Package 'branchpointer'

July 13, 2025

Type Package

Title Prediction of intronic splicing branchpoints

Version 1.35.0

Date 2017-07-14

Author Beth Signal

Maintainer Beth Signal <b.signal@garvan.org.au>

Description Predicts branchpoint probability for sites in intronic branchpoint windows. Queries can be supplied as intronic regions; or to evaluate the effects of mutations, SNPs.

License BSD_3_clause + file LICENSE

LazyData FALSE

Depends caret, R(>= 3.4)

Imports plyr, kernlab, gbm, stringr, cowplot, ggplot2, biomaRt, Biostrings, parallel, utils, stats, BSgenome.Hsapiens.UCSC.hg38, rtracklayer, GenomicRanges, GenomeInfoDb, IRanges, S4Vectors, data.table

Suggests knitr, BiocStyle

RoxygenNote 6.0.1

VignetteBuilder knitr

biocViews Software, GenomeAnnotation, GenomicVariation, MotifAnnotation

git_url https://git.bioconductor.org/packages/branchpointer

git_branch devel

git_last_commit dedb6d2

git_last_commit_date 2025-04-15

Repository Bioconductor 3.22

Date/Publication 2025-07-13

Contents

xonsToIntrons	. 2
etBranchpointSequence	. 2
etCanonical3SS	. 3
etExonDists	. 4

getBranchpointSequence

getPPT	4
getQueryLoc	5
gtfToExons	5
makeBranchpointWindowForExons	6
makeBranchpointWindowForSNP	7
plotBranchpointWindow	7
plotStructure	8
predictBranchpoints	9
predictionsToSummary	10
readQueryFile	11
	12

Index

exonsToIntrons Convert exon annotation GRanges to intron locations

Description

Converts exon annotation to intron locations overlapping the branchpoint region for exculsion of non-branchpoint region SNPs Returns a character vector of chromosome locations

Usage

exonsToIntrons(exons, maxDist = 50)

Arguments

exons	GRanges containing exon co-ordinates. Should be produced by gtfToExons()
maxDist	Maximum distance from the 3' exon to create the branchpoint region.

Value

GRanges containing intron window co-ordinates

Author(s)

Beth Signal

getBranchpointSequence

Get branchpoint sequence features Gets intronic sequence covering the branchpoint window and extracts predictive features

Description

Get branchpoint sequence features Gets intronic sequence covering the branchpoint window and extracts predictive features

getCanonical3SS

Usage

```
getBranchpointSequence(query, uniqueId = "test", queryType,
workingDirectory = ".", genome = NA, bedtoolsLocation = NA,
BSgenome = NULL, useParallel = FALSE, cores = 1, rmChr = FALSE)
```

Arguments

query	branchpointer query GenomicRanges	
uniqueId	unique string identifier for intermediate .bed and .fa files.	
queryType	type of branchpointer query. "SNP" or "region".	
workingDirecto	ry	
	directory where intermediate .bed and .fa are located	
genome	.fa genome file location	
bedtoolsLocation		
	bedtools binary location (which bedtools)	
BSgenome	BSgenome object	
useParallel	use parallelisation to speed up code?	
cores	number of cores to use in parallelisation (default = 1)	
rmChr	remove "chr" before chromosome names before writing bed file. Required if genome sequence names do not contain "chr"	

Value

GenomicRanges with all features required to predict branchpoint probability scores

Author(s)

Beth Signal

getCanonical3SS Get locations of the first five AG 3' splice site motifs

Description

Takes a variable length vector of distances to the AG motif, sorts and returns the first five. If there are less than five elements in the vector, returns the sorted vector and fills the remainder of the values with 300.

Usage

```
getCanonical3SS(ag)
```

Arguments

ag

Vector of distances to the AG splice site motif.

Value

Locations of the first five AG dinucleotides

Author(s)

Beth Signal

getExonDists Get the closest 3' and 5' exons

Description

Finds the closest annotated exons from a genomic co-ordinate. Returns the distance to the 3' exon, distance to the 5' exon, ids of the 3' and 5' exon, and if the exons are from the same parent gene

Usage

getExonDists(query, exons, queryType)

Arguments

query	GenomicRangesquery
exons	GenomicRanges containing exon co-ordinates. Should be produced by gtfToExons()
queryType	type of query. "SNP" or "region"

Value

GenomicRanges with distance to the closest 3' and 5' exons, whether these exons are part of the same gene (i.e is the location intronic), and the identifiers for the 3' and 5' exons.

Author(s)

Beth Signal

getPPT

Get the best polypyramidine tract

Description

Takes a query genomic sequence, finds all potential polypyramidine tracts (PPTs) between the test site and the annotated 3'exon. Returns the distance to the start of the longest PPT, and its length.

Usage

getPPT(attributes)

Arguments

attributes query attributes GenomicRanges

Value

distance to the start of the longest PPT, and its length

getQueryLoc

Author(s)

Beth Signal

getQueryLoc

Find the closest 3' and 5' exons to a branchpointer query

Description

Finds the closest annotated exons from genomic co-ordinates in a branchpointer query GRanges

Usage

```
getQueryLoc(query, queryType, maxDist = 50, filter = TRUE, exons)
```

Arguments

query	branchpointer query GenomicRanges must have chromosome at position 2, genomic co-ordinate at position 3, and strand at position 4.
queryType	type of query file ("SNP" or "region")
maxDist	maximum distance a SNP can be from an annotated 3' exon.
filter	remove SNP queries prior to finding finding nearest exons.
exons	data.frame containing exon co-ordinates. Should be produced by gtfToExons()

Value

GenomicRanges with the query and its location relative to the 3' and 5' exons

Author(s)

Beth Signal

gtfToExons

Convert GTF file to exon location file

Description

Converts a GTF annotation to exon locations

Usage

gtfToExons(gtf)

Arguments

gtf file containing the gtf annotation.

Value

exon annotation GRanges

Author(s)

Beth Signal

Examples

```
smallExons <- system.file("extdata","gencode.v26.annotation.small.gtf",
package = "branchpointer")
exons <- gtfToExons(smallExons)</pre>
```

makeBranchpointWindowForExons

Make branchpoint window regions

Description

Generate branchpoint window regions corresponding to annotated exon(s) within a queried gene, transcript or exon id

Usage

makeBranchpointWindowForExons(id, idType, exons, forceClosestExon = FALSE)

Arguments

id	identifier(s) for the query gene/transcript/exon id	
idType	type of id to match in the exon annotation file ("gene_id", "transcript_id", or "exon_id")	
exons	GRanges containing exon co-ordinates.	
forceClosestExon		
	Force branchpointer to find the closest exon and not the exon annotated as 5' to the query	

Value

Granges with formatted query

Author(s)

Beth Signal

Examples

```
smallExons <- system.file("extdata","gencode.v26.annotation.small.gtf",package = "branchpointer")
exons <- gtfToExons(smallExons)
windowquery <- makeBranchpointWindowForExons("ENSG00000139618.14", "gene_id", exons)
windowquery <- makeBranchpointWindowForExons("ENST00000357654.7", "transcript_id", exons)
windowquery <- makeBranchpointWindowForExons("ENSE00003518965.1", "exon_id", exons)</pre>
```

6

makeBranchpointWindowForSNP

Makes a branchpointer formatted GRanges object from refsnp ids

Description

Searches Biomart for refsnp ids, and pulls genomic location and sequence identity information Reformats alleles so each query has only one alternative allele

Usage

```
makeBranchpointWindowForSNP(refSNP, mart.snp, exons, maxDist = 50,
filter = TRUE)
```

Arguments

refSNP	Vector of refsnp ids
mart.snp	biomaRt mart object specifying the BioMart database and dataset to be used
exons	GRanges containing exon co-ordinates. Should be produced by gtfToExons()
maxDist	maximum distance a SNP can be from an annotated 3' exon.
filter	remove SNP queries prior to finding finding nearest exons?

Value

formatted SNP query GRanges

Author(s)

Beth Signal

Examples

```
smallExons <- system.file("extdata","gencode.v26.annotation.small.gtf",package = "branchpointer")
exons <- gtfToExons(smallExons)</pre>
```

mart.snp <- biomaRt::useMart("ENSEMBL_MART_SNP", dataset="hsapiens_snp", host="www.ensembl.org")
query <- makeBranchpointWindowForSNP("rs587776767", mart.snp, exons)</pre>

plotBranchpointWindow Plots branchpointer predictions

Description

Plots branchpointer predictions

Usage

```
plotBranchpointWindow(queryName, predictions, probabilityCutoff = 0.52,
    plotMutated = FALSE, plotStructure = TRUE, exons)
```

Arguments

queryName	query id used to identify the SNP or region
predictions	Granges object generated by predictBranchpoints()
probabilityCuto	ff
	probability score cutoff value for displaying U2 binding energy
plotMutated	plot alternative sequence predicitons alongside reference sequence predictions
plotStructure	plot structures for gene and 3' exon containing and skipping isoforms
exons	Granges containing exon co-ordinates. Should be produced by gtfToExons()

Value

ggplot2 plot with branchpoint features in the specified intronic region

Author(s)

Beth Signal

Examples

```
smallExons <- system.file("extdata","gencode.v26.annotation.small.gtf",
package = "branchpointer")
exons <- gtfToExons(smallExons)
g <- BSgenome.Hsapiens.UCSC.hg38::BSgenome.Hsapiens.UCSC.hg38</pre>
```

```
querySNPFile <- system.file("extdata","SNP_example.txt", package = "branchpointer")
querySNP <- readQueryFile(querySNPFile,queryType = "SNP",exons = exons, filter = FALSE)
predictionsSNP <- predictBranchpoints(querySNP,queryType = "SNP",BSgenome = g)
plotBranchpointWindow(querySNP$id[1], predictionsSNP,
plotMutated = TRUE, exons = exons)</pre>
```

plotStructure Plots transcript structures

Description

Plots transcript structures

Usage

```
plotStructure(exonID, exons, keepTranscripts = "overlapping")
```

Arguments

exonID	id of the exon to plot
exons	Granges containing exon co-ordinates.
keepTranscripts	
	which transcripts to plot ("overlapping" or "withExon") "overlapping" will plot all transcripts overlapping the exon, whereas "withExon" will plot all tran- scripts containing the exon.

predictBranchpoints

Value

ggplot2 plot transcript structures

Author(s)

Beth Signal

predictBranchpoints Predict branchpoint probability scores

Description

predicts branchpoint probability scores for each query site.

Usage

```
predictBranchpoints(query, uniqueId = "test", queryType,
workingDirectory = ".", genome = NA, bedtoolsLocation = NA,
BSgenome = NULL, useParallel = FALSE, cores = 1, rmChr = FALSE)
```

Arguments

query	branchpointer query GenomicRanges	
uniqueId	unique string identifier for intermediate .bed and .fa files.	
queryType	type of branchpointer query. "SNP" or "region".	
workingDirecto	ry	
	directory where intermediate .bed and .fa are located	
genome	.fa genome file location	
bedtoolsLocation		
	bedtools binary location (which bedtools)	
BSgenome	BSgenome object	
useParallel	use parallelisation to speed up code?	
cores	number of cores to use in parallelisation (default = 1)	
rmChr	remove "chr" before chromosome names before writing bed file. Required if genome sequence names do not contain "chr"	

Value

GenomicRanges object with branchpoint probaility scores for each site in query

Author(s)

Beth Signal

Examples

```
smallExons <- system.file("extdata","gencode.v26.annotation.small.gtf",
package = "branchpointer")
exons <- gtfToExons(smallExons)
g <- BSgenome.Hsapiens.UCSC.hg38::BSgenome.Hsapiens.UCSC.hg38
querySNPFile <- system.file("extdata","SNP_example.txt", package = "branchpointer")
querySNP <- readQueryFile(querySNPFile,queryType = "SNP",exons = exons, filter = FALSE)
predictionsSNP <- predictBranchpoints(querySNP,queryType = "SNP",BSgenome = g)</pre>
```

predictionsToSummary Convert SNP branchpoint predictions across the branchpoint window to an intronic summary

Description

Takes predictions of branchpoint probabilities from a reference and alternative SNP and summarises the effect(s) of the SNP.

Usage

```
predictionsToSummary(query, predictions, probabilityCutoff = 0.52,
    probabilityChange = 0.15)
```

Arguments

query	query GRanges containing all SNP ids to be summarised	
predictions	site-wide branchpoint proability predictions produced from predictBranchpoints()	
probabilityCuto	off	
	Value to be used as the cutoff for discriminating branchpoint sites from non- branchpoint sites (default = 0.52)	
probabilityChange		
	Minimum probability score change required to call a branchpoint site as deleted or created by a SNP (default = 0.15)	

Value

GRanges with summarised branchpoint changes occuring within the intron due to a SNP.

Author(s)

Beth Signal

Examples

```
smallExons <- system.file("extdata","gencode.v26.annotation.small.gtf", package = "branchpointer")
exons <- gtfToExons(smallExons)
g <- BSgenome.Hsapiens.UCSC.hg38::BSgenome.Hsapiens.UCSC.hg38</pre>
```

```
querySNPFile <- system.file("extdata","SNP_example.txt", package = "branchpointer")
querySNP <- readQueryFile(querySNPFile,queryType = "SNP",exons = exons, filter = FALSE)
predictionsSNP <- predictBranchpoints(querySNP,queryType = "SNP",BSgenome = g)</pre>
```

10

readQueryFile

summarySNP <- predictionsToSummary(querySNP,predictionsSNP)</pre>

readQueryFile Read a query file

Description

Reads and formats a manually generated query file, and finds realtive locations of the closest annotated exons Converts unstranded SNPs to two entries for each strand. Checks for duplicate names and replaces if found.

Usage

```
readQueryFile(queryFile, queryType, exons, maxDist = 50, filter = TRUE)
```

Arguments

queryFile	tab delimited file containing query information. For intronic regions should be in the format: region id, chromosome name, region start, region end, strand. For SNP variants should be in the format: SNP id, chromosome name, SNP position, strand, reference allele (A/T/C/G), alternative allele (A/T/C/G)
queryType	type of query file ("SNP" or "region")
exons	GRanges containing exon co-ordinates. Should be produced by gtfToExons()
maxDist	maximum distance a SNP can be from an annotated 3' exon.
filter	remove SNP queries prior to finding finding nearest exons.

Value

Formatted query GRanges

Author(s)

Beth Signal

Examples

```
smallExons <- system.file("extdata","gencode.v26.annotation.small.gtf", package = "branchpointer")
exons <- gtfToExons(smallExons)</pre>
```

```
querySNPFile <- system.file("extdata","SNP_example.txt", package = "branchpointer")
querySNP <- readQueryFile(querySNPFile, queryType = "SNP", exons)</pre>
```

```
queryIntronFile <- system.file("extdata","intron_example.txt", package = "branchpointer")
queryIntron <- readQueryFile(queryIntronFile,queryType = "region", exons)</pre>
```

Index

* internal exonsToIntrons, 2 getBranchpointSequence, 2 getCanonical3SS, 3 getExonDists, 4 getPPT, 4 getQueryLoc, 5 plotStructure, 8

exonsToIntrons, 2

getBranchpointSequence, 2
getCanonical3SS, 3
getExonDists, 4
getPPT, 4
getQueryLoc, 5
gtfToExons, 5

makeBranchpointWindowForExons, 6
makeBranchpointWindowForSNP, 7

plotBranchpointWindow, 7
plotStructure, 8
predictBranchpoints, 9
predictionsToSummary, 10

readQueryFile, 11