

Package ‘SPLINTER’

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Type Package

Title Splice Interpreter of Transcripts

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URL <https://github.com/dianalow/SPLINTER/>

BugReports <https://github.com/dianalow/SPLINTER/issues>

Description Provides tools to analyze alternative splicing sites, interpret outcomes based on sequence information, select and design primers for site validation and give visual representation of the event to guide downstream experiments.

License GPL-2

LazyData TRUE

Depends R (>= 3.6.0), grDevices, stats

Imports graphics, ggplot2, seqLogo, Biostrings, pwalgn, biomaRt, GenomicAlignments, GenomicRanges, GenomicFeatures, Gviz, IRanges, S4Vectors, Seqinfo, utils, plyr, stringr, methods, BSgenome.Mmusculus.UCSC.mm9, googleVis

biocViews ImmunoOncology, GeneExpression, RNASeq, Visualization, AlternativeSplicing

Collate primerpcr.R main_splinter.R

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VignetteBuilder knitr

Suggests txdbmaker, BiocStyle, knitr, rmarkdown

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acceptor.m

acceptor.m

Description

Acceptor site mammalian frequency matrices for GT-AG pairs from SpliceDB

Usage

```
data("acceptor.m")
```

Format

The format is: num [1:4, 1:15] 9 31.03 12.5 42.36 8.44 ... - attr(*, "dimnames")=List of 2 ..\$: chr [1:4] "A" "C" "G" "T" ..\$: chr [1:15] "V1" "V2" "V3" "V4" ...

Source

<http://www.softberry.com/spldb/SpliceDB.html>

References

Burset M., Seledtsov I., Solovyev V. (Nucl.Acids Res.,2000,28,4364-4375; Nucl. Acids Res.,2001,29,255-259)

Examples

```
data(acceptor.m)
```

addEnsemblAnnotation

addEnsemblAnnotation

Description

Adds annotation to [extractSpliceEvents](#) object (if not present)

Usage

```
addEnsemblAnnotation(data, species = "hsapiens")
```

Arguments

data	extractSpliceEvents object
species	character. biomaRt species passed to retrieve annotation. Common species include: 'hsapiens','mmusculus'

Value

[extractSpliceEvents](#) object with annotated genes under \$geneSymbol

Author(s)

Diana Low

See Also

http://asia.ensembl.org/info/data/biomart/biomart_r_package.html#biomartexamples

Examples

```
data_path<-system.file("extdata",package="SPLINTER")
splice_data<-extractSpliceEvents(data=paste(data_path,"/skipped_exons.txt",sep=""))
#splice_data<-addEnsemblAnnotation(data=splice_data,species="mmusculus")
```

 annotateEvents

annotateEvents

Description

Gives detailed description of splicing event in terms of splicing outcome post translation. Currently supports exon skipping and intron retention events.

Usage

```
annotateEvents(
  thedata,
  db,
  bsgenome,
  outputdir,
  full_output = FALSE,
  output_prefix = "results"
)
```

Arguments

thedata	list. output of extractSpliceEvents.
db	TxDb object
bsgenome	BSGenome object
outputdir	character. relative output directory to current location.
full_output	logical. writes out detailed text report and generate figures.
output_prefix	character. text prefix for full_output files.

Value

list containing information on (1) data.frame with splicing regions (2) splice event type

Author(s)

Diana LOW

callPrimer3

callPrimer3

Description

call primer3 for a given set of DNAstringSet object

Usage

```
callPrimer3(
  seq,
  size_range = "150-500",
  Tm = c(57, 59, 62),
  name = "Primer1",
  primer3 = "primer3-2.3.7/bin/primer3_core",
  thermo.param = "primer3-2.3.7/src/primer3_config/",
  sequence_target = NULL,
  settings = "primer3-2.3.7/primer3web_v4_0_0_default_settings.txt"
)
```

Arguments

seq	DNAstring object, one DNA string for the given amplicon
size_range	default: '151-500'
Tm	melting temprature parameters default:c(55,57,58)
name	name of the amplicon in chr_start_end format
primer3	primer3 path
thermo.param	thermodynamic parameters folder
sequence_target	If one or more targets is specified then a legal primer pair must flank at least one of them.
settings	text file for parameters

Details

modified to include SEQUENCE_TARGET as an option

Value

data.frame of designed primers and parameters

Author(s)

Altuna Akalin's modified Arnaud Krebs' original function further modified here by Diana Low

Examples

```
### NOT RUN ###
# primer_results<-callPrimer3(seq='')
```

checkPrimer

checkPrimer

Description

checkPrimer

Usage

```
checkPrimer(pp, genome, roi = NULL)
```

Arguments

pp	data.frame defining primers, or output of callPrimer3 . minimal columns = PRIMER_LEFT_SEQUENCE,PRIMER_RIGHT_SEQUENCE
genome	BSgenome object
roi	makeROI object

Value

list of GRanges with primer locations

Author(s)

Diana Low

Examples

```
# create a primer pair
roi
primer_pair <- data.frame(PRIMER_LEFT_SEQUENCE="agctcttgaaattggagctgac",
                          PRIMER_RIGHT_SEQUENCE="cttagaaagaacaggaaatcc",
                          stringsAsFactors=FALSE)
```

compatible_cds*compatible_cds*

Description

compatible_cds

Examples

```
data(compatible_cds)
## maybe str(compatible_cds) ; plot(compatible_cds) ...
```

compatible_tx	<i>compatible_tx</i>
---------------	----------------------

Description

compatible_tx

Examples

```
data(compatible_tx)
## maybe str(compatible_tx) ; plot(compatible_tx) ...
```

donor.m	<i>donor.m</i>
---------	----------------

Description

Donor site mammalian frequency matrices for GT-AG pairs from SpliceDB

Usage

```
data("donor.m")
```

Format

The format is: num [1:4, 1:9] 34.1 36.2 18.3 11.4 60.4 ... - attr(*, "dimnames")=List of 2 ..\$: chr [1:4] "A" "C" "G" "T" ..\$: chr [1:9] "V1" "V2" "V3" "V4" ...

Source

<http://www.softberry.com/spldb/SpliceDB.html>

References

Burset M., Seledtsov I., Solovyev V. (Nucl.Acids Res.,2000,28,4364-4375; Nucl. Acids Res.,2001,29,255-259)

Examples

```
data(donor.m)
```

eventOutcomeCompare *eventOutcomeCompare*

Description

Compares two sequences and gives differences if there's a switch from 1->2 if seq2 is NULL, assume seq1 is a list of length 2 to compare

Usage

```
eventOutcomeCompare(
  seq1,
  seq2 = NULL,
  genome,
  direction = TRUE,
  fullseq = TRUE,
  verbose = FALSE
)
```

Arguments

seq1	GRangesList
seq2	GRangesList
genome	BSGenome object
direction	logical. Report direction of sequence change.
fullseq	logical. Report full sequences.
verbose	logical. turn messages on/off.

Value

list containing
 (1) tt : PairwiseAlignmentsSingleSubject pairwise alignment
 (2) eventtypes : string detailing primary event classification

Author(s)

Diana LOW

Examples

```
suppressMessages(library(BSgenome.Mmusculus.UCSC.mm9))
bsgenome<-BSgenome.Mmusculus.UCSC.mm9
eventOutcomeCompare(seq1=compatible_cds$hits[[1]],seq2=region_minus_exon,
  genome=bsgenome,direction=TRUE)
```

`eventOutcomeTranslate` *eventOutcomeTranslate*

Description

translates sequences, reports if NMD or NTC

Usage

```
eventOutcomeTranslate(  
  seq1,  
  genome,  
  direction = FALSE,  
  fullseq = TRUE,  
  verbose = FALSE  
)
```

Arguments

<code>seq1</code>	<code>GRangesList</code>
<code>genome</code>	<code>BSGenome</code> object
<code>direction</code>	logical. Report direction of sequence change.
<code>fullseq</code>	logical. Output full AA sequence.
<code>verbose</code>	logical. turn messages on/off.

Value

list of translated sequences

Author(s)

Diana LOW

Examples

```
suppressMessages(library(BSgenome.Mmusculus.UCSC.mm9))  
bsgenome<-BSgenome.Mmusculus.UCSC.mm9  
translation_results<-eventOutcomeTranslate(compatible_cds,genome=bsgenome,  
  direction=TRUE)
```

eventPlot

*eventPlot***Description**

eventPlot

Usage

```
eventPlot(
  transcripts,
  roi_plot = NULL,
  bams = c(),
  names = c(),
  annoLabel = c("Gene A"),
  rspan = 1000,
  pfam_dom = NULL,
  showAll = TRUE
)
```

Arguments

transcripts	GRanges object
roi_plot	GRanges object region to plot
bams	character vector of bam file locations
names	character vector of name labels
annoLabel	character. annotation label
rspan	integer or NULL. number of basepairs to span from roi. if NULL, will consider whole gene of roi
pfam_dom	optional GRanges object of PFAM domains from UCSC Tables.
showAll	logical. TRUE = display splice junctions of entire view or FALSE = just roi.

Value

a Gviz plot of genomic region

Author(s)

Diana Low

Examples

```
# define BAM files
data_path<-system.file("extdata",package="SPLINTER")
mt<-paste(data_path,"/mt_chr14.bam",sep="")
wt<-paste(data_path,"/wt_chr14.bam",sep="")

# plot results
eventPlot(transcripts=valid_tx,roi_plot=roi,bams=c(wt,mt),
  names=c('wt','mt'),rspan=1000)
```

extendROI	<i>extendROI</i>
-----------	------------------

Description

extend the span of the current ROI by n number of up/downstream exon(s) by modifying roi_range within the makeROI object while retaining legacy sites by keeping \$roi and \$flank

Usage

```
extendROI(roi, tx, up = 0, down = 0, type = 1)
```

Arguments

roi	makeROI object
tx	GRangesList transcript list to pull regions from
up	integer. number of exons to extend upstream
down	integer. number of exons to extend downstream
type	integer. 1=full cassette, 2=flank only

Value

[makeROI](#) object with modified ranges

Examples

```
extendROI(roi,valid_tx,up=1)
```

extractSpliceEvents	<i>extractSpliceEvents</i>
---------------------	----------------------------

Description

Extracts the location of target, upstream and downstream splice sites Used for calculations and genome visualizations

Usage

```
extractSpliceEvents(
  data = NULL,
  filetype = "mats",
  splicetype = "SE",
  fdr = 1,
  inclusion = 1,
  start0 = TRUE
)
```

Arguments

data	character. path to file
filetype	character. type of splicing output. c('mats','custom'). see Details.
splicetype	character. c('SE', 'RI', 'MXE', 'A5SS', 'A3SS')
fdr	numeric. false discovery rate filter range [0,1]
inclusion	numeric. splicing inclusion range, takes absolute value
start0	boolean 0-base start

Details

filetype 'custom' should provide a 9-column tab-delimited text file with the following columns: ID (Ensembl gene id), Symbol (gene name), chr, strand, exonStart, exonEnd, exon2Start, exon2End, upstreamStart, upstreamEnd, downstreamStart, downstreamEnd eg. ENSG0000012345 chr1 + 3 4 5 6 1 2 7 8

Value

list containing information on
 (1) original file type
 (2) splice event type
 (3) data.frame with splicing regions

Author(s)

Diana Low

See Also

http://rnaseq-mats.sourceforge.net/user_guide.htm for MATS file definition

Examples

```
data_path<-system.file("extdata",package="SPLINTER")
splice_data<-extractSpliceEvents(data=paste(data_path,"/skipped_exons.txt",sep=""))
```

extractSpliceSites	<i>extractSpliceSites</i>
--------------------	---------------------------

Description

Extracts and formats to bed the location of target, upstream and downstream splice sites

Usage

```
extractSpliceSites(
  df,
  target = "SE",
  site = "donor",
  motif_range = c(-3, 6),
  start0 = TRUE
)
```

Arguments

df	extractSpliceEvents object
target	the target site to extract. See Details.
site	character donor or acceptor
motif_range	numeric vector of splice position to extract
start0	boolean 0-base start

Details

target : the site to extract the sequence from. It can be either the event in question (SE, RI, MXE - first exon, MXE2 - second exon, A5SSlong, A5SSshort, A3SSlong, A3SSshort, upstream or downstream). If this function is used in conjunction with [shapiroDonor](#) or [shapiroAcceptor](#) to compute scores, then most likely it will be run twice - once for the event, and the other either up- or downstream as a comparison.

Value

GRanges object

Author(s)

Diana Low

See Also

http://rnaseq-mats.sourceforge.net/user_guide.htm for MATS file definition

Examples

```
data_path<-system.file("extdata",package="SPLINTER")
splice_data<-extractSpliceEvents(data=paste(data_path, "/skipped_exons.txt", sep=""))
splice_sites<-extractSpliceSites(splice_data, target="SE")
```

findCompatibleEvents *findCompatibleEvents*

Description

Which transcript contains the event? Each event has 2 possibilities, as long as the transcript fulfills one, it passes the test Has to be exact (inner junctions)

Usage

```
findCompatibleEvents(tx, tx2 = NULL, roi, sequential = TRUE, verbose = FALSE)
```

Arguments

tx	GRangesList object of transcripts
tx2	optional GRangesList object of transcripts if tx is list of cds
roi	makeROI object containing event information
sequential	logical. Exons have to appear sequentially to be considered compatible
verbose	logical. printouts and messages.

Details

Seperates into event/region1 and 2 for the alternative case

Value

list of length 4
 (1) GRangesList
 (2) Hits status [c]=coding; [nc]=non-coding
 (3) ct - compatible transcripts
 (4) tt - total transcripts

Author(s)

Diana Low

Examples

```
compatible_cds <- findCompatibleEvents(valid_cds,roi=roi,verbose=TRUE)
```

findCompatibleExon	<i>findCompatibleExon</i>
--------------------	---------------------------

Description

Finds compatible exon in annotation with the one present in roi object

Usage

```
findCompatibleExon(tx, roi, verbose = FALSE)
```

Arguments

tx	GRangesList object of transcripts
roi	makeROI object containng event information
verbose	logical. printouts and messages.

Value

list of length 3
 (1) GRangesList hits
 (2) Number of transcripts
 (3) Original number of input transcripts

Author(s)

Diana Low

Examples

```
compatible_exons <- findCompatibleExon(valid_cds,roi)
```

findExactOverlaps	<i>findExactOverlaps</i>
-------------------	--------------------------

Description

Internal function similar to findSpliceOverlaps but only preserves internal flanks

Usage

```
findExactOverlaps(query, subject, sequential = FALSE, verbose = FALSE)
```

Arguments

query	GRanges object
subject	GRanges object
sequential	logical. TRUE if exons are sequential.
verbose	logical. report intermediate output

Value

Hits object

Author(s)

Diana Low

findTermination	<i>findTermination</i>
-----------------	------------------------

Description

Internal function to find the first stop codon that occurs in the AA sequence, returns their position and the resulting truncated protein

Usage

```
findTermination(s1)
```

Arguments

s1	character. protein sequence
----	-----------------------------

Value

list containing
(1) stop1 : stop position
(2) s1 : sequence truncated to first stop

Author(s)

Diana LOW

findTX	<i>findTX</i>
--------	---------------

Description

Given an ENSEMBL id, find all transcripts that matches id

Usage

```
findTX(id, db, tx, valid = FALSE, verbose = FALSE)
```

Arguments

id	character. transcript identification (currently ENSEMBL gene names)
db	TxDb object
tx	GRangesList
valid	logical. check if in multiples of 3 [TRUE] for CDS translation.
verbose	logical. turn messages on/off.

Value

GRangesList

Author(s)

Diana Low

Examples

```
valid_cds <- findTX(id=splice_data$data[2,]$ID, tx=thecds, db=txdb, valid=FALSE)
```

getPCRsizes	<i>getPCRsizes</i>
-------------	--------------------

Description

returns length of product given a GRanges span and GRangesList of transcripts

Usage

```
getPCRsizes(pcr_span, txlist, verbose = FALSE)
```

Arguments

pcr_span	GRanges object
txlist	GRangesList object
verbose	logical. report intermediate output.

Value

data.frame of transcript names with detected sizes in basepairs

Author(s)

Diana Low

Examples

```
suppressMessages(library(BSgenome.Mmusculus.UCSC.mm9))
bsgenome<-BSgenome.Mmusculus.UCSC.mm9
## create a primer pair
## for actual use, obtain primer pair from primer design (callPrimer3)
primer_pair <- data.frame(PRIMER_LEFT_SEQUENCE="agctcttgaaattggagctgac",
                          PRIMER_RIGHT_SEQUENCE="cttagaaagaacaggaaatcc",
                          stringsAsFactors=FALSE)

## confirm location
cp<-checkPrimer(primer_pair,bsgenome,roi)
cp

## get the PCR sizes
pcr_result1 <- getPCRsizes(cp,theexons)
```

getRegionDNA

getRegionDNA

Description

get DNA sequence give a region of interest

Usage

```
getRegionDNA(roi, genome, introns = FALSE)
```

Arguments

roi	makeROI object
genome	BSgenome object
introns	TRUE/FALSE. whether to include intronic (lowercase) DNA. By default returns only exonic (uppercase) DNA.

Value

list of
(1) DNA sequence (2) Junction start (for primer design)

Author(s)

Diana Low

Examples

```
suppressMessages(library(BSgenome.Mmusculus.UCSC.mm9))
bsgenome<-BSgenome.Mmusculus.UCSC.mm9
getRegionDNA(roi,bsgenome)
```

insertRegion	<i>insertRegion</i>
--------------	---------------------

Description

inserts a region (exon or intron) into roi object

Usage

```
insertRegion(subject, roi)
```

Arguments

subject	GrangesList
roi	makeROI object containg region of interest (to insert). refer to makeROI().

Details

in the case of intron retention, replaces exon with intron retention range reduce() the GRanges in question

Value

GRanges object

Author(s)

Diana Low

Examples

```
#Inserts the exon defined in roi GRanges object from a GRanges/GRangesList
region_minus_exon
region_with_exon<-insertRegion(region_minus_exon,roi)
```

makeROI

*makeROI***Description**

Creates an object to store information about the splice site (region of interest) including flanking regions and alternative splice outcome

Usage

```
makeROI(df, type = "SE")
```

Arguments

df data.frame object from [extractSpliceEvents](#)
 type type of splicing event c("SE", "RI", "MXE", "A5SS", "A3SS")

Value

a list containing
 (1) type : splice type
 (2) name : ID of transcript
 (3) roi : GRanges object of splice site
 (4) flank : GRanges object of flanking exons of splice site
 (5) roi_range : GRangesList of splice site and it's alternative outcome based on type

Author(s)

Diana Low

Examples

```
single_record<-splice_data$data[which(grepl("Prmt5", splice_data$data$Symbol)),]  
roi <- makeROI(single_record, type="SE")
```

makeUniqueIDs

*makeUniqueIDs***Description**

Makes unique ID names from event location

Usage

```
makeUniqueIDs(ddata)
```

Arguments

ddata extractSpliceEvents object

Value

original extractSpliceEvents list object with unique ID appended to data accessor

Author(s)

Diana Low

Examples

```
data_with_id<-makeUniqueIDs(splice_data)
```

matchExons	<i>matchExons</i>
------------	-------------------

Description

Internal function to help match the inner coordinates of a 2/3 cassette checks if reference and subject matches

Usage

```
matchExons(ref, subject)
```

Arguments

ref	GRanges object
subject	GRanges object

Value

logical. check if exons match (TRUE) or not (FALSE)

Author(s)

Diana Low

metaremove	<i>metaremove</i>
------------	-------------------

Description

helper function to remove metadata from GRanges object

Usage

```
metaremove(x)
```

Arguments

x	GRanges or GRangesList
---	------------------------

Value

GRanges or GRangesList

<code>pcr_result1</code>	<i>pcr_result1</i>
--------------------------	--------------------

Description

`pcr_result1`

Examples

```
data(pcr_result1)
```

<code>plot_seqlogo</code>	<i>plotting sequence logo</i>
---------------------------	-------------------------------

Description

Plots the sequence logo of a given set of FASTA sequences

Usage

```
plot_seqlogo(fasta_seq)
```

Arguments

`fasta_seq` DNASTringSet or path to fasta-formatted file

Value

sequence logo image

Author(s)

Diana Low

Examples

```
head(splice_fasta)
plot_seqlogo(Biostrings::DNASTringSet(splice_fasta$V2))
```

primers

primers

Description

primers designed using Primer3 for sample data

Usage

```
data("primers")
```

Format

A data frame with 5 observations on the following 28 variables.

`i` a numeric vector

`PRIMER_LEFT_SEQUENCE` a character vector

`PRIMER_RIGHT_SEQUENCE` a character vector

`PRIMER_LEFT_TM` a numeric vector

`PRIMER_RIGHT_TM` a numeric vector

`PRIMER_LEFT_pos` a numeric vector

`PRIMER_LEFT_len` a numeric vector

`PRIMER_RIGHT_pos` a numeric vector

`PRIMER_RIGHT_len` a numeric vector

`PRIMER_PAIR_PENALTY` a numeric vector

`PRIMER_LEFT_PENALTY` a numeric vector

`PRIMER_RIGHT_PENALTY` a numeric vector

`PRIMER_LEFT_GC_PERCENT` a numeric vector

`PRIMER_RIGHT_GC_PERCENT` a numeric vector

`PRIMER_LEFT_SELF_ANY_TH` a numeric vector

`PRIMER_RIGHT_SELF_ANY_TH` a numeric vector

`PRIMER_LEFT_SELF_END_TH` a numeric vector

`PRIMER_RIGHT_SELF_END_TH` a numeric vector

`PRIMER_LEFT_HAIRPIN_TH` a numeric vector

`PRIMER_RIGHT_HAIRPIN_TH` a numeric vector

`PRIMER_LEFT_END_STABILITY` a numeric vector

`PRIMER_RIGHT_END_STABILITY` a numeric vector

`PRIMER_LEFT_TEMPLATE_MISPRIMING` a numeric vector

`PRIMER_RIGHT_TEMPLATE_MISPRIMING` a numeric vector

`PRIMER_PAIR_COMPL_ANY_TH` a numeric vector

`PRIMER_PAIR_COMPL_END_TH` a numeric vector

`PRIMER_PAIR_PRODUCT_SIZE` a numeric vector

`PRIMER_PAIR_TEMPLATE_MISPRIMING` a numeric vector

Value

Dataframe of primer design results

Examples

```
data(primers)
```

psiPlot

psiPlot

Description

Plots percentage spliced in (PSI) values in terms of inclusion levels

Usage

```
psiPlot(df = NULL, type = "MATS", sample_labels = c("Sample 1", "Sample 2"))
```

Arguments

df	data.frame containing PSI values
type	character. either 'MATS' output (will read in MATS headers) or 'generic' (provide 4 or 6 column data.frame)
sample_labels	x-axis labels for the plot

Value

bar plot of PSI values

Author(s)

Diana Low

Examples

```
#we give inclusion and skipped numbers as reads
#this will be converted into percentages
df<-data.frame(inclusion1=c("6,4,6"),skipped1=c("10,12,12"),inclusion2=c("15,15,15"),
               skipped2=c("3,3,4"),stringsAsFactors = FALSE)
psiPlot(df,type='generic')
```

region_minus_exon

region_minus_exon

Description

region_minus_exon

Examples

```
data(region_minus_exon)
## maybe str(region_minus_exon) ; plot(region_minus_exon) ...
```

removeRegion	<i>removeRegion</i>
--------------	---------------------

Description

removes a region (exon) from a GRanges or GRangesList

Usage

```
removeRegion(subject, roi)
```

Arguments

subject	GRanges or GrangesList object
roi	makeROI object containing GRanges range (to remove)

Value

GRanges object

Author(s)

Diana Low

```
# Removes the exon defined in roi GRanges object from a GRanges/GRangesList compatible_cds$hits[[1]]
region_minus_exon<-removeRegion(compatible_cds$hits[[1]],roi)
```

remvalue	<i>remvalue</i>
----------	-----------------

Description

helper function to remove metadata from GRanges object used within metaremove

Usage

```
remvalue(x)
```

Arguments

x	GRanges or GRangesList
---	------------------------

Value

GRanges or GRangesList

roi	<i>roi</i>
-----	------------

Description

roi

Usage

```
data("roi")
```

Value

List containing region of interest information

Examples

```
data(roi)
```

shapiroAcceptor	<i>shapiroAcceptor</i>
-----------------	------------------------

Description

Shapiro's score of acceptor site (range is from -13 [intron] to +1 [exon]) is: $100 * ((t1 - l1)/(h1 - l1) + (t2 - l2)/(h2 - l2))/2$, where t1 is the sum of the best 8 of 10 percentages at positions -13 to -4, l1 is the sum of the lowest 8 of 10 percentages at position -13 to -4, h1 is the sum of the highest 8 of 10 percentages at positions -13 to -4, t2 is the sum of percentages at positions -3 to +1, l2 is the sum of the lowest percentages at positions -3 to +1, and h2 is the sum of the highest percentages at positions -3 to +1

Usage

```
shapiroAcceptor(reference_fasta, target_fasta)
```

Arguments

reference_fasta	vector of strings or DNASTringSet of reference splice list
target_fasta	vector of strings or DNASTringSet of fasta to score

Value

data.frame with Shapiro scores

Author(s)

Diana Low

See Also

<http://www.softberry.com/spldb/SpliceDB.html>

Examples

```
library(BSgenome.Mmusculus.UCSC.mm9)
bsgenome <- BSgenome.Mmusculus.UCSC.mm9
data_path<-system.file("extdata",package="SPLINTER")
splice_data<-extractSpliceEvents(data=paste(data_path,"/skipped_exons.txt",sep=""))
splice_sites<-extractSpliceSites(splice_data,site="acceptor")
acceptor.ss<-getSeq(bsgenome,splice_sites)
##sacceptor<-shapiroAcceptor(acceptor.m,acceptor.ss)
```

shapiroDensity

shapiroDensity

Description

convenience function for plotting Shapiro score density

Usage

```
shapiroDensity(ctrl_scores, treat_scores, sample = c(1, 2))
```

Arguments

ctrl_scores	output of shapiroDonor or shapiroAcceptor
treat_scores	output of shapiroDonor or shapiroAcceptor
sample	samplenames

Value

density plot of Shapiro scores

Author(s)

Diana Low

shapiroDonor

*shapiroDonor***Description**

Shapiro and Senapathy (1987) have developed a method to score the strength of a splice site based on percentages of each nucleotide at each position. Shapiro's score of donor site (range is from -3 [exon] to +7 [intron]) is : $100 * (t - \min) / (\max - \min)$, where t is the sum of percentages at positions -3 to +7, min is the sum of the lowest percentages at positions -3 to +7, and max is the sum of the highest percentages at positions -3 to +7.

Usage

```
shapiroDonor(reference_fasta, target_fasta)
```

Arguments

```
reference_fasta      vector of strings or DNASTringSet of reference splice list
target_fasta         vector of strings or DNASTringSet of fasta to score
```

Value

```
data.frame with Shapiro scores
```

Author(s)

```
Diana Low
```

See Also

```
http://www.softberry.com/spldb/SpliceDB.html
```

Examples

```
library(BSgenome.Mmusculus.UCSC.mm9)
bsgenome <- BSgenome.Mmusculus.UCSC.mm9
data_path<-system.file("extdata",package="SPLINTER")
splice_data<-extractSpliceEvents(data=paste(data_path,"/skipped_exons.txt",sep=""))
splice_sites<-extractSpliceSites(splice_data)
donor.ss<-getSeq(bsgenome,splice_sites)
##sdonor<-shapiroDonor(donor.m,donor.ss)
```

splice_data	<i>splice_data</i>
-------------	--------------------

Description

splice_data

Usage

```
data("splice_data")
```

Value

List containing splice event file information

Examples

```
data(splice_data)
```

splice_fasta	<i>splice_fasta</i>
--------------	---------------------

Description

splice_fasta

Usage

```
data("splice_fasta")
```

Format

A data frame with 0 observations on the following 2 variables.

V1 a numeric vector

V2 a numeric vector

Value

Dataframe of region and fasta sequence

Examples

```
data(splice_fasta)
```

`splitPCRhit`*splitPCRhit*

Description

splits the PCR alignment into the two AS conditions

Usage

```
splitPCRhit(res, hitlist)
```

Arguments

res	result from getPCRsizes
hitlist	findCompatibleEvents object

Value

list of 2 data.frame objects with isoform name (ID) and length of PCR product (bp) matching Type 1 or Type 2 transcripts

Author(s)

Diana Low

Examples

```
## as getPCRsizes gives you all PCR bands when the primers are used,  
## splitPCRhit will determine which bands are relevant to the target  
relevant_pcr_bands<-splitPCRhit(pcr_result1,compatible_tx)
```

`thecds`*thecds*

Description

thecds

Usage

```
data("thecds")
```

Value

List containing GRanges info

Examples

```
data(thecds)
```

theexons	<i>theexons</i>
----------	-----------------

Description

theexons

Usage

```
data("theexons")
```

Value

List containing GRanges info

Examples

```
data(theexons)
```

valid_cds	<i>valid_cds</i>
-----------	------------------

Description

valid_cds

Usage

```
data("valid_cds")
```

Value

GRangesList

Examples

```
data(valid_cds)
```

valid_tx	<i>valid_tx</i>
----------	-----------------

Description

valid_tx

Value

GRangesList

Examples

```
data(valid_tx)
## maybe str(valid_tx) ; plot(valid_tx) ...
```

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