Package ‘CrispRVariants’

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Type Package

Title Tools for counting and visualising mutations in a target location

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Author Helen Lindsay [aut, cre]

Maintainer Helen Lindsay <helen.lindsay@uzh.ch>

Description CrispRVariants provides tools for analysing the results of a CRISPR-Cas9 mutagenesis sequencing experiment, or other sequencing experiments where variants within a given region are of interest. These tools allow users to localize variant allele combinations with respect to any genomic location (e.g. the Cas9 cut site), plot allele combinations and calculate mutation rates with flexible filtering of unrelated variants.

biocViews CRISPR, GenomicVariation, VariantDetection, GeneticVariability, DataRepresentation, Visualization

LazyData true

Depends R (>= 3.3), ggplot2

License GPL-2

Imports AnnotationDbi, BiocParallel, Biostrings, methods,
GenomeInfoDb, GenomicAlignments, GenomicRanges, grDevices,
grid, gridExtra, IRanges, reshape2, Rsamtools, S4Vectors (>= 0.9.38), utils

Suggests BiocStyle, gdata, GenomicFeatures, knitr, rmarkdown,
trrtracklayer, sangerseqR, testthat, VariantAnnotation

VignetteBuilder knitr

NeedsCompilation no

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R topics documented:

abiToFastq ................................................. 2
addClipped .................................................. 3
addCodonFrame .......................................... 4
annotateGenePlot ........................................ 4
arrangePlots ............................................. 5
barplotAlleleFreqs ................................. 6
abifToFastq

Read a file in ab1 (Sanger) format and convert to fastq

Description

This is an R implementation of Wibowo Arindrarto’s abifpy.py trimming module, which itself implements Richard Mott’s trimming algorithm. See https://github.com/bow/abifpy for more details.

Usage

abifToFastq(seqname, fname, outfname, trim = TRUE, cutoff = 0.05, min_seq_len = 20, offset = 33, recall = FALSE)
addClipped

Arguments

- `seqname` (name of sequence, to appear in fastq file)
- `fname` (filename of sequence in ab1 format)
- `outfname` (filename to append the fastq output to)
- `trim` (should low quality bases be trimmed from the ends? TRUE or FALSE)
- `cutoff` (probability cutoff)
- `min_seq_len` (minimum number of sequenced bases required in order to trim the read)
- `offset` (phred offset for quality scores)
- `recall` (Use sangerseqR to resolve the primary sequence if two sequences are present. May cause quality scores to be ignored. (Default: FALSE))

Details

Requires Bioconductor package SangerseqR

Value

None. Sequences are appended to the outfname.

Author(s)

Helen Lindsay

Examples

```r
ab1_fname <- system.file("extdata", "IM2033.ab1", package = "CrispRVariants")
abifToFastq("IM2033", ab1_fname, "IM2033.fastq")
```

---

addClipped  

*Extrapolates mapping location from clipped, aligned reads*

Description

Extrapolates the mapping location of a read by assuming that the clipped regions should map adjacent to the mapped locations. This is not always a good assumption, particularly in the case of chimeric reads!

Usage

```r
addClipped(bam, ...)
```

## S4 method for signature 'GAlignments'

```r
addClipped(bam, ...)
```

Arguments

- `bam` (A GAlignments object)
- `...` (additional arguments)
Value

A `GRanges` representation of the extended mapping locations

Author(s)

Helen Lindsay

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**addCodonFrame**  
*Internal CrispRVariants function for indicating codon frame on an alignment tile plot*

**Description**

Adds vertical dotted lines in intervals of three nucleotides. Codon frame is supplied, alignments are assumed not to span an intron-exon junction.

**Usage**

```
addCodonFrame(p, width, codon.frame)
```

**Arguments**

- `p`  
  A ggplot object, typically from `CrispRVariants::makeAlignmentTilePlot`
- `width`  
  The number of nucleotides in the alignments
- `codon.frame`  
  The leftmost starting location of the next codon - 1, 2, or 3

**Value**

A ggplot object with added vertical lines indicating the frame

Author(s)

Helen Lindsay

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**annotateGenePlot**  
*Plots and annotates transcripts*

**Description**

Plots the gene structure, annotates this with the target location

**Usage**

```
annotateGenePlot(txdb, target, target.colour = "red", target.size = 1, gene.text.size = 10, panel.margin = grid::unit(c(0.1, 0.1, 0.1, 0.1), "lines"), plot.title = NULL, all.transcripts = TRUE)
```
**arrangePlots**

**Arguments**

- **txdb** A GenomicFeatures:TxDb object
- **target** Location of target (GRanges)
- **target.colour** Colour of box indicating target region
- **target.size** Thickness of box indicating target region
- **gene.text.size** Size for figure label
- **panel.margin** Unit object, margin size
- **plot.title** A title for the plot. If no plot.title is supplied, the title is the list of gene ids shown (default). If plot.title == FALSE, the plot will not have a title.
- **all.transcripts** If TRUE (default), all transcripts of genes overlapping the target are shown, including transcripts that do not themselves overlap the target. If FALSE, only the transcripts that overlap the target are shown.

**Value**

A ggplot2 plot of the transcript structures

---

**arrangePlots**

**Arrange plots for plotVariants:CrisprSet**

**Description**

Arranges 3 plots in two rows. The vertical margins of the left.plot and right.plot constrained to be equal

**Usage**

```r
arrangePlots(top.plot, left.plot, right.plot, fig.height = NULL, 
  col.wdth.ratio = c(2, 1), row.ht.ratio = c(1, 6), 
  left.plot.margin = grid::unit(c(0.1, 0, 3, 0.2), "lines"))
```

**Arguments**

- **top.plot** ggplot grob, placed on top of the figure, spanning the figure width
- **left.plot** ggplot, placed in the second row on the left
- **right.plot** ggplot, placed in the second row on the right. y-axis labels are removed.
- **fig.height** Actual height for the figure. If not provided, figure height is the sum of the row.ht.ratio (Default: NULL)
- **col.wdth.ratio** Vector specifying column width ratio (Default: c(2, 1))
- **row.ht.ratio** Vector specifying row height ratio (Default: c(1,6))
- **left.plot.margin** Unit object specifying margins of left.plot. Margins of right.plot are constrained by the left.plot.

**Value**

The arranged plots
barplotAlleleFreqs

Plots barplots of the spectrum of variants for a sample set

Description

For signature "matrix", this function optionally does a very naive classification of variants by size. Frameshift variant combinations are those whose sum is not divisible by three. Intron boundaries are *NOT* considered, use with caution! For signature "CrisprSet", the function uses the VariantAnnotation package to localize variant alleles with respect to annotated transcripts. Variants are annotated as "coding" when they are coding in any transcript.

signature("CrisprSet") Groups variants by size and type and produces a barplot showing the variant spectrum for each sample. Accepts all arguments accepted by barplotAlleleFreqs for signature("matrix"). Requires package "VariantAnnotation"

signature("matrix") Accepts a matrix of allele counts, with rownames being alleles and column names samples.

Usage

barplotAlleleFreqs(obj, ...)

## S4 method for signature 'CrisprSet'
barplotAlleleFreqs(obj, ..., txdb, min.freq = 0, include.chimeras = TRUE, group = NULL, palette = c("rainbow", "bluered"))

## S4 method for signature 'matrix'
barplotAlleleFreqs(obj, category.labels = NULL, group = NULL, bar.colours = NULL, group.colours = NULL, legend.text.size = 10, axis.text.size = 10, legend.symbol.size = 1, snv.label = "SNV", novar.label = "no variant", chimera.label = "Other", include.table = TRUE, classify = TRUE)

Arguments

obj The object to be plotted
...
additional arguments
txdb A transcript database object
min.freq Include variants with at frequency least min.freq in at least one sample. (Default: 0, i.e. no cutoff)
include.chimeras Should chimeric reads be included in results? (Default: TRUE)
group A grouping factor for the columns in obj. Columns in the same group will be displayed in the same text colour (Default: NULL)
palette Colour palette. Options are "rainbow", a quantitative palette (default) or "bluered", a gradient palette.
category.labels Labels for each category, corresponding to the rows of obj. Only applicable when categories are provided, i.e. "classify" is FALSE. (Default: NULL)
bar.colours  Colours for the categories in the barplot. Colours must be provided if there are more than 6 different categories.
group.colours  Colours for the text labels for the experimental groups. A set of 15 different colours is provided.
legend.text.size  The size of the legend text, in points.
axis.text.size  The size of the axis text, in points
legend.symbol.size  The size of the symbols in the legend
snv.label  The row label for single nucleotide variants
novar.label  The row label for non-variant sequences
chimera.label  The row label for chimeric (non-linearly aligned) variant alleles
include.table  Should a table of allele (variant combination) counts and total sequences be plotted? (Default: TRUE)
classify  If TRUE, performs a naive classification by size (Default: TRUE)

Value
A ggplot2 barplot of the allele distribution and optionally a table of allele counts

Author(s)
Helen Lindsay

Examples

data("gol_clutch1")
barplotAlleleFreqs(variantCounts(gol))

# Just show the barplot without the counts table:
barplotAlleleFreqs(variantCounts(gol), include.table = FALSE)

collapsePairs  \textit{Internal CrispRVariants function for collapsing pairs with concordant indels}

Description
Given a set of alignments to a target region, finds read pairs. Compares insertion/deletion locations within pairs using the cigar string. Pairs with non-identical indels are excluded. Pairs with identical indels are collapsed to a single read, taking the consensus sequence of the pairs.

Usage
collapsePairs(alns, use.consensus = TRUE, keep.unpaired = TRUE, verbose = TRUE, ...)
Arguments

- **alns**: A GAlignments object. We do not use GAlignmentPairs because amplicon-seq can result in pairs in non-standard pairing orientation. Must include BAM flag, must not include unmapped reads.
- **use.consensus**: Should the consensus sequence be used if pairs have a mismatch? Setting this to be TRUE makes this function much slower (Default: TRUE)
- **keep.unpaired**: Should unpaired and chimeric reads be included? (Default: TRUE)
- **verbose**: Report statistics on reads kept and excluded
- **...**: Additional items with the same length as alns, that should be filtered to match alns.

Value

The alignments, with non-concordant pairs removed and concordant pairs represented by a single read.

Author(s)

Helen Lindsay

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**consensusSeqs**

*Get consensus sequences for variant alleles*

Description

Return consensus sequences of variant alleles. At present, chimeric alignments are not included.

Usage

```r
consensusSeqs(obj, ...)
```

```r
## S4 method for signature 'CrisprSet'
consensusSeqs(obj, ..., top.n = NULL, min.freq = 0,
               min.count = 1)
```

Arguments

- **obj**: An object containing aligned sequences
- **...**: additional arguments
- **top.n**: (Integer n) If specified, return variants ranked at least n according to frequency across all samples (Default: 0, i.e. no cutoff)
- **min.freq**: (Float n least one sample (Default: 0)
- **min.count**: (Integer n) Return variants with count greater than n in at least one sample (Default: 0)

Value

A DNAStringSet of consensus sequences on the positive strand.
countDeletions

Author(s)
Helen Lindsay

Examples
data("gol_clutch1")
seqs <- consensusSeqs(gol, sample = 2)

Description
Counts the number of reads containing a deletion or insertion (indel) of any size in a set of aligned reads. For countDeletions and countInsertions Reads may be filtered according to whether they contain more than one indel of the same or different types.

Usage
countDeletions(alns, ...)

## S4 method for signature 'GAlignments'
countDeletions(alns, ..., multi.del = FALSE, del.and.ins = FALSE, del.ops = c("D"))

countInsertions(alns, ...)

## S4 method for signature 'GAlignments'
countInsertions(alns, ..., ins.and.del = FALSE, multi.ins = FALSE, del.ops = c("D"))

countIndels(alns)

## S4 method for signature 'GAlignments'
countIndels(alns)

indelPercent(alns)

## S4 method for signature 'GAlignments'
indelPercent(alns)

Arguments

alns The aligned reads
...
extra arguments
multi.del If TRUE, returns the exact number of deletions, i.e., if one read contains 2 deletions, it contributes 2 to the total count (default: FALSE)
del.and.ins If TRUE, counts deletions regardless of whether reads also contain insertions. If FALSE, counts reads that contain deletions but not insertions (default: FALSE)
del.ops Cigar operations counted as deletions. Default: c("D")
### CrisprRun-class

**ins.and.del**
- If TRUE, counts insertions regardless of whether reads also contain deletions
- If FALSE, counts reads that contain insertions but not deletions (default: FALSE)

**multi.ins**
- If TRUE, returns the exact number of insertions, i.e., if one read contains 2 insertions, it contributes 2 to the total count (default: FALSE)

**Value**

- countDeletions: The number of reads containing a deletion (integer)
- countInsertions: The number of reads containing an insertion (integer)
- countIndels: The number of reads containing at least one insertion
- indelPercent: The percentage of reads containing an insertion or deletion (numeric)

**Author(s)**

Helen Lindsay

**Examples**

```r
bam_fname <- system.file("extdata", "gol_F1_clutch_2_embryo_4_s.bam", package = "CrispRVariants")
bam <- GenomicAlignments::readGAlignments(bam_fname, use.names = TRUE)
countDeletions(bam)
countInsertions(bam)
countIndels(bam)
indelPercent(bam)
```

---

**Description**

A ReferenceClass container for a single sample of alignments narrowed to a target region. Typically CrisprRun objects will not be accessed directly, but if necessary via a CrisprSet class which contains a list of CrisprRun objects. Note that the CrispRVariants plotting functions don’t work on CrisprRun objects.

**Arguments**

- **bam**: a GAlignments object containing (narrowed) alignments to the target region. Filtering of the bam should generally be done before initialising a CrisprRun object.
- **target**: The target location, a GRanges object.
- **genome.ranges**: A GRangesList of genomic coordinates for the cigar operations. If bam is a standard GAlignments object, this is equivalent to cigarRangesAlongReferenceSpace + start(bam).
- **rc**: (reverse complement) Should the alignments be reverse complemented, i.e. displayed with respect to the negative strand? (Default: FALSE)
- **name**: A name for this set of reads, used in plots if present (Default: NULL)
- **chimeras**: Off-target chimeric alignments not in bam. (Default: empty)
- **verbose**: Print information about initialisation progress (Default: TRUE)
Fields

- **alns**: A GAlignments object containing the narrowed reads. Note that if the alignments are represented with respect to the reverse strand, the "start" remains with respect to the forward strand, whilst the cigar and the sequence are reverse complemented.

- **name**: The name of the sample

- **cigar_labels**: A vector of labels for the reads, based on the cigar strings, optionally renumbered with respect to a new zero point (e.g. the cut site) and shortened to only insertion and deletion locations. Set at initialisation of a CrisprSet object, but not at initialisation of a CrisprRun object.

- **chimeras**: Chimeric, off-target alignments corresponding to alignments in alns

Methods

- **getCigarLabels(target.loc, genome_to_target, ref, separate.snv = TRUE, match.label = "no variant", mismatch.label = "SNV", rc = FALSE, keep.ops = c("I", "D", "N"), upstream = 8, downstream = min(5, width(ref) - cut_site))**

  Description: Sets the "cig_labels" field, returns the cigar labels.

  Input parameters:
  - **target.loc**: The location of the cut site with respect to the target genome.
  - **genome_to_target**: A vector with names being genomic locations and values being locations with respect to the cut site.
  - **separate.snv**: Should single nucleotide variants be called? (Default: TRUE)
  - **match.label**: Label for non-variant reads (Default: no variant)
  - **mismatch.label**: Label for single nucleotide variants (Default: SNV)
  - **rc**: Should the variants be displayed with respect to the negative strand? (Default: FALSE)
  - **keep.ops**: CIGAR operations to remain in the variant label (usually indels)
  - **upstream**: distance upstream of the cut site to call SNVs
  - **downstream**: distance downstream of the cut site to call SNVs

- **getInsertionSeqs(ref_ranges, genome_ranges)**

  Description: Set the "insertions" field - a table of the locations of insertions, and the "ins_key" field which relates sequences indices to the insertions they contain.

  Input parameters:
  - **ref_ranges**: The cigar operations of the reads with respect to the reference genome.
  - **genome_ranges**: The cigar operations of the reads with respect to the genome, i.e. the reference locations shifted to their genomic start locations

- **removeSeqs(idxs)**

  Description: Remove sequences from a CrisprRun object and from the internal CrisprRun fields that store insertion locations for plotting.

  Input parameters:
  - **idxs**: Indexes of reads to remove

Author(s)

Helen Lindsay

See Also

- **CrisprSet**

Examples

```r
# readsToTarget with signature("GAlignments", "GRanges") returns a
# CrisprRun object

bam_fname <- system.file("extdata", "gol_F1_clutch_1_embryo_1_s.bam",
                        package = "CrispRVariants")

param <- Rsamtools::ScanBamParam(what = c("seq", "flag"))
alns <- GenomicAlignments::readGAlignments(bam_fname, param = param,
                                         use.names = TRUE)

reference <- Biostrings::DNAString("GGTCTCTCGCAGGATGTTGCTGG")
```
CrisprSet-class

CrisprSet class

Description

A ReferenceClass container for holding a set of narrowed alignments, each corresponding to the same target region. Individual samples are represented as CrisprRun objects. CrisprRun objects with no on-target reads are excluded. CrisprSet objects are constructed with `readsToTarget` or `readsToTargets`. For most use cases, a CrisprSet object should not be accessed directly.

Arguments

- **crispr.runs**: A list of CrisprRun objects, typically representing individual samples within an experiment.
- **reference**: The reference sequence, must be the same length as the target region.
- **target**: The target location (GRanges). Variants will be counted over this region. Need not correspond to the guide sequence.
- **rc**: Should the alignments be reverse complemented, i.e. displayed w.r.t the reverse strand? (default: FALSE)
- **short.cigars**: If TRUE, variants labels are created from the location of their insertions and deletions. For variants with no insertions or deletions, the locations of any single base mismatches are displayed (default: TRUE).
- **names**: A list of names for each of the samples, e.g. for displaying in plots. If not supplied, the names of the crispr.runs are used, which default to the filenames of the bam files if available (Default: NULL)
- **renumbered**: Should the variants be renumbered using target.loc as the zero point? If TRUE, variants are described by the location of their 5'-most base with respect to the target.loc. A 3bp deletion starting 5bp 5' of the cut site would be labelled (using short.cigars) as -5:3D (Default: TRUE)
- **target.loc**: The location of the Cas9 cut site with respect to the supplied target. (Or some other central location). Can be displayed on plots and used as the zero point for renumbering variants. For a target region with the PAM location from bases 21-23, the target.loc is base 17 (default: NA)
- **match.label**: Label for sequences with no variants (default: "no variant")
- **mismatch.label**: Label for sequences with only single nucleotide variants (default: "SNV")
- **split.snv**: Should single nucleotide variants (SNVs) be shown for reads without an insertion or deletion? (default: TRUE)
- **upstream.snv**: If split.snv = TRUE, how many bases upstream of the target.loc should SNVs be shown? (default: 8)
- **downstream.snv**: If split.snv = TRUE, how many bases downstream of the target.loc should SNVs be shown? (default: 6)
- **verbose**: If true, prints information about initialisation progress (default: TRUE)
Fields

- **crispr_runs**: A list of CrisprRun objects, typically corresponding to samples of an experiment.
- **ref**: The reference sequence for the target region, as a Biostrings::DNAString object.
- **cigar_freqs**: A matrix of counts for each variant.
- **target**: The target location, as a GRanges object.

Methods

- **classifyCodingBySize(var_type, cutoff = 10)**: Description: This is a naive classification of variants as frameshift or in-frame Coding indels are summed, and indels with sum divisible by 3 are considered frameshift. Note that this may not be correct for variants that span an intron-exon boundary. Input parameters: var_type: A vector of var_type. Only variants with var_type == "coding" are considered. Intended to work with classifyVariantsByLoc. Variants are divided into those less than and greater than "cutoff" (Default: 10). Result: A character vector with a classification for each variant allele.

- **classifyVariantsByLoc(txdb, add_chr = TRUE, verbose = TRUE, ...)**: Description: Uses the VariantAnnotation package to look up the location of the variants. VariantAnnotation allows multiple classification tags per variant, this function returns a single tag. The following preference order is used: spliceSite > coding > intron > fiveUTR > threeUTR > promoter > intergenic. Input parameters: txdb: A BSgenome transcription database. add_chr: Add "chr" to chromosome names to make compatible with UCSC (default: TRUE). verbose: Print progress (default: TRUE). ...: Filtering arguments for variantCounts. Return value: A vector of classification tags, matching the rownames of .self$cigar_freqs (the variant count table).

- **classifyVariantsByType(...)**: Description: Classifies variants as insertions, deletions, or complex (combinations). In development. Input parameters: ... Optional arguments to "variantCounts" for filtering variants before classification. Return value: A named vector classifying variant alleles as insertions, deletions, etc.

- **consensusAlleles(cig_freqs = .self$cigar_freqs, return_nms = FALSE)**: Description: Get variants by their cigar string, make the pairwise alignments for the consensus sequence for each variant allele. Input parameters: cig_freqs: A table of variant allele frequencies (by default: .self$cigar_freqs, but could also be filtered). return_nms: If true, return a list of sequences and labels (Default: FALSE). Return: A DNAStringSet of the consensus sequences for the specified alleles, or a list containing the consensus sequences and names for the labels if return_nms = TRUE.

- **filterUniqueLowQual(min_count = 2, max_n = 0, verbose = TRUE)**: Description: Deletes reads containing rare variant combinations and more than a minimum number of ambiguity characters within the target region. These are assumed to be alignment errors. Input parameters: min_count: the number of times a variant combination must occur across all samples to keep (default: 2, i.e. a variant must occur at least twice in one or more samples to keep) max_n: maximum number of ambiguity ("N") bases a read with a rare variant combination may contain. (default: 0) verbose: If TRUE, print the number of sequences removed (default: TRUE).

- **filterVariants(cig_freqs = NULL, names = NULL, columns = NULL, include.chimeras = TRUE)**: Description: Relabels specified variants in a table of variant allele counts as non-variant, e.g. variants known to exist in control samples. Accepts either a size, e.g. "1D", or a specific mutation, e.g. "+4:3D". For alleles that include one variant to be filtered and one other variant,
the other variant will be retained. If SNVs are included, these will be removed entirely, but note that SNVs are only called in reads that do not contain an insertion/deletion variant.

Input parameters:
- `cig_freqs`: A table of variant allele counts (Default: NULL, i.e. `.self$cigar_freqs`)
- `names`: Labels of variants alleles to remove (Default: NULL) columns: Indices or names of control samples. Remove all variants that occur in these columns. (Default: NULL)
- `include.chimeras`: Should chimeric reads be included? (Default: TRUE)

`heatmapCigarFreqs(as.percent = TRUE, x.size = 8, y.size = 8, x.axis.title = NULL, x.angle = 90, ...)`

Description: Internal method for CrispRVariants::plotFreqHeatmap, optionally filters the table of variants, then a table of variant counts, coloured by counts or proportions.

Input parameters:
- `as.percent`: Should colours represent the percentage of reads per sample (TRUE) or the actual counts (FALSE)? (Default: TRUE)
- `x.size`: Font size for x axis labels (Default: 8)
- `y.size`: Font size for y axis labels (Default: 8)
- `x.axis.title`: Title for x axis
- `min.freq`: Include only variants with frequency at least min.freq in at least one sample
- `min.count`: Include only variants with count at least min.count in at least one sample
- `top.n`: Include only the n most common variants
- `type`: Should labels show counts or proportions? (Default: counts)
- `header`: What should be displayed in the header of the heatmap. Default: total count for type = "counts" or proportion of reads shown in the matrix for type = "proportions".

Return value: A ggplot2 plot object. Call "print(obj)" to display

See also: CrispRVariants::plotFreqHeatmap

`makePairwiseAlns(cig_freqs = .self$cigar_freqs, ...)` Description: Get variants by their cigar string, make the pairwise alignments for the consensus sequence for each variant allele.

Input parameters:
- `cig_freqs`: A table of variant allele frequencies (by default: `.self$cigar_freqs`, but could also be filtered)
- `...`: Extra arguments for CrispRVariants::seqsToAln, e.g. which symbol should be used for representing deleted bases

`mutationEfficiency(snv = c("non_variant", "include", "exclude"), include.chimeras = TRUE, exclude.cols = NULL, group = NULL, filter.vars = NULL, filter.cols = NULL, count.alleles = FALSE, per.sample = TRUE, min.freq = 0)`

Description: Calculates summary statistics for the mutation efficiency, i.e. the percentage of reads that contain a variant. Reads that do not contain and insertion or deletion, but do contain a single nucleotide variant (snv) can be considered as mutated, non-mutated, or not included in efficiency calculations as they are ambiguous.

Input parameters:
- `snv`: One of "include" (consider reads with mismatches to be mutated), "exclude" (do not include reads with snvs in efficiency calculations), and "non_variant" (consider reads with mismatches to be non-mutated). include.chimeras: Should chimeras be counted as variants? (Default: TRUE)
- `exclude.cols`: A list of column names to exclude from calculation, e.g. if one sample is a control (default: NULL, i.e. include all columns) group: A grouping variable. Efficiency will be calculated per group, instead of for individual. Cannot be used with exclude.cols. filter.vars: Variants that should not be counted as mutations. filter.cols: Column names to be considered controls. Variants occurring in a control sample will not be counted as mutations. count.alleles: If TRUE, also report statistics about the number of alleles per sample/per group. (Default: FALSE) per.sample: Return efficiencies for each sample (Default: TRUE) min.freq: Minimum frequency for counting alleles. Does not apply to calculating efficiency. To filter when calculating efficiency, first use "variantCounts". (Default: 0, i.e. no filtering)

Return value: A vector of efficiency statistics per sample and overall, or a matrix if a group is supplied.

`plotVariants(min.freq = 0, min.count = 0, top.n = nrow(.self$cigar_freqs), renumbered = .self$pars["renumbered"], add.other = add.other, ...)` Description: Internal method for CrispRVariants::plotAlignments, optionally filters the table of variants, then plots variants with respect to the reference sequence, collapsing insertions and displaying insertion sequences below the plot.
Input parameters: min.freq: i (integer) include variants that occur at least i times in at least one sample min.count i (integer) include variants that occur at least i times in at least one sample top.n: n (integer) Plot only the n most frequent variants (default: plot all) Note that if there are ties in variant ranks, top.n only includes ties with all members ranking <= top.n renumbered: If TRUE, the x-axis is numbered with respect to the target (cut) site. If FALSE, x-axis shows genomic locations. (default: TRUE) add.other Add a blank row named "Other" for chimeric alignments, if there are any (Default: TRUE) ... additional arguments for plotAlignments

Return value: A ggplot2 plot object. Call "print(obj)" to display

Author(s)

Helen Lindsay

See Also

readsToTarget and readsToTargets for initialising a CrisprSet, CrisprRun

Examples

# Load the metadata table
md_fname <- system.file("extdata", "gol_F1_metadata_small.txt", package = "CrispRVariants")
md <- read.table(md_fname, sep = "\t", stringsAsFactors = FALSE)

# Get bam filenames and their full paths
bam_fnames <- sapply(md$bam.filename, function(fn){
  system.file("extdata", fn, package = "CrispRVariants")})

reference <- Biostrings::DNAString("GGTCTCTCGCAGGATGTTGCTGG")
gd <- GenomicRanges::GRanges("18", IRanges::IRanges(4647377, 4647399), strand = "+")

crispr_set <- readsToTarget(bam_fnames, target = gd, reference = reference,
  names = md$experiment.name, target.loc = 17)

disableDots
dispatchDots

Description

Update default values for func with values from dot args

Usage

disableDots(func, ..., call = FALSE)

Arguments

func Function to call
... dot args to pass to function
call If TRUE, call the function with the argument list and return this result (Default: FALSE)
Value

A list of arguments to pass to `func`, or if `call` is TRUE, the result of calling `func` with these arguments.

Author(s)

Helen Lindsay

Examples

```r
# Set up a function to dispatch dot arguments to:
f <- function(a=1, b=2, c=3){
  print(c(a,b,c))
}
# Set up a function for passing dots:
g <- function(...){
  CrispRVariants:::dispatchDots(f, ...)
}

g(a = 5)
g(a = 5, call = TRUE)
# Unrelated arguments will not be passed on
  g(a = 5, d = 6)
```

---

### `excludeFromBam`

**Removes reads from a bam file**

**Description**

Returns a GAlignments excluding reads based on either name and/or location

**Usage**

```r
excludeFromBam(bam, exclude.ranges = GRanges(), exclude.names = NA)
```

**Arguments**

- `bam` a GAlignments object
- `exclude.ranges` Regions to exclude, as `GRanges`
- `exclude.names` A character vector of alignments names to exclude

**Value**

The bam minus the excluded regions

**Author(s)**

Helen Lindsay
findChimeras

Find chimeric reads

Description

Find chimeric reads, assuming that the GAlignments object does not contain multimapping reads. That is, read names that appear more than ones in the file are considered chimeras. Chimeric reads are reads that cannot be mapped as a single, linear alignment. Reads from structural rearrangements such as inversions can be mapped as chimeras. Note that the indices of all chimeric reads are returned, these are not separated into individual chimeric sets.

Usage

findChimeras(bam, by.flag = FALSE)

Arguments

bam

A GAlignments object, must include names

by.flag

Can the chimeras be detected just using the supplementary alignment flag? (Default: FALSE). If TRUE, detects supplementary alignments and returns reads with the same name as a supplementary alignment (quicker). If FALSE, all alignments with duplicated names are returned.

Value

A vector of indices of chimeric sequences within the original bam

Author(s)

Helen Lindsay

See Also

plotChimeras for plotting chimeric alignment sets.

Examples

bam_fname <- system.file("extdata", "gol_F1_clutch_2_embryo_4_s.bam", package = "CrispRVariants")
bam <- GenomicAlignments::readGAlignments(bam_fname, use.names = TRUE)
chimera_indices <- findChimeras(bam)
chimeras <- bam[chimera_indices]
**findSNVs**  
*Find frequent SNVs*

**Description**  
Find single nucleotide variants (SNVs) above a specified frequency in a table of variants.

**Usage**  
```r
findSNVs(obj, ...)  
## S4 method for signature 'CrisprSet'
findSNVs(obj, ..., freq = 0.25,  
         include.chimeras = TRUE)
```

**Arguments**  
- `obj`  
  An object containing variant counts  
- `...`  
  Additional arguments  
- `freq`  
  Minimum frequency snv to return (Default: 0.25)  
- `include.chimeras`  
  Include chimeric reads when calculating SNV frequencies (Default: TRUE)

**Value**  
A vector of SNVs and their frequencies

**Author(s)**  
Helen Lindsay

---

**getChimeras**  
*Get chimeric alignments*

**Description**  
Return chimeric alignments from a collection of aligned sequences

**Usage**  
```r
getChimeras(obj, ...)
## S4 method for signature 'CrisprSet'
getChimeras(obj, ..., sample)
```

**Arguments**  
- `obj`  
  An object containing aligned sequences  
- `...`  
  Additional arguments  
- `sample`  
  The sample name or sample index to return
Value
A GAlignment object containing the chimeric read groups

Author(s)
Helen Lindsay

Examples
```r
data("gol_clutch1")
chimeras <- getChimeras(gol, sample = 2)
```

Description
This dataset is a subset of the crispant data for the golden gene used by Burger et al (submitted).

Usage
```r
data(gol_clutch1)
```

Format
A CrisprSet object countaining 8 samples

Details
- gol The variants as a CrisprSet object

Value
A CrisprSet object named "gol"

makeAlignmentTilePlot
Internal CrispRVariants function for creating the plotAlignments background

Description
Takes a matrix of characters, x and y locations and colours, creates a ggplot geom_tile plot with tiles labelled by the characters.

Usage
```r
makeAlignmentTilePlot(m, ref, xlab, plot.text.size, axis.text.size, xtick.labs, xtick.breaks, tile.height)
```
mergeCrisprSets

Arguments

- **m**: A matrix with column headings Var1: y location, Var2: x location, cols: tile fill colour, isref: transparency value text_cols: text colour
- **ref**: The reference sequence, only used for checking the number of x-tick labels when x-tick breaks are not supplied
- **xlab**: Label for the x axis
- **plot.text.size**: Size for text within plot
- **axis.text.size**: Size for text on axes
- **xtick.labs**: x axis labels
- **xtick.breaks**: Locations of x labels
- **tile.height**: Controls whitespace between tiles

Value

A ggplot object

Author(s)

Helen Lindsay

mergeCrisprSets

Merge two CrisprSets

Description

Merge two CrisprSet objects sharing a reference and target location

Usage

```r
mergeCrisprSets(x, y, ...)
```

```r
## S4 method for signature 'CrisprSet,CrisprSet'
mergeCrisprSets(x, y, ..., x.samples = NULL, y.samples = NULL, names = NULL, order = NULL)
```

Arguments

- **x**: A CrisprSet object
- **y**: A second CrisprSet object
- **...**: extra arguments
- **x.samples**: A subset of column names or indices to keep from CrispRSet x (Default: NULL, i.e. keep all)
- **y.samples**: A subset of column names or indices to keep from CrispRSet y (Default: NULL, i.e. keep all)
- **names**: New names for the merged CrisprSet object (Default: NULL)
- **order**: A list of sample names, matching the names in x and y, specifying the order of the samples in the new CrisprSet. (Not implemented yet)
**mutationEfficiency**

*Get mutation efficiency*

**Value**

A merged CrisprSet object

**Author(s)**

Helen Lindsay

**Examples**

```r
# Load the metadata table
md_fname <- system.file("extdata", "gol_F1_metadata_small.txt", package = "CrispRVariants")
md <- read.table(md_fname, sep = \"\t\", stringsAsFactors = FALSE)

# Get bam filenames and their full paths
bam_fnames <- sapply(md$bam.filename, function(fn){
  system.file("extdata", fn, package = "CrispRVariants"))

reference <- Biostrings::DNAString("GGTCTCTCGCAGGATGTTGCTGG")
gd <- GenomicRanges::GRanges("18", IRanges::IRanges(4647377, 4647399),
  strand = "+")

crispr_set1 <- readsToTarget(bam_fnames[c(1:4)], target = gd,
  reference = reference, names = md$experiment.name[c(1:4)], target.loc = 17)
crispr_set2 <- readsToTarget(bam_fnames[c(5:8)], target = gd,
  reference = reference, names = md$experiment.name[c(5:8)], target.loc = 17)
mergeCrisprSets(crispr_set1, crispr_set2)
```

---

**Description**

Returns the percentage of sequences that contain at least one mutation.

**Usage**

```r
mutationEfficiency(obj, ...)
```

```r
# S4 method for signature 'CrisprSet'

mutationEfficiency(obj, ..., snv = c("non_variant", "include", "exclude"),
  include.chimeras = TRUE, exclude.cols = NULL,
  filter.vars = NULL, filter.cols = NULL, group = NULL)
```

**Arguments**

- `obj` An object containing variant counts
- `...` additional arguments
- `snv` Single nucleotide variants (SNVs) may be considered as mutations ("include"), treated as ambiguous sequences and not counted at all ("exclude"), or treated as non-mutations, e.g. sequencing errors or pre-existing SNVs ("non_variant", default)
narrowAlignments

include.chimeras
Should chimeric alignments be counted as variants when calculating mutation efficiency (Default: TRUE)

exclude.cols
A vector of names of columns in the variant counts table that will not be considered when counting mutation efficiency

filter.vars
Variants to remove before calculating efficiency. May be either a variant size, e.g. "1D", or a particular variant/variant combination, e.g. -5:3D

filter.cols
A vector of control sample names. Any variants present in the control samples will be counted as non-variant, unless they also contain another indel. Note that this is not compatible with counting snvs as variants.

group
A grouping vector. If provided, efficiency will be calculated per group (Default: NULL)

Value
A vector of efficiency statistics per sample and overall, or a matrix of efficiency statistics per group if a group is provided

Author(s)
Helen Lindsay

Examples

data("gol_clutch1")
mutationEfficiency(gol)

---

narrowAlignments

Narrow a set of aligned reads to a target region

Description

Aligned reads are narrowed to the target region. In the case of reads with deletions spanning the boundaries of the target, reads are narrowed to the start of the deletion.

Usage

narrowAlignments(alns, target, ...)

## S4 method for signature 'GAlignments,GRanges'
narrowAlignments(alns, target, ..., reverse.complement, verbose = FALSE)

Arguments

alns
A GAlignments object including a metadata column "seq" containing the sequence

target
A GRanges object

... additional arguments

reverse.complement
Should the aligned reads be reverse complemented?

verbose
(Default: FALSE)
plotAlignments

Value

The narrowed alignments (GAlignments).

Author(s)

Helen Lindsay

Examples

bam_fname <- system.file("extdata", "gol_F1_clutch_2_embryo_4_s.bam", package = "CrispRVariants")

bam <- GenomicAlignments::readGAlignments(bam_fname, use.names = TRUE)

target <- GenomicRanges::GRanges("18", IRanges::IRanges(4647377, 4647399), strand = "+")

narrowAlignments(bam, target, reverse.complement = FALSE)

plotAlignments

Plot alignments with respect to a reference sequence

Description

(signature("CrisprSet")) Wrapper for CrisprSet$plotVariants. Optionally filters a CrisprSet frequency table, then plots variants. More information in CrisprSet

(signature("DNAString")) Plots a set of pairwise alignments to a reference sequence. Alignments should all be the same length as the reference sequences. This is achieved by removing insertions with respect to the reference, see seqsToAln. Insertions are indicated by symbols in the plot and a table showing the inserted sequences below the plot. The default options are intended for a figure 6-8 inches wide, with figure height best chosen according to the number of different variants and insertions to be displayed.

Usage

plotAlignments(obj, ...)

## S4 method for signature 'CrisprSet'
plotAlignments(obj, ..., min.freq = 0, min.count = 1,
  top.n = 50, renumbered = obj$pars["renumbered"], add.other = TRUE)

## S4 method for signature 'DNAString'
plotAlignments(obj, ..., alns, ins.sites,
  highlight.pam = TRUE, show.plot = FALSE, target.loc = 17,
  pam.start = NA, pam.end = NA, ins.size = 2, legend.cols = 3,
  xlab = NULL, xtick.labs = NULL, xtick.breaks = NULL,
  plot.text.size = 2, axis.text.size = 8, legend.text.size = 6,
  highlight.guide = TRUE, guide.loc = NULL, tile.height = 0.55,
  max.insertion.size = 20, min.insertion.freq = 5, line.weight = 1,
  legend.symbol.size = ins.size, add.other = FALSE, codon.frame = NULL)
Arguments

obj  The object to be plotted

...  Additional arguments

min.freq  i (one sample (default: 0, i.e. no frequency cutoff)

min.count  i (integer) only plot variants with count >= i in at least one sample (default: 0, i.e. no count cutoff)

top.n  (integer) Plot only the n most frequent variants (default: 50)

renumbered  If TRUE, the x-axis is numbered with respect to the target (default: TRUE)

add.other  Add a blank row labelled "Other" to the plot, for combining with plotFreqHeatmap (default: TRUE (signature "CrisprSet") FALSE (signature "matrix"))

alns  A named character vector of aligned sequences, with insertions removed

ins.sites  A table of insertion.sites, which must include cols named "start", "cigar" and "seq", for the start of the insertion in the corresponding sequence

highlight.pam  should location of PAM with respect to the target site be indicated by a box? (Default: TRUE) If TRUE, and pam.start and pam.end are not supplied, PAM is inferred from target.loc

show.plot  Should the plot be displayed (TRUE) or just returned as a ggplot object (FALSE). (Default: FALSE)

target.loc  The location of the zero point / cleavage location. Base n, where the zero point is between bases n and n+1

pam.start  The first location of the PAM with respect to the reference.

pam.end  The last location of the PAM with respect to the reference. Default is two bases after the pam.start

ins.size  The size of the symbols representing insertions within the plot.

legend.cols  The number of columns in the legend. (Default: 3)

xlab  A title for the x-axis (Default: NULL)

xtick.labs  Labels for the x-axis ticks (Default: NULL)

xtick.breaks  Locations for x-axis tick breaks (Default: NULL)

plot.text.size  The size of the text inside the plot

axis.text.size  The size of the axis labels

legend.text.size  The size of the legend labels

highlight.guide  Should the guide be indicated by a box in the reference sequence? (Default: TRUE)

guide.loc  The location of the guide region to be highlighted, as an IRanges object. Will be inferred from target.loc if highlight.guide = TRUE and no guide.loc is supplied, assuming the guide plus PAM is 23bp (Default: NULL)

tile.height  The height of the tiles within the plot. (Default: 0.55)

max.insertion.size  The maximum length of an insertion to be shown in the legend. If max.insertion.size = n, an insertion of length m > n will be annotated as "mI" in the figure. (Default: 20)
plotChimeras

min.insertion.freq
Display inserted sequences with frequency at least x amongst the sequences with an insertion of this size and length (Default: 5)

line.weight
The line thickness for the vertical line indicating the zero point (cleavage site) and the boxes for the guide and PAM. (Default: 1)

legend.symbol.size
The size of the symbols indicating insertions in the legend. (Default: ins.size)

codon.frame
Codon position of the leftmost nucleotide. If provided, codon positions in the specified frame are indicated. (Default: NULL)

Value
A ggplot2 figure

Author(s)
Helen Lindsay

See Also
seqsToAln, ggplot

Examples

#Load a CrisprSet object and plot
data("gol_clutch1")
plotAlignments(gol)

plotChimeras Display a dot plot of chimeric alignments

Description

Produces a dot plot of a set of chimeric alignments. For chimeric alignments, a single read is split into several, possibly overlapping aligned blocks. Aligned sections of chimeric reads can be separated by large genomic distances, or on separate chromosomes. plotChimeras produces a dot plot, each aligned block highlighted, and chromosomes shown in different colours. Large gaps between aligned segments are collapsed and indicated on the plot with horizontal lines. The X-axis shows each base of the entire read. Note that the mapping to the fwd strand is shown if all strands agree. The chimeric alignments must be sorted!

Usage

plotChimeras(chimeric.alns, max.gap = 10, tick.sep = 20, text.size = 10, title.size = 16, gap.pad = 20, legend.title = "Chromosome", xangle = 90, wrt.forward = FALSE, annotate.within = 20, annotations = GenomicRanges::GRanges())
plotChimeras

Arguments

chimeric.alns  A GAlignments object containing only the chimeric reads to be plotted
max.gap     If aligned segments are separated by more than max.gap,
tick.sep    How many bases should separate tick labels on plot. Default 20.
text.size   Size of X and Y tick labels on plot. Default 12
title.size  Size of X and Y axis labels on plot. Default 16
gap.pad     How much should aligned blocks be separated by? (Default: 20)
legend.title Title for the legend. Default "Chromosome"
xangle      Angle for x axis text (Default 90, i.e vertical)
wrt.forward Should chimeric alignments where all members map to the negative strand be
             displayed with respect to the forward strand, i.e. as the cigar strand is written
             (TRUE), or the negative strand (FALSE) (Default: FALSE)
annotate.within annot_aln ranges in "annotations" within n bases of a chimeric alignment (De-
                   fault 50)
annotations  A list of GRanges. Any that overlap with the chimeric alignments are highlighed
             in the plot.

Value

A ggplot2 dotplot of the chimeric alignments versus the reference sequence

Author(s)

Helen Lindsay

See Also

findChimeras for finding chimeric alignment sets.

Examples

bam_fname <- system.file("extdata", "gol_F1_clutch_2_embryo_4_s.bam",
                         package = "CrispRVariants")
bam <- GenomicAlignments::readGAlignments(bam_fname, use.names = TRUE)
# Choose a single chimeric read set to plot:
chimeras <- bam[names(bam) == "AB3092"]

# This read aligns in 3 pieces, all on chromosome 18.
# The plot shows the alignment annot_alns a small duplication and
# a long gap.
plotChimeras(chimeras)
Description

Creates a heatmap from a matrix of counts or proportions, where tiles are coloured by the proportion and labeled with the value.

Usage

plotFreqHeatmap(obj, ...)

## S4 method for signature 'matrix'
plotFreqHeatmap(obj, ..., col.sums = NULL, header = NA,
    header.name = "Total", group = NULL, group.colours = NULL,
    as.percent = TRUE, x.axis.title = NULL, x.size = 6, y.size = 8,
    x.angle = 90, legend.text.size = 6, plot.text.size = 3,
    line.width = 1, x.hjust = 1, legend.position = "right",
    x.labels = NULL, legend.key.height = grid::unit(1, "lines"))

## S4 method for signature 'CrisprSet'
plotFreqHeatmap(obj, ..., top.n = 50, min.freq = 0,
    min.count = 1, type = c("counts", "proportions"), order = NULL)

Arguments

obj A matrix of counts with rows = feature, columns = sample
...
additional arguments
col.sums Alternative column sums to be used for calculating the tile colours if as.percent = TRUE, e.g. if "obj" is a subset of a larger data set. If "NULL" (default), the column sums of "obj" are used.
header Alternative column titles, e.g. column sums for the unfiltered data set when obj is a subset. If set to "NA", column sums of obj are displayed. If "NULL", no header is displayed (Default: NA).
header.name Label for the header row (Default: "Total")
group Grouping factor for columns. If supplied, columns are ordered to match the levels (Default: NULL)
group.colours Colours for column groups, should match levels of "group". If "NULL", groups are coloured differently (Default: NULL)
as.percent Should colours represent the percentage of reads per sample (TRUE) or the actual counts (FALSE)? (Default: TRUE)
x.axis.title A title for the x-axis. (Default: NULL)
x.size Font size for x-labels (Default: 16)
y.size Font size for y-labels (Default: 16)
x.angle Angle for x-labels (Default: 90, i.e. vertical)
legend.text.size Font size for legend (Default: 16)
plotVariants

plot.text.size Font size counts within plot (Default: 8)
line.width Line thickness of title box'
x.hjust Horizontal justification of x axis labels (Default: 1)
legend.position The position of the legend (Default: right)
x.labels X-axis labels (Default: NULL, column.names of the matrix, doesn’t do anything at the moment)
legend.key.height The height of the legend key, as a "unit" object. (See unit).
top.n Show the n top ranked variants. Note that if the nth and n+1th variants have equal rank, they will not be shown. (Default: 50)
min.freq i ( one sample (default: 0, i.e no frequency cutoff)
min.count i (integer) only plot variants with count >= i in at least one sample (default: 0, i.e no count cutoff)
type Plot either "counts" or "proportions"
order A list of column names or indices specifying the order of the columns in the plot

Value

The ggplot2 plot of the variant frequencies

Examples

#Load a CrisprSet object for plotting
data("gol_clutch1")

# Plot the frequency heatmap
plotFreqHeatmap(gol)

Description

Combines a plot of transcript structure, alleles aligned with respect to a reference genome and a heatmap of counts or proportions of each allele in a set of data.

Usage

plotVariants(obj, ...)

## S4 method for signature 'CrisprSet'
plotVariants(obj, ..., txdb = NULL, add.chr = TRUE,
plotAlignments.args = list(), plotFreqHeatmap.args = list())
**rcAlns**

**rcAlns**

**Arguments**

- **obj**
  - The object to be plotted
- **...**
  - extra arguments for plot layout
- **txdb**
  - GenomicFeatures::TxDb object (default: NULL)
- **add.chr**
  - If target chromosome does not start with "chr", e.g. "chr5", add the "chr" prefix. (Default: TRUE)
- **plotAlignments.args**
  - Extra arguments for plotAlignments
- **plotFreqHeatmap.args**
  - Extra arguments for plotFreqHeatmap

**Value**

A ggplot2 plot of the variants

**See Also**

- arrangePlots for general layout options and annotateGenePlot for options relating to the transcript plot.

**Examples**

```r
#Load a CrisprSet object for plotting
data("gol_clutch1")

#Load the transcript db. This is a subset of the Ensembl Danio Rerio v73 gtf
# for the region 18:4640000-4650000 which includes the targeted gol gene

library(GenomicFeatures)
fn <- system.file("extdata", "Danio_rerio.Zv9.73.gol.sqlite", package = "CrispRVariants")
txdb <- loadDb(fn)

# Plot the variants
p <- plotVariants(gol, txdb = txdb)

#In the above plot, the bottom margin is too large, the legend is
cut off, and the text within the plots should be larger.
#These issues can be fixed with some adjustments:
p <- plotVariants(gol, txdb = txdb,
  plotAlignments.args = list(plot.text.size = 4, legend.cols = 2),
  plotFreqHeatmap.args = list(plot.text.size = 4),
  left.plot.margin = grid::unit(c(0.1,0,0.5,1), "lines"))
```

---

**rcAlns**

**Internal CrispRVariants function for determining read orientation**

**Description**

Function for determining whether reads should be oriented to the target strand, always displayed on the positive strand, or oriented to
Usage
rcAlns(target.strand, orientation)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>target.strand</td>
<td>The target strand (one of &quot;+&quot;,&quot;-&quot;,&quot;*&quot;)</td>
</tr>
<tr>
<td>orientation</td>
<td>One of &quot;target&quot;, &quot;opposite&quot; and &quot;positive&quot; (Default: &quot;target&quot;)</td>
</tr>
</tbody>
</table>

Value
A logical value indicating whether reads should be reverse complemented

Author(s)
Helen Lindsay

Description
Short reads amplified with PCR primers should start and end at defined positions. However, the ends of an aligned read may be clipped as sequencing technologies are prone to making errors at the start and end. readsByPCRPrimer extrapolates the genomic location of entire reads from their aligned sections by adding clipped sections, then finds near exact matches to a set of PCR primers. Note that this is not always a good assumption, and is misleading in the case of chimeric reads where sections clipped in one part of a chimera are aligned in another.

Usage
readsByPCRPrimer(bam, primers, ...)

## S4 method for signature 'GAlignments,GRanges'
readsByPCRPrimer(bam, primers, ..., tolerance = 0, verbose = TRUE, ignore.strand = TRUE, allow.partial = TRUE, chimera.idxs = NULL)

## S4 method for signature 'GRanges,GRanges'
readsByPCRPrimer(bam, primers, ..., tolerance = 0, verbose = TRUE, ignore.strand = TRUE, allow.partial = TRUE, chimera.idxs = NULL)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>bam</td>
<td>A set of aligned reads</td>
</tr>
<tr>
<td>primers</td>
<td>A set of ranges that the unclipped reads may map to</td>
</tr>
<tr>
<td>...</td>
<td>Additional arguments</td>
</tr>
<tr>
<td>tolerance</td>
<td>Number of bases by which reads and primers may differ at each end (Default: 0)</td>
</tr>
<tr>
<td>verbose</td>
<td>Print number of full and partial matches (Default: TRUE)</td>
</tr>
</tbody>
</table>
ignore.strand: Passed to `findOverlaps` and `disjoin`. Should strand be ignored when finding overlaps? (Default: TRUE)

allow.partial: Should reads that do not match the PCR boundaries, but map to a region covered by only one primer be considered matches? (Default: TRUE)

chimera.idxs: Indices of chimeric reads within the bam. If specified, chimeras overlapping multiple pcr primers will be removed.

Value

A ` Hits` object where "query" is the index with respect to bam and "subject" is the index with respect to the primers.

Author(s)

Helen Lindsay

See Also

`GRanges`, `GAlignments`

### Description

Trims aligned reads to one or several target regions, optionally reverse complementing the alignments.

### Usage

`readsToTarget(reads, target, ...)`

```r
## S4 method for signature 'GAlignments,GRanges'
readsToTarget(reads, target, ..., reverse.complement = TRUE, chimeras = NULL, collapse.pairs = FALSE, use.consensus = FALSE, store.chimeras = FALSE, verbose = TRUE, name = NULL, orientation = c("target", "opposite", "positive"))
```

```r
## S4 method for signature 'GAlignmentsList,GRanges'
readsToTarget(reads, target, ..., reference = reference, names = NULL, reverse.complement = TRUE, target.loc = 17, chimeras = NULL, collapse.pairs = FALSE, use.consensus = FALSE, orientation = c("target", "opposite", "positive"), verbose = TRUE)
```

```r
## S4 method for signature 'character,GRanges'
readsToTarget(reads, target, ..., reference, reverse.complement = TRUE, target.loc = 17, exclude.ranges = GRanges(), exclude.names = NA, chimeras = c("count", "exclude", "ignore", "merge"), collapse.pairs = FALSE, use.consensus = FALSE, orientation = c("target", "opposite", "positive"), names = NULL, verbose = TRUE)
```
readsToTargets(reads, targets, ...)

## S4 method for signature 'character,GRanges'
readsToTargets(reads, targets, ..., references,
primer.ranges = NULL, target.loc = 17, reverse.complement = TRUE,
collapse.pairs = FALSE, use.consensus = FALSE, ignore.strand = TRUE,
names = NULL, bpparam = BiocParallel::SerialParam(),
orientation = c("target", "opposite", "positive"), chimera.to.target = 5,
verbose = TRUE)

## S4 method for signature 'GAlignmentsList,GRanges'
readsToTargets(reads, targets, ..., references, primer.ranges = NULL, target.loc = 17,
reverse.complement = TRUE, collapse.pairs = FALSE,
use.consensus = FALSE, ignore.strand = TRUE, names = NULL,
bpparam = BiocParallel::SerialParam(), chimera.to.target = 5,
orientation = c("target", "opposite", "positive"), verbose = TRUE)

Arguments

reads A GAlignments object, or a character vector of the filenames

target A GRanges object specifying the range to narrow alignments to

... Extra arguments for initialising CrisprSet

reverse.complement

(Default: TRUE) Should the alignments be oriented to match the strand of the target? If TRUE, targets located strand and targets on the negative strand with respect to the negative strand. If FALSE, the parameter 'orientation' must be set to determine the orientation. 'reverse.complement' will be replaced by 'orientation' in a later release.

chimeras Flag to determine how chimeric reads are treated. One of "ignore", "exclude", and "merge". Default "count", "merge" not implemented yet

collapse.pairs If reads are paired, should pairs be collapsed? (Default: FALSE) Note: only collapses primary alignments, and assumes that there is only one primary alignment per read.

use.consensus Take the consensus sequence for non-matching pairs? If FALSE, the sequence of the first read is used. Can be very slow. (Default: FALSE)

store.chimeras Should chimeric reads be stored? (Default: FALSE)

verbose Print progress and statistics (Default: TRUE)

name An experiment name for the reads. (Default: NULL)

orientation One of "target" (reads are displayed on the same strand as the target) "opposite" (reads are displayed on the opposite) strand from the target or "positive" (reads are displayed on the forward strand regardless of the strand of the target) (Default:"target")

reference The reference sequence

names Experiment names for each bam file. If not supplied, filenames are used.

target.loc The zero point for renumbering (Default: 17)

exclude.ranges Ranges to exclude from consideration, e.g. homologous to a pcr primer.
includeTargetBam

exclude.names  Alignment names to exclude
targets        A set of targets to narrow reads to
references     A set of reference sequences matching the targets. References for negative
                strand targets should be on the negative strand.
primer.ranges  A set of GRanges, corresponding to the targets. Read lengths are typically
greater than target regions, and it can be that reads span multiple targets. If
primer.ranges are available, they can be used to assign such reads to the correct
target.
ignore.strand  Should strand be considered when finding overlaps? (See findOverlaps )
bpparam        A BiocParallel parameter for parallelising across reads. Default: no parallelisa-
tion. (See bpparam)
chimera.to.target  Number of bases that may separate a chimeric read set from the target.loc for it
to be assigned to the target. (Default: 5)

Value
   (signature("GAlignments", "GRanges")) A CrispRun object
   (signature("character", "GRanges")) A CrispSet object

Author(s)
   Helen Lindsay

Examples

# Load the metadata table
md_fname <- system.file("extdata", "gol_F1_metadata_small.txt", package = "CrispRVariants")
md <- read.table(md_fname, sep = "\t", stringsAsFactors = FALSE)

# Get bam filenames and their full paths
bam_fnames <- sapply(md$bam.filename, function(fn){
  system.file("extdata", fn, package = "CrispRVariants"))

reference <- Biostrings::DNAString("GGTCTCTCGCAGGATGTTGCTGG")
gd <- GenomicRanges::GRanges("18", IRanges::IRanges(4647377, 4647399),
                        strand = "+")
crispr_set <- readsToTarget(bam_fnames, target = gd, reference = reference,
                        names = md$experiment.name, target.loc = 17)
reverseCigar

Usage

```
readTargetBam(file, target, exclude.ranges = GRanges(), exclude.names = NA,
             chimera.to.target = 5, chimeras = c("count", "ignore", "exclude",
             "merge"), by.flag = TRUE, verbose = TRUE)
```

Arguments

- **file**: The name of a bam file to read in
- **target**: A GRanges object containing a single target range
- **exclude.ranges**: A GRanges object of regions that should not be counted, e.g. primer or cloning vector sequences that have a match in the genome
- **exclude.names**: A vector of read names to exclude.
- **chimera.to.target**: Maximum distance between endpoints of chimeras and target.loc for assigning chimeras to targets (default: 5)
- **chimeras**: Flag to determine how chimeric reads are treated. One of "ignore", "exclude", "count" and "merge". Default "ignore".
- **by.flag**: Is the supplementary alignment flag set? Used for identifying chimeric alignments, function is much faster if TRUE. Not all aligners set this flag. If FALSE, chimeric alignments are identified using read names (Default: TRUE)
- **verbose**: Print stats about number of alignments read and filtered. (Default: TRUE)

Value

A GenomicAlignments::GAlignment obj

---

reverseCigar

Reverses the order of operations in a cigar string

Description

For example, the string "20M5D15M" would become "15M5D20M"

Usage

```
reverseCigar(cigars)
```

Arguments

- **cigars**: the cigar strings.

Value

The reversed cigar string
**rmMultiPCRChimera**

Remove chimeric reads overlapping multiple primers

**Description**

Finds and removes sets of chimeric read alignments that overlap more than one guide, i.e. that cannot be unambiguously assigned to a single guide.

**Usage**

```r
rmMultiPCRChimera(readnames, pcrhits, chimera_idxs, ...)
```

## S4 method for signature 'character,Hits,integer'

```r
rmMultiPCRChimera(readnames, pcrhits, chimera_idxs, ..., verbose = TRUE)
```

**Arguments**

- `readnames` A set of read names, used for identifying chimeric read sets
- `pcrhits` A mapping between indices of reads and a set of PCR primers
- `chimera_idxs` location of chimeric reads within the bam
- `...` Additional arguments
- `verbose` Display information about the chimeras (Default: TRUE)

**Value**

pcrhits, with chimeric reads mapping to different primers omitted.

**Author(s)**

Helen Lindsay

---

**seqsToAln**

Creates a text alignment from a set of cigar strings

**Description**

Creates a one-to-one text alignment of a set of cigar strings with respect to the reference sequence by collapsing insertions and introducing gaps across deletions.

When genomic coordinates for the alignment start and the target region are provided, aligned sequences are cropped to the target region.

**Usage**

```r
seqsToAln(cigar, dnaseq, target, del_char = "-", aln_start = NULL, reverse_complement = FALSE)
```
setDNATileColours

Arguments

cigar  A list of cigar strings to align
dnaseq  The set of sequences corresponding to the cigars, as Biostrings::DNAStrings
target  The target region to return, as GRanges. Sequences overlapping the target region are trimmed to exactly match it.
del_char  The character to represent deleted bases. Default "-"

aln_start  Genomic start locations of aligned sequences. Should be used in conjunction with target_start and target_end.

reverse_complement  (Default: FALSE)

Value

The sequences with insertions collapsed and deletions padded

Author(s)

Helen Lindsay

---

setDNATileColours  Sets colours for plotting aligned DNA sequences.

Description

Sets tile colours for plotAlignments with a DNA alphabet. Colour names must be valid.

Usage

setDNATileColours(m)

Arguments

m  A matrix with a column named "value" of the characters at each tile position.

Value

A matrix with additional columns specifying tile and text colours

Author(s)

Helen Lindsay
transformAlnsToLong  Transform data for plotting

Description

Orders and transforms a reference sequence and a set of aligned sequences into long format, i.e. one observation (tile position) per row. Used internally by plotAlignments.

Usage

transformAlnsToLong(ref, alns, add.other = FALSE)

Arguments

ref The reference sequence
alns Character vector of aligned sequences
add.other Add a blank row labelled "Other" (Default: FALSE)

Value

A matrix of characters and plotting locations

Author(s)

Helen Lindsay

variantCounts  Get variant counts

Description

Returns a matrix of counts where rows are sequence variants and columns are samples

Usage

variantCounts(obj, ...)

## S4 method for signature 'CrisprSet'
variantCounts(obj, ..., top.n = NULL, min.freq = 0,
              min.count = 1, include.chimeras = TRUE, include.nonindel = TRUE,
              result = "counts", filter.vars = NULL)
**Arguments**

- **obj**: An object containing variant counts
- **...**: Additional arguments
- **top.n**: (Integer n) If specified, return variants ranked at least n according to frequency across all samples (Default: 0, i.e. no cutoff)
- **min.freq**: (Float n) least one sample (Default: 0)
- **min.count**: (Integer n) Return variants with count greater than n in at least one sample (Default: 0)
- **include.chimeras**: Should chimeric reads be included in the counts table? (Default: TRUE)
- **include.nonindel**: Should sequences without indels be returned? (Default: TRUE)
- **result**: Return variants as either counts ("counts", default) or proportions ("proportions")
- **filter.vars**: Labels of variants alleles to remove (Default: NULL)

**Value**

A matrix of counts where rows are variants and columns are samples

**Author(s)**

Helen Lindsay

**Examples**

```r
data("gol_clutch1")

#Return a matrix of the 5 most frequent variants
variantCounts(gol, top.n = 5)
```

---

**Description**

Used by abifToFastq to write sanger sequences to fastq format. As abifToFastq appends output to files, writeFastq checks that sequence names are unique. This function is faster with checking switched off.

**Usage**

```r
writeFastq(outf, vals, allow_spaces = FALSE, check = TRUE)
```
writeFastq

Arguments

- **outf**: Name of fastq file to append sequence
- **vals**: A list containing entries named "seq" (sequence) and "quals" (quality scores, in ASCII format)
- **allow_spaces**: Should spaces in the sequence name be substituted with underscores? TRUE or FALSE
- **check**: Check whether reads with the same name already exist in the output fastq. (Default: TRUE)

Value

None. The sequences in "vals" are written to outf

Author(s)

Helen Lindsay
Index

*Topic datasets
  * gol_clutch1, 19

abifToFastq, 2
addClipped, 3
addClipped, GAlignments-method (addClipped), 3
addCodonFrame, 4
annotateGenePlot, 4, 29
arrangePlots, 5, 29
barplotAlleleFreqs, 6
barplotAlleleFreqs, CrisprSet-method (barplotAlleleFreqs), 6
barplotAlleleFreqs, matrix-method (barplotAlleleFreqs), 6
bpparam, 33
collapsePairs, 7
consensusSeqs, 8
consensusSeqs, CrisprSet-method (consensusSeqs), 8
countDeletions, 9
countDeletions, GAlignments-method (countDeletions), 9
countIndels (countDeletions), 9
countIndels, GAlignments-method (countDeletions), 9
countInsertions (countDeletions), 9
countInsertions, GAlignments-method (countDeletions), 9
CrisprRun, 15, 33
CrisprRun (CrisprRun-class), 10
CrisprRun-class, 10
CrisprSet, 11, 23, 33
CrisprSet (CrisprSet-class), 12
CrisprSet-class, 12
disjoin, 31
dispatchDots, 15
ecludeFromBam, 16
findChimeras, 17, 26
findOverlaps, 31, 33
findSNVs, 18
findSNVs, CrisprSet-method (findSNVs), 18
GAlignments, 31
getchimeras, 18
getchimeras, CrisprSet-method (getchimeras), 18
ggplot, 25
gol (gol_clutch1), 19
gol_clutch1, 19
GRanges, 4, 16, 31

Hits, 31

indelPercent (countDeletions), 9
indelPercent, GAlignments-method (countDeletions), 9
makeAlignmentTilePlot, 19
mergeCrisprSets, 20
mergeCrisprSets, CrisprSet, CrisprSet-method (mergeCrisprSets), 20
mutationEfficiency, 21
mutationEfficiency, CrisprSet-method (mutationEfficiency), 21

narrowAlignments, 22
narrowAlignments, GAlignments, GRanges-method (narrowAlignments), 22

plotAlignments, 23, 36, 37
plotAlignments, CrisprSet-method (plotAlignments), 23
plotAlignments, DNAString-method (plotAlignments), 23
plotChimeras, 17, 25
plotFreqHeatmap, 27
plotFreqHeatmap, CrisprSet-method (plotFreqHeatmap), 27
plotFreqHeatmap, matrix-method (plotFreqHeatmap), 27
plotVariants, 28
plotVariants, CrisprSet-method (plotVariants), 28

40
rcAlns, 29
readsByPCRPrimer, 30
readsByPCRPrimer, GAlignments, GRanges-method
  (readsByPCRPrimer), 30
readsByPCRPrimer, GRanges, GRanges-method
  (readsByPCRPrimer), 30
readsToTarget, 12, 15, 31
readsToTarget, character, GRanges-method
  (readsToTarget), 31
readsToTarget, GAlignments, GRanges-method
  (readsToTarget), 31
readsToTarget, GAlignmentsList, GRanges-method
  (readsToTarget), 31
readsToTargets, 12, 15
readsToTargets (readsToTarget), 31
readsToTargets, character, GRanges-method
  (readsToTarget), 31
readsToTargets, GAlignmentsList, GRanges-method
  (readsToTarget), 31
readTargetBam, 33
reverseCigar, 34
rmMultiPCRChimera, 35
rmMultiPCRChimera, character, Hits, integer-method
  (rmMultiPCRChimera), 35
seqsToAln, 23, 25, 35
setDNATileColours, 36
transformAlnsToLong, 37
unit, 28

variantCounts, 37
variantCounts, CrispSet-method
  (variantCounts), 37

writeFastq, 38