Package 'GUIDEseq'

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Type Package

Title GUIDE-seq and PEtag-seq analysis pipeline

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Depends R (>= 3.5.0), GenomicRanges, BiocGenerics

Imports Biostrings, pwalign, CRISPRseek, ChIPpeakAnno, data.table, matrixStats, BSgenome, parallel, IRanges (>= 2.5.5), S4Vectors (>= 0.9.6), stringr, multtest, GenomicAlignments (>= 1.7.3), GenomeInfoDb, Rsamtools, hash, limma,dplyr, GenomicFeatures, rio, tidyr, tools, methods, purrr, ggplot2, openxlsx, patchwork, rlang

biocViews ImmunoOncology, GeneRegulation, Sequencing, WorkflowStep, CRISPR

Suggests knitr, RUnit, BiocStyle, BSgenome.Hsapiens.UCSC.hg19, BSgenome.Hsapiens.UCSC.hg38, TxDb.Hsapiens.UCSC.hg19.knownGene, org.Hs.eg.db, testthat (>= 3.0.0)

VignetteBuilder knitr

Description The package implements GUIDE-seq and PEtag-seq analysis workflow including functions for filtering UMI and reads with low coverage, obtaining unique insertion sites (proxy of cleavage sites), estimating the locations of the insertion sites, aka, peaks, merging estimated insertion sites from plus and minus strand, and performing off target search of the extended regions around insertion sites with mismatches and indels.

License GPL (>= 2)

LazyLoad yes

NeedsCompilation no

Config/testthat/edition 3

2 GUIDEseq-package

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Description

The package includes functions to retain one read per unique molecular identifier (UMI), filter reads lacking integration oligo sequence, identify peak locations (cleavage sites) and heights, merge peaks, perform off-target search using the input gRNA. This package leverages CRISPRseek and ChIPpeakAnno packages.

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Details

Package: GUIDEseq
Type: Package
Version: 1.0
Date: 2015-09-04

License: GPL (>= 2)

Function GUIDEseqAnalysis integrates all steps of GUIDE-seq analysis into one function call

Author(s)

Lihua Julie Zhu Maintainer:julie.zhu@umassmed.edu

References

Shengdar Q Tsai and J Keith Joung et al. GUIDE-seq enables genome-wide profiling of off-target cleavage by CRISPR-Cas nucleases. Nature Biotechnology 33, 187 to 197 (2015)

See Also

GUIDEseqAnalysis

Examples

```
if(interactive())
     library("BSgenome.Hsapiens.UCSC.hg19")
     umiFile <- system.file("extdata", "UMI-HEK293_site4_chr13.txt",</pre>
        package = "GUIDEseq")
     alignFile <- system.file("extdata", "bowtie2.HEK293_site4_chr13.sort.bam" ,</pre>
         package = "GUIDEseq")
     gRNA.file <- system.file("extdata","gRNA.fa", package = "GUIDEseq")</pre>
     guideSeqRes <- GUIDEseqAnalysis(</pre>
         alignment.inputfile = alignFile,
         umi.inputfile = umiFile, gRNA.file = gRNA.file,
         orderOfftargetsBy = "peak_score",
         descending = TRUE,
         keepTopOfftargetsBy = "predicted_cleavage_score",
         scoring.method = "CFDscore",
         BSgenomeName = Hsapiens, min.reads = 80, n.cores.max = 1)
     guideSeqRes$offTargets
}
```

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annotateOffTargets Annotate offtargets with gene name

Description

Annotate offtargets with gene name and whether it is inside an exon

Usage

```
annotateOffTargets(thePeaks, txdb, orgAnn)
```

Arguments

thePeaks Output from offTargetAnalysisOfPeakRegions

txdb TxDb object, for creating and using TxDb object, please refer to GenomicFea-

tures package. For a list of existing TxDb object, please search for annotation

package starting with Txdb at http://www.bioconductor.org/packages/release/BiocViews.html#___Annotages/release/BiocViews.html

such as TxDb.Rnorvegicus.UCSC.rn5.refGene for rat, TxDb.Mmusculus.UCSC.mm10.knownGene for mouse, TxDb.Hsapiens.UCSC.hg19.knownGene for human, TxDb.Dmelanogaster.UCSC.dm3.ensGe

for Drosophila and TxDb.Celegans.UCSC.ce6.ensGene for C.elegans

organism annotation mapping such as org.Hs.egSYMBOL in org.Hs.eg.db pack-

age for human

Value

A data frame and a tab-delimited file offTargetsInPeakRegions.xls, containing all input offtargets with potential gRNA binding sites, mismatch number and positions, alignment to the input gRNA and predicted cleavage score, and whether the offtargets are inside an exon and associated gene name.

Author(s)

Lihua Julie Zhu

See Also

GUIDEseqAnalysis

Examples

```
outputDir = getwd()
offTargets <- offTargetAnalysisOfPeakRegions(gRNA = gRNAs, peaks = peaks,
    format=c("fasta", "bed"),
    peaks.withHeader = TRUE, BSgenomeName = Hsapiens,
    upstream = 20L, downstream = 20L, PAM.size = 3L, gRNA.size = 20L,
    orderOfftargetsBy = "predicted_cleavage_score",
    PAM = "NGG", PAM.pattern = "(NGG|NAG|NGA)$", max.mismatch = 2L,
    outputDir = outputDir,
    allowed.mismatch.PAM = 3, overwrite = TRUE)
annotatedOfftargets <- annotateOffTargets(offTargets,
    txdb = TxDb.Hsapiens.UCSC.hg19.knownGene,
    orgAnn = org.Hs.egSYMBOL)
}</pre>
```

buildFeatureVectorForScoringBulge

Build Feature Vector For Scoring Offtargets with Bulge

Description

Build Feature Vector For Scoring Offtargets with Bulge

Usage

```
buildFeatureVectorForScoringBulge(
   alns,
   gRNA.size = 20,
   canonical.PAM = "NGG",
   subPAM.start = 2,
   subPAM.end = 3,
   insertion.symbol = "^",
   PAM.size = 3,
   PAM.location = "3prime"
)
```

Arguments

```
alns
                 alignments, output from getAlnWithBulge (see the example below)
                 Size of the gRNA, default to 20L
gRNA.size
                 PAM sequence, default to NGG
canonical.PAM
                 start of the subPAM, default to 2L for NGG
subPAM.start
                 End of the subPAM, default to 3L for NGG
subPAM.end
insertion.symbol
                 Symbol used to indicate bulge in DNA Default to ^
PAM.size
                 Size of the PAM, default to 3L for NGG
                 The location of the PAM, default to 3prime
PAM.location
```

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Author(s)

Lihua Julie Zhu

Examples

combineOfftargets

Combine Offtargets

Description

Merge offtargets from different samples

```
combineOfftargets(
  offtarget.folder,
  sample.name,
  remove.common.offtargets = FALSE,
  control.sample.name,
  offtarget.filename = "offTargetsInPeakRegions.xls",
  common.col = c("total.mismatch.bulge", "chromosome", "offTarget_Start",
   "offTarget_End", "offTargetStrand", "offTarget_sequence", "PAM.sequence",
    "guideAlignment2OffTarget", "mismatch.distance2PAM", "n.guide.mismatch",
   "n.PAM.mismatch", "n.DNA.bulge", "n.RNA.bulge", "pos.DNA.bulge", "DNA.bulge",
   "pos.RNA.bulge", "RNA.bulge", "gRNA.name", "gRNAPlusPAM", "predicted_cleavage_score",
    "inExon", "symbol", "entrez_id"),
  exclude.col = "",
  outputFileName,
  comparison.sample1,
  comparison.sample2,
  multiAdjMethod = "BH",
  comparison.score = c("peak_score", "n.distinct.UMIs"),
  overwrite = FALSE
)
```

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Arguments

offtarget.folder

offtarget summary output folders created in GUIDEseqAnalysis function

sample.name Sample names to be used as part of the column names in the final output file

remove.common.offtargets

Default to FALSE If set to TRUE, off-targets common to all samples will be removed.

control.sample.name

The name of the control sample for filtering off-targets present in the control sample

offtarget.filename

Default to offTargetsInPeakRegions.xls, generated in GUIDEseqAnalysis function

common.col common column names used for merge files. Default to c("total.mismatch.bulge", "chromosome",

"mismatch.distance2PAM", "n.guide.mismatch", "n.PAM.mismatch", "n.DNA.bulge", "n.RNA.bulge", "post" "RNA.bulge", "gRNA.name", "gRNAPlusPAM", "predicted_cleavage_score", "in-

"offTarget_Start","offTarget_End", "offTargetStrand", "offTarget_sequence", "PAM.sequence", "guideAlig

Exon", "symbol", "entrez_id")

exclude.col columns to be excluded before merging. Please check offTargetsInPeakRegions.xls to choose the desired columns to exclude

outputFileName The merged offtarget file

comparison.sample1

A vector of sample names to be used for comparison. For example, comparison.sample1 = c("A", "B"), comparison.sample2 = rep("Control", 2) indicates that you are interested in comparing sample A vs Control and B vs Control Please make sure the sample names specified in comparison.sample1 and comparison.sample2 are in the sample name list specified in sample.name

comparison.sample2

A vector of sample names to be used for comparison. For example, comparison.sample1 = c("A", "B"), comparison.sample2 = rep("Control", 2) indicates that you are interested in comparing sample A vs Control and B vs Control

multiAdjMethod A vector of character strings containing the names of the multiple testing procedures for which adjusted p-values are to be computed. This vector should include any of the following: "none", "Bonferroni", "Holm", "Hochberg", "SidakSS", "SidakSD", "BH", "BY", "ABH", and "TSBH". Please type ?multtest::mt.rawp2adjp for details. Default to "BH"

comparison.score

the score to be used for statistical analysis. Two options are available: "peak_score" and "n.distinct.UMIs" n.distinct.UMIs is the number of unique UMIs in the associated peak region without considering the sequence coordinates while peak_score takes into consideration of the sequence coordinates

overwrite Indicates whether to overwrite the existing file specified by outputFileName, default to FALSE.

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Details

Please note that by default, merged file will only contain peaks with offtargets found in the genome in GUIDEseqAnalysis function.

Value

a data frame containing all off-targets from all samples merged by the columns specified in common.col. Sample specific columns have sample.name concatenated to the original column name, e.g., peak score becomes sample1.peak score.

Author(s)

Lihua Julie Zhu

Examples

compareSamples

Compare Samples using Fisher's exact test

Description

Compare Samples using Fisher's exact test

```
compareSamples(
   df,
   col.count1,
   col.count2,
   total1,
   total2,
   multiAdjMethod = "BH",
   comparison.score = c("peak_score", "umi.count")
)
```

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Arguments

a data frame containing the peak score and sequence depth for each sample the score (e.g., peak_score) column used as the numerator for calculating odds ratio. For example, if the tenth column contains the score for sample 1, then set col.count1 = 10

col.count2 the score (e.g., peak_score) column used as the denominator for calculating odds

ratio. For example, if the nineteenth column contains the score for sample 1, then

set col.count2 = 19

total1 the sequence depth for sample 1 total2 the sequence depth for sample 2

multiAdjMethod A vector of character strings containing the names of the multiple testing pro-

cedures for which adjusted p-values are to be computed. This vector should include any of the following: "none", "Bonferroni", "Holm", "Hochberg", "SidakSS", "SidakSD", "BH", "BY", "ABH", and "TSBH". Please type ?multtest::mt.rawp2adjp

for details. Default to "BH"

comparison.score

the score to be used for statistical analysis. Two options are available: "peak_score" and "umi.count" umi.count is the number of unique UMIs in the associated peak region without considering the sequence coordinates while peak_score takes into consideration of the sequence coordinates

Author(s)

Lihua Julie Zhu

createBarcodeFasta

Create barcode as fasta file format for building bowtie1 index

Description

Create barcode as fasta file format for building bowtie1 index to assign reads to each library with different barcodes. The bowtie1 index has been built for the standard GUIDE-seq protocol using the standard p5 and p7 index. It can be downloaded at http://mccb.umassmed.edu/GUIDE-seq/barcode.bowtie1.index.tar.gz

```
createBarcodeFasta(
  p5.index,
  p7.index,
  reverse.p7 = TRUE,
  reverse.p5 = FALSE,
  header = FALSE,
  outputFile = "barcodes.fa"
)
```

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Arguments

p5.index	A text file with one p5 index sequence per line
p7.index	A text file with one p7 index sequence per line
reverse.p7	Indicate whether to reverse p7 index, default to TRUE for standard GUIDE-seq experiments
reverse.p5	Indicate whether to reverse p5 index, default to FALSE for standard GUIDE-seq experiments
header	Indicate whether there is a header in the p5.index and p7.index files. Default to FALSE
outputFile	Give a name to the output file where the generated barcodes are written. This file can be used to build bowtie1 index for binning reads.

Note

Create barcode file to be used to bin the reads sequenced in a pooled lane

Author(s)

Lihua Julie Zhu

Examples

getBestAlnInfo

Parse pairwise alignment

Description

Parse pairwise alignment

```
getBestAlnInfo(
  offtargetSeq,
  pa.f,
  pa.r,
  gRNA.size = 20,
  PAM = "NGG",
```

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```
PAM.size = 3,
  insertion.symbol = "^"
)
```

Arguments

offtargetSeq DNAStringSet object of length 1

pa.f Global-Local PairwiseAlignmentsSingleSubject, results of pairwiseAlignment,

alignment of pattern to subject

pa.r Global-Local PairwiseAlignmentsSingleSubject, results of pairwiseAlignment,

alignment of pattern to reverse subject

gRNA. size size of gRNA, default to 20

PAM PAM sequence, default to NGG

PAM. size PAM size, default to 3

insertion.symbol

symbol for representing bulge in offtarget, default to ^. It can also be set to

lowerCase to use lower case letter to represent insertion

Value

a dataframe with the following columns. offTarget: name of the offtarget peak_score: place holder for storing peak score gRNA.name: place holder for storing gRNA name gRNAPlusPAM: place holder for storing gRNAPlusPAM sequence offTarget_sequence: offTarget sequence with PAM in the right orientation. For PAM in the 3' prime location, offTarget is the sequence on the plus strand otherwise, is the sequence on the reverse strand seq.aligned: the aligned sequence without PAM guideAlignment2OffTarget: string representation of the alignment offTargetStrand: the strand of the offtarget mismatch.distance2PAM: mismatch distance to PAM start n.PAM.mismatch: number of mismatches in PAM n.guide.mismatch: number of mismatches in the gRNA not including PAM PAM.sequence: PAM in the offtarget offTarget_Start: offtarget start offTarget_End: offTarget end chromosome: place holder for storing offtarget chromosome pos.mismatch: mismatch positions with the correct PAM orientation, i.e., indexed form distal to proximal of PAM pos.indel: indel positions starting with deletions in the gRNA followed by those in the offtarget pos.insertion: Insertion positions in the gRNA Insertion positions are counted from distal to proximal of PAM For example, 5 means the 5th position is an insertion in gRNA pos.deletion: Deletion in the gRNA Deletion positions are counted from distal to proximal of PAM For example, 5 means the 5th position is a deletion in gRNA n.insertion: Number of insertions in the RNA. Insertions in gRNA creates bulged DNA base n.deletion: Number of deletions in the RNA. Deletions in gRNA creates bulged DNA base

Author(s)

Lihua Julie Zhu

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getPeaks

Obtain peaks from GUIDE-seq

Description

Obtain strand-specific peaks from GUIDE-seq

Usage

Arguments

GRanges with cleavage sites, output from getUniqueCleavageEvents gr window.size window size to calculate coverage step size to calculate coverage step bg.window.size window size to calculate local background min.reads minimum number of reads to be considered as a peak min.SNratio minimum signal noise ratio, which is the coverage normalized by local back-Maximum p-value to be considered as significant maxP stats Statistical test, default poisson p.adjust.methods

Adjustment method for multiple comparisons, default none

Value

peaks

GRanges with count (peak height), bg (local background), SNratio (signal noise ratio), p-value, and option adjusted p-value

summarized.count

A data frame contains the same information as peaks except that it has all the sites without filtering.

Author(s)

Lihua Julie Zhu

Examples

getUniqueCleavageEvents

Using UMI sequence to obtain the starting sequence library

Description

PCR amplification often leads to biased representation of the starting sequence population. To track the sequence tags present in the initial sequence library, a unique molecular identifier (UMI) is added to the 5 prime of each sequence in the staring library. This function uses the UMI sequence plus the first few sequence from R1 reads to obtain the starting sequence library.

```
getUniqueCleavageEvents(
  alignment.inputfile,
  umi.inputfile,
  alignment.format = c("auto", "bam", "bed"),
  umi.header = FALSE,
  read.ID.col = 1,
  umi.col = 2,
  umi.sep = "\t",
  keep.chrM = FALSE,
  keep.R1only = TRUE,
  keep.R2only = TRUE,
  concordant.strand = TRUE,
  max.paired.distance = 1000,
  min.mapping.quality = 30,
  max.R1.len = 130,
  max.R2.len = 130,
  apply.both.max.len = FALSE,
  same.chromosome = TRUE,
  distance.inter.chrom = -1,
  min.R1.mapped = 20,
```

```
min.R2.mapped = 20,
apply.both.min.mapped = FALSE,
max.duplicate.distance = 0L,
umi.plus.R1start.unique = TRUE,
umi.plus.R2start.unique = TRUE,
min.umi.count = 5L,
max.umi.count = 100000L,
min.read.coverage = 1L,
n.cores.max = 6,
outputDir,
removeDuplicate = TRUE,
ignoreTagmSite = FALSE,
ignoreUMI = FALSE
```

Arguments

alignment.inputfile

The alignment file. Currently supports bed output file with CIGAR information. Suggest run the workflow binReads.sh, which sequentially runs barcode binning, adaptor removal, alignment to genome, alignment quality filtering, and bed file conversion. Please download the workflow function and its helper scripts at http://mccb.umassmed.edu/GUIDE-seq/binReads/

umi.inputfile

A text file containing at least two columns, one is the read identifier and the other is the UMI or UMI plus the first few bases of R1 reads. Suggest use getUMI.sh to generate this file. Please download the script and its helper scripts at http://mccb.umassmed.edu/GUIDE-seq/getUMI/

alignment.format

The format of the alignment input file. Currently only bam and bed file format is supported. BED format will be deprecated soon.

umi.header Indicates whether the umi input file contains a header line or not. Default to

FALSE

read.ID.col The index of the column containing the read identifier in the umi input file,

default to 1

umi.col The index of the column containing the umi or umi plus the first few bases of

sequence from the R1 reads, default to 2

umi . sep column separator in the umi input file, default to tab

keep.chrM Specify whether to include alignment from chrM. Default FALSE

keep.R1only Specify whether to include alignment with only R1 without paired R2. Default

TRUE

keep.R2only Specify whether to include alignment with only R2 without paired R1. Default

TRUE

concordant.strand

Specify whether the R1 and R2 should be aligned to the same strand or opposite strand. Default opposite.strand (TRUE)

max.paired.distance

Specify the maximum distance allowed between paired R1 and R2 reads. Default 1000 bp

min.mapping.quality

Specify min.mapping.quality of acceptable alignments

The maximum retained R1 length to be considered for downstream analysis, default 130 bp. Please note that default of 130 works well when the read length 150 bp. Please also note that retained R1 length is not necessarily equal to the

mapped R1 length

max.R2.1en The maximum retained R2 length to be considered for downstream analysis,

default 130 bp. Please note that default of 130 works well when the read length 150 bp. Please also note that retained R2 length is not necessarily equal to the

mapped R2 length

apply.both.max.len

Specify whether to apply maximum length requirement to both R1 and R2 reads,

default FALSE same.chromosome

Specify whether the paired reads are required to align to the same chromosome,

default TRUE

distance.inter.chrom

Specify the distance value to assign to the paired reads that are aligned to differ-

ent chromosome, default -1

min.R1.mapped
The maximum mapped R1 length to be considered for downstream analysis,

default 30 bp.

min.R2.mapped The maximum mapped R2 length to be considered for downstream analysis,

default 30 bp.

apply.both.min.mapped

Specify whether to apply minimum mapped length requirement to both R1 and

R2 reads, default FALSE

max.duplicate.distance

Specify the maximum distance apart for two reads to be considered as dupli-

cates, default 0. Currently only 0 is supported

umi.plus.R1start.unique

To specify whether two mapped reads are considered as unique if both containing the same UMI and same alignment start for R1 read, default TRUE.

umi.plus.R2start.unique

To specify whether two mapped reads are considered as unique if both contain-

ing the same UMI and same alignment start for R2 read, default TRUE.

min.umi.count To specify the minimum count for a umi to be included in the subsequent analysis. Please adjust it to a higher number for deeply sequenced library and vice

versa.

max.umi.count To specify the maximum count for a umi to be included in the subsequent analysis. Please adjust it to a higher number for deeply sequenced library and vice

versa.

min.read.coverage

To specify the minimum coverage for a read UMI combination to be included in the subsequent analysis. Please note that this is different from min.umi.count which is less stringent.

n.cores.max

Indicating maximum number of cores to use in multi core mode, i.e., parallel processing, default 6. Please set it to 1 to disable multicore processing for small dataset.

outputDir

output Directory to save the figures

removeDuplicate

default to TRUE. Set it to FALSE if PCR duplicates not to be removed for testing purpose.

ignoreTagmSite

default to FALSE. To collapse reads with the same integration site and UMI but with different tagmentation site, set the option to TRUE.

ignoreUMI

default to FALSE. To collapse reads with the same integration and tagmentation site but with different UMIs, set the option to TRUE and retain the UMI that appears most frequently for each combination of integration and tagmentation site. In case of ties, randomly select one UMI.

Value

cleavage.gr

Cleavage sites with one site per UMI as GRanges with metadata column total set to 1 for each range

unique.umi.plus.R2

a data frame containing unique cleavage site from R2 reads mapped to plus strand with the following columns: seqnames (chromosome), start (cleavage/Integration site), strand, UMI (unique molecular identifier), and UMI read duplication level (min.read.coverage can be used to remove UMI-read with very low coverage)

unique.umi.minus.R2

a data frame containing unique cleavage site from R2 reads mapped to minus strand with the same columns as unique.umi.plus.R2

unique.umi.plus.R1

a data frame containing unique cleavage site from R1 reads mapped to minus strand without corresponding R2 reads mapped to the plus strand, with the same columns as unique.umi.plus.R2 $\,$

unique.umi.minus.R1

a data frame containing unique cleavage site from R1 reads mapped to plus strand without corresponding R2 reads mapped to the minus strand, with the same columns as unique.umi.plus.R2

align.umi

a data frame containing all the mapped reads with the following columns. read-Name (read ID), chr.x (chromosome of readSide.x/R1 read), start.x (start of eadSide.x/R1 read), end.x (end of eadSide.x/R1 read), mapping.qual.x (mapping quality of readSide.x/R1 read), strand.x (strand of readSide.x/R1 read), cigar.x (CIGAR of readSide.x/R1 read) , readSide.x (1/R1), chr.y (chromosome of readSide.y/R2 read) start.y (start of readSide.y/R2 read), end.y (end of readSide.y/R2 read), mapping.qual.y (mapping quality of readSide.y/R2 read), strand.y (strand of readSide.y/R2 read), cigar.y (CIGAR of readSide.y/R2 read),

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readSide.y (2/R2) R1.base.kept (retained R1 length), R2.base.kept (retained R2 length), distance (distance between mapped R1 and R2), UMI (unique molecular identifier (umi) or umi with the first few bases of R1 read)

Author(s)

Lihua Julie Zhu

References

Shengdar Q Tsai and J Keith Joung et al. GUIDE-seq enables genome-wide profiling of off-target cleavage by CRISPR-Cas nucleases. Nature Biotechnology 33, 187 to 197 (2015)

See Also

getPeaks

Examples

```
if(interactive())
{
    umiFile <- system.file("extdata", "UMI-HEK293_site4_chr13.txt",
        package = "GUIDEseq")
    alignFile <- system.file("extdata","bowtie2.HEK293_site4_chr13.sort.bam" ,
        package = "GUIDEseq")
    cleavages <- getUniqueCleavageEvents(
        alignment.inputfile = alignFile , umi.inputfile = umiFile,
        n.cores.max = 1)
    names(cleavages)
    #output a summary of duplicate counts for sequencing saturation assessment
    table(cleavages$umi.count.summary$n)
}</pre>
```

getUsedBarcodes

Create barcodes from the p5 and p7 index used for each sequencing lane

Description

Create barcodes from the p5 and p7 index for assigning reads to each barcode

```
getUsedBarcodes(
  p5.index,
  p7.index,
  header = FALSE,
  reverse.p7 = TRUE,
  reverse.p5 = FALSE,
  outputFile
)
```

Arguments

p5.index	A text file with one p5 index sequence per line
p7.index	A text file with one p7 index sequence per line
header	Indicate whether there is a header in the p5.index and p7.index files. Default to $FALSE$
reverse.p7	Indicate whether to reverse p7 index, default to TRUE for standard GUIDE-seq experiments
reverse.p5	Indicate whether to reverse p5 index, default to FALSE for standard GUIDE-seq experiments $$
outputFile	Give a name to the output file where the generated barcodes are written

Value

DNAStringSet

Note

Create barcode file to be used to bin the reads sequenced in a pooled lane

Author(s)

Lihua Julie Zhu

Examples

GUIDEseqAnalysis

Analysis pipeline for GUIDE-seq dataset

Description

A wrapper function that uses the UMI sequence plus the first few bases of each sequence from R1 reads to estimate the starting sequence library, piles up reads with a user defined window and step size, identify the insertion sites (proxy of cleavage sites), merge insertion sites from plus strand and minus strand, followed by off target analysis of extended regions around the identified insertion sites.

```
GUIDEseqAnalysis(
  alignment.inputfile,
  umi.inputfile,
  alignment.format = c("auto", "bam", "bed"),
  umi.header = FALSE,
  read.ID.col = 1L,
  umi.col = 2L,
  umi.sep = "\t",
 BSgenomeName,
  gRNA.file,
 outputDir,
  n.cores.max = 1L,
  keep.chrM = FALSE,
  keep.R1only = TRUE,
  keep.R2only = TRUE,
  concordant.strand = TRUE,
 max.paired.distance = 1000L,
 min.mapping.quality = 30L,
 max.R1.len = 130L
 max.R2.len = 130L,
 min.umi.count = 1L,
 max.umi.count = 100000L,
 min.read.coverage = 1L,
  apply.both.max.len = FALSE,
  same.chromosome = TRUE,
  distance.inter.chrom = -1L,
 min.R1.mapped = 20L,
 min.R2.mapped = 20L,
  apply.both.min.mapped = FALSE,
 max.duplicate.distance = 0L,
  umi.plus.R1start.unique = TRUE,
  umi.plus.R2start.unique = TRUE,
 window.size = 20L,
  step = 20L,
  bg.window.size = 5000L,
 min.reads = 5L,
 min.reads.per.lib = 1L,
 min.peak.score.1strandOnly = 5L,
 min.SNratio = 2,
 maxP = 0.01,
  stats = c("poisson", "nbinom"),
 p.adjust.methods = c("none", "BH", "holm", "hochberg", "hommel", "bonferroni", "BY",
    "fdr"),
  distance.threshold = 40L,
 max.overlap.plusSig.minusSig = 30L,
  plus.strand.start.gt.minus.strand.end = TRUE,
  keepPeaksInBothStrandsOnly = TRUE,
```

```
gRNA.format = "fasta",
  overlap.gRNA.positions = c(17, 18),
  upstream = 25L,
  downstream = 25L,
  PAM.size = 3L,
  gRNA.size = 20L,
  PAM = "NGG",
  PAM.pattern = "NNN$",
 max.mismatch = 6L,
  allowed.mismatch.PAM = 2L,
  overwrite = TRUE,
 weights = c(0, 0, 0.014, 0, 0, 0.395, 0.317, 0, 0.389, 0.079, 0.445, 0.508, 0.613,
    0.851, 0.732, 0.828, 0.615, 0.804, 0.685, 0.583),
 orderOfftargetsBy = c("peak_score", "predicted_cleavage_score", "n.guide.mismatch"),
  descending = TRUE,
  keepTopOfftargetsOnly = TRUE,
  keepTopOfftargetsBy = c("predicted_cleavage_score", "n.mismatch"),
  scoring.method = c("Hsu-Zhang", "CFDscore"),
 subPAM.activity = hash(AA = 0, AC = 0, AG = 0.259259259, AT = 0, CA = 0, CC = 0, CG = 0
  0.107142857, CT = 0, GA = 0.0694444444, GC = 0.0222222222, GG = 1, GT = 0.016129032, TA
    = 0, TC = 0, TG = 0.038961039, TT = 0),
  subPAM.position = c(22, 23),
  PAM.location = "3prime",
 mismatch.activity.file = system.file("extdata",
    "NatureBiot2016SuppTable19DoenchRoot.csv", package = "CRISPRseek"),
  bulge.activity.file = system.file("extdata",
    "NatureBiot2016SuppTable19DoenchRoot.xlsx", package = "GUIDEseq"),
  txdb,
  orgAnn,
 mat,
  includeBulge = FALSE,
 max.n.bulge = 2L,
 min.peak.score.bulge = 60L,
  removeDuplicate = TRUE,
  resume = FALSE,
  ignoreTagmSite = FALSE,
  ignoreUMI = FALSE
)
```

Arguments

alignment.inputfile

The alignment file. Currently supports bam and bed output file with CIGAR information. Suggest run the workflow binReads.sh, which sequentially runs barcode binning, adaptor removal, alignment to genome, alignment quality filtering, and bed file conversion. Please download the workflow function and its helper scripts at http://mccb.umassmed.edu/GUIDE-seq/binReads/

umi.inputfile A text file containing at least two columns, one is the read identifier and the other is the UMI or UMI plus the first few bases of R1 reads. Suggest use

at http://mccb.umassmed.edu/GUIDE-seq/getUMI/

	1		
alignment.format			
	The format of the alignment input file. Default bed file format. Currently only bed file format is supported, which is generated from binReads.sh		
umi.header	Indicates whether the umi input file contains a header line or not. Default to FALSE		
read.ID.col	The index of the column containing the read identifier in the umi input file, default to 1		
umi.col	The index of the column containing the umi or umi plus the first few bases of sequence from the R1 reads, default to 2		
umi.sep	column separator in the umi input file, default to tab		
BSgenomeName	BSgenome object. Please refer to available.genomes in BSgenome package. For example, BSgenome.Hsapiens.UCSC.hg19 for hg19, BSgenome.Mmusculus.UCSC.mm10 for mm10, BSgenome.Celegans.UCSC.ce6 for ce6, BSgenome.Rnorvegicus.UCSC.rn5 for rn5, BSgenome.Drerio.UCSC.danRer7 for Zv9, and BSgenome.Dmelanogaster.UCSC.dm3 for dm3		
gRNA.file	gRNA input file path or a DNAStringSet object that contains the target sequence (gRNA plus PAM)		
outputDir	the directory where the off target analysis and reports will be written to		
n.cores.max	Indicating maximum number of cores to use in multi core mode, i.e., parallel processing, default 1 to disable multicore processing for small dataset.		

getUMI.sh to generate this file. Please download the script and its helper scripts

TRUE concordant.strand

keep.chrM

keep.R1only

keep.R2only

Specify whether the R1 and R2 should be aligned to the same strand or opposite strand. Default opposite.strand (TRUE)

Specify whether to include alignment with only R1 without paired R2. Default

Specify whether to include alignment with only R2 without paired R1. Default

Specify whether to include alignment from chrM. Default FALSE

max.paired.distance

Specify the maximum distance allowed between paired R1 and R2 reads. Default 1000 bp

min.mapping.quality

Specify min.mapping.quality of acceptable alignments

max.R1.len The maximum retained R1 length to be considered for downstream analysis, default 130 bp. Please note that default of 130 works well when the read length 150 bp. Please also note that retained R1 length is not necessarily equal to the mapped R1 length

max.R2.len The maximum retained R2 length to be considered for downstream analysis, default 130 bp. Please note that default of 130 works well when the read length 150 bp. Please also note that retained R2 length is not necessarily equal to the mapped R2 length

min.umi.count

To specify the minimum total count for a umi at the genome level to be included in the subsequent analysis. For example, with min.umi.count set to 2, if a umi only has 1 read in the entire genome, then that umi will be excluded for the subsequent analysis. Please adjust it to a higher number for deeply sequenced library and vice versa.

max.umi.count

To specify the maximum count for a umi to be included in the subsequent analysis. Please adjust it to a higher number for deeply sequenced library and vice versa.

min.read.coverage

To specify the minimum coverage for a read UMI combination to be included in the subsequent analysis. Please note that this is different from min.umi.count which is less stringent.

apply.both.max.len

Specify whether to apply maximum length requirement to both R1 and R2 reads, default FALSE

same.chromosome

Specify whether the paired reads are required to align to the same chromosome, default TRUE

distance.inter.chrom

Specify the distance value to assign to the paired reads that are aligned to different chromosome, default -1

min.R1.mapped The minimum mapped R1 length to be considered for downstream analysis, default 30 bp.

min.R2.mapped The minimum mapped R2 length to be considered for downstream analysis, default 30 bp.

apply.both.min.mapped

Specify whether to apply minimum mapped length requirement to both R1 and R2 reads, default FALSE

max.duplicate.distance

Specify the maximum distance apart for two reads to be considered as duplicates, default 0. Currently only 0 is supported

umi.plus.R1start.unique

To specify whether two mapped reads are considered as unique if both containing the same UMI and same alignment start for R1 read, default TRUE.

umi.plus.R2start.unique

To specify whether two mapped reads are considered as unique if both containing the same UMI and same alignment start for R2 read, default TRUE.

window.size window size to calculate coverage

step step size to calculate coverage

bg.window.size window size to calculate local background

min.reads minimum number of reads to be considered as a peak

min.reads.per.lib

minimum number of reads in each library (usually two libraries) to be considered as a peak

min.peak.score.1strandOnly

Specify the minimum number of reads for a one-strand only peak to be included in the output. Applicable when set keepPeaksInBothStrandsOnly to FALSE and

there is only one library per sample

min. SNratio Specify the minimum signal noise ratio to be called as peaks, which is the cov-

erage normalized by local background.

maxP Specify the maximum p-value to be considered as significant

stats Statistical test, currently only poisson is implemented

p.adjust.methods

Adjustment method for multiple comparisons, default none

distance.threshold

Specify the maximum gap allowed between the plus strand and the negative strand peak, default 40. Suggest set it to twice of window.size used for peak ...

calling.

max.overlap.plusSig.minusSig

Specify the cushion distance to allow sequence error and inprecise integration Default to 30 to allow at most 10 (30-window.size 20) bp (half window) of minus-strand peaks on the right side of plus-strand peaks. Only applicable if plus.strand.start.gt.minus.strand.end is set to TRUE.

plus.strand.start.gt.minus.strand.end

Specify whether plus strand peak start greater than the paired negative strand peak end. Default to TRUE

keepPeaksInBothStrandsOnly

Indicate whether only keep peaks present in both strands as specified by plus.strand.start.gt.minus.strand.e max.overlap.plusSig.minusSig and distance.threshold.

gRNA. format Format of the gRNA input file. Currently, fasta is supported

overlap.gRNA.positions

The required overlap positions of gRNA and restriction enzyme cut site, default

17 and 18 for SpCas9.

upstream upstream offset from the peak start to search for off targets, default 25 suggest

set it to window size

downstream downstream offset from the peak end to search for off targets, default 25 suggest

set it to window size

PAM. size PAM length, default 3

gRNA. size The size of the gRNA, default 20

PAM sequence after the gRNA, default NGG

PAM. pattern Regular expression of protospacer-adjacent motif (PAM), default NNN\$. Alter-

natively set it to (NAGINGGINGA)\$ for off target search

max.mismatch Maximum mismatch to the gRNA (not including mismatch to the PAM) allowed

in off target search, default 6

allowed.mismatch.PAM

Maximum number of mismatches allowed for the PAM sequence plus the number of degenerate sequence in the PAM sequence, default to 2 for NGG PAM

overwrite overwrite the existing files in the output directory or not, default FALSE

weights

a numeric vector size of gRNA length, default c(0, 0, 0.014, 0, 0, 0.395, 0.317, 0, 0.389, 0.079, 0.445, 0.508, 0.613, 0.851, 0.732, 0.828, 0.615, 0.804, 0.685, 0.583) for SPcas9 system, which is used in Hsu et al., 2013 cited in the reference section. Please make sure that the number of elements in this vector is the same as the gRNA.size, e.g., pad 0s at the beginning of the vector.

orderOfftargetsBy

Criteria to order the offtargets, which works together with the descending pa-

descending Indicate the output order of the offtargets, i.e., in the descending or ascending order.

keepTopOfftargetsOnly

Output all offtargets or the top offtarget using the keepOfftargetsBy criteria, default to the top offtarget

keepTopOfftargetsBy

Output the top offtarget for each called peak using the keepTopOfftargetsBy criteria, If set to predicted_cleavage_score, then the offtargets with the highest predicted cleavage score will be retained If set to n.mismatch, then the offtarget with the lowest number of mismatch to the target sequence will be retained

scoring.method Indicates which method to use for offtarget cleavage rate estimation, currently two methods are supported, Hsu-Zhang and CFDscore

subPAM.activity

Applicable only when scoring.method is set to CFDscore A hash to represent the cleavage rate for each alternative sub PAM sequence relative to preferred PAM sequence

subPAM.position

Applicable only when scoring method is set to CFD score The start and end positions of the sub PAM. Default to 22 and 23 for SP with 20bp gRNA and NGG as preferred PAM

PAM.location

PAM location relative to gRNA. For example, default to 3prime for spCas9 PAM. Please set to 5prime for cpf1 PAM since it's PAM is located on the 5 prime end

mismatch.activity.file

Applicable only when scoring method is set to CFD score A comma separated (csv) file containing the cleavage rates for all possible types of single nucleotide mismatche at each position of the gRNA. By default, use the supplemental Table 19 from Doench et al., Nature Biotechnology 2016

bulge.activity.file

Used for predicting indel effect on offtarget cleavage score. An excel file with the second sheet for deletion activity and the third sheet for Insertion. By default, use the supplemental Table 19 from Doench et al., Nature Biotechnology 2016

TxDb object, for creating and using TxDb object, please refer to GenomicFea-

tures package. For a list of existing TxDb object, please search for annotation package starting with Txdb at http://www.bioconductor.org/packages/release/BiocViews.html#___ such as TxDb.Rnorvegicus.UCSC.rn5.refGene for rat, TxDb.Mmusculus.UCSC.mm10.knownGene for mouse, TxDb.Hsapiens.UCSC.hg19.knownGene for human, TxDb.Dmelanogaster.UCSC.dm3.ensGe for Drosophila and TxDb.Celegans.UCSC.ce6.ensGene for C.elegans

txdb

orgAnn organism annotation mapping such as org.Hs.egSYMBOL in org.Hs.eg.db pack-

age for human

mat nucleotide substitution matrix. Function nucleotideSubstitutionMatrix can be

used for creating customized nucleotide substitution matrix. By default, match = 1, mismatch = -1, and baseOnly = TRUE Only applicable with includeBulge

set to TRUE

includeBulge indicates whether including offtargets with indels default to FALSE

max.n.bulge offtargets with at most this number of indels to be included in the offtarget list.

Only applicable with includeBulge set to TRUE

min.peak.score.bulge

default to 60. Set it to a higher number to speed up the alignment with bulges. Any peaks with peak.score less than min.peak.score.bulge will not be included in the offtarget analysis with bulges. However, all peaks will be included in the

offtarget analysis with mismatches.

removeDuplicate

default to TRUE. Set it to FALSE if PCR duplicates not to be removed for testing

purpose

resume default to FALSE to restart the analysis. set it TRUE to resume an analysis.

ignoreTagmSite default to FALSE. To collapse reads with the same integration site and UMI but

with different tagmentation site, set the option to TRUE.

ignoreUMI default to FALSE. To collapse reads with the same integration and tagmentation

site but with different UMIs, set the option to TRUE and retain the UMI that appears most frequently for each combination of integration and tagmentation

site. In case of ties, randomly select one UMI.

Value

offTargets a data frame, containing all input peaks with potential gRNA binding sites, mis-

match number and positions, alignment to the input gRNA and predicted cleav-

age score.

merged.peaks merged peaks as GRanges with count (peak height), bg (local background),

SNratio (signal noise ratio), p-value, and option adjusted p-value

peaks GRanges with count (peak height), bg (local background), SNratio (signal noise

ratio), p-value, and option adjusted p-value

uniqueCleavages

Cleavage sites with one site per UMI as GRanges with metadata column total

set to 1 for each range

read. summary One table per input mapping file that contains the number of reads for each

chromosome location

sequence.depth sequence depth in the input alignment files

Author(s)

Lihua Julie Zhu

References

Lihua Julie Zhu, Michael Lawrence, Ankit Gupta, Herve Pages, Alper Ku- cukural, Manuel Garber and Scot A. Wolfe. GUIDEseq: a bioconductor package to analyze GUIDE-Seq datasets for CRISPR-Cas nucleases. BMC Genomics. 2017. 18:379

See Also

getPeaks

Examples

```
if(interactive())
   {
        library("BSgenome.Hsapiens.UCSC.hg19")
        umiFile <- system.file("extdata", "UMI-HEK293_site4_chr13.txt",</pre>
           package = "GUIDEseq")
        alignFile <- system.file("extdata","bowtie2.HEK293_site4_chr13.sort.bam" ,</pre>
            package = "GUIDEseq")
        gRNA.file <- system.file("extdata", "gRNA.fa", package = "GUIDEseq")</pre>
        guideSeqRes <- GUIDEseqAnalysis(</pre>
            alignment.inputfile = alignFile,
            umi.inputfile = umiFile, gRNA.file = gRNA.file,
            orderOfftargetsBy = "peak_score",
            descending = TRUE,
            keepTopOfftargetsBy = "predicted_cleavage_score",
            scoring.method = "CFDscore",
            BSgenomeName = Hsapiens, min.reads = 80, n.cores.max = 1)
        guideSeqRes$offTargets
        names(guideSeqRes)
  }
```

mergePlusMinusPeaks

Merge peaks from plus strand and minus strand

Description

Merge peaks from plus strand and minus strand with required orientation and within certain distance apart

```
mergePlusMinusPeaks(
  peaks.gr,
  peak.height.mcol = "count",
  bg.height.mcol = "bg",
  distance.threshold = 40L,
  max.overlap.plusSig.minusSig = 30L,
```

mergePlusMinusPeaks 27

```
plus.strand.start.gt.minus.strand.end = TRUE,
  output.bedfile
)
```

Arguments

peaks.gr

Specify the peaks as GRanges object, which should contain peaks from both plus and minus strand. In addition, it should contain peak height metadata column to store peak height and optionally background height.

peak.height.mcol

Specify the metadata column containing the peak height, default to count

bg.height.mcol Specify the metadata column containing the background height, default to bg distance.threshold

Specify the maximum gap allowed between the plus stranded and the negative stranded peak, default 40. Suggest set it to twice of window.size used for peak calling.

max.overlap.plusSig.minusSig

Specify the cushion distance to allow sequence error and inprecise integration Default to 30 to allow at most 10 (30-window.size 20) bp (half window) of minus-strand peaks on the right side of plus-strand peaks. Only applicable if plus.strand.start.gt.minus.strand.end is set to TRUE.

plus.strand.start.gt.minus.strand.end

Specify whether plus strand peak start greater than the paired negative strand peak end. Default to TRUE

output.bedfile Specify the bed output file name, which is used for off target analysis subsequently.

Value

output a list and a bed file containing the merged peaks a data frame of the bed format

```
mergedPeaks.gr merged peaks as GRanges
mergedPeaks.bed
merged peaks in bed format
```

Author(s)

Lihua Julie Zhu

References

Zhu L.J. et al. (2010) ChIPpeakAnno: a Bioconductor package to annotate ChIP-seq and ChIP-chip data. BMC Bioinformatics 2010, 11:237doi:10.1186/1471-2105-11-237. Zhu L.J. (2013) Integrative analysis of ChIP-chip and ChIP-seq dataset. Methods Mol Biol. 2013;1067:105-24. doi: 10.1007/978-1-62703-607-8_8.

Examples

```
if (interactive())
{
    data(peaks.gr)
    mergedPeaks <- mergePlusMinusPeaks(peaks.gr = peaks.gr,
        output.bedfile = "mergedPeaks.bed")
    mergedPeaks$mergedPeaks.gr
    head(mergedPeaks$mergedPeaks.bed)
}</pre>
```

off Target Analysis Of Peak Regions

Offtarget Analysis of GUIDE-seq peaks

Description

Finding offtargets around peaks from GUIDE-seq or around any given genomic regions

```
offTargetAnalysisOfPeakRegions(
  gRNA,
  peaks,
  format = c("fasta", "bed"),
  peaks.withHeader = FALSE,
 BSgenomeName,
  overlap.gRNA.positions = c(17, 18),
  upstream = 25L,
  downstream = 25L.
 PAM.size = 3L,
  gRNA.size = 20L,
 PAM = "NGG",
 PAM.pattern = "NNN$",
 max.mismatch = 6L,
 outputDir,
  allowed.mismatch.PAM = 2L,
 overwrite = TRUE,
 weights = c(0, 0, 0.014, 0, 0, 0.395, 0.317, 0, 0.389, 0.079, 0.445, 0.508, 0.613,
    0.851, 0.732, 0.828, 0.615, 0.804, 0.685, 0.583),
 orderOfftargetsBy = c("predicted_cleavage_score", "n.mismatch"),
 descending = TRUE,
 keepTopOfftargetsOnly = TRUE,
  scoring.method = c("Hsu-Zhang", "CFDscore"),
 subPAM.activity = hash(AA = 0, AC = 0, AG = 0.259259259, AT = 0, CA = 0, CC = 0, CG = 0
  0.107142857, CT = 0, GA = 0.069444444, GC = 0.022222222, GG = 1, GT = 0.016129032, TA
```

```
= 0, TC = 0, TG = 0.038961039, TT = 0),
subPAM.position = c(22, 23),
PAM.location = "3prime",
mismatch.activity.file = system.file("extdata",
    "NatureBiot2016SuppTable19DoenchRoot.csv", package = "CRISPRseek"),
n.cores.max = 1
)
```

Arguments

gRNA input file path or a DNAStringSet object that contains gRNA plus PAM

sequences used for genome editing

peaks peak input file path or a GenomicRanges object that contains genomic regions

to be searched for potential offtargets

format Format of the gRNA and peak input file. Currently, fasta and bed are supported

for gRNA and peak input file respectively

peaks.withHeader

Indicate whether the peak input file contains header, default FALSE

BSgenomeName BSgenome object. Please refer to available genomes in BSgenome package. For

example, BSgenome.Hsapiens.UCSC.hg19 for hg19, BSgenome.Mmusculus.UCSC.mm10 for mm10, BSgenome.Celegans.UCSC.ce6 for ce6, BSgenome.Rnorvegicus.UCSC.rn5 for rn5, BSgenome.Drerio.UCSC.danRer7 for Zv9, and BSgenome.Dmelanogaster.UCSC.dm3

for dm3

overlap.gRNA.positions

The required overlap positions of gRNA and restriction enzyme cut site, default

17 and 18 for SpCas9.

upstream upstream offset from the peak start to search for off targets, default 20

downstream downstream offset from the peak end to search for off targets, default 20

PAM. size PAM length, default 3

gRNA. size The size of the gRNA, default 20

PAM PAM sequence after the gRNA, default NGG

PAM. pattern Regular expression of protospacer-adjacent motif (PAM), default to any NNN\$.

Set it to (NAGINGGINGA)\$ if only outputs offtargets with NAG, NGA or NGG

PAM

max.mismatch Maximum mismatch allowed in off target search, default 6

outputDir the directory where the off target analysis and reports will be written to

allowed.mismatch.PAM

Number of degenerative bases in the PAM.pattern sequence, default to 2

overwrite overwrite the existing files in the output directory or not, default FALSE

weights a numeric vector size of gRNA length, default c(0, 0, 0.014, 0, 0, 0.395, 0.317,

0, 0.389, 0.079, 0.445, 0.508, 0.613, 0.851, 0.732, 0.828, 0.615, 0.804, 0.685, 0.583) for SPcas9 system, which is used in Hsu et al., 2013 cited in the reference section. Please make sure that the number of elements in this vector is the same

as the gRNA.size, e.g., pad 0s at the beginning of the vector.

orderOfftargetsBy

criteria to order the offtargets by and the top one will be kept if keepTopOfftargetsOnly is set to TRUE. If set to predicted_cleavage_score (descending order), the offtarget with the highest predicted cleavage score for each peak will be kept. If set to n.mismatch (ascending order), the offtarget with the smallest number of mismatch to the target sequence for each peak will be kept.

descending

No longer used. In the descending or ascending order. Default to order by predicted cleavage score in descending order and number of mismatch in ascending order When altering orderOfftargetsBy order, please also modify descending accordingly

keepTopOfftargetsOnly

Output all offtargets or the top offtarget per peak using the orderOfftargetsBy criteria, default to the top offtarget

scoring.method Indicates which method to use for offtarget cleavage rate estimation, currently two methods are supported, Hsu-Zhang and CFDscore

subPAM.activity

Applicable only when scoring.method is set to CFDscore A hash to represent the cleavage rate for each alternative sub PAM sequence relative to preferred PAM sequence

subPAM.position

Applicable only when scoring.method is set to CFDscore The start and end positions of the sub PAM. Default to 22 and 23 for SP with 20bp gRNA and NGG as preferred PAM

PAM.location

PAM location relative to gRNA. For example, default to 3prime for spCas9 PAM. Please set to 5prime for cpf1 PAM since it's PAM is located on the 5 prime end

mismatch.activity.file

Applicable only when scoring.method is set to CFDscore A comma separated (csv) file containing the cleavage rates for all possible types of single nucleotide mismatch at each position of the gRNA. By default, using the supplemental Table 19 from Doench et al., Nature Biotechnology 2016

n.cores.max

Indicating maximum number of cores to use in multi core mode, i.e., parallel processing, default 1 to disable multicore processing for small dataset.

Value

a tab-delimited file offTargetsInPeakRegions.tsv, containing all input peaks with potential gRNA binding sites, mismatch number and positions, alignment to the input gRNA and predicted cleavage score.

Author(s)

Lihua Julie Zhu

References

Patrick D Hsu, David A Scott, Joshua A Weinstein, F Ann Ran, Silvana Konermann, Vineeta Agarwala, Yinqing Li, Eli J Fine, Xuebing Wu, Ophir Shalem, Thomas J Cradick, Luciano A Marraffini,

Gang Bao & Feng Zhang (2013) DNA targeting specificity of rNA-guided Cas9 nucleases. Nature Biotechnology 31:827-834 Lihua Julie Zhu, Benjamin R. Holmes, Neil Aronin and Michael Brodsky. CRISPRseek: a Bioconductor package to identify target-specific guide RNAs for CRISPR-Cas9 genome-editing systems. Plos One Sept 23rd 2014 Lihua Julie Zhu (2015). Overview of guide RNA design tools for CRISPR-Cas9 genome editing technology. Frontiers in Biology August 2015, Volume 10, Issue 4, pp 289-296

See Also

GUIDEseq

Examples

```
#### the following example is also part of annotateOffTargets.Rd
if (interactive())
   library("BSgenome.Hsapiens.UCSC.hg19")
   library(GUIDEseq)
   peaks <- system.file("extdata", "T2plus1000ffTargets.bed",</pre>
        package = "CRISPRseek")
    gRNAs <- system.file("extdata", "T2.fa",
        package = "CRISPRseek")
   outputDir = getwd()
   offTargets <- offTargetAnalysisOfPeakRegions(gRNA = gRNAs, peaks = peaks,
        format=c("fasta", "bed"),
        peaks.withHeader = TRUE, BSgenomeName = Hsapiens,
        upstream = 25L, downstream = 25L, PAM.size = 3L, gRNA.size = 20L,
        orderOfftargetsBy = "predicted_cleavage_score",
        PAM = "NGG", PAM.pattern = "(NGG|NAG|NGA)$", max.mismatch = 2L,
        outputDir = outputDir,
        allowed.mismatch.PAM = 3, overwrite = TRUE
}
```

offTargetAnalysisWithBulge

offTarget Analysis With Bulges Allowed Finding offtargets around peaks from GUIDE-seq or around any given genomic regions with bulges allowed in gRNA or the DNA sequence of offTargets when aligning gRNA and DNA sequences.

Description

offTarget Analysis With Bulges Allowed Finding offtargets around peaks from GUIDE-seq or around any given genomic regions with bulges allowed in gRNA or the DNA sequence of offTargets when aligning gRNA and DNA sequences.

Usage

```
offTargetAnalysisWithBulge(
  gRNA,
  gRNA.name,
  peaks,
 BSgenomeName,
 mat,
  peaks.withHeader = FALSE,
 peaks.format = "bed",
 gapOpening = 1L,
  gapExtension = 3L,
 max.DNA.bulge = 2L,
 max.mismatch = 10L,
  allowed.mismatch.PAM = 2L,
  upstream = 20L,
  downstream = 20L,
 PAM.size = 3L,
  gRNA.size = 20L,
 PAM = "NGG",
 PAM.pattern = "NNN$",
 PAM.location = "3prime",
 mismatch.activity.file = system.file("extdata",
    "NatureBiot2016SuppTable19DoenchRoot.xlsx", package = "GUIDEseq")
)
```

Arguments

gRNA a character string containing the gRNA sequence without PAM

gRNA.name name of the gRNA

peaks peak input file path or a GenomicRanges object that contains genomic regions

to be searched for potential offtargets

BSgenomeName BSgenome object. Please refer to available.genomes in BSgenome package. For

example, BSgenome.Hsapiens.UCSC.hg19 for hg19, BSgenome.Mmusculus.UCSC.mm10 for mm10, BSgenome.Celegans.UCSC.ce6 for ce6, BSgenome.Rnorvegicus.UCSC.rn5 for rn5, BSgenome.Drerio.UCSC.danRer7 for Zv9, and BSgenome.Dmelanogaster.UCSC.dm3

for dm3

mat nucleotideSubstitutionMatrix, which can be created using nucleotideSubstitu-

tionMatrix.

peaks.withHeader

Indicate whether the peak input file contains header, default FALSE

peaks.format format of the peak file, default to bed file format. Currently, only bed format is

supported

gapOpening Gap opening penalty, default to 1L gapExtension Gap extension penalty, default to 3L

max.DNA.bulge Total number of bulges allowed, including bulges in DNA and gRNA, default to

2L

max.mismatch Maximum mismatch allowed in off target search, default 10L

allowed.mismatch.PAM

Number of degenerative bases in the PAM.pattern sequence, default to 2L

upstream upstream offset from the peak start to search for off targets, default 20

downstream downstream offset from the peak end to search for off targets, default 20

PAM. size PAM length, default 3

gRNA. size The size of the gRNA, default 20

PAM PAM sequence after the gRNA, default NGG

PAM. pattern Regular expression of protospacer-adjacent motif (PAM), default to any NNN\$.

Currently, only support NNN\$

PAM.location PAM location relative to gRNA. For example, default to 3prime for spCas9

PAM. Please set to 5prime for cpf1 PAM since it's PAM is located on the 5

prime end

mismatch.activity.file

Applicable only when scoring method is set to CFDscore A comma separated (csv) file containing the cleavage rates for all possible types of single nucleotide mismatch at each position of the gRNA. By default, using the supplemental

Table 19 from Doench et al., Nature Biotechnology 2016

Author(s)

Lihua Julie Zhu

Examples

```
if (interactive()) {
    library(GUIDEseq)
    peaks <- system.file("extdata","1450-chr14-chr2-bulge-test.bed", package = "GUIDEseq")
    mismatch.activity.file <-system.file("extdata", "NatureBiot2016SuppTable19DoenchRoot.xlsx",
        package = "GUIDEseq")

gRNA <- "TGCTTGGTCGGCACTGATAG"
gRNA.name <- "Test1450"
library(BSgenome.Hsapiens.UCSC.hg38)

temp <- offTargetAnalysisWithBulge(gRNA = gRNA, gRNA.name = gRNA.name,
        peaks = peaks, BSgenomeName = Hsapiens,
        mismatch.activity.file = mismatch.activity.file)
}</pre>
```

PEtagAnalysis

peaks.gr

example cleavage sites

Description

An example data set containing cleavage sites (peaks) from getPeaks

Format

GRanges with count (peak height), bg (local background), SNratio (signal noise ratio), p-value, and option adjusted p-value

Value

peaks.gr

GRanges with count (peak height), bg (local background), SNratio (signal noise ratio), p-value, and option adjusted p-value

Source

http://trace.ncbi.nlm.nih.gov/Traces/sra/?run=SRR1695644

Examples

```
data(peaks.gr)
names(peaks.gr)
peaks.gr
```

PEtagAnalysis

Analysis pipeline for PEtag-seq dataset

Description

A wrapper function that uses the UMI sequence plus the first few bases of each sequence from R1 reads to estimate the starting sequence library, piles up reads with a user defined window and step size, identify the insertion sites (proxy of cleavage sites), merge insertion sites from plus strand and minus strand, followed by off target analysis of extended regions around the identified insertion sites. Detailed information on additional parameters can be found in GUIDEseqAnalysis manual with help(GUIDEseqAnalysis).

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Usage

```
PEtagAnalysis(
  alignment.inputfile,
  umi.inputfile,
 BSgenomeName,
  gRNA.file,
  outputDir,
  keepPeaksInBothStrandsOnly = FALSE,
  txdb,
  orgAnn,
 PAM.size = 3L,
  gRNA.size = 20L,
  overlap.gRNA.positions = c(17, 18),
 PAM.location = "3prime",
 PBS.len = 10L,
 HA.len = 7L,
)
```

Arguments

alignment.inputfile

The alignment file. Currently supports bam and bed output file with CIGAR information. Suggest run the workflow binReads.sh, which sequentially runs barcode binning, adaptor removal, alignment to genome, alignment quality filtering, and bed file conversion. Please download the workflow function and its

helper scripts at http://mccb.umassmed.edu/GUIDE-seq/binReads/

umi.inputfile A text file containing at least two columns, one is the read identifier and the

other is the UMI or UMI plus the first few bases of R1 reads. Suggest use getUMI.sh to generate this file. Please download the script and its helper scripts

at http://mccb.umassmed.edu/GUIDE-seq/getUMI/

BSgenomeName BSgenome object. Please refer to available genomes in BSgenome package. For

> example, BSgenome. Hsapiens. UCSC.hg19 for hg19, BSgenome. Mmusculus. UCSC.mm10 for mm10, BSgenome. Celegans. UCSC.ce6 for ce6, BSgenome. Rnorvegicus. UCSC.rn5 for rn5, BSgenome.Drerio.UCSC.danRer7 for Zv9, and BSgenome.Dmelanogaster.UCSC.dm3

for dm3

gRNA.file gRNA input file path or a DNAStringSet object that contains the target sequence

(gRNA plus PAM)

outputDir the directory where the off target analysis and reports will be written to

keepPeaksInBothStrandsOnly

Indicate whether only keep peaks present in both strands as specified by plus.strand.start.gt.minus.strand.e max.overlap.plusSig.minusSig and distance.threshold. Please see GUIDEseq-

Analysis for details of additional parameters. Default to FALSE for any in vitro system, which needs to be set to TRUE for any in vivo system.

TxDb object, for creating and using TxDb object, please refer to GenomicFeatures package. For a list of existing TxDb object, please search for annotation

package starting with Txdb at http://www.bioconductor.org/packages/release/BiocViews.html#___Annota

txdb

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such as TxDb.Rnorvegicus.UCSC.rn5.refGene for rat, TxDb.Mmusculus.UCSC.mm10.knownGene for mouse, TxDb.Hsapiens.UCSC.hg19.knownGene for human, TxDb.Dmelanogaster.UCSC.dm3.ensGe

for Drosophila and TxDb.Celegans.UCSC.ce6.ensGene for C.elegans

organism annotation mapping such as org.Hs.egSYMBOL in org.Hs.eg.db pack-

age for human

PAM. size PAM length, default 3

gRNA. size The size of the gRNA, default 20

overlap.gRNA.positions

The required overlap positions of gRNA and restriction enzyme cut site, default

17 and 18 for SpCas9.

PAM.location PAM location relative to gRNA. For example, default to 3prime for spCas9

PAM. Please set to 5prime for cpf1 PAM since it's PAM is located on the 5

prime end

PBS.len Primer binding sequence length, default to 10.

HA.len Homology arm sequence length, default to 7.

... Any parameters in GUIDEseqAnalysis can be used for this function. Please type

help(GUIDEseqAnalysis for detailed information.

Value

offTargets a data frame, containing all input peaks with potential gRNA binding sites, mis-

match number and positions, alignment to the input gRNA, predicted cleavage score, PBS (primer binding sequence), and HAseq (homology arm sequence).

merged peaks as GRanges with count (peak height), bg (local background),

SNratio (signal noise ratio), p-value, and option adjusted p-value

peaks GRanges with count (peak height), bg (local background), SNratio (signal noise

ratio), p-value, and option adjusted p-value

uniqueCleavages

Cleavage sites with one site per UMI as GRanges with metadata column total

set to 1 for each range

read.summary One table per input mapping file that contains the number of reads for each

chromosome location

Author(s)

Lihua Julie Zhu

References

Lihua Julie Zhu, Michael Lawrence, Ankit Gupta, Herve Pages, Alper Ku- cukural, Manuel Garber and Scot A. Wolfe. GUIDEseq: a bioconductor package to analyze GUIDE-Seq datasets for CRISPR-Cas nucleases. BMC Genomics. 2017. 18:379

See Also

GUIDEseqAnalysis

plotAlignedOfftargets 37

Examples

```
if(!interactive())
   {
        library("BSgenome.Hsapiens.UCSC.hg19")
        library(TxDb.Hsapiens.UCSC.hg19.knownGene)
        library(org.Hs.eg.db)
        umiFile <- system.file("extdata", "UMI-HEK293_site4_chr13.txt",</pre>
           package = "GUIDEseq")
        alignFile <- system.file("extdata","bowtie2.HEK293_site4_chr13.sort.bam" ,</pre>
            package = "GUIDEseq")
        gRNA.file <- system.file("extdata", "gRNA.fa", package = "GUIDEseq")</pre>
        PET.res <- PEtagAnalysis(</pre>
            alignment.inputfile = alignFile,
            umi.inputfile = umiFile,
            gRNA.file = gRNA.file,
            orderOfftargetsBy = "peak_score",
            descending = TRUE,
            keepTopOfftargetsBy = "predicted_cleavage_score",
            scoring.method = "CFDscore",
            BSgenomeName = Hsapiens,
            txdb = TxDb.Hsapiens.UCSC.hg19.knownGene,
            orgAnn = org.Hs.egSYMBOL,
            outputDir = "PEtagTestResults",
            min.reads = 80, n.cores.max = 1,
            keepPeaksInBothStrandsOnly = FALSE,
            PBS.len = 10L,
            HA.len = 7L
            )
        PET.res$offTargets
        names(PET.res)
  }
```

plotAlignedOfftargets Plot offtargets aligned to the target sequence

Description

Plot offtargets aligned to the target sequence

Usage

```
plotAlignedOfftargets(
  offTargetFile,
  sep = "\t",
  header = TRUE,
  gRNA.size = 20L,
  input.DNA.bulge.symbol = "^",
  input.RNA.bulge.symbol = "-",
```

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```
input.match.symbol = ".",
  plot.DNA.bulge.symbol = "DNA.bulge",
 plot.RNA.bulge.symbol = "-",
 plot.match.symbol = ".",
  color.DNA.bulge = "red",
  size.symbol = 3,
 color.values = c(A = "#B5D33D", T = "#AE9CD6", C = "#6CA2EA", G = "#FED23F", `-` =
    "gray", . = "white"),
 PAM = "GGG",
 body.tile.height = 2.5,
  header.tile.height = 3.6,
 hline.offset = 3.8,
  plot.top.n,
  insertion.score.column = c("n.distinct.UMIs", "peak_score"),
  insertion.score.column.prefix,
 width. IR = 2.5,
 width.RIR = 2.5,
  family = "sans",
 hjust = "middle",
  viust = 0.5
)
```

Arguments

offTargetFile The path of the file offTargetsInPeakRegions.xls that stores the offtargets to be

plotted. This file is the output file from the function GUIDEseqAnalysis.

sep Field delimiter for the file specified as offTargetFile, default to tab dilimiter

header Indicates whether there is header in the file specified as offTargetFile, default to

TRUE

gRNA. size Size of the gRNA, default to 20 for SpCas9 system

input.DNA.bulge.symbol

The symbol used to represent DNA bulges in the file specified as offTargetFile, default to "^"

input.RNA.bulge.symbol

The symbol used to represent RNA bulges in the file specified as offTargetFile, default to "-"

input.match.symbol

The symbol used to represent matched bases in the file specified as offTargetFile, default to "."

plot.DNA.bulge.symbol

The symbol used to represent DNA bulges in the figure to be generated, default to DNA.bulge, i.e., the nucleotide in the DNA bulge. Alternatively, you can specify a symbol to represent all DNA bulges such as "I".

plot.RNA.bulge.symbol

The symbol used to represent RNA bulges in the figure to be generated, default to "-"

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plot.match.symbol

The symbol used to represent matched bases in the figure to be generated, default to " "

color.DNA.bulge

The color used to represent DNA bulges in the figure to be generated, default to

size.symbol The size used to plot the bases, and the symbols of DNA/RNA bulges, default

to 3

color.values The color used to represent different bases, DNA bulges, and RNA bulges.

PAM sequence in the target site, please update it to the exact PAM sequence in

the input target site.

body.tile.height

Specifies the height of each plotting tile around each base/symbol for offtargets, default to 2.5

header.tile.height

Specifies the height of each plotting tile around each base/symbol for the target sequence on the very top, default to 3.6

hline.offset Specifies the offset from the top border to draw the horizontal line below the gRNA sequence, default to 3.8. Increase it to move the line down and decrease

it to move the line up.

Optional. If not specified, all the offtargets in the input file specified as off-TargetFile will be included in the plot. With a very large number of offtargets, users can select the top n offtargets to be included in the plot. For example, set plot.top.n = 20 to include only top 20 offtargets in the plot. Please note offtargets are ordered by the n.distinct.UMIs or peak_score from top to bottom.

insertion.score.column

"n.distinct.UMIs" or "peak_score" to be included on

insertion.score.column.prefix

to designate sample name e.g., S1 which means that two of columns are named as S1.peak_score and S1.n.distinct.UMIs in the input file. Useful if the input file is generated by the function combineOfftargets the right side of the alignment as Insertion Events. Relative Insertion Rate (RIR) divided by ontarget peak_score/n.distinct.UMIs. For example, RIR for ontarget should be 100

width.IR For adjusting the width of the IR output

width.RIR For adjusting the width of the RIR output

font family, default to sans (Arial). Other options are serif (Times New Roman)

and mono (Courier). It is possible to use custom fonts with the extrafont package

with the following commands install.packages("extrafont") library(extrafont) font_import()

loadfonts(device = "postscript")

hjust horizontal alignment vjust vertical alignment

Value

a ggplot object

family

Author(s)

Lihua Julie Zhu

Examples

```
offTargetFilePath <- system.file("extdata/forVisualization",
   "offTargetsInPeakRegions.xls",
   package = "GUIDEseq")
fig1 <- plotAlignedOfftargets(offTargetFile = offTargetFilePath,
        plot.top.n = 20,
        plot.match.symbol = ".",
        plot.RNA.bulge.symbol = "-",
        insertion.score.column = "peak_score")
fig1

fig2 <- plotAlignedOfftargets(offTargetFile = offTargetFilePath,
        plot.top.n = 20,
        plot.match.symbol = ".",
        plot.RNA.bulge.symbol = "-",
        insertion.score.column = "n.distinct.UMIs")
fig2</pre>
```

Description

Plot offtargets from multiple samples as heatmap

Usage

```
plotHeatmapOfftargets(
   mergedOfftargets,
   min.detection.rate = 0.1,
   font.size = 12,
   on.target.predicted.score = 1,
IR.normalization = c("sequence.depth", "on.target.score", "sum.score", "none"),
   top.bottom.height.ratio = 3,
   dot.distance.breaks = c(5, 10, 20, 40, 60),
   dot.distance.scaling.factor = c(0.4, 0.6, 0.8, 1.2, 2),
   bottom.start.offset = 8,
   color.low = "white",
   color.high = "blue",
   sample.names,
   insertion.score.column = c("n.distinct.UMIs", "peak_score")
)
```

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Arguments

mergedOfftargets

a data frame from running the combineOfftargets function

min.detection.rate

minimum relative detection rate to be included in the heatmap

font.size for x labels and numbers along the y-axis.

on.target.predicted.score

Default to 1 for the CFDscore scoring method. Set it to 100 for the Hsu-Zhang scoring method.

IR.normalization

Default to sequence.depth which uses the sequencing depth for each sample in the input file to calculate the relative insertion rate (RIR). Other options are "on.target.score" and "sum.score" which use the on-target score for each sample and the sum of all on-target and off-target scores to calculate the RIR respectively. The score can be either peak.score or n.distinct.UMIs as specified by the parameter insertion.score.column

top.bottom.height.ratio

the ratio of the height of top panel vs that of the bottom panel.

dot.distance.breaks

a numeric vector for specifying the minimum number of rows in each panel to use the the corresponding distance in dot.distance.scaling.factor between consecutive dots along the y-axis. In the default setting, dot.distance.breaks and dot.distance.scaling.factor are set to c(5, 10, 20, 40, 60) and c(0.4, 0.6, 0.8, 1.2, 2) respectively, which means that if the number of rows in each panel is greater than or equal to 60, 40-59, 20-39, 10-19, 5-9, and less than 5,then the distance between consecutive dots will be plotted 2, 1.2, 0.8, 0,6, 0.4, and 0.2 (half of 0.4) units away in y-axis respectively.

dot.distance.scaling.factor

a numeric vector for specifing the distance between two consecutive dots. See dot.distance.breaks for more information.

bottom.start.offset

Default to 2, means that place the top number in the bottom panel 2 units below the top border. Increase the value will move the number away from the top border.

color.low The color used to represent the lowest indel rate, default to white

color . high The color used to represent the highest indel rate the intermediate indel rates will be colored using the color between color.low and color.high. Default to blue.

optional sample Names used to label the x-axis. If not provided, x-axis will be labeled using the sample names provided in the GUIDEseqAnalysis step.

insertion.score.column

"n.distinct.UMI" or "peak_score" to be included on the right side of the alignment as Insertion Events. Relative Insertion Rate (RIR) divided by ontarget peak_score/n.distinct.UMI. For example, RIR for ontarget should be 100

Value

a ggplot object

Author(s)

Lihua Julie Zhu

Examples

```
if (interactive())
 mergedOfftargets <-
        read.table(system.file("extdata/forVisualization",
      "mergedOfftargets.txt",
       package = "GUIDEseq"),
                   sep = "\t", header = TRUE)
 figs <- plotHeatmapOfftargets(mergedOfftargets,</pre>
                   min.detection.rate = 2.5,
                   IR.normalization = "on.target.score",
                   top.bottom.height.ratio = 12,
                   bottom.start.offset = 6,
                   dot.distance.scaling.factor = c(0.2, 0.2, 0.4, 0.4, 0.4),
                   sample.names = c("Group1", "Group2"))
                   figs[[1]]/figs[[2]] +
 plot_layout(heights = unit(c(2,1),
                             c('null', 'null')))
figs = plotHeatmapOfftargets(mergedOfftargets,
                 min.detection.rate = 1.2,
                 IR.normalization = "sum.score",
                 top.bottom.height.ratio = 12,
                 bottom.start.offset = 6,
                 dot.distance.scaling.factor = c(0.2, 0.2, 0.4, 0.4, 0.4),
                 sample.names = c("Group1", "Group2"))
                 figs[[1]]/figs[[2]] +
                 plot_layout(heights = unit(c(2,1),
                  c('null', 'null')))
 figs <- plotHeatmapOfftargets(mergedOfftargets,</pre>
   min.detection.rate = 0.2,
    IR.normalization = "sequence.depth",
    top.bottom.height.ratio = 12,
    bottom.start.offset = 6,
    dot.distance.scaling.factor = c(0.2, 0.2, 0.2, 0.2, 0.2),
    sample.names = c("Group1", "Group2"))
figs[[1]]/figs[[2]] +
   plot_layout(heights = unit(c(2,1),
    c('null', 'null')))
figs = plotHeatmapOfftargets(mergedOfftargets,
   min.detection.rate = 3,
    IR.normalization = "none",
    top.bottom.height.ratio = 12,
    bottom.start.offset = 6,
    dot.distance.scaling.factor = c(0.2, 0.2, 0.7, 0.7, 0.7),
    sample.names = c("Group1", "Group2"))
    figs[[1]]/figs[[2]]
```

plotTracks

Plot offtargets as manhantann plots or along all chromosomes with one track per chromosome, or scatter plot for two selected measurements

Description

Plot offtargets as manhantann plots or along all chromosomes with one track per chromosome, or scatter plot for two selected measurements

Usage

```
plotTracks(
  offTargetFile,
  sep = "\t",
  header = TRUE,
  gRNA.size = 20L,
  PAM.size = 3L,
  cleavage.position = 19L,
  chromosome.order = paste0("chr", c(1:22, "X", "Y", "M")),
  xlab = "Chromosome Size (bp)",
  ylab = "Peak Score",
  score.col = c("peak_score", "n.distinct.UMIs", "total.match", "gRNA.match",
    "total.mismatch.bulge", "gRNA.mismatch.bulge", "predicted_cleavage_score"),
  transformation = c("log10", "none"),
  title = "",
  axis.title.size = 12,
  axis.label.size = 8,
  strip.text.y.size = 9,
  off.target.line.size = 0.6,
  on.target.line.size = 1,
  on.target.score = 1,
  on.target.color = "red",
  off.target.color = "black",
  strip.text.y.angle = 0,
  scale.grid = c("free_x", "fixed", "free", "free_y"),
  plot.type = c("manhattan", "tracks", "scatter"),
  family = "serif",
  x.sep = 6e+06,
  plot.zero.logscale = 1e-08,
  scale.chrom = TRUE
)
```

Arguments

offTargetFile The file path containing off-targets generated from GUIDEseqAnalysis

sep The separator in the file, default to tab-delimited

header Indicates whether the input file contains a header, default to TRUE

gRNA. size The size of the gRNA, default 20

PAM. size PAM length, default 3

cleavage.position

the cleavage position of Cas nuclease, default to 19 for SpCas9.

chromosome.order

The chromosome order to plot from top to bottom

xlab The x-asix label, default to Chromosome Size (bp)

ylab The y-asix label, default to Peak Score. Change it to be consistent with the

score.col

score.col The column used as y values in the plot. Available choices are peak_score,

n.distinct.UMIs, total.match, gRNA.match, total.mismatch.bulge, gRNA.mismatch.bulge,

and predicted_cleavage_score. When plot.type is set to scatter, a vector of size two can be set. Otherwise, a scatter plot with log10 transformed n.distinct.UMIs

and log10 transformed predicted_cleavage_score will be plotted.

transformation Indicates whether plot the y-value in log10 scale or in the original scale. When

scale.col is set to total.match, gRNA.match, total.mismatch.bulge, and gRNA.mismatch.bulge,

transformation will not be applied and the data will be plotted in the original scale. When plot.type is set to "scatter", a vector of size two is required when score.col is a vector of size two. Examples are c("log10", "log10"), c("none",

"none"), c(log10", "none"), and c("none", "log10").

title The figure title, default to none.

axis.title.size

The font size for the axis labels, default to 12

axis.label.size

The font size for the tick labels, default to 8

strip.text.y.size

The font size for the strip labels, default to 9

off.target.line.size

The line size to depict the off-targets, default to 0.6

on.target.line.size

The line size to depict the on-targets, default to 1

on.target.score

The score for the on-target, default to 1 for CFD scoring system. This is the maximum score in the chosen scoring system. Change it accordingly if different off-target scoring system is used.

on.target.color

The line color to depict the on-targets, default to red

off.target.color

The line color to depict the off-targets, default to black

strip.text.y.angle

The angel for the y strip text, default to 0. Set it to 45 if angled representation is desired

scale.grid

Used to set the scales in facet_grid, default to free_x, meaning that scales vary across different x-axis, but fixed in y-axis. Other options are fixed, free, and free_y meaning that scales shared across all facets, vary across both x- and yaxises, and vary across y-axis only, respectively. For details, please type ?gg-

plot2::facet_grid

Plot type as tracks by individual chromosome or manhattan plot with all chroplot.type

mosome in one plot

family font family, default to sans (Arial). Other options are serif (Times New Roman)

and mono (Courier). It is possible to use custom fonts with the extrafont package

with the following commands install.packages("extrafont") library(extrafont) font_import()

loadfonts(device = "postscript")

For transforming the x-axis to allow sufficient spaces between small chromox.sep

somes default to 6000000

plot.zero.logscale

Specifying "none" to filter out score.col with zeros when plotting in log10 scale. Specify a very small numeric number if you intend to show the zeros in log scale in the figure. If users specify a number that's bigger than any positive score, plot.zero.logscale will be set to the minimum positive score divided by

scale.chrom Applicable to manhatann plot only. TRUE or FALSE default to TRUE to space offtargets evenly along x-axis.

Value

a ggplot object

Author(s)

Lihua Julie Zhu

Examples

```
if (interactive())
{
   offTargetFilePath <- system.file("extdata/forVisualization",
      "offTargetsInPeakRegions.xls",
       package = "GUIDEseq")
 fig1 <- plotTracks(offTargetFile = offTargetFilePath)</pre>
 fig1
 fig2 <- plotTracks(offTargetFile = offTargetFilePath,</pre>
   score.col = "total.mismatch.bulge",
   ylab = "Total Number of Mismatches and Bulges")
 fig3 <- plotTracks(offTargetFile = offTargetFilePath,</pre>
    score.col = "total.match",
    ylab = "Total Number of Matches")
```

```
fig3
 fig4 <- plotTracks(offTargetFile = offTargetFilePath,</pre>
      score.col = "gRNA.match",
      ylab = "Number of Matches in gRNA")
 fig4
  fig5 <- plotTracks(offTargetFile = offTargetFilePath,</pre>
      score.col = "gRNA.mismatch.bulge",
      ylab = "Number of Mismatches and Bulges in gRNA")
 fig5
 fig6 <- plotTracks(offTargetFile = offTargetFilePath,</pre>
     score.col = "predicted_cleavage_score",
     ylab = "CFD Score",
     scale.grid = "fixed"
     transformation = "none")
 fig6
 ## manhattan plot
 fig <- plotTracks(offTargetFile = offTargetFilePath,</pre>
        score.col = "total.mismatch.bulge", axis.title.size =9,
        plot.type = "manhattan",
        ylab = "Number of Mismatches and Bulges in gRNA Plus PAM")
  fig
 fig <- plotTracks(offTargetFile = offTargetFilePath,</pre>
       score.col = "total.match", axis.title.size =9,
       plot.type = "manhattan",
       ylab = "Number of Matches in gRNA Plus PAM")
   fig
fig <- plotTracks(offTargetFile = offTargetFilePath,</pre>
                 score.col = "gRNA.match",axis.title.size =9,
                 plot.type = "manhattan",
                 ylab = "Number of Matches in gRNA")
fig
fig <- plotTracks(offTargetFile = offTargetFilePath,</pre>
                 score.col = "gRNA.mismatch.bulge", axis.title.size =9,
                 plot.type = "manhattan",
                 ylab = "Number of Mismatches and Bulges in gRNA")
 fig
 plotTracks(offTargetFile = offTargetFilePath,
      #'score.col = "predicted_cleavage_score",
      axis.title.size =9, family = "serif", plot.zero.logscale = 1e-6,
      plot.type = "manhattan", transformation = "log10",
      ylab = "CFD Score")
 plotTracks(offTargetFile = offTargetFilePath,
       score.col = "peak_score",
       axis.title.size =9,
       plot.type = "manhattan",
       ylab = "Number of Insertion Events")
 plotTracks(offTargetFile = offTargetFilePath,
       score.col = "n.distinct.UMIs",
       axis.title.size =9,
```

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```
plot.type = "manhattan",
     ylab = "Number of Insertion Events")
# default scatter plot with blue line from fitting the entire dataset
\# and the red line from fitting the subset with CFD score > 0
plotTracks(offTargetFile = offTargetFilePath,
     axis.title.size =9, plot.zero.logscale = 1e-8,
     plot.type = "scatter")
\# select the x, y, the transformation of x and y,
# and the labels on the scatter plot
plotTracks(offTargetFile = offTargetFilePath,
     axis.title.size =9,
     score.col = c("n.distinct.UMIs", "predicted_cleavage_score"),
     transformation = c("log10", "log10"),
     plot.type = "scatter", plot.zero.logscale = 1e-8,
     xlab = "log10(Number of Insertion Events)",
     ylab = "log10(CFD score)")
}
```

uniqueCleavageEvents example unique cleavage sites

Description

An example data set containing cleavage sites with unique UMI, generated from getUniqueCleavageEvents

Value

cleavage.gr Cleavage sites with one site per UMI as GRanges with metadata column total set to 1 for each range

unique.umi.plus.R2 a data frame containing unique cleavage site from R2 reads mapped to plus strand with the following columns chr.y (chromosome of readSide.y/R2 read) chr.x (chromosome of readSide.x/R1 read) strand.y (strand of readSide.y/R2 read) strand.x (strand of readSide.x/R1 read) start.y (start of readSide.y/R2 read) end.x (start of readSide.x/R1 read) UMI (unique molecular identifier (umi) or umi with the first few bases of R1 read)

unique.umi.minus.R2 a data frame containing unique cleavage site from R2 reads mapped to minus strand with the following columns chr.y (chromosome of readSide.y/R2 read) chr.x (chromosome of readSide.x/R1 read) strand.y (strand of readSide.y/R2 read) strand.x (strand of readSide.x/R1 read) end.y (end of readSide.y/R2 read) start.x (start of readSide.x/R1 read) UMI (unique molecular identifier (umi) or umi with the first few bases of R1 read)

unique.umi.plus.R1 a data frame containing unique cleavage site from R1 reads mapped to minus strand without corresponding R2 reads mapped to the plus strand, with the following columns chr.y (chromosome of readSide.y/R2 read) chr.x (chromosome of readSide.x/R1 read) strand.y

(strand of readSide.y/R2 read) strand.x (strand of readSide.x/R1 read) start.x (start of read-Side.x/R1 read) start.y (start of readSide.y/R2 read) UMI (unique molecular identifier (umi) or umi with the first few bases of R1 read)

unique.umi.minus.R1 a data frame containing unique cleavage site from R1 reads mapped to plus strand without corresponding R2 reads mapped to the minus strand, with the following columns chr.y (chromosome of readSide.y/R2 read) chr.x (chromosome of readSide.x/R1 read) strand.y (strand of readSide.y/R2 read) strand.x (strand of readSide.x/R1 read) end.x (end of readSide.x/R1 read) end.y (end of readSide.y/R2 read) UMI (unique molecular identifier (umi) or umi with the first few bases of R1 read)

all.umi a data frame containing all the mapped reads with the following columns. readName (read ID), chr.x (chromosome of readSide.x/R1 read), start.x (start of eadSide.x/R1 read), end.x (end of eadSide.x/R1 read), mapping.qual.x (mapping quality of readSide.x/R1 read), strand.x (strand of readSide.x/R1 read), cigar.x (CIGAR of readSide.x/R1 read), readSide.x (1/R1), chr.y (chromosome of readSide.y/R2 read) start.y (start of readSide.y/R2 read), end.y (end of readSide.y/R2 read), mapping.qual.y (mapping quality of readSide.y/R2 read), strand.y (strand of readSide.y/R2 read), cigar.y (CIGAR of readSide.y/R2 read), readSide.y (2/R2) R1.base.kept (retained R1 length), R2.base.kept (retained R2 length), distance (distance between mapped R1 and R2), UMI (unique molecular identifier (umi) or umi with the first few bases of R1 read)

Source

http://trace.ncbi.nlm.nih.gov/Traces/sra/?run=SRR1695644

Examples

data(uniqueCleavageEvents)
names(uniqueCleavageEvents)
sapply(uniqueCleavageEvents, class)
uniqueCleavageEvents[[1]] # GRanges object
lapply(uniqueCleavageEvents, dim)

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