

Package ‘multicrispr’

October 18, 2022

Title Multi-locus multi-purpose Crispr/Cas design

Version 1.6.0

Encoding UTF-8

Description This package is for designing Crispr/Cas9 and Prime Editing experiments.

It contains functions to (1) define and transform genomic targets, (2) find spacers
(4) count offtarget (mis)matches, and (5) compute Doench2016/2014 targeting efficiency.
Care has been taken for multicrispr to scale well towards large target sets,
enabling the design of large Crispr/Cas9 libraries.

License GPL-2

LazyData true

RoxygenNote 7.1.2

Depends R (>= 4.0)

Imports assertive, BiocGenerics, Biostrings, BSgenome, CRISPRseek,
data.table, GenomeInfoDb, GenomicFeatures, GenomicRanges,
ggplot2, grid, karyoploteR, magrittr, methods, parallel,
plyranges, Rbowtie, reticulate, rtracklayer, stats, stringi,
tidyverse, tidyselect, utils

Suggests AnnotationHub, BiocStyle, BSgenome.Hsapiens.UCSC.hg38,
BSgenome.Mmusculus.UCSC.mm10,
BSgenome.Scerevisiae.UCSC.sacCer1, ensemblDb, IRanges, knitr,
magick, rmarkdown, testthat, TxDb.Mmusculus.UCSC.mm10.knownGene

VignetteBuilder knitr

biocViews CRISPR, Software

BugReports <https://github.com/loosolab/multicrispr/issues>

URL <https://github.com/loosolab/multicrispr>

git_url <https://git.bioconductor.org/packages/multicrispr>

git_branch RELEASE_3_15

git_last_commit 0f3e6be

git_last_commit_date 2022-04-26

Date/Publication 2022-10-18

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R topics documented:

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add_genome_matches *Add genome matches*

Description

Add genome matches

Usage

```
add_genome_matches(
  spacers,
  bsgenome = getBSgenome(genome(spacers)[1]),
  mismatches = 2,
  pam = "NGG",
  offtargetmethod = c("bowtie", "pdict")[1],
  outdir = OUTDIR,
  indexedgenomesdir = INDEXEDGENOMESDIR,
  verbose = TRUE
)
```

Arguments

spacers	GRanges
bsgenome	BSgenome
mismatches	number
pam	string
offtargetmethod	'bowtie' or 'pdict'
outdir	bowtie output directory
indexedgenomesdir	directory with indexed genomes
verbose	TRUE (default) or FALSE

Value

GRanges

Examples

```
require(magrittr)
file <- system.file('extdata/SRF.bed', package='multicrispr')
bsgenome <- BSgenome.Mmusculus.UCSC.mm10::BSgenome.Mmusculus.UCSC.mm10
targets0 <- bed_to_granges(file, 'mm10')
targets <- extend(targets0)
spacers <- find_spacers(targets, bsgenome, complement = FALSE,
                        ontargetmethod = NULL, offtargetmethod = NULL)
spacers %<% extract(1:100)
spacers %<% add_genome_matches(bsgenome)
```

`add_inverse_strand` *Add inverse strand*

Description

Add inverse strand

Usage

```
add_inverse_strand(gr, verbose = FALSE, plot = FALSE, ...)
```

Arguments

<code>gr</code>	<code>GRanges-class</code>
<code>verbose</code>	TRUE or FALSE (default)
<code>plot</code>	TRUE or FALSE (default)
<code>...</code>	<code>plot_intervals</code> arguments

Value

`GRanges-class`

Examples

```
# PE example
#-----
require(magrittr)
bsgenome <- BSgenome.Hsapiens.UCSC.hg38::BSgenome.Hsapiens.UCSC.hg38
gr <- char_to_granges(c(PRNP = 'chr20:4699600:+',           # snp
                      HBB  = 'chr11:5227002:-',          # snp
                      HEXA = 'chr15:72346580-72346583:-',    # del
                      CFTR = 'chr7:117559593-117559595:+'), # ins
                      bsgenome)
add_inverse_strand(gr, plot = TRUE)

# TFBS example
#-----
bedfile <- system.file('extdata/SRF.bed', package='multicrispr')
gr <- bed_to_granges(bedfile, genome = 'mm10')
add_inverse_strand(gr)
```

add_seq	<i>Add sequence to GRanges</i>
---------	--------------------------------

Description

Add sequence to GRanges

Usage

```
add_seq(gr, bsgenome, verbose = FALSE, as.character = TRUE)
```

Arguments

gr	GRanges-class
bsgenome	BSgenome-class
verbose	TRUE or FALSE (default)
as.character	TRUE (default) or FALSE

Value

GRanges-class

Examples

```
# PE example
#-----
require(magrittr)
bsgenome <- BSgenome.Hsapiens.UCSC.hg38::BSgenome.Hsapiens.UCSC.hg38
gr <- char_to_granges(c(PRNP = 'chr20:4699600:+',           # snp
                      HBB  = 'chr11:5227002:-',          # snp
                      HEXA = 'chr15:72346580-72346583:-',    # del
                      CFTR = 'chr7:117559593-117559595:+'), # ins
                      bsgenome)
(gr %>% add_seq(bsgenome))

# TFBS example
#-----
bsgenome <- BSgenome.Mmusculus.UCSC.mm10::BSgenome.Mmusculus.UCSC.mm10
bedfile  <- system.file('extdata/SRF.bed', package='multicrispr')
gr <- bed_to_granges(bedfile, 'mm10')
(gr %>% add_seq(bsgenome))
```

`add_target_matches` *Add target matches*

Description

Add target matches

Usage

```
add_target_matches(
  spacers,
  targets,
  bsgenome,
  mismatches = 2,
  pam = "NGG",
  outdir = OUTDIR,
  verbose = TRUE
)
```

Arguments

<code>spacers</code>	GRanges
<code>targets</code>	GRanges
<code>bsgenome</code>	BSgenome
<code>mismatches</code>	number
<code>pam</code>	string
<code>outdir</code>	bowtie output directory
<code>verbose</code>	TRUE (default) or FALSE

Value

GRanges

Examples

```
require(magrittr)
file <- system.file('extdata/SRF.bed', package='multicrispr')
bsgenome <- BSgenome.Mmusculus.UCSC.mm10::BSgenome.Mmusculus.UCSC.mm10
targets0 <- bed_to_granges(file, 'mm10')
targets <- extend(targets0)
spacers <- find_spacers(targets, bsgenome, complement = FALSE,
                        ontargetmethod = NULL, offtargetmethod = NULL)
spacers %>% add_target_matches(targets, bsgenome)
```

bed_to_granges	<i>Read bedfile into GRanges</i>
----------------	----------------------------------

Description

Read bedfile into GRanges

Usage

```
bed_to_granges(  
  bedfile,  
  genome,  
  txdb = NULL,  
  do_order = TRUE,  
  plot = TRUE,  
  verbose = TRUE  
)
```

Arguments

bedfile	file path
genome	string: UCSC genome name (e.g. 'mm10')
txdb	NULL (default) or TxDb-class (used for gene annotation)
do_order	TRUE (default) or FALSE: order on seqnames and star?
plot	TRUE (default) or FALSE: plot karyogram?
verbose	TRUE (default) or FALSE

Value

[GRanges-class](#)

See Also

[char_to_granges](#), [genes_to_granges](#)

Examples

```
bedfile <- system.file('extdata/SRF.bed', package = 'multicrispr')  
bsgenome <- BSgenome.Mmusculus.UCSC.mm10::BSgenome.Mmusculus.UCSC.mm10  
(gr <- bed_to_granges(bedfile, genome='mm10'))
```

`char_to_granges` *Convert character vector into GRanges*

Description

Convert character vector into GRanges

Usage

```
char_to_granges(x, bsgenome)
```

Arguments

<code>x</code>	character vector
<code>bsgenome</code>	BSgenome-class

Value

GRanges-class

See Also

[bed_to_granges](#), [genes_to_granges](#)

Examples

```
require(magrittr)
bsgenome <- BSgenome.Hsapiens.UCSC.hg38::BSgenome.Hsapiens.UCSC.hg38
x <- c(PRNP = 'chr20:4699600:+',           # snp
        HBB  = 'chr11:5227002:-',           # snp
        HEXA = 'chr15:72346580-72346583:-', # del
        CFTR = 'chr7:117559593-117559595:+') # ins
gr <- char_to_granges(x, bsgenome)
plot_intervals(gr, facet_var = c('targetname', 'seqnames'))
```

`double_flank` *Double flank*

Description

Double flank

Usage

```
double_flank(
  gr,
  upstart = -200,
  upend = -1,
  downstart = 1,
  downend = 200,
  strandaware = TRUE,
  plot = FALSE,
  linetype_var = "set",
  ...
)
```

Arguments

gr	GRanges-class
upstart	upstream flank start in relation to start(gr)
upend	upstream flank end in relation to start(gr)
downstart	downstream flank start in relation to end(gr)
downend	downstream flank end in relation to end(gr)
strandaware	TRUE (default) or FALSE
plot	TRUE or FALSE (default)
linetype_var	gr var mapped to linetype
...	passed to plot_intervals

Value

[GRanges-class](#)

Examples

```
# Prime Editing example
#-----
require(magrittr)
bsgenome <- BSgenome.Hsapiens.UCSC.hg38::BSgenome.Hsapiens.UCSC.hg38
gr <- char_to_granges(c(PRNP = 'chr20:4699600:+',           # snp
                      HBB  = 'chr11:5227002:-',      # snp
                      HEXA = 'chr15:72346580-72346583:-',    # del
                      CFTR = 'chr7:117559593-117559595:+'), # ins
                      bsgenome)
double_flank(gr, -10, -1, +1, +20, plot = TRUE)

# TFBS example
#-----
bedfile <- system.file('extdata/SRF.bed', package='multicrispr')
gr <- bed_to_granges(bedfile, genome = 'mm10', plot = FALSE)
double_flank(gr, plot = TRUE)
```

<i>extend_for_pe</i>	<i>Extend ranges for prime editing</i>
----------------------	--

Description

Extend target ranges to span in which to look for spacer-pam seqs

Usage

```
extend_for_pe(
  gr,
  bsgenome,
  nrt = 16,
  spacer = strrep("N", 20),
  pam = "NGG",
  plot = FALSE
)
```

Arguments

<code>gr</code>	GRanges-class
<code>bsgenome</code>	BSgenome-class
<code>nrt</code>	number: reverse transcription length
<code>spacer</code>	string: spacer pattern in extended IUPAC alphabet
<code>pam</code>	string: pam pattern in extended IUPAC alphabet
<code>plot</code>	TRUE (default) or FALSE

Details

Extend target ranges to find nearby spacers for prime editing

Value

[GRanges-class](#)

Examples

```
require(magrittr)
bsgenome <- BSgenome.Hsapiens.UCSC.hg38::BSgenome.Hsapiens.UCSC.hg38
gr <- char_to_granges(c( PRNP = 'chr20:4699600:+',           # snp
                      HBB  = 'chr11:5227002:-',          # snp
                      HEXA = 'chr15:72346580-72346583:-',   # del
                      CFTR = 'chr7:117559593-117559595:+'), # ins
                      bsgenome = bsgenome)
find_primespacers(gr, bsgenome)
(grext <- extend_for_pe(gr))
find_spacers(grext, bsgenome, complement = FALSE)
```

<code>extend_pe_to_gg</code>	<i>Extend prime editing target to find GG sites</i>
------------------------------	---

Description

Extend prime editing target to find GG sites in accessible neighbourhood

Usage

```
extend_pe_to_gg(gr, nrt = 16, plot = FALSE)
```

Arguments

<code>gr</code>	target GRanges-class
<code>nrt</code>	n RT nucleotides (default 16, recommended 10-16)
<code>plot</code>	TRUE or FALSE (default)

Details

Extends each target range to the area in which to search for a prime editing GG duplet, as shown in the sketch below.

Value

[GRanges-class](#)

Examples

```
# PE example
#-----
require(magrittr)
bsgenome <- BSgenome.Hsapiens.UCSC.hg38::BSgenome.Hsapiens.UCSC.hg38
gr <- char_to_granges(c(PRNP = 'chr20:4699600:+',           # snp
                      HBB  = 'chr11:5227002:-',          # snp
                      HEXA = 'chr15:72346580-72346583:-',    # del
                      CFTR = 'chr7:117559593-117559595:+'), # ins
                      bsgenome)
extend_pe_to_gg(gr, plot = TRUE)
```

`extract_matchranges` *Extract matching subranges*

Description

Extract subranges that match pattern

Usage

```
extract_matchranges(gr, bsgenome, pattern, plot = FALSE)
```

Arguments

<code>gr</code>	GRanges-class
<code>bsgenome</code>	BSgenome{BSgenome-class}
<code>pattern</code>	string: search pattern in extended IUPAC alphabet
<code>plot</code>	TRUE or FALSE (default)

Value

[GRanges-class](#)

Examples

```
# PE example
#-----
require(magrittr)
bsgenome <- BSgenome.Hsapiens.UCSC.hg38::BSgenome.Hsapiens.UCSC.hg38
gr <- char_to_granges(c(
  PRNP = 'chr20:4699600:+',           # snp
  HBB  = 'chr11:5227002:-',           # snp
  HEXA = 'chr15:72346580-72346583:-', # del
  CFTR = 'chr7:117559593-117559595:+'), # ins
  bsgenome)
gr %>% extend_for_pe()
pattern <- strrep('N',20) %>% paste0('NGG')
extract_matchranges(gr, bsgenome, pattern, plot = TRUE)

# TFBS examples
#-----
bsgenome <- BSgenome.Mmusculus.UCSC.mm10::BSgenome.Mmusculus.UCSC.mm10
bedfile  <- system.file('extdata/SRF.bed', package='multicrispr')
gr <- bed_to_granges(bedfile, 'mm10') %>% extend()
extract_matchranges(gr, bsgenome, pattern = strrep('N',20) %>% paste0('NGG'))
```

extract_subranges *Extract subranges*

Description

Extract subranges from a [GRanges-class](#) object

Usage

```
extract_subranges(gr, ir, plot = FALSE)
```

Arguments

gr	GRanges-class
ir	IRanges-class : subranges to be extracted
plot	TRUE or FALSE (default)

Value

[GRanges-class](#).

Examples

```
# Extract a subrange
gr <- GenomicRanges::GRanges(c(A = 'chr1:1-100:+', B = 'chr1:1-100:-'))
gr$targetname <- 'AB'
ir <- IRanges::IRanges(c(A = '1-10', A = '11-20', B = '1-10', B = '11-20'))
extract_subranges(gr, ir, plot = TRUE)

# Return empty GRanges for empty IRanges
extract_subranges(GenomicRanges::GRanges('chr1:345-456'), IRanges::IRanges())
```

find_gg *Find GG*

Description

Find GG

Usage

```
find_gg(gr)
```

Arguments

gr	GRanges-class
----	-------------------------------

Value**GRanges-class****Examples**

```
# PE example
#-----
require(magrittr)
bsgenome <- BSgenome.Hsapiens.UCSC.hg38::BSgenome.Hsapiens.UCSC.hg38
gr <- char_to_granges(c(PRNP = 'chr20:4699600:+',           # snp
                      HBB  = 'chr11:5227002:-',          # snp
                      HEXA = 'chr15:72346580-72346583:-',    # del
                      CFTR = 'chr7:117559593-117559595:+'), # ins
                      bsgenome)
gr %<>% extend_pe_to_gg(plot = TRUE) %>% add_seq(bsgenome)
find_gg(gr)
```

find_primespacers *Find prime editing spacers***Description**

Find prime editing spacers around target ranges

Usage

```
find_primespacers(
  gr,
  bsgenome,
  edits = get_plus_seq(bsgenome, gr),
  nprimer = 13,
  nrt = 16,
  ontargetmethod = c("Doench2014", "Doench2016")[1],
  offtargetmethod = c("bowtie", "pdict")[1],
  mismatches = 0,
  nickmatches = 2,
  indexedgenomesdir = INDEXEDGENOMESDIR,
  outdir = OUTDIR,
  verbose = TRUE,
  plot = TRUE,
  ...
)
```

Arguments

gr	GRanges-class
bsgenome	BSgenome-class

edits	character vector: desired edits on '+' strand. If named, names should be identical to those of gr
nprimer	n primer nucleotides (default 13, max 17)
nrt	n rev transcr nucleotides (default 16, recomm. 10-16)
ontargetmethod	'Doench2014' or 'Doench2016': on-target scoring method
offtargetmethod	'bowtie' or 'pdict'
mismatches	no of primespacer mismatches (default 0, to suppress offtarget analysis: -1)
nickmatches	no of nickspacer offtarget mismatches (default 2, to suppresses offtarget analysis: -1)
indexedgenomesdir	directory with indexed genomes (as created by index_genome)
outdir	directory where offtarget analysis output is written
verbose	TRUE (default) or FALSE
plot	TRUE (default) or FALSE
...	passed to plot_intervals

Details

Below the architecture of a prime editing site. Edits can be performed anywhere in the revtranscript area.

```
spacer pam -----== primer revtranscript -----=====
.....CC..... -----extension-----
```

Value

[GRanges-class](#) with prime editing spacer ranges and following mcols: * crisprspacer: N20 spacers * crisppam: NGG PAMs * crisprprimer: primer (on PAM strand) * crisprtranscript: reverse transcript (on PAM strand) * crisprextension: 3' extension of gRNA contains: reverse transcription template + primer binding site sequence can be found on non-PAM strand * crisprexrange: genomic range of crispr extension * Doench2016[4]: on-target efficiency scores * off0, off1, off2: number of offtargets with 0, 1, 2 mismatches * off: total number of offtargets: off = off0 + off1 + ... * nickrange: nickspacer range * nickspacer: nickspacer sequence * nickDoench2016[4]: nickspacer Doench scores * nickoff: nickspacer offtarget counts

See Also

[find_spacers](#) to find standard crispr sites

Examples

```
# Find PE spacers for 4 clinically relevant loci (Anzalone et al, 2019)
bsgenome <- BSgenome.Hsapiens.UCSC.hg38::BSgenome.Hsapiens.UCSC.hg38
gr <- char_to_granges(c(
  PRNP = 'chr20:4699600:+',           # SNP: prion disease
  HBB  = 'chr11:5227002:-',           # SNP: sickle cell anemia
```

```

HEXA = 'chr15:72346580-72346583:-', # del: tay sachs disease
CFTR = 'chr7:117559593-117559595:+'), # ins: cystic fibrosis
bsgenome)
spacers <- find_primespacers(gr, bsgenome)
spacers <- find_spacers(extend_for_pe(gr), bsgenome, complement = FALSE)

# Edit PRNP locus for resistance against prion disease (Anzalone et al, 2019)
bsgenome <- BSgenome.Hsapiens.UCSC.hg38::BSgenome.Hsapiens.UCSC.hg38
gr <- char_to_granges(c(PRNP = 'chr20:4699600:+'), bsgenome)
find_primespacers(gr, bsgenome)
find_primespacers(gr, bsgenome, edits = 'T')

```

find_spacers*Find crispr spacers in targetranges***Description**

Find crispr spacers in targetranges

Usage

```

find_spacers(
  gr,
  bsgenome,
  spacer = strrep("N", 20),
  pam = "NGG",
  complement = TRUE,
  ontargetmethod = c("Doench2014", "Doench2016")[1],
  offtargetmethod = c("bowtie", "pdict")[1],
  offtargetfilterby = character(),
  subtract_targets = FALSE,
  mismatches = 2,
  indexedgenomesdir = INDEXEDGENOMESDIR,
  outdir = OUTDIR,
  verbose = TRUE,
  plot = TRUE,
  ...
)

```

Arguments

gr	GRanges-class
bsgenome	BSgenome-class
spacer	string: spacer pattern in extended IUPAC alphabet
pam	string: pam pattern in extended IUPAC alphabet
complement	TRUE (default) or FALSE: also search in compl ranges?
ontargetmethod	'Doench2016' 'Doench2016' or NULL (no on-target score)

```

offtargetmethod
  'bowtie', 'pdict', or NULL (no offtarget analysis)
offtargetfilterby
  filter for best off-target counts by this variable
subtract_targets
  TRUE or FALSE (default): whether to subtract target (mis)matches from offtarget
  counts
mismatches
  0-3: allowed mismatches in offtargetanalysis (choose mismatch=-1 to suppress
  offtarget analysis)
indexedgenomesdir
  directory with Bowtie-indexed genomes (as produced with index\_genome)
outdir
  directory where bowtie analysis results are written to
verbose
  TRUE (default) or FALSE
plot
  TRUE (default) or FALSE
...
  passed to plot_intervals

```

Value

[GRanges-class](#)

See Also

[find_prime_spacers](#) to find prime editing spacers

Examples

```

# PE example
#-----
require(magrittr)
bsgenome <- BSgenome.Hsapiens.UCSC.hg38::BSgenome.Hsapiens.UCSC.hg38
gr <- char_to_granges(c(PRNP = 'chr20:4699600:+',           # snp
                      HBB  = 'chr11:5227002:-',           # snp
                      HEXA = 'chr15:72346580-72346583:-', # del
                      CFTR = 'chr7:117559593-117559595:+'), # ins
                      bsgenome)
plot_intervals(gr)
find_prime_spacers(gr, bsgenome)
find_spacers(extend_for_pe(gr), bsgenome, complement=FALSE, mismatches=0)
  # complement = FALSE because extend_for_pe already
  # adds reverse complements and does so in a strand-specific
  # manner

# TFBS example
#-----
bsgenome <- BSgenome.Mmusculus.UCSC.mm10::BSgenome.Mmusculus.UCSC.mm10
bedfile <- system.file('extdata/SRF.bed', package='multicrispr')
gr <- bed_to_granges(bedfile, 'mm10') %>% extend()
gr %<%> extract(1:100)
find_spacers(gr, bsgenome, subtract_targets = TRUE)

```

genes_to_granges *Convert geneids into GRanges*

Description

Convert geneids into GRanges

Usage

```
genes_to_granges(geneids, txdb, complement = TRUE, plot = TRUE, verbose = TRUE)

genefile_to_granges(file, txdb, complement = TRUE, plot = TRUE)
```

Arguments

geneids	Gene identifier vector
txdb	TxDb-class or EnsDb-class
complement	TRUE (default) or FALSE: add complementary strand?
plot	TRUE (default) or FALSE
verbose	TRUE (default) or FALSE
file	Gene identifier file (one per row)

Value

[GRanges-class](#)

See Also

[char_to_granges](#), [bed_to_granges](#)

Examples

```
# Entrez
#-----
genefile <- system.file('extdata/SRF.entrez', package='multicrispr')
geneids  <- as.character(read.table(genefile)[[1]])
txdb      <- getFromNamespace('TxDb.Mmusculus.UCSC.mm10.knownGene',
                               'TxDb.Mmusculus.UCSC.mm10.knownGene')
(gr <- genes_to_granges(geneids, txdb))
(gr <- genefile_to_granges(genefile, txdb))

# Ensembl
#-----
# txdb <- AnnotationHub::AnnotationHub()["AH75036"]
# genefile <- system.file('extdata/SRF.ensembl', package='multicrispr')
# geneids <- as.character(read.table(genefile)[[1]])
# (gr <- genes_to_granges(geneids, txdb))
# (gr <- genefile_to_granges(genefile, txdb))
```

<code>gr2dt</code>	<i>GRanges <-> data.table</i>
--------------------	-------------------------------------

Description

`GRanges <-> data.table`

Usage

```
gr2dt(gr)
```

```
dt2gr(dt, seqinfo)
```

Arguments

<code>gr</code>	<code>GRanges-class</code>
<code>dt</code>	<code>data.table</code>
<code>seqinfo</code>	<code>Seqinfo-class</code>

Value

`data.table` (`gr2dt`) or `GRanges` (`dt2gr`)

Examples

```
bsgenome <- BSgenome.Hsapiens.UCSC.hg38::BSgenome.Hsapiens.UCSC.hg38
gr <- char_to_granges(c(PRNP = 'chr20:4699600:+', # snp
                        HBB = 'chr11:5227002:-', # snp
                        HEXA = 'chr15:72346580-72346583:-', # del
                        CFTR = 'chr7:117559593-117559595:+'), # ins
                        bsgenome)
(dt <- gr2dt(gr))
(gr <- dt2gr(dt, BSgenome::seqinfo(bsgenome)))
```

<code>has Been Indexed</code>	<i>Has been indexed?</i>
-------------------------------	--------------------------

Description

Has been indexed?

Usage

```
has_Been_Indexed(bsgenome, indexedgenomesdir = INDEXEDGENOMESDIR)
```

Arguments

<code>bsgenome</code>	<code>BSgenome</code>
<code>indexedgenomesdir</code>	directory with indexed genomes

Value

TRUE or FALSE

Examples

```
bsgenome <- BSgenome.Hsapiens.UCSC.hg38::BSgenome.Hsapiens.UCSC.hg38
has_being_indexed(bsgenome)
```

<code>index_genome</code>	<i>Index genome</i>
---------------------------	---------------------

Description

Bowtie index genome

Usage

```
index_genome(
  bsgenome,
  indexedgenomesdir = INDEXEDGENOMESDIR,
  download = TRUE,
  overwrite = FALSE
)
```

Arguments

<code>bssgenome</code>	<code>BSgenome-class</code>
<code>indexedgenomesdir</code>	string: directory with bowtie-indexed genome
<code>download</code>	TRUE (default) or FALSE: whether to download pre-indexed version if available
<code>overwrite</code>	TRUE or FALSE (default)

Details

Checks whether already available locally. If not, checks whether indexed version can be downloaded from our s3 storage. If not, builds the index with bowtie. This can take a few hours, but is a one-time operation.

Value

`invisible(genomdir)`

Examples

```
bsgenome <- BSgenome.Scerevisiae.UCSC.sacCer1::Scerevisiae
index_genome(bsgenome, indexedgenomesdir = tempdir())
```

index_targets

Index targets

Description

Bowtie index targets

Usage

```
index_targets(
  targets,
  bsgenome = getBSgenome(genome(targets)[1]),
  outdir = OUTDIR,
  verbose = TRUE
)
```

Arguments

targets	GRanges-class
bsgenome	BSgenome-class
outdir	string: output directory
verbose	TRUE (default) or FALSE

Value

invisible(targetdir)

Examples

```
require(magrittr)
bsgenome <- BSgenome.Mmusculus.UCSC.mm10::BSgenome.Mmusculus.UCSC.mm10
bedfile <- system.file('extdata/SRF.bed', package = 'multicrispr')
targets <- extend(bed_to_granges(bedfile, genome = 'mm10'))
index_targets(targets, bsgenome)
```

<code>plot_intervals</code>	<i>Interval plot GRanges</i>
-----------------------------	------------------------------

Description

Interval plot GRanges

Usage

```
plot_intervals(
  gr,
  xref = "targetname",
  y = default_y(gr),
  nperchrom = 2,
  nchrom = 4,
  color_var = "targetname",
  facet_var = "seqnames",
  linetype_var = default_linetype(gr),
  size_var = default_size_var(gr),
  alpha_var = default_alpha_var(gr),
  title = NULL,
  scales = "free"
)
```

Arguments

<code>gr</code>	GRanges-class
<code>xref</code>	gr var used for scaling x axis
<code>y</code>	'names' (default) or name of gr variable
<code>nperchrom</code>	number (default 1): n head (and n tail) targets shown per chromosome
<code>nchrom</code>	number (default 6) of chromosomes shown
<code>color_var</code>	'seqnames' (default) or other gr variable
<code>facet_var</code>	NULL(default) or gr variable mapped to facet
<code>linetype_var</code>	NULL (default) or gr variable mapped to linetype
<code>size_var</code>	NULL (default) or gr variable mapped to size
<code>alpha_var</code>	NULL or gr variable mapped to alpha
<code>title</code>	NULL or string: plot title
<code>scales</code>	'free', 'fixed', etc

Value

ggplot object

See Also

[plot_karyogram](#)

Examples

```
# SRF sites
require(magrittr)
bsgenome <- BSgenome.Mmusculus.UCSC.mm10::BSgenome.Mmusculus.UCSC.mm10
bedfile <- system.file('extdata/SRF.bed', package = 'multicrispr')
targets <- bed_to_granges(bedfile, 'mm10', plot = FALSE)
plot_intervals(targets)

# PE targets
bsgenome <- BSgenome.Hsapiens.UCSC.hg38::BSgenome.Hsapiens.UCSC.hg38
gr <- char_to_granges(c(PRNP = 'chr20:4699600:+',
                         HBB = 'chr11:5227002:-',
                         HEXA = 'chr15:72346580-72346583:-',
                         CFTR = 'chr7:117559593-117559595:+'),
                         bsgenome)
spacers <- find_primespacers(gr, bsgenome, plot = FALSE)
plot_intervals(gr)
plot_intervals(extend_for_pe(gr))
plot_intervals(spacers)

# Empty gr
plot_intervals(GenomicRanges::GRanges())
```

plot_karyogram

Karyo/Interval Plot GRanges(List)

Description

Karyo/Interval Plot GRanges(List)

Usage

```
plot_karyogram(grlist, title = unique(genome(grlist)))
```

Arguments

grlist	GRanges-class
title	plot title

Value

list

See Also[plot_intervals](#)**Examples**

```
# Plot GRanges
  bedfile <- system.file('extdata/SRF.bed', package = 'multicrispr')
  gr <- bed_to_granges(bedfile, 'mm10', plot = FALSE)
  plot_karyogram(gr)

# Plot GRangesList
  flanks <- up_flank(gr, stranded=FALSE)
  grlist <- GenomicRanges::GRangesList(sites = gr, flanks = flanks)
  plot_karyogram(grlist)
```

<code>score_ontargets</code>	<i>Add on-target efficiency scores</i>
------------------------------	--

Description

Add Doench2014 or Doench2016 on-target efficiency scores

Usage

```
score_ontargets(
  spacers,
  bsgenome,
  ontargetmethod = c("Doench2014", "Doench2016")[1],
  chunksize = 10000,
  verbose = TRUE,
  plot = TRUE,
  ...
)
```

Arguments

<code>spacers</code>	<code>GRanges-class</code> : spacers
<code>bsgenome</code>	<code>BSgenome-class</code>
<code>ontargetmethod</code>	'Doench2014' (default) or 'Doench2016' (requires non-NULL argument python, virtualenv, or condaenv)
<code>chunksize</code>	Doench2016 is executed in chunks of chunksize
<code>verbose</code>	TRUE (default) or FALSE
<code>plot</code>	TRUE (default) or FALSE
<code>...</code>	passed to plot_intervals

Details

`add_ontargets` adds efficiency scores `filter_ontargets` adds efficiency scores and filters on them

Value

numeric vector

References

Doench 2014, Rational design of highly active sgRNAs for CRISPR-Cas9-mediated gene inactivation. *Nature Biotechnology*, doi: 10.1038/nbt.3026

Doench 2016, Optimized sgRNA design to maximize activity and minimize off-target effects of CRISPR-Cas9. *Nature Biotechnology*, doi: 10.1038/nbt.3437

Python module azimuth: [github/MicrosoftResearch/azimuth](https://github.com/MicrosoftResearch/azimuth)

Examples

```
# Install azimuth
#-----
## With reticulate
# require(reticulate)
# conda_create('azienv', c('python=2.7'))
# use_condaenv('azienv')
# py_install(c('azimuth', 'scikit-learn==0.17.1', 'biopython=='1.76'),
#            'azienv', pip = TRUE)

## Directly
# conda create --name azienv python=2.7
# conda activate azienv
# pip install scikit-learn==0.17.1
# pip install biopython==1.76
# pip install azimuth

# PE example
#-----
require(magrittr)
bsgenome <- BSgenome.Hsapiens.UCSC.hg38::BSgenome.Hsapiens.UCSC.hg38
targets <- char_to_granges(c(PRNP = 'chr20:4699600:+',           # snp
                           HBB   = 'chr11:5227002:-',       # snp
                           HEXA  = 'chr15:72346580-72346583:-', # del
                           CFTR  = 'chr7:117559593-117559595:+'), # ins
                           bsgenome)
spacers <- find_primespacers(targets, bsgenome, ontargetmethod=NULL,
                               offtargetmethod=NULL)
spacers %>% score_ontargets(bsgenome, 'Doench2014')
# reticulate::use_condaenv('azienv')
# reticulate::import('azimuth')
# spacers %>% score_ontargets(bsgenome, 'Doench2016')

# TFBS example
```

```
#-----
bedfile <- system.file('extdata/SRF.bed', package = 'multicrispr')
bsgenome <- BSgenome.Mmusculus.UCSC.mm10::BSgenome.Mmusculus.UCSC.mm10
targets <- extend(bed_to_granges(bedfile, 'mm10'))
spacers <- find_spacers(targets, bsgenome, ontargetmethod=NULL,
                        offtargetmethod=NULL)
spacers %<>% score_ontargets(bsgenome, 'Doench2014')
# reticulate::use_condaenv('azienv')
# reticulate::import('azimuth')
# spacers %>% score_ontargets(bsgenome, 'Doench2016')
```

up_flank*Extend or Flank GRanges***Description**

Returns extensions, upstream flanks, or downstream flanks

Usage

```
up_flank(
  gr,
  start = -200,
  end = -1,
  strandaware = TRUE,
  bsgenome = NULL,
  verbose = FALSE,
  plot = FALSE,
  linetype_var = "set",
  ...
)

down_flank(
  gr,
  start = 1,
  end = 200,
  strandaware = TRUE,
  bsgenome = NULL,
  verbose = FALSE,
  plot = FALSE,
  linetype_var = "set",
  ...
)

extend(
  gr,
  start = -22,
  end = 22,
```

```

strandaware = TRUE,
bsgenome = NULL,
verbose = FALSE,
plot = FALSE,
linetype_var = "set",
...
)

```

Arguments

gr	GRanges-class
start	number or vector (same length as gr): start definition, relative to gr start (up_flank, extend) or gr end (down_flank).
end	number or vector (same length as gr): end definition, relative to gr start (up_flank) or gr end (extend, down_flank).
strandaware	TRUE (default) or FALSE: consider strand information?
bsgenome	NULL (default) or BSgenome-class . Required to update gr\$seq if present.
verbose	TRUE or FALSE (default)
plot	TRUE or FALSE (default)
linetype_var	string: gr var mapped to linetype
...	passed to plot_intervals

Details

up_flank returns upstream flanks, in relation to start(gr). down_flank returns downstream flanks, in relation to end(gr). extend returns extensions, in relation to start(gr) and end(gr)

Value

a [GRanges-class](#)

Examples

```

# PE example
#-----
require(magrittr)
bsgenome <- BSgenome.Hsapiens.UCSC.hg38::BSgenome.Hsapiens.UCSC.hg38
gr <- char_to_granges(c(PRNP = 'chr20:4699600:+',           # snp
                      HBB  = 'chr11:5227002:-',          # snp
                      HEXA = 'chr15:72346580-72346583:-', # del
                      CFTR = 'chr7:117559593-117559595:+'),# ins
                      bsgenome = bsgenome)
gr %>% up_flank( -22, -1, plot=TRUE)
gr %>% up_flank( c(-10,-20,-30,-40), -1, plot=TRUE)
gr %>% up_flank( -22, -1, plot=TRUE, strandaware=FALSE)

gr %>% down_flank(+1, +22, plot=TRUE)
gr %>% down_flank(+1, c(10, 20, 30, 40), plot=TRUE)

```

```

gr %>% down_flank(+1, +22, plot=TRUE, strandaware=FALSE)

gr %>% extend( -10, +20, plot=TRUE)
gr %>% extend( -10, +20, plot=TRUE, strandaware=FALSE)

# TFBS example
#-----
bedfile <- system.file('extdata/SRF.bed', package='multicrispr')
gr <- bed_to_granges(bedfile, genome = 'mm10')
gr %>% extend(plot = TRUE)
gr %>% up_flank(plot = TRUE)
gr %>% down_flank(plot = TRUE)

```

write_ranges*Write GRanges to file***Description**

Write GRanges to file

Usage

```

write_ranges(gr, file, verbose = TRUE)

read_ranges(file, bsgenome)

```

Arguments

gr	GRanges-class
file	file
verbose	TRUE (default) or FALSE
bsgenome	BSgenome-class

Value

[GRanges-class](#) for read_ranges

Examples

```

# Find PE spacers for 4 clinically relevant loci (Anzalone et al, 2019)
bsgenome <- BSgenome.Hsapiens.UCSC.hg38::BSgenome.Hsapiens.UCSC.hg38
gr <- char_to_granges(c(
    PRNP = 'chr20:4699600:+',           # snp: prion disease
    HBB  = 'chr11:5227002:-',           # snp: sickle cell anemia
    HEXA = 'chr15:72346580-72346583:-', # del: tay sachs disease
    CFTR = 'chr7:117559593-117559595:+'), # ins: cystic fibrosis
    bsgenome)
file <- file.path(tempdir(), 'gr.txt')
write_ranges(gr, file)
read_ranges(file, bsgenome)

```

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