

Package ‘crisprBwa’

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Title BWA-based alignment of CRISPR gRNA spacer sequences

Depends methods

Imports BiocGenerics, BSgenome, crisprBase (>= 0.99.15), GenomeInfoDb, Rbwa, readr, stats, stringr, utils

Suggests BiocStyle, BSgenome.Hsapiens.UCSC.hg38, knitr, rmarkdown, testthat

biocViews CRISPR, FunctionalGenomics, Alignment

Description Provides a user-friendly interface to map on-targets and off-targets of CRISPR gRNA spacer sequences using bwa. The alignment is fast, and can be performed using either commonly-used or custom CRISPR nucleases. The alignment can work with any reference or custom genomes. Currently not supported on Windows machines.

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Encoding UTF-8

RoxygenNote 7.1.2

VignetteBuilder knitr

BugReports <https://github.com/crisprVerse/crisprBwa/issues>

URL <https://github.com/crisprVerse/crisprBwa>

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Author Jean-Philippe Fortin [aut, cre]

Maintainer Jean-Philippe Fortin <fortin946@gmail.com>

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<code>runBwa</code>	<i>Run BWA short-read aligner</i>
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Description

Return BWA alignments for a list of short sequences for a prebuilt BWA index.

Usage

```
runBwa(sequences, bwa_index = NULL, n_mismatches = 3)
```

Arguments

<code>sequences</code>	Character vector of DNA sequences.
<code>bwa_index</code>	String specifying path to the BWA index.
<code>n_mismatches</code>	Integer specifying maximum number of mismatches allowed between the query sequences and the index sequences.

Details

`runBwa` can be used to map short DNA sequences to a reference genome. To search for sequences while imposing constraints on PAM sequences (such as gRNA spacer sequences), see `runCrisprBwa` instead.

Value

A data.frame of the alignments with the following columns:

- `query` — string specifying query DNA sequence
- `chr` — string specifying chromosome name
- `pos` — string specifying genomic coordinate of the start of the target DNA sequence
- `strand` — string specifying strand ("+" or "-")
- `n_mismatches` — integer specifying number of mismatches between query and target sequences

Author(s)

Jean-Philippe Fortin

See Also

`link{runCrisprBwa}` to map gRNA spacer sequences.

Examples

```

fasta <- system.file(package="crisprBwa", "example/chr12.fa")
outdir <- tempdir()
index <- file.path(outdir, "chr12")
Rbwa::bwa_build_index(fasta,
                       index_prefix=index)

seqs <- c("GGAAGTTG",
          "GTGGACAC",
          "GTGTGCAA")

aln <- runBwa(seqs,
               n_mismatches=1,
               bwa_index=index)

```

runCrisprBwa

Find gRNA spacer alignments with bwa

Description

Return bwa alignments for a list of gRNA spacer sequences.

Usage

```

runCrisprBwa(
  spacers,
  bwa_index = NULL,
  bsgenome = NULL,
  crisprNuclease = NULL,
  canonical = TRUE,
  ignore_pam = FALSE,
  n_mismatches = 0,
  force_spacer_length = FALSE,
  verbose = TRUE
)

```

Arguments

<code>spacers</code>	Character vector of DNA sequences corresponding to gRNA spacer sequences. Must all be of equal length.
<code>bwa_index</code>	Path to the bwa index to be used for alignment.
<code>bsgenome</code>	BSgenome object.
<code>crisprNuclease</code>	CrisprNuclease object.
<code>canonical</code>	Should only canonical PAM sequences be considered? TRUE by default.
<code>ignore_pam</code>	If TRUE, will return all matches regardless of PAM sequence. FALSE by default.
<code>n_mismatches</code>	Integer specifying maximum number of mismatches allowed between spacer and protospacer sequences.

```

force_spacer_length
    Should the spacer length be overwritten in the crisprNuclease object? FALSE
    by default.
verbose      Should messages be printed to the consolde? TRUE by default.

```

Details

`runCrisprBwa` is similar to `runBwa`, with the addition of imposing constraints on PAM sequences such that the query sequences are valid protospacer sequences in the searched genome.

Value

`runBwa` returns spacer alignment data, including genomic coordinates and sequence.

Author(s)

Jean-Philippe Fortin

See Also

`link{runBwa}` to map general DNA sequences.

Examples

```

# Building BWA index first:
fasta <- system.file(package="crisprBwa", "example/chr12.fa")
outdir <- tempdir()
index <- file.path(outdir, "chr12")
Rbwa::bwa_build_index(fasta,
                      index_prefix=index)

# Aligning Cas9 gRNA
library(BSgenome.Hsapiens.UCSC.hg38)
seqs <- c("AGCTGTCCGTGGGGTCCGC",
         "CCCCCTGCTGCTGTGCCAGGC")
data(SpCas9, package="crisprBase")
bsgenome <- BSgenome.Hsapiens.UCSC.hg38
results <- runCrisprBwa(seqs,
                        bsgenome=bsgenome,
                        bwa_index=index,
                        n_mismatches=2,
                        crisprNuclease=SpCas9)

```

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