position with highest absolute value maximum of the window difference. Checks windows with split sites between positions -17 to -7, where 0 is TIS. Normally you expect the shift around -12 for Ribo-seq, in TCP-seq / RCP-seq it is usually a bit higher, usually because of cross-linking variations.

#### Usage

```
changePointAnalysis(
    X,
    feature = "start",
    max.pos = 40L,
    interval = seq.int(14L, 24L),
    center.pos = 12,
    info = NULL,
    verbose = FALSE
)
```

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# **Arguments**

X	a vector with count per position to analyse, assumes the zero position (TIS) is in the middle + 1 (position 0). Default it is size 60, from -30 to 29 in p-shifting
feature	(character) either "start" or "stop"
max.pos	integer, default 40L, subset x to go from index 1 to max.pos, if tail is not relevant.
interval	integer vector , default seq.int(14L, 24L). The possible shift locations, default Seperation points for upstream and downstream windows. That is $(+/-5)$ from -12) position.
center.pos	integer, default 12. Centering position for likely p-site. A first qualified guess to save time. 12 means 12 bases before TIS.
info	specify read length if wanted for verbose output.
verbose	logical, default FALSE. Report details of change point analysis.

# **Details**

For visual explanation, see the supl. data of ORFik paper: Transcript normalized means per CDS TIS region, count reads per position, divide that number per position by the total of that transcript, then sum up these numbers per position for all transcripts.

#### Value

a single numeric offset, -12 would mean p-site is 12 bases upstream

#### See Also

Other pshifting: detectRibosomeShifts(), shiftFootprints(), shiftFootprintsByExperiment(), shiftPlots(), shifts\_load(), shifts\_save()

checkRFP Helper Function to check valid RFP input

# **Description**

Helper Function to check valid RFP input

# Usage

checkRFP(class)

# Arguments

class the given class of RFP object

# Value

NULL, stop if invalid object

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#### See Also

```
Other validity: checkRNA(), is.ORF(), is.gr_or_grl(), is.grl(), is.range(), validGRL(), validSeqlevels()
```

checkRNA

Helper Function to check valid RNA input

# Description

Helper Function to check valid RNA input

#### Usage

```
checkRNA(class)
```

# **Arguments**

class

the given class of RNA object

### Value

NULL, stop if unvalid object

#### See Also

```
Other validity: checkRFP(), is.ORF(), is.gr_or_grl(), is.grl(), is.range(), validGRL(), validSeqlevels()
```

codonSumsPerGroup

Get read hits per codon

# Description

Helper for entropy function, normally not used directly Seperate each group into tuples (abstract codons) Gives sum for each tuple within each group

#### Usage

```
codonSumsPerGroup(grl, reads, weight = "score", is.sorted = FALSE)
```

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#### **Arguments**

grl a GRangesList of 5' utrs, CDS, transcripts, etc.

reads a GAlignments, GRanges, or precomputed coverage as covRle (one for each

strand) of RiboSeq, RnaSeq etc.

Weigths for scoring is default the 'score' column in 'reads'. Can also be random access paths to bigWig or fstwig file. Do not use random access for more than a few genes, then loading the entire files is usually better. File streaming is still in

beta, so use with care!

weight (default: 'score'), if defined a character name of valid meta column in subject.

GRanges("chr1", 1, "+", score = 5), would mean score column tells that this alignment region was found 5 times. Formats which loads a score column like this: Bigwig, wig, ORFik ofst, collapsed bam, bedoc and .bedo. As do CAGEr CAGE files and many other package formats. You can also assign a score col-

umn manually.

is.sorted logical (FALSE), is grl sorted. That is + strand groups in increasing ranges

(1,2,3), and - strand groups in decreasing ranges (3,2,1)

#### **Details**

Example: counts c(1,0,0,1), with reg\_len = 2, gives c(1,0) and c(0,1), these are summed and returned as data.table 10 bases, will give 3 codons, 1 base codons does not exist.

#### Value

a data.table with codon sums

codon\_usage

Codon usage

# Description

Per AA / codon, analyse the coverage, get a multitude of features. For both A sites and P-sites (Input reads must be P-sites for now) This function takes inspiration from the codonDT paper, and among others returns the negative binomial estimates, but in addition many other features.

#### Usage

```
codon_usage(
  reads,
  cds,
  mrna,
  faFile,
  filter_table,
  filter_cds_mod3 = TRUE,
  min_counts_cds_filter = max(min(quantile(filter_table, 0.5), 1000), 1000),
  with_A_sites = TRUE,
```

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```
aligned_position = "center",
code = GENETIC_CODE
)
```

#### **Arguments**

reads either a single library (GRanges, GAlignment, GAlignmentPairs), or a list of

libraries returned from outputLibs(df) with p-sites. If list, the list must have

names coresponding to the library names.

cds a GRangesList mrna a GRangesList

faFile a FaFile from genome

filter\_table a matrix / vector of length equal to cds

filter\_cds\_mod3

logical, default TRUE. Remove all ORFs that are not mod3, this speeds up the computation a lot, and usually removes malformed ORFs you would not want

anyway.

min\_counts\_cds\_filter

numeric, default: max(min(quantile(filter\_table, 0.50), 100), 100). Min-

imum number of counts from the 'filter\_table' argument.

 $\label{eq:with_Asites} \textbf{with\_A\_sites} \qquad \textbf{logical, default TRUE. Not used yet, will also return A site scores.}$ 

aligned\_position

what positions should be taken to calculate per-codon coverage. By default: "center", meaning that positions -1,0,1 will be taken. Alternative: "left", then

positions 0,1,2 are taken.

code a named character vector of size 64. Default: GENETIC\_CODE. Change if

organism does not use the standard code.

## Details

The primary column to use is "mean\_txNorm", this is the fair normalized score.

#### Value

a data.table of rows per codon / AA. All values are given per library, per site (A or P), sorted by the mean\_txNorm\_percentage column of the first library in the set, the columns are:

- variable (character): Library name
- seq (character): Amino acid:codon
- sum (integer): total counts per seq
- sum txNorm (integer): total counts per seq normalized per tx
- var (numeric): variance of total counts per seq
- N (integer): total number of codons of that type
- mean\_txNorm (numeric): Default use output, the fair codon usage, normalized both for gene and genome level for codon and read counts

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- ...
- alpha (numeric): dirichlet alpha MOM estimator (imagine mean and variance of probability in 1 value, the lower the value, the higher the variance, mean is decided by the relative value between samples)
- sum\_txNorm (integer): total counts per seq normalized per tx
- relative\_to\_max\_score (integer): Percentage use of codon
- type (factor(character)): "P" or "A"

#### References

https://www.ncbi.nlm.nih.gov/pmc/articles/PMC7196831/

#### See Also

```
Other codon: codon_usage_exp(), codon_usage_plot()
```

#### **Examples**

codon\_usage\_exp

Codon analysis for ORFik experiment

#### **Description**

Per AA / codon, analyse the coverage, get a multitude of features. For both A sites and P-sites (Input reads must be P-sites for now) This function takes inspiration from the codonDT paper, and among others returns the negative binomial estimates, but in addition many other features.

# Usage

```
codon_usage_exp(
   df,
   reads,
   cds = loadRegion(df, "cds", filterTranscripts(df)),
   mrna = loadRegion(df, "mrna", names(cds)),
   filter_cds_mod3 = TRUE,
   filter_table = assay(countTable(df, type = "summarized")[names(cds)]),
```

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```
faFile = df@fafile,
min_counts_cds_filter = max(min(quantile(filter_table, 0.5), 1000), 1000),
with_A_sites = TRUE,
code = GENETIC_CODE,
aligned_position = "center"
)
```

#### **Arguments**

df an ORFik experiment

reads either a single library (GRanges, GAlignment, GAlignmentPairs), or a list of

libraries returned from outputLibs(df) with p-sites. If list, the list must have

names coresponding to the library names.

cds a GRangesList, the coding sequences, default: loadRegion(df, "cds", filterTranscripts(df)),

longest isoform per gene.

mrna a GRangesList, the full mRNA sequences (matching by names the cds sequences),

default: loadRegion(df, "mrna", names(cds)).

filter\_cds\_mod3

logical, default TRUE. Remove all ORFs that are not mod3, this speeds up the computation a lot, and usually removes malformed ORFs you would not want

anyway.

filter\_table an numeric(integer) matrix, where rownames are the names of the full set of

mRNA transcripts. This will be subsetted to the cds subset you use. Then CDSs

are filtered from this table by the 'min\_counts\_cds\_filter' argument.

faFile FaFile, BSgenome, fasta/index file path or an ORFik experiment. This file is

usually used to find the transcript sequences from some GRangesList.

min\_counts\_cds\_filter

numeric, default: max(min(quantile(filter\_table, 0.50), 100), 100). Min-

imum number of counts from the 'filter\_table' argument.

 $\label{logical} with \verb|_A_sites| logical, default TRUE. Not used yet, will also return A site scores.$ 

code a named character vector of size 64. Default: GENETIC\_CODE. Change if

organism does not use the standard code.

aligned\_position

what positions should be taken to calculate per-codon coverage. By default: "center", meaning that positions -1,0,1 will be taken. Alternative: "left", then

positions 0,1,2 are taken.

#### **Details**

The primary column to use is "mean\_txNorm", this is the fair normalized score.

#### Value

a data.table of rows per codon / AA. All values are given per library, per site (A or P), sorted by the mean txNorm percentage column of the first library in the set, the columns are:

• variable (character): Library name

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- seq (character) : Amino acid:codon
- sum (integer): total counts per seq
- sum\_txNorm (integer): total counts per seq normalized per tx
- var (numeric): variance of total counts per seq
- N (integer): total number of codons of that type
- mean\_txNorm (numeric): Default use output, the fair codon usage, normalized both for gene and genome level for codon and read counts
- ..
- alpha (numeric): dirichlet alpha MOM estimator (imagine mean and variance of probability in 1 value, the lower the value, the higher the variance, mean is decided by the relative value between samples)
- sum\_txNorm (integer): total counts per seq normalized per tx
- relative\_to\_max\_score (integer): Percentage use of codon
- type (factor(character)): "P" or "A"

#### References

https://www.ncbi.nlm.nih.gov/pmc/articles/PMC7196831/

#### See Also

```
Other codon: codon_usage(), codon_usage_plot()
```

# **Examples**

codon\_usage\_plot

Plot codon\_usage

# **Description**

Plot codon\_usage

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#### Usage

```
codon_usage_plot(
  res,
  score_column = res$relative_to_max_score,
  ylab = "Ribo-seq library",
  legend.position = "none",
  limit = c(0, max(score_column)),
  midpoint = limit/2,
  monospace_font = TRUE
)
```

### Arguments

res a data.table of output from a codon\_usage function

score\_column numeric, default: res\$relative\_to\_max\_score. Which parameter to use as score

column.

ylab character vector, names for libraries to show on Y axis

legend.position

character, default "none", do not display legend.

limit numeric, 2 values for plot color limits. Default: c(0, max(score\_column))

midpoint numeric, default: limit/2. midpoint of color limit.

monospace\_font logical, default TRUE. Use monospace font, this does not work on systems (re-

quire specific font packages), set to FALSE if it crashes for you.

# Value

```
a ggplot object
```

#### See Also

```
Other codon: codon_usage(), codon_usage_exp()
```

# **Examples**

collapse.by.scores 37

collapse.by.scores

Merge reads by sum of existing scores

### **Description**

If you have multiple reads a same location but different read lengths, specified in meta column "size", it will sum up the scores (number of replicates) for all reads at that position

## Usage

```
collapse.by.scores(x)
```

## Arguments

Х

a GRanges object

### Value

merged GRanges object

### **Examples**

```
gr_s1 <- rep(GRanges("chr1", 1:10,"+"), 2)
gr_s2 <- GRanges("chr1", 1:12,"+")
gr2 <- GRanges("chr1", 21:40,"+")
gr <- c(gr_s1, gr_s2, gr2)
res <- convertToOneBasedRanges(gr,
    addScoreColumn = TRUE, addSizeColumn = TRUE)
ORFik:::collapse.by.scores(res)</pre>
```

collapse.fastq

Very fast fastq/fasta collapser

## Description

For each unique read in the file, collapse into 1 and state in the fasta header how many reads existed of that type. This is done after trimming usually, works best for reads < 50 read length. Not so effective for 150 bp length mRNA-seq etc.

```
collapse.fastq(
  files,
  outdir = file.path(dirname(files[1]), "collapsed"),
  header.out.format = "ribotoolkit",
  compress = FALSE,
  prefix = "collapsed_"
)
```

#### **Arguments**

files paths to fasta / fastq files to collapse. I tries to detect format per file, if file does

not have .fastq, .fastq.gz, .fq or fq.gz extensions, it will be treated as a .fasta file

format.

outdir outdir to save files, default: file.path(dirname(files[1]), "collapsed").

Inside same folder as input files, then create subfolder "collapsed", and add a

prefix of "collapsed\_" to the output names in that folder.

header.out.format

character, default "ribotoolkit", else must be "fastx". How the read header of the output fasta should be formated: ribotoolkit: ">seq1\_x55", sequence 1 has 55 duplicated reads collapsed. fastx: ">1-55", sequence 1 has 55 duplicated reads

collapsed

compress logical, default FALSE

prefix character, default "collapsed\_" Prefix to name of output file.

#### Value

invisible(NULL), files saved to disc in fasta format.

## **Examples**

```
fastq.folder <- tempdir() # <- Your fastq files
infiles <- dir(fastq.folder, "*.fastq", full.names = TRUE)
# collapse.fastq(infiles)</pre>
```

collapseDuplicatedReads

Collapse duplicated reads

### **Description**

For every GRanges, GAlignments read, with the same: seqname, start, (cigar) / width and strand, collapse and give a new meta column called "score", which contains the number of duplicates of that read. If score column already exists, will return input object!

## Usage

```
collapseDuplicatedReads(x, addScoreColumn = TRUE, ...)
```

# **Arguments**

x a GRanges, GAlignments or GAlignmentPairs object

addScoreColumn logical, default: (TRUE), if FALSE, only collapse and not keep score column of

counts for collapsed reads. Returns directly without collapsing if reuse.score.column

is FALSE and score is already defined.

.. alternative arguments for class instances. For example, see: ?'collapseDuplicatedReads, GRanges-me

#### Value

a GRanges, GAlignments, GAlignmentPairs or data.table object, same as input

## **Examples**

```
gr <- rep(GRanges("chr1", 1:10,"+"), 2)
collapseDuplicatedReads(gr)</pre>
```

```
collapseDuplicatedReads,data.table-method 

Collapse duplicated reads
```

### **Description**

For every GRanges, GAlignments read, with the same: seqname, start, (cigar) / width and strand, collapse and give a new meta column called "score", which contains the number of duplicates of that read. If score column already exists, will return input object!

### Usage

```
## S4 method for signature 'data.table'
collapseDuplicatedReads(
    x,
    addScoreColumn = TRUE,
    addSizeColumn = FALSE,
    reuse.score.column = TRUE,
    keepCigar = FALSE
)
```

### **Arguments**

x a GRanges, GAlignments or GAlignmentPairs object

addScoreColumn logical, default: (TRUE), if FALSE, only collapse and not keep score column of

counts for collapsed reads. Returns directly without collapsing if reuse.score.column

is FALSE and score is already defined.

addSizeColumn logical (FALSE), if TRUE, add a size column that for each read, that gives orig-

inal width of read. Useful if you need original read lengths. This takes care of soft clips etc. If collapsing reads, each unique range will be grouped also by

size.

reuse.score.column

logical (TRUE), if addScoreColumn is TRUE, and a score column exists, will sum up the scores to create a new score. If FALSE, will skip old score column and create new according to number of replicated reads after conversion. If

addScoreColumn is FALSE, this argument is ignored.

keepCigar logical, default FALSE. Keep the cigar information

### Value

a GRanges, GAlignments, GAlignmentPairs or data.table object, same as input

## Examples

```
gr <- rep(GRanges("chr1", 1:10,"+"), 2)
collapseDuplicatedReads(gr)</pre>
```

 ${\it collapse Duplicated Reads, GAlignment Pairs-method} \\ {\it Collapse duplicated reads}$ 

# Description

For every GRanges, GAlignments read, with the same: seqname, start, (cigar) / width and strand, collapse and give a new meta column called "score", which contains the number of duplicates of that read. If score column already exists, will return input object!

## Usage

```
## S4 method for signature 'GAlignmentPairs'
collapseDuplicatedReads(x, addScoreColumn = TRUE)
```

## **Arguments**

x a GRanges, GAlignments or GAlignmentPairs object

addScoreColumn logical, default: (TRUE), if FALSE, only collapse and not keep score column of counts for collapsed reads. Returns directly without collapsing if reuse.score.column is FALSE and score is already defined.

## Value

a GRanges, GAlignments, GAlignmentPairs or data.table object, same as input

## **Examples**

```
gr <- rep(GRanges("chr1", 1:10,"+"), 2)
collapseDuplicatedReads(gr)</pre>
```

collapseDuplicatedReads, GAlignments-method Collapse duplicated reads

#### **Description**

For every GRanges, GAlignments read, with the same: seqname, start, (cigar) / width and strand, collapse and give a new meta column called "score", which contains the number of duplicates of that read. If score column already exists, will return input object!

#### Usage

```
## S4 method for signature 'GAlignments'
collapseDuplicatedReads(x, addScoreColumn = TRUE, reuse.score.column = TRUE)
```

#### **Arguments**

a GRanges, GAlignments or GAlignmentPairs object

addScoreColumn logical, default: (TRUE), if FALSE, only collapse and not keep score column of counts for collapsed reads. Returns directly without collapsing if reuse.score.column is FALSE and score is already defined.

reuse.score.column

logical (TRUE), if addScoreColumn is TRUE, and a score column exists, will sum up the scores to create a new score. If FALSE, will skip old score column and create new according to number of replicated reads after conversion. If addScoreColumn is FALSE, this argument is ignored.

#### Value

a GRanges, GAlignments, GAlignmentPairs or data.table object, same as input

#### **Examples**

```
gr <- rep(GRanges("chr1", 1:10,"+"), 2)</pre>
collapseDuplicatedReads(gr)
```

collapseDuplicatedReads, GRanges-method Collapse duplicated reads

### **Description**

For every GRanges, GAlignments read, with the same: seqname, start, (cigar) / width and strand, collapse and give a new meta column called "score", which contains the number of duplicates of that read. If score column already exists, will return input object!

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### Usage

```
## S4 method for signature 'GRanges'
collapseDuplicatedReads(
    x,
    addScoreColumn = TRUE,
    addSizeColumn = FALSE,
    reuse.score.column = TRUE
)
```

## **Arguments**

x a GRanges, GAlignments or GAlignmentPairs object

addScoreColumn logical, default: (TRUE), if FALSE, only collapse and not keep score column of

 $counts \ for \ collapsed \ reads. \ Returns \ directly \ without \ collapsing \ if \ reuse. score. column$ 

is FALSE and score is already defined.

addSizeColumn logical (FALSE), if TRUE, add a size column that for each read, that gives orig-

inal width of read. Useful if you need original read lengths. This takes care of soft clips etc. If collapsing reads, each unique range will be grouped also by

size.

reuse.score.column

logical (TRUE), if addScoreColumn is TRUE, and a score column exists, will sum up the scores to create a new score. If FALSE, will skip old score column and create new according to number of replicated reads after conversion. If addScoreColumn is FALSE, this argument is ignored.

#### Value

a GRanges, GAlignments, GAlignmentPairs or data.table object, same as input

### **Examples**

```
gr <- rep(GRanges("chr1", 1:10,"+"), 2)
collapseDuplicatedReads(gr)</pre>
```

combn.pairs

Create all unique combinations pairs possible

### **Description**

Given a character vector, get all unique combinations of 2.

# Usage

```
combn.pairs(x)
```

### **Arguments**

x a character vector, will unique elements for you.

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#### Value

a list of character vector pairs

### **Examples**

```
df <- ORFik.template.experiment()
ORFik:::combn.pairs(df[, "libtype"])</pre>
```

computeFeatures

Get all main features in ORFik

### **Description**

If you want to get all the NGS and/or sequence features easily, you can use this function. Each feature have a link to an article describing its creation and idea behind it. Look at the functions in the feature family (in the "see also" section below) to see all of them. Example, if you want to know what the "te" column is, check out: ?translationalEff.

A short description of each feature is also shown here:

\*\* NGS features \*\* If not stated otherwise stated, the feature apply to Ribo-seq.

• countRFP: raw counts of Ribo-seq

• fpkmRFP: FPKM

• fpkmRNA: FPKM of RNA-seq

• te: Translation efficiency Ribo-seq / RNA-seq FPKM

• floss: Fragment length similarity score

• entropyRFP: Positional entropy

disengagementScores: downstream coverage from ORF

• RRS: Ribosome release score

• RSS: Ribosome staling score

• ORFScores: Periodicity score, does frame 0 have more reads

• ioScore : inside outside score: coverage ORF / coverage rest of transcript

• startCodonCoverage : Coverage over start codon + 2nt before start codon

• startRegionCoverage : Coverage over codon 2 & 3

• startRegionRelative : Peakness of TIS, startCodonCoverage / startRegionCoverage, 0-n

### \*\* Sequence features \*\*

• kozak : Similarity to kozak sequence for organism score, 0-1

• gc : GC percentage, 0-1

• StartCodons: Start codon as a string, "ATG"

• StopCodons: stop codon as a string, "TAA"

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• fractionLengths: ORF length compared to transcript, 0-1

## \*\* uORF features \*\*

- distORFCDS: Distance from ORF stop site to CDS, -n:n
- inFrameCDS: Is ORF in frame with downstream CDS, T/F
- isOverlappingCds: Is ORF overlapping with downstream CDS, T/F
- rankInTx : ORF with most upstream start codon is 1, 1-n

## Usage

```
computeFeatures(
  grl,
  RFP,
  RNA = NULL,
  Gtf,
  faFile = NULL,
  riboStart = 26,
  riboStop = 34,
  sequenceFeatures = TRUE,
  uorfFeatures = TRUE,
  grl.is.sorted = FALSE,
  weight.RFP = 1L,
  weight.RNA = 1L
)
```

## **Arguments**

grl	a GRangesList object with usually ORFs, but can also be either leaders, cds', 3' utrs, etc. This is the regions you want to score.		
RFP	RiboSeq reads as GAlignments, GRanges or GRangesList object		
RNA	RnaSeq reads as GAlignments, GRanges or GRangesList object		
Gtf	a TxDb object of a gtf file or path to gtf, gff .sqlite etc.		
faFile	a path to fasta indexed genome, an open FaFile, a BSgenome, or path to ORFik experiment with valid genome.		
riboStart	usually 26, the start of the floss interval, see ?floss		
riboStop	usually 34, the end of the floss interval		
sequenceFeatures			
	a logical, default TRUE, include all sequence features, that is: Kozak, fraction-Lengths, distORFCDS, isInFrame, isOverlapping and rankInTx. uorfFeatures = FALSE will remove the 4 last.		
uorfFeatures	a logical, default TRUE, include all uORF sequence features, that is: distOR-FCDS, isInFrame, isOverlapping and rankInTx		
grl.is.sorted	logical (F), a speed up if you know argument grl is sorted, set this to TRUE.		
weight.RFP	a vector (default: 1L). Can also be character name of column in RFP. As in translationalEff(weight = "score") for: GRanges("chr1", 1, "+", score = 5), would mean score column tells that this alignment region was found 5 times.		
weight.RNA	Same as weightRFP but for RNA weights. (default: 1L)		

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#### **Details**

If you used CageSeq to reannotate your leaders, your txDB object must contain the reassigned leaders. Use [reassignTxDbByCage()] to get the txdb.

As a note the library is reduced to only reads overlapping 'tx', so the library size in fpkm calculation is done on this subset. This will help remove rRNA and other contaminants.

Also if you have only unique reads with a weight column, explaining the number of duplicated reads, set weights to make calculations correct. See getWeights

#### Value

a data.table with scores, each column is one score type, name of columns are the names of the scores, i.g [floss()] or [fpkm()]

### See Also

```
Other features: computeFeaturesCage(), countOverlapsW(), disengagementScore(), distToCds(), distToTSS(), entropy(), floss(), fpkm(), fpkm_calc(), fractionLength(), initiationScore(), insideOutsideORF(), isInFrame(), isOverlapping(), kozakSequenceScore(), orfScore(), rankOrder(), ribosomeReleaseScore(), ribosomeStallingScore(), startRegion(), startRegionCoverage(), stopRegion(), subsetCoverage(), translationalEff()
```

### **Examples**

```
# Here we make an example from scratch
# Usually the ORFs are found in orfik, which makes names for you etc.
gtf <- system.file("extdata/references/danio_rerio", "annotations.gtf",
    package = "ORFik") ## location of the gtf file

suppressWarnings(txdb <- loadTxdb(gtf))
# use cds' as ORFs for this example
ORFs <- loadRegion(txdb, "cds")
ORFs <- makeORFNames(ORFs) # need ORF names
# make Ribo-seq data,
RFP <- unlistGrl(firstExonPerGroup(ORFs))
computeFeatures(ORFs, RFP, Gtf = txdb)
# For more details see vignettes.</pre>
```

 ${\tt compute} {\tt FeaturesCage}$ 

Get all main features in ORFik

#### Description

If you have a txdb with correctly reassigned transcripts, use: [computeFeatures()]

## Usage

```
computeFeaturesCage(
  grl,
  RFP,
 RNA = NULL,
 Gtf = NULL,
  tx = NULL,
  fiveUTRs = NULL,
  cds = NULL,
  threeUTRs = NULL,
  faFile = NULL,
  riboStart = 26,
  riboStop = 34,
  sequenceFeatures = TRUE,
  uorfFeatures = TRUE,
  grl.is.sorted = FALSE,
 weight.RFP = 1L,
 weight.RNA = 1L
)
```

### **Arguments**

grl	a GRangesList object with usually ORFs, but can also be either leaders, cds', 3' utrs, etc. This is the regions you want to score.		
RFP	RiboSeq reads as GAlignments, GRanges or GRangesList object		
RNA	RnaSeq reads as GAlignments, GRanges or GRangesList object		
Gtf	a TxDb object of a gtf file or path to gtf, gff .sqlite etc.		
tx	a GRangesList of transcripts, normally called from: exonsBy(Gtf, by = "tx", use.names = T) only add this if you are not including Gtf file If you are using CAGE, you do not need to reassign these to the cage peaks, it will do it for you.		
fiveUTRs	fiveUTRs as GRangesList, if you used cage-data to extend 5' utrs, remember to input CAGE assigned version and not original!		
cds	a GRangesList of coding sequences		
threeUTRs	a GRangesList of transcript 3' utrs, normally called from: threeUTRsByTranscript(Gtf, use.names = T)		
faFile	a path to fasta indexed genome, an open FaFile, a BSgenome, or path to ORFik experiment with valid genome.		
riboStart	usually 26, the start of the floss interval, see ?floss		
riboStop	usually 34, the end of the floss interval		
sequenceFeatures			
	a logical, default TRUE, include all sequence features, that is: Kozak, fraction-Lengths, distORFCDS, isInFrame, isOverlapping and rankInTx. uorfFeatures = FALSE will remove the 4 last.		
uorfFeatures	a logical, default TRUE, include all uORF sequence features, that is: distOR-		

FCDS, isInFrame, isOverlapping and rankInTx

```
grl.is.sorted logical (F), a speed up if you know argument grl is sorted, set this to TRUE.

weight.RFP a vector (default: 1L). Can also be character name of column in RFP. As in translationalEff(weight = "score") for: GRanges("chr1", 1, "+", score = 5), would mean score column tells that this alignment region was found 5 times.

weight.RNA Same as weightRFP but for RNA weights. (default: 1L)
```

#### **Details**

A specialized version if you don't have a correct txdb, for example with CAGE reassigned leaders while txdb is not updated. It is 2x faster for tested data. The point of this function is to give you the ability to input transcript etc directly into the function, and not load them from txdb. Each feature have a link to an article describing feature, try ?floss

#### Value

a data.table with scores, each column is one score type, name of columns are the names of the scores, i.g [floss()] or [fpkm()]

#### See Also

```
Other features: computeFeatures(), countOverlapsW(), disengagementScore(), distToCds(), distToTSS(), entropy(), floss(), fpkm(), fpkm_calc(), fractionLength(), initiationScore(), insideOutsideORF(), isInFrame(), isOverlapping(), kozakSequenceScore(), orfScore(), rankOrder(), ribosomeReleaseScore(), ribosomeStallingScore(), startRegion(), startRegionCoverage(), stopRegion(), subsetCoverage(), translationalEff()
```

### **Examples**

```
# a small example without cage-seg data:
# we will find ORFs in the 5' utrs
# and then calculate features on them
if (requireNamespace("BSgenome.Hsapiens.UCSC.hg19")) {
library(GenomicFeatures)
# Get the gtf txdb file
txdbFile <- system.file("extdata", "hg19_knownGene_sample.sqlite",</pre>
package = "GenomicFeatures")
txdb <- loadDb(txdbFile)</pre>
# Extract sequences of fiveUTRs.
fiveUTRs <- fiveUTRsByTranscript(txdb, use.names = TRUE)[1:10]</pre>
faFile <- BSgenome.Hsapiens.UCSC.hg19::Hsapiens</pre>
tx_seqs <- extractTranscriptSeqs(faFile, fiveUTRs)</pre>
# Find all ORFs on those transcripts and get their genomic coordinates
fiveUTR_ORFs <- findMapORFs(fiveUTRs, tx_seqs)</pre>
unlistedORFs <- unlistGrl(fiveUTR_ORFs)</pre>
# group GRanges by ORFs instead of Transcripts
fiveUTR_ORFs <- groupGRangesBy(unlistedORFs, unlistedORFs$names)</pre>
# make some toy ribo seq and rna seq data
```

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```
starts <- unlistGrl(ORFik:::firstExonPerGroup(fiveUTR_ORFs))
RFP <- promoters(starts, upstream = 0, downstream = 1)
score(RFP) <- rep(29, length(RFP)) # the original read widths
# set RNA seq to duplicate transcripts
RNA <- unlistGrl(exonsBy(txdb, by = "tx", use.names = TRUE))
#ORFik:::computeFeaturesCage(grl = fiveUTR_ORFs, RFP = RFP,
# RNA = RNA, Gtf = txdb, faFile = faFile)
}
# See vignettes for more examples</pre>
```

conditionNames

Get condition name variants

## Description

Used to standardize nomeclature for experiments.

Example: WT is main naming, but a variant is control will then be renamed to WT

### Usage

```
conditionNames()
```

### Value

a data.table with 2 columns, the main name, and all name variants of the main name in second column as a list.

### See Also

```
Other experiment_naming: batchNames(), cellLineNames(), cellTypeNames(), fractionNames(), inhibitorNames(), libNames(), mainNames(), repNames(), stageNames(), tissueNames()
```

config

Read directory config for ORFik experiments

## **Description**

Defines a folder for: 1. fastq files (raw data)

- 2. bam files (processed data)
- 3. references (organism annotation and STAR index)
- 4. experiments (Location to store and load all experiment .csv files) Update or use another config using config.save() function.

config.exper 49

## Usage

```
config(
  file = config_file(old_config_location = old_config_location),
  old_config_location = "~/Bio_data/ORFik_config.csv"
)
```

## **Arguments**

 $\label{location} file & location of config csv, default: config\_file (old\_config\_location = old\_config\_location) \\ old\_config\_location$ 

path, old config location before BiocFileCache implementation. Will copy this to cache directory and delete old version. This is done to follow bioc rules on not writing to user home directory.

### Value

a named character vector of length 3

## **Examples**

```
## Make with default config path
#config()
```

config.exper

Set directories for experiment

## Description

Defines a folder for: 1. fastq files (raw\_data)

- 2. bam files (processed data)
- 3. references (organism annotation and STAR index)
- 4. Experiment (name of experiment)

```
config.exper(
  experiment,
  assembly,
  type,
  config = ORFik::config(),
  sub_dir_single = file.path(type, experiment, ""),
  name_with_type_suffix = TRUE
)
```

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### Arguments

experiment short name of experiment (must be valid as a folder name)

assembly name of organism and assembly (must be valid as a folder name)

type name of sequencing type, Ribo-seq, RNA-seq, CAGE.. Can be more than one.

config a named character vector of length 3, default: ORFik::config()

sub\_dir\_single character, path. Default: file.path(type, experiment, "") The subdirectory relative to config defined main locations. If defined location should be used directly without making subdirectories, set to "".

name\_with\_type\_suffix

logical, default TRUE. Make fastq name like 'fastq RNA-seq', setting it to FALSE gives name 'fastq'. Only allowed when length(type) == 1

### Value

named character vector of paths for experiment

### **Examples**

```
# Where should files go in general?
ORFik::config()
# Paths for project: "Alexaki_Human" containing Ribo-seq and RNA-seq:
#config.exper("Alexaki_Human", "Homo_sapiens_GRCh38_101", c("Ribo-seq", "RNA-seq"))
```

config.save

Save/update directory config for ORFik experiments

## **Description**

Defines a folder for fastq files (raw\_data), bam files (processed data) and references (organism annotation and STAR index)

```
config.save(
  file = config_file(),
  fastq.dir = file.path(base.dir, "raw_data"),
  bam.dir = file.path(base.dir, "processed_data"),
  reference.dir = file.path(base.dir, "references"),
  exp.dir = file.path(base.dir, "ORFik_experiments/"),
  base.dir = "~/Bio_data",
  conf = data.frame(type = c("fastq", "bam", "ref", "exp"), directory = c(fastq.dir,
      bam.dir, reference.dir, exp.dir))
)
```

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### **Arguments**

file	location of config csv, default: config_file(old_config_location = old_config_location)	
fastq.dir	directory where ORFik puts fastq file directories, default: file.path(base.dir, "raw_data"), which is retrieved with: config()["fastq"]	
bam.dir	directory where ORFik puts bam file directories, default: file.path(base.dir, "processed_data"), which is retrieved with: config()["bam"]	
reference.dir	directory where ORFik puts reference file directories, default: file.path(base.dir, "references"), which is retrieved with: config()["ref"]	
exp.dir	directory where ORFik puts experiment csv files, default: file.path(base.dir, "ORFik_experiments/"), which is retrieved with: config()["exp"]	
base.dir	base directory for all output directories, default: "~/Bio_data"	
conf	data.frame of complete conf object, default: data.frame(type = c("fastq", "bam", "ref", "exp"), directory = c(fastq.dir, bam.dir, reference.dir, exp.dir))	

### Value

invisible(NULL), file saved to disc

## **Examples**

```
# Overwrite default config, with new base directory for files
#config.save(base.dir = "/media/Bio_data/") # Output files go here instead
# of ~/Bio_data
## Dont do this, but for understanding here is how to make a second config
#new_config_path <- config_file(query = "ORFik_config_2")
#config.save(new_config_path, "/media/Bio_data/raw_data/",
# "/media/Bio_data/processed_data", /media/Bio_data/references/)</pre>
```

config\_file

Get path for ORFik config in cache

## Description

Get path for ORFik config in cache

```
config_file(
  cache = BiocFileCache::getBFCOption("CACHE"),
  query = "ORFik_config",
  ask = interactive(),
  old_config_location = "~/Bio_data/ORFik_config.csv"
)
```

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# **Arguments**

cache path to bioc cache directory with rname from query argument. Default is:
BiocFileCache::getBFCOption("CACHE") For info, see: [BiocFileCache::BiocFileCache()]
query default: "ORFik\_config". Exact rname of the file in cache.

ask logical, default interactive().
old\_config\_location
 path, old config location before BiocFileCache implementation. Will copy this to cache directory and delete old version. This is done to follow bioc rules on

not writing to user home directory.

#### Value

a file path in cache

## **Examples**

```
config_file()
# Another config path
config_file(query = "ORFik_config_2")
```

convertLibs

Converted format of NGS libraries

### **Description**

Export as either .ofst, .wig, .bigWig,.bedo (legacy format) or .bedoc (legacy format) files:

Export files as .ofst for fastest load speed into R.

Export files as .wig / bigWig for use in IGV or other genome browsers.

The input files are checked if they exist from: envExp(df).

```
convertLibs(
   df,
   out.dir = libFolder(df),
   addScoreColumn = TRUE,
   addSizeColumn = TRUE,
   must.overlap = NULL,
   method = "None",
   type = "ofst",
   input.type = "ofst",
   reassign.when.saving = FALSE,
   envir = envExp(df),
   force = TRUE,
```

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```
library.names = bamVarName(df),
libs = outputLibs(df, type = input.type, chrStyle = must.overlap, library.names =
    library.names, output.mode = "list", force = force, BPPARAM = BPPARAM),
BPPARAM = bpparam()
)
```

#### **Arguments**

df an ORFik experiment

out.dir optional output directory, default: libFolder(df), if it is NULL, it will just reas-

sign R objects to simplified libraries. Will then create a final folder specified as: paste0(out.dir, "/", type, "/"). Here the files will be saved in format given by the

type argument.

addScoreColumn logical, default TRUE, if FALSE will not add replicate numbers as score col-

umn, see ORFik::convertToOneBasedRanges.

addSizeColumn logical, default TRUE, if FALSE will not add size (width) as size column, see

ORFik::convertToOneBasedRanges. Does not apply for (GAlignment version

of.ofst) or .bedoc. Since they contain the original cigar.

must.overlap default (NULL), else a GRanges / GRangesList object, so only reads that over-

lap (must.overlap) are kept. This is useful when you only need the reads over

transcript annotation or subset etc.

method character, default "None", the method to reduce ranges, for more info see convertToOneBasedRanges

type character, output format, default "ofst". Alternatives: "ofst", "bigWig", "wig", "bedo"

or "bedoc". Which format you want. Will make a folder within out.dir with this

name containing the files.

input.type character, input type "ofst". Remember this function uses the loaded libraries if

existing, so this argument is usually ignored. Only used if files do not already

exist.

reassign.when.saving

logical, default FALSE. If TRUE, will reassign library to converted form after

saving. Ignored when out.dir = NULL.

envir environment to save to, default envExp(df), which defaults to .GlobalEnv, but

can be set with envExp(df) <- new.env() etc.

force logical, default TRUE If TRUE, reload library files even if matching named

variables are found in environment used by experiment (see envExp) A simple way to make sure correct libraries are always loaded. FALSE is faster if data is

loaded correctly already.

library.names character vector, names of libraries, default: name decider(df, naming)

libs list, output of outputLibs as list of GRanges/GAlignments/GAlignmentPairs ob-

jects. Set input.type and force arguments to define parameters.

BPPARAM how many cores/threads to use? default: bpparam(). To see number of threads

used, do bpparam()\$workers. You can also add a time remaining bar, for a

more detailed pipeline.

#### **Details**

We advice you to not use this directly, as other function are more safe for library type conversions. See family description below. This is mostly used internally in ORFik. It is only adviced to use if large bam files are already loaded in R and conversions are wanted from those.

See export.ofst, export.wiggle, export.bedo and export.bedoc for information on file formats.

If libraries of the experiment are already loaded into environment (default: .globalEnv) is will export using those files as templates. If they are not in environment the .ofst files from the bam files are loaded (unless you are converting to .ofst then the .bam files are loaded).

### Value

invisible NULL (saves files to disc or R .GlobalEnv)

#### See Also

```
Other lib_converters: convert_bam_to_ofst(), convert_to_bigWig(), convert_to_covRle(), convert_to_covRleList()
```

#### **Examples**

```
df <- ORFik.template.experiment()
#convertLibs(df, out.dir = NULL)
# Keep only 5' ends of reads
#convertLibs(df, out.dir = NULL, method = "5prime")</pre>
```

convertToOneBasedRanges

Convert a GRanges Object to 1 width reads

### **Description**

There are 5 ways of doing this

- 1. Take 5' ends, reduce away rest (5prime)
- 2. Take 3' ends, reduce away rest (3prime)
- 3. Tile to 1-mers and include all (tileAll)
- 4. Take middle point per GRanges (middle)
- 5. Get original with metacolumns (None)

You can also do multiple at a time, then output is GRangesList, where each list group is the operation (5prime is [1], 3prime is [2] etc)

Many other ways to do this have their own functions, like startSites and stopSites etc. To retain information on original width, set addSizeColumn to TRUE. To compress data, 1 GRanges object per unique read, set addScoreColumn to TRUE. This will give you a score column with how many duplicated reads there were in the specified region.

#### Usage

```
convertToOneBasedRanges(
   gr,
   method = "5prime",
   addScoreColumn = FALSE,
   addSizeColumn = FALSE,
   after.softclips = TRUE,
   along.reference = FALSE,
   reuse.score.column = TRUE)
```

#### **Arguments**

gr GRanges, GAlignment or GAlignmentPairs object to reduce.

method character, default "5prime", the method to reduce ranges, see NOTE for more

info.

addScoreColumn logical (FALSE), if TRUE, add a score column that sums up the hits per unique

range. This will make each read unique, so that each read is 1 time, and score column gives the number of collapsed hits. A useful compression. If add-SizeColumn is FALSE, it will not differentiate between reads with same start and stop, but different length. If addSizeColumn is FALSE, it will remove it.

Collapses after conversion.

addSizeColumn logical (FALSE), if TRUE, add a size column that for each read, that gives orig-

inal width of read. Useful if you need original read lengths. This takes care of soft clips etc. If collapsing reads, each unique range will be grouped also by

ize.

after.softclips

logical (TRUE), include softclips in width. Does not apply if along.reference is

TRUE.

along.reference

logical (FALSE), example: The cigar "26MI2" is by default width 28, but if along.reference is TRUE, it will be 26. The length of the read along the reference. Also "1D20M" will be 21 if by along.reference is TRUE. Intronic regions

(cigar: N) will be removed. So: "1M200N19M" is 20, not 220.

reuse.score.column

logical (TRUE), if addScoreColumn is TRUE, and a score column exists, will sum up the scores to create a new score. If FALSE, will skip old score column and create new according to number of replicated reads after conversion. If addScoreColumn is FALSE, this argument is ignored.

#### **Details**

NOTE: Note: For cigar based ranges (GAlignments), the 5' end is the first non clipped base (neither soft clipped or hard clipped from 5'). This is following the default of bioconductor. For special case of GAlignmentPairs, 5prime will only use left (first) 5' end and read and 3prime will use only right (last) 3' end of read in pair. tileAll and middle can possibly find poinst that are not in the reads since: lets say pair is 1-5 and 10-15, middle is 7, which is not in the read.

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#### Value

Converted GRanges object

#### See Also

```
Other utils: bedToGR(), export.bed12(), export.bigWig(), export.fstwig(), export.wiggle(), fimport(), findFa(), fread.bed(), optimizeReads(), readBam(), readBigWig(), readWig()
```

## **Examples**

```
gr <- GRanges("chr1", 1:10,"+")
# 5 prime ends
convertToOneBasedRanges(gr)
# is equal to convertToOneBasedRanges(gr, method = "5prime")
# 3 prime ends
convertToOneBasedRanges(gr, method = "3prime")
# With lengths
convertToOneBasedRanges(gr, addSizeColumn = TRUE)
# With score (# of replicates)
gr <- rep(gr, 2)
convertToOneBasedRanges(gr, addSizeColumn = TRUE, addScoreColumn = TRUE)</pre>
```

## Description

Saved by default in folder "ofst" relative to default libraries of experiment. Speeds up loading of full files compared to bam by large margins.

#### Usage

```
convert_bam_to_ofst(
   df,
   in_files = filepath(df, "default"),
   out_dir = file.path(libFolder(df), "ofst"),
   verbose = TRUE,
   strandMode = rep(0, length(in_files))
)
```

### Arguments

```
an ORFik experiment, or NULL is allowed if both in_files and out_dir is spec-
ified manually.

in_files paths to input files, default: filepath(df, "default") with bam format files.

out_dir paths to output files, default file.path(libFolder(df), "cov_RLE").
```

convert\_to\_bigWig 57

verbose

logical, default TRUE, message about library output status.

strandMode

numeric, default 0. Only used for paired end bam files. One of (0: strand = \*, 1: first read of pair is +, 2: first read of pair is -). See ?strandMode. Note: Sets default to 0 instead of 1, as readGAlignmentPairs uses 1. This is to guarantee hits, but will also make mismatches of overlapping transcripts in opposite directions.

#### **Details**

If you want to keep bam files loaded or faster conversion if you already have them loaded, use ORFik::convertLibs instead

### Value

invisible(NULL), files saved to disc

### See Also

```
Other lib_converters: convertLibs(), convert_to_bigWig(), convert_to_covRle(), convert_to_covRleList()
```

### **Examples**

```
df <- ORFik.template.experiment.zf()
## Usually do default folder, here we use tmpdir
folder_to_save <- file.path(tempdir(), "ofst")
convert_bam_to_ofst(df, out_dir = folder_to_save)
fimport(file.path(folder_to_save, "ribo-seq.ofst"))</pre>
```

convert\_to\_bigWig

Convert to BigWig

## Description

Convert to BigWig

```
convert_to_bigWig(
   df,
   in_files = filepath(df, "pshifted"),
   out_dir = file.path(libFolder(df), "bigwig"),
   split.by.strand = TRUE,
   split.by.readlength = FALSE,
   seq_info = seqinfo(df),
   weight = "score",
   is_pre_collapsed = FALSE,
   verbose = TRUE
)
```

58 convert\_to\_covRle

#### **Arguments**

df an ORFik experiment, or NULL is allowed if both in\_files and out\_dir is spec-

ified manually.

in\_files paths to input files, default pshifted files: filepath(df, "pshifted") in ofst

format

out\_dir paths to output files, default file.path(libFolder(df), "bigwig").

split.by.strand

logical, default TRUE, split into forward and reverse strand RleList inside cov-

Rle object.

split.by.readlength

logical, default FALSE, split into files for each readlength, defined by read-

Widths(x) for each file.

seq\_info SeqInfo object, default seqinfo(findFa(df))

weight integer, numeric or single length character. Default "score". Use score column

in loaded in\_files.

is\_pre\_collapsed

logical, default FALSE. Have you already collapsed reads with collapse.by.scores, so each positions is only in 1 GRanges object with a score column per readlength?

Set to TRUE, only if you are sure, will give a speedup.

verbose logical, default TRUE, message about library output status.

#### Value

invisible(NULL), files saved to disc

### See Also

```
Other lib converters: convertLibs(), convert_bam_to_ofst(), convert_to_covRle(), convert_to_covRleList()
```

# Examples

convert\_to\_covRle

Convert libraries to covRle

## **Description**

Saved by default in folder "cov\_RLE" relative to default libraries of experiment

convert\_to\_covRle 59

### Usage

```
convert_to_covRle(
   df,
   in_files = filepath(df, "pshifted"),
   out_dir = file.path(libFolder(df), "cov_RLE"),
   split.by.strand = TRUE,
   split.by.readlength = FALSE,
   seq_info = seqinfo(df),
   weight = "score",
   verbose = TRUE
)
```

### **Arguments**

an ORFik experiment, or NULL is allowed if both in\_files and out\_dir is specified manually.

in\_files paths to input files, default pshifted files: filepath(df, "pshifted") in ofst

format

out\_dir paths to output files, default file.path(libFolder(df), "cov\_RLE").

split.by.strand

logical, default TRUE, split into forward and reverse strand RleList inside cov-

Rle object.

split.by.readlength

logical, default FALSE, split into files for each readlength, defined by read-

Widths(x) for each file.

seq\_info SeqInfo object, default seqinfo(findFa(df))

weight integer, numeric or single length character. Default "score". Use score column

in loaded in\_files.

verbose logical, default TRUE, message about library output status.

#### Value

invisible(NULL), files saved to disc

#### See Also

```
Other lib_converters: convertLibs(), convert_bam_to_ofst(), convert_to_bigWig(), convert_to_covRleList()
```

### **Examples**

```
df <- ORFik.template.experiment()[10,]
## Usually do default folder, here we use tmpdir
folder_to_save <- file.path(tempdir(), "cov_RLE")
convert_to_covRle(df, out_dir = folder_to_save)
fimport(file.path(folder_to_save, "RFP_Mutant_rep2.covrds"))</pre>
```

convert\_to\_covRleList

convert\_to\_covRleList Convert libraries to covRleList objects

## Description

Useful to store reads separated by readlength, for much faster coverage calculation. Saved by default in folder "cov\_RLE\_List" relative to default libraries of experiment

## Usage

```
convert_to_covRleList(
   df,
   in_files = filepath(df, "pshifted"),
   out_dir = file.path(libFolder(df), "cov_RLE_List"),
   out_dir_merged = file.path(libFolder(df), "cov_RLE"),
   split.by.strand = TRUE,
   seq_info = seqinfo(df),
   weight = "score",
   verbose = TRUE
)
```

## **Arguments**

df	an ORFik experiment, or NULL is allowed if both in_files and out_dir is specified manually.	
in_files	paths to input files, default pshifted files: $filepath(df, "pshifted")$ in ofst format	
out_dir	paths to output files, default file.path(libFolder(df), "cov_RLE_List").	
out_dir_merged	character vector of paths, default: file.path(libFolder(df), "cov_RLE"). Paths to merged output files, Set to NULL to skip making merged covRle.	
split.by.strand		
	logical, default TRUE, split into forward and reverse strand RleList inside cov-Rle object.	
seq_info	SeqInfo object, default seqinfo(findFa(df))	
weight	integer, numeric or single length character. Default "score". Use score column in loaded in_files.	
verbose	logical, default TRUE, message about library output status.	

#### Value

invisible(NULL), files saved to disc

## See Also

```
Other lib_converters: convertLibs(), convert_bam_to_ofst(), convert_to_bigWig(), convert_to_covRle()
```

convert\_to\_fstWig 61

### **Examples**

```
df <- ORFik.template.experiment()[10,]
## Usually do default folder, here we use tmpdir
folder_to_save <- file.path(tempdir(), "cov_RLE_List")
folder_to_save_merged <- file.path(tempdir(), "cov_RLE")
ORFik:::convert_to_covRleList(df, out_dir = folder_to_save,
out_dir_merged = folder_to_save_merged)
fimport(file.path(folder_to_save, "RFP_Mutant_rep2.covrds"))</pre>
```

convert\_to\_fstWig

Convert to fstwig

#### **Description**

Will split files by chromosome for faster loading for now. This feature might change in the future!

### Usage

```
convert_to_fstWig(
   df,
   in_files = filepath(df, "pshifted"),
   out_dir = file.path(libFolder(df), "fstwig"),
   split.by.strand = TRUE,
   split.by.readlength = FALSE,
   seq_info = seqinfo(df),
   weight = "score",
   is_pre_collapsed = FALSE,
   verbose = TRUE
)
```

in loaded in\_files.

# **Arguments**

df an ORFik experiment, or NULL is allowed if both in files and out dir is specified manually. paths to input files, default pshifted files: filepath(df, "pshifted") in ofst in\_files out\_dir paths to output files, default file.path(libFolder(df), "bigwig"). split.by.strand logical, default TRUE, split into forward and reverse strand RleList inside cov-Rle object. split.by.readlength logical, default FALSE, split into files for each readlength, defined by read-Widths(x) for each file. seq\_info SeqInfo object, default seqinfo(findFa(df)) integer, numeric or single length character. Default "score". Use score column weight

62 correlation.plots

```
is_pre_collapsed
```

logical, default FALSE. Have you already collapsed reads with collapse.by.scores, so each positions is only in 1 GRanges object with a score column per readlength? Set to TRUE, only if you are sure, will give a speedup.

verbose

logical, default TRUE, message about library output status.

#### Value

invisible(NULL), files saved to disc

correlation.plots

Correlation plots between all samples

### **Description**

Get correlation plot of raw counts and/or log2(count + 1) over selected region in: c("mrna", "leaders", "cds", "trailers")

Note on correlation: Pearson correlation, using pairwise observations to fill in NA values for the covariance matrix.

### Usage

```
correlation.plots(
    df,
    output.dir,
    region = "mrna",
    type = "fpkm",
    height = 400,
    width = 400,
    size = 0.15,
    plot.ext = ".pdf",
    complex.correlation.plots = TRUE,
    data_for_pairs = countTable(df, region, type = type),
    as_gg_list = FALSE,
    text_size = 4,
    method = c("pearson", "spearman")[1]
)
```

## **Arguments**

```
df an ORFik experiment

output.dir directory to save to, named: cor_plot, cor_plot_log2 and/or cor_plot_simple with either.pdf or.png

region a character (default: mrna), make raw count matrices of whole mrnas or one of (leaders, cds, trailers)
```

cor\_plot 63

which value to use, "fpkm", alternative "counts".

as\_gg\_list logical, default FALSE. Return as a list of ggplot objects instead of as a grob.

Gives you the ability to modify plots more directly.

text\_size size of correlation numbers
method c("pearson", "spearman")[1]

#### Value

type

invisible(NULL) / if as\_gg\_list is TRUE, return a list of raw plots.

cor\_plot

Get correlation between columns

### **Description**

Get correlation between columns

```
cor_plot(
  dt_cor,
  col = c(low = "blue", high = "red", mid = "white", na.value = "white"),
  limit = c(ifelse(min(dt_cor$Cor, na.rm = TRUE) < 0, -1, 0), 1),
  midpoint = mean(limit),
  label_name = "Pearson\nCorrelation",
  text_size = 4,
  legend.position = c(0.4, 0.7),
  legend.direction = "horizontal"
)</pre>
```

64 cor\_table

## Arguments

```
dt_cor
                  a data.table, with column Cor
col
                  colors c(low = "blue", high = "red", mid = "white", na.value = "white")
                  default (-1, 1), defined by: c(ifelse(min(dt_cor$Cor, na.rm = TRUE) < 0,</pre>
limit
                  -1, 0), 1)
midpoint
                  midpoint of correlation values in label coloring.
label_name
                  name of correlation method, default "Pearson Correlation" with newline af-
                  ter Pearson.
                  size of correlation numbers
text_size
legend.position
                  default c(0.4, 0.7), other: "top", "right",...
legend.direction
                  default "horizontal", or "vertical"
```

### Value

```
a ggplot (heatmap)
```

cor\_table

Get correlation between columns

## Description

Get correlation between columns

### Usage

```
cor_table(
   dt,
   method = c("pearson", "spearman")[1],
   upper_triangle = TRUE,
   decimals = 2,
   melt = TRUE,
   na.rm.melt = TRUE
)
```

## Arguments

```
dt a data.table

method c("pearson", "spearman")[1]

upper_triangle logical, default TRUE. Make lower triangle values NA.

decimals numeric, default 2. How many decimals for correlation

melt logical, default TRUE.

na.rm.melt logical, default TRUE. Remove NA values from melted table.
```

countOverlapsW 65

#### Value

a data.table with 3 columns, Var1, Var2 and Cor

countOverlapsW

CountOverlaps with weights

## Description

Similar to countOverlaps, but takes an optional weight column. This is usually the score column

### Usage

```
countOverlapsW(query, subject, weight = NULL, ...)
```

### **Arguments**

query IRanges, IRangesList, GRanges, GRangesList object. Usually transcript a tran-

script region.

subject GRanges, GRangesList, GAlignment or covRle, usually reads.

weight (default: NULL), if defined either numeric or character name of valid meta col-

umn in subject. If weight is single numeric, it is used for all. A normall weight is the score column given as weight = "score". GRanges("chr1", 1, "+", score = 5), would mean score column tells that this alignment region was found 5 times.

Ignored if subject is covRle.

... additional arguments passed to countOverlaps/findOverlaps

#### Value

a named vector of number of overlaps to subject weighted by 'weight' column.

## See Also

```
Other features: computeFeatures(), computeFeaturesCage(), disengagementScore(), distToCds(), distToTSS(), entropy(), floss(), fpkm(), fpkm_calc(), fractionLength(), initiationScore(), insideOutsideORF(), isInFrame(), isOverlapping(), kozakSequenceScore(), orfScore(), rankOrder(), ribosomeReleaseScore(), ribosomeStallingScore(), startRegion(), startRegionCoverage(), stopRegion(), subsetCoverage(), translationalEff()
```

### **Examples**

66 countTable

```
strand=c("+"),
               score=c(10, 20, 15, 5))
countOverlaps(gr1, gr2)
countOverlapsW(gr1, gr2, weight = "score")
```

countTable

Extract count table directly from experiment

### **Description**

Used to quickly load pre-created read count tables to R.

If df is experiment: Extracts by getting /QC\_STATS directory, and searching for region Requires ORFikQC to have been run on experiment, to get default count tables!

# Usage

```
countTable(
  df,
  region = "mrna",
  type = "count",
  collapse = FALSE,
  count.folder = "default"
)
```

### **Arguments**

df

an ORFik experiment or path to folder with countTable, use path if not same folder as experiment libraries. Will subset to the count tables specified if df is experiment. If experiment has 4 rows and you subset it to only 2, then only those 2 count tables will be outputted.

region

a character vector (default: "mrna"), make raw count matrices of whole mrnas

or one of (leaders, cds, trailers).

type

character, default: "count" (raw counts matrix). Which object type and normalization do you want? "summarized" (SummarizedExperiment object), "deseq" (Deseq2 experiment, design will be all valid non-unique columns except replicates, change by using DESeq2::design, normalization alternatives are: "fpkm", "log2fpkm" or "log10fpkm".

collapse

a logical/character (default FALSE), if TRUE all samples within the group SAM-PLE will be collapsed to one. If "all", all groups will be merged into 1 column called merged\_all. Collapse is defined as rowSum(elements\_per\_group) / ncol(elements\_per\_group)

count.folder

character, default "auto" (Use count tables from original bam files stored in "QC\_STATS", these are like HTseq count tables). To load your custome count tables from pshifted reads, set to "pshifted" (remember to create the pshifted tables first!). If you have custom ranges, like reads over uORFs stored in a folder called "/uORFs" relative to the bam files, set to "uORFs". Always create these custom count tables with makeSummarizedExperimentFromBam. Always make the location of the folder directly inside the bam file directory!

countTable\_regions 67

#### **Details**

If df is path to folder: Loads the file in that directory with the regex region.rds, where region is what is defined by argument, if multiple exist, see if any start with "countTable\_", if so, subset. If loaded as SummarizedExperiment or deseq, the colData will be made from ORFik.experiment information.

#### Value

a data.table/SummarizedExperiment/DESeq object of columns as counts / normalized counts per library, column name is name of library. Rownames must be unique for now. Might change.

#### See Also

Other countTable: countTable\_regions()

## **Examples**

```
# Make experiment
df <- ORFik.template.experiment()</pre>
# Make QC report to get counts ++ (not needed for this template)
# ORFikQC(df)
# Get count Table of mrnas
# countTable(df, "mrna")
# Get count Table of cds
# countTable(df, "cds")
# Get count Table of mrnas as fpkm values
# countTable(df, "mrna", type = "count")
# Get count Table of mrnas with collapsed replicates
# countTable(df, "mrna", collapse = TRUE)
# Get count Table of mrnas as summarizedExperiment
# countTable(df, "mrna", type = "summarized")
# Get count Table of mrnas as DESeq2 object,
# for differential expression analysis
# countTable(df, "mrna", type = "deseq")
```

countTable\_regions

Make a list of count matrices from experiment

### **Description**

By default will make count tables over mRNA, leaders, cds and trailers for all libraries in experiment. region

68 countTable\_regions

### Usage

```
countTable_regions(
    df,
    out.dir = libFolder(df),
    longestPerGene = FALSE,
    geneOrTxNames = "tx",
    regions = c("mrna", "leaders", "cds", "trailers"),
    type = "count",
    lib.type = "ofst",
    weight = "score",
    rel.dir = "QC_STATS",
    forceRemake = FALSE,
    library.names = bamVarName(df),
    BPPARAM = bpparam()
)
```

### **Arguments**

df	an OPFik	experiment
UT .	an Oktik	experiment

out.dir character, output directory, default: resFolder(df). Will make a folder within

this called "QC\_STATS" with all results in this directory. Warning: If you assign not default path, you will have a hazzle to load files later. Much easier to load count tables, statistics, ++ later with default. Update resFolder of df instead if

needed.

longestPerGene a logical (default FALSE), if FALSE all transcript isoforms per gene. Ignored if

"region" is not a character of either: "mRNA", "tx", "cds", "leaders" or "trailers".

geneOrTxNames a character vector (default "tx"), should row names keep trancript names ("tx")

or change to gene names ("gene")

regions a character vector, default: c("mrna", "leaders", "cds", "trailers"), make raw

count matrices of whole regions specified. Can also be a custom GRangesList

of for example uORFs or a subset of cds etc.

type default: "count" (raw counts matrix), alternative is "fpkm", "log2fpkm" or "log10fpkm"

lib.type a character(default: "default"), load files in experiment or some precomputed

variant, either "ofst", "bedo", "bedoc" or "pshifted". These are made with OR-Fik:::convertLibs() or shiftFootprintsByExperiment(). Can also be custom user

made folders inside the experiments bam folder.

weight numeric or character, a column to score overlaps by. Default "score", will check

for a metacolumn called "score" in libraries. If not found, will not use weights.

rel.dir relative output directory for out.dir, default: "QC\_STATS". For pshifted, write

"pshifted".

forceRemake logical, default FALSE. If TRUE, will not look for existing file count table files.

library.names character, default: bamVarName(df). Names to load libraries as to environment

and names to display in plots.

BPPARAM how many cores/threads to use? default: bpparam()

## Value

a list of data.table, 1 data.table per region. The regions will be the names the list elements.

### See Also

```
Other countTable: countTable()
```

## **Examples**

```
##Make experiment
df <- ORFik.template.experiment()
## Create count tables for all default regions
# countTable_regions(df)
## Pshifted reads (first create pshiftead libs)
# countTable_regions(df, lib.type = "pshifted", rel.dir = "pshifted")</pre>
```

coverageByTranscriptC coverageByTranscript with coverage input

## **Description**

Extends the function with direct genome coverage input, see coverageByTranscript for original function.

### Usage

```
coverageByTranscriptC(x, transcripts, ignore.strand = !strandMode(x))
```

## **Arguments**

```
x a covRle (one RleList for each strand in object), must have defined and correct
```

seqlengths in its SeqInfo object.

```
transcripts GRangesList
```

```
ignore.strand a logical (default: length(x) == 1)
```

### Value

```
Integer Rle of coverage, 1 per transcript
```

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coverageByTranscriptW coverageByTranscript with weights

## **Description**

Extends the function with weights, see coverageByTranscript for original function.

## Usage

```
coverageByTranscriptW(
    x,
    transcripts,
    ignore.strand = FALSE,
    weight = 1L,
    seqinfo.x.is.correct = FALSE
)
```

### **Arguments**

reads (GRanges, GAlignments)

transcripts GRangesList

ignore.strand a logical (default: FALSE)

weight a vector (default: 1L), if single number applies for all, else it must be the string

name of a defined meta column in "x", that gives number of times a read was found. GRanges("chr1", 1, "+", score = 5), would mean score column tells that

this alignment was found 5 times.

seginfo.x.is.correct

logical, default FALSE. If you know  $\boldsymbol{x}$ , has correct seqinfo, then you can save

some computation time by setting this to TRUE.

### Value

Integer Rle of coverage, 1 per transcript

coverageGroupings

Get grouping for a coverage table in ORFik

#### **Description**

Either of two groupings: GF: Gene, fraction FGF: Fraction, position, feature It finds which of these exists, and auto groups

```
coverageGroupings(logicals, grouping = "GF")
```

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## **Arguments**

logicals size 2 logical vector, the is.null checks for each column, grouping which grouping to perform, default "GF" Gene & Fraction grouping. Alternative

"FGF", Fraction & position & feature.

### **Details**

Normally not used directly!

#### Value

a quote of the grouping to pass to data.table

coverageHeatMap

Create a heatmap of coverage

## **Description**

Creates a ggplot representing a heatmap of coverage:

• Rows : Position in region

• Columns : Read length

• Index intensity: (color) coverage scoring per index.

Coverage rows in heat map is fraction, usually fractions is divided into unique read lengths (standard Illumina is 76 unique widths, with some minimum cutoff like 15.) Coverage column in heat map is score, default zscore of counts. These are the relative positions you are plotting to. Like +/- relative to TIS or TSS.

```
coverageHeatMap(
  coverage,
  output = NULL,
  scoring = "zscore",
  legendPos = "right",
  addFracPlot = FALSE,
  xlab = "Position relative to start site",
  ylab = "Protected fragment length",
  colors = "default",
  title = NULL,
  increments.y = "auto",
  gradient.max = max(coverage$score)
)
```

72 coverageHeatMap

#### **Arguments**

coverage a data.table, e.g. output of scaledWindowCoverage output character string (NULL), if set, saves the plot as pdf or png to path given. If no format is given, is save as pdf. character vector, default "zscore", Which scoring did you use to create? either scoring of zscore, transcriptNormalized, sum, mean, median, .. see ?coverageScorings for info and more alternatives. legendPos a character, Default "right". Where should the fill legend be ? ("top", "bottom", "right", "left") addFracPlot Add margin histogram plot on top of heatmap with fractions per positions the x-axis label, default "Position relative to start site" xlab the y-axis label, default "Protected fragment length" ylab colors character vector, default: "default", this gives you: c("white", "yellow2", "yellow3", "lightblue", "blue", "navy"), do "high" for more high contrasts, or specify your own colors. title a character, default NULL (no title), what is the top title of plot? increments of y axis, default "auto". Or a numeric value < max position & > increments.y min position. gradient.max numeric, defualt: max(coverage\$score). What data value should the top color be ? Good to use if you want to compare 2 samples, with the same color intensity, in that case set this value to the max score of the 2 coverage tables.

### Details

Colors: Remember if you want to change anything like colors, just return the ggplot object, and reassign like: obj + scale color brewer() etc. Standard colors are:

• 0 reads in whole readlength: gray

• few reads in position : white

• medium reads in position : yellow

• many reads in position : dark blue

### Value

a ggplot object of the coverage plot, NULL if output is set, then the plot will only be saved to location.

### See Also

Other heatmaps: heatMapL(), heatMapRegion(), heatMap\_single()
Other coveragePlot: pSitePlot(), savePlot(), windowCoveragePlot()

coveragePerTiling 73

## **Examples**

coveragePerTiling

Get coverage per group

# Description

It tiles each GRangesList group to width 1, and finds hits per position.

A range from 1:5 will split into c(1,2,3,4,5) and count hits on each. This is a safer speedup of coverageByTranscript from GenomicFeatures. It also gives the possibility to return as data.table, for faster computations.

## Usage

```
coveragePerTiling(
  grl,
  reads,
  is.sorted = FALSE,
  keep.names = TRUE,
  as.data.table = FALSE,
  withFrames = FALSE,
  weight = "score",
  drop.zero.dt = FALSE,
  fraction = NULL
)
```

## **Arguments**

grl a GRangesList of 5' utrs, CDS, transcripts, etc.

reads a GAlignments, GRanges, or precomputed coverage as covRle (one for each

strand) of RiboSeq, RnaSeq etc.

Weigths for scoring is default the 'score' column in 'reads'. Can also be random

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access paths to bigWig or fstwig file. Do not use random access for more than a few genes, then loading the entire files is usually better. File streaming is still in beta, so use with care!

is.sorted logical (FALSE), is grl sorted. That is + strand groups in increasing ranges

(1,2,3), and - strand groups in decreasing ranges (3,2,1)

keep.names logical (TRUE), keep names or not. If as.data.table is TRUE, names (genes

column) will be a factor column, if FALSE it will be an integer column (index of gene), so first input grl element is 1. Dropping names gives  $\sim 20~\%$  speedup. If drop.zero.dt is FALSE, data.table will not return names, will use index (to

avoid memory explosion).

as.data.table a logical (FALSE), return as data.table with 2 columns, position and count.

withFrames a logical (FALSE), only available if as.data.table is TRUE, return the ORF

frame, 1,2,3, where position 1 is 1, 2 is 2 and 4 is 1 etc.

weight (default: 'score'), if defined a character name of valid meta column in subject.

GRanges("chr1", 1, "+", score = 5), would mean score column tells that this alignment region was found 5 times. Formats which loads a score column like this: Bigwig, wig, ORFik ofst, collapsed bam, bedoc and .bedo. As do CAGEr CAGE files and many other package formats. You can also assign a score col-

umn manually.

drop.zero.dt logical FALSE, if TRUE and as.data.table is TRUE, remove all 0 count posi-

tions. This greatly speeds up and most importantly, greatly reduces memory usage. Will not change any plots, unless 0 positions are used in some sense.

(mean, median, zscore coverage will only scale differently)

fraction integer or character, a description column. Useful for grouping multiple outputs

together. If returned as Rle, this is added as: metadata(coverage) <- list(fraction

= fraction). If as.data.table it will be added as an additional column.

#### **Details**

NOTE: If reads contains a \$score column, it will presume that this is the number of replicates per reads, weights for the coverage() function. So delete the score column or set weight to something else if this is not wanted.

#### Value

a numeric RleList, one numeric-Rle per group with # of hits per position. Or data.table if as.data.table is TRUE, with column names c("count" [numeric or integer], "genes" [integer], "position" [integer])

# See Also

```
Other ExtendGenomicRanges: asTX(), extendLeaders(), extendTrailers(), reduceKeepAttr(), tile1(), txSeqsFromFa(), windowPerGroup()
```

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```
end = c(5, 15, 25),
               strand = "+")
grl <- GRangesList(tx1_1 = ORF)
RFP <- GRanges("1", IRanges(25, 25), "+")</pre>
coveragePerTiling(grl, RFP, is.sorted = TRUE)
# now as data.table with frames
coveragePerTiling(grl, RFP, is.sorted = TRUE, as.data.table = TRUE,
                  withFrames = TRUE)
# With score column (usually replicated reads on that position)
RFP <- GRanges("1", IRanges(25, 25), "+", score = 5)
dt <- coveragePerTiling(grl, RFP, is.sorted = TRUE,</pre>
                         as.data.table = TRUE, withFrames = TRUE)
class(dt$count) # numeric
# With integer score column (faster and less space usage)
RFP <- GRanges("1", IRanges(25, 25), "+", score = 5L)</pre>
dt <- coveragePerTiling(grl, RFP, is.sorted = TRUE,</pre>
                         as.data.table = TRUE, withFrames = TRUE)
class(dt$count) # integer
```

coverageScorings

Add a coverage scoring scheme

## **Description**

Different scorings and groupings of a coverage representation.

## Usage

```
coverageScorings(coverage, scoring = "zscore", copy.dt = TRUE)
```

# Arguments

coverage	a data.table containing at least columns (count, position), it is possible to have additionals: (genes, fraction, feature)
scoring	a character, one of (zscore, transcriptNormalized, mean, median, sum, log2sum, log10sum, sumLength, meanPos and frameSum, periodic, NULL). More info in details
copy.dt	logical TRUE, copy object, to avoid overwriting original object. Set to false to run function using reference to object, a speed up if original object is not needed.

## **Details**

Usually output of metaWindow or scaledWindowPositions is input in this function.

Content of coverage data.table: It must contain the count and position columns.

genes column: If you have multiple windows, the genes column must define which gene/transcript grouping the different counts belong to. If there is only a meta window or only 1 gene/transcript, then this column is not needed.

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fraction column: If you have coverage of i.e RNA-seq and Ribo-seq, or TCP -seq of large and small subunite, divide into fractions. Like factor(RNA, RFP)

feature column: If gene group is subdivided into parts, like gene is transcripts, and feature column can be c(leader, cds, trailer) etc.

Given a data.table coverage of counts, add a scoring scheme. per: the grouping given, if genes is defined, group by per gene in default scoring. Scorings:

- zscore (count-mean(window))/sd(window) per) # Outlier detection
- modzscore (count-median(window))/mad(window) per) # Outlier detection for signal with extreme values
- transcriptNormalized (sum(count / sum of counts per)) this is scaled per group
- mean (mean(count per))
- median (median(count per))
- sum (count per)
- log2sum (count per)
- log10sum (count per)
- sumLength (count per) / number of windows
- meanPos (mean per position per gene) used in scaledWindowPositions
- sumPos (sum per position per gene) used in scaledWindowPositions
- frameSum (sum per frame per gene) used in ORFScore
- frameSumPerL (sum per frame per read length)
- frameSumPerLG (sum per frame per read length per gene)
- fracPos (fraction of counts per position per gene) non scaled version of transcriptNormalized
- periodic (Fourier transform periodicity of meta coverage per fraction)
- NULL (no grouping, return input directly)

#### Value

a data.table with new scores (size dependent on score used)

## See Also

Other coverage: metaWindow(), regionPerReadLength(), scaledWindowPositions(), windowPerReadLength()

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coverage\_to\_dt

Convert coverage RleList to data.table

#### Description

Convert coverage RleList to data.table

## Usage

```
coverage_to_dt(
  coverage,
  keep.names = TRUE,
  withFrames = FALSE,
  weight = "score",
  drop.zero.dt = FALSE,
  fraction = NULL
)
```

## **Arguments**

coverage RleList with names

keep.names logical (TRUE), keep names or not. If as.data.table is TRUE, names (genes

column) will be a factor column, if FALSE it will be an integer column (index of gene), so first input grl element is 1. Dropping names gives ~ 20 % speedup. If drop.zero.dt is FALSE, data.table will not return names, will use index (to

avoid memory explosion).

withFrames a logical (FALSE), only available if as.data.table is TRUE, return the ORF

frame, 1,2,3, where position 1 is 1, 2 is 2 and 4 is 1 etc.

weight (default: 'score'), if defined a character name of valid meta column in subject.

GRanges("chr1", 1, "+", score = 5), would mean score column tells that this alignment region was found 5 times. Formats which loads a score column like this: Bigwig, wig, ORFik ofst, collapsed bam, bedoc and .bedo. As do CAGEr CAGE files and many other package formats. You can also assign a score col-

umn manually.

drop.zero.dt logical FALSE, if TRUE and as.data.table is TRUE, remove all 0 count posi-

tions. This greatly speeds up and most importantly, greatly reduces memory usage. Will not change any plots, unless 0 positions are used in some sense.

(mean, median, zscore coverage will only scale differently)

fraction integer or character, a description column. Useful for grouping multiple outputs

together. If returned as Rle, this is added as: metadata(coverage) <- list(fraction

= fraction). If as.data.table it will be added as an additional column.

## Value

a data.table with column names c("count" [numeric or integer], "genes" [integer], "position" [integer])

78 covRle-class

covRle

Coverage Rlelist for both strands

# **Description**

Coverage Rlelist for both strands

# Usage

```
covRle(forward = RleList(), reverse = RleList())
```

# **Arguments**

forward a RleList with defined seqinfo for forward strand counts reverse a RleList with defined seqinfo for reverse strand counts

#### Value

a covRle object

# See Also

Other covRLE: covRle-class, covRleFromGR(), covRleList, covRleList-class

# **Examples**

```
covRle()
covRle(RleList(), RleList())
chr_rle <- RleList(chr1 = Rle(c(1,2,3), c(1,2,3)))
covRle(chr_rle, chr_rle)</pre>
```

covRle-class

Coverage Rle for both strands or single

# **Description**

Given a run of coverage(x) where x are reads, this class combines the 2 strands into 1 object

#### Value

```
a covRLE object
```

## See Also

```
Other covRLE: covRle, covRleFromGR(), covRleList, covRleList-class
```

covRleFromGR 79

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Convert GRanges to covRle

# **Description**

Convert GRanges to covRle

# Usage

```
covRleFromGR(x, weight = "AUTO", ignore.strand = FALSE)
```

# Arguments

x a GRanges, GAlignment or GAlignmentPairs object. Note that coverage cal-

culation for GAlignment is slower, so usually best to call convertToOneBase-

dRanges on GAlignment object to speed it up.

weight default "AUTO", pick 'score' column if exist, else all are 1L. Can also be a

manually assigned meta column like 'score2' etc.

ignore.strand logical, default FALSE.

# Value

covRle object

# See Also

Other covRLE: covRle, covRle-class, covRleList, covRleList-class

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covRleList

Coverage Rlelist for both strands

# Description

Coverage Rlelist for both strands

## Usage

```
covRleList(list, fraction = names(list))
```

# **Arguments**

list a list or List of covRle objects of equal length and lengths

fraction character, default names (list). Names to elements of list, can be integers, as

readlengths etc.

#### Value

a covRleList object

#### See Also

```
Other covRLE: covRle, covRle-class, covRleFromGR(), covRleList-class
```

# **Examples**

```
covRleList(List(covRle()))
```

covRleList-class

List of covRle

# **Description**

Given a run of coverage(x) where x are reads, this covRle combines the 2 strands into 1 object This list can again combine these into 1 object, with accession functions and generalizations.

## Value

```
a covRleList object
```

## See Also

```
Other covRLE: covRle, covRle-class, covRleFromGR(), covRleList
```

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create.experiment

Create an ORFik experiment

## **Description**

Create a single R object that stores and controls all results relevant to a specific Next generation sequencing experiment. Click the experiment link above in the title if you are not sure what an ORFik experiment is.

By using files in a folder / folders. It will make an experiment table with information per sample, this object allows you to use the extensive API in ORFik that works on experiments.

#### Information Auto-detection:

There will be several columns you can fill in, when creating the object, if the files have logical names like (RNA-seq\_WT\_rep1.bam) it will try to auto-detect the most likely values for the columns. Like if it is RNA-seq or Ribo-seq, Wild type or mutant, is this replicate 1 or 2 etc.

You will have to fill in the details that were not auto detected. Easiest way to fill in the blanks are in a csv editor like libre Office or excel. You can also remake the experiment and specify the specific column manually. Remember that each row (sample) must have a unique combination of values. An extra column called "reverse" is made if there are paired data, like +/- strand wig files.

# Usage

```
create.experiment(
  dir,
  exper,
  saveDir = ORFik::config()["exp"],
  txdb = "".
  fa = "",
  organism = "".
  assembly = "",
  pairedEndBam = FALSE,
  viewTemplate = FALSE,
  types = c("bam", "bed", "wig", "bigWig", "ofst"),
  libtype = "auto",
  stage = "auto",
  rep = "auto",
  condition = "auto",
  fraction = "auto",
  author = "",
  files = findLibrariesInFolder(dir, types, pairedEndBam),
  result_folder = NULL,
  runIDs = extract_run_id(files)
)
```

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#### **Arguments**

dir Which directory / directories to create experiment from, must be a directory

with NGS data from your experiment. Will include all files of file type specified by "types" argument. So do not mix files from other experiments in the same

folder!

exper Short name of experiment. Will be name used to load experiment, and name

shown when running list.experiments

saveDir Directory to save experiment csv file, default: ORFik::config()["exp"], which

has default: "~/Bio\_data/ORFik\_experiments/". Set to NULL if you don't want

to save it to disc.

txdb A path to TxDb (prefered) or gff/gtf (not adviced, slower) file with transcriptome

annotation for the organism.

fa A path to fasta genome/sequences used for libraries, remember the file must

have a fasta index too.

organism character, default: "" (no organism set), scientific name of organism. Homo

sapiens, Danio rerio, Rattus norvegicus etc. If you have a SRA metadata csv file, you can set this argument to study\$ScientificName[1], where study is the

SRA metadata for all files that was aligned.

assembly character, default: "" (no assembly set). The genome assembly name, like

GRCh38 etc. Useful to add if you want detailed metadata of experiment analy-

S1S.

pairedEndBam logical FALSE, else TRUE, or a logical list of TRUE/FALSE per library you see

will be included (run first without and check what order the files will come in) 1 paired end file, then two single will be c(T, F, F). If you have a SRA metadata csv file, you can set this argument to study\$LibraryLayout == "PAIRED", where

study is the SRA metadata for all files that was aligned.

viewTemplate run View() on template when finished, default (FALSE). Usually gives you a

better view of result than using print().

types Default c("bam", "bed", "wig", "bigWig", "ofst"), which types of libraries

to allow as NGS data.

libtype character, default "auto". Library types, must be length 1 or equal length of

number of libraries. "auto" means ORFik will try to guess from file names. Example: RFP (Ribo-seq), RNA (RNA-seq), CAGE, SSU (TCP-seq 40S), LSU

(TCP-seq 80S).

stage character, default "auto". Developmental stage, tissue or cell line, must be length

1 or equal length of number of libraries. "auto" means ORFik will try to guess from file names. Example: HEK293 (Cell line), Sphere (zebrafish stage), ovary

(Tissue).

rep character, default "auto". Replicate numbering, must be length 1 or equal length

of number of libraries. "auto" means ORFik will try to guess from file names.

Example: 1 (rep 1), 2 rep(2). Insert only numbers here!

condition character, default "auto". Library conditions, must be length 1 or equal length

of number of libraries. "auto" means ORFik will try to guess from file names.

Example: WT (wild type), mutant, etc.

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fraction character, default "auto". Fractionation of library, must be length 1 or equal length of number of libraries. "auto" means ORFik will try to guess from file names. This columns is used to make experiment unique, if the other columns are not sufficient. Example: cyto (cytosolic fraction), dmso (dmso treated fraction), etc. author character, default "". Main author of experiment, usually last name is enough. When printing will state "author et al" in info. files character vector or data.table of library paths in dir. Default: findLibrariesInFolder(dir, types, pairedEndBam). Do not touch unless you want to do some subsetting, it will automatically remove files that are not of file format defined by 'type' argument. Note that sorting on number that: 10 is before 2, so 1, 2, 10, is sorted as: 1, 10, 2. If you want to fix this, you could update this argument with: ORFik:::findLibrariesInFolder()[1,3,2] to get order back to 1,2,10 etc. result\_folder character, default NULL. The folder to output analysis results like QC, count tables etc. By default the libFolder(df) folder is used, the folder of first library in experiment. If you are making a new experiment which is a collection of other experiments, set this to a new folder, to not contaminate your other experiment directories. runIDs character ids, usually SRR, ERR, or DRR identifiers, default is to search for any

of these 3 in the filename by: extract\_run\_id(files). They are optional.

#### Value

a data.frame, NOTE: this is not a ORFik experiment, only a template for it!

## See Also

```
Other ORFik_experiment: ORFik.template.experiment(), ORFik.template.experiment.zf(), bamVarName(), experiment-class, filepath(), libraryTypes(), organism, experiment-method, outputLibs(), read.experiment(), save.experiment(), validateExperiments()
```

84 defineIsoform

```
fa = fa, organism = org,
                               viewTemplate = FALSE)
## Now fix non-unique rows: either is libre office, microsoft excel, or in R
template$X5[6] <- "heart" # here a dummy example, even though data is correct
# read experiment (if you set correctly)
df <- read.experiment(template)</pre>
## Default location of experiments is ORFik::config()["exp"]
# default_experiments_path <- ORFik::config()["exp"]</pre>
# exp_path <- file.path(default_experiments_path, paste0("exper", ".csv"))</pre>
# Save with: save.experiment(df, file = exp_path)
# Then you can simply load with read.experiment(exper),
# since you saved in the default directory
## Custom location (If you work in a team, use a shared folder)
# Remember to update ORFik::config() to ripple the effect through whole
# of ORFik if you want to use this as default
new_dir <- tempdir() # Here we just use tempdir</pre>
create.experiment(dir = dir, exper, txdb = txdb,
                  saveDir = new_dir, fa = fa, organism = org)
template_loaded <- read.experiment(exper, new_dir)</pre>
# The csv template paths (from index 5) is equal to file paths of loaded exp
identical(template$X6[-seq(4)], filepath(template_loaded, "default"))
```

defineIsoform

Overlaps GRanges object with provided annotations.

#### **Description**

Overlaps GRanges object with provided annotations.

## Usage

```
defineIsoform(
  rel_orf,
  tran,
  isoform_names = c("perfect_match", "elong_START_match", "trunc_START_match",
    "elong_STOP_match", "trunc_STOP_match", "overlap_inside", "overlap_both",
    "overlap_upstream", "overlap_downstream", "upstream", "downstram", "none")
)
```

# Arguments

```
rel_orf - GRanges object of your ORF.

tran - GRanges object of annotation (transcript or cds) that overlapped in some way rel_orf.
```

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isoform\_names

- A vector of strings that will be used instead of these defaults: 'perfect\_match' - start and stop matches the tran object strand wise 'elong\_START\_match' - rel\_orf is extension from the STOP side of the tran 'trunc\_START\_match' - rel\_orf is truncation from the STOP side of the tran 'elong\_STOP\_match' - rel\_orf is extension from the START side of the tran 'trunc\_STOP\_match' - rel\_orf is truncation from the START side of the tran 'overlap\_inside' - rel\_orf is inside tran object 'overlap\_both' - rel\_orf contains tran object inside 'overlap\_upstream' - rel\_orf is overlaping upstream part of the tran 'overlap\_downstream' - rel\_orf is overlaping downstream part of the tran 'upstream' - rel\_orf is upstream towards the tran 'downstream' - rel\_orf is downstream towards the tran 'none' - when none of the above options is true

## Value

A string object of defined isoform towards transcript.

defineTrailer

Defines trailers for ORF.

# Description

Creates GRanges object as a trailer for ORFranges representing ORF, maintaining restrictions of transcriptRanges. Assumes that ORFranges is on the transcriptRanges, strands and seqlevels are in agreement. When lengthOFtrailer is smaller than space left on the transcript than all available space is returned as trailer.

## Usage

```
defineTrailer(ORFranges, transcriptRanges, lengthOftrailer = 200)
```

## Arguments

ORFranges GRanges object of your Open Reading Frame.

transcriptRanges

GRanges object of transtript.

lengthOftrailer

Numeric. Default is 10.

#### Details

It assumes that ORFranges and transcriptRanges are not sorted when on minus strand. Should be like: (200, 600) (50, 100)

#### Value

A GRanges object of trailer.

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## See Also

```
Other ORFHelpers: longestORFs(), mapToGRanges(), orfID(), startCodons(), startSites(), stopCodons(), stopSites(), txNames(), uniqueGroups(), uniqueOrder()
```

#### **Examples**

DEG.analysis

Run differential TE analysis

# Description

Expression analysis of 1 dimension, usually between conditions of RNA-seq.

Using the standardized DESeq2 pipeline flow.

Creates a DESeq model (given x is the target.contrast argument) (usually 'condition' column)

1. RNA-seq model: design =  $\sim$  x (differences between the x groups in RNA-seq)

## Usage

```
DEG.analysis(
   df,
   target.contrast = design[1],
   design = ORFik::design(df),
   p.value = 0.05,
   counts = countTable(df, "mrna", type = "summarized"),
   batch.effect = TRUE,
   pairs = combn.pairs(unlist(df[, target.contrast]))
)
```

## **Arguments**

```
df an experiment of usually RNA-seq. target.contrast
```

a character vector, default design[1]. The column in the ORFik experiment that represent the comparison contrasts. By default: the first design factor of the full experimental design. This is the factor you will do the comparison on. DESeq will normalize the counts based on the full design, but the log fold change values will be based on this contrast only. It is usually the 'condition' column.

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significant subset for p.value = 0.05 means: TE\$padj < 0.05 & Ribo\$padj < 0.05& RNA\$padj > 0.05.

a SummarizedExperiment, default: countTable(df, "mrna", type = "summacounts

rized"), all transcripts. Assign a subset if you don't want to analyze all genes. It is recommended to not subset, to give DESeq2 data for variance analysis.

batch.effect logical, default TRUE. Makes replicate column of the experiment part of the

If you believe you might have batch effects, keep as TRUE. Batch effect usually means that you have a strong variance between biological replicates. Check out pcaExperiment and see if replicates cluster together more than the design

factor, to verify if you need to set it to TRUE.

pairs list of character pairs, the experiment contrasts. Default: combn.pairs(unlist(df.rfp[,

target.contrast])

#### **Details**

#' Analysis is done between each possible combination of levels in the target contrast If target contrast is the condition column, with factor levels: WT, mut1 and mut2 with 3 replicates each. You get comparison of WT vs mut1, WT vs mut2 and mut1 vs mut2.

The respective result categories are defined as: (given a user defined p value, shown here as 0.05): Significant - p-value adjusted < 0.05 (p-value cutoff decided by 'p.value argument)

The LFC values are shrunken by lfcShrink(type = "normal").

Remember that DESeq by default can not do global change analysis, it can only find subsets with changes in LFC!

#### Value

a data.table with columns: (contrast variable, gene id, regulation status, log fold changes, p.adjust values, mean counts)

#### References

```
https://doi.org/10.1002/cpmb.108
```

#### See Also

```
Other DifferentialExpression: DEG.plot.static(), DEG_model(), DTEG.plot(), te.table(),
te_rna.plot()
```

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## **Examples**

```
## Simple example (use ORFik template, then use only RNA-seq)
df <- ORFik.template.experiment()
df.rna <- df[df$libtype == "RNA",]
design(df.rna) # The full experimental design
design(df.rna)[1] # Default target contrast
#dt <- DEG.analysis(df.rna)</pre>
```

DEG.plot.static

Plot DEG result

# **Description**

Plot setup:

X-axis: mean counts Y-axis: Log2 fold changes For explanation of plot, see DEG. analysis

## Usage

```
DEG.plot.static(
   dt,
   output.dir = NULL,
   p.value.label = 0.05,
   plot.title = "",
   plot.ext = ".pdf",
   width = 6,
   height = 6,
   dot.size = 0.4,
   xlim = "auto",
   ylim = "bidir.max",
   relative.name = paste0("DEG_plot", plot.ext)
)
```

## **Arguments**

a data.table with the results from DEG.analysis dt a character path, default NULL(no save), or a directory to save to a file. Relative output.dir name of file, specified by 'relative.name' argument. a numeric, default ifelse(!is.null(attr(dt, "p.value")), attr(dt, "p.value"), 0.05) Inp.value.label terval (0,1), use"" to not show. What p-value used for the analysis? Will be shown as a caption. title for plots, usually name of experiment etc plot.title plot.ext character, default: ".pdf". Alternatives: ".png" or ".jpg". width numeric, default 6 (in inches) height numeric, default 6 (in inches) numeric, default 0.4, size of point dots in plot. dot.size

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#### Value

a ggplot object

## See Also

Other DifferentialExpression: DEG\_model(), DTEG.analysis(), DTEG.plot(), te.table(), te\_rna.plot()

# **Examples**

```
df <- ORFik.template.experiment()
df.rna <- df[df$libtype == "RNA",]
#dt <- DEG.analysis(df.rna)
#Default scaling
#DEG.plot.static(dt)
#Manual scaling
#DEG.plot.static(dt, xlim = c(-2, 2), ylim = c(-2, 2))</pre>
```

DEG\_model

Get DESeq2 model without running results

# Description

This is the preparation step of DESeq2 analysis using ORFik::DEG.analysis. It is exported so that you can do this step in standalone, usually you want to use DEG.analysis directly.

# Usage

```
DEG_model(
   df,
   target.contrast = design[1],
   design = ORFik::design(df),
   p.value = 0.05,
   counts = countTable(df, "mrna", type = "summarized"),
   batch.effect = TRUE
)
```

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#### **Arguments**

df an experiment of usually RNA-seq.

target.contrast

a character vector, default design[1]. The column in the ORFik experiment that represent the comparison contrasts. By default: the first design factor of the full experimental design. This is the factor you will do the comparison on. DESeq will normalize the counts based on the full design, but the log fold change values will be based on this contrast only. It is usually the 'condition' column.

design

a character vector, default design(df.rfp). The full experiment design. Which factors have more than 1 level. Example: stage column are all HEK293, so it can not be a design factor. The condition column has 2 possible values, WT and mutant, so it is a factor of the experiment. Replicates column is not part of design, that is inserted later with setting batch.effect = TRUE. Library type 'libtype' column, can also no be part of initial design, it is always added inside the function, after initial setup.

p.value

counts

a numeric, default 0.05 in interval (0,1). Defines adjusted p-value to be used as significance threshold for the result groups. I.e. for exclusive translation group significant subset for p.value = 0.05 means: TE\$padj < 0.05 & Ribo\$padj < 0.05 & RNA\$padj > 0.05.

a SummarizedExperiment, default: countTable(df, "mrna", type = "summarized"), all transcripts. Assign a subset if you don't want to analyze all genes. It is recommended to not subset, to give DESeq2 data for variance analysis.

batch.effect

logical, default TRUE. Makes replicate column of the experiment part of the

design.

If you believe you might have batch effects, keep as TRUE. Batch effect usually means that you have a strong variance between biological replicates. Check out pcaExperiment and see if replicates cluster together more than the design factor, to verify if you need to set it to TRUE.

#### Value

a DESeqDataSet object with results stored as metadata columns.

# See Also

```
Other DifferentialExpression: DEG.plot.static(), DTEG.analysis(), DTEG.plot(), te.table(), te_rna.plot()
```

```
## Simple example (use ORFik template, then use only RNA-seq)
df <- ORFik.template.experiment()
df.rna <- df[df$libtype == "RNA",]
design(df.rna) # The full experimental design
target.contrast <- design(df.rna)[1] # Default target contrast
#ddsMat_rna <- DEG_model(df.rna, target.contrast)</pre>
```

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DEG\_model\_results

Get DESeq2 model results from DESeqDataSet

## **Description**

Get DESeq2 model results from DESeqDataSet

# Usage

```
DEG_model_results(ddsMat_rna, target.contrast, pairs, p.value = 0.05)
```

## **Arguments**

ddsMat\_rna a DESeqDataSet object with results stored as metadata columns.

target.contrast

a character vector, default design[1]. The column in the ORFik experiment that represent the comparison contrasts. By default: the first design factor of the full experimental design. This is the factor you will do the comparison on. DESeq will normalize the counts based on the full design, but the log fold change values

will be based on this contrast only. It is usually the 'condition' column.

pairs list of character pairs, the experiment contrasts. Default: combn.pairs(unlist(df.rfp[,

target.contrast])

p.value a numeric, default 0.05 in interval (0,1). Defines adjusted p-value to be used as

significance threshold for the result groups. I.e. for exclusive translation group significant subset for p.value = 0.05 means: TE\$padj < 0.05 & Ribo\$padj < 0.05

& RNA\$padj > 0.05.

#### Value

a data.table

```
## Simple example (use ORFik template, then use only RNA-seq)
df <- ORFik.template.experiment()
df.rna <- df[df$libtype == "RNA",]
design(df.rna) # The full experimental design
target.contrast <- design(df.rna)[1] # Default target contrast
#ddsMat_rna <- DEG_model(df.rna, target.contrast)
#pairs <- combn.pairs(unlist(df[, target.contrast]))
#dt <- DEG_model_results(ddsMat_rna, target.contrast, pairs)</pre>
```

DEG\_model\_simple

Simple Fpkm ratio test DEG

## Description

If you do not have a valid DESEQ2 experimental setup (contrast), you can use this simplified test

## Usage

```
DEG_model_simple(
  df,
  target.contrast = design[1],
  design = ORFik::design(df),
  p.value = 0.05,
  counts = countTable(df, "mrna", type = "summarized"),
  batch.effect = FALSE
)
```

#### **Arguments**

an experiment of usually RNA-seq. target.contrast

> a character vector, default design[1]. The column in the ORFik experiment that represent the comparison contrasts. By default: the first design factor of the full experimental design. This is the factor you will do the comparison on. DESeq will normalize the counts based on the full design, but the log fold change values will be based on this contrast only. It is usually the 'condition' column.

a character vector, default design(df.rfp). The full experiment design. Which design

factors have more than 1 level. Example: stage column are all HEK293, so it can not be a design factor. The condition column has 2 possible values, WT and mutant, so it is a factor of the experiment. Replicates column is not part of design, that is inserted later with setting batch. effect = TRUE. Library type 'libtype' column, can also no be part of initial design, it is always added inside

the function, after initial setup.

a numeric, default 0.05 in interval (0,1). Defines adjusted p-value to be used as

significance threshold for the result groups. I.e. for exclusive translation group significant subset for p.value = 0.05 means: TE\$padj < 0.05 & Ribo\$padj < 0.05

& RNA\$padj > 0.05.

a SummarizedExperiment, default: countTable(df, "mrna", type = "summacounts

rized"), all transcripts. Assign a subset if you don't want to analyze all genes. It is recommended to not subset, to give DESeq2 data for variance analysis.

logical, default TRUE. Makes replicate column of the experiment part of the design.

If you believe you might have batch effects, keep as TRUE. Batch effect usually

means that you have a strong variance between biological replicates. Check out pcaExperiment and see if replicates cluster together more than the design

factor, to verify if you need to set it to TRUE.

p.value

batch.effect

#### Value

a data.table of fpkm ratios

#### **Examples**

```
## Simple example (use ORFik template, then use only RNA-seq)
df <- ORFik.template.experiment()
df <- df[df$libtype == "RNA",]
#dt <- DEG_model_simple(df)</pre>
```

design, experiment-method

Get experimental design Find the column/columns that create a separation between samples, by default skips replicate and choose first that is from either: libtype, condition, stage and fraction.

# Description

Get experimental design Find the column/columns that create a separation between samples, by default skips replicate and choose first that is from either: libtype, condition, stage and fraction.

## Usage

```
## S4 method for signature 'experiment'
design(
  object,
  batch.correction.design = FALSE,
  as.formula = FALSE,
  multi.factor = TRUE
)
```

# Arguments

```
object an ORFik experiment batch.correction.design
```

logical, default FALSE. If true, add replicate as a second design factor (only if

>= 2 replicates exists).

as.formula logical, default FALSE. If TRUE, return as formula

multi.factor logical, default TRUE If FALSE, return first factor only (+ rep, if batch.correction.design

is true). Order of picking is: libtype, if not then: stage, if not then: condition, if

not then: fraction.

#### Value

a character (name of column) or a formula

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## **Examples**

```
df <- ORFik.template.experiment()
design(df) # The 2 columns that decides the design here
# If we subset it changes
design(df[df$libtype == "RFP",])
# Only single factor design, it picks first
design(df, multi.factor = FALSE)</pre>
```

# **Description**

Utilizes periodicity measurement (Fourier transform), and change point analysis to detect ribosomal footprint shifts for each of the ribosomal read lengths. Returns subset of read lengths and their shifts for which top covered transcripts follow periodicity measure. Each shift value assumes 5' anchoring of the reads, so that output offsets values will shift 5' anchored footprints to be on the p-site of the ribosome. The E-site will be shift + 3 and A site will be shift - 3. So update to these, if you rather want those.

## Usage

```
detectRibosomeShifts(
  footprints,
  txdb,
  start = TRUE,
  stop = FALSE,
  top_tx = 10L
 minFiveUTR = 30L,
 minCDS = 150L,
 minThreeUTR = if (stop) {
     30
 } else NULL,
  txNames = filterTranscripts(txdb, minFiveUTR, minCDS, minThreeUTR),
  firstN = 150L,
  tx = NULL,
 min_reads = 1000,
 min_reads_TIS = 50,
  accepted.lengths = 26:34,
 heatmap = FALSE,
 must.be.periodic = TRUE,
 strict.fft = TRUE,
  verbose = FALSE
)
```

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#### **Arguments**

GAlignments object of RiboSeq reads - footprints, can also be path to the .bam footprints /.ofst file. If GAlignment object has a meta column called "score", this will be used as replicate numbering for that read. So be careful if you have custom files with score columns, with another meaning. txdb a TxDb file, a path to one of: (.gtf,.gff, .gff2, .gff2, .db or .sqlite) or an ORFik experiment (logical) Whether to include predictions based on the start codons. Default start TRUE. stop (logical) Whether to include predictions based on the stop codons. Default FASLE. Only use if there exists 3' UTRs for the annotation. If peridicity around stop codon is stronger than at the start codon, use stop instead of start region for p-shifting. (integer), default 10. Specify which % of the top TIS coverage transcripts to use top\_tx for estimation of the shifts. By default we take top 10 top covered transcripts as they represent less noisy data-set. This is only applicable when there are more than 1000 transcripts. minFiveUTR (integer) minimum bp for 5' UTR during filtering for the transcripts. Set to NULL if no 5' UTRs exists for annotation. minCDS (integer) minimum bp for CDS during filtering for the transcripts (integer) minimum bp for 3' UTR during filtering for the transcripts. Set to minThreeUTR NULL if no 3' UTRs exists for annotation. txNames a character vector of subset of CDS to use. Default: txNames = filterTranscripts(txdb, minFiveUTR, minCDS, minThreeUTR) Example: c("ENST1000005"), will use only that transcript (You should use at least 100!). Remember that top\_tx argument, will by default specify to use top 10 % of those CDSs. Set that to 100, to use all these specified transcripts. firstN (integer) Represents how many bases of the transcripts downstream of start codons to use for initial estimation of the periodicity. a GRangesList, if you do not have 5' UTRs in annotation, send your own vertx sion. Example: extendLeaders(tx, 30) Where 30 bases will be new "leaders". Since each original transcript was either only CDS or non-coding (filtered out). default (1000), how many reads must a read-length have in total to be considered min\_reads for periodicity. min\_reads\_TIS default (50), how many reads must a read-length have in the TIS region to be considered for periodicity. accepted.lengths accepted read lengths, default 26:34, usually ribo-seq is strongest between 27:32. heatmap a logical or character string, default FALSE. If TRUE, will plot heatmap of raw reads before p-shifting to console, to see if shifts given make sense. You can

also set a filepath to save the file there.

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must.be.periodic

logical TRUE, if FALSE will not filter on periodic read lengths. (The Fourier transform filter will be skipped). This is useful if you are not going to do periodicity analysis, that is: for you more coverage depth (more read lengths) is more important than only keeping the high quality periodic read lengths.

strict.fft

logical, TRUE. Use a FFT without noise filter. This means keep only reads lengths that are "periodic for the human eye". If you want more coverage, set to FALSE, to also get read lengths that are "messy", but the noise filter detects the periodicity of 3. This should only be done when you do not need high quality periodic reads! Example would be differential translation analysis by counts over each ORF.

verbose

logical, default FALSE. Report details of analysis/periodogram. Good if you are not sure if the analysis was correct.

#### **Details**

Check out vignette for the examples of plotting RiboSeq metaplots over start and stop codons, so that you can verify visually whether this function detects correct shifts.

For how the Fourier transform works, see: isPeriodic
For how the changepoint analysis works, see: changePointAnalysis

NOTE: It will remove softclips from valid width, the CIGAR 3S30M is qwidth 33, but will remove 3S so final read width is 30 in ORFik. This is standard for ribo-seq.

#### Value

a data.table with lengths of footprints and their predicted coresponding offsets

#### References

https://bmcgenomics.biomedcentral.com/articles/10.1186/s12864-018-4912-6

#### See Also

```
Other pshifting: changePointAnalysis(), shiftFootprints(), shiftFootprintsByExperiment(), shiftPlots(), shifts_load(), shifts_save()
```

```
## Basic run
# Transcriptome annotation ->
gtf_file <- system.file("extdata/references/danio_rerio", "annotations.gtf", package = "ORFik")
# Ribo seq data ->
riboSeq_file <- system.file("extdata/Danio_rerio_sample", "ribo-seq.bam", package = "ORFik")
## Not run:
footprints <- readBam(riboSeq_file)
## Using CDS start site as reference point:
detectRibosomeShifts(footprints, gtf_file)
## Using CDS start site and stop site as 2 reference points:</pre>
```

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```
#detectRibosomeShifts(footprints, gtf_file, stop = TRUE)
## Debug and detailed information for accepted reads lengths and p-site:
detectRibosomeShifts(footprints, gtf_file, heatmap = TRUE, verbose = TRUE)
## Debug why read length 31 was not accepted or wrong p-site:
#detectRibosomeShifts(footprints, gtf_file, must.be.periodic = FALSE,
               accepted.lengths = 31, heatmap = TRUE, verbose = TRUE)
## Subset bam file
param = ScanBamParam(flag = scanBamFlag(
                       isDuplicate = FALSE,
                       isSecondaryAlignment = FALSE))
footprints <- readBam(riboSeq_file, param = param)</pre>
detectRibosomeShifts(footprints, gtf_file, stop = TRUE)
## Without 5' Annotation
library(GenomicFeatures)
txdb <- loadTxdb(gtf_file)</pre>
tx <- exonsBy(txdb, by = "tx", use.names = TRUE)
tx <- extendLeaders(tx, 30)</pre>
## Now run function, without 5' and 3' UTRs
detectRibosomeShifts(footprints, txdb, start = TRUE, minFiveUTR = NULL,
                     minCDS = 150L, minThreeUTR = NULL, firstN = 150L,
                     tx = tx)
## End(Not run)
```

detect\_ribo\_orfs

Detect ORFs by Ribosome profiling data

## Description

Finding all ORFs: 1. Find all ORFs in mRNA using ORFik findORFs, with defined parameters.

To create the candidate ORFs (all ORFs returned):

Steps (candidate set):

Define a candidate search set by these 3 rules:

1.a Allowed ORF type: uORF, NTE, etc (only keep these in candidate list)

- 1.b Must have at least x reads over whole orf (default 10 reads)
- 1.c Must have at least x reads over start site (default 3 reads)

The total list is defined by these names, and saved according to allowed ORF type/types.

To create the prediction status (TRUE/FALSE) per candidate

Steps (prediction status)

(UP\_NT is a 20nt window upstream of ORF, that stops 2NT before ORF starts) :

- 1. ORF mean reads per NT > (UP\_NT mean reads per NT \* 1.3)
- 2. ORFScore > 2.5
- 3. TIS total reads + 3 > ORF median reads per NT
- 4. Given expression above, a TRUE prediction is defined with the AND operation: 1. & 2. & 3.

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```
In code that is:
predicted <- (orfs_cov_stats$mean > upstream_cov_stats$mean*1.3) & orfs_cov_stats$ORFScores
> 2.5 & ((reads_start[candidates] + 3) > orfs_cov_stats$median)
```

## Usage

```
detect_ribo_orfs(
  df,
  out_folder,
  ORF_categories_to_keep,
 prefix_result = paste(c(ORF_categories_to_keep, gsub(" ", "_", organism(df))), collapse
 mrna = loadRegion(df, "mrna"),
 cds = loadRegion(df, "cds"),
 libraries = outputLibs(df, type = "pshifted", output.mode = "envirlist"),
 orf_candidate_ranges = findORFs(seqs = txSeqsFromFa(mrna, df, TRUE), longestORF =
    longestORF, startCodon = startCodon, stopCodon = stopCodon, minimumLength =
    minimumLength),
 orfs_gr = categorize_and_filter_ORFs(orf_candidate_ranges, ORF_categories_to_keep, cds,
    mrna),
  export_metrics_table = TRUE,
  longestORF = FALSE,
  startCodon = startDefinition(1),
  stopCodon = stopDefinition(1),
 minimumLength = 0,
 minimum_reads_ORF = 10,
 minimum_reads_start = 3
)
```

#### **Arguments**

- uORF: Upstream ORFs (Starting in 5' UTR), not overlapping CDS
- uoORF: Upstream ORFs (Starting in 5' UTR), overlapping CDS
- annotated: The defined CDS for that transcript
- NTE: 5' Start codon extension of annotated CDS
- NTT: 5' Start codon truncation of annotated CDS
- CTE: 3' stop codon extension of annotated CDS, i.e. readthrough
- CTT: 5' Start codon truncation of annotated CDS, original cds was defined with readthrough
- internal : Starting inside CDS, ending before CDS ends
- doORF: Downstream ORFs (Ending in 3' UTR), overlapping CDS

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```
• dORF: Downstream ORFs (Ending in 3' UTR), not overlapping CDS
                    • a error : Any ORF detect not in the above categories
prefix_result
                 the prefix name of output files to out_folder. Default: paste(c(ORF_categories_to_keep,
                 gsub(" ", "_", organism(df))), collapse = "_")
                 = loadRegion(df, "mrna")
mrna
                 = loadRegion(df, "cds")
cds
libraries
                 the ribo-seq libraries loaded into R as list, default: outputLibs(df, type =
                  "pshifted", output.mode = "envirlist")
orf_candidate_ranges
                  IRangesList, = findORFs(seqs = txSeqsFromFa(mrna, df, TRUE),longestORF
                 = longestORF, startCodon = startCodon, stopCodon = stopCodon, minimumLength
                 = minimumLength)
                 = categorize_and_filter_ORFs(orf_candidate_ranges, ORF_categories_to_keep,
orfs_gr
                 cds, mrna). The GRangesList set of ORFs to actually search.
export_metrics_table
                 logical, default TRUE. Export table of statistics to file with suffix: " predic-
                  tion table.rds"
longestORF
                  (logical) Default TRUE. Keep only the longest ORF per unique stopcodon: (se-
                 qname, strand, stopcodon) combination, Note: Not longest per transcript! You
                  can also use function longestORFs after creation of ORFs for same result.
                 (character vector) Possible START codons to search for. Check startDefinition
startCodon
                  for helper function. Note that it is case sensitive, so "atg" would give 0 hits for
                  a sequence with only capital "ATG" ORFs.
stopCodon
                  (character vector) Possible STOP codons to search for. Check stopDefinition
                 for helper function. Note that it is case sensitive, so "tga" would give 0 hits for
                 a sequence with only capital "TGA" ORFs.
                 (integer) Default is 0. Which is START + STOP = 6 bp. Minimum length
minimumLength
                  of ORF, without counting 3bps for START and STOP codons. For example
                 minimumLength = 8 will result in size of ORFs to be at least START + 8*3 (bp)
                 + STOP = 30 bases. Use this param to restrict search.
minimum_reads_ORF
                 numeric, default 10, orf removed if less reads overlap whole orf
minimum_reads_start
                 numeric, default 3, orf removed if less reads overlap start
```

#### Value

invisible(NULL), all ORF results saved to disc

```
# Pre requisites
# 1. Create ORFik experiment
# ORFik::create.experiment(...)
# 2. Create ORFik optimized annotation:
# makeTxdbFromGenome(gtf = ORFik:::getGtfPathFromTxdb(df), genome = df@fafile,
```

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```
organism = organism(df), optimize = TRUE)
# 3. There must exist pshifted reads, either as default files, or in a relative folder called
# "./pshifted/". See ?shiftFootprintsByExperiment
# EXAMPLE:
df <- ORFik.template.experiment()</pre>
df \leftarrow df[df$libtype == "RFP",][c(1,2),]
result_folder <- riboORFsFolder(df, tempdir())</pre>
results <- detect_ribo_orfs(df, result_folder, c("uORF", "uoORF", "annotated", "NTE"))
# Load results of annotated ORFs
table <- riboORFs(df[1,], type = "table", result_folder)</pre>
table # See all statistics
sum(table$predicted) # How many were predicted as Ribo-seq ORFs
# Load 2 results
table <- riboORFs(df[1:2,], type = "table", result_folder)</pre>
table # See all statistics
sum(table$predicted) # How many were predicted as Ribo-seq ORFs
# Load GRangesList
candidates_gr <- riboORFs(df[1,], type = "ranges_candidates", result_folder)</pre>
prediction <- riboORFs(df[1,], type = "predictions", result_folder)</pre>
predicted_gr <- riboORFs(df[1:2,], type = "ranges_predictions", result_folder)</pre>
identical(predicted_gr[[1]], candidates_gr[[1]][prediction[[1]]])
## Inspect predictions in RiboCrypt
# library(RiboCrypt)
# Inspect Predicted
view <- predicted_gr[[1]][1]</pre>
#multiOmicsPlot_ORFikExp(view, df, view, leader_extension = 100, trailer_extension = 100)
# Inspect not predicted
view <- candidates_gr[[1]][!prediction[[1]]][1]</pre>
#multiOmicsPlot_ORFikExp(view, df, view, leader_extension = 100, trailer_extension = 100)
```

disengagementScore

Disengagement score (DS)

# **Description**

Disengagement score is defined as

```
(RPFs over ORF)/(RPFs downstream to transcript end)
```

A pseudo-count of one is added to both the ORF and downstream sums.

# Usage

```
disengagementScore(
  grl,
  RFP,
  Gtf0rTx,
```

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```
RFP.sorted = FALSE,
weight = 1L,
overlapGrl = NULL
)
```

## **Arguments**

grl a GRangesList object with usually either leaders, cds', 3' utrs or ORFs.

RFP RiboSeq reads as GAlignments, GRanges or GRangesList object

Gtf0rTx If it is TxDb object transcripts will be extracted using exonsBy(Gtf, by = "tx",

use.names = TRUE). Else it must be GRangesList

RFP.sorted logical (FALSE), an optimizer, have you ran this line: RFP <- sort(RFP[countOverlaps(RFP,

tx, type = "within") > 0]) Normally not touched, for internal optimization

purposes.

weight a numeric/integer vector or metacolumn name. (default: 1L, no differential

weighting). If weight is name of defined meta column in reads object, it gives the number of times a read was found at that position. GRanges("chr1", 1, "+", score = 5), would mean "score" column tells that this alignment region was found 5 times. if 1L it means each read is weighted equal as 1, this is what among others countOverlaps() presumes, if single number (!= 1), it repeats for all ranges, if vector with length > 1, it must be equal size of the reads object.

overlapGrl an integer, (default: NULL), if defined must be countOverlaps(grl, RFP), added

for speed if you already have it

#### Value

a named vector of numeric values of scores

#### References

```
doi: 10.1242/dev.098344
```

# See Also

```
Other features: computeFeatures(), computeFeaturesCage(), countOverlapsW(), distToCds(), distToTSS(), entropy(), floss(), fpkm(), fpkm_calc(), fractionLength(), initiationScore(), insideOutsideORF(), isInFrame(), isOverlapping(), kozakSequenceScore(), orfScore(), rankOrder(), ribosomeReleaseScore(), ribosomeStallingScore(), startRegion(), startRegionCoverage(), stopRegion(), subsetCoverage(), translationalEff()
```

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distanceToFollowing

Distance to following range group

# Description

Follow means downstream GRangesList element, not including itself.

# Usage

```
distanceToFollowing(grl, grl2 = grl, ignore.strand = FALSE)
```

# **Arguments**

grl a GRangesList

gr12 a GRangesList, default 'grl'. The list that defines restrictions on extension. Can

also be another set, which is used as 'roadblocks' for extension.

ignore.strand logical, default FALSE

#### Value

numeric vector of distance

distanceToPreceding

Distance to preceding range group

# Description

Preceding means upstream GRangesList element, not including itself.

# Usage

```
distanceToPreceding(grl, grl2 = grl, ignore.strand = FALSE)
```

# **Arguments**

grl a GRangesList

grl2 a GRangesList, default 'grl'. The list that defines restrictions on extension. Can

also be another set, which is used as 'roadblocks' for extension.

ignore.strand logical, default FALSE

# Value

numeric vector of distance

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distToCds

Get distances between ORF ends and starts of their transcripts cds.

# **Description**

Will calculate distance between each ORF end and begining of the corresponding cds (main ORF). Matching is done by transcript names. This is applicable practically to the upstream (fiveUTRs) ORFs only. The cds start site, will be presumed to be on + 1 of end of fiveUTRs.

## Usage

```
distToCds(ORFs, fiveUTRs, cds = NULL)
```

# **Arguments**

ORFs orfs as GRangesList, names of orfs must be transcript names

fiveUTRs as GRangesList, remember to use CAGE version of 5' if you did

CAGE reassignment!

cds cds' as GRangesList, only add if you have ORFs going into CDS.

#### Value

an integer vector, +1 means one base upstream of cds, -1 means 2nd base in cds, 0 means orf stops at cds start.

## References

```
doi: 10.1074/jbc.R116.733899
```

## See Also

```
Other features: computeFeatures(), computeFeaturesCage(), countOverlapsW(), disengagementScore(), distToTSS(), entropy(), floss(), fpkm(), fpkm_calc(), fractionLength(), initiationScore(), insideOutsideORF(), isInFrame(), isOverlapping(), kozakSequenceScore(), orfScore(), rankOrder(), ribosomeReleaseScore(), ribosomeStallingScore(), startRegion(), startRegionCoverage(), stopRegion(), subsetCoverage(), translationalEff()
```

```
grl <- GRangesList(tx1_1 = GRanges("1", IRanges(1, 10), "+"))
fiveUTRs <- GRangesList(tx1 = GRanges("1", IRanges(1, 20), "+"))
distToCds(grl, fiveUTRs)</pre>
```

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distToTSS

Get distances between ORF Start and TSS of its transcript

# Description

Matching is done by transcript names. This is applicable practically to any region in Transcript If ORF is not within specified search space in tx, this function will crash.

## Usage

```
distToTSS(ORFs, tx)
```

## **Arguments**

```
ORFs orfs as GRangesList, names of orfs must be txname_[rank]
tx transcripts as GRangesList.
```

## Value

an integer vector, 1 means on TSS, 2 means second base of Tx.

## References

```
doi: 10.1074/jbc.R116.733899
```

# See Also

```
Other features: computeFeatures(), computeFeaturesCage(), countOverlapsW(), disengagementScore(), distToCds(), entropy(), floss(), fpkm(), fpkm_calc(), fractionLength(), initiationScore(), insideOutsideORF(), isInFrame(), isOverlapping(), kozakSequenceScore(), orfScore(), rankOrder(), ribosomeReleaseScore(), ribosomeStallingScore(), startRegion(), startRegionCoverage(), stopRegion(), subsetCoverage(), translationalEff()
```

```
grl <- GRangesList(tx1_1 = GRanges("1", IRanges(5, 10), "+"))
tx <- GRangesList(tx1 = GRanges("1", IRanges(2, 20), "+"))
distToTSS(grl, tx)</pre>
```

download.ebi 105

download.ebi

Faster download of fastq files

## **Description**

Uses ftp download from vol1 drive on EBI ftp server, for faster download of ERR, SRR or DRR files. But does not support subsetting or custom settings of files!

## Usage

```
download.ebi(
  info,
  outdir,
  rename = TRUE,
  ebiDLMethod = "auto",
  timeout = 5000,
  BPPARAM = bpparam()
)
```

## **Arguments**

1-
1

formation including the SRR numbers in a column called "Run" or "SRR". Can be SRR, ERR or DRR numbers. If only SRR numbers can not rename, since no

additional information is given.

outdir directory to store runs, files are named by default (rename = TRUE) by informa-

tion from SRA metadata table, if (rename = FALSE) named according to SRR

numbers.

rename logical or character, default TRUE (Auto guess new names). False: Skip renam-

ing. A character vector of equal size as files wanted can also be given. Priority of renaming from the metadata is to check for unique names in the Library-Name column, then the sample\_title column if no valid names in LibraryName. If new names found and still duplicates, will add "\_rep1", "\_rep2" to make them unique. If no valid names, will not rename, that is keep the SRR numbers, you

then can manually rename files to something more meaningful.

ebiDLMethod character, default "auto". Which download protocol to use in download.file

when using ebi ftp download. Sometimes "curl" is might not work (the default auto usually), in those cases use wget. See "method" argument of ?down-

load.file, for more info.

timeout 5000, how many seconds before killing download if still active? Will overwrite

global option until R session is closed. Increase value if you are on a very slow

connection or downloading a large dataset.

BPPARAM how many cores/threads to use? default: bpparam(). To see number of threads

used, do bpparam()\$workers

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#### Value

character, full filepath of downloaded files

#### See Also

```
Other sra: browseSRA(), download.SRA(), download.SRA.metadata(), get_bioproject_candidates(), install.sratoolkit(), rename.SRA.files()
```

download.SRA

Download read libraries from SRA

## **Description**

Multicore version download, see documentation for SRA toolkit for more information.

## Usage

```
download.SRA(
  info,
  outdir,
  rename = TRUE,
  fastq.dump.path = install.sratoolkit(),
  settings = paste("--skip-technical", "--split-files"),
  subset = NULL,
  compress = TRUE,
  use.ebi.ftp = is.null(subset),
  ebiDLMethod = "auto",
  timeout = 5000,
  BPPARAM = bpparam()
)
```

# Arguments

info

character vector of only SRR numbers or a data.frame with SRA metadata information including the SRR numbers in a column called "Run" or "SRR". Can be SRR, ERR or DRR numbers. If only SRR numbers can not rename, since no additional information is given.

outdir

directory to store runs, files are named by default (rename = TRUE) by information from SRA metadata table, if (rename = FALSE) named according to SRR numbers.

rename

logical or character, default TRUE (Auto guess new names). False: Skip renaming. A character vector of equal size as files wanted can also be given. Priority of renaming from the metadata is to check for unique names in the Library-Name column, then the sample\_title column if no valid names in LibraryName. If new names found and still duplicates, will add "\_rep1", "\_rep2" to make them unique. If no valid names, will not rename, that is keep the SRR numbers, you then can manually rename files to something more meaningful.

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fastq.dump.path

path to fastq-dump binary, default: path returned from install.sratoolkit()

settings a string of arguments for fastq-dump, default: paste("-gzip", "-skip-technical",

"-split-files")

subset an integer or NULL, default NULL (no subset). If defined as a integer will

download only the first n reads specified by subset. If subset is defined, will

force to use fastq-dump which is slower than ebi download.

compress logical, default TRUE. Download compressed files ".gz".

use.ebi.ftp logical, default: is.null(subset). Use ORFiks much faster download function

that only works when subset is null, if subset is defined, it uses fastqdump, it is slower but supports subsetting. Force it to use fastqdump by setting this to

FALSE.

ebiDLMethod character, default "auto". Which download protocol to use in download.file

when using ebi ftp download. Sometimes "curl" is might not work (the default auto usually), in those cases use wget. See "method" argument of ?down-

load.file, for more info.

timeout 5000, how many seconds before killing download if still active? Will overwrite

global option until R session is closed. Increase value if you are on a very slow

connection or downloading a large dataset.

BPPARAM how many cores/threads to use? default: bpparam(). To see number of threads

used, do bpparam()\$workers

#### Value

a character vector of download files filepaths

#### References

https://ncbi.github.io/sra-tools/fastq-dump.html

# See Also

```
Other sra: browseSRA(), download.SRA.metadata(), download.ebi(), get_bioproject_candidates(), install.sratoolkit(), rename.SRA.files()
```

```
SRR <- c("SRR453566") # Can be more than one

## Simple single SRR run of YEAST
outdir <- tempdir() # Specify output directory
# Download, get 5 first reads
#download.SRA(SRR, outdir, rename = FALSE, subset = 5)

## Using metadata column to get SRR numbers and to be able to rename samples
outdir <- tempdir() # Specify output directory
info <- download.SRA.metadata("SRP226389", outdir) # By study id
## Download, 5 first reads of each library and rename</pre>
```

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```
#files <- download.SRA(info, outdir, subset = 5)
#Biostrings::readDNAStringSet(files[1], format = "fastq")
## Download full libraries of experiment
## (note, this will take some time to download!)
#download.SRA(info, outdir)</pre>
```

download. SRA. metadata Downloads metadata from SRA

## **Description**

Given a experiment identifier, query information from different locations of SRA to get a complete metadata table of the experiment. It first finds Runinfo for each library, then sample info, if pubmed id is not found searches for that and searches for author through pubmed.

#### **Usage**

```
download.SRA.metadata(
    SRP,
    outdir = tempdir(),
    remove.invalid = TRUE,
    auto.detect = FALSE,
    abstract = "printsave",
    force = FALSE,
    rich.format = FALSE,
    fetch_GSE = FALSE
)
```

#### **Arguments**

SRP

character string, a study ID as either the PRJ, SRP, ERP, DRP, GSE or SRA of the study, examples would be "SRP226389" or "ERP116106". If GSE it will try to convert to the SRP to find the files. The call works as long the runs are registered on the efetch server, as their is a linked SRP link from bioproject or GSE. Example which fails is "PRJNA449388", which does not have a linking like this.

outdir

character string, directory to save file, default: tempdir(). The file will be called "SraRunInfo\_SRP.csv", where SRP is the SRP argument. We advice to use bioproject IDs "PRJNA...". The directory will be created if not existing.

remove.invalid logical, default TRUE. Remove Runs with 0 reads (spots)

auto.detect logical,

logical, default FALSE. If TRUE, ORFik will add additional columns:

LIBRARYTYPE: (is this Ribo-seq or mRNA-seq, CAGE etc),

REPLICATE: (is this replicate 1, 2 etc),

STAGE: (Which time point, cell line or tissue is this, HEK293, TCP-1, 24hpf

etc),

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CONDITION: (is this Wild type control or a mutant etc).

These values are only qualified guesses from the metadata, so always double

check!

abstract character, default "printsave". If abstract for project exists, print and save it (save

the file to same directory as runinfo). Alternatives: "print", Only print first time

downloaded, will not be able to print later.

save" save it, no print

"no" skip download of abstract

force logical, default FALSE. If TRUE, will redownload all files needed even though

they exists. Useuful if you wanted auto.detection, but already downloaded with-

out it.

rich.format logical, default FALSE. If TRUE, will fetch all Experiment and Sample at-

tributes. It means, that different studies can have different set of columns if

set to TRUE.

fetch\_GSE logical, default FALSE. Search for GSE, if exists, appends a column called

GEO. Will be included even though this study is not from GEO, then it sets

all to NA.

#### **Details**

A common problem is that the project is not linked to an article, you will then not get a pubmed id.

The algorithm works like this:

If GEO identifier, find the SRP.

Then search Entrez for project and get sample identifier.

From that extract the run information and collect into a final table.

#### Value

a data.table of the metadata, 1 row per sample, SRR run number defined in 'Run' column.

### References

```
doi: 10.1093/nar/gkq1019
```

#### See Also

```
Other sra: browseSRA(), download.SRA(), download.ebi(), get_bioproject_candidates(), install.sratoolkit(), rename.SRA.files()
```

#### **Examples**

```
## Originally on SRA
download.SRA.metadata("SRP226389")
## Now try with auto detection (guessing additional library info)
## Need to specify output dir as tempfile() to re-download
#download.SRA.metadata("SRP226389", tempfile(), auto.detect = TRUE)
## Originally on ENA (RCP-seq data)
# download.SRA.metadata("ERP116106")
```

```
## Originally on GEO (GSE) (save to directory to keep info with fastq files)
# download.SRA.metadata("GSE61011")
## Bioproject ID
# download.SRA.metadata("PRJNA231536")
```

downstreamFromPerGroup

Get rest of objects downstream (inclusive)

## Description

Per group get the part downstream of position. downstreamFromPerGroup(tx, startSites(threeUTRs, asGR = TRUE)) will return the 3' utrs per transcript as GRangesList, usually used for interesting parts of the transcripts.

### Usage

```
downstreamFromPerGroup(
   tx,
   downstreamFrom,
   is.circular = all(isCircular(tx) %in% TRUE)
)
```

#### **Arguments**

tx a GRangesList, usually of Transcripts to be changed

downstreamFrom a vector of integers, for each group in tx, where is the new start point of first

valid exon.

is.circular logical, default FALSE if not any is: all(isCircular(grl) Where grl is the ranges

checked. If TRUE, allow ranges to extend below position 1 on chromosome.

Since circular genomes can have negative coordinates.

#### **Details**

If you don't want to include the points given in the region, use downstreamOfPerGroup

## Value

a GRangesList of downstream part

#### See Also

Other GRanges: assignFirstExonsStartSite(), assignLastExonsStopSite(), downstreamOfPerGroup(), upstreamFromPerGroup(), upstreamOfPerGroup()

downstreamN 111

downstreamN Restrict GRangesList

### **Description**

Will restrict GRangesList to 'N' bp downstream from the first base.

### Usage

```
downstreamN(grl, firstN = 150L)
```

### **Arguments**

grl (GRangesList)

firstN (integer) Allow only this many bp downstream, maximum.

#### Value

a GRangesList of reads restricted to firstN and tiled by 1

downstreamOfPerGroup Get rest of objects downstream (exclusive)

### **Description**

Per group get the part downstream of position. downstreamOfPerGroup(tx, stopSites(cds, asGR = TRUE)) will return the 3' utrs per transcript as GRangesList, usually used for interesting parts of the transcripts.

### Usage

```
downstreamOfPerGroup(tx, downstreamOf)
```

### **Arguments**

tx a GRangesList, usually of Transcripts to be changed

downstreamOf a vector of integers, for each group in tx, where is the new start point of first

valid exon. Can also be a GRangesList, then stopsites will be used.

#### **Details**

If you want to include the points given in the region, use downstreamFromPerGroup

#### Value

a GRangesList of downstream part

#### See Also

Other GRanges: assignFirstExonsStartSite(), assignLastExonsStopSite(), downstreamFromPerGroup(), upstreamFromPerGroup()

DTEG.analysis

Run differential TE analysis

#### **Description**

Expression analysis of 2 dimensions, usually Ribo-seq vs RNA-seq.
Using an equal reimplementation of the deltaTE algorithm (see reference).
Creates a total of 3 DESeq models (given x is the target.contrast argument) (usually 'condition'

### \*\* The 3 differential sub models \*\*

column) and libraryType is RNA-seq and Ribo-seq):

- 1. Ribo-seq model : design =  $\sim x$  (differences between the x groups in Ribo-seq)
- 2. RNA-seq model: design =  $\sim$  x (differences between the x groups in RNA-seq)
- 3. TE model: design =  $\sim x$  + libraryType + libraryType:x (differences between the x and libraryType groups and the interaction between them)

You need at least 2 groups and 2 replicates per group. By default, the Ribo-seq counts will be over CDS and RNA-seq counts over whole mRNAs, per transcript. See notes section below for more details.

### Usage

```
DTEG.analysis(
  df.rfp,
  df.rna,
  output.dir = QCfolder(df.rfp),
  target.contrast = design[1],
  design = ORFik::design(df.rfp),
  p.value = 0.05,
  RFP_counts = countTable(df.rfp, "cds", type = "summarized"),
  RNA_counts = countTable(df.rna, "mrna", type = "summarized"),
  batch.effect = FALSE,
  pairs = combn.pairs(unlist(df.rfp[, design])),
  plot.title = "",
  plot.ext = ".pdf",
  width = 6,
  height = 6.
  dot.size = 0.4,
  relative.name = paste0("DTEG_plot", plot.ext),
  complex.categories = FALSE,
  plot_to_console = TRUE
)
```

#### **Arguments**

df.rfp a experiment of usually Ribo-seq or 80S from TCP-seq. (the numerator of the

experiment, usually having a primary role)

df.rna a experiment of usually RNA-seq. (the denominator of the experiment, usually

having a normalizing function)

output.dir character, default QCfolder(df.rfp). output.dir directory to save plots, plot

will be named "TE\_between". If NULL, will not save.

target.contrast

design

a character vector, default design[1]. The column in the ORFik experiment that represent the comparison contrasts. By default: the first design factor of the full experimental design. This is the factor you will do the comparison on. DESeq will normalize the counts based on the full design, but the log fold change values

will be based on this contrast only. It is usually the 'condition' column.

a character vector, default design(df.rfp). The full experiment design. Which

factors have more than 1 level. Example: stage column are all HEK293, so it can not be a design factor. The condition column has 2 possible values, WT and mutant, so it is a factor of the experiment. Replicates column is not part of design, that is inserted later with setting batch.effect = TRUE. Library type 'libtype' column, can also no be part of initial design, it is always added inside

the function, after initial setup.

p.value a numeric, default 0.05 in interval (0,1). Defines adjusted p-value to be used as

significance threshold for the result groups. I.e. for exclusive translation group significant subset for p.value = 0.05 means: TE\$padj < 0.05 & Ribo\$padj < 0.05

& RNA\$padj > 0.05.

RFP\_counts a SummarizedExperiment, default: countTable(df.rfp, "cds", type = "summarized"),

unshifted libraries, all transcript CDSs. If you have pshifted reads and countTables, do: countTable(df.rfp, "cds", type = "summarized", count.folder = "pshifted") Assign a subset if you don't want to analyze all genes. It is rec-

ommended to not subset, to give DESeq2 data for variance analysis.

RNA\_counts a SummarizedExperiment, default: countTable(df.rna, "mrna", type = "summa-

rized"), all transcripts. Assign a subset if you don't want to analyze all genes. It is recommended to not subset, to give DESeq2 data for variance analysis.

batch.effect logical, default TRUE. Makes replicate column of the experiment part of the

design.

If you believe you might have batch effects, keep as TRUE. Batch effect usually means that you have a strong variance between biological replicates. Check out pcaExperiment and see if replicates cluster together more than the design

factor, to verify if you need to set it to TRUE.

pairs list of character pairs, the experiment contrasts. Default: combn.pairs(unlist(df.rfp[,

target.contrast])

plot.title title for plots, usually name of experiment etc

plot.ext character, default: ".pdf". Alternatives: ".png" or ".jpg".

width numeric, default 6 (in inches) height numeric, default 6 (in inches)

dot.size numeric, default 0.4, size of point dots in plot.

relative.name character, Default: paste0("DTEG\_plot", plot.ext) Relative name of file to

be saved in folder specified in output.dir. Change to .pdf if you want pdf file

instead of png.

complex.categories

logical, default FALSE. Separate into more groups, see above for details.

plot\_to\_console

logical, default TRUE. Plot to console before returning, set to FALSE to save

some run time.

#### **Details**

Log fold changes and p-values are created from a Walds test on the comparison contrast described bellow. The RNA-seq and Ribo-seq LFC values are shrunken using DESeq2::lfcShrink(type = "normal"). Note that the TE LFC values are not shrunken (as following specifications from deltaTE paper). The adjusted p-values are created using DESEQ pAdjustMethod = "BH" (Benjamini-Hochberg correction). All other DESEQ2 arguments are default.

Analysis is done between each possible combination of levels in the target contrast If target contrast is condition column, with factor levels: WT, mut1 and mut2 with 3 replicates each. You get comparison of WT vs mut1, WT vs mut2 and mut1 vs mut2.

The respective result categories are defined through 4 main categories, first some intuition. The number of ribosomes (Ribo-seq) is significantly different between 2 contrast elements in the model if the relative counts is statistically higher/lower, for mRNA levels (RNA-seq) it is the same. So TE is then RFP / RNA which is basically how many ribosomes translated per mRNA in the sample, if contrast group 1 has TE of 2, it means 2 ribosomes per mrna fragment, while TE of 4 would be a doubling of 4 ribosomes per mRNA.

Mathematically the groups are defined by the p adjusted values as the following (te.sign means na\_safe(te.padj < p.value), na\_safe is a function where NA values are FALSE for '<=' test and TRUE for '>' test), we also use a helper function: te.sign & rfp.sign & rna.sign, all\_models\_sign := TRUE.

#### \*\* Signicant DTEG Classifications \*\*

- No change: None of the below categories
- Translation (only RFP): te.sign & rfp.sign & !rna.sign
- Expression (only RNA): !te.sign & !rfp.sign & rna.sign
- mRNA abundance : all\_models\_sign & na\_safe(te.lfc \* rna.lfc, ">", 0)
- Inverse (inverse mRNA abundance): all\_models\_sign & te.lfc \* rna.lfc, "<", 0)
- Buffering (Stable protein output): te.sign & !rfp.sign & rna.sign
- Forwarded (diagonal bottom left to top right) : !te.sign & rfp.sign & rna.sign

If complex.categories is FALSE, then Expression, Inverse and forwarded are defined 'Buffering'. mRNA abundance is called "Intensified" in original article For code, of classification, run: View(ORFik:::DTEG\_add\_regulati Feel free to redefine the categories as you want them.

See Figure 1 in the reference article for a clear definition of the groups!

If you do not need isoform variants, subset to longest isoform per gene either before or in the returned object (See examples). If you do not have RNA-seq controls, you can still use DESeq on Ribo-seq alone.

The LFC values are shrunken by lfcShrink(type = "normal").

Remember that DESeq by default can not do global change analysis, it can only find subsets with changes in LFC!

#### Value

a data.table with columns: (contrast variable, gene id, regulation status, log fold changes, p.adjust values, mean counts, significant (as logical))

#### References

```
https://doi.org/10.1002/cpmb.108
```

#### See Also

```
Other DifferentialExpression: DEG.plot.static(), DEG_model(), DTEG.plot(), te.table(), te_rna.plot()
```

#### **Examples**

```
## Simple example (use ORFik template, then split on Ribo and RNA)
df <- ORFik.template.experiment()</pre>
df.rfp <- df[df$libtype == "RFP",]</pre>
df.rna <- df[df$libtype == "RNA",]</pre>
design(df.rfp) # The experimental design, per libtype
design(df.rfp)[1] # Default target contrast
#dt <- DTEG.analysis(df.rfp, df.rna)</pre>
## If you want to use the pshifted libs for analysis:
#dt <- DTEG.analysis(df.rfp, df.rna,</pre>
                      RFP_counts = countTable(df.rfp, region = "cds",
                         type = "summarized", count.folder = "pshifted"))
## Restrict DTEGs by log fold change (LFC):
## subset to abs(LFC) < 1.5 for both rfp and rna
\#dt[abs(rfp) < 1.5 \& abs(rna) < 1.5, Regulation := "No change"]
## Only longest isoform per gene:
#tx_longest <- filterTranscripts(df.rfp, 0, 1, 0)</pre>
#dt <- dt[id %in% tx_longest,]</pre>
## Convert to gene id
#dt[, id := txNamesToGeneNames(id, df.rfp)]
## To get by gene symbol, use biomaRt conversion
## To flip directionality of contrast pair nr 2:
#design <- "condition"
#pairs <- combn.pairs(unlist(df.rfp[, design])</pre>
#pairs[[2]] <- rev(pars[[2]])</pre>
#dt <- DTEG.analysis(df.rfp, df.rna,</pre>
                      RFP_counts = countTable(df.rfp, region = "cds",
```

DTEG.plot

```
# type = "summarized", count.folder = "pshifted"),
# pairs = pairs)
```

DTEG.plot

Plot DTEG result

## **Description**

For explanation of plot catagories, see DTEG. analysis

## Usage

```
DTEG.plot(
   dt,
   output.dir = NULL,
   p.value.label = ifelse(!is.null(attr(dt, "p.value")), attr(dt, "p.value"), 0.05),
   plot.title = "",
   plot.ext = ".pdf",
   width = 6,
   height = 6,
   dot.size = 0.4,
   xlim = "bidir.max",
   ylim = "bidir.max",
   relative.name = paste0("DTEG_plot", plot.ext),
   plot_to_console = TRUE
)
```

# Arguments

dt	a data.table with the results from DTEG.analysis
output.dir	a character path, default NULL(no save), or a directory to save to a file. Relative name of file, specified by 'relative.name' argument.
p.value.label	a numeric, default ifelse(!is.null(attr(dt, "p.value")), attr(dt, "p.value"), 0.05) Interval $(0,1)$ , use"" to not show. What p-value used for the analysis? Will be shown as a caption.
plot.title	title for plots, usually name of experiment etc
plot.ext	character, default: ".pdf". Alternatives: ".png" or ".jpg".
width	numeric, default 6 (in inches)
height	numeric, default 6 (in inches)
dot.size	numeric, default 0.4, size of point dots in plot.
xlim	numeric vector or character preset, default: "bidir.max" (Equal in both $+$ / direction, using max value $+$ 0.5 of rna column in dt). If you want ggplot to decide limit, set to "auto". For numeric vector, specify min and max x limit: like $c(-5,5)$

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ylim

numeric vector or character preset, default: "bidir.max" (Equal in both + / direction, using max value + 0.5 of rfp column in dt). If you want ggplot to decide limit, set to "auto". For numeric vector, specify min and max y limit: like c(-10, 10)

relative.name

character, Default: paste@("DTEG\_plot", plot.ext) Relative name of file to be saved in folder specified in output.dir. Change to .pdf if you want pdf file instead of png.

plot\_to\_console

logical, default TRUE. Plot to console before returning, set to FALSE to save some run time.

#### Value

a ggplot object

#### See Also

```
Other DifferentialExpression: DEG.plot.static(), DEG_model(), DTEG.analysis(), te.table(), te_rna.plot()
```

## **Examples**

```
df <- ORFik.template.experiment()
df.rfp <- df[df$libtype == "RFP",]
df.rna <- df[df$libtype == "RNA",]
#dt <- DTEG.analysis(df.rfp, df.rna)
#Default scaling
#DTEG.plot(dt)
#Manual scaling
#DTEG.plot(dt, xlim = c(-2, 2), ylim = c(-2, 2))</pre>
```

entropy

Percentage of maximum entropy

# Description

Calculates percentage of maximum entropy of the 'reads' coverage over each ORF in 'grl' group. The entropy value per group is a real number in the interval (0:1), where 0 indicates no variance in reads over all codons of group For example c(0,0,0,0) has 0 entropy, since no reads overlap.

```
Interval: [0]: No reads or all reads in 1 place
Interval: [0.01-0.99]: >= 2 positions covered
Interval: [1]: all positions covered perfectly in frame
```

### Usage

```
entropy(grl, reads, weight = 1L, is.sorted = FALSE, overlapGrl = NULL)
```

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## **Arguments**

grl	a GRangesList object can be either transcripts, 5' utrs, cds', 3' utrs or ORFs as a special case (uORFs, potential new cds' etc). If regions are not spliced you can send a GRanges object.
reads	a GAlignments, GRanges or GRangesList object, usually of RiboSeq, RnaSeq, CageSeq, etc.
weight	a numeric/integer vector or metacolumn name. (default: 1L, no differential weighting). If weight is name of defined meta column in reads object, it gives the number of times a read was found at that position. GRanges("chr1", 1, "+", score = 5), would mean "score" column tells that this alignment region was found 5 times. if 1L it means each read is weighted equal as 1, this is what among others countOverlaps() presumes, if single number (!= 1), it repeats for all ranges, if vector with length > 1, it must be equal size of the reads object.
is.sorted	logical (FALSE), is grl sorted. That is $+$ strand groups in increasing ranges $(1,2,3)$ , and $-$ strand groups in decreasing ranges $(3,2,1)$
overlapGrl	an integer, (default: NULL), if defined must be countOverlaps(grl, RFP), added for speed if you already have it.

#### Value

A numeric vector containing one entropy value per element in 'grl'

#### See Also

```
Other features: computeFeatures(), computeFeaturesCage(), countOverlapsW(), disengagementScore(), distToCds(), distToTSS(), floss(), fpkm(), fpkm_calc(), fractionLength(), initiationScore(), insideOutsideORF(), isInFrame(), isOverlapping(), kozakSequenceScore(), orfScore(), rankOrder(), ribosomeReleaseScore(), ribosomeStallingScore(), startRegion(), startRegionCoverage(), stopRegion(), subsetCoverage(), translationalEff()
```

# Examples

```
# a toy example with ribo-seq p-shifted reads 

ORF <- GRangesList(tx1 = GRanges("1", IRanges(1, width = 9), "+")) 

entropy(ORF, GRanges()) # 0 

entropy(ORF, GRanges("1", IRanges(c(1)), "+")) # 0 

entropy(ORF, GRanges("1", IRanges(c(1,4,6,7)), "+")) # 0.94 

entropy(ORF, GRanges("1", IRanges(c(1,4,7)), "+", score = c(1,2,1)), 

    weight = "score") # 0.94 

entropy(ORF, GRanges("1", IRanges(c(1,4,7)), "+")) # Perfect = 1
```

envExp 119

envExp

Get ORFik experiment environment

## Description

More correctly, get the pointer reference, default is .GlobalEnv

## Usage

```
envExp(x)
```

## Arguments

Х

an ORFik experiment

#### Value

environment pointer, name of environment: pointer

```
envExp, experiment-method
```

Get ORFik experiment environment

# Description

More correctly, get the pointer reference, default is .GlobalEnv

### Usage

```
## S4 method for signature 'experiment'
envExp(x)
```

### **Arguments**

Х

an ORFik experiment

#### Value

environment pointer, name of environment: pointer

envExp<-

Set ORFik experiment environment

# Description

More correctly, set the pointer reference, default is .GlobalEnv

#### Usage

```
envExp(x) \leftarrow value
```

## **Arguments**

x an ORFik experiment

value environment pointer to assign to experiment

#### Value

an ORFik experiment with updated environment

```
envExp<-,experiment-method</pre>
```

Set ORFik experiment environment

# Description

More correctly, set the pointer reference, default is .GlobalEnv

## Usage

```
## S4 replacement method for signature 'experiment'
envExp(x) <- value</pre>
```

### **Arguments**

x an ORFik experiment

value environment pointer to assign to experiment

### Value

an ORFik experiment with updated environment

exists.ftp.dir.fast

## Description

Check if ftp directory exists

# Usage

```
exists.ftp.dir.fast(url.dir, report.error = FALSE)
```

### **Arguments**

url.dir character, url to a ftp directory.

report.error logical, FALSE. If TRUE, stop and report error.

#### Value

logical, TRUE if url directory exists

## Description

Check if ftp file exists

### Usage

```
exists.ftp.file.fast(url, report.error = FALSE)
```

### **Arguments**

url character, url to a ftp file

report.error logical, FALSE. If TRUE, stop and report error.

## Value

logical, TRUE if file exists

122 experiment-class

experiment-class

experiment class definition

### Description

It is an object that simplify and error correct your NGS workflow, creating a single R object that stores and controls all results relevant to a specific experiment.

It contains following important parts:

- filepaths: Information for each library in the experiment (for multiple file formats: bam, bed, wig, ofst, etc.)
- genome: Annotation files for the experiment (fasta genome, index, gtf, txdb)
- organism: Name (for automatic GO, sequence analysis, etc.)
- description: Author information and experiment details (use 'list.experiments()' to show all experiments made with ORFik; this makes it easy to find and load them later)
- API: ORFik supports a rich API for using the experiment, e.g., 'outputLibs(experiment, type = "wig")' to load all libraries in the wig format into R, 'loadTxdb(experiment)' to load the txdb (gtf) of the experiment, 'transcriptWindow()' to plot metacoverage for all libraries, and 'countTable(experiment)' to load count tables, etc.
- Safety: Verifies that experiments contain no duplicate, empty, or non-accessible files.

Act as a way of extension of SummarizedExperiment by allowing more ease to find not only counts, but rather information about libraries, and annotation, so that more tasks are possible. Like coverage per position in some transcript etc.

#### ## Constructor:

Simplest way to make is to call:

create.experiment(dir)

On some folder with NGS libraries (usually bam files) and see what you get. Some of the fields might be needed to fill in manually. Each resulting row must be unique (not including filepath, they are always unique), that means if it has replicates then that must be said explicit. And all filepaths must be unique and have files with size > 0.

Here all the columns in the experiment will be described: name (column info): examples

- filepaths: Information for each library in the experiment (for multiple file formats: bam, bed, wig, ofst, etc.)
- genome: Annotation files for the experiment (fasta genome, index, gtf, txdb)
- organism: Name (for automatic GO, sequence analysis, etc.)
- description: Author information and experiment details (use 'list.experiments()' to show all experiments made with ORFik; this makes it easy to find and load them later)
- API: ORFik supports a rich API for using the experiment, e.g., 'outputLibs(experiment, type = "wig")' to load all libraries in the wig format into R, 'loadTxdb(experiment)' to load the txdb (gtf) of the experiment, 'transcriptWindow()' to plot metacoverage for all libraries, and 'countTable(experiment)' to load count tables, etc.
- Safety: Verifies that experiments contain no duplicate, empty, or non-accessible files.

experiment-class 123

#### **Details**

Special rules:

Supported:

Single/paired end bam, bed, wig, ofst + compressions of these

The reverse column of the experiments says "paired-end" if bam file. If a pair of wig files, forward and reverse strand, reverse is filepath to '-' strand wig file. Paired forward / reverse wig files, must have same name except \_forward / \_reverse in name

Paired end bam, when creating experiment, set pairedEndBam = c(T, T, T, F). For 3 paired end libraries, then one single end.

Naming: Will try to guess naming for tissues / stages, replicates etc. If it finds more than one hit for one file, it will not guess. Always check that it guessed correctly.

#### Value

a ORFik experiment

#### See Also

```
Other ORFik_experiment: ORFik.template.experiment(), ORFik.template.experiment.zf(), bamVarName(), create.experiment(), filepath(), libraryTypes(), organism, experiment-method, outputLibs(), read.experiment(), save.experiment(), validateExperiments()
```

### **Examples**

```
## To see an internal ORFik example
df <- ORFik.template.experiment()</pre>
## See libraries in experiment
df
## See organism of experiment
organism(df)
## See file paths in experiment
filepath(df, "default")
## Output NGS libraries in R, to .GlobalEnv
#outputLibs(df)
## Output cds of experiment annotation
#loadRegion(df, "cds")
## This is how to make it:
## Not run:
library(ORFik)
# 1. Update path to experiment data directory (bam, bed, wig files etc)
exp_dir = "/data/processed_data/RNA-seq/Lee_zebrafish_2013/aligned/"
# 2. Set a short character name for experiment, (Lee et al 2013 -> Lee13, etc)
exper_name = "Lee13"
# 3. Create a template experiment (gtf and fasta genome)
temp <- create.experiment(exp_dir, exper_name, saveDir = NULL,</pre>
 txdb = "/data/references/Zv9_zebrafish/Danio_rerio.Zv9.79.gtf",
 fa = "/data/references/Zv9_zebrafish/Danio_rerio.Zv9.fa",
```

124 experiment.colors

```
organism = "Homo sapiens")
# 4. Make sure each row(sample) is unique and correct
# You will get a view open now, check the data.frame that it is correct:
# library type (RNA-seq, Ribo-seq), stage, rep, condition, fraction.
# Let say it did not figure out it is RNA-seq, then we do:"
temp[5:6, 1] \leftarrow "RNA" \# [row 5 and 6, col 1] are library types
# You can also do this in your spread sheet program (excel, libre office)
# Now save new version, if you did not use spread sheet.
saveName <- paste0("/data/processed_data/experiment_tables_for_R/",</pre>
exper_name,".csv")
save.experiment(temp, saveName)
# 5. Load experiment, this will validate that you actually made it correct
df <- read.experiment(saveName)</pre>
# Set experiment name not to be assigned in R variable names
df@expInVarName <- FALSE
df
## End(Not run)
```

experiment.colors

Decide color for libraries by grouping

## Description

Pick the grouping wanted for colors, by default only group by libtype. Like RNA-seq(skyblue4) and Ribo-seq(orange).

## Usage

```
experiment.colors(
   df,
   color_list = "default",
   skip.libtype = FALSE,
   skip.stage = TRUE,
   skip.replicate = TRUE,
   skip.fraction = TRUE,
   skip.condition = TRUE
```

### **Arguments**

df

an ORFik experiment

export.bed12

color_list	a character vector of colors, default "default". That is the vector c("skyblue4", 'orange', "green", "red", "gray", "yellow", "blue", "red2", "orange3"). Picks number of colors needed to make groupings have unique color
skip.libtype	a logical (FALSE), don't include libtype
skip.stage	a logical (FALSE), don't include stage in variable name.
skip.replicate	a logical (FALSE), don't include replicate in variable name.
skip.fraction	a logical (FALSE), don't include fraction
skip.condition	a logical (FALSE), don't include condition in variable name.

#### Value

a character vector of colors

port.be	d12	Export as bed12 format	

# Description

bed format for multiple exons per group, as transcripts. Can be use as alternative as a sparse .gff format for ORFs. Can be direct input for ucsc browser or IGV

# Usage

```
export.bed12(grl, file, rgb = 0)
```

### **Arguments**

grl	A GRangesList
file	a character path to valid output file name
rgb	integer vector, default (0), either single integer or vector of same size as grl to specify groups. It is adviced to not use more than 8 different groups

## **Details**

If grl has no names, groups will be named 1,2,3,4..

### Value

```
NULL (File is saved as .bed)
```

#### References

https://bedtools.readthedocs.io/en/latest/content/general-usage.html#bed-format

126 export.bedo

#### See Also

```
Other utils: bedToGR(), convertToOneBasedRanges(), export.bigWig(), export.fstwig(), export.wiggle(), fimport(), findFa(), fread.bed(), optimizeReads(), readBam(), readBigWig(), readWig()
```

## **Examples**

```
grl <- GRangesList(GRanges("1", c(1,3,5), "+"))
# export.bed12(grl, "output/path/orfs.bed")</pre>
```

export.bedo

Store GRanges object as .bedo

## Description

.bedo is .bed ORFik, an optimized bed format for coverage reads with read lengths .bedo is a text based format with columns (6 maximum):

- 1. chromosome
- 2. start
- 3. end
- 4. strand
- 5. ref width (cigar # M's, match/mismatch total)
- 6. duplicates of that read

### Usage

```
export.bedo(object, out)
```

## Arguments

object a GRanges object

out a character, location on disc (full path)

## **Details**

Positions are 1-based, not 0-based as .bed. End will be removed if all ends equals all starts. Import with import.bedo

## Value

NULL, object saved to disc

export.bedoc 127

export.bedoc

Store GAlignments object as .bedoc

### **Description**

A fast way to store, load and use bam files. (we now recommend using link{export.ofst} instead!)

.bedoc is .bed ORFik, an optimized bed format for coverage reads with cigar and replicate number. .bedoc is a text based format with columns (5 maximum):

- 1. chromosome
- 2. cigar: (cigar # M's, match/mismatch total)
- 3. start (left most position)
- 4. strand (+, -, \*)
- 5. score: duplicates of that read

## Usage

```
export.bedoc(object, out)
```

#### **Arguments**

object a GAlignments object

out a character, location on disc (full path)

#### **Details**

Positions are 1-based, not 0-based as .bed. Import with import.bedoc

## Value

NULL, object saved to disc

export.bigWig

Export as bigWig format

### **Description**

Will create 2 files, 1 for + strand (\*\_forward.bigWig) and 1 for - strand (\*\_reverse.bigWig). If all ranges are \* stranded, will output 1 file. Can be direct input for ucsc browser or IGV

128 export.bigWig

### Usage

```
export.bigWig(
   x,
   file,
   split.by.strand = TRUE,
   is_pre_collapsed = FALSE,
   seq_info = seqinfo(x)
)
```

### **Arguments**

Χ

A GRangesList, GAlignment GAlignmentPairs with score column. Will be converted to 5' end position of original range. If score column does not exist, will group ranges and give replicates as score column. Since bigWig needs a score column to represent counts!

file

a character path to valid output file name

split.by.strand

logical, default TRUE. Split bigWig into 2 files, one for each strand.

is\_pre\_collapsed

logical, default FALSE. Have you already collapsed reads with collapse.by.scores, so each positions is only in 1 GRanges object with a score column per readlength? Set to TRUE, only if you are sure, will give a speedup.

seq\_info

a Seqinfo object, default seqinfo(x). Must have non NA seqlengths defined!

## Value

```
invisible(NULL) (File is saved as 2 .bigWig files)
```

#### References

https://genome.ucsc.edu/goldenPath/help/bigWig.html

### See Also

```
Other utils: bedToGR(), convertToOneBasedRanges(), export.bed12(), export.fstwig(), export.wiggle(), fimport(), findFa(), fread.bed(), optimizeReads(), readBam(), readBigWig(), readWig()
```

## **Examples**

```
x <- c(GRanges("1", c(1,3,5), "-"), GRanges("1", c(1,3,5), "+"))
seqlengths(x) <- 10
file <- file.path(tempdir(), "rna.bigWig")
# export.bigWig(x, file)
# export.bigWig(covRleFromGR(x), file)</pre>
```

export.fstwig 129

export.fstwig

Export as fstwig (fastwig) format

### **Description**

Will create 2 files, 1 for + strand (\*\_forward.fstwig) and 1 for - strand (\*\_reverse.fstwig). If all ranges are \* stranded, will output 1 file.

### Usage

```
export.fstwig(
   x,
   file,
   by.readlength = TRUE,
   by.chromosome = TRUE,
   compress = 50
)
```

## **Arguments**

x A GRangesList, GAlignment GAlignmentPairs with score column or coverage

RLElist Will be converted to 5' end position of original range. If score column

does not exist, will group ranges and give replicates as score column.

file a character path to valid output file name

by.readlength logical, default TRUE by.chromosome logical, default TRUE

compress value in the range 0 to 100, indicating the amount of compression to use. Lower

values mean larger file sizes. The default compression is set to 50.

## Value

```
invisible(NULL) (File is saved as 2 .fstwig files)
```

## References

"TODO"

#### See Also

```
Other utils: bedToGR(), convertToOneBasedRanges(), export.bed12(), export.bigWig(), export.wiggle(), fimport(), findFa(), fread.bed(), optimizeReads(), readBam(), readBigWig(), readWig()
```

export.ofst

### **Examples**

```
x <- c(GRanges("1", c(1,3,5), "-"), GRanges("1", c(1,3,5), "+"))
x$size <- rep(c(28, 29), length.out = length(x))
x$score <- c(5,1,2,5,1,6)
seqlengths(x) <- 5
# export.fstwig(x, "~/Desktop/ribo")</pre>
```

export.ofst

Store GRanges / GAlignments object as .ofst

# Description

A much faster way to store, load and use bam files.

.ofst is ORFik fast serialized object, an optimized format for coverage reads with cigar and replicate number. It uses the fst format as back-end: fst-package.

A .ofst ribo seq file can compress the information in a bam file from 5GB down to a few MB. This new files has super fast reading time, only a few seconds, instead of minutes. It also has random index access possibility of the file.

.ofst is represented as a data.frane format with minimum 4 columns:

- 1. chromosome
- 2. start (left most position)
- 3. strand (+, -, \*)
- 4. width (not added if cigar exists)
- 5. cigar (not needed if width exists): (cigar # M's, match/mismatch total)
- 5. score: duplicates of that read
- 6. size: qwidth according to reference of read

If file is from GAlignmentPairs, it will contain a cigar1, cigar2 instead of cigar and start1 and start2 instead of start

#### Usage

```
export.ofst(x, file, ...)
```

# Arguments

```
x a GRanges, GAlignments or GAlignmentPairs object file a character, location on disc (full path)
... additional arguments for write_fst
```

#### **Details**

Other columns can be named whatever you want and added to meta columns. Positions are 1-based, not 0-based as .bed. Import with import.ofst

#### Value

NULL, object saved to disc

### **Examples**

```
## GRanges
gr <- GRanges("1:1-3:-")
# export.ofst(gr, file = "path.ofst")
## GAlignment
# Make input data.frame
df <- data.frame(seqnames = "1", cigar = "3M", start = 1L, strand = "+")
ga <- ORFik:::getGAlignments(df)
# export.ofst(ga, file = "path.ofst")</pre>
```

export.ofst, GAlignmentPairs-method

Store GRanges / GAlignments object as .ofst

### **Description**

A much faster way to store, load and use bam files.

.ofst is ORFik fast serialized object, an optimized format for coverage reads with cigar and replicate number. It uses the fst format as back-end: fst-package.

A .ofst ribo seq file can compress the information in a bam file from 5GB down to a few MB. This new files has super fast reading time, only a few seconds, instead of minutes. It also has random index access possibility of the file.

.ofst is represented as a data.frane format with minimum 4 columns:

- 1. chromosome
- 2. start (left most position)
- 3. strand (+, -, \*)
- 4. width (not added if cigar exists)
- 5. cigar (not needed if width exists): (cigar # M's, match/mismatch total)
- 5. score: duplicates of that read
- 6. size: qwidth according to reference of read

If file is from GAlignmentPairs, it will contain a cigar1, cigar2 instead of cigar and start1 and start2 instead of start

### Usage

```
## S4 method for signature 'GAlignmentPairs'
export.ofst(x, file, ...)
```

#### **Arguments**

```
x a GRanges, GAlignments or GAlignmentPairs object
file a character, location on disc (full path)
... additional arguments for write_fst
```

#### **Details**

Other columns can be named whatever you want and added to meta columns. Positions are 1-based, not 0-based as .bed. Import with import.ofst

#### Value

NULL, object saved to disc

### **Examples**

```
## GRanges
gr <- GRanges("1:1-3:-")
# export.ofst(gr, file = "path.ofst")
## GAlignment
# Make input data.frame
df <- data.frame(seqnames = "1", cigar = "3M", start = 1L, strand = "+")
ga <- ORFik:::getGAlignments(df)
# export.ofst(ga, file = "path.ofst")</pre>
```

export.ofst, GAlignments-method

Store GRanges / GAlignments object as .ofst

#### **Description**

A much faster way to store, load and use bam files.

.ofst is ORFik fast serialized object, an optimized format for coverage reads with cigar and replicate number. It uses the fst format as back-end: fst-package.

A .ofst ribo seq file can compress the information in a bam file from 5GB down to a few MB. This new files has super fast reading time, only a few seconds, instead of minutes. It also has random index access possibility of the file.

.ofst is represented as a data.frane format with minimum 4 columns:

- 1. chromosome
- 2. start (left most position)
- 3. strand (+, -, \*)
- 4. width (not added if cigar exists)
- 5. cigar (not needed if width exists): (cigar # M's, match/mismatch total)
- 5. score: duplicates of that read
- 6. size: qwidth according to reference of read

If file is from GAlignmentPairs, it will contain a cigar1, cigar2 instead of cigar and start1 and start2 instead of start

#### Usage

```
## S4 method for signature 'GAlignments'
export.ofst(x, file, ...)
```

#### **Arguments**

```
    x a GRanges, GAlignments or GAlignmentPairs object
    file a character, location on disc (full path)
    additional arguments for write_fst
```

#### **Details**

Other columns can be named whatever you want and added to meta columns. Positions are 1-based, not 0-based as .bed. Import with import.ofst

#### Value

NULL, object saved to disc

## **Examples**

```
## GRanges
gr <- GRanges("1:1-3:-")
# export.ofst(gr, file = "path.ofst")
## GAlignment
# Make input data.frame
df <- data.frame(seqnames = "1", cigar = "3M", start = 1L, strand = "+")
ga <- ORFik:::getGAlignments(df)
# export.ofst(ga, file = "path.ofst")</pre>
```

```
export.ofst, GRanges-method
```

Store GRanges / GAlignments object as .ofst

### Description

A much faster way to store, load and use bam files.

.ofst is ORFik fast serialized object, an optimized format for coverage reads with cigar and replicate number. It uses the fst format as back-end: fst-package.

A .ofst ribo seq file can compress the information in a bam file from 5GB down to a few MB. This new files has super fast reading time, only a few seconds, instead of minutes. It also has random index access possibility of the file.

.ofst is represented as a data.frane format with minimum 4 columns:

- 1. chromosome
- 2. start (left most position)
- 3. strand (+, -, \*)
- 4. width (not added if cigar exists)
- 5. cigar (not needed if width exists): (cigar # M's, match/mismatch total)
- 5. score: duplicates of that read
- 6. size: qwidth according to reference of read

If file is from GAlignmentPairs, it will contain a cigar1, cigar2 instead of cigar and start1 and start2 instead of start

134 export.wiggle

#### Usage

```
## S4 method for signature 'GRanges'
export.ofst(x, file, ...)
```

#### **Arguments**

x a GRanges, GAlignments or GAlignmentPairs object file a character, location on disc (full path)
... additional arguments for write\_fst

#### **Details**

Other columns can be named whatever you want and added to meta columns. Positions are 1-based, not 0-based as .bed. Import with import.ofst

## Value

NULL, object saved to disc

### **Examples**

```
## GRanges
gr <- GRanges("1:1-3:-")
# export.ofst(gr, file = "path.ofst")
## GAlignment
# Make input data.frame
df <- data.frame(seqnames = "1", cigar = "3M", start = 1L, strand = "+")
ga <- ORFik:::getGAlignments(df)
# export.ofst(ga, file = "path.ofst")</pre>
```

export.wiggle

Export as wiggle format

### Description

Will create 2 files, 1 for + strand (\*\_forward.wig) and 1 for - strand (\*\_reverse.wig). If all ranges are \* stranded, will output 1 file. Can be direct input for ucsc browser or IGV

## Usage

```
export.wiggle(x, file)
```

#### **Arguments**

x A GRangesList, GAlignment GAlignmentPairs with score column. Will be converted to 5' end position of original range. If score column does not exist, will group ranges and give replicates as score column.

file a character path to valid output file name

extendLeaders 135

#### Value

```
invisible(NULL) (File is saved as 2 .wig files)
```

#### References

https://genome.ucsc.edu/goldenPath/help/wiggle.html

#### See Also

```
Other utils: bedToGR(), convertToOneBasedRanges(), export.bed12(), export.bigWig(), export.fstwig(), fimport(), findFa(), fread.bed(), optimizeReads(), readBam(), readBigWig(), readWig()
```

### **Examples**

```
x \leftarrow c(GRanges("1", c(1,3,5), "-"), GRanges("1", c(1,3,5), "+"))
# export.wiggle(x, "output/path/rna.wig")
```

extendLeaders

Extend the leaders transcription start sites.

### Description

Will extend the leaders or transcripts upstream (5' end) by extension. The extension is general not relative, that means splicing will not be taken into account. Requires the grl to be sorted beforehand, use sortPerGroup to get sorted grl.

## Usage

```
extendLeaders(
  grl,
  extension = 1000L,
  cds = NULL,
  is.circular = all(isCircular(grl) %in% TRUE)
)
```

#### **Arguments**

grl usually a GRangesList of 5' utrs or transcripts. Can be used for any extension

of groups.

extension an integer, how much to max extend upstream (5' end). Either single value that

will apply for all, or same as length of grl which will give 1 update value per grl object. Or a GRangesList where start / stops by strand are the positions to use as new starts. Will not cross the chromosome boundary for non circular

chromosomes.

cds a GRangesList of coding sequences, If you want to extend 5' leaders down-

stream, to catch upstream ORFs going into cds, include it. It will add first cds exon to grl matched by names. Do not add for transcripts, as they are already

included.

136 extendLeadersUntil

is.circular

logical, default FALSE if not any is: all(isCircular(grl) Where grl is the ranges checked. If TRUE, allow ranges to extend below position 1 on chromosome. Since circular genomes can have negative coordinates.

#### Value

an extended GRangeslist

#### See Also

```
Other ExtendGenomicRanges: asTX(), coveragePerTiling(), extendTrailers(), reduceKeepAttr(), tile1(), txSeqsFromFa(), windowPerGroup()
```

#### **Examples**

```
library(GenomicFeatures)
samplefile <- system.file("extdata", "hg19_knownGene_sample.sqlite",</pre>
                           package = "GenomicFeatures")
txdb <- loadDb(samplefile)</pre>
fiveUTRs <- fiveUTRsByTranscript(txdb, use.names = TRUE) # <- extract only 5' leaders
tx <- exonsBy(txdb, by = "tx", use.names = TRUE)</pre>
cds <- cdsBy(txdb,"tx",use.names = TRUE)</pre>
## extend leaders upstream 1000
extendLeaders(fiveUTRs, extension = 1000)
## now try(extend upstream 1000, add all cds exons):
extendLeaders(fiveUTRs, extension = 1000, cds)
## when extending transcripts, don't include cds' of course,
## since they are already there
extendLeaders(tx, extension = 1000)
## Circular genome (allow negative coordinates)
circular_fives <- fiveUTRs</pre>
isCircular(circular_fives) <- rep(TRUE, length(isCircular(circular_fives)))</pre>
extendLeaders(circular_fives, extension = 32672841L)
```

extendLeadersUntil

Extend Leaders Until

#### **Description**

Extend leaders until a restriction group / position. This makes you extend until you hit another gene boundary etc.

#### **Usage**

```
extendLeadersUntil(
  grl,
  grl2 = grl,
```

extendLeadersUntil 137

```
extension = 500,
until = 200,
min_ext = 25,
is.circular = all(isCircular(grl) %in% TRUE),
...
)
```

#### **Arguments**

grl a GRangesList

grl2 a GRangesList, default 'grl'. The list that defines restrictions on extension. Can

also be another set, which is used as 'roadblocks' for extension.

extension an integer, how much to max extend upstream (5' end). Either single value that

will apply for all, or same as length of grl which will give 1 update value per grl object. Or a GRangesList where start / stops by strand are the positions to use as new starts. Will not cross the chromosome boundary for non circular

chromosomes.

until numeric, default 200. The nearest you can go to the neighbour boundaries of

grl2 (the "other" genes). Defined as boundary hit + 1, so if hit other gene with distance 22, and 'until' argument is 2, will set final extension to 22-2-1 = 19. Usually if Leaders/trailers are not defined, this makes a good pseudo leader

boundary around your other genes.

min\_ext numeric, default 25. What is the minimum extension, even though it crosses a

boundary. Will not cross the chromosome boundary for non circular chromo-

somes.

is.circular logical, default FALSE if not any is: all(isCircular(grl) Where grl is the ranges

checked. If TRUE, allow ranges to extend below position 1 on chromosome.

Since circular genomes can have negative coordinates.

... Arguments sent to distanceToPreceding

#### Value

a GRangesList of extended grl input

## **Examples**

138 extendTrailers

extendsTSSexons

Extend first exon of each transcript with length specified

### **Description**

Extend first exon of each transcript with length specified

### Usage

```
extendsTSSexons(fiveUTRs, extension = 1000)
```

## Arguments

fiveUTRs The 5' leader sequences as GRangesList

extension The number of basses to extend transcripts upstream

#### Value

GRangesList object of fiveUTRs

extendTrailers

Extend the Trailers transcription stop sites

#### Description

Will extend the trailers or transcripts downstream (3' end) by extension. The extension is general not relative, that means splicing will not be taken into account. Requires the grl to be sorted beforehand, use sortPerGroup to get sorted grl.

## Usage

```
extendTrailers(
  grl,
  extension = 1000L,
  is.circular = all(isCircular(grl) %in% TRUE)
)
```

## **Arguments**

grl

usually a GRangesList of 3' utrs or transcripts. Can be used for any extension

of groups.

extension

an integer, how much to max extend downstream (3' end). Either single value that will apply for all, or same as length of grl which will give 1 update value per grl object. Or a GRangesList where start / stops sites by strand are the positions to use as new starts. Will not cross the chromosome boundary for non circular chromosomes.

extendTrailersUntil 139

is.circular

logical, default FALSE if not any is: all(isCircular(grl) Where grl is the ranges checked. If TRUE, allow ranges to extend below position 1 on chromosome. Since circular genomes can have negative coordinates.

#### Value

an extended GRangeslist

#### See Also

```
Other ExtendGenomicRanges: asTX(), coveragePerTiling(), extendLeaders(), reduceKeepAttr(), tile1(), txSeqsFromFa(), windowPerGroup()
```

### **Examples**

extendTrailersUntil

Extend Trailers Until

## Description

Extend trailers until a restriction group / position. This makes you extend until you hit another gene boundary etc.

### Usage

```
extendTrailersUntil(
  grl,
  grl2 = grl,
  extension = 500,
  until = 200,
  min_ext = 25,
  is.circular = all(isCircular(grl) %in% TRUE),
  ...
)
```

140 extract\_run\_id

## **Arguments**

grl	a GRangesList
grl2	a GRangesList, default 'grl'. The list that defines restrictions on extension. Can also be another set, which is used as 'roadblocks' for extension.
extension	an integer, how much to max extend downstream (3' end). Either single value that will apply for all, or same as length of grl which will give 1 update value per grl object. Or a GRangesList where start / stops sites by strand are the positions to use as new starts. Will not cross the chromosome boundary for non circular chromosomes.
until	numeric, default 200. The nearest you can go to the boundary. #' Defined as boundary hit $+ 1$ , so if hit on 22, and until is 2, will set to $22+2+1=25$ . Usually if Leaders/trailers are not defined, this makes a good pseudo leader boundary around your other genes.
min_ext	numeric, default 25. What is the minimum extension, even though it crosses a boundary. Will not cross the chromosome boundary for non circular chromosomes.
is.circular	logical, default FALSE if not any is: all(isCircular(grl) Where grl is the ranges checked. If TRUE, allow ranges to extend below position 1 on chromosome. Since circular genomes can have negative coordinates.
• • •	Arguments sent to distanceToFollowing

#### Value

a GRangesList of extended grl input

# Examples

extract\_run\_id

Extract SRR/ERR/DRR run IDs from string

## Description

Extract SRR/ERR/DRR run IDs from string

# Usage

```
extract_run_id(
   x,
   search = "(SRR[0-9]+|DRR[0-9]+|ERR[0-9]+)",
   only_valid = FALSE
)
```

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# Arguments

x character vector to search through.

search the regex search, default: "(SRR[0-9]+|DRR[0-9]+|ERR[0-9]+)"

only\_valid logical, default FALSE. If TRUE, return only the hits.

### Value

a character vector of run accepted run ids according to search, if only\_valid named character vector for which indices are returned

## **Examples**

```
search <- c("SRR1230123_absdb", "SRR1241204124_asdasd", "asd_ERR1231230213",
   "DRR12412412_asdqwe", "ASDASD_ASDASD", "SRRASDASD")
ORFik:::extract_run_id(search)
ORFik:::extract_run_id(search, only_valid = TRUE)</pre>
```

f

 $strandMode\ covRle$ 

## Description

strandMode covRle

## Usage

f(x)

## Arguments

Χ

a covRle object

#### Value

the forward RleList

142 filepath

f,covRle-method

strandMode covRle

## Description

strandMode covRle

## Usage

```
## S4 method for signature 'covRle' f(x)
```

# Arguments

Χ

a covRle object

### Value

the forward RleList

filepath

Get filepaths to ORFik experiment

# Description

If other type than "default" is given and that type is not found (and 'fallback' is TRUE), it will return you ofst files, if they do not exist, then default filepaths without warning.

### Usage

```
filepath(
   df,
   type,
   basename = FALSE,
   fallback = type %in% c("pshifted", "bed", "ofst", "bedoc", "bedo"),
   suffix_stem = "AUTO",
   base_folders = libFolder(df)
)
```

filepath 143

#### **Arguments**

df an ORFik experiment

type a character(default: "default"), load files in experiment or some precomputed

variant, like "ofst" or "pshifted". These are made with ORFik:::convertLibs(), shiftFootprintsByExperiment(), etc. Can also be custom user made folders inside the experiments bam folder. It acts in a recursive manner with priority: If you state "pshifted", but it does not exist, it checks "ofst". If no .ofst files, it uses

"default", which always must exists.

Presets are (folder is relative to default lib folder, some types fall back to other formats if folder does not exist):

- "default": load the original files for experiment, usually bam.

- "ofst": loads ofst files from the ofst folder, relative to lib folder (falls back to default)

- "pshifted": loads ofst, wig or bigwig from pshifted folder (falls back to ofst, then default)

- "cov": Load covRle objects from cov\_RLE folder (fail if not found)

- "covl": Load covRleList objects, from cov\_RLE\_List folder (fail if not found)

- "bed": Load bed files, from bed folder (falls back to default)

- Other formats must be loaded directly with fimport

basename logical, default (FALSE). Get relative paths instead of full. Only use for inspec-

tion!

fallback logical, default: type If TRUE, will use type fallback, see above for info.

 $suffix\_stem$  character, default "AUTO". Which is "" for all except type = "pshifted". Then

it is "\_pshifted" appended to end of names before format. Can be vector, then it searches suffixes in priority: so if you insert c("\_pshifted", ""), it will look for

suffix \_pshifted, then the empty suffix.

base\_folders character vector, default libFolder(df), path to base folder to search for library

variant directories. If single path (length == 1), it will apply to all libraries in df. If df is a collection, an experiment where libraries are put in different folders and library variants like pshifted are put inside those respective folders,

set base\_folders = libFolder(df, mode = "all")

#### **Details**

For pshifted libraries, if "pshifted" is specified as type: if if multiple formats exist it will use a priority: ofst -> bigwig -> wig -> bed. For formats outside default, all files must be stored in the directory of the first file: base\_folder <- libFolder(df)

#### Value

a character vector of paths, or a list of character with 2 paths per, if paired libraries exists

#### See Also

```
Other ORFik_experiment: ORFik.template.experiment(), ORFik.template.experiment.zf(), bamVarName(), create.experiment(), experiment-class, libraryTypes(), organism, experiment-method, outputLibs(), read.experiment(), save.experiment(), validateExperiments()
```

### **Examples**

```
df <- ORFik.template.experiment()
filepath(df, "default")
# Subset
filepath(df[9,], "default")
# Other format path
filepath(df[9,], "ofst")
## If you have pshifted files, see shiftFootprintsByExperiment()
filepath(df[9,], "pshifted") # <- falls back to ofst</pre>
```

```
file_ext_without_compression
```

Get file extension of files without compressions

## **Description**

Get file extension of files without compressions

### Usage

```
file_ext_without_compression(x, compressions = c("gzip", "gz", "bgz", "zip"))
```

### **Arguments**

```
x character paths

compressions character vector, default: c("gzip", "gz", "bgz", "zip"). Expand if you have other formats
```

#### Value

character vector of file extensions of paths excluding suffix vector defined in compression.

### **Examples**

```
file_ext_without_compression(c("abc.bam.gz", "def.bam.zip"))
```

filterCage 145

filterCage	Filter peak of cage-data by value	

# Description

Filter peak of cage-data by value

# Usage

```
filterCage(cage, filterValue = 1, fiveUTRs = NULL, preCleanup = TRUE)
```

# Arguments

cage	Either a filePath for the CageSeq file as .bed .bam or .wig, with possible compressions (".gzip", ".gz", ".bgz"), or already loaded CageSeq peak data as GRanges or GAlignment. NOTE: If it is a .bam file, it will add a score column by running: convertToOneBasedRanges(cage, method = "5prime", addScoreColumn = TRUE) The score column is then number of replicates of read, if score column is something else, like read length, set the score column to NULL first.
filterValue	The minimum number of reads on cage position, for it to be counted as possible new tss. (represented in score column in CageSeq data) If you already filtered, set it to 0.
fiveUTRs	a GRangesList (NULL), if added will filter out cage reads by these following rules: all reads in region (-5:-1, 1:5) for each tss will be removed, removes noise.
preCleanup	logical (TRUE), if TRUE, remove all reads in region (-5:-1, 1:5) of all original tss in leaders. This is to keep original TSS if it is only +/- 5 bases from the original.

## Value

the filtered GRanges object

filterExtremePeakGenes

Filter out transcript by a median filter

# Description

For removing very extreme peaks in coverage plots, use high quantiles, like 99. Used to make your plots look better, by removing extreme peaks.

146 filterTranscripts

## Usage

```
filterExtremePeakGenes(
   tx,
   reads,
   upstream = NULL,
   downstream = NULL,
   multiplier = "0.99",
   min_cutoff = "0.999",
   pre_filter_minimum = 0,
   average = "median"
)
```

## **Arguments**

tx a GRangesList

reads a GAlignments or GRanges

upstream numeric or NULL, default NULL. if you want window of tx, instead of whole,

specify how much upstream from start of tx, 10 is include 10 bases before start

downstream numeric or NULL, default NULL. if you want window of tx, instead of whole,

specify how much downstream from start of tx, 10 is go 10 bases into tx from

start.

multiplier a character or numeric, default "0.99", either a quantile if input is string[0-1],

like "0.99", or numeric value if input is numeric. How much bigger than median

/ mean counts per gene, must a value be to be defined as extreme?

min\_cutoff a character or numeric, default "0.999", either a quantile if input is string[0-1],

like "0.999", or numeric value if input is numeric. Lowest allowed value

pre\_filter\_minimum

numeric, default 0. If value is x, will remove all positions in all genes with coverage < x, before median filter is applied. Set to 1 to remove all 0 positions.

coverage < x, before median inter is applied. Set to 1 to remove an o positions.

character, default "median". Alternative: "mean". How to scale the multiplier

argument, from median or mean of gene coverage.

#### Value

average

GRangesList (filtered)

filterTranscripts

Filter transcripts by lengths

# Description

Filter transcripts to those who have leaders, CDS, trailers of some lengths, you can also pick the longest per gene.

filterTranscripts 147

## Usage

```
filterTranscripts(
  txdb,
  minFiveUTR = 30L,
  minCDS = 150L,
  minThreeUTR = 30L,
  longestPerGene = TRUE,
  stopOnEmpty = TRUE,
  by = "tx",
  create.fst.version = FALSE
)
```

## **Arguments**

txdb a TxDb object, ORFik experiment object or a path to one of: (.gtf, .gff, .gff2,

.gff2, .db or .sqlite), Only in the loadRegion function: if it is a GRangesList, it

will return it self.

minFiveUTR (integer) minimum bp for 5' UTR during filtering for the transcripts. Set to

NULL if no 5' UTRs exists for annotation.

minCDS (integer) minimum bp for CDS during filtering for the transcripts

minThreeUTR (integer) minimum bp for 3' UTR during filtering for the transcripts. Set to

NULL if no 3' UTRs exists for annotation.

longestPerGene logical (TRUE), return only longest valid transcript per gene. NOTE: This is

by priority longest cds isoform, if equal then pick longest total transcript. So if

transcript is shorter but cds is longer, it will still be the one returned.

stopOnEmpty logical TRUE, stop if no valid transcripts are found?

by a character, default "tx" Either "tx" or "gene". What names to output region by,

the transcript name "tx" or gene names "gene". NOTE: this is not the same as cdsBy(txdb, by = "gene"), cdsBy would then only give 1 cds per Gene, loadRe-

gion gives all isoforms, but with gene names.

create.fst.version

logical, FALSE. If TRUE, creates a .fst version of the transcript length table (if it not already exists), reducing load time from ~ 15 seconds to ~ 0.01 second next time you run filterTranscripts with this txdb object. The file is stored in the

same folder as the genome this txdb is created from, with the name:

paste0(ORFik:::remove.file\_ext(metadata(txdb)[3,2]), "\_", gsub("\(.\*|

|:", "", metadata(txdb)[metadata(txdb)[,1] == "Creation time",2]), "\_txLengths.fst")

Some error checks are done to see this is a valid location, if the txdb data source

is a repository like UCSC and not a local folder, it will not be made.

#### **Details**

If a transcript does not have a trailer, then the length is 0, so they will be filtered out if you set minThreeUTR to 1. So only transcripts with leaders, cds and trailers will be returned. You can set the integer to 0, that will return all within that group.

If your annotation does not have leaders or trailers, set them to NULL, since 0 means there must exist a column called utr3\_len etc. Genes with gene\_id = NA will be be removed.

148 filterUORFs

## Value

a character vector of valid transcript names

# **Examples**

filterUORFs

Remove uORFs that are false CDS hits

# **Description**

This is a strong filtering, so that even if the cds is on another transcript, the uORF is filtered out, this is because there is no way of knowing by current ribo-seq, rna-seq experiments.

## Usage

```
filterUORFs(uorfs, cds)
```

## **Arguments**

```
uorfs (GRangesList), the uORFs to filter
```

cds (GRangesList), the coding sequences (main ORFs on transcripts), to filter against.

#### Value

```
(GRangesList) of filtered uORFs
```

# See Also

```
Other uorfs: addCdsOnLeaderEnds(), removeORFsWithSameStartAsCDS(), removeORFsWithSameStopAsCDS(), removeORFsWithStartInsideCDS(), removeORFsWithinCDS(), uORFSearchSpace()
```

149 fimport

fimport	Load any type of sequencing reads

## **Description**

Wraps around ORFik file format loaders and rtracklayer::import and tries to speed up loading with the use of data.table. Supports gzip, gz, bgz compression formats. Also safer chromosome naming with the argument chrStyle

## Usage

```
fimport(path, chrStyle = NULL, param = NULL, strandMode = 0)
```

### **Arguments**

path a character path to file (1 or 2 files), or data.table with 2 colums(forward&reverse)

> or a GRanges/Galignment/GAlignmentPairs object etc. If it is ranged object it will presume to be already loaded, so will return the object as it is, updating the

seqlevelsStyle if given.

a GRanges object, TxDb, FaFile, , a seqlevelsStyle or Seqinfo. (Default:

NULL) to get seglevelsStyle from. In addition if it is a Seginfo object, seginfo will be updated. Example of seglevelsStyle update: Is chromosome 1 called chr1 or 1, is mitocondrial chromosome called MT or chrM etc. Will use 1st seqlevel-style if more are present. Like: c("NCBI", "UCSC") -> pick "NCBI"

NULL or a ScanBamParam object. Like for scanBam, this influences what fields param

> and which records are imported. However, note that the fields specified thru this ScanBamParam object will be loaded in addition to any field required for generating the returned object (GAlignments, GAlignmentPairs, or GappedReads object), but only the fields requested by the user will actually be kept as meta-

data columns of the object.

By default (i.e. param=NULL or param=ScanBamParam()), no additional field is

loaded. The flag used is scanBamFlag(isUnmappedQuery=FALSE) for readGAlignments, readGAlignmentsList, and readGappedReads. (i.e. only records corresponding to mapped reads are loaded), and scanBamFlag(isUnmappedQuery=FALSE, isPaired=TRUE, hasUnmappedMate=FALSE) for readGAlignmentPairs (i.e. only

records corresponding to paired-end reads with both ends mapped are loaded).

numeric, default 0. Only used for paired end bam files. One of (0: strand

= \*, 1: first read of pair is +, 2: first read of pair is -). See ?strandMode. Note: Sets default to 0 instead of 1, as readGAlignmentPairs uses 1. This is to guarantee hits, but will also make mismatches of overlapping transcripts in

opposite directions.

## **Details**

NOTE: For wig/bigWig files you can send in 2 files, so that it automatically merges forward and reverse stranded objects. You can also just send 1 wig/bigWig file, it will then have "\*" as strand.

chrStyle

strandMode

150 findFa

#### Value

a GAlignments/GRanges object, depending on input.

## See Also

```
Other utils: bedToGR(), convertToOneBasedRanges(), export.bed12(), export.bigWig(), export.fstwig(), export.wiggle(), findFa(), fread.bed(), optimizeReads(), readBam(), readBigWig(), readWig()
```

## **Examples**

```
bam_file <- system.file("extdata/Danio_rerio_sample", "ribo-seq.bam", package = "ORFik")
fimport(bam_file)
# Certain chromosome naming
fimport(bam_file, "NCBI")
# Paired end bam strandMode 1:
fimport(bam_file, strandMode = 1)
# (will have no effect in this case, since it is not paired end)</pre>
```

findFa

Convenience wrapper for Rsamtools FaFile

## Description

Get fasta file object, to find sequences in file. Will load and import file if necessarry.

# Usage

```
findFa(faFile)
```

## **Arguments**

faFile

FaFile, BSgenome, fasta/index file path or an ORFik experiment. This file is usually used to find the transcript sequences from some GRangesList.

## Value

```
a FaFile or BSgenome
```

#### See Also

```
Other utils: bedToGR(), convertToOneBasedRanges(), export.bed12(), export.bigWig(), export.fstwig(), export.wiggle(), fimport(), fread.bed(), optimizeReads(), readBam(), readBigWig(), readWig()
```

findFromPath 151

## **Examples**

```
# Some fasta genome with existing fasta index in same folder
path <- system.file("extdata/references/danio_rerio", "genome_dummy.fasta", package = "ORFik")
findFa(path)</pre>
```

findFromPath

Find all candidate library types filenames

# Description

From the given experiment

## Usage

```
findFromPath(filepaths, candidates, slot = "auto")
```

# Arguments

filepaths path to all files

candidates a data.table with 2 columns, Possible names to search for, see experiment\_naming

family for candidates.

slot character, default "auto". If auto, use auto guessing of slot, else must be a char-

acter vector of length 1 or equal length as filepaths.

## Value

a candidate library types (character vector)

findLibrariesInFolder Get all library files in folder/folders of given types

## **Description**

Will try to guess paired / unpaired wig, bed, bam files.

## Usage

```
findLibrariesInFolder(dir, types, pairedEndBam = FALSE)
```

152 findMapORFs

## Arguments

dir Which directory / directories to create experiment from, must be a directory

with NGS data from your experiment. Will include all files of file type specified by "types" argument. So do not mix files from other experiments in the same

folder!

types Default c("bam", "bed", "wig", "bigWig", "ofst"), which types of libraries

to allow as NGS data.

pairedEndBam logical FALSE, else TRUE, or a logical list of TRUE/FALSE per library you see

will be included (run first without and check what order the files will come in) 1 paired end file, then two single will be c(T, F, F). If you have a SRA metadata csv file, you can set this argument to study\$LibraryLayout == "PAIRED", where

study is the SRA metadata for all files that was aligned.

#### **Details**

Set pairedEndBam if you have paired end reads as a single bam file.

#### Value

(data.table) All files found from types parameter. With 2 extra column (logical), is it wig pairs, and paired bam files.

findMapORFs

Find ORFs and immediately map them to their genomic positions.

## **Description**

This function can map spliced ORFs. It finds ORFs on the sequences of interest, but returns relative positions to the positions of 'grl' argument. For example, 'grl' can be exons of known transcripts (with genomic coordinates), and 'seq' sequences of those transcripts, in that case, this function will return genomic coordinates of ORFs found on transcript sequences.

## Usage

```
findMapORFs(
  grl,
  seqs,
  startCodon = startDefinition(1),
  stopCodon = stopDefinition(1),
  longestORF = TRUE,
  minimumLength = 0,
  groupByTx = FALSE,
  grl_is_sorted = FALSE
)
```

findMapORFs 153

## **Arguments**

grl A GRangesList of the original sequences that gave the orfs in Genomic coordinates. If grl\_is\_sorted = TRUE (default), negative exon ranges per grl object must be sorted in descending orders. segs (DNAStringSet or character vector) - DNA/RNA sequences to search for Open Reading Frames. Can be both uppercase or lowercase. Easiest call to get seqs if you want only regions from a fasta/fasta index pair is: seqs = OR-Fik:::txSeqsFromFa(grl, faFile), where grl is a GRanges/List of search regions and faFile is a FaFile. Note: Remember that if you extracted through a GRanges object, that must have been sorted with negative strand exons descending. startCodon (character vector) Possible START codons to search for. Check startDefinition for helper function. Note that it is case sensitive, so "atg" would give 0 hits for a sequence with only capital "ATG" ORFs. stopCodon (character vector) Possible STOP codons to search for. Check stopDefinition for helper function. Note that it is case sensitive, so "tga" would give 0 hits for a sequence with only capital "TGA" ORFs. longestORF (logical) Default TRUE. Keep only the longest ORF per unique stopcodon: (seqname, strand, stopcodon) combination, Note: Not longest per transcript! You can also use function longestORFs after creation of ORFs for same result. minimumLength (integer) Default is 0. Which is START + STOP = 6 bp. Minimum length of ORF, without counting 3bps for START and STOP codons. For example minimumLength = 8 will result in size of ORFs to be at least START + 8\*3 (bp) + STOP = 30 bases. Use this param to restrict search. logical (default: FALSE), should output GRangesList be grouped by exons per groupByTx ORF (TRUE) or by orfs per transcript (FALSE)? logical, default FALSE If FALSE will sort negative transcript in descending grl\_is\_sorted order for you. If you loaded ranges with default methods this is already the

## Details

This function assumes that 'seq' is in widths relative to 'grl', and that their orders match. 1st seq is 1st grl object, etc.

case, so you can set to TRUE to save some time.

See vignette for real life example.

#### Value

A GRangesList of ORFs.

## See Also

Other findORFs: findORFs(), findORFsFasta(), findUORFs(), startDefinition(), stopDefinition()

154 findMaxPeaks

## **Examples**

```
# First show simple example using findORFs
# This sequence has ORFs at 1-9 and 4-9
seqs <- DNAStringSet("ATGATGTAA") # the dna transcript sequence</pre>
findORFs(seqs)
# lets assume that this sequence comes from two exons as follows
# Then we need to use findMapORFs instead of findORFs,
# for splicing information
gr <- GRanges(seqnames = "1", # chromosome 1</pre>
              ranges = IRanges(start = c(21, 10), end = c(23, 15)),
              strand = "-", #
              names = "tx1") #From transcript 1 on chr 1
grl <- GRangesList(tx1 = gr) # 1 transcript with 2 exons</pre>
findMapORFs(grl, seqs) # ORFs are properly mapped to its genomic coordinates
grl <- c(grl, grl)</pre>
names(grl) \leftarrow c("tx1", "tx2")
findMapORFs(grl, c(seqs, seqs))
# More advanced example and how to save sequences found in vignette
```

findMaxPeaks

Find max peak for each transcript, returns as data.table, without names, but with index

## **Description**

Find max peak for each transcript, returns as data.table, without names, but with index

## Usage

```
findMaxPeaks(cageOverlaps, filteredCage)
```

# Arguments

cageOverlaps The cageOverlaps between cage and extended 5' leaders filteredCage The filtered raw cage-data used to reassign 5' leaders

#### Value

a data.table of max peaks

findNewTSS 155

findNewTSS	Finds max peaks per trancsript from reads in the cagefile
------------	---

# Description

Finds max peaks per trancsript from reads in the cagefile

# Usage

```
findNewTSS(fiveUTRs, cageData, extension, restrictUpstreamToTx)
```

## **Arguments**

fiveUTRs The 5' leader sequences as GRangesList

cageData The CAGE as GRanges object

extension The number of basses to extends transcripts upstream.

restrictUpstreamToTx

a logical (FALSE), if you want to restrict leaders to not extend closer than 5 bases from closest upstream leader, set this to TRUE.

#### Value

a Hits object

findNGSPairs Find pair of forward and reverse strand wig / bed files and paired end bam files split in two

# Description

Find pair of forward and reverse strand wig / bed files and paired end bam files split in two

## Usage

```
findNGSPairs(
  paths,
  f = c("forward", "fwd"),
  r = c("reverse", "rev"),
  format = "wig"
)
```

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## **Arguments**

paths a character path at least one .wig / .bed file

f Default (c("forward", "fwd") a character vector for forward direction regex.

r Default (c("reverse", "rev") a character vector for reverse direction regex.

format default "wig", for bed do "bed". Also searches compressions of these variants.

#### Value

if not all are paired, return original list, if they are all paired, return a data.table with matches as 2 columns

findORFs

Find Open Reading Frames.

## **Description**

Find all Open Reading Frames (ORFs) on the simple input sequences in ONLY 5'- 3' direction (+), but within all three possible reading frames. Do not use findORFs for mapping to full chromosomes, then use findMapORFs! For each sequence of the input vector IRanges with START and STOP positions (inclusive) will be returned as IRangesList. Returned coordinates are relative to the input sequences.

## Usage

```
findORFs(
  seqs,
  startCodon = startDefinition(1),
  stopCodon = stopDefinition(1),
  longestORF = TRUE,
  minimumLength = 0
)
```

## **Arguments**

seqs	(DNAStringSet or character vector) - DNA/RNA sequences to search for Open Reading Frames. Can be both uppercase or lowercase. Easiest call to get seqs if you want only regions from a fasta/fasta index pair is: seqs = OR-Fik:::txSeqsFromFa(grl, faFile), where grl is a GRanges/List of search regions and faFile is a FaFile. Note: Remember that if you extracted through a GRanges object, that must have been sorted with negative strand exons descending.
startCodon	(character vector) Possible START codons to search for. Check startDefinition for helper function. Note that it is case sensitive, so "atg" would give 0 hits for a sequence with only capital "ATG" ORFs.
stopCodon	(character vector) Possible STOP codons to search for. Check stopDefinition for helper function. Note that it is case sensitive, so "tga" would give 0 hits for a sequence with only capital "TGA" ORFs.

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longestORF (logical) Default TRUE. Keep only the longest ORF per unique stopcodon: (sequame, strand, stopcodon) combination, Note: Not longest per transcript! You can also use function longestORFs after creation of ORFs for same result.

minimumLength (integer) Default is 0. Which is START + STOP = 6 bp. Minimum length of ORF, without counting 3bps for START and STOP codons. For example minimumLength = 8 will result in size of ORFs to be at least START + 8\*3 (bp) + STOP = 30 bases. Use this param to restrict search.

#### **Details**

```
If you want antisence strand too, do: #positive strands pos <- findORFs(seqs) #negative strands (DNAStringSet only if character) neg <- findORFs(reverseComplement(DNAStringSet(seqs))) relist(c(GRanges(pos, strand = "+"), GRanges(neg, strand = "-")), skeleton = merge(pos, neg))
```

#### Value

(IRangesList) of ORFs locations by START and STOP sites grouped by input sequences. In a list of sequences, only the indices of the sequences that had ORFs will be returned, e.g. 3 sequences where only 1 and 3 has ORFs, will return size 2 IRangesList with names c("1", "3"). If there are a total of 0 ORFs, an empty IRangesList will be returned.

#### See Also

```
Other findORFs: findMapORFs(), findORFsFasta(), findUORFs(), startDefinition(), stopDefinition()
```

## **Examples**

```
## Simple examples
findORFs("ATGTAA")
findORFs("ATGTTAA") # not in frame anymore
findORFs("ATGATGTAA") # only longest of two above
findORFs("ATGATGTAA", longestORF = FALSE) # two ORFs
findORFs(c("ATGTAA", "ATGATGTAA")) # 1 ORF per transcript
## Get DNA sequences from ORFs
seq <- DNAStringSet(c("ATGTAA", "AAA", "ATGATGTAA"))</pre>
names(seq) <- c("tx1", "tx2", "tx3")
orfs <- findORFs(seq, longestORF = FALSE)</pre>
# you can get sequences like this:
gr <- unlist(orfs, use.names = TRUE)</pre>
gr <- GRanges(seqnames = names(seq)[as.integer(names(gr))],</pre>
ranges = gr, strand = "+")
# Give them some proper names:
names(gr) <- paste0("ORF_", seq.int(length(gr)), "_", seqnames(gr))</pre>
orf_seqs <- getSeq(seq, gr)</pre>
orf_seqs
# Save as .fasta (orf_seqs must be of type DNAStringSet)
```

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```
# writeXStringSet(orf_seqs, "orfs.fasta")
## Reading from file and find ORFs
#findORFs(readDNAStringSet("path/to/transcripts.fasta"))
```

findORFsFasta

Finds Open Reading Frames in fasta files.

## **Description**

Should be used for procaryote genomes or transcript sequences as fasta. Makes no sence for eukaryote whole genomes, since those contains splicing (use findMapORFs for spliced ranges). Searches through each fasta header and reports all ORFs found for BOTH sense (+) and antisense strand (-) in all frames. Name of the header will be used as seqnames of reported ORFs. Each fasta header is treated separately, and name of the sequence will be used as seqname in returned GRanges object. This supports circular genomes.

#### **Usage**

```
findORFsFasta(
  filePath,
  startCodon = startDefinition(1),
  stopCodon = stopDefinition(1),
  longestORF = TRUE,
  minimumLength = 0,
  is.circular = FALSE
)
```

#### **Arguments**

C:1.D.41

startCodon

filePath	(character) Path to the fasta file. Can be both uppercase or lowercase. Or a
	already loaded R object of either types: "BSgenome" or "DNAStringSet" with
	named sequences

(character vector) Possible START codons to search for. Check startDefinition for helper function. Note that it is case sensitive, so "atg" would give 0 hits for

a sequence with only capital "ATG" ORFs.

stopCodon (character vector) Possible STOP codons to search for. Check stopDefinition

for helper function. Note that it is case sensitive, so "tga" would give 0 hits for

a sequence with only capital "TGA" ORFs.

longestORF (logical) Default TRUE. Keep only the longest ORF per unique stopcodon: (se-

quame, strand, stopcodon) combination, Note: Not longest per transcript! You can also use function longestORFs after creation of ORFs for same result.

minimumLength (integer) Default is 0. Which is START + STOP = 6 bp. Minimum length

of ORF, without counting 3bps for START and STOP codons. For example minimumLength = 8 will result in size of ORFs to be at least START + 8\*3 (bp)

+ STOP = 30 bases. Use this param to restrict search.

findPeaksPerGene 159

is.circular

(logical) Whether the genome in filePath is circular. Prokaryotic genomes are usually circular. Be carefull if you want to extract sequences, remember that seqlengths must be set, else it does not know what last base in sequence is before loop ends!

#### **Details**

Remember if you have a fasta file of transcripts (transcript coordinates), delete all negative stranded ORFs afterwards by: orfs <- orfs[strandBool(orfs)] # negative strand orfs make no sense then. Seqnames are created from header by format: >name info, so name must be first after "biggern than" and space between name and info. Also make sure your fasta file is valid (no hidden spaces etc), as this might break the coordinate system!

## Value

(GRanges) object of ORFs mapped from fasta file. Positions are relative to the fasta file.

#### See Also

```
Other findORFs: findMapORFs(), findORFs(), findUORFs(), startDefinition(), stopDefinition()
```

# **Examples**

```
# location of the example fasta file
example_genome <- system.file("extdata/references/danio_rerio", "genome_dummy.fasta",
    package = "ORFik")
orfs <- findORFsFasta(example_genome)
# To store ORF sequences (you need indexed genome .fai file):
fa <- FaFile(example_genome)
names(orfs) <- paste0("ORF_", seq.int(length(orfs)), "_", seqnames(orfs))
orf_seqs <- getSeq(fa, orfs)
# You sequences (fa), needs to have isCircular(fa) == TRUE for it to work
# on circular wrapping ranges!
# writeXStringSet(DNAStringSet(orf_seqs), "orfs.fasta")</pre>
```

findPeaksPerGene

Find peaks per gene

## **Description**

For finding the peaks (stall sites) per gene, with some default filters. A peak is basically a position of very high coverage compared to its surrounding area, as measured using zscore.

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## Usage

```
findPeaksPerGene(
   tx,
   reads,
   top_tx = 0.5,
   min_reads_per_tx = 20,
   min_reads_per_peak = 10,
   type = "max",
   gene_ids = names(tx),
   coverage = coveragePerTiling(tx, reads, TRUE, as.data.table = TRUE)
)
```

#### **Arguments**

tx a GRangesList

reads a GAlignments or GRanges, must be 1 width reads like p-shifts, or other reads

that is single positioned. It will work with non 1 width bases, but you then get

larger areas for peaks.

top\_tx numeric, default 0.50 (only use 50% top transcripts by read counts).

min\_reads\_per\_tx

numeric, default 20. Gene must have at least 20 reads, applied before type filter.

min\_reads\_per\_peak

numeric, default 10. Peak must have at least 10 reads.

type character, default "max". Get only max peak per gene. Alternatives: "all", all

peaks passing the input filter will be returned. "median", only peaks that is higher than the median of all peaks. "maxmedian": get first "max", then median

of those.

gene\_ids character vector, names of genes, default names(tx)

coverage a data.table of coverage, with columns position, score and genes

#### **Details**

For more details see reference, which uses a slightly different method by zscore of a sliding window instead of over the whole tx.

#### Value

a data.table of gene\_id, position, counts of the peak, zscore and standard deviation of the peak compared to rest of gene area.

#### References

doi: 10.1261/rna.065235.117

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## **Examples**

```
df <- ORFik.template.experiment()
cds <- loadRegion(df, "cds")
# Load ribo seq from ORFik
rfp <- fimport(df[3,]$filepath)
# All transcripts passing filter
findPeaksPerGene(cds, rfp, top_tx = 0)
# Top 50% of genes
findPeaksPerGene(cds, rfp)</pre>
```

findUORFs

Find upstream ORFs from transcript annotation

## **Description**

Procedure: 1. Create a new search space starting with the 5' UTRs. 2. Redefine TSS with CAGE if wanted. 3. Add the whole of CDS to search space to allow uORFs going into cds. 4. find ORFs on that search space. 5. Filter out wrongly found uORFs, if CDS is included. The CDS, alternative CDS, uORFs starting within the CDS etc.

# Usage

```
findUORFs(
   fiveUTRs,
   fa,
   startCodon = startDefinition(1),
   stopCodon = stopDefinition(1),
   longestORF = TRUE,
   minimumLength = 0,
   cds = NULL,
   cage = NULL,
   extension = 1000,
   filterValue = 1,
   restrictUpstreamToTx = FALSE,
   removeUnused = FALSE
)
```

## **Arguments**

fiveUTRs (GRangesList) The 5' leaders or full transcript sequences

a FaFile. With fasta sequences corresponding to fiveUTR annotation. Usually loaded from the genome of an organism with fa = ORFik:::findFa("path/to/fasta/genome")

startCodon (character vector) Possible START codons to search for. Check startDefinition for helper function. Note that it is case sensitive, so "atg" would give 0 hits for a sequence with only capital "ATG" ORFs.

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stopCodon (character vector) Possible STOP codons to search for. Check stopDefinition

for helper function. Note that it is case sensitive, so "tga" would give 0 hits for

a sequence with only capital "TGA" ORFs.

longestORF (logical) Default TRUE. Keep only the longest ORF per unique stopcodon: (se-

quame, strand, stopcodon) combination, Note: Not longest per transcript! You can also use function longestORFs after creation of ORFs for same result.

can also use function longestures after creation of OREs for same result.

minimumLength (integer) Default is 0. Which is START + STOP = 6 bp. Minimum length

of ORF, without counting 3bps for START and STOP codons. For example minimumLength = 8 will result in size of ORFs to be at least START + 8\*3 (bp)

+ STOP = 30 bases. Use this param to restrict search.

cds (GRangesList) CDS of relative fiveUTRs, applicable only if you want to extend

5' leaders downstream of CDS's, to allow upstream ORFs that can overlap into

CDS's.

cage Either a filePath for the CageSeq file as .bed .bam or .wig, with possible com-

pressions (".gzip", ".gz", ".bgz"), or already loaded CageSeq peak data as GRanges or GAlignment. NOTE: If it is a .bam file, it will add a score column by running: convertToOneBasedRanges(cage, method = "5prime", addScoreColumn = TRUE) The score column is then number of replicates of read, if score column

is something else, like read length, set the score column to NULL first.

extension The maximum number of basses upstream of the TSS to search for CageSeq

peak.

filterValue The minimum number of reads on cage position, for it to be counted as possible

new tss. (represented in score column in CageSeq data) If you already filtered,

set it to 0.

restrictUpstreamToTx

a logical (FALSE). If TRUE: restrict leaders to not extend closer than 5 bases

from closest upstream leader, set this to TRUE.

removeUnused logical (FALSE), if False: (standard is to set them to original annotation), If

TRUE: remove leaders that did not have any cage support.

### **Details**

From default a filtering process is done to remove "fake" uORFs, but only if cds is included, since uORFs that stop on the stop codon on the CDS is not a uORF, but an alternative cds by definition, etc.

#### Value

A GRangesList of uORFs, 1 granges list element per uORF.

#### See Also

Other findORFs: findMapORFs(), findORFs(), findORFsFasta(), startDefinition(), stopDefinition()

findUORFs\_exp 163

## **Examples**

findUORFs\_exp

Find upstream ORFs from transcript annotation

## **Description**

Procedure: 1. Create a new search space starting with the 5' UTRs. 2. Redefine TSS with CAGE if wanted. 3. Add the whole of CDS to search space to allow uORFs going into cds. 4. find ORFs on that search space. 5. Filter out wrongly found uORFs, if CDS is included. The CDS, alternative CDS, uORFs starting within the CDS etc.

#### Usage

```
findUORFs_exp(
  df.
  faFile = findFa(df),
  leaders = loadRegion(txdb, "leaders"),
  startCodon = startDefinition(1),
  stopCodon = stopDefinition(1),
  longestORF = TRUE,
 minimumLength = 0,
  overlappingCDS = FALSE,
  cage = NULL,
  extension = 1000,
  filterValue = 1,
  restrictUpstreamToTx = FALSE,
  removeUnused = FALSE,
  save_optimized = FALSE
)
```

findUORFs\_exp

#### **Arguments**

df a txdb or experiment

faFile FaFile of genome, default findFa(df). Default only works for ORFik experi-

ments, if TxDb, input manually like: findFa(genome path)

leaders GRangesList, default: loadRegion(txdb, "leaders"). If you do not have any good

leader annotation, a hack is to use ORFik:::groupGRangesBy(startSites(loadRegion(txdb,

"cds"), asGR = TRUE, keep.names = TRUE, is.sorted = TRUE))

startCodon (character vector) Possible START codons to search for. Check startDefinition

for helper function. Note that it is case sensitive, so "atg" would give 0 hits for

a sequence with only capital "ATG" ORFs.

stopCodon (character vector) Possible STOP codons to search for. Check stopDefinition

for helper function. Note that it is case sensitive, so "tga" would give 0 hits for

a sequence with only capital "TGA" ORFs.

longestORF (logical) Default TRUE. Keep only the longest ORF per unique stopcodon: (se-

quame, strand, stopcodon) combination, Note: Not longest per transcript! You can also use function longestORFs after creation of ORFs for same result.

minimumLength (integer) Default is 0. Which is START + STOP = 6 bp. Minimum length

of ORF, without counting 3bps for START and STOP codons. For example minimumLength = 8 will result in size of ORFs to be at least START + 8\*3 (bp)

+ STOP = 30 bases. Use this param to restrict search.

overlappingCDS logical, default FALSE. Include uORFs that overlap CDS.

cage Either a filePath for the CageSeq file as .bed .bam or .wig, with possible com-

pressions (".gzip", ".gz", ".bgz"), or already loaded CageSeq peak data as GRanges or GAlignment. NOTE: If it is a .bam file, it will add a score column by running: convertToOneBasedRanges(cage, method = "5prime", addScoreColumn = TRUE) The score column is then number of replicates of read, if score column

is something else, like read length, set the score column to NULL first.

extension The maximum number of basses upstream of the TSS to search for CageSeq

peak.

filterValue The minimum number of reads on cage position, for it to be counted as possible

new tss. (represented in score column in CageSeq data) If you already filtered,

set it to 0.

restrictUpstreamToTx

a logical (FALSE). If TRUE: restrict leaders to not extend closer than 5 bases

from closest upstream leader, set this to TRUE.

removeUnused logical (FALSE), if False: (standard is to set them to original annotation), If

TRUE: remove leaders that did not have any cage support.

save\_optimized logical, default FALSE. If TRUE, save in the optimized folder for the exper-

iment. You must have made this directory before running this function (call

makeTxdbFromGenome first if not).

#### **Details**

From default a filtering process is done to remove "fake" uORFs, but only if cds is included, since uORFs that stop on the stop codon on the CDS is not a uORF, but an alternative cds by definition, etc.

find\_url\_ebi

#### Value

A GRangesList of uORFs, 1 granges list element per uORF.

## See Also

```
Other findORFs: findMapORFs(), findORFs(), findORFsFasta(), startDefinition(), stopDefinition()
```

## **Examples**

```
df <- ORFik.template.experiment()
# Without cds overlapping, no 5' leader extension
findUORFs_exp(df, extension = 0)
# Without cds overlapping, extends 5' leaders by 1000 (good for yeast etc)
findUORFs_exp(df)
# Include cds overlapping uorfs
findUORFs_exp(df, overlappingCDS = TRUE)</pre>
```

find\_url\_ebi

Locates and check if fastq files exists in ebi

## Description

Look for files in ebi following url: ftp://ftp.sra.ebi.ac.uk/vol1/fastq Paired end and single end fastq files.

EBI uses 3 ways to organize data inside vol1/fastq:

- 1: Most common: SRR(3 first)/0(2 last)/whole
- 2: less common: SRR(3 first)/00(1 last)/whole
- 3: least common SRR(3 first)/whole

# Usage

```
find_url_ebi(SRR, stop.on.error = FALSE, study = NULL)
```

## **Arguments**

SRR character, SRR, ERR or DRR numbers.

stop.on.error logical FALSE, if TRUE will stop if all files are not found. If FALSE returns

empty character vector if error is catched.

study default NULL, optional PRJ (study id) to speed up search for URLs.

#### Value

full url to fastq files, same length as input (2 urls for paired end data). Returns empty character() if all files not found.

find\_url\_ebi\_safe

## **Examples**

```
# Test the 3 ways to get fastq files from EBI
# Both single end and paired end data
# Most common: SRR(3 first)/0(2 last)/whole
# Single
ORFik:::find_url_ebi("SRR10503056")
# Paired
ORFik:::find_url_ebi("SRR10500056")
# less common: SRR(3 first)/00(1 last)/whole
# Single
#ORFik:::find_url_ebi("SRR1562873")
# Paired
#ORFik:::find_url_ebi("SRR1560083")
# least common SRR(3 first)/whole
# Single
#ORFik:::find_url_ebi("SRR105687")
# Paired
#ORFik:::find_url_ebi("SRR105788")
```

find\_url\_ebi\_safe

Find URL for EBI fastq files

# Description

Safer version

## Usage

```
find_url_ebi_safe(accession, SRR = NULL, stop.on.error = FALSE)
```

## Arguments

accession character: (PRJ, SRP, ERP, DRP, SRX, SRR, ERR,...). For studies or samples, it

returns all runs per study or sample.

SRR character, which SRR numbers to subset by (can also be ERR or DRR numbers)

stop.on.error logical FALSE, if TRUE will stop if all files are not found. If FALSE returns

empty character vector if error is catched.

## Value

character (1 element per SRR number)

firstEndPerGroup 167

firstEndPerGroup

Get first end per granges group

## **Description**

```
grl must be sorted, call ORFik:::sortPerGroup if needed
```

# Usage

```
firstEndPerGroup(grl, keep.names = TRUE)
```

# Arguments

```
grl a GRangesList
keep.names a boolean, keep names or not, default: (TRUE)
```

### Value

```
a Rle(keep.names = T), or integer vector(F)
```

## **Examples**

```
 \begin{split} \text{gr\_plus} <&- \text{GRanges}(\text{seqnames} = \text{c("chr1", "chr1")}, \\ &\quad \text{ranges} = \text{IRanges}(\text{c(7, 14), width} = 3), \\ &\quad \text{strand} = \text{c("+", "+")}) \\ \text{gr\_minus} <&- \text{GRanges}(\text{seqnames} = \text{c("chr2", "chr2")}, \\ &\quad \text{ranges} = \text{IRanges}(\text{c(4, 1), c(9, 3)}), \\ &\quad \text{strand} = \text{c("-", "-")}) \\ \text{grl} <&- \text{GRangesList}(\text{tx1} = \text{gr\_plus}, \text{tx2} = \text{gr\_minus}) \\ \text{firstEndPerGroup}(\text{grl}) \end{aligned}
```

firstExonPerGroup

Get first exon per GRangesList group

# Description

```
grl must be sorted, call ORFik:::sortPerGroup if needed
```

# Usage

```
firstExonPerGroup(grl)
```

## **Arguments**

```
grl a GRangesList
```

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## Value

a GRangesList of the first exon per group

# **Examples**

```
 \begin{split} \text{gr\_plus} <& - \text{GRanges}(\text{seqnames} = \text{c("chr1", "chr1")}, \\ & \text{ranges} = \text{IRanges}(\text{c(7, 14), width} = 3), \\ & \text{strand} = \text{c("+", "+")}) \\ \text{gr\_minus} <& - \text{GRanges}(\text{seqnames} = \text{c("chr2", "chr2")}, \\ & \text{ranges} = \text{IRanges}(\text{c(4, 1), c(9, 3)}), \\ & \text{strand} = \text{c("-", "-")}) \\ \text{grl} <& - \text{GRangesList}(\text{tx1} = \text{gr\_plus}, \text{tx2} = \text{gr\_minus}) \\ \text{firstExonPerGroup}(\text{grl}) \end{aligned}
```

firstStartPerGroup

Get first start per granges group

# **Description**

grl must be sorted, call ORFik:::sortPerGroup if needed

## Usage

```
firstStartPerGroup(grl, keep.names = TRUE)
```

## **Arguments**

```
grl a GRangesList
keep.names a boolean, keep names or not, default: (TRUE)
```

## Value

```
a Rle(keep.names = TRUE), or integer vector(FALSE)
```

## **Examples**

```
 \begin{split} \text{gr\_plus} <&- \text{GRanges}(\text{seqnames} = \text{c("chr1", "chr1")}, \\ &\quad \text{ranges} = \text{IRanges}(\text{c(7, 14), width} = 3), \\ &\quad \text{strand} = \text{c("+", "+")}) \\ \text{gr\_minus} <&- \text{GRanges}(\text{seqnames} = \text{c("chr2", "chr2")}, \\ &\quad \text{ranges} = \text{IRanges}(\text{c(4, 1), c(9, 3)}), \\ &\quad \text{strand} = \text{c("-", "-")}) \\ \text{grl} <&- \text{GRangesList}(\text{tx1} = \text{gr\_plus}, \text{tx2} = \text{gr\_minus}) \\ \text{firstStartPerGroup}(\text{grl}) \end{aligned}
```

fix\_malformed\_gff 169

fix\_malformed\_gff

Fix a malformed gff file

## **Description**

Basically removes all info lines with character length > 32768 and save that new file.

# Usage

```
fix_malformed_gff(gff)
```

## **Arguments**

gff

character, path to gtf, can not be gzipped!

## Value

path of fixed gtf

# **Examples**

```
# fix_malformed_gff("my_bad_gff.gff")
```

flankPerGroup

Get flanks per group

# Description

For a GRangesList, get start and end site, return back as GRangesList.

## Usage

```
flankPerGroup(grl)
```

## **Arguments**

grl

 $a\;\mathsf{GRangesList}$ 

# Value

a GRangesList, 1 GRanges per group with: start as minimum start of group and end as maximum per group.

## **Examples**

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floss

Fragment Length Organization Similarity Score

## **Description**

This feature is usually calcualted only for RiboSeq reads. For reads of width between 'start' and 'end', sum the fraction of RiboSeq reads (per read widths) that overlap ORFs and normalize by CDS read width fractions. So if all read length are width 34 in ORFs and CDS, value is 1. If width is 33 in ORFs and 34 in CDS, value is 0. If width is 33 in ORFs and 50/50 (33 and 34) in CDS, values will be 0.5 (for 33).

## Usage

```
floss(grl, RFP, cds, start = 26, end = 34, weight = 1L)
```

## **Arguments**

_	
grl	a GRangesList object can be either transcripts, 5' utrs, cds', 3' utrs or ORFs as a special case (uORFs, potential new cds' etc). If regions are not spliced you can send a GRanges object.
RFP	ribosomal footprints, given as GAlignments or GRanges object, must be already shifted and resized to the p-site. Requires a \$size column with original read lengths.
cds	a GRangesList of coding sequences, cds has to have names as grl so that they can be matched
start	usually 26, the start of the floss interval (inclusive)
end	usually 34, the end of the floss interval (inclusive)
weight	a numeric/integer vector or metacolumn name. (default: 1L, no differential weighting). If weight is name of defined meta column in reads object, it gives the number of times a read was found at that position. GRanges("chr1", 1, "+", score = 5), would mean "score" column tells that this alignment region was found 5 times. if 1L it means each read is weighted equal as 1, this is what among others countOverlaps() presumes, if single number (!= 1), it repeats for all ranges, if vector with length > 1, it must be equal size of the reads object.

#### **Details**

Pseudo explanation of the function:

```
SUM[start to stop]((grl[start:end][name]/grl) / (cds[start:end][name]/cds))
```

Where 'name' is transcript names.

Please read more in the article.

## Value

a vector of FLOSS of length same as grl, 0 means no RFP reads in range, 1 is perfect match.

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#### References

```
doi: 10.1016/j.celrep.2014.07.045
```

#### See Also

```
Other features: computeFeatures(), computeFeaturesCage(), countOverlapsW(), disengagementScore(), distToCds(), distToTSS(), entropy(), fpkm(), fpkm_calc(), fractionLength(), initiationScore(), insideOutsideORF(), isInFrame(), isOverlapping(), kozakSequenceScore(), orfScore(), rankOrder(), ribosomeReleaseScore(), ribosomeStallingScore(), startRegion(), startRegionCoverage(), stopRegion(), subsetCoverage(), translationalEff()
```

## **Examples**

```
ORF1 <- GRanges(seqnames = "1",
               ranges = IRanges(start = c(1, 12, 22),
               end = c(10, 20, 32)),
               strand = "+")
grl \leftarrow GRangesList(tx1_1 = ORF1)
# RFP is 1 width position based GRanges
RFP <- GRanges("1", IRanges(c(1, 25, 35, 38), width = 1), "+")
RFP$size <- c(28, 28, 29) # original width in size col
cds <- GRangesList(tx1 = GRanges("1", IRanges(35, 44), "+"))</pre>
\# grl must have same names as cds + \_1 etc, so that they can be matched.
floss(grl, RFP, cds)
# or change ribosome start/stop, more strict
floss(grl, RFP, cds, 28, 28)
# With repeated alignments in score column
ORF2 <- GRanges(seqnames = "1",
               ranges = IRanges(start = c(12, 22, 36),
               end = c(20, 32, 38)),
               strand = "+")
grl \leftarrow GRangesList(tx1_1 = ORF1, tx1_2 = ORF2)
score(RFP) \leftarrow c(5, 10, 5, 10)
floss(grl, RFP, cds, weight = "score")
```

## **Description**

For internal use only!

## Usage

```
footprints.analysis(rw, heatmap, region = "start of CDS")
```

172 fpkm

## Arguments

rw a data.table of position, score and fraction (read length) of either TIS or TES

(translation end site, around 3' UTR)

heatmap a logical or character string, default FALSE. If TRUE, will plot heatmap of raw

reads before p-shifting to console, to see if shifts given make sense. You can

also set a filepath to save the file there.

region a character string, default "start of CDS"

#### Value

invisible(NULL)

fpkm

Create normalizations of overlapping read counts.

## Description

FPKM is short for "Fragments Per Kilobase of transcript per Million fragments in library". When calculating RiboSeq data FPKM over ORFs, use ORFs as 'grl'. When calculating RNASeq data FPKM, use full transcripts as 'grl'. It is equal to RPKM given that you do not have paired end reads.

## Usage

```
fpkm(grl, reads, pseudoCount = 0, librarySize = "full", weight = 1L)
```

#### **Arguments**

grl a GRangesList object can be either transcripts, 5' utrs, cds', 3' utrs or ORFs as

a special case (uORFs, potential new cds' etc). If regions are not spliced you

can send a GRanges object.

reads a GAlignments, GRanges or GRangesList object, usually of RiboSeq, RnaSeq,

CageSeq, etc.

pseudoCount a numeric, default 0, set it to 1 if you want to avoid NA and inf values.

librarySize either numeric value or character vector. Default ("full"), number of align-

ments in library (reads). If you just have a subset, you can give the value by librarySize = length(wholeLib) or sum(wholeLib\$score), if you want lib size to be only number of reads overlapping grl, do: librarySize = "overlapping" sum(countOverlaps(reads, grl) > 0), if reads[1] has 3 hits in grl, and reads[2] has 2 hits, librarySize will be 2, not 5. You can also get the inverse overlap, if you want lib size to be total number of overlaps, do: librarySize = "DESeq" This is standard fpkm way of DESeq2::fpkm(robust = FALSE) sum(countOverlaps(grl, reads)) if grl[1] has 3 reads and grl[2] has 2 reads, librarySize is 5, not 2.

fpkm 173

weight

a numeric/integer vector or metacolumn name. (default: 1L, no differential weighting). If weight is name of defined meta column in reads object, it gives the number of times a read was found at that position. GRanges("chr1", 1, "+", score = 5), would mean "score" column tells that this alignment region was found 5 times. if 1L it means each read is weighted equal as 1, this is what among others countOverlaps() presumes, if single number (!= 1), it repeats for all ranges, if vector with length > 1, it must be equal size of the reads object.

#### **Details**

Note also that you must consider if you will use the whole read library or just the reads overlapping 'grl' for library size. A normal question here is, does it make sense to include rRNA in library size? If you only want overlapping grl, do: librarySize = "overlapping"

#### Value

a numeric vector with the fpkm values

#### References

doi: 10.1038/nbt.1621

#### See Also

```
Other features: computeFeatures(), computeFeaturesCage(), countOverlapsW(), disengagementScore(), distToCds(), distToTSS(), entropy(), floss(), fpkm_calc(), fractionLength(), initiationScore(), insideOutsideORF(), isInFrame(), isOverlapping(), kozakSequenceScore(), orfScore(), rankOrder(), ribosomeReleaseScore(), ribosomeStallingScore(), startRegion(), startRegionCoverage(), stopRegion(), subsetCoverage(), translationalEff()
```

#### **Examples**

174 fractionLength

_			_	
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Create normalizations of read counts

## **Description**

A helper for [fpkm()] Normally use function [fpkm()], if you want unusual normalization , you can use this. Short for: Fragments per kilobase of transcript per million fragments Normally used in Translations efficiency calculations

## Usage

```
fpkm_calc(counts, lengthSize, librarySize)
```

#### **Arguments**

counts a list, # of read hits per group
lengthSize a list of lengths per group

librarySize a numeric of size 1, the # of reads in library

#### Value

a numeric vector

#### References

doi: 10.1038/nbt.1621

## See Also

Other features: computeFeatures(), computeFeaturesCage(), countOverlapsW(), disengagementScore(), distToCds(), distToTSS(), entropy(), floss(), fpkm(), fractionLength(), initiationScore(), insideOutsideORF(), isInFrame(), isOverlapping(), kozakSequenceScore(), orfScore(), rankOrder(), ribosomeReleaseScore(), ribosomeStallingScore(), startRegion(), startRegionCoverage(), stopRegion(), subsetCoverage(), translationalEff()

fractionLength

Fraction Length

## **Description**

```
Fraction Length is defined as
```

```
(widths of grl)/tx_len
```

so that each group in the grl is divided by the corresponding transcript.

fractionLength 175

## Usage

```
fractionLength(grl, tx_len = widthPerGroup(tx, TRUE), tx = NULL)
```

## Arguments

grl	a GRangesList object with usually either leaders, cds', 3' utrs or ORFs. ORFs are a special case, see argument tx_len
tx_len	the transcript lengths of the transcripts, a named (tx names) vector of integers. If you have the transcripts as GRangesList, call 'ORFik:::widthPerGroup(tx, TRUE)'.
	If you used CageSeq to reannotate leaders, then the tss for the the leaders have changed, therefore the tx lengths have changed. To account for that call: 'tx_len <- widthPerGroup(extendLeaders(tx, cageFiveUTRs))' and calculate fraction length using 'fractionLength(grl, tx_len)'.
tx	default NULL, a GRangesList object of transcript to get lengths from. Pass in

default NULL, a GRangesList object of transcript to get lengths from. Pass in for wrapping to widths inside the function.

## Value

a numeric vector of ratios

### References

```
doi: 10.1242/dev.098343
```

## See Also

```
Other features: computeFeatures(), computeFeaturesCage(), countOverlapsW(), disengagementScore(), distToCds(), distToTSS(), entropy(), floss(), fpkm(), fpkm_calc(), initiationScore(), insideOutsideORF(), isInFrame(), isOverlapping(), kozakSequenceScore(), orfScore(), rankOrder(), ribosomeReleaseScore(), ribosomeStallingScore(), startRegion(), startRegionCoverage(), stopRegion(), subsetCoverage(), translationalEff()
```

# **Examples**

176 fread.bed

fractionNames

Get cell fraction name variants

## **Description**

Used to standardize nomeclature for experiments. Example: cytosolic, mitochondrial, specific gene knock down

## Usage

fractionNames()

#### Value

a data.table with 2 columns, the main name, and all name variants of the main name in second column as a list.

#### See Also

```
Other experiment_naming: batchNames(), cellLineNames(), cellTypeNames(), conditionNames(), inhibitorNames(), libNames(), mainNames(), repNames(), stageNames(), tissueNames()
```

fread.bed

Load bed file as GRanges

## **Description**

Wraps around import.bed and tries to speed up loading with the use of data.table. Supports gzip, gz, bgz and bed formats. Also safer chromosome naming with the argument chrStyle

## Usage

```
fread.bed(filePath, chrStyle = NULL)
```

## **Arguments**

filePath The location of the bed file

chrStyle a GRanges object, TxDb, FaFile, , a seqlevelsStyle or Seqinfo. (Default:

NULL) to get seqlevelsStyle from. In addition if it is a Seqinfo object, seqinfo will be updated. Example of seqlevelsStyle update: Is chromosome 1 called chr1 or 1, is mitocondrial chromosome called MT or chrM etc. Will use 1st seqlevel-style if more are present. Like: c("NCBI", "UCSC") -> pick "NCBI"

## Value

a GRanges object

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## See Also

```
Other utils: bedToGR(), convertToOneBasedRanges(), export.bed12(), export.bigWig(), export.fstwig(), export.wiggle(), fimport(), findFa(), optimizeReads(), readBam(), readBigWig(), readWig()
```

## **Examples**

gcContent

Get GC content

## **Description**

0.5 means 50

#### Usage

```
gcContent(seqs, fa = NULL)
```

## **Arguments**

seqs a character vector of sequences, or ranges as GRangesList

fa fasta index file .fai file, either path to it, or the loaded FaFile, default (NULL),
only set if you give ranges as GRangesList

## Value

a numeric vector of gc content scores

## **Examples**

178 geneToSymbol

geneToSymbol

Get gene symbols from Ensembl gene ids

## **Description**

If your organism is not in this list of supported organisms, manually assign the input arguments. There are 2 main fetch modes:

By gene ids (Single accession per gene)

By tx ids (Multiple accessions per gene)

Run the mode you need depending on your required attributes.

Will check for already existing table of all genes, and use that instead of re-downloading every time (If you input valid experiment or txdb and have run makeTxdbFromGenome with symbols = TRUE, you have a file called gene\_symbol\_tx\_table.fst) will load instantly. If df = NULL, it can still search cache to load a bit slower.

## Usage

```
geneToSymbol(
    df,
    organism_name = organism(df),
    gene_ids = filterTranscripts(df, by = "gene", 0, 0, 0),
    org.dataset = paste0(tolower(substr(organism_name, 1, 1)), gsub(".* ", replacement =
        "", organism_name), "_gene_ensembl"),
    ensembl = biomaRt::useEnsembl("ensembl", dataset = org.dataset),
    attribute = "external_gene_name",
    include_tx_ids = FALSE,
    uniprot_id = FALSE,
    force = FALSE,
    verbose = TRUE
)
```

# Arguments

df	an ORFik experiment or TxDb object with defined organism slot. If set will look for file at path of txdb / experiment reference path named: 'gene_symbol_tx_table.fst' relative to the txdb/genome directory. Can be set to NULL if gene_ids and organism is defined manually.
organism_name	default, organism(df). Scientific name of organism, like ("Homo sapiens"), remember capital letter for first name only!
gene_ids	default, filterTranscripts(df, by = "gene", 0, 0, 0). Ensembl gene IDs to search for (default all transcripts coding and noncoding) To only get coding do: filterTranscripts(df, by = "gene", 0, 1, 0)
org.dataset	default, paste0(tolower(substr(organism_name, 1, 1)), gsub(".*", replacement = "", organism_name), "_gene_ensembl") the ensembl dataset to use. For Homo sapiens, this converts to default as: hsapiens_gene_ensembl

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ensembl default, useEnsembl("ensembl",dataset=org.dataset) .The mart connec-

tion.

attribute default, "external\_gene\_name", the biomaRt column / columns default(primary

gene symbol names). These are always from specific database, like hgnc symbol

for human, and mgi symbol for mouse and rat, sgd for yeast etc.

include\_tx\_ids logical, default FALSE, also match tx ids, which then returns as the 3rd column.

Only allowed when 'df' is defined. If

uniprot\_id logical, default FALSE. Include uniprotsptrembl and/or uniprotswissprot. If in-

clude\_tx\_ids you will get per isoform if available, else you get canonical uniprot id per gene. If both uniprotsptrembl and uniprotswissprot exists, it will make a merged uniprot id column with rule: if id exists in uniprotswissprot, keep. If

not, use uniprotsptrembl column id.

force logical FALSE, if TRUE will not look for existing file made through makeTxdbFromGenome

corresponding to this txdb / ORFik experiment stored with name "gene\_symbol\_tx\_table.fst".

verbose logical TRUE, if FALSE, do not output messages.

#### Value

data.table with 2, 3 or 4 columns: gene\_id, gene\_symbol, tx\_id and uniprot\_id named after attribute, sorted in order of gene\_ids input. (example: returns 3 columns if include\_tx\_ids is TRUE), and more if additional columns are specified in 'attribute' argument.

#### **Examples**

```
## Without ORFik experiment input
gene_id_ATF4 <- "ENSG00000128272"
#geneToSymbol(NULL, organism_name = "Homo sapiens", gene_ids = gene_id_ATF4)
# With uniprot canonical isoform id:
#geneToSymbol(NULL, organism_name = "Homo sapiens", gene_ids = gene_id_ATF4, uniprot_id = TRUE)
## All genes from Organism using ORFik experiment
# df <- read.experiment("some_experiment)
# geneToSymbol(df)

## Non vertebrate species (the ones not in ensembl, but in ensemblGenomes mart)
#txdb_ylipolytica <- loadTxdb("txdb_path")
#dt2 <- geneToSymbol(txdb_ylipolytica, include_tx_ids = TRUE,
# ensembl = useEnsemblGenomes(biomart = "fungi_mart", dataset = "ylipolytica_eg_gene"))</pre>
```

getGAlignments

Internal GAlignments loader from fst data.frame

## Description

Internal GAlignments loader from fst data.frame

180 getGAlignmentsPairs

## Usage

```
getGAlignments(df, seqinfo = NULL)
```

## **Arguments**

df a data.frame with columns minimum 4 columns: seqnames, start ("pos" in final

GA object), cigar and strand.

Additional columns will be assigned as meta columns

seqinfo Seqinfo object, defaul NULL (created from ranges). Add to avoid warnings later

on differences in seqinfo.

## Value

GAlignments object

getGAlignmentsPairs Internal GAlignmentPairs loader from fst data.frame

## **Description**

Internal GAlignmentPairs loader from fst data.frame

#### Usage

```
getGAlignmentsPairs(df, strandMode = 0, seqinfo = NULL)
```

#### **Arguments**

df a data.frame with columns minimum 6 columns: seqnames, start1/start2 (inte-

gers), cigar1/cigar2 and strand

Additional columns will be assigned as meta columns

strandMode numeric, default 0. Only used for paired end bam files. One of (0: strand

= \*, 1: first read of pair is +, 2: first read of pair is -). See ?strandMode. Note: Sets default to 0 instead of 1, as readGAlignmentPairs uses 1. This is to guarantee hits, but will also make mismatches of overlapping transcripts in

opposite directions.

seqinfo Seqinfo object, defaul NULL (created from ranges). Add to avoid warnings later

on differences in seqinfo.

#### Value

GAlignmentPairs object

getGenomeAndAnnotation

Download genome (fasta), annotation (GTF) and contaminants

#### **Description**

This function automatically downloads (if files not already exists) genomes and contaminants specified for genome alignment. By default, it will use ensembl reference, upon completion, the function will store a file called file.path(output.dir, "outputs.rds") with the output paths of your completed genome/annotation downloads. For most non-model nonvertebrate organisms, you need my fork of biomartr for it to work: remotes::install\_github("Roleren/biomartr) If you misspelled something or crashed, delete wrong files and run again.

Do remake = TRUE, to do it all over again.

#### Usage

```
getGenomeAndAnnotation(
  organism,
  output.dir,
  db = "ensembl",
 GTF = TRUE,
  genome = TRUE,
 merge_contaminants = TRUE,
  phix = FALSE,
  ncRNA = FALSE,
  tRNA = FALSE,
  rRNA = FALSE,
  gunzip = TRUE,
  remake = FALSE,
  assembly_type = c("primary_assembly", "toplevel"),
  optimize = FALSE,
  gene_symbols = FALSE,
  uniprot_id = FALSE,
  pseudo_5UTRS_if_needed = NULL,
  remove_annotation_outliers = TRUE,
  notify_load_existing = TRUE,
 assembly = organism,
  refseq_genbank_format = c("gtf", "gff3")[1]
)
```

### **Arguments**

```
organism scientific name of organism, Homo sapiens, Danio rerio, Mus musculus, etc. See biomartr:::get.ensembl.info() for full list of supported organisms.

output.dir directory to save downloaded data
```

db

database to use for genome and GTF, default adviced: "ensembl" (remember to set assembly\_type to "primary\_assembly", else it will contain haplotypes, very large file!). Alternatives: "refseq" (reference assemblies) and "genbank" (all assemblies)

**GTF** 

logical, default: TRUE, download gtf of organism specified in "organism" argument. If FALSE, check if the downloaded file already exist. If you want to use a custom gtf from you hard drive, set GTF = FALSE, and assign:

annotation <- getGenomeAndAnnotation(gtf = FALSE)

annotation["gtf"] = "path/to/gtf.gtf".

If db is not "ensembl", you will instead get a gff file.

genome

logical, default: TRUE, download genome of organism specified in "organism" argument. If FALSE, check if the downloaded file already exist. If you want to use a custom gtf from you hard drive, set GTF = FALSE, and assign: annotation <- getGenomeAndAnnotation(genome = FALSE) annotation["genome"] = "path/to/genome.fasta".

Will download the primary assembly from Ensembl.

merge\_contaminants

logical, default TRUE. Will merge the contaminants specified into one fasta file, this considerably saves space and is much quicker to align with STAR than each contaminant on it's own. If no contaminants are specified, this is ignored.

phix

logical, default FALSE, download phiX sequence to filter out Illumina control reads. ORFik defines Phix as a contaminant genome. Phix is used in Illumina sequencers for sequencing quality control. Genome is: refseq, Escherichia phage phiX174. If sequencing facility created fastq files with the command bcl2fastq, then there should be very few phix reads left in the fastq files recieved.

ncRNA

logical or character, default FALSE (not used, no download), if TRUE or defined path, ncRNA is used as a contaminant reference. If TRUE, will try to find ncRNA sequences from the gtf file, usually represented as lncRNA (long noncoding RNA's). Will let you know if no ncRNA sequences were found in gtf. If not found try character input:

Alternatives; "auto": Will try to find ncRNA file on NONCODE from organism, Homo sapiens -> human etc. "auto" will not work for all, then you must specify the name used by NONCODE, go to the link below and find it. If not "auto" / "" it must be a character vector of species common name (not scientific name) Homo sapiens is human, Rattus norwegicus is rat etc, download ncRNA sequence to filter out with. From NONCODE online server, if you cant find common name see: http://www.noncode.org/download.php/

tRNA

logical or character, default FALSE (not used, no download), tRNA is used as a contaminant genome. If TRUE, will try to find tRNA sequences from the gtf file, usually represented as Mt\_tRNA (mature tRNA's). Will let you know if no tRNA sequences were found in gtf. If not found try character input:

if not "" it must be a character vector to valid path of mature tRNAs fasta file to remove as contaminants on your disc. Find and download your wanted mtRNA at: http://gtrnadb.ucsc.edu/, or run trna-scan on you genome.

rRNA

logical or character, default FALSE (not used, no download), rRNA is used as a contaminant reference If TRUE, will try to find rRNA sequences from the gtf

file, usually represented as rRNA (ribosomal RNA's). Will let you know if no rRNA sequences were found in gtf. If not found you can try character input: If "silva" will download silva SSU & LSU sequences for all species (250MB file) and use that. If you want a smaller file go to https://www.arb-silva.de/ If not "" or "silva" it must be a character vector to valid path of mature rRNA fasta file to remove as contaminants on your disc.

gunzip

logical, default TRUE, uncompress downloaded files that are zipped when downloaded, should be TRUE!

remake

logical, default: FALSE, if TRUE remake everything specified

assembly\_type

character, default c("primary\_assembly", "toplevel"). Used for ensembl only, specifies the genome assembly type. Searches for both primary and toplevel, and if both are found, uses the first by order (so primary is prioritized by default). The Primary assembly should usually be used if it exists. The "primary assembly" contains all the top-level sequence regions, excluding alternative haplotypes and patches. If the primary assembly file is not present for a species (only defined for standard model organisms), that indicates that there were no haplotype/patch regions, and in such cases, the 'toplevel file is used. For more details see: ensembl tutorial

optimize

logical, default FALSE. Create a folder within the output folder (defined by txdb\_file\_out\_path), that includes optimized objects to speed up loading of annotation regions from up to 15 seconds on human genome down to 0.1 second. ORFik will then load these optimized objects instead. Currently optimizes filterTranscript() function and loadRegion() function for 5' UTRs, 3' UTRs, CDS, mRNA (all transcript with CDS) and tx (all transcripts).

gene\_symbols

logical default FALSE. If TRUE, will download and store all gene symbols for all transcripts (coding and noncoding)- In a file called: "gene\_symbol\_tx\_table.fst" in same folder as txdb. hgcn for human, mouse symbols for mouse and rat, more to be added.

uniprot\_id

logical default FALSE. If TRUE, will download and store all uniprot id for all transcripts (coding and noncoding)- In a file called: "gene\_symbol\_tx\_table.fst" in same folder as txdb.

pseudo\_5UTRS\_if\_needed

integer, default NULL. If defined > 0, will add pseudo 5' UTRs of maximum this length if 'minimum\_5UTR\_percentage" (default 30 mRNAs (coding transcripts) do not have a leader. (NULL and 0 are both the ignore command)

remove\_annotation\_outliers

logical, default TRUE. Only for refseq. shall outlier lines be removed from the input annotation\_file? If yes, then the initial annotation\_file will be overwritten and the removed outlier lines will be stored at tempdir for further exploration. Among others Aridopsis refseq contains malformed lines, where this is needed

notify\_load\_existing

logical, default TRUE. If annotation exists (defined as: locally (a file called outputs.rds) exists in outputdir), print a small message notifying the user it is not redownloading. Set to FALSE, if this is not wanted

assembly

character, default is assembly = organism, which means getting the first assembly in list, otherwise the name of the assembly wanted, like "GCA\_00005845"

will get ecoli substrain k12, which is the most used ones for references. Usually ignore this for non bacterial species.

refseq\_genbank\_format

= c("gtf", "gff3")[1] Gtf format files are usually more secure from bugs downstream, so we highly advice to use them. GFF3 files can sometimes include information you might not find in the gtf, so sometimes it makes sense to use it.

#### **Details**

Some files that are made after download:

- A fasta index for the genome
- A TxDb to speed up GTF/GFF reading
- Seperat of merged contaminant files

Files that can be made:

- Gene symbols (hgnc, etc)
- Uniprot ids (For name of protein structures)

If you want custom genome or gtf from you hard drive, assign existing paths like this: annotation <- getGenomeAndAnnotation(GTF = "path/to/gtf.gtf", genome = "path/to/genome.fasta")

#### Value

a named character vector of path to genomes and gtf downloaded, and additional contaminants if used. If merge\_contaminants is TRUE, will not give individual fasta files to contaminants, but only the merged one.

#### References

https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4919035/

#### See Also

```
Other STAR: STAR.align.folder(), STAR.align.single(), STAR.allsteps.multiQC(), STAR.index(), STAR.install(), STAR.multiQC(), STAR.remove.crashed.genome(), install.fastp()
```

```
## Get Saccharomyces cerevisiae genome and gtf (create txdb for R)
#getGenomeAndAnnotation("Saccharomyces cerevisiae", tempdir(), assembly_type = "toplevel")
## Download and add pseudo 5' UTRs
#getGenomeAndAnnotation("Saccharomyces cerevisiae", tempdir(), assembly_type = "toplevel",
# pseudo_5UTRS_if_needed = 100)
## Get Danio rerio genome and gtf (create txdb for R)
#getGenomeAndAnnotation("Danio rerio", tempdir())

output.dir <- "/Bio_data/references/zebrafish"
## Get Danio rerio and Phix contamints to deplete during alignment
#getGenomeAndAnnotation("Danio rerio", output.dir, phix = TRUE)
## Optimize for ORFik (speed up for large annotations like human or zebrafish)</pre>
```

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```
#getGenomeAndAnnotation("Danio rerio", tempdir(), optimize = TRUE)
# Drosophila melanogaster (toplevel exists only)
#getGenomeAndAnnotation("drosophila melanogaster", output.dir = file.path(config["ref"],
# "Drosophila_melanogaster_BDGP6"), assembly_type = "toplevel")
## How to save malformed refseq gffs:
## First run function and let it crash:
#annotation <- getGenomeAndAnnotation(organism = "Arabidopsis thaliana",</pre>
# output.dir = "~/Desktop/test_plant/",
# assembly_type = "primary_assembly", db = "refseq")
## Then apply a fix (example for linux, too long rows):
# fixed_gff <- fix_malformed_gff("~/Desktop/test_plant/Arabidopsis_thaliana_genomic_refseq.gff")
## Then updated arguments:
# annotation <- c(fixed_gff, "~/Desktop/test_plant/Arabidopsis_thaliana_genomic_refseq.fna")
# names(annotation) <- c("gtf", "genome")</pre>
# Then make the txdb (for faster R use)
# makeTxdbFromGenome(annotation["gtf"], annotation["genome"], organism = "Arabidopsis thaliana")
```

getGRanges

Internal GRanges loader from fst data.frame

### Description

Internal GRanges loader from fst data.frame

#### Usage

```
getGRanges(df, seqinfo = NULL)
```

# Arguments

df a data.frame with columns minimum 4 columns: seqnames, start, strand

Additional specific columns are:

- width (if not set, width is set to 1 for all reads) Additional columns will be assigned as meta columns

seqinfo Seqinfo object, defaul NULL (created from ranges). Add to avoid warnings later

on differences in seqinfo.

#### Value

GRanges object

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getGtfPathFromTxdb

Get path of GTF that created txdb

# Description

Will crash and report proper error if no gtf is found

# Usage

```
getGtfPathFromTxdb(txdb, stop.error = TRUE)
```

# Arguments

txdb a loaded TxDb object

stop.error logical TRUE, stop if Txdb does not have a gtf. If FALSE, return NULL.

# Value

a character file path, returns NULL if not valid and stop.error is FALSE.

getNGenesCoverage

Get number of genes per coverage table

# Description

Used to count genes in ORFik meta plots

# Usage

```
getNGenesCoverage(coverage)
```

# Arguments

coverage

a data.table with coverage

# Value

number of genes in coverage

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getWeights

Get weights from a subject GenomicRanges object

### **Description**

Get weights from a subject GenomicRanges object

# Usage

```
getWeights(subject, weight = 1L)
```

### Arguments

subject

a GRanges, IRanges, GAlignment, GAlignmentPairs or covRle object

weight

a numeric/integer vector or metacolumn name. (default: 1L, no differential weighting). If weight is name of defined meta column in reads object, it gives the number of times a read was found at that position. GRanges("chr1", 1, "+", score = 5), would mean "score" column tells that this alignment region was found 5 times. if 1L it means each read is weighted equal as 1, this is what among others countOverlaps() presumes, if single number (!= 1), it repeats for all ranges, if vector with length > 1, it must be equal size of the reads object.

# Value

a numeric vector of weights of equal size to subject

```
get_bioproject_candidates
```

Query eutils for bioproject IDs

# Description

The default query of Ribosome Profiling human, will result in internal entrez search of: Ribosome[All Fields] AND Profiling[All Fields] AND ("Homo sapiens"[Organism] OR human[All Fields])

# Usage

```
get_bioproject_candidates(
  term = "Ribosome Profiling human",
  as_accession = TRUE,
  add_study_title = FALSE,
  RetMax = 10000
)
```

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#### **Arguments**

term character, default "Ribosome Profiling human". A space is translated into AND,

that means "Ribosome AND Profiling AND human", will give same as above.

To do OR operation, do: "Ribosome OR profiling OR human".

as\_accession logical, default TRUE. Get bioproject accessions: PRJNA, PRJEB, PRJDB val-

ues, or IDs (FALSE), numbers only. Accessions are usually the thing needed for

most tools.

add\_study\_title

logical, default FALSE. If TRUE, return as data table with 2 columns: id: ID or

accessions. title: The title of the study.

RetMax integer, default 10000. How many IDs to return maximum

### Value

character vector of Accessions or IDs. If add\_study\_title is TRUE, returns a data.table.

#### References

https://www.ncbi.nlm.nih.gov/books/NBK25501/

### See Also

```
Other sra: browseSRA(), download.SRA(), download.SRA.metadata(), download.ebi(), install.sratoolkit(), rename.SRA.files()
```

# **Examples**

```
term <- "Ribosome Profiling Saccharomyces cerevisiae"
# get_bioproject_candidates(term)</pre>
```

get\_genome\_fasta

Download genome (fasta), annotation (GTF) and contaminants

# Description

This function automatically downloads (if files not already exists) genomes and contaminants specified for genome alignment. By default, it will use ensembl reference, upon completion, the function will store a file called file.path(output.dir, "outputs.rds") with the output paths of your completed genome/annotation downloads. For most non-model nonvertebrate organisms, you need my fork of biomartr for it to work: remotes::install\_github("Roleren/biomartr) If you misspelled something or crashed, delete wrong files and run again.

Do remake = TRUE, to do it all over again.

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#### Usage

```
get_genome_fasta(
  genome,
  output.dir,
  organism,
  assembly,
  assembly_type,
  db,
  gunzip
)
```

# **Arguments**

genome logical, default: TRUE, download genome of organism specified in "organism"

argument. If FALSE, check if the downloaded file already exist. If you want to

use a custom gtf from you hard drive, set GTF = FALSE, and assign: annotation <- getGenomeAndAnnotation(genome = FALSE)

annotation["genome"] = "path/to/genome.fasta".
Will download the primary assembly from Ensembl.

output.dir directory to save downloaded data

organism scientific name of organism, Homo sapiens, Danio rerio, Mus musculus, etc.

See biomartr:::get.ensembl.info() for full list of supported organisms.

assembly character, default is assembly = organism, which means getting the first assem-

bly in list, otherwise the name of the assembly wanted, like "GCA\_000005845" will get ecoli substrain k12, which is the most used ones for references. Usually

ignore this for non bacterial species.

assembly\_type character, default c("primary\_assembly", "toplevel"). Used for ensembl only,

specifies the genome assembly type. Searches for both primary and toplevel, and if both are found, uses the first by order (so primary is prioritized by default). The Primary assembly should usually be used if it exists. The "primary assembly" contains all the top-level sequence regions, excluding alternative haplotypes and patches. If the primary assembly file is not present for a species (only defined for standard model organisms), that indicates that there were no haplotype/patch regions, and in such cases, the 'toplevel file is used. For more

details see: ensembl tutorial

db database to use for genome and GTF, default adviced: "ensembl" (remember to

set assembly\_type to "primary\_assembly", else it will contain haplotypes, very large file!). Alternatives: "refseq" (reference assemblies) and "genbank" (all

assemblies)

gunzip logical, default TRUE, uncompress downloaded files that are zipped when down-

loaded, should be TRUE!

### **Details**

Some files that are made after download:

- A fasta index for the genome
- A TxDb to speed up GTF/GFF reading

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- Seperat of merged contaminant files

Files that can be made:

- Gene symbols (hgnc, etc)
- Uniprot ids (For name of protein structures)

If you want custom genome or gtf from you hard drive, assign existing paths like this: annotation <- getGenomeAndAnnotation(GTF = "path/to/gtf.gtf", genome = "path/to/genome.fasta")

#### Value

a named character vector of path to genomes and gtf downloaded, and additional contaminants if used. If merge\_contaminants is TRUE, will not give individual fasta files to contaminants, but only the merged one.

#### References

https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4919035/

#### See Also

```
Other STAR: STAR.align.folder(), STAR.align.single(), STAR.allsteps.multiQC(), STAR.index(), STAR.install(), STAR.multiQC(), STAR.remove.crashed.genome(), install.fastp()
```

```
## Get Saccharomyces cerevisiae genome and gtf (create txdb for R)
#getGenomeAndAnnotation("Saccharomyces cerevisiae", tempdir(), assembly_type = "toplevel")
## Download and add pseudo 5' UTRs
#getGenomeAndAnnotation("Saccharomyces cerevisiae", tempdir(), assembly_type = "toplevel",
# pseudo_5UTRS_if_needed = 100)
## Get Danio rerio genome and gtf (create txdb for R)
#getGenomeAndAnnotation("Danio rerio", tempdir())
output.dir <- "/Bio_data/references/zebrafish"</pre>
## Get Danio rerio and Phix contamints to deplete during alignment
#getGenomeAndAnnotation("Danio rerio", output.dir, phix = TRUE)
## Optimize for ORFik (speed up for large annotations like human or zebrafish)
#getGenomeAndAnnotation("Danio rerio", tempdir(), optimize = TRUE)
# Drosophila melanogaster (toplevel exists only)
#getGenomeAndAnnotation("drosophila melanogaster", output.dir = file.path(config["ref"],
# "Drosophila_melanogaster_BDGP6"), assembly_type = "toplevel")
## How to save malformed refseq gffs:
## First run function and let it crash:
#annotation <- getGenomeAndAnnotation(organism = "Arabidopsis thaliana",
# output.dir = "~/Desktop/test_plant/",
# assembly_type = "primary_assembly", db = "refseq")
## Then apply a fix (example for linux, too long rows):
# fixed_gff <- fix_malformed_gff("~/Desktop/test_plant/Arabidopsis_thaliana_genomic_refseq.gff")
## Then updated arguments:
# annotation <- c(fixed_gff, "~/Desktop/test_plant/Arabidopsis_thaliana_genomic_refseq.fna")
```

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```
# names(annotation) <- c("gtf", "genome")
# Then make the txdb (for faster R use)
# makeTxdbFromGenome(annotation["gtf"], annotation["genome"], organism = "Arabidopsis thaliana")</pre>
```

get\_genome\_gtf

Download genome (fasta), annotation (GTF) and contaminants

# **Description**

This function automatically downloads (if files not already exists) genomes and contaminants specified for genome alignment. By default, it will use ensembl reference, upon completion, the function will store a file called file.path(output.dir, "outputs.rds") with the output paths of your completed genome/annotation downloads. For most non-model nonvertebrate organisms, you need my fork of biomartr for it to work: remotes::install\_github("Roleren/biomartr) If you misspelled something or crashed, delete wrong files and run again.

Do remake = TRUE, to do it all over again.

# Usage

```
get_genome_gtf(
   GTF,
   output.dir,
   organism,
   assembly,
   db,
   gunzip,
   genome,
   optimize = FALSE,
   uniprot_id = FALSE,
   gene_symbols = FALSE,
   pseudo_5UTRS_if_needed = NULL,
   remove_annotation_outliers = TRUE,
   refseq_genbank_format = c("gtf", "gff3")[1]
)
```

# Arguments

GTF	logical, default: TRUE, download gtf of organism specified in "organism" argument. If FALSE, check if the downloaded file already exist. If you want to use a custom gtf from you hard drive, set GTF = FALSE, and assign: annotation <- getGenomeAndAnnotation(gtf = FALSE) annotation["gtf"] = "path/to/gtf.gtf".  If db is not "ensembl", you will instead get a gff file.
output.dir	directory to save downloaded data
organism	scientific name of organism, Homo sapiens, Danio rerio, Mus musculus, etc. See biomartr:::get.ensembl.info() for full list of supported organisms.

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assembly character, default is assembly = organism, which means getting the first assem-

bly in list, otherwise the name of the assembly wanted, like "GCA\_000005845" will get ecoli substrain k12, which is the most used ones for references. Usually

ignore this for non bacterial species.

db database to use for genome and GTF, default adviced: "ensembl" (remember to

set assembly\_type to "primary\_assembly", else it will contain haplotypes, very large file!). Alternatives: "refseq" (reference assemblies) and "genbank" (all

assemblies)

gunzip logical, default TRUE, uncompress downloaded files that are zipped when down-

loaded, should be TRUE!

genome character path, default NULL. Path to fasta genome, corresponding to the gtf.

must be indexed (.fai file must exist there). If you want to make sure chromosome naming of the GTF matches the genome and correct seqlengths. If value

is NULL or FALSE, it will be ignored.

optimize logical, default FALSE. Create a folder within the output folder (defined by

txdb\_file\_out\_path), that includes optimized objects to speed up loading of annotation regions from up to 15 seconds on human genome down to 0.1 second. ORFik will then load these optimized objects instead. Currently optimizes filterTranscript() function and loadRegion() function for 5' UTRs, 3' UTRs, CDS,

mRNA (all transcript with CDS) and tx (all transcripts).

uniprot\_id logical default FALSE. If TRUE, will download and store all uniprot id for all

transcripts (coding and noncoding)- In a file called: "gene\_symbol\_tx\_table.fst"

in same folder as txdb.

gene\_symbols logical default FALSE. If TRUE, will download and store all gene symbols for

all transcripts (coding and noncoding)- In a file called: "gene\_symbol\_tx\_table.fst" in same folder as txdb. hgcn for human, mouse symbols for mouse and rat, more

to be added.

pseudo\_5UTRS\_if\_needed

integer, default NULL. If defined > 0, will add pseudo 5' UTRs of maximum this length if 'minimum\_5UTR\_percentage" (default 30 mRNAs (coding tran-

scripts) do not have a leader. (NULL and 0 are both the ignore command)

remove\_annotation\_outliers

logical, default TRUE. Only for refseq. shall outlier lines be removed from the input annotation\_file? If yes, then the initial annotation\_file will be overwritten and the removed outlier lines will be stored at tempdir for further exploration.

Among others Aridopsis refseq contains malformed lines, where this is needed

refseq\_genbank\_format

= c("gtf", "gff3")[1] Gtf format files are usually more secure from bugs downstream, so we highly advice to use them. GFF3 files can sometimes include information you might not find in the gtf, so sometimes it makes sense to use it.

Details

Some files that are made after download:

- A fasta index for the genome
- A TxDb to speed up GTF/GFF reading

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- Seperat of merged contaminant files

Files that can be made:

- Gene symbols (hgnc, etc)
- Uniprot ids (For name of protein structures)

If you want custom genome or gtf from you hard drive, assign existing paths like this: annotation <- getGenomeAndAnnotation(GTF = "path/to/gtf.gtf", genome = "path/to/genome.fasta")

#### Value

a named character vector of path to genomes and gtf downloaded, and additional contaminants if used. If merge\_contaminants is TRUE, will not give individual fasta files to contaminants, but only the merged one.

#### References

https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4919035/

#### See Also

```
Other STAR: STAR.align.folder(), STAR.align.single(), STAR.allsteps.multiQC(), STAR.index(), STAR.install(), STAR.multiQC(), STAR.remove.crashed.genome(), install.fastp()
```

```
## Get Saccharomyces cerevisiae genome and gtf (create txdb for R)
#getGenomeAndAnnotation("Saccharomyces cerevisiae", tempdir(), assembly_type = "toplevel")
## Download and add pseudo 5' UTRs
#getGenomeAndAnnotation("Saccharomyces cerevisiae", tempdir(), assembly_type = "toplevel",
# pseudo_5UTRS_if_needed = 100)
## Get Danio rerio genome and gtf (create txdb for R)
#getGenomeAndAnnotation("Danio rerio", tempdir())
output.dir <- "/Bio_data/references/zebrafish"</pre>
## Get Danio rerio and Phix contamints to deplete during alignment
#getGenomeAndAnnotation("Danio rerio", output.dir, phix = TRUE)
## Optimize for ORFik (speed up for large annotations like human or zebrafish)
#getGenomeAndAnnotation("Danio rerio", tempdir(), optimize = TRUE)
# Drosophila melanogaster (toplevel exists only)
#getGenomeAndAnnotation("drosophila melanogaster", output.dir = file.path(config["ref"],
# "Drosophila_melanogaster_BDGP6"), assembly_type = "toplevel")
## How to save malformed refseq gffs:
## First run function and let it crash:
#annotation <- getGenomeAndAnnotation(organism = "Arabidopsis thaliana",
# output.dir = "~/Desktop/test_plant/",
# assembly_type = "primary_assembly", db = "refseq")
## Then apply a fix (example for linux, too long rows):
# fixed_gff <- fix_malformed_gff("~/Desktop/test_plant/Arabidopsis_thaliana_genomic_refseq.gff")
## Then updated arguments:
# annotation <- c(fixed_gff, "~/Desktop/test_plant/Arabidopsis_thaliana_genomic_refseq.fna")
```

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```
# names(annotation) <- c("gtf", "genome")
# Then make the txdb (for faster R use)
# makeTxdbFromGenome(annotation["gtf"], annotation["genome"], organism = "Arabidopsis thaliana")</pre>
```

get\_noncoding\_rna

Download genome (fasta), annotation (GTF) and contaminants

### **Description**

This function automatically downloads (if files not already exists) genomes and contaminants specified for genome alignment. By default, it will use ensembl reference, upon completion, the function will store a file called file.path(output.dir, "outputs.rds") with the output paths of your completed genome/annotation downloads. For most non-model nonvertebrate organisms, you need my fork of biomartr for it to work: remotes::install\_github("Roleren/biomartr) If you misspelled something or crashed, delete wrong files and run again.

Do remake = TRUE, to do it all over again.

### Usage

```
get_noncoding_rna(ncRNA, output.dir, organism, gunzip)
```

# **Arguments**

ncRNA

logical or character, default FALSE (not used, no download), if TRUE or defned path, ncRNA is used as a contaminant reference. If TRUE, will try to find ncRNA sequences from the gtf file, usually represented as lncRNA (long noncoding RNA's). Will let you know if no ncRNA sequences were found in gtf. If not found try character input:

Alternatives; "auto": Will try to find ncRNA file on NONCODE from organism, Homo sapiens -> human etc. "auto" will not work for all, then you must specify the name used by NONCODE, go to the link below and find it. If not "auto" / "" it must be a character vector of species common name (not scientific name) Homo sapiens is human, Rattus norwegicus is rat etc, download ncRNA sequence to filter out with. From NONCODE online server, if you cant find common name see: http://www.noncode.org/download.php/

output.dir directory to save downloaded data

organism scientific name of organism, Homo sapiens, Danio rerio, Mus musculus, etc.

See biomartr:::get.ensembl.info() for full list of supported organisms.

gunzip logical, default TRUE, uncompress downloaded files that are zipped when down-

loaded, should be TRUE!

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#### **Details**

Some files that are made after download:

- A fasta index for the genome
- A TxDb to speed up GTF/GFF reading
- Seperat of merged contaminant files

Files that can be made:

- Gene symbols (hgnc, etc)
- Uniprot ids (For name of protein structures)

If you want custom genome or gtf from you hard drive, assign existing paths like this: annotation <- getGenomeAndAnnotation(GTF = "path/to/gtf.gtf", genome = "path/to/genome.fasta")

#### Value

a named character vector of path to genomes and gtf downloaded, and additional contaminants if used. If merge\_contaminants is TRUE, will not give individual fasta files to contaminants, but only the merged one.

#### References

https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4919035/

#### See Also

```
Other STAR: STAR.align.folder(), STAR.align.single(), STAR.allsteps.multiQC(), STAR.index(), STAR.install(), STAR.multiQC(), STAR.remove.crashed.genome(), install.fastp()
```

```
## Get Saccharomyces cerevisiae genome and gtf (create txdb for R)
#getGenomeAndAnnotation("Saccharomyces cerevisiae", tempdir(), assembly_type = "toplevel")
## Download and add pseudo 5' UTRs
#getGenomeAndAnnotation("Saccharomyces cerevisiae", tempdir(), assembly_type = "toplevel",
# pseudo_5UTRS_if_needed = 100)
## Get Danio rerio genome and gtf (create txdb for R)
#getGenomeAndAnnotation("Danio rerio", tempdir())
output.dir <- "/Bio_data/references/zebrafish"</pre>
## Get Danio rerio and Phix contamints to deplete during alignment
#getGenomeAndAnnotation("Danio rerio", output.dir, phix = TRUE)
## Optimize for ORFik (speed up for large annotations like human or zebrafish)
#getGenomeAndAnnotation("Danio rerio", tempdir(), optimize = TRUE)
# Drosophila melanogaster (toplevel exists only)
#getGenomeAndAnnotation("drosophila melanogaster", output.dir = file.path(config["ref"],
# "Drosophila_melanogaster_BDGP6"), assembly_type = "toplevel")
## How to save malformed refseq gffs:
## First run function and let it crash:
#annotation <- getGenomeAndAnnotation(organism = "Arabidopsis thaliana",</pre>
# output.dir = "~/Desktop/test_plant/",
```

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```
# assembly_type = "primary_assembly", db = "refseq")
## Then apply a fix (example for linux, too long rows):
# fixed_gff <- fix_malformed_gff("~/Desktop/test_plant/Arabidopsis_thaliana_genomic_refseq.gff")
## Then updated arguments:
# annotation <- c(fixed_gff, "~/Desktop/test_plant/Arabidopsis_thaliana_genomic_refseq.fna")
# names(annotation) <- c("gtf", "genome")
# Then make the txdb (for faster R use)
# makeTxdbFromGenome(annotation["gtf"], annotation["genome"], organism = "Arabidopsis thaliana")</pre>
```

get\_phix\_genome

Download genome (fasta), annotation (GTF) and contaminants

### Description

This function automatically downloads (if files not already exists) genomes and contaminants specified for genome alignment. By default, it will use ensembl reference, upon completion, the function will store a file called file.path(output.dir, "outputs.rds") with the output paths of your completed genome/annotation downloads. For most non-model nonvertebrate organisms, you need my fork of biomartr for it to work: remotes::install\_github("Roleren/biomartr) If you misspelled something or crashed, delete wrong files and run again.

Do remake = TRUE, to do it all over again.

#### **Usage**

```
get_phix_genome(phix, output.dir, gunzip)
```

#### **Arguments**

phix

logical, default FALSE, download phiX sequence to filter out Illumina control reads. ORFik defines Phix as a contaminant genome. Phix is used in Illumina sequencers for sequencing quality control. Genome is: refseq, Escherichia phage phiX174. If sequencing facility created fastq files with the command bcl2fastq, then there should be very few phix reads left in the fastq files recieved.

output.dir

directory to save downloaded data

gunzip

logical, default TRUE, uncompress downloaded files that are zipped when down-

loaded, should be TRUE!

#### **Details**

Some files that are made after download:

- A fasta index for the genome
- A TxDb to speed up GTF/GFF reading
- Seperat of merged contaminant files

Files that can be made:

- Gene symbols (hgnc, etc)

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```
- Uniprot ids (For name of protein structures)

If you want custom genome or gtf from you hard drive, assign existing paths like this:

annotation <- getGenomeAndAnnotation(GTF = "path/to/gtf.gtf", genome = "path/to/genome.fasta")
```

#### Value

a named character vector of path to genomes and gtf downloaded, and additional contaminants if used. If merge\_contaminants is TRUE, will not give individual fasta files to contaminants, but only the merged one.

#### References

https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4919035/

#### See Also

```
Other STAR: STAR.align.folder(), STAR.align.single(), STAR.allsteps.multiQC(), STAR.index(), STAR.install(), STAR.multiQC(), STAR.remove.crashed.genome(), install.fastp()
```

```
## Get Saccharomyces cerevisiae genome and gtf (create txdb for R)
#getGenomeAndAnnotation("Saccharomyces cerevisiae", tempdir(), assembly_type = "toplevel")
## Download and add pseudo 5' UTRs
#getGenomeAndAnnotation("Saccharomyces cerevisiae", tempdir(), assembly_type = "toplevel",
# pseudo_5UTRS_if_needed = 100)
## Get Danio rerio genome and gtf (create txdb for R)
#getGenomeAndAnnotation("Danio rerio", tempdir())
output.dir <- "/Bio_data/references/zebrafish"</pre>
## Get Danio rerio and Phix contamints to deplete during alignment
#getGenomeAndAnnotation("Danio rerio", output.dir, phix = TRUE)
## Optimize for ORFik (speed up for large annotations like human or zebrafish)
#getGenomeAndAnnotation("Danio rerio", tempdir(), optimize = TRUE)
# Drosophila melanogaster (toplevel exists only)
#getGenomeAndAnnotation("drosophila melanogaster", output.dir = file.path(config["ref"],
# "Drosophila_melanogaster_BDGP6"), assembly_type = "toplevel")
## How to save malformed refseq gffs:
## First run function and let it crash:
#annotation <- getGenomeAndAnnotation(organism = "Arabidopsis thaliana",
# output.dir = "~/Desktop/test_plant/",
# assembly_type = "primary_assembly", db = "refseq")
## Then apply a fix (example for linux, too long rows):
# fixed_gff <- fix_malformed_gff("~/Desktop/test_plant/Arabidopsis_thaliana_genomic_refseq.gff")
## Then updated arguments:
# annotation <- c(fixed_gff, "~/Desktop/test_plant/Arabidopsis_thaliana_genomic_refseq.fna")
# names(annotation) <- c("gtf", "genome")</pre>
# Then make the txdb (for faster R use)
# makeTxdbFromGenome(annotation["gtf"], annotation["genome"], organism = "Arabidopsis thaliana")
```

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get\_silva\_rRNA

Download Silva SSU & LSU sequences

### **Description**

Version downloaded is 138.1. NR99\_tax (non redundant)

### Usage

```
get_silva_rRNA(output.dir)
```

# Arguments

output.dir

directory to save downloaded data

# **Details**

If it fails from timeout, set higher timeout: options(timeout = 200)

#### Value

filepath to downloaded file

### **Examples**

```
output.dir <- tempdir()
# get_silva_rRNA(output.dir)</pre>
```

groupGRangesBy

Group GRanges

# **Description**

It will group / split the GRanges object by the argument 'other'. For example if you would like to to group GRanges object by gene, set other to gene names.

If 'other' is not specified function will try to use the names of the GRanges object. It will then be similar to 'split(gr, names(gr))'.

# Usage

```
groupGRangesBy(gr, other = NULL)
```

# Arguments

gr a GRanges object

other a vector of unique names to group by (default: NULL)

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#### **Details**

It is important that all intended groups in 'other' are uniquely named, otherwise duplicated group names will be grouped together.

# Value

a GRangesList named after names(GRanges) if other is NULL, else names are from unique(other)

# **Examples**

```
ORFranges <- GRanges(seqnames = Rle(rep("1", 3)),
                      ranges = IRanges(start = c(1, 10, 20),
                                        end = c(5, 15, 25),
                      strand = "+")
ORFranges2 <- GRanges("1",
                       ranges = IRanges(start = c(20, 30, 40),
                                         end = c(25, 35, 45)),
                       strand = "+")
names(ORFranges) = rep("tx1_1", 3)
names(ORFranges2) = rep("tx1_2", 3)
grl <- GRangesList(tx1_1 = ORFranges, tx1_2 = ORFranges2)</pre>
gr <- unlist(grl, use.names = FALSE)</pre>
## now recreate the grl
## group by orf
grltest <- groupGRangesBy(gr) # using the names to group</pre>
identical(grl, grltest) ## they are identical
## group by transcript
names(gr) <- txNames(gr)</pre>
grltest <- groupGRangesBy(gr)</pre>
identical(grl, grltest) ## they are not identical
```

groupings

Get number of ranges per group as an iteration

# Description

Get number of ranges per group as an iteration

#### **Usage**

```
groupings(grl)
```

# **Arguments**

grl

GRangesList

200 hasHits

#### Value

an integer vector

### **Examples**

gSort

Sort a GRangesList, helper.

### **Description**

A helper for [sortPerGroup()]. A faster, more versatile reimplementation of GenomicRanges::sort() Normally not used directly. Groups first each group, then either decreasing or increasing (on starts if byStarts == T, on ends if byStarts == F)

### Usage

```
gSort(grl, decreasing = FALSE, byStarts = TRUE)
```

# Arguments

grl a GRangesList

decreasing should the first in each group have max(start(group)) ->T or min-> default(F)?

byStarts a logical T, should it order by starts or ends F.

#### Value

an equally named GRangesList, where each group is sorted within group.

hasHits

Hits from reads

# Description

Finding GRanges groups that have overlap hits with reads Similar to

# Usage

```
hasHits(grl, reads, keep.names = FALSE, overlaps = NULL)
```

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# **Arguments**

```
grl a GRangesList or GRanges object
reads a GRanges, GAlignment or GAlignmentPairs object
keep.names logical (F), keep names or not
overlaps default NULL, if not null must be countOverlaps(grl, reads), input if you have it already.
```

# Value

```
a list of logicals, T == hit, F == no hit
```

heatMapL

Coverage heatmap of multiple libraries

# **Description**

Coverage heatmap of multiple libraries

# Usage

```
heatMapL(
  region,
  tx,
  df,
  outdir,
  scores = "sum",
  upstream,
  downstream,
  zeroPosition = upstream,
  acceptedLengths = NULL,
  type = "ofst",
  legendPos = "right",
  colors = "default",
  addFracPlot = TRUE,
  location = "TIS",
  shifting = NULL,
  skip.last = FALSE,
  plot.ext = ".pdf",
  plot.together = TRUE,
  title = TRUE,
  scale_x = 5.5,
  scale_y = 15.5,
  gradient.max = "default",
 BPPARAM = BiocParallel::SerialParam()
)
```

202 heatMapL

#### **Arguments**

#' a GRangesList object of region, usually either leaders, cds', 3' utrs or ORFs, region start region, stop regions etc. This is the region that will be mapped in heatmap default NULL, a GRangesList of transcripts or (container region), names of tx tx must contain all grl names. The names of grl can also be the ORFik orf names. that is "txName\_id" df an ORFik experiment outdir a character path to directory to save plot, will be named from ORFik experiment columns character vector, default c("transcriptNormalized", "sum"), either of zsscores core, transcriptNormalized, sum, mean, median, .. see ?coverageScorings for info and more alternatives. upstream 1 or 2 integers, default c(50, 30), how long upstream from 0 should window extend (first index is 5' end extension, second is 3' end extension). If only 1 shifting, only 1 value should be given, if two are given will use first. downstream 1 or 2 integers, default c(29, 69), how long upstream from 0 should window extend (first index is 5' end extension, second is 3' end extension). If only 1 shifting, only 1 value should be given, if two are given will use first. zeroPosition an integer DEFAULT (upstream), what is the center point? Like leaders and cds combination, then 0 is the TIS and -1 is last base in leader. NOTE!: if windows have different widths, this will be ignored. acceptedLengths an integer vector (NULL), the read lengths accepted. Default NULL, means all lengths accepted. character, default: "ofst". Type of library: either "default", usually bam format type (the one you gave to experiment), "pshifted" pshifted reads, "ofst", "bed", "bedo" optimized bed, or "wig" legendPos a character, Default "right". Where should the fill legend be ? ("top", "bottom", "right", "left") colors character vector, default: "default", this gives you: c("white", "yellow2", "yellow3", "lightblue", "blue", "navy"), do "high" for more high contrasts, or specify your own colors. addFracPlot Add margin histogram plot on top of heatmap with fractions per positions location a character, default "start site", will make xlabel of heatmap be Position relative to "start site" or alternative given.

shifting a character, default c("5prime", "3prime"), can also be NULL (no shifting of

reads). If NULL, will use first index of 'upstream' and 'downstream' argument.

skip.last skip top(highest) read length, default FALSE plot.ext a character, default ".pdf", alternative ".png"

title a character, default NULL (no title), what is the top title of plot?

logical (default: FALSE), plot all in 1 plot (if TRUE)

scale\_x numeric, how should the width of the single plots be scaled, bigger the number,

the bigger the plot

plot.together

heatMapRegion 203

scale\_y numeric, how should the height of the plots be scaled, bigger the number, the

bigger the plot

gradient.max numeric or character, default: "default", which is: max(coverage\$score), the

max coverage over all readlengths. If you want all plots to use same reference point for max scaling, then first detect this point, look at max in plot etc, and use

that value, to get all plots to have same max point.

BPPARAM a core param, default: single thread: BiocParallel::SerialParam(). Set to

BiocParallel::bpparam() to use multicore. Be aware, this uses a lot of extra

ram (40GB+) for larger human samples!

#### Value

invisible(NULL), plots are saved

#### See Also

Other heatmaps: coverageHeatMap(), heatMapRegion(), heatMap\_single()

heatMapRegion

Create coverage heatmaps of specified region

# Description

Simplified input space for easier abstraction of coverage heatmaps Pick your transcript region and plot directly Input CAGE file if you use TSS and want improved 5' annotation.

#### Usage

```
heatMapRegion(
  df,
  region = "TIS",
 outdir = "default",
  scores = c("transcriptNormalized", "sum"),
  type = "ofst",
  cage = NULL,
  plot.ext = ".pdf",
  acceptedLengths = 21:75,
  upstream = c(50, 30),
  downstream = c(29, 69),
  shifting = c("5prime", "3prime"),
  longestPerGene = TRUE,
  colors = "default",
  scale_x = 5.5,
  scale_y = 15.5,
 gradient.max = "default",
 BPPARAM = BiocParallel::SerialParam()
)
```

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#### **Arguments**

df an ORFik experiment a character, default "TIS". The centering point for the heatmap (what is position region 0, beween -50 and 50 etc), can be any combination of the set: c("TSS", "TIS", "TTS", "TES"), which are: - Transcription start site (5' end of mrna) - Translation initation site (5' end of CDS) - Translation termination site (5' end of 3' UTRs) - Transcription end site (3' end of 3' UTRs) outdir a character path, default: "default", saves to: file.path(QCfolder(df), "heatmaps/"), a created folder within the ORFik experiment data folder for plots. Change if you want custom location. character vector, default c("transcriptNormalized", "sum"), either of zsscores core, transcriptNormalized, sum, mean, median, .. see ?coverageScorings for character, default: "ofst". Type of library: either "default", usually bam format type (the one you gave to experiment), "pshifted" pshifted reads, "ofst", "bed", "bedo" optimized bed, or "wig" a character path to library file or a GRanges, GAlignments preloaded file of cage CAGE data. Only used if "TSS" is defined as region, to redefine 5' leaders. plot.ext a character, default ".pdf", alternative ".png" acceptedLengths an integer vector (NULL), the read lengths accepted. Default NULL, means all lengths accepted. upstream 1 or 2 integers, default c(50, 30), how long upstream from 0 should window extend (first index is 5' end extension, second is 3' end extension). If only 1 shifting, only 1 value should be given, if two are given will use first. 1 or 2 integers, default c(29, 69), how long upstream from 0 should window downstream extend (first index is 5' end extension, second is 3' end extension). If only 1 shifting, only 1 value should be given, if two are given will use first. shifting a character, default c("5prime", "3prime"), can also be NULL (no shifting of reads). If NULL, will use first index of 'upstream' and 'downstream' argument. logical, default TRUE. Use only longest transcript isoform per gene. This will longestPerGene speed up your computation. colors character vector, default: "default", this gives you: c("white", "yellow2", "yellow3", "lightblue", "blue", "navy"), do "high" for more high contrasts, or specify your own colors.

numeric, how should the width of the single plots be scaled, bigger the number,

scale\_x

the bigger the plot

scale\_y numeric, how should the height of the plots be scaled, bigger the number, the

bigger the plot

gradient.max numeric or character, default: "default", which is: max(coverage\$score), the

max coverage over all readlengths. If you want all plots to use same reference point for max scaling, then first detect this point, look at max in plot etc, and use

that value, to get all plots to have same max point.

heatMap\_single 205

**BPPARAM** 

a core param, default: single thread: BiocParallel::SerialParam(). Set to BiocParallel::bpparam() to use multicore. Be aware, this uses a lot of extra ram (40GB+) for larger human samples!

#### Value

```
invisible(NULL), plots are saved
```

#### See Also

```
Other heatmaps: coverageHeatMap(), heatMapL(), heatMap_single()
```

### **Examples**

```
# Toy example, will not give logical output, but shows how it works
df <- ORFik.template.experiment()[9:10,] # Subset to 2 Ribo-seq libs
#heatMapRegion(df, "TIS", outdir = "default")
#
# Do also TSS, add cage for specific TSS
# heatMapRegion(df, c("TSS", "TIS"), cage = "path/to/cage.bed")
# Do on pshifted reads instead of original files
remove.experiments(df) # Remove loaded experiment first
# heatMapRegion(df, "TIS", type = "pshifted")</pre>
```

heatMap\_single

Coverage heatmap of single libraries

# Description

Coverage heatmap of single libraries

### Usage

```
heatMap_single(
  region,
  tx,
  reads,
  outdir,
  scores = "sum",
  upstream,
  downstream,
  zeroPosition = upstream,
  returnCoverage = FALSE,
  acceptedLengths = NULL,
  legendPos = "right",
  colors = "default",
  addFracPlot = TRUE,
```

206 heatMap\_single

```
location = "start site",
shifting = NULL,
skip.last = FALSE,
title = NULL,
gradient.max = "default"
)
```

#### **Arguments**

region #' a GRangesList object of region, usually either leaders, cds', 3' utrs or ORFs,

start region, stop regions etc. This is the region that will be mapped in heatmap

tx default NULL, a GRangesList of transcripts or (container region), names of tx

must contain all grl names. The names of grl can also be the ORFik orf names.

that is "txName\_id"

reads a GAlignments, GRanges, or precomputed coverage as covRleList (where

names of covRle objects are readlengths) of RiboSeq, RnaSeq etc.

Weigths for scoring is default the 'score' column in 'reads'. Can also be random access paths to bigWig or fstwig file. Do not use random access for more than a

few genes, then loading the entire files is usually better.

outdir a character path to save file as: not just directory, but full name.

scores character vector, default "sum", either of zscore, transcriptNormalized, sum,

mean, median, .. see ?coverageScorings for info and more alternatives.

upstream an integer, relative region to get upstream from.
downstream an integer, relative region to get downstream from

zeroPosition an integer DEFAULT (upstream), what is the center point? Like leaders and cds

combination, then 0 is the TIS and -1 is last base in leader. NOTE!: if windows

have different widths, this will be ignored.

returnCoverage logical, default: FALSE, return coverage, if FALSE returns plot instead.

acceptedLengths

an integer vector (NULL), the read lengths accepted. Default NULL, means all

lengths accepted.

legendPos a character, Default "right". Where should the fill legend be ? ("top", "bottom",

"right", "left")

colors character vector, default: "default", this gives you: c("white", "yellow2", "yel-

low3", "lightblue", "blue", "navy"), do "high" for more high contrasts, or specify

your own colors.

addFracPlot Add margin histogram plot on top of heatmap with fractions per positions

location a character, default "start site", will make xlabel of heatmap be Position relative

to "start site" or alternative given.

shifting a character, default NULL (no shifting), can also be either of c("5prime", "3prime")

skip.last skip top(highest) read length, default FALSE

title a character, default NULL (no title), what is the top title of plot?

gradient.max numeric or character, default: "default", which is: max(coverage\$score), the

max coverage over all readlengths. If you want all plots to use same reference point for max scaling, then first detect this point, look at max in plot etc, and use

that value, to get all plots to have same max point.

import.bedo 207

# Value

```
ggplot2 grob (default), data.table (if returnCoverage is TRUE)
```

# See Also

Other heatmaps: coverageHeatMap(), heatMapL(), heatMapRegion()

import.bedo

Load GRanges object from .bedo

# **Description**

.bedo is .bed ORFik, an optimized bed format for coverage reads with read lengths .bedo is a text based format with columns (6 maximum):

- 1. chromosome
- 2. start
- 3. end
- 4. strand
- 5. ref width (cigar # M's, match/mismatch total)
- 6. duplicates of that read

# Usage

```
import.bedo(path)
```

# **Arguments**

path

a character, location on disc (full path)

# **Details**

Positions are 1-based, not 0-based as .bed. export with export.bedo

# Value

GRanges object

208 import.fstwig

import.bedoc

Load GAlignments object from .bedoc

# Description

A much faster way to store, load and use bam files.

.bedoc is .bed ORFik, an optimized bed format for coverage reads with cigar and replicate number. .bedoc is a text based format with columns (5 maximum):

- 1. chromosome
- 2. cigar: (cigar # M's, match/mismatch total)
- 3. start (left most position)
- 4. strand (+, -, \*)
- 5. score: duplicates of that read

# Usage

```
import.bedoc(path)
```

# **Arguments**

path

a character, location on disc (full path)

#### **Details**

Positions are 1-based, not 0-based as .bed. export with export.bedo

# Value

GAlignments object

import.fstwig

Import region from fastwig

# **Description**

Import region from fastwig

# Usage

```
import.fstwig(gr, dir, id = "", readlengths = "all")
```

import.ofst 209

#### **Arguments**

gr a GRanges object of exons

dir prefix to filepath for file strand and chromosome will be added

id id to column type, not used currently!

readlengths integer / character vector, default "all". Or a subset of readlengths.

#### Value

a data.table with columns specified by readlengths

import.ofst

Load GRanges / GAlignments object from .ofst

# **Description**

A much faster way to store, load and use bam files.

.ofst is ORFik fast serialized object, an optimized format for coverage reads with cigar and replicate number. It uses the fst format as back-end: fst-package.

A .ofst ribo seq file can compress the information in a bam file from 5GB down to a few MB. This new files has super fast reading time, only a few seconds, instead of minutes. It also has random index access possibility of the file.

.ofst is represented as a data.frane format with minimum 4 columns:

- 1. chromosome
- 2. start (left most position)
- 3. strand (+, -, \*)
- 4. width (not added if cigar exists)
- 5. cigar (not needed if width exists): (cigar # M's, match/mismatch total)
- 5. score: duplicates of that read
- 6. size: qwidth according to reference of read

If file is from GAlignmentPairs, it will contain a cigar1, cigar2 instead of cigar and start1 and start2 instead of start

### Usage

```
import.ofst(file, strandMode = 0, seqinfo = NULL)
```

#### **Arguments**

file a path to a .ofst file

strandMode numeric, default 0. Only used for paired end bam files. One of (0: strand

= \*, 1: first read of pair is +, 2: first read of pair is -). See ?strandMode. Note: Sets default to 0 instead of 1, as readGAlignmentPairs uses 1. This is to guarantee hits, but will also make mismatches of overlapping transcripts in

opposite directions.

seqinfo Seqinfo object, defaul NULL (created from ranges). Add to avoid warnings later

on differences in seqinfo.

210 importGtfFromTxdb

#### **Details**

Other columns can be named whatever you want and added to meta columns. Positions are 1-based, not 0-based as .bed. Import with import.ofst

#### Value

a GAlignment, GAlignmentPairs or GRanges object, dependent of if cigar/cigar1 is defined in .ofst file.

# **Examples**

```
## GRanges
gr <- GRanges("1:1-3:-")
tmp <- file.path(tempdir(), "path.ofst")
# export.ofst(gr, file = tmp)
# import.ofst(tmp)
## GAlignment
# Make input data.frame
df <- data.frame(seqnames = "1", cigar = "3M", start = 1L, strand = "+")
ga <- ORFik:::getGAlignments(df)
# export.ofst(ga, file = tmp)
# import.ofst(tmp)</pre>
```

importGtfFromTxdb

Import the GTF / GFF that made the txdb

#### **Description**

Import the GTF / GFF that made the txdb

# Usage

```
importGtfFromTxdb(txdb, stop.error = TRUE)
```

#### **Arguments**

txdb a TxDb, path to txdb / gff or ORFik experiment object stop.error logical TRUE, stop if Txdb does not have a gtf. If FALSE, return NULL.

#### Value

data.frame, the gtf/gff object imported with rtracklayer::import. Or NULL, if stop.error is FALSE, and no GTF file found.

inhibitorNames 211

	inhibitorNames	Get translocation inhibitor name variants	
--	----------------	---	--

# Description

Used to standardize nomeclature for experiments. Example: cycloheximide, lactimidomycin, harringtonine

# Usage

```
inhibitorNames()
```

#### Value

a data.table with 2 columns, the main name, and all name variants of the main name in second column as a list.

# See Also

```
Other experiment_naming: batchNames(), cellLineNames(), cellTypeNames(), conditionNames(), fractionNames(), libNames(), mainNames(), repNames(), stageNames(), tissueNames()
```

initiationScore

Get initiation score for a GRangesList of ORFs

# **Description**

initiationScore tries to check how much each TIS region resembles, the average of the CDS TIS regions.

### Usage

```
initiationScore(grl, cds, tx, reads, pShifted = TRUE, weight = "score")
```

# Arguments

grl	a GRangesList object with ORFs
cds	a GRangesList object with coding sequences
tx	a GRangesList of transcripts covering grl.
reads	ribo seq reads as GAlignments, GRanges or GRangesList object
pShifted	a logical (TRUE), are riboseq reads p-shifted?

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weight

a numeric/integer vector or metacolumn name. (default: 1L, no differential weighting). If weight is name of defined meta column in reads object, it gives the number of times a read was found at that position. GRanges("chr1", 1, "+", score = 5), would mean "score" column tells that this alignment region was found 5 times. if 1L it means each read is weighted equal as 1, this is what among others countOverlaps() presumes, if single number (!= 1), it repeats for all ranges, if vector with length > 1, it must be equal size of the reads object.

#### **Details**

Since this features uses a distance matrix for scoring, values are distributed like this:

As result there is one value per ORF:

0.000: means that ORF had no reads

-1.000: means that ORF is identical to average of CDS

1.000: means that orf is maximum different than average of CDS

If a score column is defined, it will use it as weights, see getWeights

#### Value

an integer vector, 1 score per ORF, with names of grl

#### References

doi: 10.1186/s12915-017-0416-0

#### See Also

```
Other features: computeFeatures(), computeFeaturesCage(), countOverlapsW(), disengagementScore(), distToCds(), distToTSS(), entropy(), floss(), fpkm(), fpkm_calc(), fractionLength(), insideOutsideORF(), isInFrame(), isOverlapping(), kozakSequenceScore(), orfScore(), rankOrder(), ribosomeReleaseScore(), ribosomeStallingScore(), startRegion(), startRegionCoverage(), stopRegion(), subsetCoverage(), translationalEff()
```

insideOutsideORF 213

insideOutsideORF

Inside/Outside score (IO)

# Description

Inside/Outside score is defined as

```
(reads over ORF)/(reads outside ORF and within transcript)
```

A pseudo-count of one is added to both the ORF and outside sums.

### Usage

```
insideOutsideORF(
  grl,
  RFP,
  GtfOrTx,
  ds = NULL,
  RFP.sorted = FALSE,
  weight = 1L,
  overlapGrl = NULL
)
```

# Arguments

grl	a GRangesList object with usually either leaders, cds', 3' utrs or ORFs.
RFP	RiboSeq reads as GAlignments, GRanges or GRangesList object
Gtf0rTx	If it is TxDb object transcripts will be extracted using exonsBy(Gtf, by = "tx", use.names = TRUE). Else it must be GRangesList
ds	numeric vector (NULL), disengagement score. If you have already calculated disengagementScore, input here to save time.
RFP.sorted	logical (FALSE), an optimizer, have you ran this line: RFP <- sort(RFP[countOverlaps(RFP, tx, type = "within") > 0]) Normally not touched, for internal optimization purposes.
weight	a numeric/integer vector or metacolumn name. (default: 1L, no differential weighting). If weight is name of defined meta column in reads object, it gives the number of times a read was found at that position. GRanges("chr1", 1, "+",

score = 5), would mean "score" column tells that this alignment region was

214 insideOutsideORF

found 5 times. if 1L it means each read is weighted equal as 1, this is what among others countOverlaps() presumes, if single number (!= 1), it repeats for all ranges, if vector with length > 1, it must be equal size of the reads object.

overlapGrl

an integer, (default: NULL), if defined must be countOverlaps(grl, RFP), added for speed if you already have it

#### Value

a named vector of numeric values of scores

#### References

doi: 10.1242/dev.098345

#### See Also

```
Other features: computeFeatures(), computeFeaturesCage(), countOverlapsW(), disengagementScore(), distToCds(), distToTSS(), entropy(), floss(), fpkm(), fpkm_calc(), fractionLength(), initiationScore(), isInFrame(), isOverlapping(), kozakSequenceScore(), orfScore(), rankOrder(), ribosomeReleaseScore(), ribosomeStallingScore(), startRegion(), startRegionCoverage(), stopRegion(), subsetCoverage(), translationalEff()
```

```
# Check inside outside score of a ORF within a transcript
ORF <- GRanges("1",
               ranges = IRanges(start = c(20, 30, 40),
                                  end = c(25, 35, 45)),
               strand = "+")
grl <- GRangesList(tx1_1 = ORF)
tx1 <- GRanges(seqnames = "1",
               ranges = IRanges(start = c(1, 10, 20, 30, 40, 50),
                                 end = c(5, 15, 25, 35, 45, 200)),
               strand = "+")
tx \leftarrow GRangesList(tx1 = tx1)
RFP <- GRanges(seqnames = "1",
                  ranges = IRanges(start = c(1, 4, 30, 60, 80, 90),
                                    end = c(30, 33, 63, 90, 110, 120)),
                  strand = "+")
insideOutsideORF(grl, RFP, tx)
```

install.fastp 215

install.fastp

Download and prepare fastp trimmer

# **Description**

```
On Linux, will not run "make", only use precompiled fastp file.
On Mac OS it will use precompiled binaries.
For windows must be installed through WSL (Windows Subsystem Linux)
```

# Usage

```
install.fastp(folder = "~/bin")
```

# **Arguments**

folder

path to folder for download, file will be named "fastp", this should be most recent version. On mac it will search for a folder called fastp-master inside folder given. Since there is no precompiled version of fastp for Mac OS.

#### Value

path to runnable fastp

### References

https://www.ncbi.nlm.nih.gov/pmc/articles/PMC6129281/

#### See Also

```
Other STAR: STAR.align.folder(), STAR.align.single(), STAR.allsteps.multiQC(), STAR.index(), STAR.install(), STAR.multiQC(), STAR.remove.crashed.genome(), getGenomeAndAnnotation()
```

```
## With default folder:
#install.fastp()

## Or set manual folder:
folder <- "~/I/WANT/IT/HERE/"
#install.fastp(folder)</pre>
```

216 install.sratoolkit

install.sratoolkit Download sra toolkit

# Description

Currently supported for Linux (64 bit centos and ubunutu is tested to work) and Mac-OS(64 bit). If other linux distro, centos binaries will be used.

# Usage

```
install.sratoolkit(folder = "~/bin", version = "2.11.3")
```

# Arguments

folder default folder, "~/bin" version a string, default "2.11.3"

#### Value

path to fastq-dump in sratoolkit

#### References

https://ncbi.github.io/sra-tools/fastq-dump.html

### See Also

```
Other sra: browseSRA(), download.SRA(), download.SRA.metadata(), download.ebi(), get_bioproject_candidates rename.SRA.files()
```

```
# install.sratoolkit()
## Custom folder and version (not adviced)
folder <- "/I/WANT/IT/HERE/"
# install.sratoolkit(folder, version = "2.10.9")</pre>
```

is.grl 217

is.grl

Helper function to check for GRangesList

# Description

Helper function to check for GRangesList

# Usage

```
is.grl(class)
```

## **Arguments**

class

the class you want to check if is GRL, either a character from class or the object

### Value

a boolean

### See Also

```
Other validity: checkRFP(), checkRNA(), is.ORF(), is.gr_or_grl(), is.range(), validGRL(), validSeqlevels()
```

is.gr\_or\_grl

Helper function to check for GRangesList or GRanges class

# **Description**

Helper function to check for GRangesList or GRanges class

### Usage

```
is.gr_or_grl(class)
```

## **Arguments**

class

the class you want to check if is GRL or GR, either a character from class or the object itself.

## Value

a boolean

### See Also

```
Other validity: checkRFP(), checkRNA(), is.ORF(), is.grl(), is.range(), validGRL(), validSeqlevels()
```

is.range

is.ORF

Check if all requirements for an ORFik ORF is accepted.

# Description

Check if all requirements for an ORFik ORF is accepted.

## Usage

```
is.ORF(grl)
```

## **Arguments**

grl

a GRangesList or GRanges to check

#### Value

```
a logical (TRUE/FALSE)
```

### See Also

```
Other validity: checkRFP(), checkRNA(), is.gr_or_grl(), is.grl(), is.range(), validGRL(), validSeqlevels()
```

is.range

Helper function to check for ranged object

## **Description**

Helper function to check for ranged object

## Usage

```
is.range(x)
```

### **Arguments**

Χ

the object to check is a ranged object. Either GRangesList, GRanges, IRangesList, IRanges.

# Value

a boolean

### See Also

```
Other validity: checkRFP(), checkRNA(), is.ORF(), is.gr_or_grl(), is.grl(), validGRL(), validSeqlevels()
```

isInFrame 219

isInFrame

Find frame for each orf relative to cds

### **Description**

Input of this function, is the output of the function [distToCds()], or any other relative ORF frame.

### Usage

```
isInFrame(dists)
```

## **Arguments**

dists

a vector of integer distances between ORF and cds. 0 distance means equal frame

#### **Details**

```
possible outputs: 0: orf is in frame with cds 1: 1 shifted from cds 2: 2 shifted from cds
```

#### Value

a logical vector

#### References

```
doi: 10.1074/jbc.R116.733899
```

#### See Also

```
Other features: computeFeatures(), computeFeaturesCage(), countOverlapsW(), disengagementScore(), distToCds(), distToTSS(), entropy(), floss(), fpkm(), fpkm_calc(), fractionLength(), initiationScore(), insideOutsideORF(), isOverlapping(), kozakSequenceScore(), orfScore(), rankOrder(), ribosomeReleaseScore(), ribosomeStallingScore(), startRegion(), startRegionCoverage(), stopRegion(), subsetCoverage(), translationalEff()
```

```
# simple example
isInFrame(c(3,6,8,11,15))

# GRangesList example
grl <- GRangesList(tx1_1 = GRanges("1", IRanges(1,10), "+"))
fiveUTRs <- GRangesList(tx1 = GRanges("1", IRanges(1,20), "+"))
dist <- distToCds(grl, fiveUTRs)
isInFrame <- isInFrame(dist)</pre>
```

220 isOverlapping

isOverlapping

Find frame for each orf relative to cds

## **Description**

Input of this function, is the output of the function [distToCds()]

## Usage

```
isOverlapping(dists)
```

## **Arguments**

dists

a vector of distances between ORF and cds

#### Value

a logical vector

### References

```
doi: 10.1074/jbc.R116.733899
```

### See Also

```
Other features: computeFeatures(), computeFeaturesCage(), countOverlapsW(), disengagementScore(), distToCds(), distToTSS(), entropy(), floss(), fpkm(), fpkm_calc(), fractionLength(), initiationScore(), insideOutsideORF(), isInFrame(), kozakSequenceScore(), orfScore(), rankOrder(), ribosomeReleaseScore(), ribosomeStallingScore(), startRegion(), startRegionCoverage(), stopRegion(), subsetCoverage(), translationalEff()
```

```
# simple example
isOverlapping(c(-3,-6,8,11,15))
# GRangesList example
grl <- GRangesList(tx1_1 = GRanges("1", IRanges(1,10), "+"))
fiveUTRs <- GRangesList(tx1 = GRanges("1", IRanges(1,20), "+"))
dist <- distToCds(grl, fiveUTRs)
isOverlapping <- isOverlapping(dist)</pre>
```

isPeriodic 221

isPeriodic	Find if there is a periodicity of 3 in the vector	

## **Description**

It uses Fourier transform + periodogram for finding periodic vectors on the transcript normalized counts over all CDS regions from position 0 (TIS) to 149 (or other max position if increased by the user.

Checks if there is a periodicity and if the periodicity is 3, more precisely between 2.9 and 3.1.

# Usage

```
isPeriodic(x, info = NULL, verbose = FALSE, strict.fft = TRUE)
```

## Arguments

info specify read length if wanted for verbose output.

verbose logical, default FALSE. Report details of analysis/periodogram. Good if you are

not sure if the analysis was correct.

strict.fft logical, TRUE. Use a FFT without noise filter. This means keep only reads

lengths that are "periodic for the human eye". If you want more coverage, set to FALSE, to also get read lengths that are "messy", but the noise filter detects the periodicity of 3. This should only be done when you do not need high quality periodic reads! Example would be differential translation analysis by counts

over each ORF.

#### **Details**

Input data:

Transcript normalized means per CDS TIS region, count reads per position, divide that number per position by the total of that transcript, then sum up these numbers per position for all transcripts. Detection method:

The maximum dominant Fourier frequencies is found by finding which period has the highest spectrum density (using a 10

#### Value

a logical, if it is periodic.

222 kozakHeatmap

ko:	zakl	Hea	4 t i	man

Make sequence region heatmap relative to scoring

## **Description**

Given sequences, DNA or RNA. And some score, ribo-seq fpkm, TE etc. Create a heatmap divided per letter in seqs, by how strong the score is.

### Usage

```
kozakHeatmap(
   seqs,
   rate,
   start = 1,
   stop = max(nchar(seqs)),
   center = ceiling((stop - start + 1)/2),
   min.observations = ">q1",
   skip.startCodon = FALSE,
   xlab = "TIS",
   type = "ribo-seq"
)
```

# **Arguments**

seqs the sequences (character vector, DNAStringSet)
rate a scoring vector (equal size to seqs)

start position in seqs to start at (first is 1), default 1.

stop position in seqs to stop at (first is 1), default max(nchar(seqs)), that is the longest

sequence length

center position in seqs to center at (first is 1), center will be +1 in heatmap

min.observations

How many observations per position per letter to accept? numeric or quantile, default (">q1", bigger than quartile 1 (25 percentile)). You can do (10), to get

all with more than 10 observations.

skip.startCodon

startCodon is defined as after centering (position 1, 2 and 3). Should they be skipped? default (FALSE). Not relevant if you are not doing Translation initiation.

tion sites (TIS).

xlab Region you are checking, default (TIS)

type What type is the rate scoring? default (ribo-seq)

#### Details

It will create blocks around the highest rate per position

kozakSequenceScore 223

#### Value

a ggplot of the heatmap

## **Examples**

```
## Not run:
if (requireNamespace("BSgenome.Hsapiens.UCSC.hg19")) {
         txdbFile <- system.file("extdata", "hg19_knownGene_sample.sqlite",</pre>
                                                                                                                    package = "GenomicFeatures")
        #Extract sequences of Coding sequences.
        cds <- loadRegion(txdbFile, "cds")</pre>
        tx <- loadRegion(txdbFile, "mrna")</pre>
        # Get region to check
        kozak Regions <- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens. UCSC.hg19:: Hsapiens +- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \ tx, \ BSgenome. Hsapiens +- \ start Region String (cds, \ tx, \
                                                                                                                                                                  , upstream = 4, 5)
        # Some toy ribo-seq fpkm scores on cds
        set.seed(3)
         fpkm <- sample(1:115, length(cds), replace = TRUE)</pre>
        kozakHeatmap(kozakRegions, fpkm, 1, 9, skip.startCodon = F)
}
## End(Not run)
```

kozakSequenceScore

Make a score for each ORFs start region by proximity to Kozak

## **Description**

The closer the sequence is to the Kozak sequence the higher the score, based on the experimental pwms from article referenced. Minimum score is 0 (worst correlation), max is 1 (the best base per column was chosen).

# Usage

```
kozakSequenceScore(grl, tx, faFile, species = "human", include.N = FALSE)
```

## **Arguments**

```
grl a GRangesList grouped by ORF

tx a GRangesList, the reference area for ORFs, each ORF must have a coresponding tx.

faFile FaFile, BSgenome, fasta/index file path or an ORFik experiment. This file is usually used to find the transcript sequences from some GRangesList.
```

224 kozakSequenceScore

("human"), which species to use, currently supports human (Homo sapiens), zebrafish (Danio rerio) and mouse (Mus musculus). Both scientific or common name for these species will work. You can also specify a pfm for your own species. Syntax of pfm is an rectangular integer matrix, where all columns must sum to the same value, normally 100. See example for more information. Rows are in order: c("A", "C", "G", "T")

include.N logical (F), if TRUE, allow N bases to be counted as hits, score will be average of the other bases. If True, N bases will be added to pfm, automaticly, so dont include them if you make your own pfm.

#### **Details**

Ranges that does not have minimum 15 length (the kozak requirement as a sliding window of size 15 around grl start), will be set to score 0. Since they should not have the posibility to make an efficient ribosome binding.

### Value

```
a numeric vector with values between 0 and 1 an integer vector, one score per orf
```

#### References

doi: https://doi.org/10.1371/journal.pone.0108475

#### See Also

```
Other features: computeFeatures(), computeFeaturesCage(), countOverlapsW(), disengagementScore(), distToCds(), distToTSS(), entropy(), floss(), fpkm(), fpkm_calc(), fractionLength(), initiationScore(), insideOutsideORF(), isInFrame(), isOverlapping(), orfScore(), rankOrder(), ribosomeReleaseScore(), ribosomeStallingScore(), startRegion(), startRegionCoverage(), stopRegion(), subsetCoverage(), translationalEff()
```

kozak\_IR\_ranking 225

```
kozakSequenceScore(ORFs, tx, faFile)
# For more details see vignettes.
```

kozak\_IR\_ranking

Rank kozak initiation sequences

# Description

Defined as region (-4, -1) relative to TIS

## Usage

```
kozak_IR_ranking(cds_k, mrna, dt.ir, faFile, group.min = 10, species = "human")
```

# Arguments

cds_k	cds ranges (GRangesList)
mrna	mrna ranges (GRangesList)
dt.ir	data.table with a column called IR, initiation rate
faFile	FaFile, BSgenome, fasta/index file path or an ORFik experiment. This file is usually used to find the transcript sequences from some GRangesList.
group.min	numeric, default 10. Minimum transcripts per initation group to be included
species	("human"), which species to use, currently supports human (Homo sapiens), zebrafish (Danio rerio) and mouse (Mus musculus). Both scientific or common name for these species will work. You can also specify a pfm for your own species. Syntax of pfm is an rectangular integer matrix, where all columns must sum to the same value, normally 100. See example for more information. Rows are in order: c("A", "C", "G", "T")

# Value

a ggplot grid object

lastExonEndPerGroup Get last end per granges group

# Description

Get last end per granges group

```
lastExonEndPerGroup(grl, keep.names = TRUE)
```

226 lastExonPerGroup

### **Arguments**

```
grl a GRangesList
keep.names a boolean, keep names or not, default: (TRUE)
```

### Value

```
a Rle(keep.names = T), or integer vector(F)
```

### **Examples**

lastExonPerGroup

Get last exon per GRangesList group

## **Description**

grl must be sorted, call ORFik:::sortPerGroup if needed

### Usage

```
lastExonPerGroup(grl)
```

## **Arguments**

grl

 $a \; \mathsf{GRangesList}$ 

#### Value

a GRangesList of the last exon per group

```
 \begin{split} \text{gr\_plus} <& - \text{GRanges}(\text{seqnames} = \text{c("chr1", "chr1")}, \\ & \text{ranges} = \text{IRanges}(\text{c(7, 14), width} = 3), \\ & \text{strand} = \text{c("+", "+")}) \\ \text{gr\_minus} <& - \text{GRanges}(\text{seqnames} = \text{c("chr2", "chr2")}, \\ & \text{ranges} = \text{IRanges}(\text{c(4, 1), c(9, 3)}), \\ & \text{strand} = \text{c("-", "-")}) \\ \text{grl} <& - \text{GRangesList}(\text{tx1} = \text{gr\_plus}, \text{tx2} = \text{gr\_minus}) \\ \text{lastExonPerGroup}(\text{grl}) \end{aligned}
```

lastExonStartPerGroup 227

lastExonStartPerGroup Get last start per granges group

# Description

Get last start per granges group

# Usage

```
lastExonStartPerGroup(grl, keep.names = TRUE)
```

## **Arguments**

```
grl a GRangesList
keep.names a boolean, keep names or not, default: (TRUE)
```

#### Value

```
a Rle(keep.names = T), or integer vector(F)
```

## **Examples**

length,covRle-method length covRle

# Description

Number of chromosomes

## Usage

```
## S4 method for signature 'covRle'
length(x)
```

### **Arguments**

x a covRle object

## Value

an integer, number of chromosomes in covRle object

```
length,covRleList-method
```

length covRleList

## Description

Number of covRle objects

# Usage

```
## S4 method for signature 'covRleList'
length(x)
```

# Arguments

Χ

a covRleList object

## Value

an integer, number of covRle objects

lengths,covRle-method lengths covRle

# Description

Lengths of each chromosome

## Usage

```
## S4 method for signature 'covRle'
lengths(x)
```

# Arguments

Χ

a covRle object

## Value

a named integer vector of chromosome lengths

lengths,covRleList-method

lengths covRleList

# Description

Lengths of each chromosome

# Usage

```
## S4 method for signature 'covRleList'
lengths(x)
```

# Arguments

Х

a covRle object

## Value

a named integer vector of chromosome lengths

libFolder

Get ORFik experiment library folder

# Description

Get ORFik experiment library folder

## Usage

```
libFolder(x, mode = "first")
```

# Arguments

x an ORFik experiment

mode character, default "first". Alternatives: "unique", "all".

## Value

a character path

230 libNames

```
libFolder, experiment-method
```

Get ORFik experiment library folder

## **Description**

Get ORFik experiment library folder

## Usage

```
## S4 method for signature 'experiment'
libFolder(x, mode = "first")
```

# Arguments

x an ORFik experiment

mode character, default "first". Alternatives: "unique", "all".

#### Value

a character path

libNames

Get library name variants

## **Description**

Used to standardize nomeclature for experiments.

Example: RFP is main naming, but a variant is ribo-seq ribo-seq will then be renamed to RFP

### Usage

```
libNames()
```

#### Value

a data.table with 2 columns, the main name, and all name variants of the main name in second column as a list.

### See Also

```
Other experiment_naming: batchNames(), cellLineNames(), cellTypeNames(), conditionNames(), fractionNames(), inhibitorNames(), mainNames(), repNames(), stageNames(), tissueNames()
```

libraryTypes 231

libraryTypes

Which type of library type in experiment?

## **Description**

Which type of library type in experiment?

## Usage

```
libraryTypes(df, uniqueTypes = TRUE)
```

### **Arguments**

```
df an ORFik experiment uniqueTypes logical, default TRUE. Only return unique lib types.
```

#### Value

library types (character vector)

### See Also

```
Other ORFik_experiment: ORFik.template.experiment(), ORFik.template.experiment.zf(), bamVarName(), create.experiment(), experiment-class, filepath(), organism, experiment-method, outputLibs(), read.experiment(), save.experiment(), validateExperiments()
```

### **Examples**

```
df <- ORFik.template.experiment()
libraryTypes(df)
libraryTypes(df, uniqueTypes = FALSE)</pre>
```

list.experiments

List current experiment available

### **Description**

Will only search .csv extension, also exclude any experiment with the word template.

```
list.experiments(
  dir = ORFik::config()["exp"],
  pattern = "*",
  libtypeExclusive = NULL,
  validate = TRUE,
  BPPARAM = bpparam()
)
```

232 list.genomes

## **Arguments**

dir directory for ORFik experiments: default: ORFik::config()["exp"], which by

default is: "~/Bio\_data/ORFik\_experiments/"

pattern allowed patterns in experiment file name: default ("\*", all experiments)

libtypeExclusive

search for experiments with exclusivly this libtype, default (NULL, all)

validate logical, default TRUE. Abort if any library files does not exist. Do not set this

to FALSE, unless you know what you are doing!

BPPARAM how many cores/threads to use? default: bpparam()

#### Value

a data.table, 1 row per experiment with columns:

- experiment (name),

- organism
- author
- libtypes
- number of samples

## Examples

```
## Make your experiments
df <- ORFik.template.experiment(TRUE)
df2 <- df[1:6,] # Only first 2 libs
## Save them
# save.experiment(df, "~/Bio_data/ORFik_experiments/exp1.csv")
# save.experiment(df2, "~/Bio_data/ORFik_experiments/exp1_subset.csv")
## List all experiment you have:
## Path above is default path, so no dir argument needed
#list.experiments()
#list.experiments(pattern = "subset")
## For non default directory experiments
#list.experiments(dir = "MY/CUSTOM/PATH)</pre>
```

list.genomes

List genomes created with ORFik

## Description

Given the reference.folder, list all valid references. An ORFik genome is defined as a folder with a file called output.rds that is a named R vector with names gtf and genome, where the values are character paths to those files inside that folder. This makes sure that this reference was made by ORFik and not some other program.

```
list.genomes(reference.folder = ORFik::config()["ref"])
```

loadRegion 233

## **Arguments**

```
reference.folder character path, default: ORFik::config()["ref"].
```

### Value

- a data.table with 5 columns:
- character (name of folder)
- logical (does it have a gtf)
- logical (does it have a fasta genome)
- logical (does it have a STAR index)
- logical (only displayed if some are TRUE, does it have protein structure predictions of ORFs from alphafold etc, in folder called 'protein\_structure\_predictions')
- logical (only displayed if some are TRUE, does it have gene symbol fst file from bioMart etc, in file called 'gene\_symbol\_tx\_table.fst')

## **Examples**

```
## Run with default config path
#list.genomes()
## Run with custom config path
list.genomes(tempdir())
## Get the path to fasta genome of first organism in list
#readRDS(file.path(config()["ref"], list.genomes()$name, "outputs.rds")[1])["genome"]
```

 ${\tt loadRegion}$ 

Load transcript region

# Description

Usefull to simplify loading of standard regions, like cds' and leaders. Adds another safety in that seqlevels will be set

```
loadRegion(
  txdb,
  part = "tx",
  names.keep = NULL,
  by = "tx",
  skip.optimized = FALSE
)
```

234 loadRegions

### **Arguments**

txdb a TxDb object, ORFik experiment object or a path to one of: (.gtf ,.gff, .gff2,

.gff2, .db or .sqlite), Only in the loadRegion function: if it is a GRangesList, it

will return it self.

part a character, one of: tx, ncRNA, mrna, leader, cds, trailer, intron, NOTE: dif-

ference between tx and mrna is that tx are all transcripts, while mrna are all

transcripts with a cds, respectivly ncRNA are all tx without a cds.

names.keep a character vector of subset of names to keep. Example: loadRegions(txdb,

names = "ENST1000005"), will return only that transcript. Remember if you

set by to "gene", then this list must be with gene names.

by a character, default "tx" Either "tx" or "gene". What names to output region by,

the transcript name "tx" or gene names "gene". NOTE: this is not the same as cdsBy(txdb, by = "gene"), cdsBy would then only give 1 cds per Gene, loadRe-

gion gives all isoforms, but with gene names.

skip.optimized logical, default FALSE. If TRUE, will not search for optimized rds files to load

created from ORFik::makeTxdbFromGenome(..., optimize = TRUE). The opti-

mized files are ~ 100x faster to load for human genome.

#### **Details**

Load as GRangesList if input is not already GRangesList.

#### Value

a GRangesList of region

### **Examples**

loadRegions

Get all regions of transcripts specified to environment

#### Description

By default loads all parts to .GlobalEnv (global environemnt) Useful to not spend time on finding the functions to load regions.

loadRegions 235

### Usage

```
loadRegions(
  txdb,
  parts = c("mrna", "leaders", "cds", "trailers"),
  extension = "",
  names.keep = NULL,
  by = "tx",
  skip.optimized = FALSE,
  envir = .GlobalEnv
)
```

#### **Arguments**

txdb

experiment the transcript parts you want, default: c("mrna", "leaders", "cds", "trailers"). parts See ?loadRegion for more info on this argument. What to add on the name after leader, like: B -> leadersB extension a character vector of subset of names to keep. Example: loadRegions(txdb, names.keep names = "ENST1000005"), will return only that transcript. Remember if you set by to "gene", then this list must be with gene names. a character, default "tx" Either "tx" or "gene". What names to output region by, by the transcript name "tx" or gene names "gene". NOTE: this is not the same as cdsBy(txdb, by = "gene"), cdsBy would then only give 1 cds per Gene, loadRegion gives all isoforms, but with gene names. skip.optimized logical, default FALSE. If TRUE, will not search for optimized rds files to load

mized files are ~ 100x faster to load for human genome.

Which environment to save to, default: .GlobalEnv

a TxDb file, a path to one of: (.gtf,.gff, .gff2, .gff2, .db or .sqlite) or an ORFik

created from ORFik::makeTxdbFromGenome(..., optimize = TRUE). The opti-

#### Value

envir

invisible(NULL) (regions saved in envir)

236 loadTxdb

loadTranscriptType Load transcripts of given biotype

### **Description**

Like rRNA, snoRNA etc. NOTE: Only works on gtf/gff, not .db object for now. Also note that these anotations are not perfect, some rRNA annotations only contain 5S rRNA etc. If your gtf does not contain evertyhing you need, use a resource like repeatmasker and download a gtf: https://genome.ucsc.edu/cgi-bin/hgTables

### Usage

```
loadTranscriptType(object, part = "rRNA", tx = NULL)
```

### **Arguments**

object a TxDb, ORFik experiment or path to gtf/gff,

part a character, default rRNA. Can also be: snoRNA, tRNA etc. As long as that

biotype is defined in the gtf.

tx a GRangesList of transcripts (Optional, default NULL, all transcript of that

type), else it must be names a list to subset on.

### Value

a GRangesList of transcript of that type

#### References

```
doi: 10.1002/0471250953.bi0410s25
```

## **Examples**

```
gtf <- "path/to.gtf"
#loadTranscriptType(gtf, part = "rRNA")
#loadTranscriptType(gtf, part = "miRNA")</pre>
```

loadTxdb

General loader for txdb

## **Description**

Useful to allow fast TxDb loader like .db

```
loadTxdb(txdb, chrStyle = NULL, organism = NA, chrominfo = NULL)
```

longestORFs 237

## **Arguments**

txdb a TxDb file, a path to one of: (.gtf ,.gff2, .gff2, .db or .sqlite) or an ORFik

experiment

chrStyle a GRanges object, TxDb, FaFile, , a seqlevelsStyle or Seqinfo. (Default:

NULL) to get seqlevelsStyle from. In addition if it is a Seqinfo object, seqinfo will be updated. Example of seqlevelsStyle update: Is chromosome 1 called chr1 or 1, is mitocondrial chromosome called MT or chrM etc. Will use 1st seqlevel-style if more are present. Like: c("NCBI", "UCSC") -> pick "NCBI"

organism character, default NA. Scientific name of organism. Only used if input is path to

gff.

chrominfo Seqinfo object, default NULL. Only used if input is path to gff.

#### Value

a TxDb object

### **Examples**

longestORFs

Get longest ORF per stop site

### **Description**

Rule: if seqname, strand and stop site is equal, take longest one. Else keep. If IRangesList or IRanges, seqnames are groups, if GRanges or GRangesList seqnames are the seqlevels (e.g. chromosomes/transcripts)

### Usage

```
longestORFs(grl)
```

## **Arguments**

grl a GRangesList/IRangesList, GRanges/IRanges of ORFs

## Value

```
a GRangesList/IRangesList, GRanges/IRanges (same as input)
```

238 mainNames

### See Also

```
Other ORFHelpers: defineTrailer(), mapToGRanges(), orfID(), startCodons(), startSites(), stopCodons(), stopSites(), txNames(), uniqueGroups(), uniqueOrder()
```

## **Examples**

```
ORF1 = GRanges("1", IRanges(10,21), "+")
ORF2 = GRanges("1", IRanges(1,21), "+") # <- longest
grl <- GRangesList(ORF1 = ORF1, ORF2 = ORF2)
longestORFs(grl) # get only longest</pre>
```

mainNames

Get main name from variant name

## **Description**

Used to standardize nomeclature for experiments.

Example: RFP is main naming, but a variant is ribo-seq ribo-seq will then be renamed to RFP

### Usage

```
mainNames(names, dt)
```

### **Arguments**

names a character vector of names that must exist in dt\$allNames

dt a data.table with 2 columns (mainName, allNames)

## Value

a data.table with 2 columns, the main name, and all name variants of the main name in second column as a list.

### See Also

```
Other experiment_naming: batchNames(), cellLineNames(), cellTypeNames(), conditionNames(), fractionNames(), inhibitorNames(), libNames(), repNames(), stageNames(), tissueNames()
```

makeExonRanks 239

s Make grouping by exons ranks
0 1 0 2

## **Description**

There are two ways to make vector of exon ranking: 1. Iterate per exon in ORF, byTranscript = FALSE 2. Iterate per ORF in transcript, byTranscript = TRUE.

## Usage

```
makeExonRanks(grl, byTranscript = FALSE)
```

## **Arguments**

grl a GRangesList

byTranscript logical (default: FALSE), groups orfs by transcript name or ORF name, if ORfs

are by transcript, check duplicates.

#### **Details**

Either by transcript or by original groupings. Must be ordered, so that same transcripts are ordered together.

### Value

an integer vector of indices for exon ranks

	makeORFNames	Make ORF names per orf	
--	--------------	------------------------	--

# **Description**

grl must be grouped by transcript If a list of orfs are grouped by transcripts, but does not have ORF names, then create them and return the new GRangesList

## Usage

```
makeORFNames(grl, groupByTx = TRUE)
```

## **Arguments**

grl a GRangesList

groupByTx logical (T), should output GRangesList be grouped by transcripts (T) or by

ORFs (F)?

### Value

(GRangesList) with ORF names, grouped by transcripts, sorted.

### **Examples**

```
 \begin{split} \text{gr\_plus} &<\text{- GRanges}(\text{seqnames} = \text{c("chr1", "chr1")}, \\ &\quad \text{ranges} = \text{IRanges}(\text{c(7, 14), width} = 3), \\ &\quad \text{strand} = \text{c("+", "+")}) \\ \text{gr\_minus} &<\text{- GRanges}(\text{seqnames} = \text{c("chr2", "chr2")}, \\ &\quad \text{ranges} = \text{IRanges}(\text{c(4, 1), c(9, 3)}), \\ &\quad \text{strand} = \text{c("-", "-")}) \\ \text{grl} &<\text{- GRangesList(tx1 = gr\_plus, tx2 = gr\_minus)} \\ \text{makeORFNames}(\text{grl}) \end{aligned}
```

makeSummarizedExperimentFromBam

Make a count matrix from a library or experiment

### **Description**

Make a summerizedExperiment / matrix object from bam files or other library formats sepcified by lib.type argument. Works like HTSeq, to give you count tables per library.

### Usage

```
makeSummarizedExperimentFromBam(
    df,
    saveName = NULL,
    longestPerGene = FALSE,
    geneOrTxNames = "tx",
    region = "mrna",
    type = "count",
    lib.type = "ofst",
    weight = "score",
    forceRemake = FALSE,
    force = TRUE,
    library.names = bamVarName(df),
    BPPARAM = BiocParallel::SerialParam()
)
```

## **Arguments**

df an ORFik experiment

saveName a character (default NULL), if set save experiment to path given. Always saved as .rds., it is optional to add .rds, it will be added for you if not present. Also

used to load existing file with that name.

longestPerGene a logical (default FALSE), if FALSE all transcript isoforms per gene. Ignored if "region" is not a character of either: "mRNA", "tx", "cds", "leaders" or "trailers". a character vector (default "tx"), should row names keep trancript names ("tx") geneOrTxNames or change to gene names ("gene") region a character vector (default: "mrna"), make raw count matrices of whole mrnas or one of (leaders, cds, trailers). Can also be a GRangesList, then it uses this region directly. Can then be uORFs or a subset of CDS etc. type default: "count" (raw counts matrix), alternative is "fpkm", "log2fpkm" or "log10fpkm" a character(default: "default"), load files in experiment or some precomputed lib.type variant, either "ofst", "bedo", "bedoc" or "pshifted". These are made with OR-Fik:::convertLibs() or shiftFootprintsByExperiment(). Can also be custom user made folders inside the experiments bam folder. weight numeric or character, a column to score overlaps by. Default "score", will check for a metacolumn called "score" in libraries. If not found, will not use weights. forceRemake logical, default FALSE. If TRUE, will not look for existing file count table files. logical, default TRUE If TRUE, reload library files even if matching named force variables are found in environment used by experiment (see envExp) A simple way to make sure correct libraries are always loaded. FALSE is faster if data is loaded correctly already. character, default: bamVarName(df). Names to load libraries as to environment library.names and names to display in plots. **BPPARAM** how many cores/threads to use? default: BiocParallel::SerialParam()

## **Details**

If txdb or gtf path is added, it is a rangedSummerizedExperiment NOTE: If the file called saveName exists, it will then load file, not remake it!

There are different ways of counting hits on transcripts, ORFik does it as pure coverage (if a single read aligns to a region with 2 genes, both gets a count of 1 from that read). This is the safest way to avoid false negatives (genes with no assigned hits that actually have true hits).

#### Value

a SummarizedExperiment object or data.table if "type" is not "count, with rownames as transcript / gene names.

```
##Make experiment
df <- ORFik.template.experiment()
# makeSummarizedExperimentFromBam(df)
## Only cds (coding sequences):
# makeSummarizedExperimentFromBam(df, region = "cds")
## FPKM instead of raw counts on whole mrna regions
# makeSummarizedExperimentFromBam(df, type = "fpkm")
## Make count tables of pshifted libraries over uORFs
uorfs <- GRangesList(uorf1 = GRanges("chr23", 17599129:17599156, "-"))</pre>
```

242 makeTxdbFromGenome

```
#saveName <- file.path(dirname(df$filepath[1]), "uORFs", "countTable_uORFs")
#makeSummarizedExperimentFromBam(df, saveName, region = uorfs)
## To load the uORFs later
# countTable(df, region = "uORFs", count.folder = "uORFs")</pre>
```

makeTxdbFromGenome

Make txdb from genome

## **Description**

Make a Txdb with defined seqlevels and seqlevelsstyle from the fasta genome. This makes it more fail safe than standard Txdb creation. Example is that you can not create a coverage window outside the chromosome boundary, this is only possible if you have set the seqlengths.

# Usage

```
makeTxdbFromGenome(
  gtf,
  genome = NULL,
  organism,
  optimize = FALSE,
  gene_symbols = FALSE,
  uniprot_id = FALSE,
  pseudo_5UTRS_if_needed = NULL,
  minimum_5UTR_percentage = 30,
  return = is.null(txdb_file_out_path),
  txdb_file_out_path = paste0(gtf, ".db"),
  symbols_file_out_path = file.path(dirname(gtf), "gene_symbol_tx_table.fst")
)
```

## **Arguments**

gtf	path to gtf file
genome	character, default NULL. Path to fasta genome corresponding to the gtf. If NULL, can not set seqlevels. If value is NULL or FALSE, it will be ignored.
organism	Scientific name of organism, first letter must be capital! Example: Homo sapiens. Will force first letter to capital and convert any "_" (underscore) to " " (space)
optimize	logical, default FALSE. Create a folder within the output folder (defined by txdb_file_out_path), that includes optimized objects to speed up loading of annotation regions from up to 15 seconds on human genome down to 0.1 second. ORFik will then load these optimized objects instead. Currently optimizes filterTranscript() function and loadRegion() function for 5' UTRs, 3' UTRs, CDS, mRNA (all transcript with CDS) and tx (all transcripts).

mapToGRanges 243

gene\_symbols

logical default FALSE. If TRUE, will download and store all gene symbols for all transcripts (coding and noncoding)- In a file called: "gene\_symbol\_tx\_table.fst" in same folder as txdb. hgcn for human, mouse symbols for mouse and rat, more to be added.

uniprot\_id

logical default FALSE. If TRUE, will download and store all uniprot id for all transcripts (coding and noncoding)- In a file called: "gene\_symbol\_tx\_table.fst" in same folder as txdb.

pseudo\_5UTRS\_if\_needed

integer, default NULL. If defined > 0, will add pseudo 5' UTRs of maximum this length if 'minimum\_5UTR\_percentage" (default 30 mRNAs (coding transcripts) do not have a leader. (NULL and 0 are both the ignore command)

minimum\_5UTR\_percentage

numeric, default 30. What minimum percentage of mRNAs most have a 5' UTRs (leaders), to not do the pseudo\_UTR addition. If percentage is higher, addition is ignored, set to 101 to always do it.

return

 $logical, default\ FALSE.\ If\ TRUE,\ return\ TXDB\ object,\ else\ invisible (NULL).$ 

txdb\_file\_out\_path

character path, default paste0(gtf, ".db"). Set to NULL to not write file to disc.

symbols\_file\_out\_path

character path, default file.path(dirname(gtf), "gene\_symbol\_tx\_table.fst"). Must be defined as character if "gene\_symbols" is TRUE. Ignored if "gene\_symbols" is FALSE.

#### Value

logical, default is.null(txdb\_file\_out\_path), Txdb saved to disc named default paste0(gtf, ".db"). Set 'return' argument to TRUE, to also get txdb back as an object.

### **Examples**

 ${\tt mapToGRanges}$ 

Map orfs to genomic coordinates

# Description

Creates GRangesList from the results of ORFs\_as\_List and the GRangesList used to find the ORFs

244 matchColors

### Usage

```
mapToGRanges(grl, result, groupByTx = TRUE, grl_is_sorted = FALSE)
```

## **Arguments**

grl A GRangesList of the original sequences that gave the orfs in Genomic coor-

dinates. If grl\_is\_sorted = TRUE (default), negative exon ranges per grl object

must be sorted in descending orders.

result IRangesList A list of the results of finding uorfs list syntax is: Per list group in

IRangesList is per grl index. In transcript coordinates. The names are grl index

as character.

groupByTx logical (T), should output GRangesList be grouped by transcripts (T) or by

ORFs (F)?

grl\_is\_sorted logical, default FALSE If FALSE will sort negative transcript in descending

order for you. If you loaded ranges with default methods this is already the

case, so you can set to TRUE to save some time.

### **Details**

There is no check on invalid matches, so be carefull if you use this function directly.

#### Value

A GRangesList of ORFs.

### See Also

```
Other ORFHelpers: defineTrailer(), longestORFs(), orfID(), startCodons(), startSites(), stopCodons(), stopSites(), txNames(), uniqueGroups(), uniqueOrder()
```

matchColors Match coloring of coverage plot
---

### **Description**

Check that colors match with the number of fractions.

# Usage

```
matchColors(coverage, colors)
```

## **Arguments**

coverage a data.table with coverage colors a character vector of colors

matchNaming 245

### Value

number of genes in coverage

matchNaming

Match naming of GRangesList

## Description

Given a GRangesList and a reference, make the naming convention and the number of metacolumns equal to reference

### Usage

```
matchNaming(gr, reference)
```

### **Arguments**

gr a GRangesList or GRanges object reference a GRangesList of a reference

#### Value

a GRangesList

matchSeqStyle

A wrapper for seqlevelsStyle

## Description

To make sure chromosome naming is correct (chr1 vs 1 vs I etc)

#### **Usage**

```
matchSeqStyle(range, chrStyle = NULL)
```

# Arguments

range a ranged object, (GRanges, GAlignment etc)

chrStyle a GRanges object, TxDb, FaFile, , a seqlevelsStyle or Seqinfo. (Default:

NULL) to get seqlevelsStyle from. In addition if it is a Seqinfo object, seqinfo will be updated. Example of seqlevelsStyle update: Is chromosome 1 called chr1 or 1, is mitocondrial chromosome called MT or chrM etc. Will use 1st seqlevel-style if more are present. Like: c("NCBI", "UCSC") -> pick "NCBI"

### Value

a GAlignment/GRanges object depending on input.

246 mergeFastq

mergeFastq	Merge groups of Fastq /Fasta files	
------------	------------------------------------	--

## **Description**

Will use multithreading to speed up process. Only works for Unix OS (Linux and Mac)

### Usage

```
mergeFastq(in_files, out_files, BPPARAM = bpparam())
```

### **Arguments**

in_files	character specify the full path to the individual fastq.gz files. Seperated by space per file in group: For 2 output files from 4 input files: in_files <- c("file1.fastq file2.fastq". "file3.fastq file4.fastq")
out_files	character specify the path to the FASTQ directory For 2 output files: out_files <- c("/merged/file1&2.fastq", "/merged/file3&4.fastq")
BPPARAM	how many cores/threads to use? default: bpparam(). To see number of threads used, do bpparam()\$workers

#### Value

invisible(NULL).

```
fastq.folder <- tempdir() # <- Your fastq files</pre>
infiles <- dir(fastq.folder, "*.fastq", full.names = TRUE)</pre>
## Not run:
# Seperate files into groups (here it is 4 output files from 12 input files)
in_files <- c(paste0(grep(infiles, pattern = paste0("ribopool-",</pre>
               seq(11, 14), collapse = "|"), value = TRUE), collapse = " "),
              paste0(grep(infiles, pattern = paste0("ribopool-",
               seq(18, 19), collapse = "|"), value = TRUE), collapse = " "),
              paste0(grep(infiles, pattern = paste0("C11-",
               seg(11, 14), collapse = "|"), value = TRUE), collapse = " "),
              paste0(grep(infiles, pattern = paste0("C11-",
               seq(18, 19), collapse = "|"), value = TRUE), collapse = " "))
out_files <- paste0(c("SSU_ribopool", "LSU_ribopool", "SSU_WT", "LSU_WT"), ".fastq.gz")</pre>
merged.fastq.folder <- file.path(fastq.folder, "merged/")</pre>
out_files <- file.path(merged.fastq.folder, out_files)</pre>
mergeFastq(in_files, out_files)
## End(Not run)
```

mergeLibs 247

mergeLibs

Merge and save libraries of experiment

## **Description**

Aggregate count of reads (from the "score" column) by making a merged library. Only allowed for .ofst files!

# Usage

```
mergeLibs(
   df,
   out_dir = file.path(libFolder(df), "ofst_merged"),
   mode = "all",
   type = "ofst",
   keep_all_scores = TRUE,
   paths = filepath(df, type),
   lib_names_full = bamVarName(df, skip.libtype = FALSE),
   max_splits = 20
)
```

#### **Arguments**

df

an ORFik experiment

out\_dir

Ouput directory, default file.path(dirname(df\$filepath[1]), "ofst\_merged"), saved as "all.ofst" in this folder if mode is "all". Use a folder called pshifted\_merged, for default Ribo-seq ofst files.

mode

character, default "all". Merge all or "rep" for collapsing replicates only, or "lib" for collapsing all per library type.

type

a character(default: "default"), load files in experiment or some precomputed variant, like "ofst" or "pshifted". These are made with ORFik:::convertLibs(), shiftFootprintsByExperiment(), etc. Can also be custom user made folders inside the experiments bam folder. It acts in a recursive manner with priority: If you state "pshifted", but it does not exist, it checks "ofst". If no .ofst files, it uses "default", which always must exists.

Presets are (folder is relative to default lib folder, some types fall back to other formats if folder does not exist):

- "default": load the original files for experiment, usually bam.
- "ofst": loads ofst files from the ofst folder, relative to lib folder (falls back to default)
- "pshifted": loads ofst, wig or bigwig from pshifted folder (falls back to ofst, then default)
- "cov": Load covRle objects from cov\_RLE folder (fail if not found)
- "covl": Load covRleList objects, from cov\_RLE\_List folder (fail if not found)
- "bed": Load bed files, from bed folder (falls back to default)
- Other formats must be loaded directly with fimport

248 metadata.autnaming

keep\_all\_scores

logical, default TRUE, keep all library scores in the merged file. These score

columns are named the libraries full name from bamVarName(df).

paths character vector, the filpaths to use, default filepath(df, type). Change type

argument if not correct. If that is not enough, then you can also update this

argument. But be careful about using this directly.

lib\_names\_full character vector, default: bamVarName(df, skip.libtype = FALSE). Name to

assign to single libraries inside merged file, only kept if mode != "all"

max\_splits integer, default 20. If number of rows to merge > 2^31, how many times can

you allow split merging to try to "rescue" the merging process?

#### Value

NULL, files saved to disc. A data.table with a score column that now contains the sum of scores per merge setting.

### **Examples**

```
df2 <- ORFik.template.experiment()
df2 <- df2[df2$libtype == "RFP",]
# Merge all
#mergeLibs(df2, tempdir(), mode = "all", type = "default")
# Read as GRanges with mcols
#fimport(file.path(tempdir(), "all.ofst"))
# Read as direct fst data.table
#read_fst(file.path(tempdir(), "all.ofst"))
# Collapse replicates
#mergeLibs(df2, tempdir(), mode = "rep", type = "default")
# Collapse by lib types
#mergeLibs(df2, tempdir(), mode = "lib", type = "default")</pre>
```

metadata.autnaming

Guess SRA metadata columns

### **Description**

Guess SRA metadata columns

#### Usage

```
metadata.autnaming(file)
```

# **Arguments**

file a data.table of SRA metadata

# Value

a data.table of SRA metadata with additional columns: LIBRARYTYPE, REPLICATE, STAGE, CONDITION, INHIBITOR

metaWindow 249

metaWindow

Calculate meta-coverage of reads around input GRanges/List object.

# Description

Sums up coverage over set of GRanges objects as a meta representation.

## Usage

```
metaWindow(
    x,
    windows,
    scoring = "sum",
    withFrames = FALSE,
    zeroPosition = NULL,
    scaleTo = 100,
    fraction = NULL,
    feature = NULL,
    forceUniqueEven = !is.null(scoring),
    forceRescale = TRUE,
    weight = "score",
    drop.zero.dt = FALSE,
    append.zeroes = FALSE
)
```

## **Arguments**

fraction

X	GRanges/GAlignment object of your reads. Remember to resize them beforehand to width of 1 to focus on 5' ends of footprints etc, if that is wanted.
windows	GRangesList or GRanges of your ranges
scoring	a character, default: "sum", one of (zscore, transcriptNormalized, mean, median, sum, sumLength, NULL), see ?coverageScorings for info and more alternatives.
withFrames	a logical (TRUE), return positions with the 3 frames, relative to zeroPosition. zeroPosition is frame $0$ .
zeroPosition	an integer DEFAULT (NULL), the point if all windows are equal size, that should be set to position 0. Like leaders and cds combination, then 0 is the TIS and -1 is last base in leader. NOTE!: if not all windows have equal width, this will be ignored. If all have equal width and zeroPosition is NULL, it is set to as.integer(width / 2).
scaleTo	an integer (100), if windows have different size, a meta window can not directly be created, since a meta window must have equal size for all windows. Rescale (bin) all windows to scaleTo. i.e $c(1,2,3)$ -> size 2 -> coverage of position $c(1, mean(2,3))$ etc.

for large sub-unit TCP-seq.

a character/integer (NULL), the fraction i.e (27) for read length 27, or ("LSU")

250 metaWindow

feature a character string, info on region. Usually either gene name, transcript part like

cds, leader, or CpG motifs etc.

forceUniqueEven

a logical (TRUE), if TRUE; require that all windows are of same width and even.

To avoid bugs. FALSE if score is NULL.

forceRescale logical, default TRUE. If TRUE, if unique(widthPerGroup(windows)) has

length > 1, it will force all windows to width of the scaleTo argument, making

a binned meta coverage.

weight (default: 'score'), if defined a character name of valid meta column in subject.

GRanges("chr1", 1, "+", score = 5), would mean score column tells that this alignment region was found 5 times. Formats which loads a score column like this: Bigwig, wig, ORFik ofst, collapsed bam, bedoc and .bedo. As do CAGEr CAGE files and many other package formats. You can also assign a score col-

umn manually.

drop.zero.dt logical FALSE, if TRUE and as.data.table is TRUE, remove all 0 count posi-

tions. This greatly speeds up and most importantly, greatly reduces memory usage. Will not change any plots, unless 0 positions are used in some sense.

(mean, median, zscore coverage will only scale differently)

append.zeroes logical, default FALSE. If TRUE and drop.zero.dt is TRUE and all windows

have equal length, it will add back 0 values after transformation. Sometimes needed for correct plots, if TRUE, will call abort if not all windows are equal

length!

#### Value

A data.table with scored counts (score) of reads mapped to positions (position) specified in windows along with frame (frame) per gene (genes) per library (fraction) per transcript region (feature). Column that does not apply is not given, but position and (score/count) is always returned.

#### See Also

Other coverage: coverageScorings(), regionPerReadLength(), scaledWindowPositions(), windowPerReadLength()

```
model.matrix,experiment-method
```

Get experiment design model matrix

## **Description**

The function extends stats::model.matrix.

#### Usage

```
## S4 method for signature 'experiment'
model.matrix(object, design_formula = design(object, as.formula = TRUE))
```

# **Arguments**

```
object an ORFik experiment
```

design\_formula the experiment design, as formula, subset columns, to change the model.matrix,

default: design(object, as.formula = TRUE)

#### Value

a matrix with design and level attributes

# **Examples**

```
df <- ORFik.template.experiment()
model.matrix(df)</pre>
```

name

Get name of ORFik experiment

# Description

Get name of ORFik experiment

## Usage

```
name(x)
```

## **Arguments**

Χ

an ORFik experiment

#### Value

character, name of experiment

name, experiment-method

Get name of ORFik experiment

# Description

Get name of ORFik experiment

# Usage

```
## S4 method for signature 'experiment'
name(x)
```

## **Arguments**

Х

an ORFik experiment

#### Value

character, name of experiment

```
nrow, experiment-method
```

Internal nrow function for ORFik experiment Number of runs in experiment

# Description

Internal nrow function for ORFik experiment Number of runs in experiment

## Usage

```
## S4 method for signature 'experiment'
nrow(x)
```

## **Arguments**

Χ

an ORFik experiment

## Value

number of rows in experiment (integer)

numCodons 253

numCodons

Get number of codons

# Description

Length of object / 3. Choose either only whole codons, or with stubs. ORF stubs are not relevant, since there are no correctly defined ORFs that are 17 bases long etc.

## Usage

```
numCodons(grl, as.integer = TRUE, keep.names = FALSE)
```

## Arguments

grl a GRangesList object

as.integer a logical (TRUE), remove stub codons

keep.names a logical (FALSE)

## Value

an integer vector

numExonsPerGroup

Get list of the number of exons per group

# Description

Can also be used generaly to get number of GRanges object per GRangesList group

## Usage

```
numExonsPerGroup(grl, keep.names = TRUE)
```

## **Arguments**

grl a GRangesList

keep. names a logical, keep names or not, default: (TRUE)

#### Value

an integer vector of counts

254 ofst\_merge

### **Examples**

```
 \begin{split} \text{gr\_plus} &<\text{-} \text{ GRanges}(\text{seqnames} = \text{c("chr1", "chr1")}, \\ &\quad \text{ranges} = \text{IRanges}(\text{c(7, 14), width} = 3), \\ &\quad \text{strand} = \text{c("+", "+")}) \\ \text{gr\_minus} &<\text{-} \text{ GRanges}(\text{seqnames} = \text{c("chr2", "chr2")}, \\ &\quad \text{ranges} = \text{IRanges}(\text{c(4, 1), c(9, 3)}), \\ &\quad \text{strand} = \text{c("-", "-")}) \\ \text{grl} &<\text{-} \text{ GRangesList}(\text{tx1} = \text{gr\_plus}, \text{tx2} = \text{gr\_minus}) \\ \text{numExonsPerGroup}(\text{grl}) \end{aligned}
```

ofst\_merge

Merge multiple ofst file

#### **Description**

Collapses and sums the score column of each ofst file It is required that each file is of same ofst type. That is if one file has cigar information, all must have it.

### Usage

```
ofst_merge(
  file_paths,
  lib_names = sub(pattern = "\\.ofst$", replacement = "", basename(file_paths)),
  keep_all_scores = TRUE,
  keepCigar = TRUE,
  sort = TRUE,
  max_splits = 20
)
```

### **Arguments**

Full path to .ofst files wanted to merge file\_paths lib\_names character, the name to give the resulting score columns. Default: sub(pattern = "\.ofst\$", replacement = "", basename(file\_paths)) keep\_all\_scores logical, default TRUE, keep all library scores in the merged file. These score columns are named the libraries full name from bamVarName(df). logical, default TRUE. If CIGAR is defined, keep column. Setting to FALSE keepCigar compresses the file much more usually. sort logical, default TRUE. Sort the ranges. Will make the file smaller and faster to load, but some additional merging time is added. max\_splits integer, default 20. If number of rows to merge  $> 2^31$ , how many times can you allow split merging to try to "rescue" the merging process?

### Value

a data.table of merged result, it is merged on all columns except "score". The returned file will contain the scores of each file + the aggregate sum score.

```
optimizedTranscriptLengths
```

Load length and names of all transcripts

## **Description**

A speedup wrapper around transcriptLengths, default load time of lengths is ~ 15 seconds, if ORFik fst optimized lengths object has been made, load that file instead: load time reduced to ~ 0.1 second.

### Usage

```
optimizedTranscriptLengths(
   txdb,
   with.utr5_len = TRUE,
   with.utr3_len = TRUE,
   create.fst.version = FALSE,
   optimized_path = optimized_txdb_path(txdb, stop.error = FALSE)
)
```

#### **Arguments**

txdb

a TxDb object, ORFik experiment object or a path to one of: (.gtf ,.gff, .gff2, .gff2, .db or .sqlite), Only in the loadRegion function: if it is a GRangesList, it will return it self.

will return it self.
with.utr5\_len logical TRUE, include

logical TRUE, include length of 5' UTRs, ignored if .fst exists

with.utr3\_len logical TRUE, include length of 3' UTRs, ignored if .fst exists

create.fst.version

logical, FALSE. If TRUE, creates a .fst version of the transcript length table (if it not already exists), reducing load time from ~ 15 seconds to ~ 0.01 second next time you run filterTranscripts with this txdb object. The file is stored in the same folder as the genome this txdb is created from, with the name:

paste0(ORFik:::remove.file\_ext(metadata(txdb)[3,2]), "\_", gsub(" \(.\*|
|:", "", metadata(txdb)[metadata(txdb)[,1] == "Creation time",2]), "\_txLengths.fst")

Some error checks are done to see this is a valid location, if the txdb data source is a repository like UCSC and not a local folder, it will not be made.

optimized\_path

character, path to optimized txdb objects, default: optimized\_txdb\_path(txdb, stop.error = FALSE). If no existing file, will be slower and load lengths through transcriptLengths.

#### Value

a data.table of loaded lengths 8 columns, 1 row per transcript isoform.

256 optimized\_txdb\_path

### **Examples**

```
dt <- optimizedTranscriptLengths(ORFik.template.experiment())
dt
dt[cds_len > 0,] # All mRNA
```

optimized\_txdb\_path

Get path for optimization files for txdb

# Description

Get path for optimization files for txdb

### Usage

```
optimized_txdb_path(
   txdb,
   create.dir = FALSE,
   stop.error = TRUE,
   gtf_path = getGtfPathFromTxdb(txdb, stop.error = stop.error)
)
```

## **Arguments**

txdb a TxDb object, ORFik experiment object or a path to one of: (.gtf ,.gff, .gff2, .gff2, .db or .sqlite), Only in the loadRegion function: if it is a GRangesList, it

will return it self.

create.dir logical FALSE, if TRUE create the optimization directory, this should only be

called first time used.

stop.error logical TRUE

gtf\_path path to gtf where output should be stored in subfolder "./ORFik\_optimized"

## Value

a character file path, returns NULL if not valid and stop.error is FALSE.

optimizeReads 257

optimizeReads

Find optimized subset of valid reads

# **Description**

Keep only the ones that overlap within the grl ranges. Also sort them in the end

## Usage

```
optimizeReads(grl, reads)
```

## **Arguments**

grl a GRangesList or GRanges object

reads a GRanges, GAlignment or GAlignmentPairs object

## Value

the reads as GRanges, GAlignment or GAlignmentPairs

#### See Also

```
Other utils: bedToGR(), convertToOneBasedRanges(), export.bed12(), export.bigWig(), export.fstwig(), export.wiggle(), fimport(), findFa(), fread.bed(), readBam(), readBigWig(), readWig()
```

optimizeTranscriptRegions

Make optimized GRangesList objects saved to disc

## **Description**

Much faster to load

### Usage

```
optimizeTranscriptRegions(
  txdb,
  base_path = optimized_txdb_path(txdb, create.dir = TRUE),
  regions = c("tx", "mrna", "leaders", "cds", "trailers", "ncRNA")
)
```

258 orfFrameDistributions

#### **Arguments**

txdb a TxDb object, ORFik experiment object or a path to one of: (.gtf ,.gff, .gff2,

.gff2, .db or .sqlite), Only in the loadRegion function: if it is a GRangesList, it

will return it self.

base\_path Directy and file prefix for files, will append "\_region.rds", where region is spe-

cific region.

regions character, default: c("tx", "mrna", "leaders", "cds", "trailers", "ncRNA"). Valid

options specified by loadRegion.

### Value

invisible(NULL)

orfFrameDistributions Find shifted Ribo-seq frame distributions

### **Description**

Per library: get coverage over CDS per frame per readlength Return as data.datable with information and best frame found. Can be used to automize re-shifting of read lengths (find read lengths where frame 0 is not the best frame over the entire cds)

## Usage

```
orfFrameDistributions(
   df,
   type = "pshifted",
   weight = "score",
   orfs = loadRegion(df, part = "cds"),
   BPPARAM = BiocParallel::bpparam()
)
```

### **Arguments**

df an ORFik experiment

type type of library loaded, default pshifted, warning if not pshifted might crash if

too many read lengths!

weight which column in reads describe duplicates, default "score".

orfs GRangesList, default loadRegion(df, part = "cds")

BPPARAM how many cores/threads to use? default: bpparam(). To see number of threads

used, do bpparam() \$workers. You can also add a time remaining bar, for a

more detailed pipeline.

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## Value

data.table with columns: fraction (library) frame (0, 1, 2) score (coverage) length (read length) percent (coverage percentage of library) percent\_length (coverage percentage of library and length) best\_frame (TRUE/FALSE, is this the best frame per length)

# Examples

```
df <- ORFik.template.experiment()[3,]
dt <- orfFrameDistributions(df, BPPARAM = BiocParallel::SerialParam())
## Check that frame 0 is best frame for all
all(dt[frame == 0,]$best_frame)</pre>
```

orfID

Get id's for each orf

## **Description**

These id's can be uniqued by isoform etc, this is not supported by GenomicRanges.

### Usage

```
orfID(grl, with.tx = FALSE)
```

### **Arguments**

grl a GRangesList

with.tx a boolean, include transcript names, if you want unique orfs, so that they dont

have duplicates from different isoforms, set it to FALSE.

## Value

```
a character vector of ids, 1 per orf
```

### See Also

```
Other ORFHelpers: defineTrailer(), longestORFs(), mapToGRanges(), startCodons(), startSites(), stopCodons(), stopSites(), txNames(), uniqueGroups(), uniqueOrder()
```

ORFik.template.experiment

An ORFik experiment to see how it looks

## **Description**

Toy-data created to resemble human genes:

Number of genes: 6

Genome size: 1161nt x 6 chromosomes = 6966 nt Experimental design (2 replicates, Wild type vs Mutant)

CAGE: 4 libraries PAS (poly-A): 4 libraries Ribo-seq: 4 libraries RNA-seq: 4 libraries

### Usage

```
ORFik.template.experiment(as.temp = FALSE)
```

# Arguments

as.temp

logical, default FALSE, load as ORFik experiment. If TRUE, loads as data.frame template of the experiment.

# Value

an ORFik experiment

## See Also

```
Other ORFik_experiment: ORFik.template.experiment.zf(), bamVarName(), create.experiment(), experiment-class, filepath(), libraryTypes(), organism, experiment-method, outputLibs(), read.experiment(), save.experiment(), validateExperiments()
```

## **Examples**

```
ORFik.template.experiment()
```

 ${\tt ORFik.template.experiment.zf}$ 

An ORFik experiment to see how it looks

#### **Description**

Toy-data created to resemble Zebrafish genes:

Number of genes: 150 Ribo-seq: 1 library

### Usage

```
ORFik.template.experiment.zf(as.temp = FALSE)
```

### **Arguments**

as.temp

logical, default FALSE, load as ORFik experiment. If TRUE, loads as data.frame template of the experiment.

#### Value

an ORFik experiment

### See Also

```
Other ORFik_experiment: ORFik.template.experiment(), bamVarName(), create.experiment(), experiment-class, filepath(), libraryTypes(), organism, experiment-method, outputLibs(), read.experiment(), save.experiment(), validateExperiments()
```

### **Examples**

```
ORFik.template.experiment.zf()
```

ORFikQC

A post Alignment quality control of reads

## **Description**

The ORFik QC uses the aligned files (usually bam files), fastp and STAR log files combined with annotation to create relevant statistics.

This report consists of several steps:

- 1. Convert bam file / Input files to ".ofst" format, if not already done. This format is around 400x faster to use in R than the bam format. Files are also outputted to R environment specified by envExp(df)
- 2. From this report you will get a summary csv table, with distribution of aligned reads and overlap

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counts over transcript regions like: leader, cds, trailer, lincRNAs, tRNAs, rRNAs, snoRNAs etc. It will be called STATS.csv. And can be imported with QCstats function.

- 3. It will also make correlation plots and meta coverage plots, so you get a good understanding of how good the quality of your NGS data production + aligner step were.
- 4. Count tables are produced, similar to HTseq count tables. Over mrna, leader, cds and trailer separately. This tables are stored as SummarizedExperiment, for easy loading into DEseq, conversion to normalized fpkm values, or collapsing replicates in an experiment. And can be imported with countTable function.

Everything will be outputed in the directory of your NGS data, inside the folder ./QC\_STATS/, relative to data location in 'df'. You can specify new out location with out.dir if you want.

To make a ORFik experiment, see ?ORFik::experiment

To see some normal mrna coverage profiles of different RNA-seq protocols: https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4310221/figure/F6/

#### Usage

```
ORFikQC(
    df,
    out.dir = resFolder(df),
    plot.ext = ".pdf",
    create.ofst = TRUE,
    complex.correlation.plots = TRUE,
    library.names = bamVarName(df),
    use_simplified_reads = TRUE,
    BPPARAM = bpparam()
)
```

#### **Arguments**

df an ORFik experiment

out.dir character, output directory, default: resFolder(df). Will make a folder within

this called "QC\_STATS" with all results in this directory. Warning: If you assign not default path, you will have a hazzle to load files later. Much easier to load count tables, statistics, ++ later with default. Update resFolder of df instead if

needed.

plot.ext character, default: ".pdf". Alternatives: ".png" or ".jpg". Note that in pdf format

the complex correlation plots become very slow to load!

create.ofst logical, default TRUE. Create ".ofst" files from the input libraries, ofst is much

faster to load in R, for later use. Stored in ./ofst/ folder relative to experiment

main folder.

complex.correlation.plots

logical, default TRUE. Add in addition to simple correlation plot two computationally heavy dots + correlation plots. Useful for deeper analysis, but takes longer time to run, especially on low-quality gpu computers. Set to FALSE to

skip these.

 $\label{library.names} \mbox{ character, default: } bamVarName (df). \mbox{ Names to load libraries as to environment}$ 

and names to display in plots.

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use\_simplified\_reads

logical, default TRUE. For count tables and coverage plots a speed up for GAlignments is to use 5' ends only. This will lose some detail for splice sites, but is usually irrelevant. Note: If reads are precollapsed GRanges, set to FALSE to avoid recollapsing.

**BPPARAM** 

how many cores/threads to use? default: bpparam(). To see number of threads used, do bpparam()\$workers. You can also add a time remaining bar, for a more detailed pipeline.

### Value

```
invisible(NULL) (objects are stored to disc)
```

### See Also

```
Other QC report: QCplots(), QCstats()
```

## **Examples**

```
# Load an experiment
df <- ORFik.template.experiment()
# Run QC
#QCreport(df, tempdir())
# QC on subset
#QCreport(df[9,], tempdir())</pre>
```

orfScore

Get ORFscore for a GRangesList of ORFs

# Description

ORFscore tries to check whether the first frame of the 3 possible frames in an ORF has more reads than second and third frame. IMPORTANT: Only use p-shifted libraries, see (detectRibosomeShifts). Else this score makes no sense.

# Usage

```
orfScore(
  grl,
  RFP,
  is.sorted = FALSE,
  weight = "score",
  overlapGrl = NULL,
  coverage = NULL,
  stop3 = TRUE
)
```

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#### **Arguments**

grl	a GRangesList of 5' utrs, CDS, transcripts, etc.
RFP	ribosomal footprints, given as GAlignments or GRanges object, must be already shifted and resized to the p-site. Requires a \$size column with original read lengths.
is.sorted	logical (FALSE), is grl sorted. That is $+$ strand groups in increasing ranges $(1,2,3)$ , and $-$ strand groups in decreasing ranges $(3,2,1)$
weight	(default: 'score'), if defined a character name of valid meta column in subject. GRanges("chr1", 1, "+", score = 5), would mean score column tells that this alignment region was found 5 times. Formats which loads a score column like this: Bigwig, wig, ORFik ofst, collapsed bam, bedoc and .bedo. As do CAGEr CAGE files and many other package formats. You can also assign a score column manually.
overlapGrl	an integer, (default: NULL), if defined must be countOverlaps(grl, RFP), added for speed if you already have it.
coverage	a data.table from coveragePerTiling of length same as 'grl' argument. Save time if you have already computed it.
stop3	logical, default TRUE. Stop if any input is of width < 3.

#### **Details**

Pseudocode: assume rff - is reads fraction in specific frame

```
ORFScore = log(rff1 + rff2 + rff3)
```

If rff2 or rff3 is bigger than rff1, negate the resulting value.

```
ORFScore[rff1Smaller] <- ORFScore[rff1Smaller] * -1</pre>
```

As result there is one value per ORF: - Positive values say that the first frame have the most reads, - zero values means it is uniform: (ORFscore between -2.5 and 2.5 can be considered close to uniform), - negative values say that the first frame does not have the most reads. NOTE non-pshifted reads: If reads are not of width 1, then a read from 1-4 on range of 1-4, will get scores frame1 = 2, frame2 = 1, frame3 = 1. What could be logical is that only the 5' end is important, so that only frame1 = 1, to get this, you first resize reads to 5'end only.

General NOTES: 1. p shifting is not exact, so some functional ORFs will get a bad ORF score.

2. If a score column is defined, it will use it as weights, set to weight = 1L if you don't have weight, and score column is something else.

3. If needed a test for significance and critical values, use chi-squared. There are 3 degrees of freedom (3 frames), so critical 0.05 (3-1 degrees of freedm = 2), value is: log2(6) = 2.58 see getWeights

### Value

a data.table with 4 columns, the orfscore (ORFScores) and score of each of the 3 tiles (frame\_zero\_RP, frame\_one\_RP, frame\_two\_RP)

### References

```
doi: 10.1002/embj.201488411
```

#### See Also

```
Other features: computeFeatures(), computeFeaturesCage(), countOverlapsW(), disengagementScore(), distToCds(), distToTSS(), entropy(), floss(), fpkm(), fpkm_calc(), fractionLength(), initiationScore(), insideOutsideORF(), isInFrame(), isOverlapping(), kozakSequenceScore(), rankOrder(), ribosomeReleaseScore(), ribosomeStallingScore(), startRegion(), startRegionCoverage(), stopRegion(), subsetCoverage(), translationalEff()
```

### **Examples**

organism, experiment-method

Get ORFik experiment organism

### Description

If not defined directly, checks the txdb / gtf organism information, if existing.

## Usage

```
## S4 method for signature 'experiment'
organism(object)
```

### **Arguments**

```
object an ORFik experiment
```

### Value

character, name of organism

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### See Also

```
Other ORFik_experiment: ORFik.template.experiment(), ORFik.template.experiment.zf(), bamVarName(), create.experiment(), experiment-class, filepath(), libraryTypes(), outputLibs(), read.experiment(), save.experiment(), validateExperiments()
```

## **Examples**

```
# if you have set organism in txdb of ORFik experiment:
df <- ORFik.template.experiment()
organism(df)

#' If you have not set the organism you can do:
#gtf <- "pat/to/gff_or_gff"

#txdb_path <- paste0(gtf, ".db") # This file is created in next step
#txdb <- makeTxdbFromGenome(gtf, genome, organism = "Homo sapiens",
# optimize = TRUE, return = TRUE)
# then use this txdb in you ORFik experiment and load:
# create.experiment(exper = "new_experiment",
# txdb = txdb_path) ...
# organism(read.experiment("new-experiment))</pre>
```

outputLibs

Output NGS libraries to R as variables

### Description

By default loads the original files of the experiment into the global environment, named by the rows of the experiment required to make all libraries have unique names.

Uses multiple cores to load, defined by multicoreParam

## Usage

```
outputLibs(
   df,
   type = "default",
   paths = filepath(df, type),
   param = NULL,
   strandMode = 0,
   naming = "minimum",
   library.names = name_decider(df, naming),
   output.mode = "envir",
   chrStyle = NULL,
   envir = envExp(df),
   verbose = TRUE,
   force = TRUE,
   BPPARAM = bpparam()
)
```

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#### **Arguments**

df

an ORFik experiment

type

a character(default: "default"), load files in experiment or some precomputed variant, like "ofst" or "pshifted". These are made with ORFik:::convertLibs(), shiftFootprintsByExperiment(), etc. Can also be custom user made folders inside the experiments bam folder. It acts in a recursive manner with priority: If you state "pshifted", but it does not exist, it checks "ofst". If no .ofst files, it uses "default", which always must exists.

Presets are (folder is relative to default lib folder, some types fall back to other formats if folder does not exist):

- "default": load the original files for experiment, usually bam.
- "ofst": loads ofst files from the ofst folder, relative to lib folder (falls back to default)
- "pshifted": loads ofst, wig or bigwig from pshifted folder (falls back to ofst, then default)
- "cov": Load covRle objects from cov RLE folder (fail if not found)
- "covl": Load covRleList objects, from cov RLE List folder (fail if not found)
- "bed": Load bed files, from bed folder (falls back to default)
- Other formats must be loaded directly with fimport

paths

character vector, the filpaths to use, default filepath(df, type). Change type argument if not correct. If that is not enough, then you can also update this argument. But be careful about using this directly.

param

NULL or a ScanBamParam object. Like for scanBam, this influences what fields and which records are imported. However, note that the fields specified thru this ScanBamParam object will be loaded in addition to any field required for generating the returned object (GAlignments, GAlignmentPairs, or GappedReads object), but only the fields requested by the user will actually be kept as metadata columns of the object.

By default (i.e. param=NULL or param=ScanBamParam()), no additional field is loaded. The flag used is scanBamFlag(isUnmappedQuery=FALSE) for readGAlignments, readGAlignmentsList, and readGappedReads. (i.e. only records corresponding to mapped reads are loaded), and scanBamFlag(isUnmappedQuery=FALSE, isPaired=TRUE, hasUnmappedMate=FALSE) for readGAlignmentPairs (i.e. only records corresponding to paired-end reads with both ends mapped are loaded).

strandMode

numeric, default 0. Only used for paired end bam files. One of (0: strand = \*, 1: first read of pair is +, 2: first read of pair is -). See ?strandMode. Note: Sets default to 0 instead of 1, as readGAlignmentPairs uses 1. This is to guarantee hits, but will also make mismatches of overlapping transcripts in opposite directions.

naming

a character (default: "minimum"). Name files as minimum information needed to make all files unique. Set to "full" to get full names. Set to "fullexp", to get full name with experiment name as prefix, the last one guarantees uniqueness.

library.names

character vector, names of libraries, default: name\_decider(df, naming)

character, default "envir". Output libraries to environment. Alternative: "list", return as list. "envirlist", output to envir and return as list. If output is list format, the list elements are named from: bamVarName(df.rfp) (Full or minimum naming based on 'naming' argument)

output.mode

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chrStyle	a GRanges object, TxDb, FaFile, , a seqlevelsStyle or Seqinfo. (Default: NULL) to get seqlevelsStyle from. In addition if it is a Seqinfo object, seqinfo will be updated. Example of seqlevelsStyle update: Is chromosome 1 called chr1 or 1, is mitocondrial chromosome called MT or chrM etc. Will use 1st seqlevel-style if more are present. Like: c("NCBI", "UCSC") -> pick "NCBI"
envir	environment to save to, default envExp(df), which defaults to .GlobalEnv, but can be set with envExp(df) <- new.env() etc.
verbose	logical, default TRUE, message about library output status.
force	logical, default TRUE If TRUE, reload library files even if matching named variables are found in environment used by experiment (see envExp) A simple way to make sure correct libraries are always loaded. FALSE is faster if data is loaded correctly already.
BPPARAM	how many cores/threads to use? default: bpparam(). To see number of threads used, do bpparam()\$workers. You can also add a time remaining bar, for a more detailed pipeline.

#### **Details**

The functions checks if the total set of libraries have already been loaded: i.e. Check if all names from 'library.names' exists as S4 objects in environment of experiment.

#### Value

NULL (libraries set by envir assignment), unless output.mode is "list" or "envirlist": Then you get a list of the libraries.

## See Also

```
Other ORFik_experiment: ORFik.template.experiment(), ORFik.template.experiment.zf(), bamVarName(), create.experiment(), experiment-class, filepath(), libraryTypes(), organism, experiment-methread.experiment(), save.experiment(), validateExperiments()
```

# Examples

```
## Load a template ORFik experiment
df <- ORFik.template.experiment()
## Default library type load, usually bam files
# outputLibs(df, type = "default")
## .ofst file load, if ofst files does not exists
## it will load default
# outputLibs(df, type = "ofst")
## .wig file load, if wiggle files does not exists
## it will load default
# outputLibs(df, type = "wig")
## Load as list
outputLibs(df, output.mode = "list")
## Load libs to new environment (called ORFik in Global)
# outputLibs(df, envir = assign(name(df), new.env(parent = .GlobalEnv)))
## Load to hidden environment given by experiment</pre>
```

pasteDir 269

```
# envExp(df) <- new.env()
# outputLibs(df)</pre>
```

pasteDir

A paste function for directories Makes sure slashes are corrected, and not doubled.

### **Description**

A paste function for directories Makes sure slashes are corrected, and not doubled.

## Usage

```
pasteDir(...)
```

## **Arguments**

... any amount of arguments that are possible to convert to characters

## Value

the pasted string

pcaExperiment

Simple PCA analysis

### **Description**

Detect outlier libraries with PCA analysis. Will output PCA plot of PCA component 1 (x-axis) vs PCA component 2 (y-axis) for each library (colored by library), shape by replicate. Will be extended to allow batch correction in the future.

## Usage

```
pcaExperiment(
   df,
   output.dir = NULL,
   table = countTable(df, "cds", type = "fpkm"),
   title = "PCA analysis by CDS fpkm",
   subtitle = paste("Numer of genes/regions:", nrow(table)),
   plot.ext = ".pdf",
   return.data = FALSE,
   color.by.group = TRUE
)
```

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## **Arguments**

df an ORFik experiment output.dir default NULL, else character path to directory. File saved as "PCAplot\_(experiment name)(plot.ext)" table data.table, default countTable(df, "cds", type = "fpkm"), a data.table of counts per column (default normalized fpkm values). title character, default "CDS fpkm". subtitle character, default: paste("Numer of genes:", nrow(table)) plot.ext character, default: ".pdf". Alternatives: ".png" or ".jpg". Note that in pdf format the complex correlation plots become very slow to load! logical, default FALSE. Return data instead of plot return.data color.by.group logical, default TRUE. Colors in PCA plot represent unique library groups, if FALSE. Color each sample in seperate color (harder to distinguish for > 10

#### Value

ggplot or invisible(NULL) if output.dir is defined or < 3 samples. Returns data.table with PCA analysis if return.data is TRUE.

### **Examples**

### **Description**

```
50 \rightarrow 0.5 etc, if length cds > minimum.cds
```

samples)

## Usage

```
percentage_to_ratio(top_tx, cds, minimum.cds = 1000)
```

### **Arguments**

top\_tx numeric

cds GRangesList object minimum.cds numeric, default 1000

### Value

```
numeric, as ratio of 1
```

plotHelper 271

plotHelper

Helper function for coverage plots

## **Description**

Should only be used internally

### Usage

```
plotHelper(
  coverage,
  df,
  outdir,
  scores,
  returnCoverage = FALSE,
  title = "coverage metaplot",
  plot.ext = ".pdf",
  colors = c("skyblue4", "orange"),
  plotFunction = "windowCoveragePlot"
)
```

### **Arguments**

coverage a data.table containing at least columns (count/score, position), it is possible to

have additionals: (genes, fraction, feature)

df an ORFik experiment

outdir directory to save to (default: NULL, no saving)

scores scoring function (default: c("sum", "transcriptNormalized")), see ?coverageScor-

ings for possible scores.

returnCoverage (defualt: FALSE), return the ggplot object (TRUE) or NULL (FALSE).

title Title to give plot

plot.ext character, default: ".pdf". Alternatives: ".png" or ".jpg".

colors Which colors to use, default auto color from function experiment.colors, new

color per library type. Else assign colors yourself.

plotFunction Which plot function, default: windowCoveragePlot

### Value

NULL (or ggplot object if returnCoverage is TRUE)

pmapFromTranscriptF Faster pmapFromTranscript

## **Description**

Map range coordinates between features in the transcriptome and genome (reference) space. The length of x must be the same as length of transcripts. Only exception is if x have integer names like (1, 3, 3, 5), so that x[1] maps to 1, x[2] maps to transcript 3 etc.

# Usage

```
pmapFromTranscriptF(x, transcripts, removeEmpty = FALSE)
```

# Arguments

x IRangesList/IRanges/GRanges to map to genomic coordinates
transcripts a GRangesList to map against (the genomic coordinates)
removeEmpty a logical, remove non hit exons, else they are set to 0. That is all exons in the reference that the transcript coordinates do not span.

#### Details

This version tries to fix the short commings of GenomicFeature's version. Much faster and uses less memory. Implemented as dynamic program optimized c++ code.

### Value

a GRangesList of mapped reads, names from ranges are kept.

### **Examples**

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Faster pmapToTranscript

# Description

Map range coordinates between features in the transcriptome and genome (reference) space. The length of x must be the same as length of transcripts. Only exception is if x have integer names like (1, 3, 3, 5), so that x[1] maps to 1, x[2] maps to transcript 3 etc.

# Usage

```
pmapToTranscriptF(
    x,
    transcripts,
    ignore.strand = FALSE,
    x.is.sorted = TRUE,
    tx.is.sorted = TRUE
)
```

## Arguments

X	$GRangesList/GRanges/IRangesList/IRanges\ to\ map\ to\ transcriptomic\ coordinates$
transcripts	a GRangesList/GRanges/IRangesList/IRanges to map against (the genomic coordinates). Must be of lower abstraction level than $x$ . So if $x$ is GRanges, transcripts can not be IRanges etc.
ignore.strand	When ignore.strand is TRUE, strand is ignored in overlaps operations (i.e., all strands are considered "+") and the strand in the output is '*'.  When ignore.strand is FALSE (default) strand in the output is taken from the transcripts argument. When transcripts is a GRangesList, all inner list elements of a common list element must have the same strand or an error is thrown.  Mapped position is computed by counting from the transcription start site (TSS) and is not affected by the value of ignore.strand.
x.is.sorted	if x is a GRangesList object, are "-" strand groups pre-sorted in decreasing order within group, default: TRUE $$
tx.is.sorted	if transcripts is a GRangesList object, are "-" strand groups pre-sorted in decreasing order within group, default: TRUE

## **Details**

This version tries to fix the shortcommings of GenomicFeature's version. Much faster and uses less memory. Implemented as dynamic program optimized c++ code.

## Value

object of same class as input x, names from ranges are kept.

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### **Examples**

```
library(GenomicFeatures)
# Need 2 ranges object, the target region and whole transcript
# x is target region
x \leftarrow GRanges("chr1", IRanges(start = c(26, 29), end = c(27, 29)), "+")
names(x) \leftarrow rep("tx1_ORF1", length(x))
x <- groupGRangesBy(x)</pre>
# tx is the whole region
tx_gr \leftarrow GRanges("chr1", IRanges(c(5, 29), c(27, 30)), "+")
names(tx_gr) <- rep("tx1", length(tx_gr))</pre>
tx <- groupGRangesBy(tx_gr)</pre>
pmapToTranscriptF(x, tx)
pmapToTranscripts(x, tx)
# Reuse names for matching
x \leftarrow GRanges("chr1", IRanges(start = c(26, 29, 5), end = c(27, 29, 18)), "+")
names(x) < c(rep("tx1_1", 2), "tx1_2")
x <- groupGRangesBy(x)</pre>
tx1_2 <- GRanges("chr1", IRanges(c(4, 28), c(26, 31)), "+")
names(tx1_2) <- rep("tx1", 2)
tx <- c(tx, groupGRangesBy(tx1_2))</pre>
a <- pmapToTranscriptF(x, tx[txNames(x)])</pre>
b <- pmapToTranscripts(x, tx[txNames(x)])</pre>
identical(a, b)
seqinfo(a)
# A note here, a & b only have 1 seqlength, even though the 2 "tx1"
# are different in size. This is an artifact of using duplicated names.
## Also look at the asTx for a similar useful function.
```

prettyScoring

Prettify scoring name

### **Description**

Prettify scoring name

#### Usage

```
prettyScoring(scoring)
```

### **Arguments**

```
scoring a character (the scoring)
```

#### Value

a new scoring name or the same if pretty

pseudo.transform 275

pseudo.transform

Transform object

### **Description**

Similar to normal transform like log2 or log10. But keep 0 values as 0, to avoid Inf values and negtive values are made as -scale(abs(x)), to avoid NaN values.

### Usage

```
pseudo.transform(x, scale = log2, by.reference = FALSE)
```

## **Arguments**

x a numeric vector or data.frame/data.table of numeric columns
scale a function, default log2, which function to transform with.
by.reference logical, FALSE. if TRUE, update object by reference if it is data.table.

#### Value

same object class as x, with transformed values

pSitePlot

Plot area around TIS as histogram

## **Description**

Usefull to validate p-shifting is correct Can be used for any coverage of region around a point, like TIS, TSS, stop site etc.

## Usage

```
pSitePlot(
  hitMap,
  length = unique(hitMap$fraction),
  region = "start",
  output = NULL,
  type = "canonical CDS",
  scoring = "Averaged counts",
  forHeatmap = FALSE,
  title = "auto",
  facet = FALSE,
  frameSum = FALSE
)
```

pSitePlot

### **Arguments**

hitMap	a data.frame/data.table, given from metaWindow (must have columns: position, (score or count) and frame)
length	an integer (29), which read length is this for?
region	a character (start), either "start or "stop"
output	character (NULL), if set, saves the plot as pdf or png to path given. If no format is given, is save as pdf.
type	character (canonical CDS), type for plot
scoring	character, default: (Averaged counts), which scoring did you use ? see ?coverageScorings for info and more alternatives.
forHeatmap	a logical (FALSE), should the plot be part of a heatmap? It will scale it differently. Removing title, x and y labels, and truncate spaces between bars.
title	character, title of plot. Default "auto", will make it: paste("Length", length, "over", region, "of", type). Else set your own (set to NULL to remove all together).
facet	logical, default FALSE. If you input multiple read lengths, specified by fraction column of hitMap, it will split the plots for each read length, putting them under each other. Ignored if forHeatmap is TRUE.
frameSum	logical default FALSE. If TRUE, add an addition plot to the right, sum per frame over all positions per length.

### **Details**

The region is represented as a histogram with different colors for the 3 frames. To make it easy to see patterns in the reads. Remember if you want to change anything like colors, just return the ggplot object, and reassign like: obj + scale\_color\_brewer() etc.

### Value

a ggplot object of the coverage plot, NULL if output is set, then the plot will only be saved to location.

### See Also

Other coveragePlot: coverageHeatMap(), savePlot(), windowCoveragePlot()

# **Examples**

QCfolder 277

```
# See vignette for more examples
```

QCfolder

Get ORFik experiment QC folder path

# Description

Get ORFik experiment QC folder path

# Usage

```
QCfolder(x)
```

# Arguments

Х

an ORFik experiment

### Value

a character path

```
QCfolder, experiment-method
```

Get ORFik experiment QC folder path

# Description

Get ORFik experiment QC folder path

# Usage

```
## S4 method for signature 'experiment'
QCfolder(x)
```

## Arguments

Х

an ORFik experiment

### Value

a character path

278 QCplots

QCplots

Correlation and coverage plots for ORFikQC

## **Description**

Correlation plots default to mRNA covering reads. Meta plots defaults to leader, cds, trailer. Output will be stored in same folder as the libraries in df.

Correlation plots will be fpkm correlation and log2(fpkm + 1) correlation between samples.

### Usage

```
QCplots(
   df,
   region = "mrna",
   stats_folder = QCfolder(df),
   plot.ext = ".pdf",
   complex.correlation.plots = TRUE,
   library.names = bamVarName(df),
   force = TRUE,
   BPPARAM
)
```

#### **Arguments**

df an ORFik experiment

region a character (default: mrna), make raw count matrices of whole mrnas or one of

(leaders, cds, trailers)

stats\_folder directory to save, default: QCfolder(df)

plot.ext character, default: ".pdf". Alternatives: ".png" or ".jpg". Note that in pdf format

the complex correlation plots become very slow to load!

complex.correlation.plots

logical, default TRUE. Add in addition to simple correlation plot two computationally heavy dots + correlation plots. Useful for deeper analysis, but takes longer time to run, especially on low-quality gpu computers. Set to FALSE to

skip these.

library.names character vector, names of libraries, default: name\_decider(df, naming)

force logical, default TRUE If TRUE, reload library files even if matching named

variables are found in environment used by experiment (see envExp) A simple way to make sure correct libraries are always loaded. FALSE is faster if data is

loaded correctly already.

BPPARAM how many cores/threads to use? default: bpparam(). To see number of threads

used, do bpparam()\$workers. You can also add a time remaining bar, for a

more detailed pipeline.

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#### **Details**

```
Is part of QCreport
```

#### Value

invisible(NULL) (objects stored to disc)

#### See Also

```
Other QC report: QCreport(), QCstats()
```

QCreport

A post Alignment quality control of reads

# Description

The ORFik QC uses the aligned files (usually bam files), fastp and STAR log files combined with annotation to create relevant statistics.

This report consists of several steps:

- 1. Convert bam file / Input files to ".ofst" format, if not already done. This format is around 400x faster to use in R than the bam format. Files are also outputted to R environment specified by envExp(df)
- 2. From this report you will get a summary csv table, with distribution of aligned reads and overlap counts over transcript regions like: leader, cds, trailer, lincRNAs, tRNAs, rRNAs, snoRNAs etc. It will be called STATS.csv. And can be imported with QCstats function.
- 3. It will also make correlation plots and meta coverage plots, so you get a good understanding of how good the quality of your NGS data production + aligner step were.
- 4. Count tables are produced, similar to HTseq count tables. Over mrna, leader, cds and trailer separately. This tables are stored as SummarizedExperiment, for easy loading into DEseq, conversion to normalized fpkm values, or collapsing replicates in an experiment. And can be imported with countTable function.

Everything will be outputed in the directory of your NGS data, inside the folder ./QC\_STATS/, relative to data location in 'df'. You can specify new out location with out.dir if you want.

To make a ORFik experiment, see ?ORFik::experiment

To see some normal mrna coverage profiles of different RNA-seq protocols: https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4310221/figure/F6/

# Usage

```
QCreport(
   df,
   out.dir = resFolder(df),
   plot.ext = ".pdf",
   create.ofst = TRUE,
   complex.correlation.plots = TRUE,
```

280 QCreport

```
library.names = bamVarName(df),
use_simplified_reads = TRUE,
BPPARAM = bpparam()
)
```

#### **Arguments**

df an ORFik experiment

out.dir character, output directory, default: resFolder(df). Will make a folder within

this called "QC\_STATS" with all results in this directory. Warning: If you assign not default path, you will have a hazzle to load files later. Much easier to load count tables, statistics, ++ later with default. Update resFolder of df instead if

needed.

plot.ext character, default: ".pdf". Alternatives: ".png" or ".jpg". Note that in pdf format

the complex correlation plots become very slow to load!

create.ofst logical, default TRUE. Create ".ofst" files from the input libraries, ofst is much

faster to load in R, for later use. Stored in ./ofst/ folder relative to experiment

main folder.

complex.correlation.plots

logical, default TRUE. Add in addition to simple correlation plot two computationally heavy dots + correlation plots. Useful for deeper analysis, but takes longer time to run, especially on low-quality gpu computers. Set to FALSE to

skip these.

 $\label{library.names} \mbox{ character, default: } bamVarName(df). \ Names \ to \ load \ libraries \ as \ to \ environment$ 

and names to display in plots.

use\_simplified\_reads

logical, default TRUE. For count tables and coverage plots a speed up for GAlignments is to use 5' ends only. This will lose some detail for splice sites, but is usually irrelevant. Note: If reads are precollapsed GRanges, set to FALSE to

avoid recollapsing.

BPPARAM how many cores/threads to use? default: bpparam(). To see number of threads

used, do bpparam() \$workers. You can also add a time remaining bar, for a

more detailed pipeline.

#### Value

```
invisible(NULL) (objects are stored to disc)
```

### See Also

```
Other QC report: QCplots(), QCstats()
```

# Examples

```
# Load an experiment
df <- ORFik.template.experiment()
# Run QC
#QCreport(df, tempdir())</pre>
```

QCstats 281

```
# QC on subset
#QCreport(df[9,], tempdir())
```

QCstats

Load ORFik QC Statistics report

# Description

Loads the pre / post alignment statistcs made in ORFik.

## Usage

```
QCstats(df, path = file.path(QCfolder(df), "STATS.csv"))
```

## **Arguments**

df an ORFik experiment

path to QC statistics report, default: file.path(dirname(df\$filepath[1]), "/QC\_STATS/STATS.csv")

### **Details**

The ORFik QC uses the aligned files (usually bam files), fastp and STAR log files combined with annotation to create relevant statistics.

#### Value

```
data.table of QC report or NULL if not exists
```

# See Also

```
Other QC report: QCplots(), QCreport()
```

# **Examples**

```
df <- ORFik.template.experiment()
## First make QC report
# QCreport(df)
# stats <- QCstats(df)</pre>
```

QC\_count\_tables

QCStats.p10t Make piot of OKPik QCTeport	QCstats.plot	Make plot of ORFik QCreport	
--	--------------	-----------------------------	--

# Description

From post-alignment QC relative to annotation, make a plot for all samples. Will contain among others read lengths, reads overlapping leaders, cds, trailers, mRNA / rRNA etc.

# Usage

```
QCstats.plot(stats, output.dir = NULL, plot.ext = ".pdf", as_gg_list = FALSE)
```

## **Arguments**

stats	the experiment object or path to custom ORFik QC folder where a file called "STATS.csv" is located.
output.dir	NULL or character path, default: NULL, plot not saved to disc. If defined saves plot to that directory with the name "/STATS_plot.pdf".
plot.ext	character, default: ".pdf". Alternatives: ".png" or ".jpg".
as_gg_list	logical, default FALSE. Return as a list of ggplot objects instead of as a grob. Gives you the ability to modify plots more directly.

# Value

the plot object, a grob of ggplot objects of the the statistics data

# **Examples**

```
df <- ORFik.template.experiment()[3,]
## First make QC report
# QCreport(df)
## Now you can get plot
# QCstats.plot(df)</pre>
```

QC_count_tables	Create count table info for QC report	
-----------------	---------------------------------------	--

# Description

The better the annotation / gtf used, the more results you get.

QC\_count\_tables 283

### Usage

```
QC_count_tables(
    df,
    out.dir,
    type = "ofst",
    use_simplified_reads = TRUE,
    force = TRUE,
    library.names = bamVarName(df),
    BPPARAM = bpparam()
)
```

#### **Arguments**

df

an ORFik experiment

out.dir

character, output directory, default: resFolder(df). Will make a folder within this called "QC\_STATS" with all results in this directory. Warning: If you assign not default path, you will have a hazzle to load files later. Much easier to load count tables, statistics, ++ later with default. Update resFolder of df instead if needed.

type

a character(default: "default"), load files in experiment or some precomputed variant, like "ofst" or "pshifted". These are made with ORFik:::convertLibs(), shiftFootprintsByExperiment(), etc. Can also be custom user made folders inside the experiments bam folder. It acts in a recursive manner with priority: If you state "pshifted", but it does not exist, it checks "ofst". If no .ofst files, it uses "default", which always must exists.

Presets are (folder is relative to default lib folder, some types fall back to other formats if folder does not exist):

- "default": load the original files for experiment, usually bam.
- "ofst": loads ofst files from the ofst folder, relative to lib folder (falls back to default)
- "pshifted": loads ofst, wig or bigwig from pshifted folder (falls back to ofst, then default)
- "cov": Load covRle objects from cov\_RLE folder (fail if not found)
- "covl": Load covRleList objects, from cov\_RLE\_List folder (fail if not found)
- "bed": Load bed files, from bed folder (falls back to default)
- Other formats must be loaded directly with fimport

# use\_simplified\_reads

logical, default TRUE. For count tables and coverage plots a speed up for GAlignments is to use 5' ends only. This will lose some detail for splice sites, but is usually irrelevant. Note: If reads are precollapsed GRanges, set to FALSE to avoid recollapsing.

force

logical, default TRUE If TRUE, reload library files even if matching named variables are found in environment used by experiment (see envExp) A simple way to make sure correct libraries are always loaded. FALSE is faster if data is loaded correctly already.

library.names

character vector, names of libraries, default: name\_decider(df, naming)

284 r,covRle-method

**BPPARAM** 

how many cores/threads to use? default: bpparam(). To see number of threads used, do bpparam()\$workers. You can also add a time remaining bar, for a more detailed pipeline.

### Value

a data.table of the count info

r

strandMode covRle

# Description

strandMode covRle

# Usage

r(x)

## **Arguments**

Х

a covRle object

#### Value

the forward RleList

r,covRle-method

strandMode covRle

# Description

strandMode covRle

## Usage

```
## S4 method for signature 'covRle' r(x)
```

# Arguments

Χ

a covRle object

## Value

the forward RleList

rankOrder 285

rankOrder

ORF rank in transcripts

## **Description**

Creates an ordering of ORFs per transcript, so that ORF with the most upstream start codon is 1, second most upstream start codon is 2, etc. Must input a grl made from ORFik, txNames\_2 -> 2.

### Usage

```
rankOrder(grl)
```

### **Arguments**

grl

a GRangesList object with ORFs

### Value

a numeric vector of integers

### References

```
doi: 10.1074/jbc.R116.733899
```

### See Also

```
Other features: computeFeatures(), computeFeaturesCage(), countOverlapsW(), disengagementScore(), distToCds(), distToTSS(), entropy(), floss(), fpkm(), fpkm_calc(), fractionLength(), initiationScore(), insideOutsideORF(), isInFrame(), isOverlapping(), kozakSequenceScore(), orfScore(), ribosomeReleaseScore(), ribosomeStallingScore(), startRegion(), startRegionCoverage(), stopRegion(), subsetCoverage(), translationalEff()
```

## **Examples**

286 read.experiment

read.experiment

Read ORFik experiment

## **Description**

Read in runs / samples from an experiment as a single R object. To read an ORFik experiment, you must of course make one first. See create.experiment The file must be csv and be a valid ORFik experiment

# Usage

```
read.experiment(
   file,
   in.dir = ORFik::config()["exp"],
   validate = TRUE,
   output.env = .GlobalEnv
)
```

## **Arguments**

file	relative path to a ORFik experiment. That is a .csv file following ORFik experiment style ("," as seperator). , or a template data.frame from create.experiment. Can also be full path to file, then in.dir argument is ignored.
in.dir	Directory to load experiment csv file from, default: ORFik::config()["exp"], which has default "~/Bio_data/ORFik_experiments/" Set to NULL if you don't want to save it to disc. Does not apply if file argument is not a path (can also be a data.frame). Also does not apply if file argument was given as full path.
validate	logical, default TRUE. Abort if any library files does not exist. Do not set this to FALSE, unless you know what you are doing!
output.env	an environment, default .GlobalEnv. Which environment should ORFik output libraries to (if this is done), can be updated later with envExp(df) <- new.env().

### Value

```
an ORFik experiment
```

## See Also

```
Other ORFik_experiment: ORFik.template.experiment(), ORFik.template.experiment.zf(), bamVarName(), create.experiment(), experiment-class, filepath(), libraryTypes(), organism, experiment-methoutputLibs(), save.experiment(), validateExperiments()
```

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### **Examples**

```
# From file
## Not run:
# Read from file
df <- read.experiment(filepath) # <- valid ORFik .csv file

## End(Not run)
## Read from (create.experiment() template)
df <- ORFik.template.experiment()

## To save it, do:
# save.experiment(df, file = "path/to/save/experiment")
## You can then do:
# read.experiment("path/to/save/experiment")
# or (identical):
# read.experiment("experiment", in.dir = "path/to/save/")</pre>
```

readBam

Custom bam reader

### Description

Read in Bam file from either single end or paired end. Safer combined version of readGalignments and readGalignmentPairs that takes care of some common errors.

If QNAMES of the aligned reads are from collapsed fasta files (if the names are formated from collapsing in either (ORFik, ribotoolkit or fastx)), the bam file will contain a meta column called "score" with the counts of duplicates per read. Only works for single end reads, as perfect duplication events for paired end is more rare and therefor not supported!.

### Usage

```
readBam(path, chrStyle = NULL, param = NULL, strandMode = 0)
```

### **Arguments**

path

a character / data.table with path to .bam file. There are 3 input file possibilities.

- single end : a character path (length 1)
- paired end (1 file): Either a character path (length of 2), where path[2] is "paired-end", or a data.table with 2 columns, forward = path & reverse = "paired-end"
- paired end (2 files): Either a character path (length of 2), where path[2] is path to R2, or a data.table with 2 columns, forward = path to R1 & reverse = path to R2. (This one is not used often)

chrStyle

a GRanges object, TxDb, FaFile, , a seqlevelsStyle or Seqinfo. (Default: NULL) to get seqlevelsStyle from. In addition if it is a Seqinfo object, seqinfo will be updated. Example of seqlevelsStyle update: Is chromosome 1 called

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chr1 or 1, is mitocondrial chromosome called MT or chrM etc. Will use 1st seqlevel-style if more are present. Like: c("NCBI", "UCSC") -> pick "NCBI"

param

NULL or a ScanBamParam object. Like for scanBam, this influences what fields and which records are imported. However, note that the fields specified thru this ScanBamParam object will be loaded *in addition* to any field required for generating the returned object (GAlignments, GAlignmentPairs, or GappedReads object), but only the fields requested by the user will actually be kept as metadata columns of the object.

By default (i.e. param=NULL or param=ScanBamParam()), no additional field is loaded. The flag used is scanBamFlag(isUnmappedQuery=FALSE) for readGAlignments, readGAlignmentsList, and readGappedReads. (i.e. only records corresponding to mapped reads are loaded), and scanBamFlag(isUnmappedQuery=FALSE, isPaired=TRUE, hasUnmappedMate=FALSE) for readGAlignmentPairs (i.e. only records corresponding to paired-end reads with both ends mapped are loaded).

strandMode

numeric, default 0. Only used for paired end bam files. One of (0: strand = \*, 1: first read of pair is +, 2: first read of pair is -). See ?strandMode. Note: Sets default to 0 instead of 1, as readGAlignmentPairs uses 1. This is to guarantee hits, but will also make mismatches of overlapping transcripts in opposite directions.

#### **Details**

In the future will use a faster .bam loader for big .bam files in R.

#### Value

a GAlignments or GAlignmentPairs object of bam file

#### See Also

Other utils: bedToGR(), convertToOneBasedRanges(), export.bed12(), export.bigWig(), export.fstwig(), export.wiggle(), fimport(), findFa(), fread.bed(), optimizeReads(), readBigWig(), readWig()

### **Examples**

```
bam_file <- system.file("extdata/Danio_rerio_sample", "ribo-seq.bam", package = "ORFik")
readBam(bam_file, "UCSC")</pre>
```

readBigWig

Custom bigWig reader

#### **Description**

Given 2 bigWig files (.bw, .bigWig), first is forward second is reverse. Merge them and return as GRanges object. If they contain name reverse and forward, first and second order does not matter, it will search for forward and reverse.

readLengthTable 289

### Usage

```
readBigWig(path, chrStyle = NULL, as = "GRanges")
```

## Arguments

path a character path to two .bigWig files, or a data.table with 2 columns, (forward,

filepath) and reverse, only 1 row.

chrStyle a GRanges object, TxDb, FaFile, , a seqlevelsStyle or Seqinfo. (Default:

NULL) to get seqlevelsStyle from. In addition if it is a Seqinfo object, seqinfo will be updated. Example of seqlevelsStyle update: Is chromosome 1 called chr1 or 1, is mitocondrial chromosome called MT or chrM etc. Will use 1st seqlevel-style if more are present. Like: c("NCBI", "UCSC") -> pick "NCBI"

as Specifies the class of the return object. Default is GRanges, which has one range

per range in the file, and a score column holding the value for each range. For NumericList, one numeric vector is returned for each range in the selection argument. For RleList, there is one Rle per sequence, and that Rle spans the

entire sequence.

#### Value

a GRanges object of the file/s

#### See Also

```
Other utils: bedToGR(), convertToOneBasedRanges(), export.bed12(), export.bigWig(), export.fstwig(), export.wiggle(), fimport(), findFa(), fread.bed(), optimizeReads(), readBam(), readWig()
```

readLengthTable

Make table of readlengths

### Description

Summarizing all libraries in experiment, make a table of proportion of read lengths.

## Usage

```
readLengthTable(
   df,
   output.dir = NULL,
   type = "ofst",
   force = TRUE,
   library.names = bamVarName(df),
   BPPARAM = bpparam()
)
```

290 readWidths

#### **Arguments**

df an ORFik experiment

output.dir NULL or character path, default: NULL, plot not saved to disc. If defined saves

plot to that directory with the name "./readLengths.csv".

type character, default: "ofst". Type of library: either "default", usually bam format

(the one you gave to experiment), "pshifted" pshifted reads, "ofst", "bed", "bedo"

optimized bed, or "wig"

force logical, default TRUE If TRUE, reload library files even if matching named

variables are found in environment used by experiment (see envExp) A simple way to make sure correct libraries are always loaded. FALSE is faster if data is

loaded correctly already.

library.names character vector, names of libraries, default: name\_decider(df, naming)

BPPARAM a core param, default: single thread: BiocParallel::SerialParam(). Set to

BiocParallel::bpparam() to use multicore. Be aware, this uses a lot of extra

ram (40GB+) for larger human samples!

#### Value

a data.table object of the the read length data with columns: c("sample", "sample\_id", "read length", "counts", "counts\_per\_sample", "perc\_of\_counts\_per\_sample")

readWidths

Get read widths

### **Description**

Input any reads, e.g. ribo-seq object and get width of reads, this is to avoid confusion between width, qwidth and meta column containing original read width.

### Usage

```
readWidths(reads, after.softclips = TRUE, along.reference = FALSE)
```

#### **Arguments**

reads a GRanges, GAlignment, GAlignmentPairs or covRleList object.

after.softclips

logical (TRUE), include softclips in width. Does not apply if along reference is

TRUE.

along.reference

logical (FALSE), example: The cigar "26MI2" is by default width 28, but if along.reference is TRUE, it will be 26. The length of the read along the reference. Also "1D20M" will be 21 if by along.reference is TRUE. Intronic regions

(cigar: N) will be removed. So: "1M200N19M" is 20, not 220.

readWig 291

#### **Details**

If input is p-shifted and GRanges, the "\$size" or "\$score" colum" must exist, and the column must contain the original read widths. In ORFik "\$size" have higher priority than "\$score" for defining length. ORFik P-shifting creates a \$size column, other softwares like shoelaces creates a score column.

Remember to think about how you define length. Like the question: is a Illumina error mismatch sufficient to reduce size of read and how do you know what is biological variance and what are Illumina errors?

#### Value

an integer vector of widths

### **Examples**

```
gr <- GRanges("chr1", 1)
readWidths(gr)

# GAlignment with hit (1M) and soft clipped base (1S)
ga <- GAlignments(seqnames = "1", pos = as.integer(1), cigar = "1M1S",
    strand = factor("+", levels = c("+", "-", "*")))
readWidths(ga) # Without soft-clip bases

readWidths(ga, after.softclips = FALSE) # With soft-clip bases</pre>
```

readWig

Custom wig reader

# Description

Given 2 wig files, first is forward second is reverse. Merge them and return as GRanges object. If they contain name reverse and forward, first and second order does not matter, it will search for forward and reverse.

#### Usage

```
readWig(path, chrStyle = NULL)
```

## **Arguments**

path

a character path to two .wig files, or a data.table with 2 columns, (forward, filepath) and reverse, only 1 row.

chrStyle

a GRanges object, TxDb, FaFile, , a seqlevelsStyle or Seqinfo. (Default: NULL) to get seqlevelsStyle from. In addition if it is a Seqinfo object, seqinfo will be updated. Example of seqlevelsStyle update: Is chromosome 1 called chr1 or 1, is mitocondrial chromosome called MT or chrM etc. Will use 1st seqlevel-style if more are present. Like: c("NCBI", "UCSC") -> pick "NCBI"

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### Value

a GRanges object of the file/s

#### See Also

Other utils: bedToGR(), convertToOneBasedRanges(), export.bed12(), export.bigWig(), export.fstwig(), export.wiggle(), fimport(), findFa(), fread.bed(), optimizeReads(), readBam(), readBigWig()

reassignTSSbyCage

Reassign all Transcript Start Sites (TSS)

### **Description**

Given a GRangesList of 5' UTRs or transcripts, reassign the start sites using max peaks from CageSeq data. A max peak is defined as new TSS if it is within boundary of 5' leader range, specified by 'extension' in bp. A max peak must also be higher than minimum CageSeq peak cutoff specified in 'filterValue'. The new TSS will then be the positioned where the cage read (with highest read count in the interval). If removeUnused is TRUE, leaders without cage hits, will be removed, if FALSE the original TSS will be used.

### Usage

```
reassignTSSbyCage(
  fiveUTRs,
  cage,
  extension = 1000,
  filterValue = 1,
  restrictUpstreamToTx = FALSE,
  removeUnused = FALSE,
  preCleanup = TRUE,
  cageMcol = FALSE
)
```

set it to 0.

### **Arguments**

fiveUTRs	(GRangesList) The 5' leaders or full transcript sequences
cage	Either a filePath for the CageSeq file as .bed .bam or .wig, with possible compressions (".gzip", ".gz", ".bgz"), or already loaded CageSeq peak data as GRanges or GAlignment. NOTE: If it is a .bam file, it will add a score column by running: convertToOneBasedRanges(cage, method = "5prime", addScoreColumn = TRUE) The score column is then number of replicates of read, if score column is something else, like read length, set the score column to NULL first.
extension	The maximum number of basses upstream of the TSS to search for CageSeq peak.
filterValue	The minimum number of reads on cage position, for it to be counted as possible new tss. (represented in score column in CageSeq data) If you already filtered,

reassignTSSbyCage 293

restrictUpstreamToTx

a logical (FALSE). If TRUE: restrict leaders to not extend closer than 5 bases

from closest upstream leader, set this to TRUE.

removeUnused logical (FALSE), if False: (standard is to set them to original annotation), If

TRUE: remove leaders that did not have any cage support.

preCleanup logical (TRUE), if TRUE, remove all reads in region (-5:-1, 1:5) of all original

tss in leaders. This is to keep original TSS if it is only +/- 5 bases from the

original.

cageMcol a logical (FALSE), if TRUE, add a meta column to the returned object with the

raw CAGE counts in support for new TSS.

### **Details**

Note: If you used CAGEr, you will get reads of a probability region, with always score of 1. Remember then to set filterValue to 0. And you should use the 5' end of the read as input, use: ORFik:::convertToOneBasedRanges(cage) NOTE on filtervalue: To get high quality TSS, set filtervalue to median count of reads overlapping per leader. This will make you discard a lot of new TSS positions though. I usually use 10 as a good standard.

TIP: do summary(countOverlaps(fiveUTRs, cage)) so you can find a good cutoff value for noise.

#### Value

a GRangesList of newly assigned TSS for fiveUTRs, using CageSeq data.

#### See Also

```
Other CAGE: assignTSSByCage(), reassignTxDbByCage()
```

### **Examples**

```
# example 5' leader, notice exon_rank column
fiveUTRs <- GenomicRanges::GRangesList(</pre>
 GenomicRanges::GRanges(seqnames = "chr1",
                          ranges = IRanges::IRanges(1000, 2000),
                          strand = "+",
                          exon_rank = 1)
names(fiveUTRs) <- "tx1"</pre>
# make fake CAGE data from promoter of 5' leaders, notice score column
cage <- GenomicRanges::GRanges(</pre>
 seqnames = "1",
 ranges = IRanges::IRanges(500, width = 1),
 strand = "+",
 score = 10) # <- Number of tags (reads) per position</pre>
# notice also that seqnames use different naming, this is fixed by ORFik
# finally reassign TSS for fiveUTRs
reassignTSSbyCage(fiveUTRs, cage)
# See vignette for example using gtf file and real CAGE data.
```

reassignTxDbByCage

Input a txdb and reassign the TSS for each transcript by CAGE

#### **Description**

Given a TxDb object, reassign the start site per transcript using max peaks from CageSeq data. A max peak is defined as new TSS if it is within boundary of 5' leader range, specified by 'extension' in bp. A max peak must also be higher than minimum CageSeq peak cutoff specified in 'filter-Value'. The new TSS will then be the positioned where the cage read (with highest read count in the interval).

# Usage

```
reassignTxDbByCage(
  txdb,
  cage,
  extension = 1000,
  filterValue = 1,
  restrictUpstreamToTx = FALSE,
  removeUnused = FALSE,
  preCleanup = TRUE
)
```

#### **Arguments**

txdb a TxDb file, a path to one of: (.gtf, .gff2, .gff2, .db or .sqlite)	ite) or an ORFik
--	------------------

experiment

cage Either a filePath for the CageSeq file as .bed .bam or .wig, with possible com-

pressions (".gzip", ".gz", ".bgz"), or already loaded CageSeq peak data as GRanges or GAlignment. NOTE: If it is a .bam file, it will add a score column by running: convertToOneBasedRanges(cage, method = "5prime", addScoreColumn = TRUE) The score column is then number of replicates of read, if score column

is something else, like read length, set the score column to NULL first.

extension The maximum number of basses upstream of the TSS to search for CageSeq

peak.

filterValue The minimum number of reads on cage position, for it to be counted as possible

new tss. (represented in score column in CageSeq data) If you already filtered,

set it to 0.

restrictUpstreamToTx

a logical (FALSE). If TRUE: restrict leaders to not extend closer than 5 bases

from closest upstream leader, set this to TRUE.

removeUnused logical (FALSE), if False: (standard is to set them to original annotation), If

TRUE: remove leaders that did not have any cage support.

preCleanup logical (TRUE), if TRUE, remove all reads in region (-5:-1, 1:5) of all original

tss in leaders. This is to keep original TSS if it is only +/- 5 bases from the

original.

reduceKeepAttr 295

### **Details**

Note: If you used CAGEr, you will get reads of a probability region, with always score of 1. Remember then to set filterValue to 0. And you should use the 5' end of the read as input, use: ORFik:::convertToOneBasedRanges(cage)

### Value

a TxDb obect of reassigned transcripts

### See Also

```
Other CAGE: assignTSSByCage(), reassignTSSbyCage()
```

## **Examples**

```
## Not run:
library(GenomicFeatures)
# Get the gtf txdb file
txdbFile <- system.file("extdata", "hg19_knownGene_sample.sqlite",
package = "GenomicFeatures")
cagePath <- system.file("extdata", "cage-seq-heart.bed.bgz",
package = "ORFik")
reassignTxDbByCage(txdbFile, cagePath)
## End(Not run)</pre>
```

reduceKeepAttr

Reduce GRanges / GRangesList

## **Description**

Reduce away all GRanges elements with 0-width.

# Usage

```
reduceKeepAttr(
  grl,
  keep.names = FALSE,
  drop.empty.ranges = FALSE,
  min.gapwidth = 1L,
  with.revmap = FALSE,
  with.inframe.attrib = FALSE,
  ignore.strand = FALSE,
  min.strand.decreasing = TRUE
)
```

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#### **Arguments**

```
a GRangesList or GRanges object
grl
                 (FALSE) keep the names and meta columns of the GRangesList
keep.names
drop.empty.ranges
                  (FALSE) if a group is empty (width 0), delete it.
                 (1L) how long gap can it be between two ranges, to merge them.
min.gapwidth
with.revmap
                  (FALSE) return info on which mapped to which
with.inframe.attrib
                 (FALSE) For internal use.
ignore.strand
                 (FALSE), can different strands be reduced together.
min.strand.decreasing
                  (TRUE), if GRangesList, return minus strand group ranges in decreasing order
                  (1-5, 30-50) \rightarrow (30-50, 1-5)
```

### **Details**

Extends function reduce by trying to keep names and meta columns, if it is a GRangesList. It also does not lose sorting for GRangesList, since original reduce sorts all by ascending position. If keep.names == FALSE, it's just the normal GenomicRanges::reduce with sorting negative strands descending for GRangesList.

#### Value

A reduced GRangesList

### See Also

```
Other ExtendGenomicRanges: asTX(), coveragePerTiling(), extendLeaders(), extendTrailers(), tile1(), txSeqsFromFa(), windowPerGroup()
```

## **Examples**

regionPerReadLength 297

regionPerReadLength

Find proportion of reads per position per read length in region

### Description

This is defined as: Given some transcript region (like CDS), get coverage per position. By default only returns positions that have hits, set drop.zero.dt to FALSE to get all 0 positions.

### Usage

```
regionPerReadLength(
  grl,
  reads,
  acceptedLengths = NULL,
  withFrames = TRUE,
  scoring = "transcriptNormalized",
  weight = "score",
  exclude.zero.cov.grl = TRUE,
  drop.zero.dt = TRUE,
  BPPARAM = bpparam()
)
```

#### **Arguments**

grl a GRangesList object with usually either leaders, cds', 3' utrs or ORFs

reads a GAlignments, GRanges, or precomputed coverage as covRleList (where

names of covRle objects are readlengths) of RiboSeq, RnaSeq etc.

Weigths for scoring is default the 'score' column in 'reads'. Can also be random access paths to bigWig or fstwig file. Do not use random access for more than a

few genes, then loading the entire files is usually better.

acceptedLengths

an integer vector (NULL), the read lengths accepted. Default NULL, means all

lengths accepted.

withFrames logical TRUE, add ORF frame (frame 0, 1, 2), starting on first position of every

grl.

scoring a character (transcriptNormalized), which meta coverage scoring? one of (zs-

core, transcriptNormalized, mean, median, sum, sumLength, fracPos), see ?coverageScorings for more info. Use to decide a scoring of hits per position for metacoverage etc. Set to NULL if you do not want meta coverage, but instead

want per gene per position raw counts.

weight (default: 'score'), if defined a character name of valid meta column in subject.

GRanges("chr1", 1, "+", score = 5), would mean score column tells that this alignment region was found 5 times. Formats which loads a score column like this: Bigwig, wig, ORFik ofst, collapsed bam, bedoc and .bedo. As do CAGEr CAGE files and many other package formats. You can also assign a score col-

umn manually.

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exclude.zero.cov.grl

logical, default TRUE. Do not include ranges that does not have any coverage

(0 reads on them), this makes it faster to run.

drop.zero.dt logical, default TRUE. If TRUE and as.data.table is TRUE, remove all 0 count

positions. This greatly speeds up and most importantly, greatly reduces memory usage. Will not change any plots, unless 0 count positions are used in some

sense.

BPPARAM how many cores/threads to use? default: bpparam()

#### Value

a data.table with lengths by coverage.

### See Also

Other coverage: coverageScorings(), metaWindow(), scaledWindowPositions(), windowPerReadLength()

## **Examples**

```
# Raw counts per gene per position
cds <- GRangesList(tx1 = GRanges("1", 100:129, "+"))
reads <- GRanges("1", seq(79,129, 3), "+")
reads$size <- 28 # <- Set read length of reads
regionPerReadLength(cds, reads, scoring = NULL)
## Sum up reads in each frame per read length per gene
regionPerReadLength(cds, reads, scoring = "frameSumPerLG")</pre>
```

remakeTxdbExonIds

Get new exon ids after update of txdb

## Description

Get new exon ids after update of txdb

### Usage

```
remakeTxdbExonIds(txList)
```

#### **Arguments**

```
txList a list, call of as.list(txdb)
```

### Value

a new valid ordered list of exon ids (integer)

remove.experiments 299

remove.experiments

Remove ORFik experiment libraries load in R

# Description

Variable names defined by df, in envir defined

## Usage

```
remove.experiments(df, envir = envExp(df))
```

## **Arguments**

df an ORFik experiment

envir environment to save to, default envExp(df), which defaults to .GlobalEnv, but

can be set with envExp(df) <- new.env() etc.

### Value

NULL (objects removed from envir specified)

# **Examples**

```
df <- ORFik.template.experiment()
# Output to .GlobalEnv with:
# outputLibs(df)
# Then remove them with:
# remove.experiments(df)</pre>
```

remove.file\_ext

Remove file extension of path

# Description

Allows removal of compression

## Usage

```
remove.file_ext(path, basename = FALSE)
```

## Arguments

path character path (allows multiple paths)

basename relative path (TRUE) or full path (FALSE)? (default: FALSE)

### Value

character path without file extension

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removeMetaCols

Removes meta columns

# Description

Removes meta columns

# Usage

```
removeMetaCols(grl)
```

### **Arguments**

grl

a GRangesList or GRanges object

### Value

same type and structure as input without meta columns

removeORFsWithinCDS

Remove ORFs that are within cds

# Description

Remove ORFs that are within cds

# Usage

```
removeORFsWithinCDS(grl, cds)
```

## **Arguments**

grl

(GRangesList), the ORFs to filter

cds

(GRangesList), the coding sequences (main ORFs on transcripts), to filter against.

## Value

(GRangesList) of filtered uORFs

## See Also

Other uorfs: addCdsOnLeaderEnds(), filterUORFs(), removeORFsWithSameStartAsCDS(), removeORFsWithSameStopAremoveORFsWithStartInsideCDS(), uORFSearchSpace()

removeORFsWithSameStartAsCDS

Remove ORFs that have same start site as the CDS

# Description

Remove ORFs that have same start site as the CDS

#### Usage

```
removeORFsWithSameStartAsCDS(grl, cds)
```

### **Arguments**

grl (GRangesList), the ORFs to filter

cds (GRangesList), the coding sequences (main ORFs on transcripts), to filter against.

#### Value

(GRangesList) of filtered uORFs

#### See Also

Other uorfs: addCdsOnLeaderEnds(), filterUORFs(), removeORFsWithSameStopAsCDS(), removeORFsWithStartInsid removeORFsWithinCDS(), uORFSearchSpace()

removeORFsWithSameStopAsCDS

Remove ORFs that have same stop site as the CDS

## **Description**

Remove ORFs that have same stop site as the CDS

# Usage

```
removeORFsWithSameStopAsCDS(grl, cds)
```

### **Arguments**

grl (GRangesList), the ORFs to filter

cds (GRangesList), the coding sequences (main ORFs on transcripts), to filter against.

### Value

(GRangesList) of filtered uORFs

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### See Also

Other u or fs: add Cds On Leader Ends (), filter UORFs (), remove ORFs With Same Start AsCDS (), remove ORFs With Start Insiremove ORFs Within CDS (), u ORFS earch Space ()

removeORFsWithStartInsideCDS

Remove ORFs that have start site within the CDS

# Description

Remove ORFs that have start site within the CDS

# Usage

```
removeORFsWithStartInsideCDS(grl, cds)
```

## **Arguments**

grl (GRangesList), the ORFs to filter

cds (GRangesList), the coding sequences (main ORFs on transcripts), to filter against.

### Value

(GRangesList) of filtered uORFs

## See Also

Other uorfs: addCdsOnLeaderEnds(), filterUORFs(), removeORFsWithSameStartAsCDS(), removeORFsWithSameStopAremoveORFsWithinCDS(), uORFSearchSpace()

removeTxdbExons

Remove exons in txList that are not in fiveUTRs

# Description

Remove exons in txList that are not in fiveUTRs

## Usage

```
removeTxdbExons(txList, fiveUTRs)
```

## Arguments

txList a list, call of as.list(txdb) fiveUTRs a GRangesList of 5' leaders

removeTxdbTranscripts

### Value

a list, modified call of as.list(txdb)

removeTxdbTranscripts Remove specific transcripts in txdb List

## **Description**

Remove all transcripts, except the ones in fiveUTRs.

### Usage

```
removeTxdbTranscripts(txList, fiveUTRs)
```

## Arguments

txList a list, call of as.list(txdb) fiveUTRs a GRangesList of 5' leaders

## Value

a txList

rename.SRA.files

Rename SRA files from metadata

# Description

Rename SRA files from metadata

### Usage

```
rename.SRA.files(files, new_names)
```

## **Arguments**

files a character vector, with full path to all the files

new\_names a character vector of new names or a data.table with metadata to use to rename

(usually from SRA metadata). Priority of renaming from the metadata is to check for unique names in the LibraryName column, then the sample\_title column if no valid names in LibraryName. If found and still duplicates, will add "\_rep1", "\_rep2" to make them unique. Paired end data will get a extension of \_p1 and \_p2. If no valid names, will not rename, that is keep the SRR numbers,

you then can manually rename files to something more meaningful.

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### Value

a character vector of new file names

### See Also

Other sra: browseSRA(), download.SRA(), download.SRA.metadata(), download.ebi(), get\_bioproject\_candidates install.sratoolkit()

repNames

Get replicate name variants

# Description

Used to standardize nomeclature for experiments.

Example: 1 is main naming, but a variant is rep1 rep1 will then be renamed to 1

### Usage

repNames()

### Value

a data.table with 2 columns, the main name, and all name variants of the main name in second column as a list.

#### See Also

Other experiment\_naming: batchNames(), cellLineNames(), cellTypeNames(), conditionNames(), fractionNames(), inhibitorNames(), libNames(), mainNames(), stageNames(), tissueNames()

resFolder

Get ORFik experiment main output directory

## **Description**

Get ORFik experiment main output directory

## Usage

resFolder(x)

## **Arguments**

Х

an ORFik experiment

### Value

a character path

resFolder, experiment-method

Get ORFik experiment main output directory

### **Description**

Get ORFik experiment main output directory

# Usage

```
## S4 method for signature 'experiment'
resFolder(x)
```

#### **Arguments**

Х

an ORFik experiment

#### Value

a character path

restrictTSSByUpstreamLeader

Restrict extension of 5' UTRs to closest upstream leader end

## **Description**

Basicly this function restricts all startSites, to the upstream GRangesList objects end. Usually leaders, for CAGE. Example: leader1: start on 10, leader2: stop on 8, extend leader1 to 5 -> this function will resize leader1 to 9, to be outside leader2, so that CAGE reads can not wrongly overlap.

### Usage

```
restrictTSSByUpstreamLeader(fiveUTRs, shiftedfiveUTRs)
```

## **Arguments**

```
fiveUTRs The 5' leader sequences as GRangesList shiftedfiveUTRs
```

The 5' leader sequences as GRangesList shifted by CAGE

#### Value

GRangesList object of restricted fiveUTRs

revElementsF

Reverse elements within list

# Description

A faster version of S4Vectors::revElements

# Usage

```
revElementsF(x)
```

# Arguments

Χ

RleList

## Value

a RleList (reversed inside list elements)

 ${\tt reverseMinusStrandPerGroup}$ 

Reverse minus strand

# Description

Reverse minus strand per group in a GRangesList Only reverse if minus strand is in increasing order

# Usage

```
reverseMinusStrandPerGroup(grl, onlyIfIncreasing = TRUE)
```

## **Arguments**

```
 \begin{array}{ll} & \text{grl} & \text{a GRangesList} \\ & \text{onlyIfIncreasing} \\ & & \text{logical, default (TRUE), only reverse if decreasing} \end{array}
```

## Value

```
a \; \mathsf{GRangesList}
```

riboORFs 307

riboORFs	Load Predicted translons	

## **Description**

Load Predicted translons

### Usage

```
riboORFs(df, type = "table", folder = riboORFsFolder(df))
```

### **Arguments**

df ORFik experiment

type default "table", alternatives: c("table", "ranges\_candidates", "ranges\_predictions",

"predictions")

folder base folder to check for computed results, default: riboORFsFolder(df)

#### Value

a data.table, GRangesList or list of logical vector depending on input

## **Examples**

```
df <- ORFik.template.experiment()
df <- df[df$libtype == "RFP",][c(1,2),]
# riboORFs(df) # Works when you have run prediction</pre>
```

riboORFsFolder

Define folder for prediction output

## Description

Define folder for prediction output

## Usage

```
riboORFsFolder(df, parrent_dir = resFolder(df))
```

### **Arguments**

df ORFik experiment

parrent\_dir Parrent directory of computed study results, default: resFolder(df)

308 RiboQC.plot

### Value

```
a file path (full path)
```

### **Examples**

```
df <- ORFik.template.experiment()
df <- df[df$libtype == "RFP",][c(1,2),]
riboORFsFolder(df)
riboORFsFolder(df, tempdir())</pre>
```

RiboQC.plot

Quality control for pshifted Ribo-seq data

# Description

Combines several statistics from the pshifted reads into a plot:

- -1 Coding frame distribution per read length
- -2 Alignment statistics
- -3 Biotype of non-exonic pshifted reads
- -4 mRNA localization of pshifted reads

## Usage

```
RiboQC.plot(
   df,
   output.dir = QCfolder(df),
   width = 6.6,
   height = 4.5,
   plot.ext = ".pdf",
   type = "pshifted",
   weight = "score",
   bar.position = "dodge",
   as_gg_list = FALSE,
   BPPARAM = BiocParallel::SerialParam(progressbar = TRUE)
)
```

## **Arguments**

```
df an ORFik experiment

output.dir NULL or character path, default: NULL, plot not saved to disc. If defined saves plot to that directory with the name "/STATS_plot.pdf".

width width of plot, default 6.6 (in inches)

height height of plot, default 4.5 (in inches)

plot.ext character, default: ".pdf". Alternatives: ".png" or ".jpg".
```

ribosomeReleaseScore 309

type type of library loaded, default pshifted, warning if not pshifted might crash if

too many read lengths!

weight which column in reads describe duplicates, default "score".

bar.position character, default "dodge". Should Ribo-seq frames per read length be posi-

tioned as "dodge" or "stack" (on top of each other).

as\_gg\_list logical, default FALSE. Return as a list of ggplot objects instead of as a grob.

Gives you the ability to modify plots more directly.

BPPARAM how many cores/threads to use? default: bpparam(). To see number of threads

used, do bpparam() \$workers. You can also add a time remaining bar, for a

more detailed pipeline.

#### Value

the plot object, a grob of ggplot objects of the the data

# Examples

```
df <- ORFik.template.experiment()
df <- df[9,] #lets only p-shift RFP sample at index 9
#shiftFootprintsByExperiment(df)
#RiboQC.plot(df, tempdir())</pre>
```

ribosomeReleaseScore Ribosome Release Score (RRS)

## Description

Ribosome Release Score is defined as

```
(RPFs over ORF)/(RPFs over 3' utrs)
```

and additionally normalized by lengths. If RNA is added as argument, it will normalize by RNA counts to justify location of 3' utrs. It can be understood as a ribosome stalling feature. A pseudocount of one was added to both the ORF and downstream sums.

### Usage

```
ribosomeReleaseScore(
  grl,
  RFP,
  GtfOrThreeUtrs,
  RNA = NULL,
  weight.RFP = 1L,
  weight.RNA = 1L,
  overlapGrl = NULL
)
```

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### **Arguments**

grl	a GRangesList object with usually either leaders, cds', 3' utrs or ORFs.
RFP	RiboSeq reads as GAlignments, GRanges or GRangesList object
GtfOrThreeUtrs	if Gtf: a TxDb object of a gtf file transcripts is called from: 'threeUTRsByTranscript(Gtf, use.names = TRUE)', if object is GRangesList, it is presumed to be the 3' utrs
RNA	RnaSeq reads as GAlignments, GRanges or GRangesList object
weight.RFP	a vector (default: 1L). Can also be character name of column in RFP. As in translationalEff(weight = "score") for: GRanges("chr1", 1, "+", score = 5), would mean score column tells that this alignment region was found 5 times.
weight.RNA	Same as weightRFP but for RNA weights. (default: 1L)
overlapGrl	an integer, (default: NULL), if defined must be countOverlaps(grl, RFP), added for speed if you already have it

### Value

a named vector of numeric values of scores, NA means that no 3' utr was found for that transcript.

#### References

```
doi: 10.1016/j.cell.2013.06.009
```

#### See Also

```
Other features: computeFeatures(), computeFeaturesCage(), countOverlapsW(), disengagementScore(), distToCds(), distToTSS(), entropy(), floss(), fpkm(), fpkm_calc(), fractionLength(), initiationScore(), insideOutsideORF(), isInFrame(), isOverlapping(), kozakSequenceScore(), orfScore(), rankOrder(), ribosomeStallingScore(), startRegion(), startRegionCoverage(), stopRegion(), subsetCoverage(), translationalEff()
```

# **Examples**

ribosomeStallingScore 311

```
ribosomeStallingScore Ribosome Stalling Score (RSS)
```

# Description

```
Is defined as
```

```
(RPFs over ORF stop sites)/(RPFs over ORFs)
```

and normalized by lengths A pseudo-count of one was added to both the ORF and downstream sums.

### Usage

```
ribosomeStallingScore(grl, RFP, weight = 1L, overlapGrl = NULL)
```

### **Arguments**

grl	a GRangesList obj	ect with usually	v either leaders.	cds', 3'	utrs or ORFs.
0			,	• • • •	or or or or

RFP RiboSeq reads as GAlignments, GRanges or GRangesList object

weight a numeric/integer vector or metacolumn name. (default: 1L, no differential

weighting). If weight is name of defined meta column in reads object, it gives the number of times a read was found at that position. GRanges("chr1", 1, "+", score = 5), would mean "score" column tells that this alignment region was found 5 times. if 1L it means each read is weighted equal as 1, this is what among others countOverlaps() presumes, if single number (!= 1), it repeats for all ranges, if vector with length > 1, it must be equal size of the reads object.

overlapGrl an integer, (default: NULL), if defined must be countOverlaps(grl, RFP), added

for speed if you already have it

#### Value

a named vector of numeric values of RSS scores

### References

```
doi: 10.1016/j.cels.2017.08.004
```

#### See Also

```
Other features: computeFeatures(), computeFeaturesCage(), countOverlapsW(), disengagementScore(), distToCds(), distToTSS(), entropy(), floss(), fpkm(), fpkm_calc(), fractionLength(), initiationScore(), insideOutsideORF(), isInFrame(), isOverlapping(), kozakSequenceScore(), orfScore(), rankOrder(), ribosomeReleaseScore(), startRegion(), startRegionCoverage(), stopRegion(), subsetCoverage(), translationalEff()
```

ribo\_fft

### **Examples**

ribo\_fft

Get periodogram data per read length

### **Description**

A data.table of periods and amplitudes, great to detect ribosomal read lengths. Uses 5' end of reads to detect periodicity. Works both before and after p-shifting. Plot results with ribo\_fft\_plot.

## Usage

```
ribo_fft(footprints, cds, read_lengths = 26:34, firstN = 150)
```

# Arguments

footprints	Ribosome footprints in either GAlignments or GRanges
cds	a GRangesList of coding sequences. Length must match length of argument mrna, and all must have length > arugment firstN.
read_lengths	integer vector, default: 26:34, which read length to check for. Will exclude all read_lengths that does not exist for footprints.
firstN	(integer) Represents how many bases of the transcripts downstream of start codons to use for initial estimation of the periodicity.

### Value

a data.table with read\_length, amplitude and periods

### **Examples**

```
## Note, this sample data is not intended to be strongly periodic.
## Real data should have a cleaner peak for x = 3 (periodicity)
# Load sample data
df <- ORFik.template.experiment()
# Load annotation
loadRegions(df, "cds", names.keep = filterTranscripts(df))
# Select a riboseq library
df <- df[df$libtype == "RFP", ]
footprints <- fimport(filepath(df[1,], "default"))
fft_dt <-ribo_fft(footprints, cds)
ribo_fft_plot(fft_dt)</pre>
```

ribo\_fft\_plot 313

ribo\_fft\_plot

Get periodogram plot per read length

### **Description**

Get periodogram plot per read length

## Usage

```
ribo_fft_plot(fft_dt, period_window = c(0, 6))
```

### **Arguments**

```
fft_dt a data.table with read_length, amplitude and periods period_window x axis limits, default c(0,6)
```

### Value

a ggplot, geom\_line plot facet by read length.

### **Examples**

```
## Note, this sample data is not intended to be strongly periodic.
## Real data should have a cleaner peak for x = 3 (periodicity)
# Load sample data
df <- ORFik.template.experiment()
# Load annotation
cds <- loadRegion(df, "cds", names.keep = filterTranscripts(df))
# Select a riboseq library
df <- df[df$libtype == "RFP", ]
footprints <- fimport(filepath(df[1,], "default"))
fft_dt <-ribo_fft(footprints, cds)
ribo_fft_plot(fft_dt)</pre>
```

rnaNormalize

Normalize a data.table of coverage by RNA seq per position

# Description

Normalizes per position per gene by this function: (reads at position / min(librarysize, 1) \* number of genes) / fpkm of that gene's RNA-seq

## Usage

```
rnaNormalize(coverage, df, dfr = NULL, tx, normalizeMode = "position")
```

314 runIDs

### **Arguments**

coverage a data.table containing at least columns (count/score, position), it is possible to

have additionals: (genes, fraction, feature)

df an ORFik experiment

dfr an ORFik experiment of RNA-seq to normalize against. Will add RNA nor-

malized to plot name if this is done.

tx a GRangesList of mrna transcripts

normalizeMode a character (default: "position"), how to normalize library against rna library.

Either on "position", normalize by number of genes, sum of reads and RNA seq, on tx "region" or "feature": same as position but RNA is split into the feature groups to normalize. Useful if you have a list of targets and background genes.

#### **Details**

Good way to compare libraries

### Value

a data.table of normalized transcripts by RNA.

runIDs

Get SRR/DRR/ERR run ids from ORFik experiment

## **Description**

Get SRR/DRR/ERR run ids from ORFik experiment

### Usage

runIDs(x)

## **Arguments**

x an ORFik experiment

#### Value

a character vector of runIDs, "" if not existing.

```
runIDs, experiment-method
```

Get SRR/DRR/ERR run ids from ORFik experiment

### **Description**

Get SRR/DRR/ERR run ids from ORFik experiment

## Usage

```
## S4 method for signature 'experiment'
runIDs(x)
```

### **Arguments**

Χ

an ORFik experiment

### Value

a character vector of runIDs, "" if not existing.

save.experiment

Save experiment to disc

# Description

Save experiment to disc

## Usage

```
save.experiment(df, file)
```

## Arguments

df an ORFik experiment file name of file to save df as

# Value

NULL (experiment save only)

### See Also

```
Other ORFik_experiment: ORFik.template.experiment(), ORFik.template.experiment.zf(), bamVarName(), create.experiment(), experiment-class, filepath(), libraryTypes(), organism, experiment-methoutputLibs(), read.experiment(), validateExperiments()
```

316 savePlot

### **Examples**

```
df <- ORFik.template.experiment()
## Save with:
#save.experiment(df, file = "path/to/save/experiment.csv")
## Identical (.csv not needed, can be added):
#save.experiment(df, file = "path/to/save/experiment")</pre>
```

savePlot

Helper function for writing plots to disc

## **Description**

Helper function for writing plots to disc

## Usage

```
savePlot(
  plot,
  output = NULL,
  width = 200,
  height = 150,
  plot.ext = ".pdf",
  dpi = 300,
  limitsize = FALSE
)
```

## **Arguments**

plot the ggplot to save

output character string (NULL), if set, saves the plot as pdf or png to path given. If no

format is given, is save as specified by plot.ext argument.

width width of output in mm height height of output in mm

plot.ext character, default: ".pdf". Alternatives: ".png" or ".jpg".

dpi (300) dpi of plot

limitsize logical, default FALSE. If TRUE, activate ggplot max size restriction.

## Value

a ggplot object of the coverage plot, NULL if output is set, then the plot will only be saved to location.

#### See Also

```
Other\ coverage Plot:\ coverage Heat Map(),\ pSitePlot(),\ window Coverage Plot()
```

scaledWindowPositions 317

scaledWindowPositions Scale (bin) windows to a meta window of given size

#### **Description**

For example scale a coverage table of a all human CDS to width 100

## Usage

```
scaledWindowPositions(
  grl,
  reads,
  scaleTo = 100,
  scoring = "meanPos",
  weight = "score",
  is.sorted = FALSE,
  drop.zero.dt = FALSE)
```

#### Arguments

grl	a GRangesList of 5' utrs, CDS, transcripts, etc.
	CA1:

reads a GAlignments, GRanges, or precomputed coverage as covRle (one for each

strand) of RiboSeq, RnaSeq etc.

Weigths for scoring is default the 'score' column in 'reads'. Can also be random access paths to bigWig or fstwig file. Do not use random access for more than a few genes, then loading the entire files is usually better. File streaming is still in

beta, so use with care!

scaleTo an integer (100), if windows have different size, a meta window can not directly

be created, since a meta window must have equal size for all windows. Rescale all windows to scaleTo. i.e c(1,2,3) -> size 2 -> c(1, mean(2,3)) etc. Can also be

a vector, 1 number per grl group.

scoring a character, one of (zscore, transcriptNormalized, mean, median, sum, log2sum,

 $log10 sum, sumLength, meanPos \ and \ frameSum, periodic, NULL). \ More \ info \ in$ 

details

weight (default: 'score'), if defined a character name of valid meta column in subject.

GRanges("chr1", 1, "+", score = 5), would mean score column tells that this alignment region was found 5 times. Formats which loads a score column like this: Bigwig, wig, ORFik ofst, collapsed bam, bedoc and .bedo. As do CAGEr CAGE files and many other package formats. You can also assign a score col-

umn manually.

is.sorted logical (FALSE), is grl sorted. That is + strand groups in increasing ranges

(1,2,3), and - strand groups in decreasing ranges (3,2,1)

drop.zero.dt logical FALSE, if TRUE and as.data.table is TRUE, remove all 0 count posi-

tions. This greatly speeds up and most importantly, greatly reduces memory usage. Will not change any plots, unless 0 positions are used in some sense.

(mean, median, zscore coverage will only scale differently)

#### **Details**

Nice for making metaplots, the score will be mean of merged positions.

#### Value

A data.table with scored counts (counts) of reads mapped to positions (position) specified in windows along with frame (frame).

#### See Also

Other coverage: coverageScorings(), metaWindow(), regionPerReadLength(), windowPerReadLength()

#### **Examples**

```
library(GenomicRanges)
windows <- GRangesList(GRanges("chr1", IRanges(1, 200), "-"))
x <- GenomicRanges::GRanges(
    seqnames = "chr1",
    ranges = IRanges::IRanges(c(1, 100, 199), c(2, 101, 200)),
    strand = "-")
scaledWindowPositions(windows, x, scaleTo = 100)</pre>
```

scoreSummarizedExperiment

 $Helper\ function\ for\ make Summarized Experiment From Bam$ 

### **Description**

If txdb or gtf path is added, it is a rangedSummerizedExperiment For FPKM values, DESeq2::fpkm(robust = FALSE) is used

### Usage

```
scoreSummarizedExperiment(
  final,
  score = "transcriptNormalized",
  collapse = FALSE
)
```

### **Arguments**

final ranged summarized experiment object
score default: "transcriptNormalized" (row norma

default: "transcriptNormalized" (row normalized raw counts matrix), alternative

is "fpkm", "log2fpkm" or "log10fpkm"

collapse a logical/character (default FALSE), if TRUE all samples within the group SAM-

PLE will be collapsed to one. If "all", all groups will be merged into 1 column called merged\_all. Collapse is defined as rowSum(elements\_per\_group) /

ncol(elements\_per\_group)

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## Value

a DEseq summerizedExperiment object (transcriptNormalized) or matrix (if fpkm input)

```
seqinfo, covRle-method Seqinfo covRle Extracted from forward RleList
```

## **Description**

Seqinfo covRle Extracted from forward RleList

## Usage

```
## S4 method for signature 'covRle'
seqinfo(x)
```

## **Arguments**

x a covRle object

# Value

integer vector with names

```
seqinfo,covRleList-method
```

Seqinfo covRle Extracted from forward RleList

# Description

Seqinfo covRle Extracted from forward RleList

## Usage

```
## S4 method for signature 'covRleList'
seqinfo(x)
```

# Arguments

x a covRle object

## Value

integer vector with names

```
seqinfo, experiment-method
```

Seqinfo ORFik experiment Extracted from fasta genome index

# Description

Seqinfo ORFik experiment Extracted from fasta genome index

# Usage

```
## S4 method for signature 'experiment'
seqinfo(x)
```

# Arguments

Χ

an ORFik experiment

## Value

integer vector with names

```
seqlevels, covRle-method
```

Seqlevels covRle Extracted from forward RleList

# Description

Seqlevels covRle Extracted from forward RleList

## Usage

```
## S4 method for signature 'covRle'
seqlevels(x)
```

# Arguments

Χ

a covRle object

## Value

integer vector with names

```
seqlevels, covRleList-method
```

Seglevels covRleList Extracted from forward RleList

## **Description**

Seqlevels covRleList Extracted from forward RleList

# Usage

```
## S4 method for signature 'covRleList'
seqlevels(x)
```

# Arguments

Χ

a covRle object

## Value

integer vector with names

```
seqlevels, experiment-method
```

Seqlevels ORFik experiment Extracted from fasta genome index

# Description

Seqlevels ORFik experiment Extracted from fasta genome index

## Usage

```
## S4 method for signature 'experiment'
seqlevels(x)
```

# Arguments

Х

an ORFik experiment

## Value

integer vector with names

322 seqnamesPerGroup

```
seqnames, experiment-method
```

Seqnames ORFik experiment Extracted from fasta genome index

# Description

Seqnames ORFik experiment Extracted from fasta genome index

## Usage

```
## S4 method for signature 'experiment'
seqnames(x)
```

# Arguments

Х

an ORFik experiment

# Value

integer vector with names

seqnamesPerGroup

Get list of seqnames per granges group

# Description

Get list of seqnames per granges group

## Usage

```
seqnamesPerGroup(grl, keep.names = TRUE)
```

# Arguments

grl a GRangesList

keep.names a boolean, keep names or not, default: (TRUE)

### Value

```
a character vector or Rle of seqnames(if seqnames == T)
```

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### **Examples**

```
 \begin{split} \text{gr\_plus} &<\text{- GRanges}(\text{seqnames} = \text{c("chr1", "chr1")}, \\ &\quad \text{ranges} = \text{IRanges}(\text{c(7, 14), width} = 3), \\ &\quad \text{strand} = \text{c("+", "+")}) \\ \text{gr\_minus} &<\text{- GRanges}(\text{seqnames} = \text{c("chr2", "chr2")}, \\ &\quad \text{ranges} = \text{IRanges}(\text{c(4, 1), c(9, 3)}), \\ &\quad \text{strand} = \text{c("-", "-")}) \\ \text{grl} &<\text{- GRangesList}(\text{tx1} = \text{gr\_plus}, \text{tx2} = \text{gr\_minus}) \\ \text{seqnamesPerGroup}(\text{grl}) \end{aligned}
```

shiftFootprints

Shift footprints by selected offsets

### **Description**

Function shifts footprints (GRanges) using specified offsets for every of the specified lengths. Reads that do not conform to the specified lengths are filtered out and rejected. Reads are resized to single base in 5' end fashion, treated as p site. This function takes account for junctions and soft clips in cigars of the reads. Length of the footprint is saved in size' parameter of GRanges output. Footprints are also sorted according to their genomic position, ready to be saved as a ofst, covRle, bed or wig file.

### Usage

```
shiftFootprints(footprints, shifts, sort = TRUE)
```

## **Arguments**

footprints	Galignments object of RiboSeq reads - footprints, can also be path to the .bam /.ofst file. If Galignment object has a meta column called "score", this will be used as replicate numbering for that read. So be careful if you have custom files with score columns, with another meaning.
shifts	a data.frame / data.table with minimum 2 columns, fraction (selected read lengths) and offsets_start (relative position in nt). Output from detectRibosomeShifts. Run ORFik::shifts_load(df)[[1]] for an example of input.
sort	logical, default TRUE. If False will keep original order of reads, and not sort output reads in increasing genomic location per chromosome and strand.

#### **Details**

The two columns in the shift data.frame/data.table argument are:

- fraction Numeric vector of lengths of footprints you select for shifting.
- offsets\_start Numeric vector of shifts for corresponding selected\_lengths. eg. c(-10, -10) with selected\_lengths of c(31, 32) means length of 31 will be shifted left by 10. Footprints of length 32 will be shifted right by 10.

NOTE: It will remove softclips from valid width, the CIGAR 3S30M is qwidth 33, but will remove 3S so final read width is 30 in ORFik.

#### Value

A GRanges object of shifted footprints, sorted and resized to 1bp of p-site, with metacolumn "size" indicating footprint size before shifting and resizing, sorted in increasing order.

#### References

https://bmcgenomics.biomedcentral.com/articles/10.1186/s12864-018-4912-6

#### See Also

```
Other pshifting: changePointAnalysis(), detectRibosomeShifts(), shiftFootprintsByExperiment(), shiftPlots(), shifts_load(), shifts_save()
```

## **Examples**

```
## Basic run
# Transcriptome annotation ->
gtf_file <- system.file("extdata/references/danio_rerio", "annotations.gtf", package = "ORFik")
# Ribo seq data ->
riboSeq_file <- system.file("extdata/Danio_rerio_sample", "ribo-seq.bam", package = "ORFik")
## Not run:
footprints <- readBam(riboSeq_file)

# detect the shifts automagically
shifts <- detectRibosomeShifts(footprints, gtf_file)
# shift the RiboSeq footprints
shiftedReads <- shiftFootprints(footprints, shifts)

## End(Not run)</pre>
```

shiftFootprintsByExperiment

Shift footprints of each file in experiment

### **Description**

A function that combines the steps of periodic read length detection, p-site shift detection and p-shifting into 1 function. For more details, see: detectRibosomeShifts

Saves files to a specified location as .ofst and .wig, The .ofst file will include a score column containing read width.

The .wig files, will be saved in pairs of  $\pm$ -strand, and score column will be replicates of reads starting at that position, score = 5 means 5 reads.

Remember that different species might have different default Ribosome read lengths, for human, mouse etc, normally around 27:30.

# Usage

```
shiftFootprintsByExperiment(
 df,
 out.dir = pasteDir(libFolder(df), "/pshifted/"),
 start = TRUE,
 stop = FALSE,
  top_tx = 10L,
 minFiveUTR = 30L,
 minCDS = 150L,
 minThreeUTR = if (stop) {
     30
} else NULL,
 firstN = 150L,
 min_reads = 1000,
 min_reads_TIS = 50,
 accepted.lengths = 26:34,
 output_format = c("ofst", "wig"),
 BPPARAM = bpparam(),
  tx = NULL,
  shift.list = NULL,
 log = TRUE,
 heatmap = FALSE,
 must.be.periodic = TRUE,
 strict.fft = TRUE,
 verbose = FALSE
)
```

# Arguments

df	an ORFik experiment
out.dir	output directory for files, default: pasteDir(libFolder(df), "/pshifted/"), making a /pshifted folder inside default bam file location
start	(logical) Whether to include predictions based on the start codons. Default TRUE.
stop	(logical) Whether to include predictions based on the stop codons. Default FASLE. Only use if there exists 3' UTRs for the annotation. If peridicity around stop codon is stronger than at the start codon, use stop instead of start region for p-shifting.
top_tx	(integer), default 10. Specify which % of the top TIS coverage transcripts to use for estimation of the shifts. By default we take top 10 top covered transcripts as they represent less noisy data-set. This is only applicable when there are more than 1000 transcripts.
minFiveUTR	(integer) minimum bp for 5' UTR during filtering for the transcripts. Set to NULL if no 5' UTRs exists for annotation.
minCDS	(integer) minimum bp for CDS during filtering for the transcripts
minThreeUTR	(integer) minimum bp for 3' UTR during filtering for the transcripts. Set to NULL if no 3' UTRs exists for annotation.

firstN (integer) Represents how many bases of the transcripts downstream of start

codons to use for initial estimation of the periodicity.

min\_reads default (1000), how many reads must a read-length have in total to be considered

for periodicity.

min\_reads\_TIS default (50), how many reads must a read-length have in the TIS region to be

considered for periodicity.

accepted.lengths

accepted read lengths, default 26:34, usually ribo-seq is strongest between 27:32.

 $output\_format \quad default \ c("ofst", "wig"), use \ export.ofst \ or \ wiggle \ format \ (wig) \ using \ export.wiggle \ format \ (wig) \ using \ (wig) \ using \ (wig) \ using \ (wig) \ using \ (wig) \$ 

? Default is both.

Options are: c("ofst", "bigWig", "wig", "bed", "bedo") For future coverage per nucleotide, we advice to do here ofst and bigWig for other genome browsers,

then call convert\_to\_covRleList to get much faster R objects.

The wig format version can be used in IGV, the score column is counts of that read with that read length, the cigar reference width is lost, ofst is much faster to save and load in R, and retain cigar reference width, but can not be used in IGV.

Also for larger tracks, you can use "bigWig".

BPPARAM how many cores/threads to use? default: bpparam()

tx a GRangesList, if you do not have 5' UTRs in annotation, send your own ver-

sion. Example: extendLeaders(tx, 30) Where 30 bases will be new "leaders". Since each original transcript was either only CDS or non-coding (filtered out).

shift.list default NULL, or a list containing named data.frames / data.tables with mini-

mum 2 columns, fraction (selected read lengths) and offsets\_start (relative posi-

tion in nt). 1 named data.frame / data.table per library. Output from detectRibosomeShifts.

Run ORFik::shifts\_load(df) for an example of input. The names of the list must be the file paths of the Ribo-seq libraries. Use this to edit the shifts, if you

suspect some of them are wrong in an experiment.

logical, default (TRUE), output a log file with parameters used and a .rds file

with all shifts per library (can be loaded with shifts\_load)

heatmap a logical or character string, default FALSE. If TRUE, will plot heatmap of raw

reads before p-shifting to console, to see if shifts given make sense. You can

also set a filepath to save the file there.

must.be.periodic

logical TRUE, if FALSE will not filter on periodic read lengths. (The Fourier transform filter will be skipped). This is useful if you are not going to do periodicity analysis, that is: for you more coverage depth (more read lengths) is more

important than only keeping the high quality periodic read lengths.

strict.fft logical, TRUE. Use a FFT without noise filter. This means keep only reads

lengths that are "periodic for the human eye". If you want more coverage, set to FALSE, to also get read lengths that are "messy", but the noise filter detects the periodicity of 3. This should only be done when you do not need high quality periodic reads! Example would be differential translation analysis by counts

over each ORF.

verbose logical, default FALSE. Report details of analysis/periodogram. Good if you are

not sure if the analysis was correct.

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#### Value

NULL (Objects are saved to out.dir/pshited/"name\_pshifted.ofst", wig, bedo or .bedo)

#### References

https://bmcgenomics.biomedcentral.com/articles/10.1186/s12864-018-4912-6

#### See Also

```
Other pshifting: changePointAnalysis(), detectRibosomeShifts(), shiftFootprints(), shiftPlots(), shifts_load(), shifts_save()
```

#### **Examples**

```
df <- ORFik.template.experiment.zf()
df <- df[1,] #lets only p-shift first RFP sample
## Output files as both .ofst and .wig(can be viewed in IGV/UCSC)
shiftFootprintsByExperiment(df)
# If you only need in R, do: (then you get no .wig files)
#shiftFootprintsByExperiment(df, output_format = "ofst")
## With debug info:
#shiftFootprintsByExperiment(df, verbose = TRUE)
## Re-shift, if you think some are wrong
## Here as an example we update library 1, third read length to shift 12
shift.list <- shifts_load(df)
shift.list[[1]]$offsets_start[3] <- -12
#shiftFootprintsByExperiment(df, shift.list = shift.list)
## For additional speedup in R for nucleotide coverage (coveragePerTiling etc)</pre>
```

shiftPlots

Plot shifted heatmaps per library

#### **Description**

Around CDS TISs, plot coverage. A good validation for you p-shifting, to see shifts are corresponding and close to the CDS TIS.

# Usage

```
shiftPlots(
    df,
    output = NULL,
    title = "Ribo-seq",
    scoring = "transcriptNormalized",
    pShifted = TRUE,
    upstream = if (pShifted) 5 else 20,
    downstream = if (pShifted) 20 else 5,
```

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```
type = "bar",
addFracPlot = TRUE,
plot.ext = ".pdf",
dpi = ifelse(nrow(df) < 22, 300, 200),
height_scaler = ifelse(type == "heatmap", 85, 95),
BPPARAM = bpparam()
)</pre>
```

# Arguments

df	an ORFik experiment
output	name to save file, full path. (Default NULL) No saving. Sett to "auto" to save to QC_STATS folder of experiment named: "pshifts_barplots.png" or "pshifts_heatmaps.png" depending on type argument. Folder must exist!
title	Title for top of plot, default "Ribo-seq". A more informative name could be "Ribo-seq zebrafish Chew et al. 2013"
scoring	which scoring scheme to use for heatmap, default "transcriptNormalized". Some alternatives: "sum", "zscore".
pShifted	a logical (TRUE), are Ribo-seq reads p-shifted to size 1 width reads? If upstream and downstream is set, this argument is irrelevant. So set to FALSE if this is not p-shifted Ribo-seq.
upstream	an integer (5), relative region to get upstream from. Default: ifelse(!is.null(tx), ifelse(pShifted, 5, 20), min(ifelse(pShifted, 5, 20), 0))
downstream	an integer (20), relative region to get downstream from. Default: ifelse(pShifted, 20, 5)
type	character, default "bar". Plot as faceted bars, gives more detailed information of read lengths, but harder to see patterns over multiple read lengths. Alternative: "heatmap", better overview of patterns over multiple read lengths.
addFracPlot	logical, default TRUE, add positional sum plot on top per heatmap.
plot.ext	default ".pdf". Alternative ".png". Only added if output is "auto".
dpi	numeric, default: ifelse(nrow(df) < 22, 300, 200)
height_scaler	numeric default: ifelse(type == "heatmap", 85, 95). The total height of plot in unit "mm" is (length(res) -1) * height_scaler. Increase if many readlengths are used.
BPPARAM	how many cores/threads to use? default: bpparam()

#### Value

```
a ggplot2 grob object
```

# See Also

```
Other pshifting: changePointAnalysis(), detectRibosomeShifts(), shiftFootprints(), shiftFootprintsByExperishifts_load(), shifts_save()
```

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### **Examples**

```
df <- ORFik.template.experiment.zf()
df <- df[df$libtype == "RFP",][1,] #lets only p-shift first RFP sample
#shiftFootprintsByExperiment(df, output_format = "bedo)
#grob <- shiftPlots(df, title = "Ribo-seq Human ORFik et al. 2020")
#plot(grob) #Only plot in RStudio for small amount of files!</pre>
```

shifts\_load

Load the shifts from experiment

# Description

When you p-shift using the function shiftFootprintsByExperiment, you will get a list of shifts per library. To automatically load them, you can use this function. Defaults to loading pshifts, if you made a-sites or e-sites, change the path argument to ashifted/eshifted folder instead.

#### Usage

```
shifts_load(
   df,
   path = file.path(libFolder(df), "pshifted", "shifting_table.rds")
)
```

#### **Arguments**

```
df an ORFik experiment

path path, default file.path(libFolder(df), "pshifted", "shifting_table.rds"). Path to
.rds file containing the shifts as a list, one list element per shifted bam file.
```

#### Value

a list of the shifts, one list element per shifted bam file.

### See Also

```
Other pshifting: changePointAnalysis(), detectRibosomeShifts(), shiftFootprints(), shiftFootprintsByExperishiftPlots(), shifts_save()
```

# **Examples**

```
df <- ORFik.template.experiment()
# subset on Ribo-seq
df <- df[df$libtype == "RFP",][1,]
#shiftFootprintsByExperiment(df)
#shifts_load(df)</pre>
```

shifts\_save

shifts\_save

Save shifts for Ribo-seq

#### **Description**

Should be stored in pshifted folder relative to default files

#### Usage

```
shifts_save(shifts, folder)
```

### **Arguments**

shifts a list of data.table/data.frame objects. Must be named with the full path to

ofst/bam files that defines the shifts.

folder directory to save file, Usually: file.path(libFolder(df), "pshifted"), where df is

the ORFik experiment / or your path of default file types. It will be named file.path(folder, "shifting\_table.rds"). For ORFik to work optimally, the folder

should be the /pshifted/ folder relative to default files.

#### Value

invisible(NULL), file saved to disc as "shifting\_table.rds".

#### See Also

Other pshifting: changePointAnalysis(), detectRibosomeShifts(), shiftFootprints(), shiftFootprintsByExperishiftPlots(), shifts\_load()

#### **Examples**

```
df <- ORFik.template.experiment.zf()
shifts <- shifts_load(df)
original_shifts <- file.path(libFolder(df), "pshifted", "shifting_table.rds")
# Move to temp
new_shifts_path <- file.path(tempdir(), "shifting_table.rds")
new_shifts <- c(shifts, shifts)
names(new_shifts)[2] <- file.path(tempdir(), "RiboSeqTemp.ofst")
saveRDS(new_shifts, new_shifts_path)
new_shifts[[1]][1,2] <- -10
# Now update the new shifts, here we input only first
shifts_save(new_shifts[1], tempdir())
readRDS(new_shifts_path) # You still get 2 outputs</pre>
```

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show,covRle-method

covRle show definition

# Description

Show a simplified version of the covRle

# Usage

```
## S4 method for signature 'covRle'
show(object)
```

# Arguments

object

acovRle

#### Value

print state of covRle

```
show,covRleList-method
```

covRleList show definition

# Description

Show a simplified version of the covRleList.

# Usage

```
## S4 method for signature 'covRleList'
show(object)
```

# Arguments

object

acovRleList

# Value

print state of covRleList

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```
show, experiment-method
```

experiment show definition

### **Description**

Show a simplified version of the experiment. The show function simplifies the view so that any column of data (like replicate or stage) is not shown, if all values are identical in that column. Filepaths are also never shown.

#### Usage

```
## S4 method for signature 'experiment'
show(object)
```

# Arguments

object

an ORFik experiment

#### Value

print state of experiment

simpleLibs

Converted format of NGS libraries

### Description

Export as either .ofst, .wig, .bigWig,.bedo (legacy format) or .bedoc (legacy format) files:

Export files as .ofst for fastest load speed into R.

Export files as .wig / bigWig for use in IGV or other genome browsers.

The input files are checked if they exist from: envExp(df).

### Usage

```
simpleLibs(
   df,
   out.dir = libFolder(df),
   addScoreColumn = TRUE,
   addSizeColumn = TRUE,
   must.overlap = NULL,
   method = "None",
   type = "ofst",
   input.type = "ofst",
```

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```
reassign.when.saving = FALSE,
envir = envExp(df),
force = TRUE,
library.names = bamVarName(df),
libs = outputLibs(df, type = input.type, chrStyle = must.overlap, library.names =
    library.names, output.mode = "list", force = force, BPPARAM = BPPARAM),
BPPARAM = bpparam()
)
```

#### **Arguments**

df an ORFik experiment

out.dir optional output directory, default: libFolder(df), if it is NULL, it will just reas-

sign R objects to simplified libraries. Will then create a final folder specified as: paste0(out.dir, "/", type, "/"). Here the files will be saved in format given by the

type argument.

addScoreColumn logical, default TRUE, if FALSE will not add replicate numbers as score col-

umn, see ORFik::convertToOneBasedRanges.

addSizeColumn logical, default TRUE, if FALSE will not add size (width) as size column, see

ORFik::convertToOneBasedRanges. Does not apply for (GAlignment version

of.ofst) or .bedoc. Since they contain the original cigar.

must.overlap default (NULL), else a GRanges / GRangesList object, so only reads that over-

lap (must.overlap) are kept. This is useful when you only need the reads over

transcript annotation or subset etc.

method character, default "None", the method to reduce ranges, for more info see convertToOneBasedRanges

type character, output format, default "ofst". Alternatives: "ofst", "bigWig", "wig", "bedo"

or "bedoc". Which format you want. Will make a folder within out.dir with this

name containing the files.

input . type character, input type "ofst". Remember this function uses the loaded libraries if

existing, so this argument is usually ignored. Only used if files do not already

exist.

reassign.when.saving

logical, default FALSE. If TRUE, will reassign library to converted form after

saving. Ignored when out.dir = NULL.

envir environment to save to, default envExp(df), which defaults to .GlobalEnv, but

can be set with envExp(df) <- new.env() etc.

force logical, default TRUE If TRUE, reload library files even if matching named

variables are found in environment used by experiment (see envExp) A simple way to make sure correct libraries are always loaded. FALSE is faster if data is

loaded correctly already.

library.names character vector, names of libraries, default: name\_decider(df, naming)

libs list, output of outputLibs as list of GRanges/GAlignments/GAlignmentPairs ob-

jects. Set input.type and force arguments to define parameters.

BPPARAM how many cores/threads to use? default: bpparam(). To see number of threads

used, do bpparam()\$workers. You can also add a time remaining bar, for a

more detailed pipeline.

334 sortPerGroup

#### **Details**

We advice you to not use this directly, as other function are more safe for library type conversions. See family description below. This is mostly used internally in ORFik. It is only adviced to use if large bam files are already loaded in R and conversions are wanted from those.

See export.ofst, export.wiggle, export.bedo and export.bedoc for information on file formats.

If libraries of the experiment are already loaded into environment (default: .globalEnv) is will export using those files as templates. If they are not in environment the .ofst files from the bam files are loaded (unless you are converting to .ofst then the .bam files are loaded).

#### Value

invisible NULL (saves files to disc or R .GlobalEnv)

#### See Also

```
Other lib_converters: convert_bam_to_ofst(), convert_to_bigWig(), convert_to_covRle(), convert_to_covRleList()
```

### **Examples**

```
df <- ORFik.template.experiment()
#convertLibs(df, out.dir = NULL)
# Keep only 5' ends of reads
#convertLibs(df, out.dir = NULL, method = "5prime")</pre>
```

sortPerGroup

Sort a GRangesList

#### **Description**

A faster, more versatile reimplementation of sort.GenomicRanges for GRangesList, needed since the original works poorly for more than 10k groups. This function sorts each group, where "+" strands are increasing by starts and "-" strands are decreasing by ends.

#### Usage

```
sortPerGroup(grl, ignore.strand = FALSE, quick.rev = FALSE)
```

#### **Arguments**

grl a GRangesList

ignore.strand a boolean, (default FALSE): should minus strands be sorted from highest to

lowest ends. If TRUE: from lowest to highest ends.

quick.rev default: FALSE, if TRUE, given that you know all ranges are sorted from min

to max for both strands, it will only reverse coordinates for minus strand groups,

and only if they are in increasing order. Much quicker

splitIn3Tx 335

### **Details**

Note: will not work if groups have equal names.

#### Value

an equally named GRangesList, where each group is sorted within group.

# **Examples**

```
 \begin{split} \text{gr\_plus} <& - \text{GRanges}(\text{seqnames} = \text{c("chr1", "chr1")}, \\ & \text{ranges} = \text{IRanges}(\text{c(14, 7), width} = 3), \\ & \text{strand} = \text{c("+", "+")}) \\ \text{gr\_minus} <& - \text{GRanges}(\text{seqnames} = \text{c("chr2", "chr2")}, \\ & \text{ranges} = \text{IRanges}(\text{c(1, 4), c(3, 9)}), \\ & \text{strand} = \text{c("-", "-")}) \\ \text{grl} <& - \text{GRangesList}(\text{tx1} = \text{gr\_plus}, \text{tx2} = \text{gr\_minus}) \\ \text{sortPerGroup}(\text{grl}) \end{aligned}
```

splitIn3Tx

Create binned coverage of transcripts, split into the 3 parts.

# Description

The 3 parts of transcripts are the leaders, the cds' and trailers. Per transcript part, bin them all to windowSize (default 100), and make a data.table, rows are positions, useful for plotting with ORFik and ggplot2.

### Usage

```
splitIn3Tx(
  leaders,
  cds,
  trailers,
  reads,
  windowSize = 100,
  fraction = "1",
  weight = "score",
  is.sorted = FALSE,
  drop.zero.dt = FALSE,
  BPPARAM = BiocParallel::SerialParam()
)
```

336 stageNames

#### **Arguments**

leaders a GRangesList of leaders (5' UTRs) cds a GRangesList of coding sequences trailers a GRangesList of trailers (3' UTRs) reads GRanges or GAlignment of reads windowSize an integer (100), size of windows (columns). All genes with region smaller than this size are filter out for metacoverage. fraction a character (1), info on reads (which read length, or which type (RNA seq)) (row names) weight (default: 'score'), if defined a character name of valid meta column in subject. GRanges("chr1", 1, "+", score = 5), would mean score column tells that this alignment region was found 5 times. Formats which loads a score column like this: Bigwig, wig, ORFik ofst, collapsed bam, bedoc and .bedo. As do CAGEr CAGE files and many other package formats. You can also assign a score column manually. is.sorted logical (FALSE), is grl sorted. That is + strand groups in increasing ranges (1,2,3), and - strand groups in decreasing ranges (3,2,1)drop.zero.dt logical FALSE, if TRUE and as.data.table is TRUE, remove all 0 count positions. This greatly speeds up and most importantly, greatly reduces memory usage. Will not change any plots, unless 0 positions are used in some sense.

(mean, median, zscore coverage will only scale differently)

how many cores/threads to use? default: bpparam()

# Value

**BPPARAM** 

a data.table with columns position, score

	stageNames	Get stage name variants	
--	------------	-------------------------	--

### **Description**

Used to standardize nomeclature for experiments.

Example: Find timepoints 2 hours, 4 hours etc. Example: If using zebrafish stages as TRUE, 64Cell stage is same as 2 hours post fertilization, so all 2hpf will be converted to 64Cell etc.

### Usage

```
stageNames(zebrafish.stages = FALSE)
```

# Arguments

zebrafish.stages

logical, FALSE. If true, convert time points to stages.

#### Value

a data.table with 2 columns, the main name, and all name variants of the main name in second column as a list.

#### References

https://www.mbl.edu/zebrafish/files/2013/03/Kimmel\_stagingseries1.pdf

#### See Also

```
Other experiment_naming: batchNames(), cellLineNames(), cellTypeNames(), conditionNames(), fractionNames(), inhibitorNames(), libNames(), mainNames(), repNames(), tissueNames()
```

STAR.align.folder

Align all libraries in folder with STAR

### **Description**

Does either all files as paired end or single end, so if you have mix, split them in two different folders.

If STAR halts at .... loading genome, it means the STAR index was aborted early, then you need to run: STAR.remove.crashed.genome(), with the genome that crashed, and rerun.

# Usage

```
STAR.align.folder(
  input.dir,
  output.dir,
  index.dir,
  star.path = STAR.install(),
  fastp = install.fastp(),
  paired.end = FALSE,
  steps = "tr-ge",
  adapter.sequence = "auto",
  quality.filtering = FALSE,
 min.length = 20,
 mismatches = 3,
  trim.front = 0,
 max.multimap = 10,
  alignment.type = "Local",
  allow.introns = TRUE,
 max.cpus = min(90, BiocParallel::bpparam()$workers),
 wait = TRUE,
  include.subfolders = "n",
  resume = NULL,
 multiQC = TRUE,
  keep.contaminants = FALSE,
```

```
keep.contaminants.type = c("bam", "fastq")[1],
keep.unaligned.genome = FALSE,
keep.index.in.memory = FALSE,
script.folder = system.file("STAR_Aligner", "RNA_Align_pipeline_folder.sh", package =
    "ORFik"),
script.single = system.file("STAR_Aligner", "RNA_Align_pipeline.sh", package = "ORFik")
)
```

#### **Arguments**

input.dir

path to fast files to align, the valid input files will be search for from formats: (".fasta", ".fastq", ".fq", or ".fa") with or without compression of .gz. Also either paired end or single end reads. Pairs will automatically be detected from similarity of naming, separated by something as .1 and .2 in the end. If files are renamed, where pairs are not similarily named, this process will fail to find correct pairs!

output.dir

directory to save indices, default: paste0(dirname(arguments[1]), "/STAR\_index/"), where arguments is the arguments input for this function.

index.dir

path to STAR index folder. Path returned from ORFik function STAR.index, when you created the index folders.

star.path

path to STAR, default: STAR.install(), if you don't have STAR installed at default location, it will install it there, set path to a runnable star if you already have it.

fastp

path to fastp trimmer, default: install.fastp(), if you have it somewhere else already installed, give the path. Only works for unix (linux or Mac OS), if not on unix, use your favorite trimmer and give the output files from that trimmer as input.dir here.

paired.end

a logical: default FALSE, alternative TRUE. If TRUE, will auto detect pairs by names. Can not be a combination of both TRUE and FALSE!

If running in folder mode: The folder must then contain an even number of files and they must be named with the same prefix and sufix of either \_1 and \_2, 1 and 2, etc. If SRR numbers are used, it will start on lowest and match with second lowest etc.

steps

a character, default: "tr-ge", trimming then genome alignment steps of depletion and alignment wanted: The posible candidates you can use are:

- tr : trim reads
- co: contamination merged depletion
- ph: phix depletion
- rR: rrna depletion
- nc : ncrna depletion
- tR: trna depletion (Mature tRNA, so no intron checks done)
- ge : genome alignment
- all: run steps: "tr-co-ge" or "tr-ph-rR-nc-tR-ge", depending on if you have merged contaminants or not

If not "all", a subset of these ("tr-co-ph-rR-nc-tR-ge")

If co (merged contaminants) is used, non of the specific contaminants can be specified, since they should be a subset of co.

The step where you align to the genome is usually always included, unless you are doing pure contaminant analysis or only trimming. For Ribo-seq and TCP(RCP-seq) you should do rR (ribosomal RNA depletion), so when you made the STAR index you need the rRNA step, either use rRNA from .gtf or manual download. (usually just download a Silva rRNA database for SSU&LSU at: https://www.arb-silva.de/) for your species.

#### adapter.sequence

character, default: "auto". Auto detect adapter using fastp adapter auto detection, checking first 1.5M reads. (Auto detection of adapter will not work 100% of the time (if the library is of low quality), then you must rerun this function with specified adapter from fastp adapter analysis. , using FASTQC or other adapter detection tools, else alignment will most likely fail!). If already trimmed or trimming not wanted: adapter.sequence = "disable" .You can manually assign adapter like: "ATCTCGTATGCCGTCTTCTGCTTG" or "AAAAAAAAAAAA". You can also specify one of the three presets:

- illumina (TrueSeq ~75/100 bp sequencing) : AGATCGGAAGAGC
- small\_RNA (standard for ~50 bp sequencing): TGGAATTCTCGG
- nextera: CTGTCTCTTATA

Paired end auto detection uses overlap sequence of pairs, to use the slower more secure paired end adapter detection, specify as: "autoPE".

### quality.filtering

logical, default FALSE. Not needed for modern library prep of RNA-seq, Riboseq etc (usually  $< \sim 0.5$  If you are aligning bad quality data, set this to TRUE. These filters will then be applied (default of fastp), filter if:

- Number of N bases in read : > 5
- Read quality : > 40% of bases in the read are <Q15

min.length 20, minimum length of aligned read without mismatches to pass filter. Anything under 20 is dangerous, as chance of random hits will become high!

mismatches 3, max non matched bases. Excludes soft-clipping, this only filters reads that have defined mismatches in STAR. Only applies for genome alignment step.

trim. front 0, default trim 0 bases 5'. For Ribo-seq use default 0. Ignored if tr (trim) is not one of the arguments in "steps"

numeric, default 10. If a read maps to more locations than specified, will skip the read. Set to 1 to only get unique mapping reads. Only applies for genome alignment step. The depletions are allowing for multimapping.

alignment.type default: "Local": standard local alignment with soft-clipping allowed, "End-ToEnd" (global): force end-to-end read alignment, does not soft-clip.

allow.introns logical, default TRUE. Allow large gaps of N in reads during genome alignment, if FALSE: sets –alignIntronMax to 1 (no introns). NOTE: You will still get some spliced reads if you assigned a gtf at the index step.

max.cpus

integer, default: min(90, BiocParallel:::bpparam()\$workers), number of threads to use. Default is minimum of 90 and maximum cores - 2. So if you have 8 cores it will use 6. Note: FASTP will use maximum 16 threads as from testing I see performance actually degrades using anything higher. From testing I also see STAR gets no performance gain after ~50 threads. I do suspect this will change when hard drives gets better in the future.

wait

a logical (not NA) indicating whether the R interpreter should wait for the command to finish, or run it asynchronously. This will be ignored (and the interpreter will always wait) if intern = TRUE. When running the command asynchronously, no output will be displayed on the Rgui console in Windows (it will be dropped, instead).

#### include.subfolders

"n" (no), do recursive search downwards for fast files if "y".

resume

default: NULL, continue from step, lets say steps are "tr-ph-ge": (trim, phix depletion, genome alignment) and resume is "ge", you will then use the assumed already trimmed and phix depleted data and start at genome alignment, useful if something crashed. Like if you specified wrong STAR version, but the trimming step was completed. Resume mode can only run 1 step at the time.

multiQC

logical, default TRUE. Do mutliQC comparison of STAR alignment between all the samples. Outputted in aligned/LOGS folder. See ?STAR.multiQC

#### keep.contaminants

logical, default FALSE. Create and keep contaminant aligning bam files, default is to only keep unaliged fastq reads, which will be further processed in "ge" genome alignment step. Useful if you want to do further processing on contaminants, like specific coverage of specific tRNAs etc.

### keep.contaminants.type

logical, default "bam". If aligned files of contaminants are kept, which format to output as, only supports "bam" for now. Fasta / Fastq will be implemented later.

#### keep.unaligned.genome

logical, default FALSE. Create and keep reads that did not align at the genome alignment step, default is to only keep the aliged bam file. Useful if you want to do further processing on plasmids/custom sequences.

#### keep.index.in.memory

logical or character, default FALSE (i.e. LoadAndRemove). For STAR.align.single: If TRUE, will keep index in memory, useful if you need to loop over single calls, instead of using STAR.align.folder (remember last run should use FALSE, to remove index). For STAR.align.folder:

Only applies to last library, will always keep for all libraries before last. Alternative useful for MAC machines especially is "noShared", for machines that do not support shared memory index, usually gives error: "abort trap 6".

script.folder

location of STAR index script, default internal ORFik file. You can change it and give your own if you need special alignments.

script.single

location of STAR single file alignment script, default internal ORFik file. You can change it and give your own if you need special alignments.

#### **Details**

Can only run on unix systems (Linux, Mac and WSL (Windows Subsystem Linux)), and requires a minimum of 30GB memory on genomes like human, rat, zebrafish etc.

If for some reason the internal STAR alignment bash script will not work for you, like if you want more customization of the STAR/fastp arguments. You can copy the internal alignment script, edit it and give that as the script used for this function.

The trimmer used is fastp (the fastest I could find), also works on (Linux, Mac and WSL (Windows Subsystem Linux)). If you want to use your own trimmer set file1/file2 to the location of the trimmed files from your program.

A note on trimming from creator of STAR about trimming: "adapter trimming it definitely needed for short RNA sequencing. For long RNA-seq, I would agree with Devon that in most cases adapter trimming is not advantageous, since, by default, STAR performs local (not end-to-end) alignment, i.e. it auto-trims." So trimming can be skipped for longer reads.

#### Value

output.dir, can be used as as input in ORFik::create.experiment

#### See Also

```
Other STAR: STAR.align.single(), STAR.allsteps.multiQC(), STAR.index(), STAR.install(), STAR.multiQC(), STAR.remove.crashed.genome(), getGenomeAndAnnotation(), install.fastp()
```

#### **Examples**

```
# First specify directories wanted (temp directory here)
config_file <- tempfile()</pre>
#config.save(config_file, base.dir = tempdir())
#config <- ORFik::config(config_file)</pre>
## Yeast RNA-seq samples (small genome)
#project <- ORFik::config.exper("chalmers_2012", "Saccharomyces_cerevisiae", "RNA-seq", config)</pre>
#annotation.dir <- project["ref"]</pre>
#fastq.input.dir <- project["fastq RNA-seq"]</pre>
#bam.output.dir <- project["bam RNA-seq"]</pre>
## Download some SRA data and metadata (subset to 50k reads)
# info <- download.SRA.metadata("SRP012047", outdir = conf["fastq RNA-seq"])</pre>
# info <- info[1:2,] # Subset to 2 first libraries</pre>
# download.SRA(info, fastq.input.dir, rename = FALSE, subset = 50000)
## No contaminant depletion:
# annotation <- getGenomeAndAnnotation("Saccharomyces cerevisiae", annotation.dir)
# index <- STAR.index(annotation)</pre>
# STAR.align.folder(fastq.input.dir, bam.output.dir,
                     index, paired.end = FALSE) # Trim, then align to genome
## Human Ribo-seq sample (NB! very large genome and libraries!)
## Requires >= 32 GB memory
#project <- ORFik::config.exper("subtelny_2014", "Homo_sapiens", "Ribo-seq", config)</pre>
#annotation.dir <- project["ref"]</pre>
```

```
#fastq.input.dir <- project["fastq Ribo-seq"]</pre>
#bam.output.dir <- project["bam Ribo-seq"]</pre>
## Download some SRA data and metadata (full libraries)
# info <- download.SRA.metadata("DRR041459", fastq.input.dir)</pre>
# download.SRA(info, fastq.input.dir, rename = FALSE)
## Now align 2 different ways, without and with contaminant depletion
## No contaminant depletion:
# annotation <- getGenomeAndAnnotation("Homo sapiens", annotation.dir)</pre>
# index <- STAR.index(annotation)</pre>
# STAR.align.folder(fastq.input.dir, bam.output.dir,
                     index, paired.end = FALSE)
## All contaminants merged:
# annotation <- getGenomeAndAnnotation(</pre>
    organism = "Homo_sapiens",
     phix = TRUE, ncRNA = TRUE, tRNA = TRUE, rRNA = TRUE,
     output.dir = annotation.dir
# index <- STAR.index(annotation)</pre>
# STAR.align.folder(fastq.input.dir, bam.output.dir,
                     index, paired.end = FALSE,
                     steps = "tr-ge")
```

STAR.align.single

Align single or paired end pair with STAR

#### **Description**

Given a single NGS fastq/fasta library, or a paired setup of 2 mated libraries. Run either combination of fastq trimming, contamination removal and genome alignment. Works for (Linux, Mac and WSL (Windows Subsystem Linux))

# Usage

```
STAR.align.single(
   file1,
   file2 = NULL,
   output.dir,
   index.dir,
   star.path = STAR.install(),
   fastp = install.fastp(),
   steps = "tr-ge",
   adapter.sequence = "auto",
   quality.filtering = FALSE,
   min.length = 20,
   mismatches = 3,
   trim.front = 0,
```

```
max.multimap = 10,
alignment.type = "Local",
allow.introns = TRUE,
max.cpus = min(90, BiocParallel::bpparam()$workers),
wait = TRUE,
resume = NULL,
keep.contaminants = FALSE,
keep.unaligned.genome = FALSE,
keep.index.in.memory = FALSE,
script.single = system.file("STAR_Aligner", "RNA_Align_pipeline.sh", package = "ORFik")
```

#### **Arguments**

file1 library file, if paired must be R1 file. Allowed formats are: (.fasta, .fastq, .fq, or.fa) with or without compression of .gz. This filename usually contains a suffix of .1 file2 default NULL, set if paired end to R2 file. Allowed formats are: (.fasta, .fastq, .fq, or.fa) with or without compression of .gz. This filename usually contains a suffix of .2 directory to save indices, default: paste0(dirname(arguments[1]), "/STAR\_index/"), output.dir where arguments is the arguments input for this function. index.dir path to STAR index folder. Path returned from ORFik function STAR.index, when you created the index folders. path to STAR, default: STAR.install(), if you don't have STAR installed at destar.path fault location, it will install it there, set path to a runnable star if you already have it. fastp path to fastp trimmer, default: install.fastp(), if you have it somewhere else already installed, give the path. Only works for unix (linux or Mac OS), if not on unix, use your favorite trimmer and give the output files from that trimmer as input.dir here. steps a character, default: "tr-ge", trimming then genome alignment steps of depletion and alignment wanted: The posible candidates you can use

• tr: trim reads

are:

- co: contamination merged depletion
- ph : phix depletion
- rR: rrna depletion
- nc : ncrna depletion
- tR: trna depletion (Mature tRNA, so no intron checks done)
- ge : genome alignment
- all: run steps: "tr-co-ge" or "tr-ph-rR-nc-tR-ge", depending on if you have merged contaminants or not

If not "all", a subset of these ("tr-co-ph-rR-nc-tR-ge")

If co (merged contaminants) is used, non of the specific contaminants can be specified, since they should be a subset of co.

The step where you align to the genome is usually always included, unless you are doing pure contaminant analysis or only trimming. For Ribo-seq and TCP(RCP-seq) you should do rR (ribosomal RNA depletion), so when you made the STAR index you need the rRNA step, either use rRNA from .gtf or manual download. (usually just download a Silva rRNA database for SSU&LSU at: https://www.arb-silva.de/) for your species.

#### adapter.sequence

character, default: "auto". Auto detect adapter using fastp adapter auto detection, checking first 1.5M reads. (Auto detection of adapter will not work 100% of the time (if the library is of low quality), then you must rerun this function with specified adapter from fastp adapter analysis. , using FASTQC or other adapter detection tools, else alignment will most likely fail!). If already trimmed or trimming not wanted: adapter.sequence = "disable" . You can manually assign adapter like: "ATCTCGTATGCCGTCTTCTGCTTG" or "AAAAAAAAAAAA.". You can also specify one of the three presets:

- illumina (TrueSeq ~75/100 bp sequencing) : AGATCGGAAGAGC
- small\_RNA (standard for ~50 bp sequencing): TGGAATTCTCGG
- nextera: CTGTCTCTTATA

Paired end auto detection uses overlap sequence of pairs, to use the slower more secure paired end adapter detection, specify as: "autoPE".

### quality.filtering

logical, default FALSE. Not needed for modern library prep of RNA-seq, Riboseq etc (usually < ~ 0.5 If you are aligning bad quality data, set this to TRUE. These filters will then be applied (default of fastp), filter if:

- Number of N bases in read : > 5
- Read quality : > 40% of bases in the read are < Q15

min.length 20, minimum length of aligned read without mismatches to pass filter. Anything under 20 is dangerous, as chance of random hits will become high!

mismatches 3, max non matched bases. Excludes soft-clipping, this only filters reads that have defined mismatches in STAR. Only applies for genome alignment step.

trim.front 0, default trim 0 bases 5'. For Ribo-seq use default 0. Ignored if tr (trim) is not one of the arguments in "steps"

numeric, default 10. If a read maps to more locations than specified, will skip the read. Set to 1 to only get unique mapping reads. Only applies for genome alignment step. The depletions are allowing for multimapping.

alignment.type default: "Local": standard local alignment with soft-clipping allowed, "End-ToEnd" (global): force end-to-end read alignment, does not soft-clip.

logical, default TRUE. Allow large gaps of N in reads during genome alignment, if FALSE: sets -alignIntronMax to 1 (no introns). NOTE: You will still get some spliced reads if you assigned a gtf at the index step.

max.multimap

allow.introns

max.cpus integer, default: min(90, BiocParallel:::bpparam()\$workers), number of

threads to use. Default is minimum of 90 and maximum cores - 2. So if you have 8 cores it will use 6. Note: FASTP will use maximum 16 threads as from testing I see performance actually degrades using anything higher. From testing I also see STAR gets no performance gain after ~50 threads. I do suspect this

will change when hard drives gets better in the future.

wait a logical (not NA) indicating whether the R interpreter should wait for the com-

mand to finish, or run it asynchronously. This will be ignored (and the interpreter will always wait) if intern = TRUE. When running the command asynchronously, no output will be displayed on the Rgui console in Windows (it will

be dropped, instead).

resume default: NULL, continue from step, lets say steps are "tr-ph-ge": (trim, phix

depletion, genome alignment) and resume is "ge", you will then use the assumed already trimmed and phix depleted data and start at genome alignment, useful if something crashed. Like if you specified wrong STAR version, but the trimming

step was completed. Resume mode can only run 1 step at the time.

keep.contaminants

logical, default FALSE. Create and keep contaminant aligning bam files, default is to only keep unaliged fastq reads, which will be further processed in "ge" genome alignment step. Useful if you want to do further processing on

contaminants, like specific coverage of specific tRNAs etc.

keep.unaligned.genome

logical, default FALSE. Create and keep reads that did not align at the genome alignment step, default is to only keep the aliged bam file. Useful if you want to

do further processing on plasmids/custom sequences.

keep.index.in.memory

logical or character, default FALSE (i.e. LoadAndRemove). For STAR.align.single: If TRUE, will keep index in memory, useful if you need to loop over single calls, instead of using STAR.align.folder (remember last run should use FALSE, to re-

move index). For STAR.align.folder:

Only applies to last library, will always keep for all libraries before last. Alternative useful for MAC machines especially is "noShared", for machines that do

not support shared memory index, usually gives error: "abort trap 6".

script.single location of STAR single file alignment script, default internal ORFik file. You

can change it and give your own if you need special alignments.

### **Details**

Can only run on unix systems (Linux, Mac and WSL (Windows Subsystem Linux)), and requires a minimum of 30GB memory on genomes like human, rat, zebrafish etc.

If for some reason the internal STAR alignment bash script will not work for you, like if you want more customization of the STAR/fastp arguments. You can copy the internal alignment script, edit it and give that as the script used for this function.

The trimmer used is fastp (the fastest I could find), also works on (Linux, Mac and WSL (Windows Subsystem Linux)). If you want to use your own trimmer set file1/file2 to the location of the trimmed files from your program.

A note on trimming from creator of STAR about trimming: "adapter trimming it definitely needed

for short RNA sequencing. For long RNA-seq, I would agree with Devon that in most cases adapter trimming is not advantageous, since, by default, STAR performs local (not end-to-end) alignment, i.e. it auto-trims." So trimming can be skipped for longer reads.

#### Value

output.dir, can be used as as input in ORFik::create.experiment

#### See Also

```
Other STAR: STAR.align.folder(), STAR.allsteps.multiQC(), STAR.index(), STAR.install(), STAR.multiQC(), STAR.remove.crashed.genome(), getGenomeAndAnnotation(), install.fastp()
```

### **Examples**

```
## Specify output libraries (using temp config)
config_file <- tempfile()</pre>
#config.save(config_file, base.dir = tempdir())
#config <- ORFik::config(config_file)</pre>
#project <- ORFik::config.exper("yeast_1", "Saccharomyces_cerevisiae", "RNA-seq", config)</pre>
# Get genome of yeast (quite small)
# arguments <- getGenomeAndAnnotation("Saccharomyces cerevisiae", project["ref"])</pre>
# index <- STAR.index(arguments)</pre>
## Make fake reads
#genome <- readDNAStringSet(arguments["genome"])</pre>
#which_chromosomes <- sample(seq_along(genome), 1000, TRUE, prob = width(genome))</pre>
#nt50_windows <- lapply(which_chromosomes, function(x)</pre>
# {window <- sample(width(genome[x]) - 51, 1); genome[[x]][seq(window, window+49)]})</pre>
#nt50_windows <- DNAStringSet(nt50_windows)</pre>
#names(nt50_windows) <- paste0("read_", seq_along(nt50_windows))</pre>
#dir.create(project["fastq RNA-seq"], recursive = TRUE)
#fake_fasta <- file.path(project["fastq RNA-seq"], "fake-RNA-seq.fasta")</pre>
#writeXStringSet(nt50_windows, fake_fasta, format = "fasta")
## Align the fake reads and import bam
# STAR.align.single(fake_fasta, NULL, project["bam RNA-seq"], index, steps = "ge")
#bam_file <- list.files(file.path(project["bam RNA-seq"], "aligned"),</pre>
# pattern = "\.bam$", full.names = TRUE)
#fimport(bam_file)
```

STAR.allsteps.multiQC Create STAR multiQC plot and table

#### Description

Takes a folder with multiple Log.final.out files from STAR, and create a multiQC report. This is automatically run with STAR.align.folder function.

STAR.index 347

#### Usage

```
STAR.allsteps.multiQC(
  folder,
  steps = "auto",
  plot.ext = ".pdf",
  output.file = file.path(folder, "full_process.csv")
)
```

### **Arguments**

path to main output folder of STAR run. The folder that contains /aligned/, 
"/trim/, "contaminants\_depletion" etc. To find the LOGS folders in, to use for 
summarized statistics.

steps a character, default "auto". Find which steps you did. If manual, a combination 
of "tr-co-ge". See STAR alignment functions for description.

plot.ext character, default ".pdf". Which format to save QC plot. Alternative: ".png".

output.file character, file path, default: file.path(folder, "full\_process.csv")

#### Value

data.table of main statistics, plots and data saved to disc. Named: "/00\_STAR\_LOG\_plot.pdf" and "/00\_STAR\_LOG\_table.csv"

#### See Also

```
Other STAR: STAR.align.folder(), STAR.align.single(), STAR.index(), STAR.install(), STAR.multiQC(), STAR.remove.crashed.genome(), getGenomeAndAnnotation(), install.fastp()
```

### **Examples**

```
process_dir <- system.file("extdata/test_processing/", package = "ORFik")
STAR.allsteps.multiQC(process_dir)
STAR.allsteps.multiQC(process_dir, steps = "tr-ge")</pre>
```

STAR.index

Create STAR genome index

#### **Description**

Used as reference when aligning data
Get genome and gtf by running getGenomeAndFasta()

348 STAR.index

#### Usage

```
STAR.index(
  arguments,
  output.dir = paste0(dirname(arguments[1]), "/STAR_index/"),
  star.path = STAR.install(),
 max.cpus = min(90, BiocParallel::bpparam()$workers),
 max.ram = 30,
  SAsparse = 1,
  tmpDirStar = "-",
  wait = TRUE,
  remake = FALSE,
 script = system.file("STAR_Aligner", "STAR_MAKE_INDEX.sh", package = "ORFik"),
  notify_load_existing = TRUE
)
```

#### **Arguments**

arguments a named character vector containing paths wanted to use for index creation.

They must be named correctly: names must be a subset of: c("gtf", "genome",

"contaminants", "phix", "rRNA", "tRNA", "ncRNA")

output.dir directory to save indices, default: paste0(dirname(arguments[1]), "/STAR\_index/"),

where arguments is the arguments input for this function.

path to STAR, default: STAR.install(), if you don't have STAR installed at destar.path

fault location, it will install it there, set path to a runnable star if you already

have it.

max.cpus integer, default: min(90, BiocParallel:::bpparam()\$workers), number of

> threads to use. Default is minimum of 90 and maximum cores - 2. So if you have 8 cores it will use 6. Note: FASTP will use maximum 16 threads as from testing I see performance actually degrades using anything higher. From testing I also see STAR gets no performance gain after ~50 threads. I do suspect this

will change when hard drives gets better in the future.

integer, default 30, in Giga Bytes (GB). Maximum amount of RAM allowed for max.ram

> STAR limitGenomeGenerateRAM argument. RULE: idealy 10x genome size, but do not set too close to machine limit. Default fits well for human genome

size (3 GB \* 10 = 30 GB)

int > 0, default 1. If you do not have at least 64GB RAM, you might need to SAsparse

> set this to 2. suffux array sparsity, i.e. distance between indices: use bigger numbers to decrease needed RAM at the cost of mapping speed reduction. Only

applies to genome, not conaminants.

character, default "-". STAR automatic temp folder creation, deleted when done. tmpDirStar

> The directory can not exists, as a safety STAR must make it!. If you are on a NFS file share drive, and you have a non NFS tmp dir, set this to tempfile() or

the manually specified folder to get a considerable speedup!

a logical (not NA) indicating whether the R interpreter should wait for the com-

mand to finish, or run it asynchronously. This will be ignored (and the interpreter will always wait) if intern = TRUE. When running the command asyn-

wait

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chronously, no output will be displayed on the Rgui console in Windows (it will

be dropped, instead).

remake logical, default: FALSE, if TRUE remake everything specified

script location of STAR index script, default internal ORFik file. You can change it

and give your own if you need special alignments.

notify\_load\_existing

logical, default TRUE. If annotation exists (defined as: locally (a file called outputs.rds) exists in outputdir), print a small message notifying the user it is

not redownloading. Set to FALSE, if this is not wanted

#### **Details**

Can only run on unix systems (Linux and Mac), and requires minimum 30GB memory on genomes like human, rat, zebrafish etc.

If for some reason the internal STAR index bash script will not work for you, like if you have a very small genome. You can copy the internal index script, edit it and give that as the Index script used for this function. It is recommended to run through the RStudio local job tab, to give full info about the run. The system console will not stall, as can happen in happen in normal RStudio console.

#### Value

output.dir, can be used as as input for STAR.align..

#### See Also

```
Other STAR: STAR.align.folder(), STAR.align.single(), STAR.allsteps.multiQC(), STAR.install(), STAR.multiQC(), STAR.remove.crashed.genome(), getGenomeAndAnnotation(), install.fastp()
```

### **Examples**

```
## Manual way, specify all paths yourself.
#arguments <- c(path.GTF, path.genome, path.phix, path.rrna, path.trna, path.ncrna)
#names(arguments) <- c("gtf", "genome", "phix", "rRNA", "tRNA","ncRNA")
#STAR.index(arguments, "output.dir")

## Or use ORFik way:
output.dir <- "/Bio_data/references/Human"
# arguments <- getGenomeAndAnnotation("Homo sapiens", output.dir)
# STAR.index(arguments, output.dir)</pre>
```

STAR.install

Download and prepare STAR

#### **Description**

Will not run "make", only use precompiled STAR file.

Can only run on unix systems (Linux and Mac), and requires minimum 30GB memory on genomes like human, rat, zebrafish etc.

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#### Usage

```
STAR.install(folder = "~/bin", version = "2.7.4a")
```

### **Arguments**

folder path to folder for download, fille will be named "STAR-version", where version

is version wanted.

version default "2.7.4a"

#### **Details**

ORFik for now only uses precompiled STAR binaries, so if you already have a STAR version it is adviced to redownload the same version, since STAR genome indices usually does not work between STAR versions.

#### Value

path to runnable STAR

#### References

https://www.ncbi.nlm.nih.gov/pubmed/23104886

#### See Also

```
Other STAR: STAR.align.folder(), STAR.align.single(), STAR.allsteps.multiQC(), STAR.index(), STAR.multiQC(), STAR.remove.crashed.genome(), getGenomeAndAnnotation(), install.fastp()
```

### **Examples**

```
## Default folder install:
#STAR.install()
## Manual set folder:
folder <- "/I/WANT/IT/HERE"
#STAR.install(folder, version = "2.7.4a")</pre>
```

STAR.multiQC

Create STAR multiQC plot and table

### Description

Takes a folder with multiple Log.final.out files from STAR, and create a multiQC report

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### Usage

```
STAR.multiQC(
  folder,
  type = "aligned",
  plot.ext = ".pdf",
  log_files = dir(folder, "Log.final.out", full.names = TRUE),
  simplified_table = TRUE
)
```

# **Arguments**

folder	path to LOGS folder of ORFik STAR runs. Can also be the path to the aligned/ (parent directory of LOGS), then it will move into LOG from there. Only if no files with pattern Log.final.out are found in parent directory. If no LOGS folder is found it can check for a folder /aligned/LOGS/ so to go 2 folders down.
type	a character path, default "aligned". Which subfolder to check for. If you want log files for contamination do type = "contaminants_depletion"
plot.ext	character, default ".pdf". Which format to save QC plot. Alternative: ".png".
log_files	character, path to "Log.final.out" STAR files, default: dir(folder, "Log.final.out", full.names = TRUE)
simplified_table	
	logical default TDIJE Subset columns to the most important ones

logical, default TRUE. Subset columns, to the most important ones.

#### Value

a data.table with all information from STAR runs, plot and data saved to disc. Named: " $/00\_STAR\_LOG\_plot.pdf$ " and " $/00\_STAR\_LOG\_table.csv$ "

# See Also

```
Other STAR: STAR.align.folder(), STAR.align.single(), STAR.allsteps.multiQC(), STAR.index(), STAR.install(), STAR.remove.crashed.genome(), getGenomeAndAnnotation(), install.fastp()
```

### **Examples**

```
#' @examples
process_dir <- system.file("extdata/test_processing/", package = "ORFik")
STAR.multiQC(process_dir)</pre>
```

352 startCodons

```
STAR.remove.crashed.genome
```

Remove crashed STAR genome

# **Description**

This happens if you abort STAR run early, and it halts at: ..... loading genome

#### Usage

```
STAR.remove.crashed.genome(index.path, star.path = STAR.install())
```

### **Arguments**

index.path path to index folder of genome

star.path path to STAR, default: STAR.install(), if you don't have STAR installed at de-

fault location, it will install it there, set path to a runnable star if you already

have it.

#### Value

return value from system call, 0 if all good.

### See Also

```
Other STAR: STAR.align.folder(), STAR.align.single(), STAR.allsteps.multiQC(), STAR.index(), STAR.install(), STAR.multiQC(), getGenomeAndAnnotation(), install.fastp()
```

#### **Examples**

```
index.path = "/home/data/human_GRCh38/STAR_INDEX/genomeDir/"
# STAR.remove.crashed.genome(index.path = index.path)
## If you have the index argument from STAR.index function:
# index.path <- STAR.index()
# STAR.remove.crashed.genome(file.path(index.path, "genomeDir"))
# STAR.remove.crashed.genome(file.path(index.path, "contaminants_genomeDir"))</pre>
```

startCodons Get the Start codons

Get the Start codons(3 bases) from a GRangesList of orfs grouped by orfs

# **Description**

In ATGTTTTGA, get the positions ATG. It takes care of exons boundaries, with exons < 3 length.

startDefinition 353

#### Usage

```
startCodons(grl, is.sorted = FALSE)
```

#### **Arguments**

```
grl a GRangesList object
is.sorted a boolean, a speedup if you know the ranges are sorted
```

#### Value

a GRangesList of start codons, since they might be split on exons

#### See Also

```
Other ORFHelpers: defineTrailer(), longestORFs(), mapToGRanges(), orfID(), startSites(), stopCodons(), stopSites(), txNames(), uniqueGroups(), uniqueOrder()
```

### **Examples**

startDefinition

Returns start codon definitions

# Description

According to: <a href="http://www.ncbi.nlm.nih.gov/Taxonomy/taxono

### Usage

```
startDefinition(transl_table)
```

### **Arguments**

```
transl_table numeric. NCBI genetic code number for translation.
```

### Value

A string of START sites separatd with "I".

354 startRegion

#### See Also

```
Other findORFs: findMapORFs(), findORFs(), findORFsFasta(), findUORFs(), stopDefinition()
```

### **Examples**

```
startDefinition
startDefinition(1)
```

startRegion

Start region as GRangesList

#### **Description**

Get the start region of each ORF. If you want the start codon only, set upstream = 0 or just use startCodons. Standard is 2 upstream and 2 downstream, a width 5 window centered at start site. since p-shifting is not 100 usually the reads from the start site.

# Usage

```
startRegion(grl, tx = NULL, is.sorted = TRUE, upstream = 2L, downstream = 2L)
```

# Arguments

grl	a GRangesList object with usually either leaders, cds', 3' utrs or ORFs
tx	default NULL, a GRangesList of transcripts or (container region), names of tx
	must contain all grl names. The names of grl can also be the ORFik orf names.
	that is "txName_id"
is.sorted	logical (TRUE), is grl sorted.
unstream	an integer (2) relative region to get unstream from

upstream an integer (2), relative region to get upstream from. downstream an integer (2), relative region to get downstream from

# **Details**

If tx is null, then upstream will be forced to 0 and downstream to a maximum of grl width (3' UTR end for mRNAs). Since there is no reference for splicing.

### Value

a GRanges, or GRangesList object if any group had > 1 exon.

#### See Also

```
Other features: computeFeatures(), computeFeaturesCage(), countOverlapsW(), disengagementScore(), distToCds(), distToTSS(), entropy(), floss(), fpkm(), fpkm_calc(), fractionLength(), initiationScore(), insideOutsideORF(), isInFrame(), isOverlapping(), kozakSequenceScore(), orfScore(), rankOrder(), ribosomeReleaseScore(), ribosomeStallingScore(), startRegionCoverage(), stopRegion(), subsetCoverage(), translationalEff()
```

startRegionCoverage 355

### **Examples**

startRegionCoverage

Start region coverage

### **Description**

Get the number of reads in the start region of each ORF. If you want the start codon coverage only, set upstream = 0. Standard is 2 upstream and 2 downstream, a width 5 window centered at start site. since p-shifting is not 100 start site.

#### Usage

```
startRegionCoverage(
  grl,
  RFP,
  tx = NULL,
  is.sorted = TRUE,
  upstream = 2L,
  downstream = 2L,
  weight = 1L
)
```

#### Arguments

grl a GRangesList object with usually either leaders, cds', 3' utrs or ORFs

RFP ribo seq reads as GAlignments, GRanges or GRangesList object

tx default NULL, a GRangesList of transcripts or (container region), names of tx

must contain all grl names. The names of grl can also be the ORFik orf names.

that is "txName id"

is.sorted logical (TRUE), is grl sorted.

upstream an integer (2), relative region to get upstream from.

downstream an integer (2), relative region to get downstream from

weight a numeric/integer vector or metacolumn name. (de:

a numeric/integer vector or metacolumn name. (default: 1L, no differential weighting). If weight is name of defined meta column in reads object, it gives the number of times a read was found at that position. GRanges("chr1", 1, "+", score = 5), would mean "score" column tells that this alignment region was found 5 times. if 1L it means each read is weighted equal as 1, this is what among others countOverlaps() presumes, if single number (!= 1), it repeats for all ranges, if vector with length > 1, it must be equal size of the reads object.

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#### **Details**

If tx is null, then upstream will be force to 0 and downstream to a maximum of grl width. Since there is no reference for splicing.

#### Value

a numeric vector of counts

#### See Also

```
Other features: computeFeatures(), computeFeaturesCage(), countOverlapsW(), disengagementScore(), distToCds(), distToTSS(), entropy(), floss(), fpkm(), fpkm_calc(), fractionLength(), initiationScore(), insideOutsideORF(), isInFrame(), isOverlapping(), kozakSequenceScore(), orfScore(), rankOrder(), ribosomeReleaseScore(), ribosomeStallingScore(), startRegion(), stopRegion(), subsetCoverage(), translationalEff()
```

# Examples

startRegionString

Get start region as DNA-strings per GRanges group

#### Description

One window per start site, if upstream and downstream are both 0, then only the startsite is returned.

# Usage

```
startRegionString(grl, tx, faFile, upstream = 20, downstream = 20)
```

### **Arguments**

```
grl a GRangesList of ranges to find regions in.

tx a GRangesList of transcripts or (container region), names of tx must contain all gr names. The names of gr can also be the ORFik orf names. that is "tx-Name_id".
```

startSites 357

faFile FaFile, BSgenome, fasta/index file path or an ORFik experiment. This file is usually used to find the transcript sequences from some GRangesList.

upstream an integer, default (0), relative region to get upstream end from. (0 means start site, +1 is one upstream, -1 is one downstream)

downstream an integer, default (0), relative region to get downstream end from (0 means start site, +1 is one downstream, -1 is one upstream)

#### Value

a character vector of start regions

startSites Get the start sites from a GRangesList of orfs grouped by orfs

# Description

In ATGTTTTGG, get the position of the A.

#### Usage

```
startSites(grl, asGR = FALSE, keep.names = FALSE, is.sorted = FALSE)
```

### **Arguments**

```
grl a GRangesList object
asGR a boolean, return as GRanges object
keep.names a logical (FALSE), keep names of input.
is.sorted a speedup, if you know the ranges are sorted
```

#### Value

if asGR is False, a vector, if True a GRanges object

### See Also

```
Other ORFHelpers: defineTrailer(), longestORFs(), mapToGRanges(), orfID(), startCodons(), stopCodons(), stopSites(), txNames(), uniqueGroups(), uniqueOrder()
```

### **Examples**

```
 \begin{split} \text{gr\_plus} <&- \text{GRanges}(\text{seqnames} = \text{c("chr1", "chr1")}, \\ &\quad \text{ranges} = \text{IRanges}(\text{c(7, 14), width} = 3), \\ &\quad \text{strand} = \text{c("+", "+")}) \\ \text{gr\_minus} <&- \text{GRanges}(\text{seqnames} = \text{c("chr2", "chr2")}, \\ &\quad \text{ranges} = \text{IRanges}(\text{c(4, 1), c(9, 3)}), \\ &\quad \text{strand} = \text{c("-", "-")}) \\ \text{grl} <&- \text{GRangesList}(\text{tx1} = \text{gr\_plus}, \text{tx2} = \text{gr\_minus}) \\ \text{startSites}(\text{grl, is.sorted} = \text{FALSE}) \\ \end{split}
```

358 stopCodons

stopCodons	Get the Stop codons (3 bases) from a GRangesList of orfs grouped by orfs
------------	--

### **Description**

In ATGTTTTGA, get the positions TGA. It takes care of exons boundaries, with exons < 3 length.

### Usage

```
stopCodons(grl, is.sorted = FALSE)
```

# **Arguments**

```
grl a GRangesList object
is.sorted a boolean, a speedup if you know the ranges are sorted
```

### Value

a GRangesList of stop codons, since they might be split on exons

#### See Also

```
Other ORFHelpers: defineTrailer(), longestORFs(), mapToGRanges(), orfID(), startCodons(), startSites(), stopSites(), txNames(), uniqueGroups(), uniqueOrder()
```

### **Examples**

stopDefinition 359

stopDefinition

Returns stop codon definitions

### **Description**

According to: <a href="http://www.ncbi.nlm.nih.gov/Taxonomy/taxono

### Usage

```
stopDefinition(transl_table)
```

### **Arguments**

transl\_table numeric. NCBI genetic code number for translation.

### Value

A string of STOP sites separatd with "I".

#### See Also

```
Other findORFs: findMapORFs(), findORFs(), findORFsFasta(), findUORFs(), startDefinition()
```

# **Examples**

```
stopDefinition
stopDefinition(1)
```

stopRegion

Stop region as GRangesList

#### **Description**

Get the stop region of each ORF / region. If you want the stop codon only, set downstream = 0 or just use stopCodons. Standard is 2 upstream and 2 downstream, a width 5 window centered at stop site.

# Usage

```
stopRegion(grl, tx = NULL, is.sorted = TRUE, upstream = 2L, downstream = 2L)
```

360 stopSites

### **Arguments**

grl	a GRangesList object with usually either leaders, cds', 3' utrs or ORFs
tx	default NULL, a GRangesList of transcripts or (container region), names of tx must contain all grl names. The names of grl can also be the ORFik orf names. that is "txName_id"
is.sorted	logical (TRUE), is grl sorted.
upstream	an integer (2), relative region to get upstream from.
downstream	an integer (2), relative region to get downstream from

#### **Details**

If tx is null, then downstream will be forced to 0 and upstream to a minimum of -grl width (to the TSS). . Since there is no reference for splicing.

#### Value

a GRanges, or GRangesList object if any group had > 1 exon.

#### See Also

```
Other features: computeFeatures(), computeFeaturesCage(), countOverlapsW(), disengagementScore(), distToCds(), distToTSS(), entropy(), floss(), fpkm(), fpkm_calc(), fractionLength(), initiationScore(), insideOutsideORF(), isInFrame(), isOverlapping(), kozakSequenceScore(), orfScore(), rankOrder(), ribosomeReleaseScore(), ribosomeStallingScore(), startRegion(), startRegionCoverage(), subsetCoverage(), translationalEff()
```

# Examples

stopSites

Get the stop sites from a GRangesList of orfs grouped by orfs

### **Description**

In ATGTTTTGC, get the position of the C.

### Usage

```
stopSites(grl, asGR = FALSE, keep.names = FALSE, is.sorted = FALSE)
```

strandBool 361

#### **Arguments**

```
grl a GRangesList object
asGR a boolean, return as GRanges object
keep.names a logical (FALSE), keep names of input.
is.sorted a speedup, if you know the ranges are sorted
```

#### Value

if asGR is False, a vector, if True a GRanges object

## See Also

```
Other ORFHelpers: defineTrailer(), longestORFs(), mapToGRanges(), orfID(), startCodons(), startSites(), stopCodons(), txNames(), uniqueGroups(), uniqueOrder()
```

### **Examples**

```
 \begin{split} \text{gr\_plus} <&- \text{GRanges}(\text{seqnames} = \text{c("chr1", "chr1")}, \\ &\quad \text{ranges} = \text{IRanges}(\text{c(7, 14), width} = 3), \\ &\quad \text{strand} = \text{c("+", "+")}) \\ \text{gr\_minus} <&- \text{GRanges}(\text{seqnames} = \text{c("chr2", "chr2")}, \\ &\quad \text{ranges} = \text{IRanges}(\text{c(4, 1), c(9, 3)}), \\ &\quad \text{strand} = \text{c("-", "-")}) \\ \text{grl} <&- \text{GRangesList}(\text{tx1} = \text{gr\_plus}, \text{tx2} = \text{gr\_minus}) \\ \text{stopSites}(\text{grl}, \text{is.sorted} = \text{FALSE}) \\ \end{split}
```

strandBool

Get logical list of strands

# Description

Helper function to get a logical list of True/False, if GRangesList group have + strand = T, if - strand = F Also checks for \* strands, so a good check for bugs

### Usage

```
strandBool(grl)
```

## **Arguments**

grl a GRangesList or GRanges object

### Value

a logical vector

## **Examples**

strandMode,covRle-method

strandMode covRle

## **Description**

strandMode covRle

## Usage

```
## S4 method for signature 'covRle'
strandMode(x)
```

## **Arguments**

Х

a covRle object

### Value

integer vector with names

```
strand \verb|Mode,covRleList-method|\\ strand \verb|Mode covRle|\\
```

## **Description**

strandMode covRle

### Usage

```
## S4 method for signature 'covRleList'
strandMode(x)
```

# Arguments

Х

a covRle object

#### Value

integer vector with names

strandPerGroup 363

strandPerGroup

Get list of strands per granges group

## **Description**

Get list of strands per granges group

### Usage

```
strandPerGroup(grl, keep.names = TRUE)
```

## Arguments

```
grl a GRangesList
```

keep.names a boolean, keep names or not, default: (TRUE)

#### Value

a vector named/unnamed of characters

## **Examples**

```
 \begin{split} \text{gr\_plus} <&- \text{GRanges}(\text{seqnames} = \text{c("chr1", "chr1")}, \\ &\quad \text{ranges} = \text{IRanges}(\text{c(7, 14), width} = 3), \\ &\quad \text{strand} = \text{c("+", "+")}) \\ \text{gr\_minus} <&- \text{GRanges}(\text{seqnames} = \text{c("chr2", "chr2")}, \\ &\quad \text{ranges} = \text{IRanges}(\text{c(4, 1), c(9, 3)}), \\ &\quad \text{strand} = \text{c("-", "-")}) \\ \text{grl} <&- \text{GRangesList}(\text{tx1} = \text{gr\_plus}, \text{tx2} = \text{gr\_minus}) \\ \text{strandPerGroup}(\text{grl}) \end{aligned}
```

subsetCoverage

Subset GRanges to get coverage.

## **Description**

GRanges object should be beforehand tiled to size of 1. This subsetting takes account for strand.

## Usage

```
subsetCoverage(cov, y)
```

## Arguments

cov A coverage object from coverage()

y GRanges object for which coverage should be extracted

364 subsetToFrame

#### Value

numeric vector of coverage of input GRanges object

#### See Also

```
Other features: computeFeatures(), computeFeaturesCage(), countOverlapsW(), disengagementScore(), distToCds(), distToTSS(), entropy(), floss(), fpkm(), fpkm_calc(), fractionLength(), initiationScore(), insideOutsideORF(), isInFrame(), isOverlapping(), kozakSequenceScore(), orfScore(), rankOrder(), ribosomeReleaseScore(), ribosomeStallingScore(), startRegion(), startRegionCoverage(), stopRegion(), translationalEff()
```

subsetToFrame

Subset GRanges to get desired frame.

## **Description**

Usually used for ORFs to get specific frame (0-2): frame 0, frame 1, frame 2

## Usage

```
subsetToFrame(x, frame)
```

## **Arguments**

x A tiled to size of 1 GRanges object

frame A numeric indicating which frame to extract

# **Details**

GRanges object should be beforehand tiled to size of 1. This subsetting takes account for strand.

#### Value

GRanges object reduced to only first frame

# **Examples**

```
subsetToFrame(GRanges("1", IRanges(1:10, width = 1), "+"), 2)
```

sum,covRle-method 365

sum,covRle-method

sum covRle

# Description

Sum coverage per chromosome

# Usage

```
## S4 method for signature 'covRle'
sum(x)
```

# Arguments

Χ

a covRle object

# Value

an integer, sum of coverage per chromosomes in covRle object

 ${\it symbols}$ 

Get ORFik experiment QC folder path

# Description

Get ORFik experiment QC folder path

# Usage

```
symbols(x)
```

# **Arguments**

Х

an ORFik experiment

## Value

a data.table with gene id, gene symbols and tx ids (3 columns)

366 te.plot

```
symbols, experiment-method
```

Get ORFik experiment QC folder path

## **Description**

Get ORFik experiment QC folder path

## Usage

```
## S4 method for signature 'experiment'
symbols(x)
```

#### **Arguments**

Χ

an ORFik experiment

#### Value

a character path

te.plot

Translational efficiency plots

# Description

Create 2 TE plots of:

- Within sample (TE log2 vs mRNA fpkm) ("default")
- Between all combinations of samples (x-axis: rna1fpkm rna2fpkm, y-axis rfp1fpkm rfp2fpkm)

## Usage

```
te.plot(
   df.rfp,
   df.rna,
   output.dir = QCfolder(df.rfp),
   type = c("default", "between"),
   filter.rfp = 1,
   filter.rna = 1,
   collapse = FALSE,
   plot.title = "",
   plot.ext = ".pdf",
   width = 6,
   height = "auto"
)
```

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## **Arguments**

df.rfp	a experiment of Ribo-seq or 80S from TCP-seq.
df.rna	a experiment of RNA-seq
output.dir	directory to save plots, plots will be named "TE_between.pdf" and "TE_within.pdf"
type	which plots to make, default: c("default", "between"). Both plots.
filter.rfp	numeric, default 1. minimum fpkm value to be included in plots
filter.rna	numeric, default 1. minimum fpkm value to be included in plots
collapse	a logical/character (default FALSE), if TRUE all samples within the group SAM-PLE will be collapsed to one. If "all", all groups will be merged into 1 column called merged_all. Collapse is defined as rowSum(elements_per_group) / ncol(elements_per_group)
plot.title	title for plots, usually name of experiment etc
plot.ext	character, default: ".pdf". Alternatives: ".png" or ".jpg".
width	numeric, default 6 (in inches)
height	numeric or character, default "auto", which is: 3 + (ncol(RFP_CDS_FPKM)-2). Else a numeric value of height (in inches)

## **Details**

Ribo-seq and RNA-seq must have equal nrows, with matching samples. Only exception is if RNA-seq is 1 single sample. Then it will use that for each of the Ribo-seq samples. Same stages, conditions etc, with a unique pairing 1 to 1. If not you can run collapse = "all". It will then merge all and do combined of all RNA-seq vs all Ribo-seq

# Value

a data.table with TE values, fpkm and log fpkm values, library samples melted into rows with split variable called "variable".

# **Examples**

```
##
# df.rfp <- read.experiment("zf_baz14_RFP")
# df.rna <- read.experiment("zf_baz14_RNA")
# te.plot(df.rfp, df.rna)
## Collapse replicates:
# te.plot(df.rfp, df.rna, collapse = TRUE)</pre>
```

368 te.table

te.table

Create a TE table

# Description

```
Creates a data.table with 6 columns, column names are: variable, rfp_log2, rna_log2, rna_log10, TE_log2, id
```

## Usage

```
te.table(df.rfp, df.rna, filter.rfp = 1, filter.rna = 1, collapse = FALSE)
```

# Arguments

df.rfp	a experiment of usually Ribo-seq or 80S from TCP-seq. (the numerator of the experiment, usually having a primary role)
df.rna	a experiment of usually RNA-seq. (the denominator of the experiment, usually having a normalizing function)
filter.rfp	numeric, default 1. What is the minimum fpkm value?
filter.rna	numeric, default 1. What is the minimum fpkm value?
collapse	a logical/character (default FALSE), if TRUE all samples within the group SAM-PLE will be collapsed to one. If "all", all groups will be merged into 1 column called merged_all. Collapse is defined as rowSum(elements_per_group) / ncol(elements_per_group)

#### Value

a data.table with 6 columns

# See Also

```
Other\ Differential Expression:\ DEG.plot.static(), DEG\_model(), DTEG.analysis(), DTEG.plot(), \\ te\_rna.plot()
```

# **Examples**

```
df <- ORFik.template.experiment()
df.rfp <- df[df$libtype == "RFP",]
df.rna <- df[df$libtype == "RNA",]
#te.table(df.rfp, df.rna)</pre>
```

te\_rna.plot

te\_rna.plot

Translational efficiency plots

## Description

```
Create TE plot of:
```

- Within sample (TE log2 vs mRNA fpkm)

### Usage

```
te_rna.plot(
   dt,
   output.dir = NULL,
   filter.rfp = 1,
   filter.rna = 1,
   plot.title = "",
   plot.ext = ".pdf",
   width = 6,
   height = "auto",
   dot.size = 0.4,
   xlim = c(filter.rna, filter.rna + 2.5)
)
```

# Arguments

```
dt
                  a data.table with the results from te.table
output.dir
                  a character path, default NULL(no save), or a directory to save to a file will be
                  called "TE_within.pdf"
filter.rfp
                  numeric, default 1. What is the minimum fpkm value?
filter.rna
                  numeric, default 1. What is the minimum fpkm value?
plot.title
                  title for plots, usually name of experiment etc
                  character, default: ".pdf". Alternatives: ".png" or ".jpg".
plot.ext
width
                  numeric, default 6 (in inches)
height
                  a numeric, width of plot in inches. Default "auto".
                  numeric, default 0.4, size of point dots in plot.
dot.size
xlim
                  numeric vector of length 2. X-axis limits. Default: c(filter.rna, filter.rna
                  +2.5)
```

#### Value

```
a ggplot object
```

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### See Also

```
Other DifferentialExpression: DEG.plot.static(), DEG_model(), DTEG.analysis(), DTEG.plot(), te.table()
```

## **Examples**

```
df <- ORFik.template.experiment()
df.rfp <- df[df$libtype == "RFP",]
df.rna <- df[df$libtype == "RNA",]
#dt <- te.table(df.rfp, df.rna)
#te_rna.plot(dt, filter.rfp = 0, filter.rna = 5, dot.size = 1)</pre>
```

tile1

Tile each GRangesList group to 1-base resolution.

## Description

Will tile a GRangesList into single by resolution, each group of the list will be splited by positions of 1. Returned values are sorted as the same groups as the original GRangesList, except they are in by resolutions. This is not supported originally by GenomicRanges for GRangesList.

## Usage

```
tile1(grl, sort.on.return = TRUE, matchNaming = TRUE, is.sorted = TRUE)
```

## **Arguments**

grl	a GRangesList object with names.
sort.on.return	logical (TRUE), should the groups be sorted before return (Negative ranges should be in decreasing order). Makes it a bit slower, but much safer for downstream analysis.
matchNaming	logical (TRUE), should groups keep unlisted names and meta data.(This make the list very big, for $> 100 \rm K$ groups)
is.sorted	logical (TRUE), grl is presorted (negative coordinates are decreasing). Set to FALSE if they are not, else output will most likely be wrong!

#### Value

a GRangesList grouped by original group, tiled to 1. Groups with identical names will be merged.

### See Also

```
Other ExtendGenomicRanges: asTX(), coveragePerTiling(), extendLeaders(), extendTrailers(), reduceKeepAttr(), txSeqsFromFa(), windowPerGroup()
```

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#### **Examples**

tissueNames

Get tissue name variants

## **Description**

Used to standardize nomeclature for experiments.

Example: testis is main naming, but a variant is testicles. testicles will then be renamed to testis.

#### Usage

```
tissueNames()
```

### Value

a data.table with 2 columns, the main name, and all name variants of the main name in second column as a list.

### See Also

```
Other experiment_naming: batchNames(), cellLineNames(), cellTypeNames(), conditionNames(), fractionNames(), inhibitorNames(), libNames(), mainNames(), repNames(), stageNames()
```

TOP.Motif.ecdf

TOP Motif ecdf plot

### Description

Given sequences, DNA or RNA. And some score, scanning efficiency (SE), ribo-seq fpkm, TE etc.

TOP.Motif.ecdf

## Usage

```
TOP.Motif.ecdf(
   seqs,
   rate,
   start = 1,
   stop = max(nchar(seqs)),
   xlim = c("q10", "q99"),
   type = "Scanning efficiency",
   legend.position.1st = c(0.75, 0.28),
   legend.position.motif = c(0.75, 0.28))
```

## **Arguments**

seqs	the sequences (character vector, DNAStringSet), of 5' UTRs (leaders). See example below for input.	
rate	a scoring vector (equal size to seqs)	
start	position in seqs to start at (first is 1), default 1.	
stop	position in seqs to stop at (first is 1), default max(nchar(seqs)), that is the longest sequence length	
×lim	What interval of rate values you want to show type: numeric or quantile of length 2, 1. default c("q10","q99"). bigger than 10 percentile and less than 99 percentile. 2. Set to numeric values, like c(5, 1000), 3. Set to NULL if you want all values. Backend uses coord_cartesian.	
type	What type is the rate scoring? default ("Scanning efficiency")	
legend.position.1st		
	adjust left plot label position, default $c(0.75, 0.28)$ , ("none", "left", "right", "bottom", "top", or two-element numeric vector)	
legend.position.motif		
	adjust right plot label position, default c(0.75, 0.28), ("none", "left", "right", "bottom", "top", or two-element numeric vector)	

### **Details**

Top motif defined as a TSS of C and 4 T's or C's (pyrimidins) downstream of TSS C.

The right plot groups: C nucleotide, TOP motif (C, then 4 pyrimidines) and OTHER (all other TSS variants).

#### Value

```
a ggplot gtable of the TOP motifs in 2 plots
```

# **Examples**

```
## Not run:
if (requireNamespace("BSgenome.Hsapiens.UCSC.hg19")) {
  txdbFile <- system.file("extdata", "hg19_knownGene_sample.sqlite",</pre>
```

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```
package = "GenomicFeatures")
 #Extract sequences of Coding sequences.
 leaders <- loadRegion(txdbFile, "leaders")</pre>
 # Should update by CAGE if not already done
 cageData <- system.file("extdata", "cage-seq-heart.bed.bgz",</pre>
                           package = "ORFik")
 leadersCage <- reassignTSSbyCage(leaders, cageData)</pre>
 # Get region to check
 seqs <- startRegionString(leadersCage, NULL,</pre>
        BSgenome.Hsapiens.UCSC.hg19::Hsapiens, 0, 4)
 # Some toy ribo-seq fpkm scores on cds
 set.seed(3)
 fpkm <- sample(1:115, length(leadersCage), replace = TRUE)</pre>
 # Standard arguments
 TOP.Motif.ecdf(seqs, fpkm, type = "ribo-seq FPKM",
                 legend.position.1st = "bottom",
                 legend.position.motif = "bottom")
 # with no zoom on x-axis:
 TOP.Motif.ecdf(seqs, fpkm, xlim = NULL,
                 legend.position.1st = "bottom",
                 legend.position.motif = "bottom")
}
## End(Not run)
```

topMotif

TOP Motif detection

## **Description**

Per leader, detect if the leader has a TOP motif at TSS (5' end of leader) TOP motif defined as: (C, then 4 pyrimidines)

## Usage

```
topMotif(seqs, start = 1, stop = max(nchar(seqs)), return.sequence = TRUE)
```

### **Arguments**

seqs the sequences (character vector, DNAStringSet), of 5' UTRs (leaders) start re-

gion. seqs must be of minimum widths start - stop + 1 to be included.

See example below for input.

start position in seqs to start at (first is 1), default 1.

stop position in seqs to stop at (first is 1), default max(nchar(seqs)), that is the longest

sequence length

return.sequence

logical, default TRUE, return as data.table with sequence as columns in addition

to TOP class. If FALSE, return character vector.

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#### Value

default: return.sequence == FALSE, a character vector of either TOP, C or OTHER. C means leaders started on C, Other means not TOP and did not start on C. If return.sequence == TRUE, a data.table is returned with the base per position in the motif is included as additional columns (per position called seq1, seq2 etc) and a id column called X.gene\_id (with names of seqs).

## **Examples**

transcriptWindow

Make 100 bases size meta window for all libraries in experiment

### Description

Gives you binned meta coverage plots, either saved seperatly or all in one.

#### Usage

```
transcriptWindow(
  leaders,
  cds,
  trailers,
  df,
  outdir = NULL,
  scores = c("sum", "transcriptNormalized"),
  allTogether = TRUE,
  colors = experiment.colors(df),
  title = "Coverage metaplot",
  windowSize = min(100, min(widthPerGroup(leaders, FALSE)), min(widthPerGroup(cds, FALSE))),
```

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```
returnPlot = is.null(outdir),
  dfr = NULL,
  idName = "",
  plot.ext = ".pdf",
  type = "ofst",
  is.sorted = FALSE,
  drop.zero.dt = TRUE,
  verbose = TRUE,
  force = TRUE,
  library.names = bamVarName(df),
  BPPARAM = bpparam()
)
```

#### **Arguments**

leaders a GRangesList of leaders (5' UTRs)
cds a GRangesList of coding sequences
trailers a GRangesList of trailers (3' UTRs)

df an ORFik experiment

outdir directory to save to (default: NULL, no saving)

scores scoring function (default: c("sum", "transcriptNormalized")), see ?coverageScor-

ings for possible scores.

allTogether plot all coverage plots in 1 output? (defualt: TRUE)

colors Which colors to use, default auto color from function experiment.colors, new

color per library type. Else assign colors yourself.

title title of ggplot

windowSize size of binned windows, default: 100

returnPlot return plot from function, default is.null(outdir), so TRUE if outdir is not de-

fined.

dfr an ORFik experiment of RNA-seq to normalize against. Will add RNA nor-

malized to plot name if this is done.

idName A character ID to add to saved name of plot, if you make several plots in the

same folder, and same experiment, like splitting transcripts in two groups like

targets / nontargets etc. (default: "")

plot.ext character, default: ".pdf". Alternatives: ".png" or ".jpg".

type a character(default: "default"), load files in experiment or some precomputed

variant, like "ofst" or "pshifted". These are made with ORFik:::convertLibs(), shiftFootprintsByExperiment(), etc. Can also be custom user made folders inside the experiments bam folder. It acts in a recursive manner with priority: If you state "pshifted", but it does not exist, it checks "ofst". If no .ofst files, it uses

"default", which always must exists.

Presets are (folder is relative to default lib folder, some types fall back to other

formats if folder does not exist):

- "default": load the original files for experiment, usually bam.

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- "ofst": loads ofst files from the ofst folder, relative to lib folder (falls back to default)

- "pshifted": loads ofst, wig or bigwig from pshifted folder (falls back to ofst, then default)

- "cov": Load covRle objects from cov\_RLE folder (fail if not found)

- "covl": Load covRleList objects, from cov\_RLE\_List folder (fail if not found)

- "bed": Load bed files, from bed folder (falls back to default)

- Other formats must be loaded directly with fimport

is.sorted logical (FALSE), is grl sorted. That is + strand groups in increasing ranges

(1,2,3), and - strand groups in decreasing ranges (3,2,1)

drop.zero.dt logical FALSE, if TRUE and as.data.table is TRUE, remove all 0 count posi-

tions. This greatly speeds up and most importantly, greatly reduces memory usage. Will not change any plots, unless 0 positions are used in some sense.

(mean, median, zscore coverage will only scale differently)

verbose logical, default TRUE, message about library output status.

force logical, default TRUE If TRUE, reload library files even if matching named

variables are found in environment used by experiment (see envExp) A simple way to make sure correct libraries are always loaded. FALSE is faster if data is

loaded correctly already.

library.names character vector, names of libraries, default: name\_decider(df, naming)

BPPARAM how many cores/threads to use? default: bpparam()

### Value

NULL, or ggplot object if returnPlot is TRUE

# See Also

Other experiment plots: transcriptWindow1(), transcriptWindowPer()

### **Examples**

```
df <- ORFik.template.experiment()[3,] # Only third library
loadRegions(df) # Load leader, cds and trailers as GRangesList
#transcriptWindow(leaders, cds, trailers, df, outdir = "directory/to/save")</pre>
```

transcriptWindow1

Meta coverage over all transcripts

### Description

Given as single window

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### Usage

```
transcriptWindow1(
   df,
   outdir = NULL,
   scores = c("sum", "zscore"),
   colors = experiment.colors(df),
   title = "Coverage metaplot",
   windowSize = 100,
   returnPlot = is.null(outdir),
   dfr = NULL,
   idName = "",
   plot.ext = ".pdf",
   type = "ofst",
   drop.zero.dt = drop.zero.dt,
   BPPARAM = bpparam()
)
```

### **Arguments**

df an ORFik experiment

outdir directory to save to (default: NULL, no saving)

scores scoring function (default: c("sum", "transcriptNormalized")), see ?coverageScor-

ings for possible scores.

colors Which colors to use, default auto color from function experiment . colors, new

color per library type. Else assign colors yourself.

title title of ggplot

windowSize size of binned windows, default: 100

returnPlot return plot from function, default is.null(outdir), so TRUE if outdir is not de-

fined.

dfr an ORFik experiment of RNA-seq to normalize against. Will add RNA nor-

malized to plot name if this is done.

idName A character ID to add to saved name of plot, if you make several plots in the

same folder, and same experiment, like splitting transcripts in two groups like

targets / nontargets etc. (default: "")

plot.ext character, default: ".pdf". Alternatives: ".png" or ".jpg".

type a character(default: "default"), load files in experiment or some precomputed

variant, like "ofst" or "pshifted". These are made with ORFik:::convertLibs(), shiftFootprintsByExperiment(), etc. Can also be custom user made folders inside the experiments bam folder. It acts in a recursive manner with priority: If you state "pshifted", but it does not exist, it checks "ofst". If no .ofst files, it uses

"default", which always must exists.

Presets are (folder is relative to default lib folder, some types fall back to other formats if folder does not exist):

- "default": load the original files for experiment, usually bam.

- "ofst": loads ofst files from the ofst folder, relative to lib folder (falls back to

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default)

- "pshifted": loads ofst, wig or bigwig from pshifted folder (falls back to ofst, then default)

- "cov": Load covRle objects from cov\_RLE folder (fail if not found)

- "covl": Load covRleList objects, from cov\_RLE\_List folder (fail if not found)

- "bed": Load bed files, from bed folder (falls back to default)

- Other formats must be loaded directly with fimport

drop.zero.dt

logical FALSE, if TRUE and as.data.table is TRUE, remove all 0 count positions. This greatly speeds up and most importantly, greatly reduces memory usage. Will not change any plots, unless 0 positions are used in some sense. (mean, median, zscore coverage will only scale differently)

**BPPARAM** 

how many cores/threads to use? default: bpparam()

#### Value

NULL, or ggplot object if returnPlot is TRUE

#### See Also

Other experiment plots: transcriptWindow(), transcriptWindowPer()

transcriptWindowPer

Helper function for transcriptWindow

## **Description**

Make 100 bases size meta window for one library in experiment

### Usage

```
transcriptWindowPer(
  leaders,
  cds,
  trailers,
  df,
  outdir = NULL,
  scores = c("sum", "zscore"),
  reads,
  returnCoverage = FALSE,
  windowSize = 100,
  drop.zero.dt = TRUE,
  BPPARAM = bpparam()
)
```

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### **Arguments**

leaders a GRangesList of leaders (5' UTRs)

cds a GRangesList of coding sequences

trailers a GRangesList of trailers (3' UTRs)

df an ORFik experiment

outdir directory to save to (default: NULL, no saving)

scores scoring function (default: c("sum", "transcriptNormalized")), see ?coverageScor-

ings for possible scores.

reads a GRanges / GAligment object of reads, can also be a list of those.

returnCoverage return data.table with coverage (default: FALSE)

windowSize size of binned windows, default: 100

drop.zero.dt logical FALSE, if TRUE and as.data.table is TRUE, remove all 0 count posi-

tions. This greatly speeds up and most importantly, greatly reduces memory usage. Will not change any plots, unless 0 positions are used in some sense.

(mean, median, zscore coverage will only scale differently)

BPPARAM how many cores/threads to use? default: bpparam()

#### **Details**

Gives you binned meta coverage plots, either saved seperatly or all in one.

#### Value

NULL, or ggplot object if returnPlot is TRUE

### See Also

Other experiment plots: transcriptWindow(), transcriptWindow1()

translationalEff Translational efficiency

## Description

Uses RnaSeq and RiboSeq to get translational efficiency of every element in 'grl'. Translational efficiency is defined as:

(density of RPF within ORF) / (RNA expression of ORFs transcript)

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#### Usage

```
translationalEff(
  grl,
  RNA,
  RFP,
  tx,
  with.fpkm = FALSE,
  pseudoCount = 0,
  librarySize = "full",
  weight.RFP = 1L,
  weight.RNA = 1L
)
```

#### Arguments

grl	a GRangesList object can be either transcripts, 5' utrs, cds', 3' utrs or ORFs as
	a special case (uORFs, potential new cds' etc). If regions are not spliced you

can send a GRanges object.

RNA RnaSeq reads as GAlignments, GRanges or GRangesList object

RFP RiboSeq reads as GAlignments, GRanges or GRangesList object

tx a GRangesList of the transcripts. If you used cage data, then the tss for the the

leaders have changed, therefor the tx lengths have changed. To account for that call: 'translationalEff(grl, RNA, RFP, tx = extendLeaders(tx, cageFiveUTRs))

' where cageFiveUTRs are the reannotated by CageSeq data leaders.

with.fpkm logical, default: FALSE, if true return the fpkm values together with transla-

tional efficiency as a data.table

pseudoCount a numeric, default 0, set it to 1 if you want to avoid NA and inf values.

librarySize either numeric value or character vector. Default ("full"), number of align-

ments in library (reads). If you just have a subset, you can give the value by librarySize = length(wholeLib) or sum(wholeLib\$score), if you want lib size to be only number of reads overlapping grl, do: librarySize = "overlapping" sum(countOverlaps(reads, grl) > 0), if reads[1] has 3 hits in grl, and reads[2] has 2 hits, librarySize will be 2, not 5. You can also get the inverse overlap, if you want lib size to be total number of overlaps, do: librarySize = "DESeq" This is standard fpkm way of DESeq2::fpkm(robust = FALSE) sum(countOverlaps(grl,

reads)) if grl[1] has 3 reads and grl[2] has 2 reads, librarySize is 5, not 2.

weight.RFP a vector (default: 1L). Can also be character name of column in RFP. As in trans-

lationalEff(weight = "score") for: GRanges("chr1", 1, "+", score = 5), would

mean score column tells that this alignment region was found 5 times.

weight.RNA Same as weightRFP but for RNA weights. (default: 1L)

### Value

a numeric vector of fpkm ratios, if with fpkm is TRUE, return a data table with te and fpkm values (total 3 columns then)

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#### References

```
doi: 10.1126/science.1168978
```

#### See Also

```
Other features: computeFeatures(), computeFeaturesCage(), countOverlapsW(), disengagementScore(), distToCds(), distToTSS(), entropy(), floss(), fpkm(), fpkm_calc(), fractionLength(), initiationScore(), insideOutsideORF(), isInFrame(), isOverlapping(), kozakSequenceScore(), orfScore(), rankOrder(), ribosomeReleaseScore(), ribosomeStallingScore(), startRegion(), startRegionCoverage(), stopRegion(), subsetCoverage()
```

### **Examples**

trimming.table

Create trimming table

### **Description**

From fastp runs in ORFik alignment process

### Usage

```
trimming.table(
  trim_folder,
  raw_libraries = dir(trim_folder, "\\.json", full.names = TRUE)
)
```

### **Arguments**

```
folder folder folder of trimmed files, only reads fastp .json files. Can be NULL if raw_libraries
is defined

raw_libraries character, default: dir(trim_folder, "\.json", full.names = TRUE), file paths
of all json file paths.
```

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#### Value

a data.table with 6 columns, raw\_library (names of library), raw\_reads (numeric, number of raw reads), trim\_reads (numeric, number of trimmed reads), raw\_mean\_length (numeric, raw mean read length), trim\_mean\_length (numeric, trim mean read length).

## **Examples**

```
# Location of fastp trimmed .json files
trimmed_file <- system.file("extdata/test_processing/trim",
   "output_template.json", package = "ORFik")
trimmed_folder <- dirname(trimmed_file)
trimming.table(trimmed_folder)
trimming.table(NULL, trimmed_file)</pre>
```

trim\_detection

Add trimming info to QC report

### **Description**

Only works if alignment was done using ORFik with STAR.

#### Usage

```
trim_detection(df, finals, alignment_folder = libFolder(df, "unique"))
```

#### **Arguments**

### Value

a data.table of the update finals object with trim info

txNames 383

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Get transcript names from orf names

## Description

Using the ORFik definition of orf name, which is: example ENSEMBL:

tx name: ENST0909090909090 orf id: \_1 (the first of on that tx) orf\_name: ENST0909090909090\_1

So therefor txNames("ENST09090909090901") = ENST09090909090

#### Usage

```
txNames(grl, ref = NULL, unique = FALSE)
```

#### **Arguments**

grl a GRangesList grouped by ORF, GRanges object or IRanges object.

ref a reference GRangesList. The object you want grl to subset by names. Add to

make sure naming is valid.

unique a boolean, if true unique the names, used if several orfs map to same transcript

and you only want the unique groups

#### **Details**

The names must be extracted from a column called names, or the names of the grl object. If it is already tx names, it returns the input

NOTE! Do not use \_123 etc in end of transcript names if it is not ORFs. Else you will get errors. Just \_ will work, but if transcripts are called ENST\_123124124000 etc, it will crash, so substitute "\_" with "." gsub("\_", ".", names)

# Value

a character vector of transcript names, without \_\* naming

#### See Also

```
Other ORFHelpers: defineTrailer(), longestORFs(), mapToGRanges(), orfID(), startCodons(), startSites(), stopCodons(), stopSites(), uniqueGroups(), uniqueOrder()
```

384 txNamesToGeneNames

#### **Examples**

```
 \begin{split} \text{gr\_plus} <& \text{- GRanges}(\text{seqnames} = \text{c("chr1", "chr1")}, \\ & \text{ranges} = \text{IRanges}(\text{c(7, 14), width} = 3), \\ & \text{strand} = \text{c("+", "+")}) \\ \text{gr\_minus} <& \text{- GRanges}(\text{seqnames} = \text{c("chr2", "chr2")}, \\ & \text{ranges} = \text{IRanges}(\text{c(4, 1), c(9, 3)}), \\ & \text{strand} = \text{c("-", "-")}) \\ \text{grl} <& \text{- GRangesList}(\text{tx1\_1} = \text{gr\_plus, tx2\_1} = \text{gr\_minus}) \\ \text{\# there are 2 orfs, both the first on each transcript} \\ \text{txNames}(\text{grl}) \end{aligned}
```

txNamesToGeneNames

Convert transcript names to gene names

# Description

Works for ensembl, UCSC and other standard annotations.

#### Usage

```
txNamesToGeneNames(txNames, txdb)
```

## Arguments

txNames character vector, the transcript names to convert. Can also be a named object

with tx names (like a GRangesList), will then extract names.

txdb the transcript database to use or gtf/gff path to it.

### Value

character vector of gene names

# **Examples**

```
df <- ORFik.template.experiment()
txdb <- loadTxdb(df)
loadRegions(txdb, "cds") # using tx names
txNamesToGeneNames(cds, txdb)
# Identical to:
loadRegions(txdb, "cds", by = "gene")</pre>
```

txSeqsFromFa 385

txSeqsFromFa	Get transcript sequence from a GRangesList and a faFile or BSgenome

# Description

For each GRanges object, find the sequence of it from faFile or BSgenome.

## Usage

```
txSeqsFromFa(grl, faFile, is.sorted = FALSE, keep.names = TRUE)
```

## **Arguments**

grl	a GRangesList object
faFile	FaFile, BSgenome, fasta/index file path or an ORFik experiment. This file is usually used to find the transcript sequences from some GRangesList.
is.sorted	a speedup, if you know the grl ranges are sorted
keep.names	a logical, default (TRUE), if FALSE: return as character vector without names.

### **Details**

A wrapper around extractTranscriptSeqs that works for DNAStringSet and ORFik experiment input. For debug of errors do: which(!(unique(seqnamesPerGroup(grl, FALSE))) This happens usually when the grl contains chromsomes that the fasta file does not have. A normal error is that mitocondrial chromosome is called MT vs chrM even though they have same seqlevelsStyle. The above line will give you which chromosome it is missing.

## Value

```
a DNAStringSet of the transcript sequences
```

#### See Also

```
Other ExtendGenomicRanges: asTX(), coveragePerTiling(), extendLeaders(), extendTrailers(), reduceKeepAttr(), tile1(), windowPerGroup()
```

386 uniqueOrder

uniqueGroups

Get the unique set of groups in a GRangesList

# Description

Sometimes GRangesList groups might be identical, for example ORFs from different isoforms can have identical ranges. Use this function to reduce these groups to unique elements in GRangesList grl, without names and metacolumns.

### Usage

```
uniqueGroups(grl)
```

#### Arguments

```
grl a GRangesList
```

#### Value

a GRangesList of unique orfs

#### See Also

```
Other ORFHelpers: defineTrailer(), longestORFs(), mapToGRanges(), orfID(), startCodons(), startSites(), stopCodons(), stopSites(), txNames(), uniqueOrder()
```

## **Examples**

```
gr1 <- GRanges("1", IRanges(1,10), "+")
gr2 <- GRanges("1", IRanges(20, 30), "+")
# make a grl with duplicated ORFs (gr1 twice)
grl <- GRangesList(tx1_1 = gr1, tx2_1 = gr2, tx3_1 = gr1)
uniqueGroups(grl)</pre>
```

uniqueOrder

Get unique ordering for GRangesList groups

## Description

This function can be used to calculate unique numerical identifiers for each of the GRangesList elements. Elements of GRangesList are unique when the GRanges inside are not duplicated, so ranges differences matter as well as sorting of the ranges.

### Usage

```
uniqueOrder(grl)
```

unlistGrl 387

# **Arguments**

```
grl a GRangesList
```

#### Value

an integer vector of indices of unique groups

#### See Also

```
uniqueGroups
```

```
Other ORFHelpers: defineTrailer(), longestORFs(), mapToGRanges(), orfID(), startCodons(), startSites(), stopCodons(), stopSites(), txNames(), uniqueGroups()
```

## **Examples**

```
gr1 <- GRanges("1", IRanges(1,10), "+")
gr2 <- GRanges("1", IRanges(20, 30), "+")
# make a grl with duplicated ORFs (gr1 twice)
grl <- GRangesList(tx1_1 = gr1, tx2_1 = gr2, tx3_1 = gr1)
uniqueOrder(grl) # remember ordering

# example on unique ORFs
uniqueORFs <- uniqueGroups(grl)
# now the orfs are unique, let's map back to original set:
reMappedGrl <- uniqueORFs[uniqueOrder(grl)]</pre>
```

unlistGrl

Safe unlist

# Description

Same as [AnnotationDbi::unlist2()], keeps names correctly. Two differences is that if grl have no names, it will not make integer names, but keep them as null. Also if the GRangesList has names , and also the GRanges groups, then the GRanges group names will be kept.

### Usage

```
unlistGrl(grl)
```

## **Arguments**

grl

a GRangesList

### Value

```
a GRanges object
```

388 uORFSearchSpace

### **Examples**

unlistToExtremities

Get flanks as GRanges

## Description

For a GRangesList, get start and end site, return back as GRanges.

# Usage

```
unlistToExtremities(grl)
```

### **Arguments**

```
grl a GRangesList
```

#### Value

a GRangs object with meta column "group", which gives

uORFSearchSpace

Create search space to look for uORFs

## **Description**

Given a GRangesList of 5' UTRs or transcripts, reassign the start sites using max peaks from CageSeq data (if CAGE is given). A max peak is defined as new TSS if it is within boundary of 5' leader range, specified by 'extension' in bp. A max peak must also be higher than minimum CageSeq peak cutoff specified in 'filterValue'. The new TSS will then be the positioned where the cage read (with highest read count in the interval). If you want to include uORFs going into the CDS, add this argument too.

uORFSearchSpace 389

#### Usage

```
uORFSearchSpace(
  fiveUTRs,
  cage = NULL,
  extension = 1000,
  filterValue = 1,
  restrictUpstreamToTx = FALSE,
  removeUnused = FALSE,
  cds = NULL
)
```

#### **Arguments**

fiveUTRs (GRangesList) The 5' leaders or full transcript sequences

cage Either a filePath for the CageSeq file as .bed .bam or .wig, with possible com-

pressions (".gzip", ".gz", ".bgz"), or already loaded CageSeq peak data as GRanges or GAlignment. NOTE: If it is a .bam file, it will add a score column by running: convertToOneBasedRanges(cage, method = "5prime", addScoreColumn = TRUE) The score column is then number of replicates of read, if score column

is something else, like read length, set the score column to NULL first.

extension The maximum number of basses upstream of the TSS to search for CageSeq

peak.

filterValue The minimum number of reads on cage position, for it to be counted as possible

new tss. (represented in score column in CageSeq data) If you already filtered,

set it to 0.

restrictUpstreamToTx

a logical (FALSE). If TRUE: restrict leaders to not extend closer than 5 bases

from closest upstream leader, set this to TRUE.

removeUnused logical (FALSE), if False: (standard is to set them to original annotation), If

TRUE: remove leaders that did not have any cage support.

cds (GRangesList) CDS of relative fiveUTRs, applicable only if you want to extend

5' leaders downstream of CDS's, to allow upstream ORFs that can overlap into

CDS's.

#### Value

a GRangesList of newly assigned TSS for fiveUTRs, using CageSeq data.

## See Also

```
Other uorfs: addCdsOnLeaderEnds(), filterUORFs(), removeORFsWithSameStartAsCDS(), removeORFsWithSameStopAremoveORFsWithStartInsideCDS(), removeORFsWithinCDS()
```

# Examples

```
# example 5' leader, notice exon_rank column
fiveUTRs <- GenomicRanges::GRangesList(</pre>
```

390 updateTxdbStartSites

updateTxdbRanks

Update exon ranks of exon data.frame inside txdb object

# Description

Update exon ranks of exon data.frame inside txdb object

## Usage

```
updateTxdbRanks(exons)
```

#### **Arguments**

exons

a data.frame, call of as.list(txdb)\$splicings

## Value

a data.frame, modified call of as.list(txdb)

# Description

Update start sites of leaders

### Usage

```
updateTxdbStartSites(txList, fiveUTRs, removeUnused)
```

### **Arguments**

txList a list, call of as.list(txdb)
fiveUTRs a GRangesList of 5' leaders

removeUnused logical (FALSE), remove leaders that did not have any cage support. (standard

is to set them to original annotation)

#### Value

a list, modified call of as.list(txdb)

upstreamFromPerGroup Get rest of objects upstream (inclusive)

## **Description**

Per group get the part upstream of position. upstreamFromPerGroup(tx, stopSites(fiveUTRs, asGR = TRUE)) will return the 5' utrs per transcript as GRangesList, usually used for interesting parts of the transcripts.

## Usage

```
upstreamFromPerGroup(tx, upstreamFrom)
```

## **Arguments**

tx a GRangesList, usually of Transcripts to be changed

upstreamFrom a vector of integers, for each group in tx, where is the new start point of first

valid exon.

#### **Details**

If you don't want to include the points given in the region, use upstreamOfPerGroup

#### Value

a GRangesList of upstream part

#### See Also

 $Other\ GRanges:\ assign First Exons Start Site(), assign Last Exons Stop Site(), downstream From Per Group(), downstream Of Per Group(), upstream Of Per Group()$ 

upstreamOfPerGroup

Get rest of objects upstream (exclusive)

# Description

Per group get the part upstream of position upstreamOfPerGroup(tx, startSites(cds, asGR = TRUE)) will return the 5' utrs per transcript, usually used for interesting parts of the transcripts.

# Usage

```
upstreamOfPerGroup(
   tx,
   upstreamOf,
   allowOutside = TRUE,
   is.circular = all(isCircular(tx) %in% TRUE)
)
```

## **Arguments**

tx	a GRangesList, usually of Transcripts to be changed
upstreamOf	a vector of integers, for each group in tx, where is the the base after the new stop point of last valid exon.
allowOutside	a logical (T), can upstreamOf extend outside range of tx, can set boundary as a false hit, so beware.
is.circular	logical, default FALSE if not any is: all(isCircular(grl) Where grl is the ranges checked. If TRUE, allow ranges to extend below position 1 on chromosome. Since circular genomes can have negative coordinates.

## Value

a GRangesList of upstream part

#### See Also

 $Other \ GRanges: assign First Exons Start Site(), assign Last Exons Stop Site(), downstream From Per Group(), downstream Great Group(), upstream From Per Group(), upstream From Per$ 

validateExperiments 393

validateExperiments

Validate ORFik experiment

## **Description**

Check for valid existing, non-empty and all unique. A good way to see if your experiment is valid.

## Usage

```
validateExperiments(df, library.names = bamVarName(df))
```

### **Arguments**

df an ORFik experiment
library.names character vector, names of libraries, default: name\_decider(df, naming)

#### Value

NULL (Stops if failed)

#### See Also

Other ORFik\_experiment: ORFik.template.experiment(), ORFik.template.experiment.zf(), bamVarName(), create.experiment(), experiment-class, filepath(), libraryTypes(), organism, experiment-methoutputLibs(), read.experiment(), save.experiment()

validGRL

Helper Function to check valid GRangesList input

## **Description**

Helper Function to check valid GRangesList input

### Usage

```
validGRL(class, type = "grl", checkNULL = FALSE)
```

### **Arguments**

class as character vector the given class of supposed GRangesList object

type a character vector, is it gtf, cds, 5', 3', for messages.

checkNULL should NULL classes be checked and return indeces of these?

### Value

```
either NULL or indices (checkNULL == TRUE)
```

394 widthPerGroup

#### See Also

```
Other validity: checkRFP(), checkRNA(), is.ORF(), is.gr_or_grl(), is.grl(), is.range(), validSeqlevels()
```

validSeqlevels

Helper function to find overlaping seglevels

### **Description**

Keep only seqnames in reads that are in grl Useful to avoid seqname warnings in bioC

### Usage

```
validSeqlevels(grl, reads)
```

## **Arguments**

grl a GRangesList or GRanges object

reads a GRanges, GAlignment or GAlignmentPairs object

#### Value

a character vector of valid seqlevels

#### See Also

```
Other validity: checkRFP(), checkRNA(), is.ORF(), is.gr_or_grl(), is.grl(), is.range(), validGRL()
```

widthPerGroup

Get list of widths per granges group

# Description

Get list of widths per granges group

## Usage

```
widthPerGroup(grl, keep.names = TRUE)
```

## Arguments

grl a GRangesList

keep.names a boolean, keep names or not, default: (TRUE)

windowCoveragePlot 395

## Value

an integer vector (named/unnamed) of widths

## **Examples**

```
\begin{split} \text{gr\_plus} <& - \text{GRanges}(\text{seqnames} = \text{c("chr1", "chr1")}, \\ & \text{ranges} = \text{IRanges}(\text{c(7, 14), width} = 3), \\ & \text{strand} = \text{c("+", "+")}) \\ \text{gr\_minus} <& - \text{GRanges}(\text{seqnames} = \text{c("chr2", "chr2")}, \\ & \text{ranges} = \text{IRanges}(\text{c(4, 1), c(9, 3)}), \\ & \text{strand} = \text{c("-", "-")}) \\ \text{grl} <& - \text{GRangesList}(\text{tx1} = \text{gr\_plus}, \text{tx2} = \text{gr\_minus}) \\ \text{widthPerGroup}(\text{grl}) \end{split}
```

windowCoveragePlot

Get meta coverage plot of reads

## **Description**

Spanning a region like a transcripts, plot how the reads distribute.

## Usage

```
windowCoveragePlot(
  coverage,
  output = NULL,
  scoring = "zscore",
  colors = c("skyblue4", "orange"),
  title = "Coverage metaplot",
  type = "transcripts",
  scaleEqual = FALSE,
  setMinToZero = FALSE
)
```

# Arguments

coverage	a data.table, e.g. output of scaledWindowCoverage
output	character string (NULL), if set, saves the plot as pdf or png to path given. If no format is given, is save as pdf.
scoring	character vector, default "zscore", either of zscore, transcriptNormalized, sum, mean, median, or NULL. Set NULL if already scored. see ?coverageScorings for info and more alternatives.
colors	character vector colors to use in plot, will fix automaticly, using binary splits with colors c('skyblue4', 'orange').
title	a character (metaplot) (what is the title of plot?)

type a character (transcripts), what should legends say is the whole region? Tran-

scripts, genes, non coding rnas etc.

scaleEqual a logical (FALSE), should all fractions (rows), have same max value, for easy

comparison of max values if needed.

setMinToZero a logical (FALSE), should minimum y-value be 0 (TRUE). With FALSE mini-

mum value is minimum score at any position. This parameter overrides scaleE-

qual.

#### **Details**

If coverage has a column called feature, this can be used to subdivide the meta coverage into parts as (5' UTRs, cds, 3' UTRs) These are the columns in the plot. The fraction column divide sequence libraries. Like ribo-seq and rna-seq. These are the rows of the plot. If you return this function without assigning it and output is NULL, it will automaticly plot the figure in your session. If output is assigned, no plot will be shown in session. NULL is returned and object is saved to output.

Colors: Remember if you want to change anything like colors, just return the ggplot object, and reassign like: obj + scale\_color\_brewer() etc.

#### Value

a ggplot object of the coverage plot, NULL if output is set, then the plot will only be saved to location.

#### See Also

```
Other coveragePlot: coverageHeatMap(), pSitePlot(), savePlot()
```

### **Examples**

windowPerGroup 397

# Description

Per GRanges input (gr) of single position inputs (center point), create a GRangesList window output of specified upstream, downstream region relative to some transcript "tx".

If downstream is 20, it means the window will start 20 downstream of gr start site (-20 in relative transcript coordinates.) If upstream is 20, it means the window will start 20 upstream of gr start site (+20 in relative transcript coordinates.) It will keep exon structure of tx, so if -20 is on next exon, it jumps to next exon.

### Usage

```
windowPerGroup(gr, tx, upstream = 0L, downstream = 0L)
```

## **Arguments**

gr	a GRanges/IRanges object (startSites or others, must be single point per in genomic coordinates)
tx	a $GRangesList$ of transcripts or (container region), names of tx must contain all gr names. The names of gr can also be the ORFik orf names. that is "tx-Name_id".
upstream	an integer, default $(0)$ , relative region to get upstream end from. $(0 \text{ means start site}, +1 \text{ is one upstream}, -1 \text{ is one downstream})$
downstream	an integer, default (0), relative region to get downstream end from (0 means start site, +1 is one downstream, -1 is one upstream)

#### **Details**

If a region has a part that goes out of bounds, E.g if you try to get window around the CDS start site, goes longer than the 5' leader start site, it will set start to the edge boundary (the TSS of the transcript in this case). If region has no hit in bound, a width 0 GRanges object is returned. This is useful for things like countOverlaps, since 0 hits will then always be returned for the correct object index. If you don't want the 0 width windows, use reduce() to remove 0-width windows.

# Value

a GRanges, or GRangesList object if any group had > 1 exon.

## See Also

```
Other ExtendGenomicRanges: asTX(), coveragePerTiling(), extendLeaders(), extendTrailers(), reduceKeepAttr(), tile1(), txSeqsFromFa()
```

## **Examples**

```
# find 2nd codon of an ORF on a spliced transcript
ORF <- GRanges("1", c(3), "+") # start site
names(ORF) <- "tx1_1" # ORF 1 on tx1
tx <- GRangesList(tx1 = GRanges("1", c(1,3,5,7,9,11,13), "+"))
windowPerGroup(ORF, tx, upstream = 0, downstream = 0) # <- TIS
windowPerGroup(ORF, tx, upstream = 0, downstream = 1) # <- first and second base
windowPerGroup(ORF, tx, upstream = -1, downstream = 1) # <- second base
# find 2nd codon of an ORF on a spliced transcript
windowPerGroup(ORF, tx, upstream = -3, downstream = 5) # <- 2nd codon
# With multiple extensions downstream
ORF <- rep(ORF, 2)
names(ORF)[2] <- "tx1_2"
windowPerGroup(ORF, tx, upstream = 0, downstream = c(2, 5))
# The last one gives 2nd for first ORF and (1st and 2nd) codon for
# second ORF, returned as two groups of class GRanges/GRangsList</pre>
```

windowPerReadLength Find proportion of reads per position per read length in window

# Description

This is defined as: Fraction of reads per read length, per position in whole window (defined by upstream and downstream) If tx is not NULL, it gives a metaWindow, centered around startSite of grl from upstream and downstream. If tx is NULL, it will use only downstream, since it has no reference on how to find upstream region. The exception is when upstream is negative, that is, going into downstream region of the object.

#### Usage

#### **Arguments**

grl a GRangesList object with usually either leaders, cds', 3' utrs or ORFs

tx default NULL, a GRangesList of transcripts or (container region), names of tx

must contain all grl names. The names of grl can also be the ORFik orf names.

that is "txName\_id"

reads a GAlignments, GRanges, or precomputed coverage as covRleList (where

names of covRle objects are readlengths) of RiboSeq, RnaSeq etc.

Weigths for scoring is default the 'score' column in 'reads'. Can also be random access paths to bigWig or fstwig file. Do not use random access for more than a

few genes, then loading the entire files is usually better.

pShifted a logical (TRUE), are Ribo-seq reads p-shifted to size 1 width reads? If upstream

and downstream is set, this argument is irrelevant. So set to FALSE if this is not

p-shifted Ribo-seq.

upstream an integer (5), relative region to get upstream from. Default: ifelse(!is.null(tx),

ifelse(pShifted, 5, 20), min(ifelse(pShifted, 5, 20), 0))

downstream an integer (20), relative region to get downstream from. Default: ifelse(pShifted,

20, 5)

acceptedLengths

an integer vector (NULL), the read lengths accepted. Default NULL, means all

lengths accepted.

zeroPosition an integer DEFAULT (upstream), what is the center point? Like leaders and cds

combination, then 0 is the TIS and -1 is last base in leader. NOTE!: if windows

have different widths, this will be ignored.

scoring a character (transcriptNormalized), which meta coverage scoring? one of (zs-

core, transcriptNormalized, mean, median, sum, sumLength, fracPos), see ?coverageScorings for more info. Use to decide a scoring of hits per position for metacoverage etc. Set to NULL if you do not want meta coverage, but instead

want per gene per position raw counts.

weight (default: 'score'), if defined a character name of valid meta column in subject.

GRanges("chr1", 1, "+", score = 5), would mean score column tells that this alignment region was found 5 times. Formats which loads a score column like this: Bigwig, wig, ORFik ofst, collapsed bam, bedoc and .bedo. As do CAGEr CAGE files and many other package formats. You can also assign a score col-

umn manually.

drop.zero.dt logical FALSE, if TRUE and as.data.table is TRUE, remove all 0 count posi-

tions. This greatly speeds up and most importantly, greatly reduces memory usage. Will not change any plots, unless 0 positions are used in some sense.

(mean, median, zscore coverage will only scale differently)

append.zeroes logical, default FALSE. If TRUE and drop.zero.dt is TRUE and all windows

have equal length, it will add back 0 values after transformation. Sometimes needed for correct plots, if TRUE, will call abort if not all windows are equal

length!

windows the GRangesList windows to actually check, default: startRegion(grl, tx,

TRUE, upstream, downstream).

#### **Details**

Careful when you create windows where not all transcripts are long enough, this function usually is used first with filterTranscripts to make sure they are of all of valid length!

#### Value

a data.table with 4 columns: position (in window), score, fraction (read length). If score is NULL, will also return genes (index of grl). A note is that if no coverage is found, it returns an empty data.table.

#### See Also

Other coverage: coverageScorings(), metaWindow(), regionPerReadLength(), scaledWindowPositions()

## **Examples**

```
cds <- GRangesList(tx1 = GRanges("1", 100:129, "+"))
tx <- GRangesList(tx1 = GRanges("1", 80:129, "+"))
reads <- GRanges("1", seq(79,129, 3), "+")
windowPerReadLength(cds, tx, reads, scoring = "sum")
windowPerReadLength(cds, tx, reads, scoring = "transcriptNormalized")</pre>
```

windowPerTranscript

Get a binned coverage window per transcript

#### **Description**

Per transcript (or other regions), bin them all to windowSize (default 100), and make a data.table, rows are positions, useful for plotting with ORFik and ggplot2.

## Usage

```
windowPerTranscript(
  txdb,
  reads,
  splitIn3 = TRUE,
  windowSize = 100,
  fraction = "1",
  weight = "score",
  drop.zero.dt = FALSE,
  BPPARAM = bpparam()
)
```

xAxisScaler 401

#### **Arguments**

txdb a TxDb object or a path to gtf/gff/db file.

reads GRanges or GAlignment of reads

splitIn3 a logical(TRUE), split window in 3 (leader, cds, trailer)

windowSize an integer (100), size of windows (columns). All genes with region smaller than

this size are filter out for metacoverage.

fraction a character (1), info on reads (which read length, or which type (RNA seq)) (row

names)

weight (default: 'score'), if defined a character name of valid meta column in subject.

GRanges("chr1", 1, "+", score = 5), would mean score column tells that this alignment region was found 5 times. Formats which loads a score column like this: Bigwig, wig, ORFik ofst, collapsed bam, bedoc and .bedo. As do CAGEr CAGE files and many other package formats. You can also assign a score col-

umn manually.

drop.zero.dt logical FALSE, if TRUE and as.data.table is TRUE, remove all 0 count posi-

tions. This greatly speeds up and most importantly, greatly reduces memory usage. Will not change any plots, unless 0 positions are used in some sense.

(mean, median, zscore coverage will only scale differently)

BPPARAM how many cores/threads to use? default: bpparam()

#### **Details**

NOTE: All ranges with smaller width than windowSize, will of course be removed. What is the 100th position on a 1 width object?

#### Value

a data.table with columns position, score

xAxisScaler Scale x axis correctly

#### **Description**

Works for all coverage plots, that need 0 position aligning

## Usage

xAxisScaler(covPos)

#### **Arguments**

covPos a numeric vector of positions in coverage

402 yAxisScaler

## **Details**

It basicly bins the x axis on floor(length of x axis / 20) or 1 if x < 20

#### Value

a numeric vector from the seq() function, aligned to 0.

yAxisScaler

Scale y axis correctly

# Description

Works for all coverage plots.

# Usage

```
yAxisScaler(covPos, increments.y = "auto")
```

# Arguments

covPos a levels object from a factor of y axis

increments.y increments of y axis, default "auto". Or a numeric value < max position & >

min position.

#### Value

a character vector from the seq() function, aligned to 0.

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