Package 'bambu'

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Type Package

Title Context-Aware Transcript Quantification from Long Read RNA-Seq data

Version 3.6.0

Description bambu is a R package for multi-sample transcript discovery and quantification using long read RNA-Seq data. You can use bambu after read alignment to obtain expression estimates for known and novel transcripts and genes. The output from bambu can directly be used for visualisation and downstream analysis such as differential gene expression or transcript usage.

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Enhances parallel

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FeatureExtraction, GeneExpression, GenomeAnnotation, GenomeAssembly, ImmunoOncology, LongRead, MultipleComparison,

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Transcription, Transcriptomics

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URL https://github.com/GoekeLab/bambu

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Description

This function takes bam file of genomic alignments and performs isoform recontruction and gene and transcript expression quantification. It also allows saving of read class files of alignments, extending provided annotations, and quantification based on extended annotations. When multiple samples are provided, extended annotations will be combined across samples to allow comparison.

Usage

```
bambu(
  reads,
  annotations = NULL,
  genome = NULL,
 NDR = NULL,
  opt.discovery = NULL,
  opt.em = NULL,
  rcOutDir = NULL,
  discovery = TRUE,
  quant = TRUE,
  stranded = FALSE,
  ncore = 1,
  yieldSize = NULL,
  trackReads = FALSE,
  returnDistTable = FALSE,
  lowMemory = FALSE,
  fusionMode = FALSE,
  verbose = FALSE
)
```

Arguments

reads A string or a vector of strings specifying the paths of bam files for genomic

alignments, or a BamFile object or a BamFileList object (see Rsamtools). Alternatively string or a vector of strings specifying the read class files that are

saved during previous run of bambu.

annotations A path to a .gtf file or a TxDb object or a GRangesList object obtained by

prepareAnnotations.

genome A path to a fasta file or a BSGenome object.

NDR specifying the maximum NDR rate to novel transcript output from detected read

classes, defaults to an automatic recommendation

opt.discovery A list of controlling parameters for isoform reconstruction process:

remove.subsetTx indicating whether filter to remove read classes which are a subset of known transcripts(), defaults to TRUE

min.readCount specifying minimum read count to consider a read class valid in a sample, defaults to 2

min.readFractionByGene specifying minimum relative read count per gene, highly expressed genes will have many high read count low relative abundance transcripts that can be filtered, defaults to 0.05

min.sampleNumber specifying minimum sample number with minimum read count, defaults to 1

min.exonDistance specifying minum distance to known transcript to be considered valid as new, defaults to 35bp

min.exonOverlap specifying minimum number of bases shared with annotation to be assigned to the same gene id, defaults to 10bp

min.primarySecondaryDist specifying the minimum number of distance threshold, defaults to 5bp

min.primarySecondaryDistStartEnd1 specifying the minimum number of distance threshold, used for extending annotation, defaults to 5bp

min.primarySecondaryDistStartEnd2 specifying the minimum number of distance threshold, used for estimating distance to annotation, defaults to 5bp

min.txScore.multiExon specifying the minimum transcript level threshold for multi-exon transcripts during sample combining, defaults to 0

min.txScore.singleExon specifying the minimum transcript level threshold for single-exon transcripts during sample combining, defaults to 1

fitReadClassModel A boolean specifying if Bambu should attempt to train a transcript discovery model for all samples. Defaults to TRUE

defaultModels A model object obtained by codetrainBambu or when return-Model is TRUE

returnModel A boolean specifying if the trained model is output with the readclass files. Defaults to FALSE

baselineFDR A number between 0 - 1, specifying the false discovery rate used during NDR recomendation. Defaults to 0.1

min.readFractionByEqClass indicating the minimum relative read count of a subset transcript compared to all superset transcripts (ie the relative read count within the minimum equivalent class). This filter is applied on the set of annotations across all samples using the total read count, this is not a per-sample filter. Please use with caution. defaults to 0

prefix specifying prefix for new gene Ids (genePrefix.number), defaults to "Bambu"

opt.em A list of controlling parameters for quantification algorithm estimation process:

maxiter specifying maximum number of run iterations, defaults to 10000

degradationBias correcting for degradation bias, defaults to TRUE

conv specifying the covergence threshold control, defaults to 0.0001

minvalue specifying the minvalue for convergence consideration, defaults to 0.00000001

sig.digit specifying the maximum significant digits of the reported estimates

rcOutDir A string variable specifying the path to where read class files will be saved.

discovery A logical variable indicating whether annotations are to be extended. Defaults

to TRUE

quant A logical variable indicating whether quantification will be performed. If false

the output type will change. Defaults to TRUE

stranded A boolean for strandedness, defaults to FALSE.

ncore specifying number of cores used when parallel processing is used, defaults to 1.

yieldSize see Rsamtools.

trackReads When TRUE read names will be tracked and output as metadata in the final out-

put as readToTranscriptMaps detailing. the assignment of reads to transcripts.

The output is a list with an entry for each sample.

returnDistTable

When TRUE the calculated distance table between read classes and annotations will be output as metadata as distTables. The output is a list with an entry for each sample.

lowMemory Read classes will be processed by chromosomes when lowMemory is specified.

This option provides an efficient way to process big samples.

fusionMode A logical variable indicating whether run in fusion mode

verbose A logical variable indicating whether processing messages will be printed.

Details

Main function

Value

bambu will output different results depending on whether *quant* mode is on. By default, *quant* is set to TRUE, so bambu will generate a *SummarizedExperiment* object that contains the transcript expression estimates. Transcript expression estimates can be accessed by *counts()*, including the following variables

counts expression estimates

CPM sequencing depth normalized estimates

fullLengthCounts estimates of read counts mapped as full length reads for each transcript **uniqueCounts** counts of reads that are uniquely mapped to each transcript

Output annotations that are usually the annotations with/without novel transcripts/genes added, depending on whether *discovery* mode is on can be accessed by *rowRanges()* Transcript to gene map can be accessed by *rowData()*, with *eqClass* that defining equivalent class for each transcript

In the case when *quant* is set to FALSE, i.e., only transcript discovery is performed, bambu will report the *grangeslist* of the extended annotations.

Examples

6 compareTranscripts

compareTranscripts compare alternatively-spliced transcripts

Description

compare alternatively-spliced transcripts

Usage

```
compareTranscripts(query, subject)
```

Arguments

query a GRangesList of transcripts subject a GRangesList of transcripts

Details

This function compares two alternatively-spliced transcripts and returns a tibble object that determines the type of the alternative splicing between the query and the subject transcript. Alternative splicing includes exons skipping, intron retention, alternative 5' exon start site, alternative 3' exon end site, alternative first exon, alternative last exon, alternative transcription start site (TSS), alternative transcription end site (TES), internal first exon, internal last exon, or a mixed combination of them.

Value

a tibble object with the following columns:

Remark: two exons, one each from query and subject transcript are defined to be equivalent if one of the exons fully covers the another exon.

- alternativeFirstExon: FALSE if query transcript and subject transcript have equivalent first exon. TRUE otherwise.
- alternativeTSS: +k if the query initiates the transcription k sites earlier than the subject transcript, -k if the query initiates the transcription k sites later than the subject transcript.
- internalFirstExon.query: TRUE if the first exon of the query transcript is equivalent to one of the exon of the subject transcript (except the first exon). FALSE otherwise.
- internalFirstExon.subject: TRUE if the first exon of the subject transcript is equivalent to one of the exon of the query transcript (except the first exon). FALSE otherwise.
- alternativeLastExon: FALSE if query transcript and subject transcript have equivalent last exon. TRUE otherwise.
- alternativeTES: +k if the query transcript ends the transcription k sites later than the subject transcript, -k if the query transcript ends the transcription k sites earlier than the subject transcript.

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• internalLastExon.query: TRUE if the last exon of the query transcript is equivalent to one of the exon in the subject transcript (except the last exon). FALSE otherwise.

- internalLastExon.subject: TRUE if the last exon of the subject transcript is equivalent to one of the exon in the query transcript (except the last exon). FALSE otherwise.
- intronRetention.subject: k if there is/are k intron(s) in the subject transcript relative to the query transcript.
- intronRetention.query: k if there is/are k intron(s) in the query transcript relative to the subject transcript.
- exonSkipping.query: k if there is/are k exon(s) in the subject transcript (except the first and last) not in the query transcript.
- exonSkipping.subject: k if there is/are k exon(s) in the query transcript (except the first and last) not in the subject transcript.
- exon5prime (splicing): k if there is/are k equivalent exon(s) having different 5' start site (except the 5' start site of first exon).
- exon3prime (splicing): k if there is/are k equivalent exon(s) having different 3' end site (except the 3' end site of the last exon).

See Also

Value for more details about each of the alternative splicing events.

Examples

plotBambu

plot.bambu

Description

plotSEOuptut

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Usage

```
plotBambu(
    se,
    group.variable = NULL,
    type = c("annotation", "pca", "heatmap"),
    gene_id = NULL,
    transcript_id = NULL
)
```

Arguments

An summarized experiment object obtained from bambu or transcriptToGeneExpression.

Wariable for grouping in plot, has be to provided if choosing to plot PCA.

type plot type variable, a values of annotation for a single gene with heatmap for isoform expressions, pca, or heatmap, see details.

gene_id specifying the gene_id for plotting gene annotation, either gene_id or transcript_id has to be provided when type = "annotation".

transcript_id has to be provided when type = "annotation, either gene_id or transcript_id has to be provided when type = "annotation"

Details

type indicates the type of plots to be plotted. There are two types of plots can be chosen, PCA or heatmap.

Value

A heatmap plot for all samples

Examples

```
se <- readRDS(system.file("extdata",
"seOutputCombined_SGNex_A549_directRNA_replicate5_run1_chr9_1_1000000.rds",
package = "bambu"))
plotBambu(se, type = "PCA")</pre>
```

prepareAnnotations

prepare reference annotations for long read RNA-Seq analysis with Bambu

Description

prepare reference annotations for long read RNA-Seq analysis with Bambu

Usage

```
prepareAnnotations(x)
```

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Arguments

Х

A path to gtf file or a TxDb object.

Details

This function creates a reference annotation object which is used for transcript discovery and quantification in Bambu. prepareAnnotations can use a path to a gtf file or a TxDB object as input, and returns a annotation object that stores additional information about transcripts which is used in Bambu. For each transcript, exons are ranked from first to last exon in direction of transcription.

Value

A GRangesList object with additional details for each exon and transcript that are required by Bambu. Exons are ranked by the exon_rank column, corresponding to the rank in direction of transcription (from first to last exon). In addition to exon rank, gene id, transcript id, and the minimum transcript equivalent class is stored as well (a transcript equivalence class of a transcript x is the collection of transcripts where their exon junctions contain, in a continuous way, the exon junctions of the transcript x). The object is designed to be used by Bambu, and the direct access of the metadata is not recommended.

See Also

[bambu()] for transcript discovery and quantification from long read RNA-Seq.

Examples

```
gtf.file <- system.file("extdata",
    "Homo_sapiens.GRCh38.91_chr9_1_1000000.gtf",
    package = "bambu"
)
annotations <- prepareAnnotations(x = gtf.file)

# run bambu
test.bam <- system.file("extdata",
    "SGNex_A549_directRNA_replicate5_run1_chr9_1_1000000.bam",
    package = "bambu")
fa.file <- system.file("extdata",
    "Homo_sapiens.GRCh38.dna_sm.primary_assembly_chr9_1_1000000.fa",
    package = "bambu")
se <- bambu(reads = test.bam, annotations = annotations,
    genome = fa.file, discovery = TRUE, quant = TRUE)</pre>
```

readFromGTF

convert a GTF file into a GRangesList

Description

Outputs GRangesList object from reading a GTF file

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Usage

```
readFromGTF(file, keep.extra.columns = NULL)
```

Arguments

Value

grlist a GRangesList object, with two columns

TXNAME specifying prefix for new gene Ids (genePrefix.number), defaults to empty

GENEID indicating whether filter to remove read classes which are a subset of known transcripts(), defaults to TRUE

Examples

```
gtf.file <- system.file("extdata",
    "Homo_sapiens.GRCh38.91_chr9_1_1000000.gtf",
    package = "bambu"
)
readFromGTF(gtf.file)</pre>
```

trainBambu

Function to train a model for use on other data

Description

This function train a model for use on other data

Usage

```
trainBambu(
  rcFile = NULL,
  min.readCount = 2,
  nrounds = 50,
  NDR.threshold = 0.1,
  verbose = TRUE
)
```

Arguments

rcFile summerized experiment object with read classes/ranges produced from bambu(quant

= FALSE, discovery = FALSE) or rcOutdir

min.readCount the minimum number of reads a read class is required to be have to be used for

training

nrounds xgboost hyperparameter - the number of decision trees in the final mode

NDR. threshold the effective NDR threshold that bambu will try and match on other samples

when using this model

verbose if additional messages should be output Output - A list containing 6 objects

which is passed directly into bambu(opt.discovery=list(defaultModels=trainBambu())) transcriptModelME - the model for multi-exon transcripts transcriptModelSE - the model for single-exon transcripts txScoreBaseline - the txScore used for NDR calibration for multi-exon transcripts txScoreBaselineSE - [DEPRECATED] the txScore used for NDR calibration for single-exon transcripts lmNDR = lm-NDR - the linear model of the reletionship between txScore and NDR used to calculate the baseline for multi-exon transcripts lmNDR.SE = lmNDR.SE - the linear model of the reletionship between txScore and NDR used to calculate the baseline for single-exon transcripts NDR.threshold - the NDR threshold used to

calculate the txScoreBaseline on the lmNDR (baselineFDR)

Details

Function to train a model for use on other data

Value

It returns a model object to use in bambu

transcriptToGeneExpression

transcript to gene expression

Description

Reduce transcript expression to gene expression

Usage

transcriptToGeneExpression(se)

Arguments

se a summarizedExperiment object from bambu

Value

A SummarizedExperiment object

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Examples

writeBambuOutput

Write Bambu results to GTF and transcript/gene-count files

Description

Write Bambu results to GTF and transcript/gene-count files

Usage

```
writeBambuOutput(se, path, prefix = "")
```

Arguments

se a SummarizedExperiment object from bambu.

path the destination of the output files (gtf, transcript counts, and gene counts)

prefix the prefix of the output files

Details

The function will write the output from Bambu to files. The annotations will be written to a .gtf file, transcript counts (total counts, CPM, full-length counts, partial-length counts, and unique counts) and gene counts will be written to .txt files.

Examples

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writeToGTF	write GRangeslist into GTF file	

Description

Write annotation GRangesList into a GTF file

Usage

```
writeToGTF(annotation, file, geneIDs = NULL)
```

Arguments

annotation a GRangesList object file the output gtf file name

geneIDs an optional dataframe of geneIDs (column 2) with the corresponding transcrip-

tIDs (column 1)

Value

```
gtf a GTF dataframe
```

Examples

```
outputGtfFile <- tempfile()
gr <- readRDS(system.file("extdata",
          "annotationGranges_txdbGrch38_91_chr9_1_1000000.rds",
          package = "bambu"
))
writeToGTF(gr, outputGtfFile)</pre>
```

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