# Package 'VaSP'

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Type Package

Version 1.21.0

- **Title** Quantification and Visualization of Variations of Splicing in Population
- **Description** Discovery of genome-wide variable alternative splicing events from short-read RNA-seq data and visualizations of gene splicing information for publication-quality multi-panel figures in a population. (Warning: The visualizing function is removed due to the dependent package Sushi deprecated. If you want to use it, please change back to an older version.)
- URL https://github.com/yuhuihui2011/VaSP

BugReports https://github.com/yuhuihui2011/VaSP/issues

License GPL (>= 2.0)

**Depends** R (>= 4.0), ballgown

**Imports** IRanges, GenomicRanges, S4Vectors, parallel, matrixStats, GenomicAlignments, GenomeInfoDb, Rsamtools, cluster, stats, graphics, methods

Suggests knitr, rmarkdown

VignetteBuilder knitr

**biocViews** RNASeq, AlternativeSplicing, DifferentialSplicing, StatisticalMethod, Visualization, Preprocessing, Clustering, DifferentialExpression, KEGG, ImmunoOncology

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```
BMfinder
```

Discover bimodal distrubition features

# Description

Find bimodal distrubition features and divide the samples into 2 groups by k-means clustering.

# Usage

# Arguments

х	a numeric matrix with feature rows and sample columns, e.g., splicing score matrix from spliceGenome or spliceGene function.
p.value	p.value threshold for bimodal distrubition test
maf	minor allele frequency threshold in k-means clustering
miss	missing grouping rate threshold in k-means clustering
fold	fold change threshold between the two groups
log	whether the scores are to be logarithmic. If TRUE, all the scores are log2 tran- formed before k-means clustering: $x = log2(x+1)$ .
cores	threads to be used. This value is passed to <b>?mclapply</b> in <b>parallel</b> package

# Details

The matrix contains 1, 2 and NA, and values of 'x' in group 2 are larger than group 1.

# Value

a matrix with feature rows and sample columns.

# Examples

```
data(rice.bg)
score<-spliceGene(rice.bg,'MSTRG.183',junc.type='score')
score<-round(score,2)
as<-BMfinder(score,cores=1) # 4 bimodal distrubition features found
##compare
as
score[rownames(score)%in%rownames(as),]</pre>
```

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getDepth

#### Description

Get read depth from a BAM file (in bedgraph format)

# Usage

```
getDepth(x, chrom, start, end)
```

# Arguments

х	path to a BAM file
chrom	chromosome of a region to be searched
start	start position
end	end position

# Value

a data.frame in bedgraph file format.

# Examples

getGeneinfo Get Gene Informaton from a ballgown object

#### Description

Get gene informaton from a ballgown object by genes or by genomic regions

# Usage

#### Arguments

genes	a character vector specifying gene IDs in 'bg'. Any values other than NA over- ride genomic region (chrom, start, stop)
bg	ballgown object
chrom	chromosome of a region
start	start postion
end	stop postion
samples	names of samples. The transcrpts in these samples are subjected to 'trans.select'
trans.select	logical expression-like string, indicating transcript rows to select from a matrix of transcript coverages: NA value keeps all transcripts.

#### Value

a data.frame in bed-like file format

#### Examples

```
data(rice.bg)
unique(geneIDs(rice.bg))
gene_id <- c('MSTRG.181', 'MSTRG.182', 'MSTRG.183')</pre>
geneinfo <- getGeneinfo(genes=gene_id,rice.bg)</pre>
trans <- table(geneinfo$name) # show how many exons each transcript has</pre>
trans
# library(Sushi)
# chrom = geneinfo$chrom[1]
# chromstart = min(geneinfo$start) - 1e3
# chromend = max(geneinfo$stop) + 1e3
# color = rep(SushiColors(2)(length(trans)), trans)
# par(mar=c(3,1,1,1))
# plotGenes(geneinfo, chrom, chromstart, chromend, col = color, bheight = 0.2,
            bentline = FALSE, plotgenetype = 'arrow', labeloffset = 0.5)
#
# labelgenome(chrom, chromstart , chromend, side = 1, n = 5, scale = 'Kb')
```

rice.bg

Rice ballgown object

#### Description

Small ballgown object created with a subset of rice RNAseq data, for demonstration purposes

#### Format

a ballgown object with 33 transcripts and 6 samples

#### Details

The raw RNA-seq data were screened and trimmed using Trimmomatic (Bolger et al., 2014) and RNA-seq mapping, transcript assembly, and quantification were conducted with HISAT, StringTie, and Ballgown by following the method described by Pertea et al. (Pertea et al., 2016). The rice.bg is a subset ballgown object with 33 transcripts and 6 samples (Yu et al., 2021).

#### spliceGene

#### Source

The raw RNA-seq data were from the project of variation in transcriptional responses to salt stress in rice (SRA Accession: SRP106054)

#### References

Yu, H., Du, Q., Campbell, M., Yu, B., Walia, H. and Zhang, C. (2021), Genome-wide discovery of natural variation in pre-mRNA splicing and prioritising causal alternative splicing to salt stress response in rice. New Phytol. https://doi.org/10.1111/nph.17189

Bolger, A.M., Lohse, M., and Usadel, B. (2014). Trimmomatic: a flexible trimmer for Illumina sequence data. Bioinformatics 30, 2114-2120.

Pertea, M., Kim, D., Pertea, G.M., Leek, J.T., and Salzberg, S.L. (2016). Transcript-level expression analysis of RNA-seq experiments with HISAT, StringTie and Ballgown. Nat Protoc 11, 1650-1667.

#### Examples

data(rice.bg)
rice.bg
# ballgown instance with 33 transcripts and 6 samples

spliceGene

Calculate Splicing Scores for One Gene

# Description

Calculate splicing Scores from ballgown object for a given gene. This function can only calculate one gene. Please use function spliceGenome to obtain genome-wide splicing scores.

#### Usage

#### Arguments

bg	ballgown object
gene	a character string specifying gene id
samples	names of samples
junc.type	type of junction estimate ('score' for junction score; 'count' for junction read count)
trans.select	logical expression-like string, indicating transcript rows to select from a matrix of transcript coverages: NA value keeps all transcripts. e.g. use trans.select='rowMaxs(x)>=1' to filter the transcripts with the maximium coverage among all the samples less than 1.
junc.select	logical expression-like string, indicating junction rows to select from a matrix of junction counts: NA value keeps all junctions. e.g. use junc.select='rowMaxs(x)>=5' to filter the junctions with the maximium read count among all the samples less than 5.

#### Details

score = junction count/gene-level per base read coverage. Row functions for matrices are useful to select transcripts and junctions. See matrixStats package.

#### Value

a matrix of junction scores with intron rows and sample columns.

#### References

Yu, H., Du, Q., Campbell, M., Yu, B., Walia, H. and Zhang, C. (2021), Genome-wide discovery of natural variation in pre-mRNA splicing and prioritising causal alternative splicing to salt stress response in rice. New Phytol. https://doi.org/10.1111/nph.17189

# See Also

spliceGenome, which calculates splicing scores in whole genome.

#### Examples

```
data(rice.bg)
rice.bg
head(geneIDs(rice.bg))
score<-spliceGene(rice.bg,'MSTRG.183',junc.type='score')
count<-spliceGene(rice.bg,'MSTRG.183',junc.type='count')
## compare
tail(score)
tail(count)
## get intron structrue
intron<-structure(rice.bg)$intron
intron[intron$id%in%rownames(score)]</pre>
```

spliceGenome

```
Calculate Genome-wide Splicing Scores
```

### Description

Calculate splicing scores from ballgown objects for all genes.

# Usage

#### spliceGenome

#### Arguments

bg	ballgown object
gene.select	logical expression-like string, indicating genes to select from a matrix of gene- level coverages: NA value keeps all genes. e.g. gene.select = 'rowQuantiles(x,probs = $0.05$ )>=1' keeps the genes with the read coverage greater than or equal to 1 in at least 95 (0.05 quantile). Used to filter low expressed genes.
intron.select	logical expression-like string, indicating introns to select from a matrix of junction counts: NA value keeps all introns. e.g. intron.select = 'rowQuantiles(x,probs = $0.95$ )>=5' keeps the introns with the read count greater than or euqal to 5 in at least 5 (0.95 quantile). Used to filter introns with very few junction reads supporting.

#### Details

score = junction count/gene-level per base read coverage. Row functions for matrixStats package are useful to select genes and introns.

# Value

a list of two elelments: 'score' is matrix of intron splicing scores with intron rows and sample columns and 'intron' is a GRanges object of intron structure. See structure in **ballgown** package

#### References

Yu, H., Du, Q., Campbell, M., Yu, B., Walia, H. and Zhang, C. (2021), Genome-wide discovery of natural variation in pre-mRNA splicing and prioritising causal alternative splicing to salt stress response in rice. New Phytol. https://doi.org/10.1111/nph.17189

#### See Also

spliceGene, which calculates splicing scores in one gene.

#### Examples

```
data(rice.bg)
rice.bg
```

```
splice<-spliceGenome(rice.bg,gene.select=NA,intron.select=NA)
names(splice)</pre>
```

head(splice\$score)
splice\$intron

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