Package ‘RIPSeeker’

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Type Package

Title RIPSeeker: a statistical package for identifying protein-associated transcripts from RIP-seq experiments

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Author Yue Li

Maintainer Yue Li <yueli@cs.toronto.edu>

Description Infer and discriminate RIP peaks from RIP-seq alignments using two-state HMM with negative binomial emission probability. While RIPSeeker is specifically tailored for RIP-seq data analysis, it also provides a suite of bioinformatics tools integrated within this self-contained software package comprehensively addressing issues ranging from post-alignments processing to visualization and annotation.

Depends R (>= 2.15), methods, S4Vectors (>= 0.9.25), IRanges, GenomicRanges, SummarizedExperiment, Rsamtools, GenomicAlignments, rtracklayer

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License GPL-2

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RIPSeeker-package .................................................. 2
addDummyProb ...................................................... 3
addPseudoAlignment .............................................. 5
annotateRIP ......................................................... 6
binCount .......................................................... 9
combineAlignGals ............................................... 10
combineRIP ........................................................ 11
computeLogOdd ................................................... 13
computeRPKM ..................................................... 14
RIPSeeker-package

RIPSeeker: a statistical package for identifying protein-associated transcripts from RIP-seq experiments

Description

RIPSeeker infers and discriminates RIP peaks from RIP-seq alignments using two-state HMM with negative binomial emission probability. While RIPSeeker is specifically tailored for RIP-seq data analysis, it also provides a suite of bioinformatics tools integrated within this self-contained software package comprehensively addressing issues ranging from post-alignments processing to visualization and annotation. In addition, a rule-based approach is provided as an additional function named `rulebaseRIPSeek` for user to obtain RPKM/FPKM (and fold-change) for the gene/transcripts expressions in RIP (and control) based on automatically retrieved online Ensembl annotation given single or paired-end alignments.
Details

Package: RIPSeeker
Type: Package
Version: 1.4.0
Date: 2012-11-06
License: GPL-2

The front-end main function `ripSeek` suffices for most applications. The function takes as the only required argument the path to alignment files (BAM/BED/SAM) and outputs predicted RIP regions. Optionally, user may indicate via `cNAME` which file(s) in the first file argument list is/are control to enable empirical false discover rate (eFDR) computation. If the arguments `biomaRt_dataset` and/or `goAnno` are set, ripSeek will return the annotated RIP predictions and the enriched GO terms corresponding to the genomic context of the RIP predictions. User can also specify the thresholds for statistical significance scores via `logOddCutoff`, `pvalCutoff`, `pvalAdjCutoff`, `eFDRcutoff`.

Author(s)

Yue Li <yueli@cs.toronto.edu>

References


See Also

ripSeek, rulebaseRIPSeek

Examples

library(RIPSeeker)
ls("package:RIPSeeker")

---

`addDummyProb` Create a dummy GRanges object as a placeholder in case nbh_em fails (Internal function)

Description

This function is used to generate a place holder in cases the EM fails to converge on a chromosome due to too few number of reads mapped to that chromosome. This is an internal function not expected to be directly called by the user.
addDummyProb

**Usage**

```r
call = addDummyProb(alignGR, K = 2, randomProb = FALSE, runViterbi = FALSE, ...)
```

**Arguments**

- `alignGR`  
  GRanges object derived from RIP-seq alignment inputs.
- `K`  
  Number of hidden states (Default: 2).
- `randomProb`  
  A binary value to indicate whether to use random probability as a place holder to present posterior probabilities. If set FALSE, (by default), equal probability is used for all states.
- `runViterbi`  
  A binary value to indicate whether to generate place holder for the Viterbi state sequence (Default: FALSE).
- `...`  
  Additional arguments expected to contain the bin size used for computing the bin counts in `binCount` function, and any other extra arguments are ignored.

**Details**

A privatie function to fall back in case HMM fails to converge mostly due to too many zero counts in the input vector. When that occurs, a GRanges place holder object needs to be returned to keep consistent with the remaining GRanges for each chromosome. Thus, all information slot will be generated as place holder to properly create the GRangesList for the predictions on all chromosomes (each as an GRanges item in the list).

**Value**

- `GRanges`  
  A GRanges object containing the read count (in the defined bin size), alpha, beta, TRANS dummy values for the HMM

**Author(s)**

Yue Li

**See Also**

- `mainSeekSingleChrom`

**Examples**

```r
# Retrieve system files
extdata.dir <- system.file("extdata", package="RIPSeeker")
bamFiles <- list.files(extdata.dir, ".bam$", recursive=TRUE, full.names=TRUE)
bamFiles <- grep("PRC2", bamFiles, value=TRUE)
alignGal <- getAlignGal(bamFiles[1], reverseComplement=TRUE, genomeBuild="mm9")
alignGR <- as(alignGal, "GRanges")
alignGRLList <- GRangesList(as.list(split(alignGR, seqnames(alignGR))))
x <- addDummyProb(alignGRLList$chrX, binSize=10000)
```
addPseudoAlignment

Add a psuedoalignment as a placeholder for the chromosome (Internal function)

Description
Check whether chromosome has at least one alignment to prevent abnormal behaviour of the subsequent functions. In case no alignment is found on an entire chromosome, add a pseudo-alignment as a placeholder for that chromosome.

Usage
addPseudoAlignment(alignGR)

Arguments
alignGR GRanges object containing the alignment information.

Details
In case no alignment is found on an entire chromosome, add an alignment with start 1 and end 20 as a placeholder for the chromosome. This step is necessary to maintain the chromosome information.

Value
alignGR Original or augmented input GRanges object with pseudoreads, depending on whether there exists empty chromosome(s).

Author(s)
Yue Li

See Also
combineAlignGals, readGAlignments, readGAlignmentPairs, import

Examples
# Retrieve system files
extdata.dir <- system.file("extdata", package="RIPSeeker")
bamFiles <- list.files(extdata.dir, ".bam$", recursive=TRUE, full.names=TRUE)
bamFiles <- grep("PRC2", bamFiles, value=TRUE)
alignGal <- getAlignGal(bamFiles[1], reverseComplement=TRUE, genomeBuild="mm9")
alignGR <- as(alignGal, "GRanges")
alignGR
annotateRIP <- addPseudoAlignment(alignGR)

---

**annotateRIP**

**Annotate RIP peaks with genomic information and perform GO enrichment**

**Description**

Given the genomic coordinates of each predicted RIP regions, query the Ensembl database whether each region is nearby or overlaps any known (noncoding) genes.

**Usage**

annotateRIP(sigGRanges, biomaRt_dataset, featureType = "TSS", goAnno, strandSpecific = FALSE, exportFormat = "txt", hasGOdb = !missing(goAnno), goPval = 0.1, outDir, ...)

**Arguments**

- **sigGRanges:** GRanges object indicating the chromosomal coordinates of each RIP peaks.
- **biomaRt_dataset:** Ensembl dataset available from biomaRt (See `listDatasets`). For instance, the human and mouse annotations are `hsapiens_gene_ensembl` and `mmusculus_gene_ensembl`, respectively.
- **featureType:** TSS, miRNA, Exon, 5'UTR, 3'UTR, transcript or Exon plus UTR defined in `getAnnotation`.
- **goAnno:** Optional argument that specifies a GO dataset used for GO enrichment analysis performed by `getEnrichedGO`. For instance, the human and mouse GO datasets are `org.Hs.eg.db` and `org.Mm.eg.db`.
- **strandSpecific:** Indicate whether the annotations should be strand-specific (Default: FALSE)
- **exportFormat:** Format to export using `exportGRanges` (Default: "txt", i.e. tab-delim file).
- **hasGOdb:** A binary flag that indicates whether GO enrichment is performed in order to export the results. hasGOdb can be FALSE either because goAnno is not specify or because the GO database does not exist.
- **goPval:** P-value cutoff to determine the significance of enriched GO terms by `getEnrichedGO`.
- **outDir:** Output directory.
- **...** Extra arguments passed to `useMart` to specify the database and to passed `getEnrichedGO` to specify the GO enrichment procedure.
Details

To access the up-to-date Ensembl database, RIPSeeker employs `useMart` and `getAnnotation` from biomaRt and ChIPpeakAnno Bioconductor packages to dynamically establish internet connection to the database and retrieve the up-to-date annotations. Then, `annotatePeakInBatch` from ChIPpeakAnno is used to efficiently annotate all of the predicted regions based on the Ensembl annotation. A predicted region may overlap multiple genes, all of which will be reported as separate records. Moreover, `getEnrichedGO` from ChIPpeakAnno is applied to the annotated predictions to discover enriched Gene Ontology (GO) terms involving the protein-associated transcriptome.

In order to use old annotation (e.g., mm9 v.s. mm10), user also needs to specify the host and biomart arguments accepted within `useMart`. To access to mouse annotation from Ensembl version 65, for instance, user needs to call `annotateRIP(..., dataset="mmusculus_gene_ensembl", biomart="ENSEMBL_MART_ENSEMBL", host="dec2011.archive.ensembl.org", ...)`, which will run `useMart(dataset="mmusculus_gene_ensembl", biomart="ENSEMBL_MART_ENSEMBL", host="dec2011.archive.ensembl.org", ...)` to get the mm9 annotation from Ensembl (v65).

Value

- `sigGRangesAnnotated`: `sigGRanges` augmented with genomic information including "ensembl_gene_id", "external_gene_id", and "description"
- `enrichedGO`: Output from `getEnrichedGO`. All three main GO categories ("Biological Process", "Molecular Function", "Cellular Component") are combined together and returned. The argument is only returned when `hasGOdb` is TRUE.

If `outDir` is specified, then the above `sigGRangesAnnotated` is saved as `RIPregions_annotated.txt` and `RIPregions_annotated.RData`, and `enrichedGO` as `RIPregions_enrichedGO.txt` in the `outDir` directory.

Author(s)

Yue Li

References


Lihua Julie Zhu, Herve Pages, Claude Gazin, Nathan Lawson, Jianhong Ou, Simon Lin, David Lapointe and Michael Green (2012). ChIPpeakAnno: Batch annotation of the peaks identified from either ChIP-seq, ChIP-chip experiments or any experiments resulted in large number of chromosome ranges.. R package version 2.4.0.

See Also

`useMart`, `getAnnotation`, `getEnrichedGO`
annotateRIP

Examples

```
if(interactive()) { # need internet connection
# Retrieve system files
extdata.dir <- system.file("extdata", package="RIPSeeker")

bamFiles <- list.files(extdata.dir, ".bam", recursive=TRUE, full.names=TRUE)

bamFiles <- grep("PRC2", bamFiles, value=TRUE)

# Parameters setting
binSize <- NULL # automatically determine bin size
minBinSize <- 10000 # min bin size in automatic bin size selection
maxBinSize <- 12000 # max bin size in automatic bin size selection
multicore <- FALSE # use multicore
strandType <- "-" # set strand type to minus strand

biomart <- "ENSEMBL_MART_ENSEMBL" # use archive to get ensembl 65
dataset <- "mmusculus_gene_ensembl" # mouse dataset id name
host <- "dec2011.archive.ensembl.org" # use ensembl 65 for annotation
goAnno <- "org.Mm.eg.db"

# use defined binSize from RIP
RIPBinSize <- lapply(mainSeekOutputRIP$nbhGRList, function(x) median(width(x)))

mainSeekOutputRIP <- mainSeek(bamFiles=grep(pattern="SRR039214", bamFiles, value=TRUE, invert=TRUE),
binSize=binSize, minBinSize = minBinSize,
maxBinSize = maxBinSize, strandType=strandType,
reverseComplement=TRUE, genomeBuild="mm9",
uniqueHit = TRUE, assignMultihits = TRUE,
rerunWithDisambiguatedMultihits = TRUE,
multicore=multicore, silentMain=FALSE, verbose=TRUE)

mainSeekOutputCTL <- mainSeek(bamFiles=grep(pattern="SRR039214", bamFiles, value=TRUE, invert=FALSE),
binSize=RIPBinSize, strandType=strandType,
reverseComplement=TRUE, genomeBuild="mm9",
uniqueHit = TRUE, assignMultihits = TRUE,
rerunWithDisambiguatedMultihits = TRUE,
multicore=multicore, silentMain=FALSE, verbose=TRUE)

# significance test on Viterbi predicted peaks
ripGR <- seekRIP(mainSeekOutputRIP$nbhGRList$chrX, mainSeekOutputCTL$nbhGRList)

# Annotate peaks
annotatedRIPGR <- annotateRIP(sigGRanges = ripGR,
biomaRt_dataset = dataset, goAnno = goAnno,
strandSpecific = !is.null(strandType),
host=host, biomart=biomart)
```
**binCount**

```
head(annotatedRIPGR$sigGRangesAnnotated)
```

---

**binCount**  
*Count reads in nonoverlapping bins across a chromosome*

---

**Description**

Stratify chromosome into nonoverlapping bins of the same size and count the number of reads that fall within each bin.

**Usage**

```
binCount(alignGR, binSize, returnBinCountOnly = FALSE)
```

**Arguments**

- `alignGR`: GRanges object containing the alignments for a single chromosome.
- `binSize`: An integer for the bin size.
- `returnBinCountOnly`: Binary indicator. If TRUE, only the integer read count is returned; if FALSE, GRanges of bins with value slot saved as the corresponding read counts is returned.

**Details**

The function is designed to operate to a single chromosome to facilitate parallel computing on multiple chromosomes independently. The function is used in `evalBinSize` to select the optimal bin size based on the read counts and in `mainSeekSingleChrom` to provide the read count as input for the HMM.

**Value**

If `returnBinCountOnly` is TRUE, then the integer read count is returned; if `returnBinCountOnly` is FALSE, then the GRanges of bins with value slot saved for the corresponding read counts is returned.

**Author(s)**

Yue Li

**References**


**See Also**

- `selectBinSize`
- `evalBinSize`
# Retrieve system files
extdata.dir <- system.file("extdata", package="RIPSeeker")

bamFiles <- list.files(extdata.dir, ".bam", recursive=TRUE, full.names=TRUE)
bamFiles <- grep("PRC2", bamFiles, value=TRUE)

alignGal <- getAlignGal(bamFiles[1], reverseComplement=TRUE, genomeBuild="mm9")

alignGR <- as(alignGal, "GRanges")

alignGRList <- GRangesList(as.list(split(alignGR, seqnames(alignGR))))

binSize <- 1000

binGR <- binCount(alignGRList$chrX, binSize)

## Examples

# Retrieve system files
extdata.dir <- system.file("extdata", package="RIPSeeker")

bamFiles <- list.files(extdata.dir, ".bam", recursive=TRUE, full.names=TRUE)
bamFiles <- grep("PRC2", bamFiles, value=TRUE)

alignGal <- getAlignGal(bamFiles[1], reverseComplement=TRUE, genomeBuild="mm9")

alignGR <- as(alignGal, "GRanges")

alignGRList <- GRangesList(as.list(split(alignGR, seqnames(alignGR))))

binSize <- 1000

binGR <- binCount(alignGRList$chrX, binSize)

## Description

Import and process individual BAM/SAM/BED alignment files using `getAlignGal` and combine them into a single GAlignments object.

## Usage

combineAlignGals(bamFiles, ...)

## Arguments

- `bamFiles`: A list of paths to the alignment files.
- `...`: Arguments passed to `getAlignGal`.

## Details

If there is only one BAM file, then simply return the output from `getAlignGal`; otherwise, all processed alignments are pooled to form a single GAlignments object.

## Value

- `combinedGal`: GAlignments object containing the (combined) processed alignments with the values slot saved for the "uniqueHits" binary flag defined in `getAlignGal` and metadata saved as a list containing argument setting for `reverseComplement`, `returnDuplicate`, `flagMultiHits`, `returnOnlyUniqueHits` defined in `getAlignGal`.
Note

User are recommended to pool technical replicates but keep biological replicate separate for confirmation.

Author(s)

Yue Li

See Also

getAlignGal, readGAlignments, readGAlignmentPairs, import

Examples

# Retrieve system files
extdata.dir <- system.file("extdata", package="RIPSeeker")

bamFiles <- list.files(extdata.dir, ".bam$", recursive=TRUE, full.names=TRUE)

bamFiles <- grep("PRC2", bamFiles, value=TRUE)

# combine the alignments for technical replicates
alignGal <-
combineAlignGals(bamFiles=grep(pattern="SRR039214", bamFiles, value=TRUE, invert=TRUE),
reverseComplement=TRUE, genomeBuild="mm9")

combineRIP

Combined predictions from (presumably) biological replicates.

Description

A simple helper function that combines multiple prediction lists from biological replicates into a single list.

Usage

combineRIP(ripPath, pattern="gff3$",
combineOption="intersect",
 pvalCutoff=1, pvalAdjCutoff=1, eFDR_cutoff=1,
 logOddCutoff=-Inf, maxgap=1e3, minIntersect, genomeBuild)

Arguments

ripPath

Pattern for the names of the prediction files to combine. The file names are expected to have a common extension such as "bed", "gff3", "gtf", but this is not enforced. Default: "gff3$" (i.e. the default output RIPregions.gff3 from ripSeek).
combineOption  Options on how to combine the peaks including:
"intersect": is selected (default), only peaks in each biological replicate list that overlap with or are adjacent within maxgap nucleotides to at least minIntersect other replicates will be kept. If minIntersect is unspecified, then only the peaks that consistently predicted in all replicates are kept.
"merge": All overlapping peaks from the replicates will be merged into one peak.
"union": All overlapping peaks from the replicates will be merged into one peak.

pvalCutoff  Threshold for the p-value cutoff. Only peaks with p-value less than the logOddCutoff will be reported. Default: 1 (i.e. no cutoff).

pvalAdjCutoff  Threshold for the adjusted p-value cutoff. Only peaks with adjusted p-value less than the logOddCutoff will be reported. Default: 1 (i.e. no cutoff).

efDRcutoff  Threshold for the empirical false discovery rate (eFDR). Only peaks with eFDR less than the eFDRcutoff will be reported. Default: 1 (i.e. no cutoff).

logOddCutoff  Threshold for the log odd ratio of posterior for the RIP over the background states (See seekRIP). Only peaks with logOdd score greater than the logOddCutoff will be reported. Default: -Inf (i.e. no cutoff).

maxgap  Maximum gap allowed to determine two peaks agree with each other.

minIntersect  Minimum number of replicates required to have peaks either intersect or are adjacent to the peak in other replicate.

genoBuild  Genome build used to obtain the chromosome information from online UCSC database to assign chromosome length to the GRanges object created as the combined peak list.

Value

gr  GRanges object containing chromosome locations of the combined peaks.

Note

Please run ripSeek first on all biological replicates and renamed each "RIPregions.gff3" output to correspond to different biological replicates and place all of the files into a single folder. The path of this folder can then be used as the input argument for ripPath.

Author(s)

Yue Li

References


Michael Lawrence, Vince Carey and Robert Gentleman. rtracklayer: R interface to genome browsers and their annotation tracks. R package version 1.16.3.

See Also

combineAlignGals, ripSeek, import, import, reduce, countOverlaps
computeLogOdd

Examples

# Retrieve system files
ripath <- system.file("extdata/RIPregions", package="RIPSeeker")
gr1 <- combineRIP(ripath, combineOption="intersect", genomeBuild="mm9")
gr2 <- combineRIP(ripath, combineOption="merge", genomeBuild="mm9")
gr3 <- combineRIP(ripath, combineOption="union", genomeBuild="mm9")
length(gr1)
length(gr2)
length(gr3)

computeLogOdd(nbhGR)  
Compute the log odd ratio of RIP over background.

Description

The RIPScore is computed as the log odd ratio of the posterior for the RIP state \( z_i = 2 \) over the posterior for the background state \( z_i = 1 \).

Usage

computeLogOdd(nbhGR)

Arguments

nbhGR  
GRanges of bins with the value slot saved for the posterior probabilities for the background and RIP state.

Details

To assess the statistical significance of the RIP predictions, we assign each bin a RIPScore defined as the log odd ratio of the posterior for the RIP state \( z_i = 2 \) over the posterior for the background state \( z_i = 1 \). When control is available, the RIPScore is updated as the difference between the RIPScores evaluated separately for RIP and control libraries. The scoring system captures the model confidence for the RIP state of each bin in the RIP library penalized by the false confidence for the RIP state of the same bin in the control library. In addition, RIPScore obviates scaling of read counts. Since sequencing depth usually differs between RIP and control libraries, scaling is necessary if the statistical score were derived from the read count differences. On the other hand, simplistic linear scaling may distort the data.

Value

A vector of log odd scores for each bin in nbhGR.
computeRPKM

## Author(s)

Yue Li

## See Also

seekRIP, scoreMergedBins, logScoreWithoutControl, logScoreWithControl

## Examples

```r
# Retrieve system files
extdata.dir <- system.file("extdata", package="RIPSeeker")

bamFiles <- list.files(extdata.dir, ".bam$", recursive=TRUE, full.names=TRUE)
bamFiles <- grep("PRC2", bamFiles, value=TRUE)

# Retrieve system files
extdata.dir <- system.file("extdata", package="RIPSeeker")

bamFiles <- list.files(extdata.dir, ".bam$", recursive=TRUE, full.names=TRUE)
bamFiles <- grep("PRC2", bamFiles, value=TRUE)

alignGal <- getAlignGal(bamFiles[1], reverseComplement=TRUE, genomeBuild="mm9")

alignGR <- as(alignGal, "GRanges")

alignGRLList <- GRangesList(as.list(split(alignGR, seqnames(alignGR))))

## run main function for HMM inference on a single chromosome

nbhGR <- mainSeekSingleChrom(alignGR=alignGRLList$chrX, K = 2, binSize=1e5)

ripscore <- computeLogOdd(nbhGR)
```

### Description

Given a list of single-end or paired-end read alignment files in BAM/SAM/BED format, compute the read counts and normalized read counts as expression of annotated transcript in the unit of "reads per kilobase of exon per million mapped reads" (RPKM).

### Usage

```r
computeRPKM(bamFiles, RIPSeekerRead = TRUE, paired = FALSE, countMode = "IntersectionNotEmpty", featureGRanges, idType = "ensembl_transcript_id", featureType = "exon", ignore.strand = FALSE, txDbName = "biomart", moreGeneInfo = FALSE, saveData, justRPKM = TRUE, ...)
```
computeRPKM

Arguments

- **bamFiles**: A list of one or more BAM/SAM/BED alignment files.
- **RIPSeekerRead**: Binary flag. If TRUE, then import and process the alignment files using the built-in function `combineAlignGals` from RIPSeeker package; if FALSE, then import the files by directly calling the required functions. The flag makes using the function outside of RIPSeeker package become possible.
- **paired**: Binary to indicate whether the alignment files are paired-end. The alignment file must be either paired-end or single-end but not both.
- **countMode**: An argument used to set the mode argument in the underlying function `summarizeOverlaps` employed to compute the read counts for each feature. The possible mode includes "Union", "IntersectionStrict", and "IntersectionNotEmpty". All three modes avoid double counting the reads by either discarding reads that completely fall into multiple features or counting the read only once for the feature that uniquely and completely includes it. Please refer to `summarizeOverlaps` for details.
- **featureGRanges**: GRanges of features as an optional argument for function to compute RPKM/FPKM just for those features without retrieving online annotations.
- **idType**: A character string that specifies the type of the annotations, which can "ensembl_transcript_id", "ensembl_gene_id", "ucsc", etc. Refer to `listFilters` for more information.
- **featureType**: Features that will be grouped by genes/transcripts in a GRangesList. The available options are "exon" (Default), "intron", "fiveUTR", "threeUTR", and "CDS" corresponding to the functions `exonsBy`, `cdsBy`, `intronsByTranscript`, `fiveUTRsByTranscript`, `threeUTRsByTranscript`, and `cdsBy`, respectively.
- **ignore.strand**: Whether to ignore strand when counting the reads (Default: FALSE).
- **txDbName**: Name of the transcript database to use to retrieve the annotation. The available options are "biomart" (Default) or "UCSC" corresponding to the functions `makeTxDbFromBiomart` and `makeTxDbFromUCSC`, respectively.
- **moreGeneInfo**: Binary indicator to indicate whether to download more information for each genes/transcripts rather than having only the gene/transcript IDs (Default: FALSE).
- **saveData**: Path of output file.
- **justRPKM**: Binary for whether to return only the `RangedSummarizedExperiment`.
- **...**: Extra arguments passed to functions `makeTxDbFromBiomart`, `makeTxDbFromUCSC`, `useMart`, `combineAlignGals`.

Details

The function is a wrapper function making use of several external functions from several well maintained and freely available Bioconductor packages including GenomicFeatures, GenomicRanges, biomaRt and Rsamtools packages. The paired-end alignments are converted into single-end using function `galp2gal` and then subject to read count computation by `summarizeOverlaps`, which does not yet directly support paired-end alignments.

Value

- **rpkmSEobject**: A `RangedSummarizedExperiment` object with assays slot saved for counts, `rowRanges` holds the features, `metadata` for RPKM/FPKM (normalized) gene expression.
computeRPKM

rpkmDF
Data frame with or without the detailed gene information columns depending on whether moreGeneInfo is TRUE or FALSE. rpkmDF is only returned within in a list when justRPKM is FALSE.

featureGRanges
The features in GRanges object that are used to compute the gene expression. featureGRanges is only returned within in a list when justRPKM is FALSE.

Note
Also works for RNA-seq alignments.

Author(s)
Yue Li

References


See Also
makeTxDbFromBiomart, makeTxDbFromUCSC, useMart, exonsBy, cdsBy, intronsByTranscript, fiveUTRsByTranscript

Examples
if(interactive()) { # need internet connection

# Retrieve system files
extdata.dir <- system.file("extdata", package="RIPSeeker")
bamFiles <- list.files(extdata.dir, ".bam$", recursive=TRUE, full.names=TRUE)
bamFiles <- grep("PRC2", bamFiles, value=TRUE)

# use biomart
txDBName <- "biomart"
biomart <- "ENSEMBL_MART_ENSEMBL" # use archive to get ensembl 65
dataset <- "mmusculus_gene_ensembl"
host <- "de2011.archive.ensembl.org" # use ensembl 65 for annotation
resultlist <- computeRPKM(bamFiles=grep(pattern="SRR039214", bamFiles, value=TRUE, invert=TRUE), #featureGRanges=featureGRanges,
Among multiple alignments of the same read (i.e. multihit), select the alignment corresponding to the bin with the maximum posterior for the enriched state.

**Usage**

```r
disambiguateMultihits(alignGal, nbhGRList, postprobCutoff = 0)
```

**Arguments**

- `alignGal` GAlignments object with an additional column in the `values` slot that indicates whether the read corresponding to the current alignment is a unique hit (i.e., read mapped uniquely to a single loci) or multihit (i.e., read mapped to multiple loci).
- `nbhGRList` GRangesList each item containing the HMM training results on a single chromosome. Importantly, the posterior probabilities for the background and enriched states need to be present in the `metadata` slot and used to disambiguate multihits, which is done by `mainSeekSingleChrom`.
- `postprobCutoff` Posterior cutoff for returning only the reads with maximum posterior that is greater than the threshold (Default: 0; i.e., no cutoff).

**Details**

Each multihit (i.e., read aligned to multiple loci) flagged in the `getAlignGal` function are assigned to a unique locus corresponding to the \( j^{\text{th}} \) bin with the highest posterior or responsibility from the RIP state. Intuitively, the RIP state corresponds to the read-enriched loci. Disambiguating multihits in this way will potentially improve the power of detecting more RIP regions but may also introduce certain bias towards the idea of "rich gets richer". After this step, RIPSeeker will rerun the functions from `selectBinSize` to `nbh` to improve the HMM model estimation with augmented read count data. Optionally, user can choose not to reiterate the training process to go straight to the next step to detect RIP regions (See `seekRIP`).

**Value**

GAlignments with each read mapped uniquely to a single locus.

**Author(s)**

Yue Li
empiricalFDR

Compute empirical false discovery rate

Description

At a p-value, find the number of regions in RIP library (denoted as "trueCount") and the number of regions in control library (denoted as "falseCount"). The empirical false discovery rate (eFDR) is estimated as the ratio of the falseCount over the trueCount.

See Also

getAlignGal, ripSeek, mainSeek, mainSeekSingleChrom

Examples

```r
# Retrieve system files
extdata.dir <- system.file("extdata", package="RIPSeeker")

bamFiles <- list.files(extdata.dir, ".bam$", recursive=TRUE, full.names=TRUE)
bamFiles <- grep("PRC2", bamFiles, value=TRUE)

# Parameters setting
binSize <- 1e5  # use a large fixed bin size for demo only
minBinSize <- NULL  # turn off min bin size in automatic bin size selection
maxBinSize <- NULL  # turn off max bin size in automatic bin size selection
multicore <- FALSE  # use multicore
strandType <- "-"  # set strand type to minus strand

# Retrieve system files
extdata.dir <- system.file("extdata", package="RIPSeeker")
bamFiles <- list.files(extdata.dir, ".bam$", recursive=TRUE, full.names=TRUE)
bamFiles <- grep("PRC2", bamFiles, value=TRUE)

alignGal <- combineAlignGals(bamFiles=grep(pattern="SRR039214", bamFiles, value=TRUE, invert=TRUE), reverseComplement=TRUE, genomeBuild="mm9")
alignGR <- as(alignGal, "GRanges")
alignGR <- addPseudoAlignment(alignGR)
alignGRL <- GRangesList(as.list(split(alignGR, seqnames(alignGR))))

# run mainSeekSingleChrom function for HMM inference on a single chromosome
nbhGRL <- lapply(alignGRL, mainSeekSingleChrom, K = 2, binSize=binSize,
                  minBinSize = minBinSize, maxBinSize = maxBinSize, runViterbi=FALSE)
nbhGRL <- GRangesList(nbhGRL)
alignGalFiltered <- disambiguateMultihits(alignGal, nbhGRL)
```
**empiricalFDR**

**Usage**

```r
empiricalFDR(pval, pvalRIP, pvalCTL)
```

**Arguments**

- `pval`: A scalar p-value.
- `pvalRIP`: A column vector of p-values for the peaks identified from RIP v.s. control comparison.
- `pvalCTL`: A column vector of p-values for the peaks identified from control v.s. RIP comparison.

**Details**

Only when the control is available, is an empirical false discovery rate (eFDR) estimated based on the idea of "sample swap" inspired by MACS (a ChIP-seq algorithm from Zhang et al. (2008). At each p-value, RIPSeeker finds the number of significant RIP-regions over control (CTL) based on `pvalRIP` and the number of significant control regions over RIP based on `pvalCTL`. The eFDR is defined as the ratio of the number of "RIP" (false positive) regions identified from CTL-RIP comparison over the number of RIP regions from the RIP-CTL comparison. The maximum value for eFDR is 1 and minimum value for eFDR is max(p-value, 0). The former takes care of the case where the numerator is bigger than the denominator, and the latter for zero numerator.

**Value**

A scalar probability value that represents the eFDR.

**Note**

This is an internal function used in `seekRIP`.

**Author(s)**

Yue Li

**References**


**See Also**

`logScoreWithControl`, `seekRIP`, `computeLogOdd`, `scoreMergedBins`

**Examples**

```r
pvalRIP <- runif(100)
pvalCTL <- runif(100)
eFDR <- empiricalFDR(pvalRIP[1], pvalRIP, pvalCTL)
pvalRIP[1]
```
```
eFDR

# more significant pval
pvalRIP[1] <- 1e-4

eFDR <- empiricalFDR(pvalRIP[1], pvalRIP, pvalCTL)

pvalRIP[1]
eFDR
```

---

**evalBinSize**  
*Evaluate bin size using Shimazaki cost function*

**Description**

Given a bin size and a GRanges alignment object, the function computes the bin count and returns the cost of the bin size based on Shimazaki cost function.

**Usage**

evalBinSize(binSize, alignGR)

**Arguments**

- `binSize` An integer that indicates the bin size applied to the binning of the chromosome.
- `alignGR` GRanges object of alignments to a single chromosome.

**Details**

The function implements the algorithm developed by Shimazaki and Shinomoto (2007), which is based on the goodness of the fit of the time histogram to estimate the rate of neural response of an animal to certain stimuli in a spike-in experiment. The algorithm involves four simple steps:

1. Divide chromosome sequence into \( N \) bins of width \( b \).
2. Count number of read counts \( x_i \) that enter the \( i \)'th bin.
3. Compute: \( \bar{x} = \frac{1}{N} \sum_{i=1}^{N} x_i \) and \( v = \frac{1}{N} \sum_{i=1}^{N} (x_i - \bar{x})^2 \).
4. Compute: \( C(b) = \frac{2\bar{x} - v}{b^2} \)

**Value**

- `cost` A scalar value for the cost of the bin size.

**Author(s)**

Yue Li

**References**

See Also

selectBinSize, binCount

Examples

# Retrieve system files
extdata.dir <- system.file("extdata", package="RIPSeeker")

bamFiles <- list.files(extdata.dir, ".bam", recursive=TRUE, full.names=TRUE)
bamFiles <- grep("PRC2", bamFiles, value=TRUE)

alignGal <- getAlignGal(bamFiles[1], reverseComplement=TRUE, genomeBuild="mm9")

alignGR <- as(alignGal, "GRanges")

alignGRList <- GRangesList(as.list(split(alignGR, seqnames(alignGR))))

binSize <- 1000

costs <- evalBinSize(binSize, alignGRList$chrX)

---

exportGRanges  

Export GRanges object in a specified format

Description

A wrapper function of export with additional support for exporting tab-delimited format with no re-arrangement of the original GRanges output.

Usage

exportGRanges(gRanges, outfile, exportFormat)

Arguments

  gRanges  GRanges object to export.
  outfile  File path for output.
  exportFormat  Desirable format including "txt" and other formats specified in export.

Value

Output the text to the file stream defined in outfile.

Note

The function is used in ripSeek to export desired format and can be used as general purpose function.

Author(s)

Yue Li
Convert GAlignmentPairs to GAlignments

Description

Convert GAlignmentPairs to GAlignments using CIGAR to mark flanked portion of the pairs as 'N'.

Usage

galp2gal(galp)

Arguments

galp

Details

Each proper read pairs is combined into a single alignment record making use of the CIGAR flag 'N' to indicate the number of bases between the mate pairs (i.e., the difference between the start of the right mate pair and the end of the left mate pair). In other words, the paired-end alignments are treated as gapped alignments of long fragments. The function is used within getAlignGal but can be used as a stand-alone function as well.
Import and process single-end or paired-end alignments in a BAM/SAM/BED file to retain valid alignments defined by the arguments below. Multihits (same read mapped to multiple loci) are flagged for the subsequent disambiguation with function `disambiguateMultihits`. The final output is a GAlignments object.

### Usage

```r
getAlignGal(alignFilePath, format, genomeBuild, deleteGeneratedBAM = FALSE, reverseComplement = FALSE, returnDuplicate = FALSE, flagMultiHits = TRUE, returnOnlyUniqueHits = FALSE, paired = FALSE, ...)
```
getAlignGal

Arguments

alignFilePath Path to the alignment file.

format The alignment format can be determined automatically from the file extension or specified by the user. The supported formats are BAM, SAM, and BED.

genomeBuild Genome build used to obtain the chromosome information from online UCSC database in order to construct GAlignments object. Since the BAM/SAM header provides the chromosome information, the argument needs to be set only in the absence of the header information for some BAM/SAM files or when BED file is used. Examples for the common genomeBuild are "mm9" for mouse or "hg19" for human reference genomes. Note that an appropriate genome build that has been used in the alignment is important for desirable outcome. For instance, user should use "mm10" if the alignments are based on "mm10" rather than "mm9" genome build.

deleteGeneratedBAM Binary indicator to indicate whether the converted BAM from the original SAM input file needs to be deleted from the local disk (Default: FALSE).

reverseComplement Binary indicator to indicate whether the reads were sequenced from the opposite strand of the original RNA molecule. reverseComplement only applies to strand-specific sequencing in which case only the strand generated during second strand synthesis is sequenced. Thus, if reverseComplement=TRUE, the strand signs of the alignments are switched (i.e. + to -, - to +, and * unchanged); otherwise (reverseComplement=FALSE) retain the original the strand signs.

returnDuplicate Indicator (TRUE, FALSE, NA) to instruct whether the duplicate alignments need to be returned (Default: FALSE). Duplicate reads are a set of reads that align to exactly the same genomic coordinate. Because transcripts are usually hundreds or thousands of base pairs long and thus much longer than the read (25-100 nt), the chance that the same 25-100 nt portion of the transcript being sequenced multiple times is very small and may very likely be due to PCR artifact. This argument is actually passed to 'isDuplicate' in scanBamFlag.

flagMultiHits Binary indicator for whether to add additional binary column named "unique-Hits" to indicate whether the corresponding aligned reads are unique hit (unique-Hits==TRUE) or multihit (uniqueHits==FALSE). Multihits represent multiple alignments of the same read due to gene duplications or repetitive elements of the genome. The multihits typically constitute a substantial proportion of the total mapped reads. Rather than being removed, these multihits are flagged (flagMultiHits=TRUE by default) and in the later step assigned to a unique region by (disambiguateMultihits).

returnOnlyUniqueHits Binary indicator to return only the unique hits and discard all of the multihits (Default: FALSE).

paired Binary indicator to indicate whether the alignments are paired-end (Default: FALSE). For paired-end alignments, properly paired reads are combined into a single alignment record making use of the CIGAR flag ‘N’ to indicate the number of bases between the mate pairs (i.e., the length of the insert fragment). In other words, the paired-end alignments are treated as gapped alignments of long fragments (See galp2gal).

... Extra arguments are ignored.
**Details**

The BAM file is imported using `readGAlignments` for single-end or `readGAlignmentPairs` for paired-end alignments. The SAM file is converted to BAM first and then imported as above. The BED file is first imported by `import` as GRanges object and subsequently converted to GAlignments via the constructor function `GAlignments`.

**Value**

alignGal  
GAlignments object containing the processed alignments with the values slot saved for the "uniqueHits" binary flag (See `flagMultiHits` above) and metadata saved as a list containing argument setting for `reverseComplement`, `returnDuplicate`, `flagMultiHits`, `returnOnlyUniqueHits`.

**Author(s)**

Yue Li

**References**


Michael Lawrence, Vince Carey and Robert Gentleman. rtracklayer: R interface to genome browsers and their annotation tracks. R package version 1.16.3.

**See Also**

`combineAlignGals`, `readGAlignments`, `readGAlignmentPairs`, `import`

**Examples**

```r
# Retrieve system files
extdata.dir <- system.file("extdata", package="RIPSeeker")
bamFiles <- list.files(extdata.dir, ".bam$", recursive=TRUE, full.names=TRUE)
bamFiles <- grep("PRC2", bamFiles, value=TRUE)
alignGal <- getAlignGal(bamFiles[1], reverseComplement=TRUE, genomeBuild="mm9")
```

---

**logScoreWithControl**  
*Compute RIPScore based on RIP and control posteriors and test for significance*

**Description**

Compute the RIPScore using both RIP and control posteriors for each bins, merge and summarize the scores for the merged bins, and finally compute the p-value and adjusted p-value for the summary RIPScore.

**Usage**

```r
logScoreWithControl(nbhGRRIP, nbhGRCTL, padjMethod = "BH", getControlStats = TRUE)
```
logScoreWithControl

Arguments

nbhGRRIP  GRanges object for the RIP library created from mainSeek containing the posterior probabilities of the hidden states for each observed read count.
nbhGRCTL  An optional argument as a GRanges object for the control library created from mainSeek containing the posterior probabilities of the hidden states for each observed read count.
padjMethod  Method used to adjust multiple testing performed in p.adjust (Default: "BH").
getControlStats  Binary indicator to whether return statistics including computed for the control library alone. If TRUE, then all control specific score columns will be reported with a prefix "CTL".

Details

The RIPScore is computed in computeLogOdd as the log odd ratio of the posterior for the RIP state ($z_i = 2$) over the posterior for the background state ($z_i = 1$) in RIP library subtracted by the log odd ratio computed from the control library. The adjacent bins with hidden states predicted by nbh_vit as the enriched state (corresponding to the NB with larger mean) are merged. The RIPScores are averaged over the merged bins. To assess the statistical significance of the RIPScore for each region, we assume that the RIPScore follows a Gaussian (Normal) distribution with mean and standard deviation estimated using the RIPScores over all of the bins. The rationale is based on the assumption that most of the RIPScores correspond to the background state and together contribute to a stable estimate of the test statistics (TS) and p-value computed using the R built-in function pnorm. The p-value is adjusted by p.adjust with BH method by default. The same procedure is applied optionally to the control library.

Value

GRanges of merged bins with values slot saved for RIPScore (lodOdd), p-value (pval), adjusted p-value (pvalAdj) for RIP and optionally for control.

Note

Internal function used by seekRIP.

Author(s)

Yue Li

See Also

logScoreWithoutControl, seekRIP, computeLogOdd, scoreMergedBins

Examples

if(interactive()) { # check the example in seekRIP
# Retrieve system files
extdata.dir <- system.file("extdata", package="RIPSeeker")
bamFiles <- list.files(extdata.dir, ".bam\$", recursive=TRUE, full.names=TRUE)
bamFiles <- grep("PRC2", bamFiles, value=TRUE)
logScoreWithoutControl

Compute RIPScore based on RIP posteriors alone and test for significance

Description

Compute the RIPScore using only the RIP (typically when control is unavailable) posteriors for each bins, merge and summarize the scores for the merged bins, and finally compute the p-value and adjusted p-value for the summary RIPScore.

Usage

logScoreWithoutControl(nbhGRRIP, padjMethod = "BH")

Arguments

nbhGRRIP  
GRanges object for the RIP library created from `mainSeek` containing the posteriors probabilities of the hidden states for each observed read count.

pajdMethod  
Method used to adjust multiple testing performed in `p.adjust` (Default: "BH").
The RIPScore is computed in `computeLogOdd` as the log odd ratio of the posterior for the RIP state \((z_i = 2)\) over the posterior for the background state \((z_i = 1)\) in RIP library alone (typically when control is unavailable). The adjacent bins with hidden states predicted by `nbh_vit` as the enriched state (corresponding to the NB with larger mean) are merged. The RIPScores are averaged over the merged bins. To assess the statistical significance of the RIPScore for each region, we assume that the RIPScores follow a Gaussian (Normal) distribution with mean and standard deviation estimated using the RIPScores over all of the bins. The rationale is based on the assumption that most of the RIPScores correspond to the background state and together contribute to a stable estimate of the test statistics (TS) and p-value computed using the R built-in function `pnorm`. The p-value is adjusted by `p.adjust` with BH method by default.

**Value**

GRanges of merged bins with values slot saved for RIPScore (lodOdd), p-value (pval), adjusted p-value (pvalAdj) for RIP

**Note**

Internal function used by `seekRIP`.

**Author(s)**

Yue Li

**See Also**

`logScoreWithControl`, `seekRIP`, `computeLogOdd`, `scoreMergedBins`

**Examples**

```r
if(interactive()) {
  # check the example in seekRIP
  # Retrieve system files
  extdata.dir <- system.file("extdata", package="RIPSeeker")

  bamFiles <- list.files(extdata.dir, ".bam$", recursive=TRUE, full.names=TRUE)

  bamFiles <- grep("PRC2", bamFiles, value=TRUE)

  # Parameters setting
  binSize <- 1e5 # use a large fixed bin size for demo only
  multicore <- FALSE # use multicore
  strandType <- "-" # set strand type to minus strand

  # run main function for HMM inference on all chromosomes
  mainSeekOutputRIP <- mainSeek(bamFiles=grep(pattern="SRR039214", bamFiles, value=TRUE, invert=TRUE),
                                binSize=binSize, strandType=strandType,
                                reverseComplement=TRUE, genomeBuild="mm9",
                                uniqueHit = TRUE, assignMultihits = TRUE,
                                rerunWithDisambiguatedMultihits = FALSE,
                                multicore=multicore, silentMain=FALSE, verbose=TRUE)

  # Compute log score and test for significance WITHOUT control
```
mainSeek $\quad$ Train HMM parameters on each chromosome independently from the alignments.

Description

A back-end function used by the front-end function ripSeek to train HMM parameters on all of the chromosomes independently. This function in turn calls another function mainSeekSingleChrom to compute HMM parameters on each chromosome separately or in parallel (if multicore is TRUE).

Usage

```r
mainSeek(bamFiles, reverseComplement = FALSE, genomeBuild = "mm9", uniqueHit = TRUE, assignMultihits = TRUE, strandType = NULL, paired = FALSE, rerunWithDisambiguatedMultihits = TRUE, silentMain = FALSE, multicore = TRUE, returnAllResults = TRUE, ...)```

Arguments

- **bamFiles** A list of paths to individual BAM files. BED and SAM files are also accepted.
- **reverseComplement** Whether the reads came from the original or the opposite strand of the RNA being sequenced. If former, then reverseComplement should be FALSE; otherwise TRUE, in which case the strand signs will be switched from + to -, - to +, and * is unchanged.
- **genomeBuild** When the input alignment format is BED, genomeBuild is only required in `getAlignGal` to determine the chromosome lengths for the GAlignments object using function `SeqinfoForUCSCGenome`. BAM and SAM header have chromosome information, and thus genomeBuild is not needed.
- **uniqueHit** Binary indicator. If uniqueHit=TRUE, only reads mapped to single unique loci are used to train the HMM. Otherwise, all of the reads including multihits will be used for the HMM. A multihit is a read mapped to more than one loci. The flags for uniqueHits and multihits are the metadata values of GAlignments object constructed in `getAlignGal`.
- **assignMultihits** Binary indicator used by ripSeek to tell the function whether disambiguate multihits by assigning them to unique loci with the maximum posterior probability obtained from running HMM (See `nbh_em`)
- **strandType** A character variable indicate which strand the RIPSeeker needs to operate on. The options are NULL, '+', '-', '*'. If NULL or '*', then all of the reads will be used (preferable for non-strand specific sequencing). If '+' or '-', only reads from '+' or '-' strand will be used, respectively. Note that the sign is assumed to be THE SAME AS the strand sign of the processed alignment object and will be the opposite sign if reverseComplement is TRUE (See reverseComplement above).
paired Binary to indicate whether the library is paired-end (TRUE) or single-end (FALSE by default) (see getAlignGal).

rerunWithDisambiguatedMultihits After multihits have been assigned to unique loci, rerunWithDisambiguatedMultihits (Default: TRUE) indicates whether to re-run the HMM on the augmented read alignment data. If FALSE, the HMM step will not be re-run, and the workflow will proceed to RIP detection (See seekRIP) using the nondisambiguated alignments, which can either be the alignments containing only the uniqueHits (if uniqueHit=TRUE) or the alignments containing both the uniqueHits and multihits (if uniqueHit=FALSE).

silentMain Binary indicator to indicate whether to disable the verbose from the mainSeekSingleChrom function. If FALSE (by default), the EM training process will be output to the console for user to keep track of the training progress.

multicore Binary indicator to indicate whether to use mclapply function to compute HMM on chromosomes in parallel. The multicore function will speed up the computation by a factor proportional to the total number of CPU cores on the machine but may impose larger memory overhead than the single-threading approach.

returnAllResults Binary indicator to indicate whether to return all (HMM trained parameters, original, and disambiguated GAlignments) or just the HMM results.

Value A list containing:

- nbhGRLList GRangesList each item containing the HMM training results on a single chromosome.
- alignGal Original alignment data in GAlignments object
- alignGalFiltered Disambiguated alignment data with multihits assigned to unique loci.

Author(s) Yue Li

See Also ripSeek, mainSeekSingleChrom, mclapply

Examples

```r
# Retrieve system files
extdata.dir <- system.file("extdata", package="RIPSeeker")
bamFiles <- list.files(extdata.dir, ".bam$", recursive=TRUE, full.names=TRUE)
bamFiles <- grep("PRC2", bamFiles, value=TRUE)

# Parameters setting
binSize <- 1e5 # use a large fixed bin size for demo only
minBinSize <- NULL # min bin size in automatic bin size selection
maxBinSize <- NULL # max bin size in automatic bin size selection
```
multicore <- FALSE # use multicore
strandType <- "-" # set strand type to minus strand

######################################################################### run mainSeekSingleChrom function for HMM inference on all chromosomes #########################################################################
mainSeekOut <- mainSeek(bamFiles=grep(pattern="SRR039214", bamFiles, value=TRUE, invert=TRUE),
                           binSize=binSize, minBinSize = minBinSize,
                           maxBinSize = maxBinSize, strandType=strandType,
                           reverseComplement=TRUE, genomeBuild="mm9",
                           uniqueHit = TRUE, assignMultihits = TRUE,
                           rerunWithDisambiguatedMultihits = FALSE,
                           multicore=multicore, silentMain=FALSE, verbose=TRUE)

mainSeekSingleChrom  Automatic bin size selection, bin count, and HMM parameters optimization on read count vector from a single chromosome (Internal function)

Description

This an internal function used by mainSeek to accomplish three major tasks on a single chromosome: automatically select bin size, compute read counts within the bins, and obtain optimal HMM parameters.

Usage

mainSeekSingleChrom(alignGR, K = 2, binSize = NULL, minReadCount = 10,
                    backupNumBins = 10, minBinSize = 200, maxBinSize = 1200,
                    increment = 5, pathToSavePlotsOfBinSizesVersusCosts,
                    verbose = TRUE, allowSecondAttempt = TRUE, ...)

Arguments

alignGR  GRanges containing the alignments on a single chromosome.
K  Number of hidden states (Default: 2). By default, state 1 specifies the background and state 2 the RIP regions. The two states are recognized by the means for the two distributions (See nbh_em).
binSize  Size to use for binning the read counts across each chromosome. If NULL, optimal bin size within a range (default: minBinSize=200, maxBinSize=1200) will be automatically selected (See selectBinSize).
minReadCount  Minimum aligned read counts needed for HMM to converge (Default: 10). Note that HMM may not converge some times when majority of the read counts are zero even if some read count > 10. When that happens, a back-up function addDummyProb comes in to create a placeholder for the corresponding chromosome in GRangeList to maintain the data structure to preserve all information (successfully) obtained from other chromosomes.
backupNumBins  If read count is less than minReadCount, then use backupNumBins (Default: 10) to bin the chromosome.
**minBinSize**  
Minimum bin size to start with the bin selection (See `selectBinSize`). Default to 200, common minimum band size selected in RIP or RNA-seq library construction.

**maxBinSize**  
Maximum bin size to stop with the bin selection (See `selectBinSize`). Default: 1200.

**increment**  
Step-wise increment in bin size selection (See `selectBinSize`). Default: 5.

**pathToSavePlotsOfBinSizesVersusCosts**  
Directory used to save the diagnostic plots for bin size selection.

**verbose**  
Binary indicator for disable (FALSE) or enable (TRUE) HMM training message from function `nbh` to output to the console.

**allowSecondAttempt**  
In case HMM fails to converge due to malformed parameters in EM iteration, re-iterating the HMM process each time with a different suboptimal bin size in attempt to succeed in some trial. If all yield nothing, fall back up to `addDummyProb` to return the placeholder for the chromosome.

...  
Arguments passed to `nbh`.

**Value**

**nbhGR**  
GRanges object containing the optimized HMM parameters (and the Viterbi hidden state sequence) accompanied with the read count vector following the (automatic) binning scheme.

**Note**

Unless a highly customized workflow is needed, `ripSeek` is the high-level front-end main function that should be used in most cases.

**Author(s)**

Yue Li

**See Also**

ripSeek, mainSeek, nbh_em

**Examples**

```r
# Retrieve system files
extdata.dir <- system.file("extdata", package="RIPSeeker")

bamFiles <- list.files(extdata.dir, ".bam$", recursive=TRUE, full.names=TRUE)

bamFiles <- grep("PRC2", bamFiles, value=TRUE)

# Parameters setting
binSize <- 1e5  # use a large fixed bin size for demo only
minBinSize <- NULL  # min bin size in automatic bin size selection
maxBinSize <- NULL  # max bin size in automatic bin size selection
multicore <- FALSE  # use multicore
strandType <- "-"  # set strand type to minus strand

# Retrieve system files
```
extdata.dir <- system.file("extdata", package="RIPSeeker")
bamFiles <- list.files(extdata.dir, ".bam$", recursive=TRUE, full.names=TRUE)
bamFiles <- grep("PRC2", bamFiles, value=TRUE)
alignGal <- getAlignGal(bamFiles[1], reverseComplement=TRUE, genomeBuild="mm9")
alignGR <- as(alignGal, "GRanges")
alignGRList <- GRangesList(as.list(split(alignGR, seqnames(alignGR))))

# run main function for HMM inference on a single chromosome
nbhGR <- mainSeekSingleChrom(alignGR=alignGRList$chrX, K = 2, binSize=binSize,
minBinSize = minBinSize, maxBinSize = maxBinSize)
nbhGR

---

**nbh**  
*Generic function of negative binomial HMM*

**Description**  
Generic function for nbh.GRanges and nbh.integer

**Usage**  
nbh(x, ...)

**Arguments**

- **x**: Object of class Integer or GRanges.
- **...**: Extra arguments passed to either nbh.GRanges or nbh.integer.

**Author(s)**
Yue Li

**References**


**See Also**

mainSeekSingleChrom, nbh.integer, nbh.GRanges
Examples

# Retrieve system files
extdata.dir <- system.file("extdata", package="RIPSeeker")

bamFiles <- list.files(extdata.dir, ".bam", recursive=TRUE, full.names=TRUE)
bamFiles <- grep("PRC2", bamFiles, value=TRUE)

alignGal <- getAlignGal(bamFiles[1], reverseComplement=TRUE, genomeBuild="mm9")

alignGR <- as(alignGal, "GRanges")

alignGRList <- GRangesList(as.list(split(alignGR, seqnames(alignGR))))

binSize <- 1e5 # use a large fixed bin size for demo only

binGR <- binCount(alignGRList$chrX, binSize)

# test on GRanges object
nbhGR <- nbh(binGR, 2, runViterbi=TRUE)

# test on integer object
nbhList <- nbh(values(binGR)$count, 2, runViterbi=TRUE)

---

### nbh.GRanges

Optimize HMM parameters based on the read counts on a chromosome

#### Description

Inheritance function from nbh that receives an object of GRanges class with additional column of read counts (for each strand) and call nbh.integer to derive the most probable sequence of hidden states

#### Usage

```
## S3 method for class 'GRanges'
nbh(x, K, ...)
```

#### Arguments

- **x**: GRanges with 'values' slot used for bin counts in 1D vector of integers.
- **K**: Number of hidden states.
- **...**: Extra arguments passed to nbh.integer for the actual HMM computation.
Value

binGR    GRanges of bin counts with metadata slot saved for the optimized HMM parameters including alpha, beta for the K negative binomial mixture components and TRANS (the transition probabilities)

Author(s)

Yue Li

References


See Also

mainSeekSingleChrom, nbh.integer

Examples

if(interactive()) ?nbh # see nbh for example of nbh running on GRanges object

---

### nbh.integer

**HMM posterior decoding and NB parameter optimization**

Description

Inheritance function from nbh that receives a vector of integers and compute optimal HMM parameters via EM algorithm.

Usage

```r
## S3 method for class 'integer'
nbh(x, K, NBM_NIT_MAX = 250,
    NBM_TOL = 0.01, NBH_NIT_MAX = 250,
    NBH_TOL = 0.001, runViterbi = FALSE, ...)
```

Arguments

- `x` A vector of integers, conceptually representing the read counts within bins of chromosome.
- `K` Number of hidden states.
- `NBM_NIT_MAX` Maximum number of EM iterations (Default: 250) for the negative binomial mixture model (NBM) initialization step (See nbm_em).
**NBM_TOL** Threshold as fraction of increase in likelihood (given the current NBM parameters) comparing with the likelihood from the last iteration. EM for the NBM stops when the improvement is below the threshold (Default: 0.01).

**NBH_NIT_MAX** Maximum number of EM iterations (Default: 250) for the negative binomial hidden Markov model (NBH).

**NBH_TOL** Threshold as fraction of increase in likelihood (given the current NBH parameters) comparing with the likelihood from the last iteration. EM for the NBH stops when the improvement is below the threshold (Default: 0.001).

**runViterbi** Binary indicator. If TRUE, Viterbi algorithm will be applied to derive the maximum likelihood hidden state sequence using the optimized HMM parameters obtained from the EM (See *nbh_em*).

... Extra arguments are ignored.

**Details**

The function consists of three major steps: (1) negative binomial mixture model used to initialize HMM parameters; (2) optimization of HMM parameters using EM algorithm; (3) Viterbi maximum-likelihood estimation of hidden state sequence. Step (1) involves optimization of NBM parameters assuming the data points are independently sampled from a mixture of K NB distributions (See *nbh_init*). Given the optimized parameters for K-NBM, step (2) drops the independence assumption by introducing the transition probability between hidden variables, which is initialized as the mixing proportions of NBM (See *nbh_init*). Given the optimized HMM parameters, step (3) derives the maximum likelihood hidden state sequence using Viterbi algorithm. Step (3) is run only when runViterbi is TRUE.

**Value**

A list containing:

- **initAlpha** Initialized alpha of NBM from *nbh_init*.
- **initBeta** Initialized beta of NBM from *nbh_init*.
- **initTRANS** Initialized mixing proportion of NBM from *nbh_init*.
- **postprob** Posteriors of the K hidden states for each observed count derived from *nbh_em* (e.g., posteriors of background and enriched state in a two-state HMM).
- **alpha** Optimized alpha of the NB mixture components in the HMM using *nbh_em*.
- **TRANS** Optimized transition probability of the HMM using *nbh_em*.
- **viterbi_state** Sequence of discrete values representing the hidden states derived from the maximum likelihood estimation using Viterbi algorithm (See *nbh_vit*).

**Author(s)**

Yue Li

**References**


nbh_chk

See Also
mainSeekSingleChrom, nbh, nbh.GRanges

Examples
if(interactive()) ?nbh # see nbh for example of nbh running on integer object

---

nbh_chk  

Check the parameters of the negative binomial HMM

Description
The function verifies the numerical range and dimension of the NBH paramters alpha, beta, and TRANS and returns the number of hidden states. It is used in nbh_em before running EM.

Usage
nbh_chk(TRANS, alpha, beta)

Arguments
TRANS  

Expected a squared matrix of probabilities (0 \leq p \leq 1) with row and column length equal to that of alpha and beta and row sum and column sum both equal to 1 (within some numerical deviation of 1e-6).

alpha

Expected a vector of positive values with length equal to that of beta and the row/column of TRANS.

beta

Expected a vector of positive values with length equal to that of alpha and the row/column of TRANS.

Value
N   

Number of components or equivalently the length of alpha, beta, or wgth.

Author(s)
Yue Li

References

See Also
nbh_em, nbm_chk
Examples

```r
# two hidden states
TRANS <- matrix(c(0.9, 0.1, 0.3, 0.7), nrow=2, byrow=TRUE)
alpha <- c(2, 4)
beta <- c(1, 0.25)
nbh_chk(TRANS, alpha, beta)
```

                       **nbh_em**  

*Expectation conditional maximization of negative binomial HMM parameters using forward-backward algorithm*

Description

Given an input read count vector of integers, the function optimizes the parameters for the negative binomial HMM of K hidden states using expectation conditional maximization with forward-backward algorithm to achieve the exact inference.

Usage

```r
nbh_em(count, TRANS, alpha, beta, NBH_NIT_MAX = 250, 
        NBH_TOL = 1e-05, MAXALPHA = 1e+07, MAXBETA = 1e+07)
```

Arguments

- **count** A vector of integers, conceptually representative of the read counts within bins of chromosome.
- **TRANS** Transition probability matrix, a squared matrix of probabilities \(0 \leq p \leq 1\) with row and column length equal to that of alpha and beta and row sum and column sum both equal to 1 (within some numerical deviation of \(1e^{-6}\)).
- **alpha** Shape parameter of the NB as a vector of positive values with length equal to that of beta and the row/column of TRANS.
- **beta** Inverse scale parameter of the NB as a vector of positive values with length equal to that of beta and the row/column of TRANS.
- **NBH_NIT_MAX** Maximum number of EM iterations (Default: 250) for the negative binomial hidden Markov model (NBH).
- **NBH_TOL** Threshold as fraction of increase in likelihood (given the current NBH parameters) comparing with the likelihood from the last iteration. EM for the NBH stops when the improvement is below the threshold (Default: 0.001).
- **MAXALPHA** The maximum value of alpha in case the update goes beyond the numerical upper limit of the system. Once alpha becomes larger than MAXALPHA, the EM iteration is prematurely terminated to prevent malfunction.
- **MAXBETA** The maximum value of beta in case the update goes beyond the numerical upper limit of the system. Once beta becomes larger than MAXBETA, the EM iteration is prematurely terminated to prevent malfunction.
Details

Given a K-state HMM with NB emission (NBH), the goal is to maximize the likelihood function with respect to the parameters comprising of $\alpha_k$ and $\beta_k$ for the K NB components and the transition probabilities $A_{jk}$ between any state $j$ and $k$, which are the priors $p(z = k)$. Because there is no analytical solution for the maximum likelihood (ML) estimators of the above quantities, a modified EM procedures called Expectation Conditional Maximization is employed (Meng and Rubin, 1994).

In E-step, the posterior probability is evaluated by forward-backward algorithm using NB density functions with initialized alpha, beta, and TRANS. In the CM step, $A_{jk}$ is evaluated first followed by Newton updates of $\alpha_k$ and $\beta_k$. EM iteration terminates when the percentage of increase of log likelihood drop below NBH_TOL, which is deterministic since EM is guaranteed to converge. For more details, please see the manuscript of RIPSeeker.

Value

A list containing:

- **alpha**: optimized alpha_k for NB at state K
- **beta**: optimized beta_k for NB at state K
- **TRANS**: optimized transition probability matrix
- **logl**: Log likelihood in each EM iteration.
- **postprob**: Posterior probabilities for each observed data point at the last EM iteration.
- **dens**: the negative binomial probabilities computed at the last EM iteration

Author(s)

Yue Li

References


See Also

- nbh_init, nbh, nbh.GRanges, nbh_vit,nbm_em

Examples

```r
# Simulate data
TRANS_s <- matrix(c(0.9, 0.1, 0.3, 0.7), nrow=2, byrow=TRUE)
alpha_s <- c(2, 4)
beta_s <- c(1, 0.25)
Total <- 100
```
nbh_gen

Simulate data from a negative binomial HMM.

Description

Generate count data and the hidden states based on the user-supplied HMM parameters.
**nbh_gen**

**Usage**

nbh_gen(TRANS, alpha, beta, Total)

**Arguments**

TRANS  
Expected a squared matrix of probabilities \((0 \leq p \leq 1)\) with row and column length equal to that of alpha and beta and row sum and column sum both equal to 1 (within some numerical deviation of 1e-6).

alpha  
Expected a vector of positive values with length equal to that of beta and the row/column of TRANS.

beta  
Expected a vector of positive values with length equal to that of alpha and the row/column of TRANS.

Total  
Total number of data points to generate.

**Value**

A list containing:

- count  
  Simulation count data.
- label  
  Hidden states associated with the simulated data.

**Author(s)**

Yue Li

**References**


**See Also**

nbh_em, nbm_chk, randindx

**Examples**

# Simulate data using user-supplied transition prob, alpha and beta for the NB HMM parameters
TRANS_s <- matrix(c(0.9, 0.1, 0.3, 0.7), nrow=2, byrow=TRUE)
alpha_s <- c(2, 4)
beta_s <- c(1, 0.25)
Total <- 100
x <- nbh_gen(TRANS_s, alpha_s, beta_s, Total)
nbh_init

Initialize negative binomial HMM parameters using negative binomial mixture model

Description

The function finds a sensible set of initial NB HMM parameters by fitting a NB mixture model of K components using the read count data.

Usage

nbh_init(count, K, NBM_NIT_MAX = 250, NBM_TOL = 0.001)

Arguments

count
A vector of integers, conceptually representing the read counts within bins of chromosome.

K
Number of hidden states.

NBM_NIT_MAX
Maximum number of EM iterations (Default: 250) for the negative binomial mixture model (NBM) initialization step (See nbm_em).

NBM_TOL
Threshold as fraction of increase in likelihood (given the current NBM parameters) comparing with the likelihood from the last iteration. EM for the NBM stops when the improvement is below the threshold (Default: 0.01).

Details

Because the EM algorithm in HMM tends to fall into local optimal with poor initialization, NB mixture model with K mixture components (K-NBM) is first applied to the data to obtain a reasonable estimate for the HMM parameters. Given the read count vector, the function applied the lower level function nbm_em (NB mixture model) to find alpha, beta, and mixing proportion of the K NB mixture components. Alpha and beta are the parameters of the NB mixture components initialized as the last K quantiles of the nonzero read counts and 1, respectively. The mixing proportions or component weights (wght) of the NB distributions are first initialized as uniform and after EM optimization are used to form a symmetrical transition probability matrix such that probability of state 1 transitioning to state 2 is equal to the probability of state 2 transitioning to state 1. Such matrix is used as the initial transition probability for the HMM model training (See nbh_em).

Value

A list containing:

alpha
Alpha parameter of the K NB components optimized using nbm_em

beta
Beta parameter of the K NB components optimized using nbm_em

TRANS
Transition probability initialized as a symmetrical matrix of mixing proportion of the K NB components optimized using nbm_em.

Author(s)

Yue Li
References


See Also

nbm_em, nbh, nbh.GRanges, nbh_em

Examples

# Simulate data
Total_train <- 1000
Total_test <- 200
TRANS_s <- matrix(c(0.9, 0.1, 0.5, 0.5), nrow=2, byrow=TRUE)
alpha_s <- c(2, 2)
beta_s <- c(1, 0.25)
train <- nbh_gen(TRANS_s, alpha_s, beta_s, Total_train)
test <- nbh_gen(TRANS_s, alpha_s, beta_s, Total_test)
nbhInit <- nbh_init(train$count, ncol(TRANS_s))
count <- train$count
label <- train$label
# NBH initialization
nbhInit <- nbh_init(count, ncol(TRANS_s))
TRANS0 <- nbhInit$TRANS
alpha0 <- nbhInit$alpha
beta0 <- nbhInit$beta
# NBH EM
nbh <- nbh_em(count, TRANS0, alpha0, beta0)
map.accuracy <- length(which(max.col(nbh$postprob) == label))/Total_train
vit <- nbh_vit(count, nbh$TRANS, nbh$alpha, nbh$beta)
vit.accuracy <- length(which(vit$class == label))/Total_train
vit_test <- nbh_vit(test$count, nbh$TRANS, nbh$alpha, nbh$beta)
vit_test.accuracy <- length(which(vit_test$class == test$label))/Total_test
nbh_wt_KMLinit <- list(mapAccuracy_train=map.accuracy, vitAccuracy_train=vit.accuracy, vitLogl_train=vit$logl, vitAccuracy_test=vit_test.accuracy, vitLogl_test=vit_test$logl)
nbh_vit

Derive maximum likelihood hidden state sequence using Viterbi algorithm

Description

Given read counts and HMM parameters (optimized by nbh_em), derive the sequence of hidden states that maximizes the joint likelihood of observed and latent data.

Usage

nbh_vit(count, TRANS, alpha, beta)

Arguments

count A vector of integers, conceptually representative of the read counts within bins of chromosome.

TRANS Optimized transition probability matrix, a squared matrix of probabilities ($0 \leq p \leq 1$) with row and column length equal to that of alpha and beta and row sum and column sum both equal to 1 (within some numerical deviation of 1e-6).

alpha Optimized shape parameter of the NB as a vector of positive values with length equal to that of beta and the row/column of TRANS.

beta Optimized inverse scale parameter of the NB as a vector of positive values with length equal to that of beta and the row/column of TRANS.

Details

Given a K-state HMM with NB emission (NBH), the goal is to find the latent states corresponding to the observed data that maximize the joint likelihood $lnp(X, Z) = lnp(x_1, \ldots, x_N, z_1, \ldots, z_N)$. The optimal solution is obtained via Viterbi algorithm, which essentially belongs to the more general framework of Dynamic Programming.

Briefly, starting from the second node of the Markov chain, we select state of the first node that maximizes $lnp(x_1, x_2, z_2|z_1)$ for every state of $z_2$. Then, we move on to the next node and the next until reaching to the last node. In the end, we make choice for the state of the last node that together leads to the maximum $lnp(X, Z)$. Finally, we backtrack to find the choices of states in all of the intermediate nodes to form the final solution.

Value

A list containing:

class ML sequence of latent states

logl Log-likelihood corresponding to the latent states class

Note

The function is expected to run after learning the model parameters of HMM using nbh_em and (optionally) disambiguating the multihits using nbh_vit. However, nothing prevents user from running it with a random set of HMM parameters. Also, note that Viterbi algorithm finds the most probable sequence of states, which is not the same as maximizing the posterior probabilities for all the individual latent variables. For instance, a observed data point may be classified as from state 2 in the most probable chain in spite its marginal posterior probability for state 2 is zero.
Author(s)

Yue Li

References


See Also

nbh_init, nbh, nbh.GRanges, nbh_em, nbm_em

Examples

```r
# Simulate data
TRANS_s <- matrix(c(0.9, 0.1, 0.3, 0.7), nrow=2, byrow=TRUE)
alpha_s <- c(2, 4)
beta_s <- c(1, 0.25)
Total <- 100

x <- nbh_gen(TRANS_s, alpha_s, beta_s, Total);

count <- x$count
label <- x$label

Total <- length(count)

# dummy initialization
TRANS0 <- matrix(rep(0.5,4), 2)
alpha0 <- c(1, 20)

beta0 <- c(1, 1)

NIT_MAX <- 50
TOL <- 1e-100

nbh <- nbh_em(count, TRANS0, alpha0, beta0, NIT_MAX, TOL)

map.accuracy <- length(which(max.col(nbh$postprob) == label))/Total

vit <- nbh_vit(count, nbh$TRANS, nbh$alpha, nbh$beta)

vit.accuracy <- length(which(vit$class == label))/Total

# Plots
par(mfrow=c(2,2), cex.lab=1.2, cex.main=1.2)
```
nbm_chk

Check the parameters of the negative binomial mixture model

Description

The function verifies the numerical range and dimension of the NBM parameters alpha, beta, and wght and returns the number of components. It is used in nbm_em before running EM.

Usage

nbm_chk(alpha, beta, wght)

Arguments

alpha Expected a vector of positive values with length equal to that of beta and wght.
beta Expected a vector of positive values with length equal to that of alpha and wght.
wght Expected a vector of probabilities ($0 \leq p \leq 1$) with length equal to that of alpha and beta and summed to 1 (within some numerical deviation of $1e-6$).

Value

N Number of components or equivalently the length of alpha, beta, or wght.

Author(s)

Yue Li

References


### Examples

```r
# two mixing components
wght <- c(0.5, 0.5)
alpha <- c(1, 20)
beta <- c(1, 1)
nbm_chk(alpha, beta, wght)
```

### Description

Given an input read count vector of integers, the function optimizes the parameters for the negative binomial mixture model of K components using expectation conditional maximization.

### Usage

```r
nbm_em(count, alpha, beta, wght, NBM_NIT_MAX = 250, NBM_TOL = 0.01)
```

### Arguments

- **count**: A vector of integers, conceptually representing the read counts within bins of chromosome.
- **alpha**: Initial values for $\alpha_k$ for all K NB.
- **beta**: Initial values for $\beta_k$ for all K NB.
- **wght**: Initial values for $\pi_k$ for all K NB.
- **NBM_NIT_MAX**: Maximum number of EM iterations (Default: 250).
- **NBM_TOL**: Threshold as fraction of increase in likelihood (given the current NBM parameters) comparing with the likelihood from the last iteration. EM for the NBM stops when the improvement is below the threshold (Default: 0.01).

### Details

Given a $K$-NBM, the goal is to maximize the likelihood function with respect to the parameters comprising of $\alpha_k$ and $\beta_k$ for the K NB components and the mixing coefficients $\pi_k$, which are the priors $p(z = k)$. Because there is no analytical solution for the maximum likelihood (ML) estimators of the above quantities, a modified EM procedures called Expectation Conditional Maximization is employed (Meng and Rubin, 1994).

In E-step, the posterior probability is evaluated using NB density functions with initialized $\alpha_k$, $\beta_k$, and $\pi_k$. In the CM step, $\pi_k$ is evaluated first followed by Newton updates of $\alpha_k$ and $\beta_k$. EM iteration terminates when the percetnage of increase of log likelihood drop below `NBM_TOL`, which is deterministic since EM is guaranteed to converge. For more details, please see the manuscript of RIPSeeker.
Value

A list containing:

- `alpha`: `alpha_k` for all K components of NB.
- `beta`: `beta_k` for all K components of NB.
- `wght`: `pi_k` for all K components of NB.
- `logl`: Log likelihood in each EM iteration.
- `postprob`: Posterior probabilities for each observed data point in the last EM iteration.

Author(s)

Yue Li

References


See Also

- `nbh_init`, `nbh`, `nbh.GRanges`, `nbh_em`

Examples

```r
# Simulate data
TRANS_s <- matrix(c(0.9, 0.1, 0.3, 0.7), nrow=2, byrow=TRUE)
alpha_s <- c(2, 4)
beta_s <- c(1, 0.25)
Total <- 1000
x <- nbh_gen(TRANS_s, alpha_s, beta_s, Total);
N <- 2

cnt <- x$count
label <- x$label

Total <- length(cnt)

# dummy initialization
wght0 <- c(0.5, 0.5)
alpha0 <- c(1, 20)
beta0 <- c(1, 1)

NIT_MAX <- 50
TOL <- 1e-100
```
plotCoverage

# initialize param with nbm

nbm <- nbm_em(cnt, alpha0, beta0, wght0, NIT_MAX, TOL)

map.accuracy <- length(which(max.col(nbm$postprob) == label))/Total

print(map.accuracy)

plotCoverage      Plot read coverage for a GRanges object

Description

An internal function used by plotStrandedCoverage to plot read counts within each fixed bin across the entire chromosome.

Usage

plotCoverage(x, plotLegend = FALSE, legend.cex = 1, ...)

Arguments

x          GRanges object with values slot saved for read counts within the corresponding ranges.
plotLegend  Binary indicator. If TRUE, legend will be plotted on the top left the plot. Legend is expected to be the chromosome name and length, which must be available in the GRange object argument.
legend.cex  Font size of the legend.
...         Extra arguments passed to either the plot or the legend.

Details

The read counts is plotted in blue bars as positive integer across the x-axis as the sorted positions across the chromosome. The plot can be used to examine the overall alignment properties for each chromosome.

Note

Users are not recommended run this function directly but rather via a much more user friendly function plotStrandedCoverage.

Author(s)

Yue Li

References

See Also

plotStrandedCoverage, plot, legend

Examples

# Retrieve system files
extdata.dir <- system.file("extdata", package="RIPSeeker")

bamFiles <- list.files(extdata.dir, ".bam", recursive=TRUE, full.names=TRUE)
bamFiles <- grep("PRC2", bamFiles, value=TRUE)
alignGal <- getAlignGal(bamFiles[1], reverseComplement=TRUE, genomeBuild="mm9")
alignGR <- as(alignGal, "GRanges")
alignGRList <- GRangesList(as.list(split(alignGR, seqnames(alignGR))))

binSize <- 1000
binGR <- binCount(alignGRList$chrX, binSize)

plotCoverage(binGR, plotLegend=TRUE)

plotStrandedCoverage  Plot strand-specific read coverage for a GRanges object

Description

Plot read counts within fixed bin across the entire chromosome.

Usage

plotStrandedCoverage(gr, binSize = 1000, plotLegend = FALSE, ylim, ...)

Arguments

gr  
GRanges object containing the alignments.

binSize  
Integer indicate the size of the bin used to compute and plot the read counts.

plotLegend  
Binary indicator. If TRUE, legend will be plotted on the top left the plot. Legend is expected to be the chromosome name and length, which must be available in the GRanges object argument.

ylim  
A two element scale on the y-axis, indicating the maximum read counts on the + and - strand to be plotted (e.g., ylim=c(-200, 200)).

...  
Extra arguments passed to plotCoverage.
Details

Read count on + and - strand are displayed as red and blue bars on the positive and negative y-axis, respectively. The x-axis indicates the positions across the chromosome. The plot can be used to examine for each chromosome the overall alignment properties such as strand specificity (expected in non-strand-specific sequencing) and aggregation of reads.

Author(s)

Yue Li

References


See Also

plotCoverage, plot, legend

Examples

# Retrieve system files
extdata.dir <- system.file("extdata", package="RIPSeeker")

bamFiles <- list.files(extdata.dir, ".bam$", recursive=TRUE, full.names=TRUE)
bamFiles <- grep("PRC2", bamFiles, value=TRUE)
alignGal <- getAlignGal(bamFiles[1], reverseComplement=TRUE, genomeBuild="mm9")
alignGR <- as(alignGal, "GRanges")
alignGRList <- GRangesList(as.list(split(alignGR, seqnames(alignGR))))
binSize <- 1000
plotStrandedCoverage(gr=alignGRList$chrX, binSize=binSize, xlab="", ylab="", plotLegend=TRUE, box.lty=0, legend.cex=2 )

---

randindx

Generates random indexes with a specified probability distribution

Description

Returns an array of T indexes distributed as specified by p (which should be a normalized probability vector).

Usage

randindx(p, Total, NO_CHK)
Arguments

\texttt{p} \quad \text{A row vector of normalized probabilities that dictate the transition probability from the current state to the next state. For example, } p = [0.2, 0.8] \text{ indicates that the current state transitions to state 1 at 0.2 and 2 at 0.8. The current state itself can either be the state 1 or 2.}

\texttt{Total} \quad \text{Total number of states needed to be generated using the input transition vector.}

\texttt{NO_CHK} \quad \text{Check whether the first argument is a valid row vector of normalized probabilities.}

Details

The function is used by \texttt{nbh_gen} to generate random data point based on the user-supplied transition probability matrix.

Value

\texttt{I} \quad \text{Index/Indices or state(s) sampled following the transition.}

Author(s)

Yue Li

References


See Also

\texttt{nbh_gen}

Examples

```r
# Total contains the length of data to simulate
Total <- 100

# number of states
N <- 2

# transition probabilities between states
TRANS <- matrix(c(0.9, 0.1, 0.3, 0.7), nrow=2, byrow=TRUE)

label <- matrix(0, Total, 1)

# Simulate initial state
label[1] <- randindx(matrix(1, ncol=N)/N, 1, 1)

# Use Markov property for the following time index
for(t in 2:Total) {
  label[t] <- randindx(TRANS[label[t-1],], 1, 1)
}
```
Description

This function is the main interface to most essential functions of RIPSeeker package.

Usage

```r
ripSeek(bamPath, cNAME, binSize = NULL, strandType = NULL, paired=FALSE, biomaRt_dataset, goAnno, exportFormat = "gff3", annotateFormat = "txt", annotateType = "TSS", outDir, padjMethod = "BH", logOddCutoff = 0, pvalCutoff = 1, pvalAdjCutoff = 1, eFDRCutoff = 1, ...)
```

Arguments

- **bamPath**: Either a path to all of the bam files or a list of paths to individual BAM files. BED and SAM files are also accepted.
- **cNAME**: An identifier pattern found in the control alignment files. Once specified, these files will be used as control and the remaining files as RIP for discriminative analysis (see `seekRIP`).
- **binSize**: Size to use for binning the read counts across each chromosome. If NULL, optimal bin size within a range (default: minBinSize=200, maxBinSize=1200) will be automatically selected (see `selectBinSize`).
- **strandType**: Type of strand can be +, -, or * as in GAlignments, GAlignmentPairs, or GRanges (see `GenomicRanges`).
- **paired**: Binary to indicate whether the library is paired-end (TRUE) or single-end (FALSE by default) (see `getAlignGal`).
- **biomaRt_dataset**: The dataset name used in biomaRt for retrieving genomic information for a given species name (see `annotateRIP`).
- **goAnno**: GO dataset name used for GO enrichment analysis (See `annotateRIP`).
- **exportFormat**: Format to export the RIP predictions. The commonly used ones are GFF and BED, which can be directly imported as a track to a genomic viewer such as Integrative Genomic Viewer, SAVANT or UCSC browser.
- **annotateFormat**: Format to export the annotated RIP predictions. The default "txt" is a tab-delimited format, recommended for viewing in Excel.
- **annotateType**: Type of genomic information in association with the RIP predictions that can be retrieved from Ensembl database (Default: TSS; See `annotateRIP`).
- **outDir**: Output directory to save the results. The output data include ...
- **padjMethod**: Method to adjust multiple testing (Benjamini-Hocherge method by default).
- **logOddCutoff**: Threshold for the log odd ratio of posterior for the RIP over the background states (See `seekRIP`). Only peaks with logOdd score greater than the logOddCutoff will be reported. Default: 1.
pvalCutoff: Threshold for the p-value for the logOdd score. Only peaks with p-value less than the pvalCutoff will be reported. Default: 1 (i.e. no cutoff).

pvalAdjCutoff: Threshold for the adjusted p-value for the logOdd score. Only peaks with adjusted p-value less than the pvalAdjCutoff will be reported. Default: 1 (i.e. no cutoff).

eFDRcutoff: Threshold for the empirical false discovery rate (eFDR). Only peaks with eFDR less than the eFDRcutoff will be reported. Default: 1 (i.e. no cutoff).

Details

This is the main front-end function of RIPSeeker and in many cases the only function that users need to get RIP predictions and all relevant information.

Value

A list is returned with the following items:

mainSeekOutputRIP

A (inner) list comprising three items:

nbhGRList: GRangesList of the HMM trained parameters for each chromosome on RIP.

alignGal, alignGalFiltered: GAlignments objects of the RIP alignment outputs from combineAlignGals and disambiguateMultihits, respectively. The former may contain multiple alignments due to the same reads whereas the latter contains a one-to-one mapping from read to alignment after disambiguating the multihits.

mainSeekOutputCTL

Same as mainSeekOutputRIP but for the control library (if available).

RIPGRList

The results as GRangesList generated from the RIP peak detection. Each list item represents the RIP peaks on a chromosome accompanied with statistical scores including (read) count, logOddScore, pval, pvalAdj, eFDR for the RIP and control (if available). Please refer to seekRIP for more details.

annotatedRIPGR

If annotatedRIPGR is TRUE, the additional genomic information will be retrieved according to the genomic coordinates of the peaks in RIPGRList. The results are saved in this separate GRanges object as the final results that user will find the most useful.

Note

You may only want to know the expression/abundance of known transcripts/genes or the foldchange between two conditions. In that case, use rulebaseRIPSeek and computeRPKM, respectively. Both singl-end and paired-end alignments are supported in these functions.

Author(s)

Yue Li
References


The RIPSeeker manuscript has been submitted to NAR for review.

See Also

rulebaseRIPSeek

Examples

if(interactive()) { # need internet connection

# Retrieve system files
extdata.dir <- system.file("extdata", package="RIPSeeker")

bamFiles <- list.files(extdata.dir, ".bam$", recursive=TRUE, full.names=TRUE)
bamFiles <- grep("PRC2", bamFiles, value=TRUE)
cNAME <- "SRR039214" # specify control name

# output file directory
outDir <- paste(getwd(), "ripSeek_example", sep="/"")

# Parameters setting
binSize <- NULL # automatically determine bin size
minBinSize <- 10000 # min bin size in automatic bin size selection
maxBinSize <- 12000 # max bin size in automatic bin size selection
multicore <- TRUE # use multicore
strandType <- "-" # set strand type to minus strand

biomart <- "ENSEMBL_MART_ENSEMBL" # use archive to get ensembl 65
dataset <- "mmusculus_gene_ensembl" # mouse dataset id name
host <- "dec2011.archive.ensembl.org" # use ensembl 65 for annotation
goAnno <- "org.Mm.eg.db"

# run main function ripSeek to predict RIP
seekOut <- ripSeek(bamPath=bamFiles, cNAME=cNAME,
binSize=binSize, minBinSize = minBinSize,
maxBinSize = maxBinSize, strandType=strandType,
outDir=outDir, silentMain=FALSE,
verbose=TRUE, reverseComplement=TRUE, genomeBuild="mm9",
biomart=biomart, host=host,
biomaRt_dataset = dataset,
goAnno = goAnno,
uniqueHit = TRUE, assignMultihits = TRUE,
rerunWithDisambiguatedMultihits = TRUE, multicore=multicore)

# visualization
viewRIP(seekOut$RIPGRList$chrX, seekOut$mainSeekOutputRIP$alignGalFiltered,
rulebaseRIPSeek

Compute the RPKM and foldchange between two conditions for the annotated genes

Description

The function takes alignments in two conditions (with replicates) as input and computes the gene expression in the two conditions in the unit of RPKM (reads per kilobase of exon per million mapped reads) or FPKM for paired-end alignments (where "F" stands for the fragment the mate-pairs are derived from), and then the foldchange ratio of the RPKM of each gene in RIP or treatment condition in general over control condition. The control files (i.e. the denominator in the foldchange ratio) is specified by user in the "cNAME" argument.

Usage

rulebaseRIPSeek(bamFiles, cNAME, featureGRanges, rpkmCutoff = 0.4, fcCutoff = 3, moreRIPGeneInfo = TRUE, idType = "ensembl_transcript_id", myMin = .Machine$double.xmin, saveRData, ...)

Arguments

bamFiles A list of one or more BAM/SAM/BED alignment files.
cNAME An identifier pattern found in the control alignment files. Once specified, these files will be used as control as the denominator of the foldchange ratio and the remaining files as RIP, the numerator of the foldchange ratio.
featureGRanges GRanges of features as an optional argument for function to compute RPKM/FPKM just for those features without retrieving online annotations.
rpkmCutoff Cutoff for RPKM in RIP above which the genes will be reported if the fcCutoff is also satisfied (Default: 0.4).
fCutoff Cutoff for foldchange in RIP relative to the control above which the genes will be reported if the rpkmCutoff is also satisfied (Default: 3).
moreRIPGeneInfo Binary indicator to indicate whether to download more information for each genes/transcripts rather than having only the gene/transcript IDs (Default: TRUE).
idType A character string that specifies the type of the annotations, which can "ensembl_transcript_id" (Default), "ensembl_gene_id", "ucsc", etc. Refer to listFilters for more information.
myMin Add a small value to both the numerator and denominator as "pseudocount" to prevent the case where the denominator is zero and the ratio becomes infinity regardless the value of the numerator (Default: .Machine$double.xmin).
saveRData Path of output RData and tab-delim results.
... Extra arguments passed to computeRPKM and/oruseMart.
Details

The function uses `computeRPKM` to download annotation and compute RPKM/FPKM of the annotated genes in the list of files. The alignments file are separated into control as identified by the "cNAME" and the RIP (or any treatment) that do not have the cNAME in their file names. The alignments in either group are pooled together. If `moreRIPGeneInfo` is specified, the function will query the Ensembl database. The chromosome ID in the numerical format used in Ensembl is prefixed with "chr" and the strand 1 and -1 converted to + and - for convenience.

Value

A list containing the following items:

- `nRPKM`: RPKM of genes in RIP or treatment condition ('n' stands for numerator in the foldchange ratio).
- `dRPKM`: RPKM of genes in control condition ('d' stands for denominator in the foldchange ratio).
- `rpkmDF`: Data frame containing read count, RPKM for the RIP (or treatment) and control, foldchange, and optional gene information including "chromosome_name", "start_position", "end_position", "strand", "external_gene_id", "ensembl_transcript_id", "ensembl_gene_id", "ucsc", "description".
- `rpkmCutoff`: Cutoff used for RPKM as book keeping value.
- `fcCutoff`: Cutoff used for foldchange as book keeping value.
- `featureGRanges`: `GRanges` object of the features for which the RPKM and foldchange are computed.

Note

Also works for RNA-seq alignments.

Author(s)

Yue Li

References


**scoreMergedBins**

Average log odd scores over bins being merged into a single region

**Description**

Sum, normalize the read counts, and average the logOdd score over the bins being merged into a single enriched region.

**Usage**

```r
scoreMergedBins(findOverlapsHits, unmergedGRAll, mergedGRAll)
```

**Arguments**

- **findOverlapsHits**
  - Output from `findOverlaps` as two columns indices with the first column containing the indices for unmerged GRanges and the second column the indices of the matched merged GRanges.

- **unmergedGRAll**
  - GRanges before merging.

- **mergedGRAll**
  - GRanges after merging.
Details

The consecutive RIP-bins predicted by the Viterbi function (See nbh_vit) are merged into a single RIP region. An aggregate RIPScore as the averaged RIPScores over the associated merged bins is assigned to each merged RIP region. In the RIPSeeker workflow, the averaged RIPScore then becomes the representative score for the region and subject to significance test carried out in seekRIP.

Value

A merged GRanges each with scores including summed read count, averaged log odd scores, and FPK (fragment per kilobase of region length). The latter score represents a normalized read count.

Note

This function is expected to be called only from logScoreWithoutControl and logScoreWithControl.

Author(s)

Yue Li

See Also

seekRIP, computeLogOdd logScoreWithControl, logScoreWithoutControl

Examples

```r
if(interactive()) { # see example in seekRIP
  # Retrieve system files
  extdata.dir <- system.file("extdata", package="RIPSeeker")
  bamFiles <- list.files(extdata.dir, ".bam$", recursive=TRUE, full.names=TRUE)
  bamFiles <- grep("PRC2", bamFiles, value=TRUE)
  bamFiles <- grep("SRR039214", bamFiles, value=TRUE, invert=TRUE)

  # Parameters setting
  binSize <- 1e5 # use a large fixed bin size for demo only
  multicore <- FALSE # use multicore
  strandType <- "-" # set strand type to minus strand

  mainSeekOutputRIP <- mainSeek(bamFiles=grep(pattern="SRR039214", bamFiles, value=TRUE, invert=TRUE),
                                binSize=binSize, strandType=strandType,
                                reverseComplement=TRUE, genomeBuild="mm9",
                                uniqueHit = TRUE, assignMultihits = TRUE,
                                rerunWithDisambiguatedMultihits = TRUE,
                                multicore=multicore, silentMain=FALSE, verbose=TRUE)

  nbhGRRIP <- mainSeekOutputRIP$nbhGRList$chrX

  logOddScore <- computeLogOdd(nbhGRRIP)

  values(nbhGRRIP) <- cbind(as.data.frame(values(nbhGRRIP)), logOddScore)
}
```
seekRIP <- which(values(nbhGRRIP)$viterbi_state == 2)
unmergedRIP <- nbhGRRIP[enrichIdx]
mergedRIP <- reduce(unmergedRIP, min.gapwidth = median(width(unmergedRIP)))
overlapIdx <- findOverlaps(mergedRIP, unmergedRIP)

# a list with query hits as names and subject hits as items
findOverlapsHits <- split(overlapIdx, queryHits(overlapIdx))

# get the score for the first merged region
x <- scoreMergedBins(findOverlapsHits[[1]], unmergedRIP, mergedRIP)

# get scores for all of the merged regions
mergedRIPList <- lapply(split(overlapIdx, queryHits(overlapIdx)),
scoreMergedBins, unmergedRIP, mergedRIP)

names(mergedRIPList) <- NULL
mergedRIP <- do.call(c, mergedRIPList)

# logOddScore is the averaged logOddScore across merged bins
mergedRIP

seekRIP  
Identify significant peaks

Description

Based on the posteriors derived from HMM by mainSeek, find the significant RIP regions derived from merging the adjacent RIP bins. The significance test makes use of the log odd ratio of the posteriors for RIP over the background states.

Usage

seekRIP(nbhGRRIP, nbhGRCTL = NULL, padjMethod = "BH", logOddCutoff = -Inf, pvalCutoff = 1, pvalAdjCutoff = 1, eFDRcutoff = 1)

Arguments

nbhGRRIP  
GRanges object for the RIP library created from mainSeek containing the posterior probabilities of the hidden states for each observed read count.

nbhGRCTL  
An optional argument as a GRanges object for the control library created from mainSeek containing the posterior probabilities of the hidden states for each observed read count.

padjMethod  
Method used to adjust multiple testing performed in p.adjust (Default: "BH").

logOddCutoff  
Threshold for the log odd ratio of posterior for the RIP over the background states (See seekRIP). Only peaks with logOdd score greater than the logOddCutoff will be reported. Default: -Inf (i.e. no cutoff).
pvalCutoff  Threshold for the adjusted p-value for the logOdd score. Only peaks with adjusted p-value less than the pvalAdjCutoff will be reported. Default: 1 (i.e. no cutoff).

pvalAdjCutoff  Threshold for the adjusted p-value for the logOdd score. Only peaks with adjusted p-value less than the pvalAdjCutoff will be reported. Default: 1 (i.e. no cutoff).

eFDRcutoff  Threshold for the empirical false discovery rate (eFDR). Only peaks with eFDR less than the pvalAdjCutoff will be reported. Default: 1 (i.e. no cutoff).

Details

The RIPScore is computed in `computeLogOdd` as the log odd ratio of the posterior for the RIP state ($z_i = 2$) over the posterior for the background state ($z_i = 1$) in RIP library. When control is available, the RIPScore is updated by the difference between the RIPScores between RIP and control. The adjacent bins with hidden states predicted by `nbh_vit` as the enriched state (corresponding to the NB with larger mean) are merged. The RIPScores are averaged over the merged bins. To assess the statistical significance of the RIPScore for each region, we assume that the RIPScore follows a Gaussian (Normal) distribution with mean and standard deviation estimated using the RIPScores over all of the bins. The rationale is based on the assumption that most of the RIPScores correspond to the background state and together contribute to a stable estimate of the test statistics (TS) and p-value computed using the R built-in function `pnorm`.

The p-value is adjusted by `p.adjust` with BH method by default. The same procedure is applied optionally to the control library. Only when the control is available, is an empirical false discovery rate (eFDR) estimated based on the idea of "sample swap" inspired by MACS (a ChIP-seq algorithm from Zhang et al. (2008). At each p-value, RIPSeeker finds the number of significant RIP-regions over control (CTL) based on pvalRIP and the number of significant control regions over RIP based on pvalCTL. The eFDR is defined as the ratio of the number of "RIP" (false positive) regions identified from CTL-RIP comparison over the number of RIP regions from the RIP-CTL comparison. The maximum value for eFDR is 1 and minimum value for eFDR is max(p-value, 0). The former takes care of the case where the numerator is bigger than the denominator, and the latter for zero numerator.

Value

`GRanges` object containing the merged bins with values slot saved for RIPScore (lodOdd), p-value (pval), adjusted p-value (pvalAdj) for RIP and optionally for control.

Note

Internal function used by `ripSeek`.

Author(s)

Yue Li

References

selectBinSize

Select optimal bin size based on Shimazaki formula

Description
The function iteratively estimates the cost of increasing bin size within a defined range and finally selects the bin size with minimum cost.

Usage
```r
selectBinSize(alignGR, minBinSize, maxBinSize = 1000, increment = 5, getFullResults = FALSE)
```

Examples
```r
if(interactive()) {
  # Retrieve system files
  extdata.dir <- system.file("extdata", package="RIPSeeker")
  bamFiles <- list.files(extdata.dir, "\.".bam\", recursive=TRUE, full.names=TRUE)
  bamFiles <- grep("PRC2", bamFiles, value=TRUE)
  # Parameters setting
  binSize <- 1e5 # use a large fixed bin size for demo only
  multicore <- FALSE # use multicore
  strandType <- "-" # set strand type to minus strand
  mainSeekOutputRIP <- mainSeek(bamFiles=grep(pattern="SRR039214", bamFiles, value=TRUE, invert=TRUE), binSize=binSize, strandType=strandType, reverseComplement=TRUE, genomeBuild="mm9", uniqueHit = TRUE, assignMultihits = TRUE, rerunWithDisambiguatedMultihits = FALSE, multicore=multicore, silentMain=FALSE, verbose=TRUE)
  mainSeekOutputCTL <- mainSeek(bamFiles=grep(pattern="SRR039214", bamFiles, value=TRUE, invert=FALSE), binSize=binSize, strandType=strandType, reverseComplement=TRUE, genomeBuild="mm9", uniqueHit = TRUE, assignMultihits = TRUE, rerunWithDisambiguatedMultihits = FALSE, multicore=multicore, silentMain=FALSE, verbose=TRUE)
  # with control
  ripGR.wicontrol <- seekRIP(mainSeekOutputRIP$nbhGRList$chrX, mainSeekOutputCTL$nbhGRList)
  # without control
  ripGR.wocontrol <- seekRIP(mainSeekOutputRIP$nbhGRList$chrX)
}
```
selectBinSize

Arguments

alignGR  GRanges object of alignments on a single chromosome
minBinSize Minimum bin size to start with (Default: 5 * read length)
maxBinSize Maximum bin size to end with (Default: 1000).
increment Number of bases to increment the bin size in searching for the optimal bin size within the defined range (Default: 5).
getFullResults Binary indicator. If TRUE, the optimal bin size (with the minimum cost), minimum cost, and all of the bin sizes considered and their costs are returned. If FALSE, only the optimal bin size is returned.

Details

Based on the preprocessed alignments for a chromosome, RIPSeeker divides the chromosome into bins of equal size \( b \) and compute the number of reads that \( b \) needs to be determined either empirically (e.g., based on the gel-selected length of the RNA fragment) or computationally. If the bin size is too small, the read counts fluctuates greatly, making it difficult to discern the underlying read count distribution. Additionally, input size to HMM increases as bin size decreases. A very small bin size results in a very long Markov chain of read counts to model, making the computation inefficient. On the other hand, if a bin size is too large, resolution becomes poor. Consequently, one cannot detect the local RIP region with subtle but intrinsic difference from the background, and the RIP regions tend to be too wide for designing specific primer for validation.

Intuitively, selecting an appropriate bin size for each chromosome is metaphorically equivalent to choosing an optimal intervals for building a histogram (Song, 2011). Here we implement the algorithm developed by Shimazaki and Shinomoto (2007), which is based on the goodness of the fit of the time histogram to estimate the rate of neural response of an animal to certain stimuli in a spike-in experiment. This approach has been successfully applied in a recently developed ChIP-seq program (Song and Smith, 2011). Algorithm 1 describes the pseudocode adapted from Shimazaki and Shinomoto (2007) that iteratively estimates the cost \( C \) of increasing bin size \( b \) within a defined range is outlined as follows.

For \( b = \text{minBinSize} \) to \( \text{maxBinSize} \); do
1. Divide chromosome sequence into \( N \) bins of width \( b \).
2. Count number of read counts \( x_i \) that enter the \( i \)’th bin.
3. Compute: \( \bar{x} = \frac{1}{N} \sum_{i=1}^{N} x_i \) and \( v = \frac{1}{N} \sum_{i=1}^{N} (x_i - \bar{x})^2 \).
4. Compute: \( C(b) = \frac{2\bar{x} - v}{b} \).
End For

Choose \( b \) that minimize \( C(b) \)

Value

When \( \text{getFullResults} \) is TRUE, return a list containing:
- bestBinSize the optimal bin size (with the minimum cost)
- minCosts cost of the optimal bin size
- binSizes all of the bin sizes considered
- costs all of the costs

When \( \text{getFullResults} \) is FALSE, only the optimal bin size (bestBinSize) is returned.
selectBinSize

Author(s)

Yue Li

References


See Also

evalBinSize, binCount

Examples

# Retrieve system files
extdata.dir <- system.file("extdata", package="RIPSeeker")

bamFiles <- list.files(extdata.dir, ".bam $", recursive=TRUE, full.names=TRUE)
bamFiles <- grep("PRC2", bamFiles, value=TRUE)

alignGal <- getAlignGal(bamFiles[1], reverseComplement=TRUE, genomeBuild="mm9")

alignGR <- as(alignGal, "GRanges")

alignGRList <- GRangesList(as.list(split(alignGR, seqnames(alignGR))))

minBinSize <- 200

maxBinSize <- 1200

gr <- alignGRList$chrX

b <- selectBinSize(gr, minBinSize, maxBinSize, increment=100, getFullResults=TRUE)

plot(b$binSizes, b$costs)

chrname <- as.character(runValue(seqnames(gr)))

chrlen <- seqlengths(gr)[chrname]

legend("topright", box.lty=0,

sprintf("%s: 1-%d; Total mapped reads: %d; Optimal bin size = %d bp", chrname, chrlen, length(gr), b$bestBinSize))
**statdis**

*Returns the stationary distribution of a Markov chain.*

**Description**

Given a transition matrix $A$, returns the stationary distribution of a Markov chain by computing the eigen vectors of $A$.

**Usage**

```r
statdis(A)
```

**Arguments**

- **A**: Transition probability matrix, a squared matrix of probabilities ($0 \leq p \leq 1$) with row and column length equal to that of alpha and beta and row sum and column sum both equal to 1 (within some numerical deviation of 1e-6).

**Value**

- **w**: Stationary weights for the distributions of K components based on the transition probability matrix.

**Author(s)**

Yue Li

**References**


**See Also**

`nbh_em`

**Examples**

```r
# Simulate data
TRANS_s <- matrix(c(0.9, 0.1, 0.3, 0.7), nrow=2, byrow=TRUE)
alpha_s <- c(2, 4)
beta_s  <- c(1, 0.25)
Total <- 100
x <- nbh_gen(TRANS_s, alpha_s, beta_s, Total);

count <- x$count
label <- x$label

Total <- length(count)
# dummy initialization
```
TRANS0 <- matrix(rep(0.5,4), 2)

alpha0 <- c(1, 20)

beta0 <- c(1, 1)

NIT_MAX <- 50

TOL <- 1e-100

nbh <- nbh_em(count, TRANS0, alpha0, beta0, NIT_MAX, TOL)

map.accuracy <- length(which(max.col(nbh$postprob) == label))/Total

vit <- nbh_vit(count, nbh$TRANS, nbh$alpha, nbh$beta)

vit.accuracy <- length(which(vit$class == label))/Total

# Plot the marginal distribution (in the stationary regime)
# Compute negative binomial distributions for all model states

t <- 0:max(count)

tmp <- nbh_em(t, nbh$TRANS, nbh$alpha, nbh$beta, 1)

dens <- tmp$dens

w <- statdis(nbh$TRANS)

# Plot estimate of marginal probabilities
marprob <- apply(t(dens) * (t(w) %*% matrix(1, ncol=length(t))), 2, sum)

plot(t, marprob, pch=8, col="blue", main="Estimated marginal distribution")

# Plot empirical estimated probabilities
dhist <- matrix(0, ncol=length(t))

for(i in t){
dhist[1+i] <- sum(count == i)/Total
}

points(t, dhist, pch=3, col="red")

---

viewRIP

**Visualize peaks from UCSC genome browser.**

**Description**

Upload alignments, peaks, statistical scores to UCSC genome browser for comparative visualization of the results and data available in the UCSC database.

**Usage**

```r
viewRIP(seekedRIP, alignGR, alignGRCTL,
        binGR = seekedRIP, scoreType = "eFDR",
        cutoffLine = 0.001, displayALLChr = FALSE, ...)
```
Arguments

- **seekedRIP** (`GRangesList`) obtained from `ripSeek`. Each list item represents the RIP peaks on a chromosome accompanied with statistical scores including (read) count, logOddScore, pval, pvalAdj, eFDR for the RIP and control (if available). Please refer to `seekRIP` for more details.

- **alignGR** (`GRanges`) of read alignments for the RIP.

- **alignGRCTL** (`GRanges`) of read alignments for the control.

- **binGR** (`GRanges`) containing read count column corresponding to the peaks. By default, `alignGR` is used as `binGR` to display the read count in RIP condition.

- **scoreType** Type of statistical score to display as another track in the browser (Default: eFDR). eFDR/pval/pvalAdj is displayed at -log10 scale.

- **cutoffLine** Draw a cutoffline in the browser to indicate the significance level above which the peaks are considered significant.

- **displayALLChr** Binary indicator when TRUE upload and display the information for only one chromosome rather than upload all chromosomes (Default: TRUE).

Details

The function is a wrapper function of `browserSession`, `track`, and `browserView`.

Note

If input contain multiple chromosomes, then multiple browser window will be open to display each chromosome. A more user-friendly way is to upload all of the information to UCSC and open a single browser for visualization, which may become one of the new features in future release.

Author(s)

Yue Li

References

Michael Lawrence, Vince Carey and Robert Gentleman (). rtracklayer: R interface to genome browsers and their annotation tracks. R package version 1.16.3.

See Also

`ripSeek`, `browserSession`, `track`, `browserView`

Examples

```r
if(interactive()) { # need internet connection
  # Retrieve system files
  extdata.dir <- system.file("extdata", package="RIPSeeker")
  bamFiles <- list.files(extdata.dir, ".bam$", recursive=TRUE, full.names=TRUE)
  bamFiles <- grep("PRC2", bamFiles, value=TRUE)
  cNAME <- "SRR039214" # specify control name
```
# Parameters setting
binSize <- NULL # automatically determine bin size
minBinSize <- 10000 # min bin size in automatic bin size selection
maxBinSize <- 12000 # max bin size in automatic bin size selection
multicore <- TRUE # use multicore
strandType <- "-" # set strand type to minus strand

################ run main function ripSeek to predict RIP ###############
seekOut <- ripSeek(bamPath=bamFiles, cNAME=cNAME,
                   binSize=binSize, minBinSize = minBinSize,
                   maxBinSize = maxBinSize, strandType=strandType,
                   silentMain=TRUE, verbose=FALSE,
                   reverseComplement=TRUE, genomeBuild="mm9",
                   uniqueHit = TRUE, assignMultihits = TRUE,
                   rerunWithDisambiguatedMultihits = TRUE, multicore=multicore)

################ visualization ###############
viewRIP(seekOut$RIPGRList$chrX, seekOut$mainSeekOutputRIP$alignGalFiltered,
         seekOut$mainSeekOutputCTL$alignGalFiltered, scoreType="eFDR")
Index

*Topic Hidden Markov model  
RIPSeeker-package, 2
*Topic RIP-seq  
RIPSeeker-package, 2
*Topic high-throughput sequencing analysis  
RIPSeeker-package, 2
*Topic package  
RIPSeeker-package, 2
addDummyProb, 3, 31, 32
addPseudoAlignment, 5
annotatePeakInBatch, 7
annotateRIP, 6, 53
binCount, 4, 9, 21, 64
browserSession, 67
browserView, 67
cdsBy, 15, 16, 58
combineAlignGals, 5, 10, 12, 15, 16, 23, 25, 54, 58
combineRIP, 11
computeLogOdd, 13, 19, 26, 28, 59, 61, 62
computeRPKM, 14, 54, 56, 57
countOverlaps, 12
disambiguateMultihits, 17, 23, 24, 54
downstreamIntervals, 30, 54
draw, 51
empiricalFDR, 18, 62
evalBinSize, 9, 20, 64
exonsBy, 15, 16, 58
export, 21, 22
exportGRanges, 6, 21
findOverlaps, 58
fiveUTRsByTranscript, 15, 16, 58
GAAlignments, 25, 54
galp2gal, 15, 22, 24
GenomicRanges, 53
getAlignGal, 10, 11, 17, 18, 22, 23, 29, 30, 53
getAnnotation, 6, 7
getEnrichedGO, 6, 7
GRanges, 6, 12, 21, 26, 28, 50, 57, 60, 61, 67
GRangesList, 54, 67
import, 5, 11, 12, 23, 25
intronsByTranscript, 15, 16, 58
legend, 49–51
listDatasets, 6
listFilters, 15, 56
logScoreWithControl, 14, 19, 25, 28, 59, 62
logScoreWithoutControl, 14, 26, 27, 59, 62
mainSeek, 18, 26, 27, 29, 31, 32, 54, 60
mainSeekSingleChrom, 4, 9, 17, 18, 30, 31, 33, 35, 37
makeDbFromBiomart, 15, 16, 58
makeDbFromUCSC, 15, 16, 58
mclapply, 30
nbh, 17, 32, 33, 34, 35, 37, 39, 43, 45, 48
nbh.GRanges, 33, 34, 37, 39, 43, 45, 48
nbh.integer, 33–35, 35
nbh_chk, 37, 47
nbh_em, 29, 31, 32, 36, 37, 38, 41–45, 48, 65
nbh_gen, 40, 52
nbh_init, 36, 39, 42, 45, 48
nbh_vit, 26, 28, 36, 39, 44, 44, 59, 61
nbm_chk, 37, 41, 46
nbm_em, 35, 39, 42, 43, 45–47, 47
p.adjust, 26–28, 60, 61
plot, 49–51
plotCoverage, 49, 50, 51
plotStrandedCoverage, 49, 50, 50
pnorm, 26, 28, 61
randindx, 41, 51
RangedSummarizedExperiment, 15
readAlignmentPairs, 5, 11, 16, 23, 25, 58
readAlignments, 5, 11, 16, 23, 25, 58
reduce, 12
ripSeek, 3, 11, 12, 18, 29, 30, 32, 53, 61, 62, 67

69
RIPSeeker, 15
RIPSeeker (RIPSeeker-package), 2
RIPSeeker-package, 2
rulebaseRIPSeek, 2, 3, 54, 55, 56

scanBamFlag, 24
ScanBamParam, 16, 58
scoreMergedBins, 14, 19, 26, 28, 58, 62
seekRIP, 12, 14, 17, 19, 26, 28, 30, 53, 54, 59, 60, 60, 67
selectBinSize, 9, 17, 21, 31, 32, 53, 62
SeqinfoForUCSCGenome, 29
statdis, 65
summarizeOverlaps, 15, 16, 58

threeUTRsByTranscript, 15, 16, 58
track, 67

useMart, 6, 7, 15, 16, 56, 58

viewRIP, 66