Package ‘derfinder’

March 26, 2015

Type Package

Title Annotation-agnostic differential expression analysis of RNA-seq data at base-pair resolution

Version 1.1.17

Date 2015-03-12

Depends R(>= 3.2)

Imports AnnotationDbi (>= 1.27.9), BiocParallel, bumphunter (>= 1.7.6), derfinderHelper (>= 1.1.0), GenomeInfoDb (>= 1.3.3), GenomicAlignments, GenomicFeatures, GenomicFiles, GenomicRanges (>= 1.17.40), Hmisc, IRanges (>= 1.99.28), qvalue, Rsamtools, rtracklayer, S4Vectors (>= 0.2.3)

Suggests biovizBase, devtools (>= 1.6), derfinderData (>= 0.99.0), ggplot2, knitr, knitrBootstrap (>= 0.9.0), rmarkdown (>= 0.3.3), testthat, TxDb.Hsapiens.UCSC.hg19.knownGene

VignetteBuilder knitr

Description Annotation-agnostic differential expression analysis of RNA-seq data by calculating F-statistics at base-pair resolution

License Artistic-2.0

LazyData true

URL https://github.com/lcolladotor/derfinder

BugReports https://github.com/lcolladotor/derfinder/issues

biocViews DifferentialExpression, Sequencing, RNASeq, Software

NeedsCompilation no

Author Leonardo Collado-Torres [aut, cre], Alyssa C. Frazee [aut], Andrew E. Jaffe [aut], Jeffrey T. Leek [aut, ths]

Maintainer Leonardo Collado-Torres <lcollado@jhu.edu>
**R topics documented:**

- derfinder-package ................................................................. 2
- advancedArg ............................................................................. 3
- analyzeChr ............................................................................... 4
- annotateRegions ....................................................................... 6
- calculatePvalues ....................................................................... 7
- calculateStats ........................................................................... 10
- coerceGR .................................................................................. 11
- collapseFullCoverage ................................................................. 12
- coverageToExon ......................................................................... 13
- createBw ................................................................................... 15
- createBwSample ......................................................................... 16
- extendedMapSeqlevels ................................................................. 17
- filterData .................................................................................... 18
- findRegions ................................................................................ 20
- fullCoverage .............................................................................. 22
- genomeData ............................................................................... 23
- genomeDataRaw ......................................................................... 24
- genomeFstats ............................................................................. 24
- genomeInfo ................................................................................ 25
- genomeRegions ......................................................................... 26
- genomicState .............................................................................. 26
- getRegionCoverage ..................................................................... 27
- loadCoverage ............................................................................. 28
- makeGenomicState ..................................................................... 30
- makeModels ............................................................................... 32
- mergeResults ............................................................................. 33
- preprocessCoverage .................................................................. 35
- rawFiles .................................................................................... 37
- regionMatrix ............................................................................. 38
- sampleDepth .............................................................................. 39

**Index**

- derfinder-package 41

---

**Description**

Fast differential expression analysis of RNA-seq data at base-pair resolution from multiple samples. The analysis pipeline involves loading the sample BAM files using `rawFiles` and `loadCoverage`, preprocessing the data by using `preprocessCoverage`, calculating the F-statistics (while adjusting for some confounders) using `makeModels` and `calculateStats`, calculating the p-values and finding the regions of interest using `calculatePvalues`, and finally annotating them using `matchGenes` from the bumphunter package.
advancedArg

Author(s)
Leonardo Collado-Torres <lcollado@jhu.edu>

References

Rafael A. Irizarry, Martin Aryee, Hector Corrada Bravo, Kasper D. Hansen and Harris A. Jaffee.

advancedArg  List advanced arguments

Description
Find in GitHub the documentation for the advanced arguments of a given function.

Usage
advancedArg(fun, package = "derfinder", browse = interactive())

Arguments
  fun                     The name of a function(s) that has advanced arguments in package.
  package                The name of the package where the function is stored. Only 'derfinder', 'derfinderPlot', and 'regionReport' are accepted.
  browse                 Whether to open the URLs in a browser.

Details
If you are interested on the default options used for functions that run on multiple cores, check https://github.com/lcolladotor/derfinder/blob/master/R/utils.R Note that in general, SnowParam is more memory efficient than link[BiocParallel]MulticoreParam. If you so desire, use your favorite cluster type by specifying BPPARAM.

Value
A vector of URLs with the GitHub search queries.

Author(s)
Leonardo Collado-Torres
Examples

```r
## Open the advanced argument docs for loadCoverage()
if(interactive()) {
    advancedArg('loadCoverage')
} else {
    (advancedArg('loadCoverage', browse = FALSE))
}
```

---

**analyzechr**  
*Run the derfinder analysis on a chromosome*

**Description**

This is a major wrapper for running several key functions from this package. It is meant to be used after `loadCoverage` has been used for a specific chromosome. The steps run include `makeModels`, `preprocessCoverage`, `calculateStats`, `calculatePvalues` and annotating with `annotateTranscripts` and `matchGenes`.

**Usage**

```r
analyzeChr(chr, coverageInfo, models, cutoffPre = 5, cutoffFstat = 1e-08,
           cutoffType = "theoretical", nPermute = 1, seeds = as.integer(gsub("-", 
                          
           "", Sys.Date())) + seq_len(nPermute), groupInfo, txdb = NULL,
           writeOutput = TRUE, runAnnotation = TRUE, lowMemDir = file.path(chr,
                          "chunksDir"), ...)"
```

**Arguments**

- **chr**  
  Used for naming the output files when `writeOutput=TRUE` and the resulting GRanges object.

- **coverageInfo**  
  A list containing a `data.frame` -- coverage -- with the coverage data and a logical Rle -- position -- with the positions that passed the cutoff. This object is generated using `loadCoverage`.

- **models**  
  The output from `makeModels`.

- **cutoffPre**  
  This argument is passed to `preprocessCoverage` (cutoff).

- **cutoffFstat**  
  This is used to determine the cutoff argument of `calculatePvalues` and it’s behaviour is determined by `cutoffType`.

- **cutoffType**  
  If set to `empirical`, the `cutoffFstat` (example: 0.99) quantile is used via `quantile`. If set to `theoretical`, the theoretical `cutoffFstats` (example: 1e-08) is calculated via `qf`. If set to `manual`, `cutoffFstats` is passed to `calculatePvalues` without any other calculation.

- **nPermute**  
  The number of permutations. Note that for a full chromosome, a small amount (10) of permutations is sufficient. If set to 0, no permutations are performed and thus no null regions are used, however, the $regions component is created.
seeds  An integer vector of length nPermute specifying the seeds to be used for each permutation. If NULL no seeds are used.

groupInfo  A factor specifying the group membership of each sample that can later be used with the plotting functions in the derfinderPlot package.

txdb  This argument is passed to annotateTranscripts. If NULL, TxDb.Hsapiens.UCSC.hg19.knownGene is used.

writeOutput  If TRUE, output Rdata files are created at each step inside a directory with the chromosome name (example: 'chr21' if chrnum='21'). One Rdata file is created for each component described in the return section.

runAnnotation  If TRUE annotateTranscripts and matchGenes are run. Otherwise these steps are skipped.

lowMemDir  If specified, each chunk is saved into a separate Rdata file under lowMemDir and later loaded in fstats.apply when running calculateStats and calculatePvalues. Using this option helps reduce the memory load as each fork in bplapply loads only the data needed for the chunk processing. The downside is a bit longer computation time due to input/output.

Arguments passed to other methods and/or advanced arguments.

Details

If you are working with data from an organism different from 'Homo sapiens' specify so by setting the global 'species' and 'chrsStyle' options. For example:

```r
optionsHspecies = \'arabidopsis_thaliana\'
optionsHchrsStyle = \'NCBI\'
```

Value

If returnOutput=TRUE, a list with six components:

- **timeinfo**  The wallclock timing information for each step.
- **optionsStats**  The main options used when running this function.
- **coveragePrep**  The output from preprocessCoverage.
- **fstats**  The output from calculateStats.
- **regions**  The output from calculatePvalues.
- **annotation**  The output from matchGenes.

These are the same components that are written to Rdata files if writeOutput=TRUE.

Author(s)

Leonardo Collado-Torres

See Also

makeModels, preprocessCoverage, calculateStats, calculatePvalues, annotateTranscripts, matchGenes
Examples

```r
## Collapse the coverage information
collapsedFull <- collapseFullCoverage(list(genomeData$coverage),
                                        verbose = TRUE)

## Calculate library size adjustments
sample Depths <- sampleDepth(collapsedFull, probs = c(0.5), nonzero = TRUE,
                               verbose = TRUE)

## Build the models
groupInfo <- genomeInfo$pop
adjustvars <- data.frame(genomeInfo$gender)
models <- makeModels(sampleDepths, testvars = groupInfo, adjustvars = adjustvars)

## Analyze the chromosome
results <- analyzeChr(chr = '21', coverageInfo = genomeData, models = models,
                       cutoffFstat = 1, cutoffType = 'manual', groupInfo = groupInfo, mc.cores = 1,
                       writeOutput = FALSE, returnOutput = TRUE, method = 'regular',
                       runAnnotation = FALSE)
names(results)
```

---

**annotateRegions**

Assign genomic states to regions

Description

This function takes the regions found in `calculatePvalues` and assigns them genomic states constructed with `makeGenomicState`. The main workhorse functions are `countOverlaps` and `findOverlaps`.

Usage

`annotateRegions(regions, genomicState, annotate = TRUE, ...)`

Arguments

- **regions**: The $regions output from `calculatePvalues`.
- **genomicState**: A GRanges object created with `makeGenomicState`. It can be either the genomicState$fullGenome or genomicState$codingGenome component.
- **annotate**: If TRUE then the regions are annotated by the genomic state. Otherwise, only the overlaps between the regions and the genomic states are computed.
- ... Arguments passed to other methods and/or advanced arguments.

Details

You might want to specify arguments such as `minoverlap` to control how the overlaps are determined. See `findOverlaps` for further details.
calculatePvalues

Value

A list with elements countTable and annotationList (only if annotate=TRUE).

- **countTable**: This is a data.frame with the number of overlaps from the regions vs the genomic states with one type per column. For example, if fullOrCoding='full' then the columns are exon, intergenic and intron.

- **annotationList**: This is a GRangesList with the genomic states that overlapped with the regions. The names of this GRangesList correspond to the region index in regions.

Author(s)

Andrew Jaffe, Leonardo Collado-Torres

See Also

makeGenomicState, calculatePvalues

Examples

```r
## Annotate regions, first two only
annotatedRegions <- annotateRegions(regions=genomeRegions$regions[1:2],
                                   genomicState=genomicState$fullGenome, minOverlap=1)
annotatedRegions
```

---

calculatePvalues    Calculate p-values and identify regions

Description

First, this function finds the regions of interest according to specified cutoffs. Then it permutes the samples and re-calculates the F-statistics. The area of the statistics from these segments are then used to calculate p-values for the original regions.

Usage

```r
calculatePvalues(coveragePrep, models, fstats, nPermute = 1L, 
                 seeds = as.integer(gsub("-", ",", Sys.Date())) + seq_len(nPermute), 
                 chr, 
                 cutoff = quantile(fstats, 0.99), significantCut = c(0.05, 0.1), 
                 lowMemDir = NULL, ...)
```

Arguments

- **coveragePrep**: A list with $coverageProcessed, $mclapplyIndex, and $position normally generated using preprocessCoverage.
- **models**: A list with $mod and $mod0 normally generated using makeModels.
- **fstats**: A numerical Rle with the F-statistics normally generated using calculateStats.
calculatePvalues

nPermuted The number of permutations. Note that for a full chromosome, a small amount (10) of permutations is sufficient. If set to 0, no permutations are performed and thus no null regions are used, however, the regions component is created.

seeds An integer vector of length nPermuted specifying the seeds to be used for each permutation. If NULL no seeds are used.

chr A single element character vector specifying the chromosome name. This argument is passed to findRegions.

cutoff F-statistic cutoff to use to determine segments.

significantCut A vector of length two specifying the cutoffs used to determine significance. The first element is used to determine significance for the p-values and the second element is used for the q-values.

lowMemDir The directory where the processed chunks are saved when using preprocessCoverage with a specified lowMemDir.

... Arguments passed to other methods and/or advanced arguments.

Value

A list with four components:

regions is a GRanges with metadata columns given by findRegions with the additional metadata column pvalues: p-value of the region calculated via permutations of the samples; qvalues: the qvalues calculated using qvalue; significant: whether the p-value is less than 0.05 (by default); significantQval: whether the q-value is less than 0.10 (by default). It also includes the mean coverage of the region (mean from the mean coverage at each base calculated in preprocessCoverage). Furthermore, if groupInfo was not NULL in preprocessCoverage, then the group mean coverage is calculated as well as the log 2 fold change (using group 1 as the reference).

nullStats is a numeric Rle with the mean of the null statistics by segment.

nullWidths is a numeric Rle with the length of each of the segments in the null distribution. The area can be obtained by multiplying the absolute nullstats by the corresponding lengths.

nullPermutation is a Rle with the permutation number from which the null region originated from.

Author(s)

Leonardo Collado-Torres

See Also

findRegions, fstats.apply, qvalue

Examples

## Collapse the coverage information
collapsedFull <- collapseFullCoverage(list(genomeData$coverage),
   verbose = TRUE)

## Calculate library size adjustments
```r
sampleDepths <- sampleDepth(collapsedFull, probs = c(0.5), verbose = TRUE)

## Build the models
group <- genomeInfo$pop
adjustvars <- data.frame(genomeInfo$gender)
models <- makeModels(sampleDepths, testvars = group, adjustvars = adjustvars)

## Preprocess the data
## Automatic chunksize used to then compare 1 vs 4 cores in the 'do not run'
## section
prep <- preprocessCoverage(genomeData, groupInfo = group, cutoff = 0,
                         scalefac = 32, chunksize = NULL, colsubset = NULL, mc.cores = 4)

## Get the F statistics
fstats <- genomeFstats

## We recommend determining the cutoff to use based on the F-distribution
## although you could also based it on the observed F-statistics.

## In this example we use a low cutoff used for illustrative purposes
cutoff <- 1

## Calculate the p-values and define the regions of interest.
regsWithP <- calculatePvalues(prep, models, fstats, nPermute = 1, seeds = 1,
                           chr = 'chr21', cutoff = cutoff, mc.cores = 1, method = 'regular')

## Not run:
## Calculate again, but with 10 permutations instead of just 1
regsWithP <- calculatePvalues(prep, models, fstats, nPermute = 10, seeds = 1:10,
                           chr = 'chr21', cutoff = cutoff, mc.cores = 2, method = 'regular')

## Check that they are the same as the previously calculated regions
library(testthat)
expect_that(regsWithP, equals(genomeRegions))

## Histogram of the theoretical p-values by region
hist(pf(regsWithP$regions$value, df1 - df0, n - df1), main = 'Distribution
original p-values by region', freq = FALSE)

## Histogram of the permuted p-values by region
hist(regsWithP$regions$pvalues, main = 'Distribution permuted p-values by
region', freq = FALSE)

## MA style plot
library('ggplot2')
ma <- data.frame(mean = regsWithP$regions$meanCoverage,
                log2FoldChange = regsWithP$regions$log2FoldChangeVRIvsCEU)
ggplot(ma, aes(x = log2(mean), y = log2FoldChange)) + geom_point() +
ylab('Fold Change (log2)') + xlab('Mean coverage (log2)') +
labs(title = 'MA style plot')

## Annotate the results
```
library('bumhunter')
library('TxDb.Hsapiens.UCSC.hg19.knownGene')
genes <- annotateTranscripts(TxDB.Hsapiens.UCSC.hg19.knownGene)
annotation <- matchGenes(regsWithP$regions, genes)
head(annotation)

## End(Not run)

calculateStats  

Calculate F-statistics at base pair resolution from a loaded BAM files

Description

After defining the models of interest (see makeModels) and pre-processing the data (see preprocessCoverage), use calculateStats to calculate the F-statistics at base-pair resolution.

Usage

calculateStats(coveragePrep, models, lowMemDir = NULL, ...)

Arguments

coveragePrep  A list with $coverageProcessed, $mclapplyIndex, and $position normally generated using preprocessCoverage.
models       A list with $mod and $mod0 normally generated using makeModels.
lowMemDir    The directory where the processed chunks are saved when using preprocessCoverage with a specified lowMemDir.
...          Arguments passed to other methods and/or advanced arguments.

Value

A numeric Rle with the F-statistics per base pair that passed the cutoff.

Author(s)

Leonardo Collado-Torres

See Also

makeModels, preprocessCoverage
coerceGR

Examples

```r
## Collapse the coverage information
collapsedFull <- collapseFullCoverage(list(genomeData$coverage),
    verbose = TRUE)

## Calculate library size adjustments
sampleDepths <- sampleDepth(collapsedFull, probs=c(0.5), verbose = TRUE)

## Build the models
group <- genomeInfo$pop
adjustvars <- data.frame(genomeInfo$gender)
models <- makeModels(sampleDepths, testvars = group, adjustvars = adjustvars)

## Preprocess the data
prep <- preprocessCoverage(genomeData, cutoff = 0, scalefac = 32,
    chunksize=le3, colsubset=NULL)

## Run the function
fstats <- calculateStats(prep, models, verbose=TRUE, method='regular')
fstats

## Not run:
## Compare vs pre-packaged F-statistics
library('testthat')
expect_that(fstats, is_equivalent_to(genomeFstats))
## End(Not run)
```

---

c coerceGR  

Coerce the coverage to a GRanges object for a given sample

Description

Given the output of fullCoverage, coerce the coverage to a GRanges object.

Usage

coerceGR(sample, fullCov, ...)

Arguments

- **sample**: The name or integer index of the sample of interest to coerce to a GRanges object.
- **fullCov**: A list where each element is the result from loadCoverage used with returnCoverage = TRUE. Can be generated using fullCoverage.
- **...**: Arguments passed to other methods and/or advanced arguments.
Value

A GRanges object with score metadata vector containing the coverage information for the specified sample. The ranges reported are only those for regions of the genome with coverage greater than zero.

Author(s)

Leonardo Collado-Torres

See Also

GRanges

Examples

```r
## Create a small fullCov object with data only for chr21
fullCov <- list('chr21' = genomeDataRaw)

## Coerce to a GRanges the first sample
gr <- createBwSample('ERR009101', fullCov = fullCov,
  seqlengths = c('chr21' = 48129895))

## Explore the output
gr
collapseFullCoverage
```

Description

For a given data set this function collapses the full coverage information for each sample from all the chromosomes. The resulting information per sample is the number of bases with coverage 0, 1, etc. It is similar to using `table()` on a regular vector. This information is then used by `sampleDepth` for calculating the sample depth adjustments. The data set can loaded to R using (see `fullCoverage`) and optionally filtered using `filterData`.

Usage

```r
collapseFullCoverage(fullCov, colsubset = NULL, save = FALSE, ...)
```

Arguments

- `fullCov` A list where each element is the result from `loadCoverage` used with cutoff=NULL. Can be generated using `fullCoverage`.
- `colsubset` Which columns of coverageInfo$coverage to use.
- `save` If TRUE, the result is saved as 'collapsedFull.Rdata'.
- `...` Arguments passed to other methods and/or advanced arguments.
Value

A list with one element per sample. Then per sample, a list with two vector elements: values and weights. The first one is the coverage value and the second one is the number of bases with that value.

Author(s)

Leonardo Collado-Torres

See Also

fullCoverage, sampleDepth

Examples

```r
## Collaps the coverage information for the filtered data
collapsedFull <- collapseFullCoverage(list(genomeData),
   verbose=TRUE)
collapsedFull

## Not run:
## You can also collapsed the raw data
collapsedFullRaw <- collapseFullCoverage(list(genomeDataRaw), verbose=TRUE)

## End(Not run)
```

---

coverageToExon  
Extract coverage information for exons

Description

This function extracts the coverage information calculated by fullCoverage for a set of exons determined by makeGenomicState. The underlying code is similar to getRegionCoverage with additional tweaks for calculating RPKM values.

Usage

```r
coverageToExon(fullCov = NULL, genomicState, L = NULL, returnType = "raw",
   files = NULL, ...)
```

Arguments

- **fullCov**  
  A list where each element is the result from loadCoverage used with returnCoverage = TRUE. Can be generated using fullCoverage. Alternatively, specify files to extract the coverage information from the regions of interest. This can be helpful if you do not wish to store fullCov for memory reasons.

- **genomicState**  
  A GRanges object created with makeGenomicState. It can be either the genomicState$fullGenome or genomicState$codingGenome component.
coverageToExon

L

The width of the reads used.

returnType

If raw, then the raw coverage information per exon is returned. If rpkm, RPKM values are calculated for each exon.

files

A character vector with the full path to the sample BAM files (or BigWig files). The names are used for the column names of the DataFrame. Check rawFiles for constructing files. files can also be a BamFileList object created with BamFileList or a BigWigFileList object created with BigWigFileList.

... Arguments passed to other methods and/or advanced arguments.

Details

Parallelization is used twice. First, it is used by strand. Second, for processing the exons by chromosome. So there is no gain in using mc.cores greater than the maximum of the number of strands and number of chromosomes.

If fullcov is NULL and files is specified, this function will attempt to read the coverage from the files. Note that if you used 'totalMapped' and 'targetSize' before, you will have to specify them again to get the same results.

See also advancedArg with fun='loadCoverage' for other details.

Value

A matrix (nrow = number of exons in genomicState corresponding to the chromosomes in fullCov, ncol = number of samples) with the number of reads (or RPKM) per exon. The row names correspond to the row indexes of genomicState$fullGenome (if fullOrcoding='full') or genomicState$codingGenome (if fullOrcoding='coding').

Author(s)

Andrew Jaffe, Leonardo Collado-Torres

See Also

fullCoverage, getRegionCoverage

Examples

## Obtain fullCov object
fullCov <- list('21'=genomeDataRaw$coverage)

## Use only the first two exons
smallGenomicState <- genomicState
smallGenomicState$fullGenome <- smallGenomicState$fullGenome[
    which(smallGenomicState$fullGenome$theRegion == 'exon')[1:2] ]

## Finally, get the coverage information for each exon
exonCov <- coverageToExon(fullCov=fullCov, 
                          genomicState=smallGenomicState$fullGenome, L=36)
createBw

Export coverage to BigWig files

Description
Using output from fullCoverage, export the coverage from all the samples to BigWig files using createBwSample.

Usage
createBw(fullCov, path = ".", keepGR = TRUE, ...)

Arguments
- fullCov: A list where each element is the result from loadCoverage used with returnCoverage = TRUE. Can be generated using fullCoverage.
- path: The path where the BigWig files will be created.
- keepGR: If TRUE, the GRanges objects created by coerceGR grouped into a GRangesList are returned. Otherwise they are discarded.
- ...: Arguments passed to other methods and/or advanced arguments.

Details
Use at most one core per chromosome.

Value
If keepGR = TRUE, then a GRangesList with the output for coerceGR for each of the samples.

Author(s)
Leonardo Collado-Torres

See Also
GRangesList, export, createBwSample, coerceGR

Examples
```r
## Create a small fullCov object with data only for chr21
fullCov <- list('chr21' = genomeDataRaw)

## Keep only 2 samples
fullCov$chr21$coverage <- fullCov$chr21$coverage[c(1, 31)]

## Create the BigWig files for all samples in a test dir
dir.create('createBw-example')
bws <- createBw(fullCov, 'createBw-example')
```
### Explore the output

\texttt{bws}

### First sample

\texttt{bws[[1]]}

### Note that if a sample has no bases with coverage \( > 0 \), the \texttt{GRanges} object is empty and no BigWig file is created for that sample.

\texttt{bws[[2]]}

---

**createBwSample**

Create a BigWig file with the coverage information for a given sample

**Description**

Given the output of `fullCoverage`, this function coerces the coverage to a \texttt{GRanges} object using `coerceGR` and then exports the coverage to a BigWig file using `export`.

**Usage**

\[
\text{createBwSample}(\text{sample, path} = ".", \text{fullCov, keepGR} = \text{TRUE}, \ldots)
\]

**Arguments**

- **sample**: The name or integer index of the sample of interest to coerce to a \texttt{GRanges} object.
- **path**: The path where the BigWig file will be created.
- **fullCov**: A list where each element is the result from `loadCoverage` used with `returnCoverage = \text{TRUE}`. Can be generated using `fullCoverage`.
- **keepGR**: If \text{TRUE}, the \texttt{GRanges} object created by `coerceGR` is returned. Otherwise it is discarded.
- \ldots: Arguments passed to other methods and/or advanced arguments.

**Value**

Creates a BigWig file with the coverage information (regions with coverage greater than zero) for a given sample. If `keepGR` it returns the output from `coerceGR`.

**Author(s)**

Leonardo Collado-Torres

**See Also**

\texttt{GRanges, export, linkcoerceGR}
Examples

```r
## Create a small fullCov object with data only for chr21
fullCov <- list(chr21 = genomeDataRaw)

## Create a BigWig for the first sample in a test directory
dir.create('createBwSample-example')
bw <- createBwSample('ERR09101', 'createBwSample-example',
                    fullCov = fullCov, seqlengths = c(chr21 = 48129895))

## Explore the output
bw
```

extendedMapSeqlevels  

Change naming style for a set of sequence names

Description

If available, use the information from GenomeInfoDb for your species of interest to map the sequence names from the style currently used to another valid style. For example, for Homo sapiens map '2' (NCBI style) to 'chr2' (UCSC style). If the information from GenomeInfoDb is not available, the original sequence names will be returned.

Usage

```r
extendedMapSeqlevels(seqnames, style = getOption("chrsStyle", "UCSC"),
                      species = getOption("species", "homo_sapiens"), currentStyle = NULL, ...)
```

Arguments

- `seqnames` A character vector with the sequence names.
- `style` A single character vector specifying the naming style to use for renaming the sequence names.
- `species` A single character vector with the species of interest: it has to match the valid species names supported in GenomeInfoDb. See `names(GenomeInfoDb::genomestyles())`. If `species = NULL`, a guess will be made using the available information in GenomeInfoDb.
- `currentStyle` A single character vector with the currently used naming style. If NULL, a guess will be made from the naming styles supported by species.
- `...` Arguments passed to other methods and/or advanced arguments.

Details

This function is inspired from `mapSeqlevels` with the difference that it will return the original sequence names if the species, current naming style or target naming style are not supported in GenomeInfoDb.
Value

A vector of sequence names using the specified naming style.

Author(s)

L. Collado-Torres

Examples

```r
## Without guessing any information
extendedMapSeqlevels("2", 'UCSC', 'Homo sapiens', 'NCBI')

## Guessing the current naming style
extendedMapSeqlevels("2", 'UCSC', 'Homo sapiens')

## Guess species and current style
extendedMapSeqlevels("2", 'NCBI')

## Guess species while supplying the current style.
## Probably an uncommon use-case
extendedMapSeqlevels("2", 'NCBI', currentStyle = 'TAIR10')

## Sequence names are unchanged when using an unsupported species
extendedMapSeqlevels('seq2', 'NCBI', 'toyOrganism')

## Not run:
## Set global species and style option
options('chrsStyle' = 'UCSC')
options('species' = 'homo_sapiens')

## Run using global options
extendedMapSeqlevels('2')

## End(Not run)
```

filterData

| Filter the positions of interest |

Description

For a group of samples this function reads the coverage information for a specific chromosome directly from the BAM files. It then merges them into a DataFrame and removes the bases that do not pass the cutoff. This is a helper function for `loadCoverage` and `preprocessCoverage`.

Usage

```r
filterData(data, cutoff = NULL, index = NULL, filter = "one",
            totalMapped = NULL, targetSize = 8e+07, ...)
```
filterData

Arguments

data Either a list of Rle objects or a DataFrame with the coverage information.
cutoff The base-pair level cutoff to use. It's behavior is controlled by filter.
index A logical Rle with the positions of the chromosome that passed the cutoff. If NULL it is assumed that this is the first time using filterData and thus no previous index exists.
filter Has to be either 'one' (default) or 'mean'. In the first case, at least one sample has to have coverage above cutoff. In the second case, the mean coverage has to be greater than cutoff.
totalMapped The total number of reads mapped for each sample. Providing this data adjusts the coverage to reads in targetSize library prior to filtering. By default, to reads per 80 million reads.
targetSize The target library size to adjust the coverage to. Used only when totalMapped is specified.
... Arguments passed to other methods and/or advanced arguments.

Details

If cutoff is NULL then the data is grouped into DataFrame without applying any cutoffs. This can be useful if you want to use loadCoverage to build the coverage DataFrame without applying any cutoffs for other downstream purposes like plotting the coverage values of a given region. You can always specify the colsubset argument in preprocessCoverage to filter the data before calculating the F statistics.

Value

A list with up to three components.

coverage is a DataFrame object where each column represents a sample. The number of rows depends on the number of base pairs that passed the cutoff and the information stored is the coverage at that given base. Included only when returnCoverage = TRUE.

position is a logical Rle with the positions of the chromosome that passed the cutoff.

meanCoverage is a numeric Rle with the mean coverage at each base. Included only when returnMean = TRUE.

Author(s)

Leonardo Collado-Torres

See Also

loadCoverage, preprocessCoverage
findRegions

Find non-zero regions in a Rle

Description

Find genomic regions for which a numeric vector is above (or below) predefined thresholds. In other words, this function finds the candidate Differentially Expressed Regions (candidate DERs). This is similar to regionFinder and is a helper function for calculatePvalues.

Usage

findRegions(position = NULL, fstats, chr, oneTable = TRUE,
maxClusterGap = 300L, cutoff = quantile(fstats, 0.99), segmentIR = NULL,
...)

Arguments

position A logical Rle of genomic positions. This is generated in loadCoverage. Note that it gets updated in preprocessCoverage if colsubset is not NULL.

fstats A numeric Rle with the F-statistics. Usually obtained using calculateStats.

chr A single element character vector specifying the chromosome name.

oneTable If TRUE only one GRanges is returned. Otherwise, a GRangesList with two components is returned: one for the regions with positive values and one for the negative values.

maxClusterGap This determines the maximum gap between candidate DERs. It should be greater than maxRegionGap (0 by default).

cutoff Threshold applied to the fstats used to determine the regions.

segmentIR An IRanges object with the genomic positions that are potentials DERs. This is used in calculatePvalues to speed up permutation calculations.

... Arguments passed to other methods and/or advanced arguments.
**findRegions**

Details

- `regionFinder` adapted to Rle world.

Value

- Either a GRanges or a GRangesList as determined by `onetable`. Each of them has the following metadata variables.
  - **value** The mean of the values of `y` for the given region.
  - **area** The absolute value of the sum of the values of `y` for the given region.
  - **indexStart** The start position of the region in terms of the index for `y`.
  - **indexEnd** The end position of the region in terms of the index for `y`.
  - **cluster** The cluster ID.
  - **clusterL** The total length of the cluster.

Author(s)

- Leonardo Collado-Torres

References


See Also

- `calculatePvalues`

Examples

```
## Preprocess the data
prep <- preprocessCoverage(genomeData, cutoff=0, scalefac=32, chunksize=1e3,
                          colsubset=NULL)

## Get the F statistics
fstats <- genomeFstats

## Find the regions
regs <- findRegions(prep$position, fstats, 'chr21', verbose=TRUE)

## Not run:
## Once you have the regions you can proceed to annotate them
library('bumphunter')
library('TxDb.Hsapiens.UCSC.hg19.knownGene')
genesis <- annotateTranscripts(TxDB.Hsapiens.UCSC.hg19.knownGene)
anotation <- matchGenes(regs, genesis)
anotation

## End(Not run)
```
fullCoverage

Description

For a group of samples this function reads the coverage information for several chromosomes directly from the BAM files. Per chromosome, it merges the unfiltered coverage by sample into a DataFrame. The end result is a list with one such DataFrame objects per chromosome.

Usage

fullCoverage(files, chrs, bai = NULL, chrlens = NULL, outputs = NULL, cutoff = NULL, ...)

Arguments

files A character vector with the full path to the sample BAM files (or BigWig files). The names are used for the column names of the DataFrame. Check rawFiles for constructing files. files can also be a BamFileList object created with BamFileList or a BigWigFileList object created with BigWigFileList.

chrs The chromosome of the files to read. The format has to match the one used in the input files.

bai The full path to the BAM index files. If NULL it is assumed that the BAM index files are in the same location as the BAM files and that they have the .bai extension. Ignored if files is a BamFileList object.

chrlens The chromosome lengths in base pairs. If it’s NULL, the chromosome length is extracted from the BAM files. Otherwise, it should have the same length as chrs.

outputs This argument is passed to the output argument of loadCoverage. If NULL or 'auto' it is then recycled.

cutoff This argument is passed to filterData.

... Arguments passed to other methods and/or advanced arguments.

Value

A list with one element per chromosome.

Each element is a DataFrame with the coverage information produced by loadCoverage.

Author(s)

Leonardo Collado-Torres

See Also

loadCoverage, filterData
Examples

datadir <- system.file('extdata', 'genomeData', package='derfinder')
files <- rawFiles(datadir=datadir, samplepatt='*.accepted_hits.bam$',
  fileterm=NULL)
## Shorten the column names
names(files) <- gsub('.accepted_hits.bam', '', names(files))

## Read and filter the data, only for 1 file
fullCov <- fullCoverage(files=files[1], chrs=c('21', '22'))
fullCov

## Not run:
## You can then use filterData() to filter the data if you want to.
## Use bplapply() if you want to do so with multiple cores as shown below.
library('BiocParallel')
p <- SnowParam(2L, outfile = Sys.getenv('SGE_STDERR_PATH'))
bplapply(fullCov, function(x) {
  library('derfinder'); filterData(x, cutoff=0 ), BPPARAM = p
})

## End(Not run)

---

**genomeData**

*Genome samples processed data*

**Description**

10kb region from chr21 processed for 31 RNA-seq samples described in `genomeInfo`. The TopHat BAM files are included in the package and this is the output of `loadCoverage` applied to it. For more information check the example of `loadCoverage`.

**Format**

A list with two components.

- **coverage** is a DataFrame object where each column represents a sample.
- **position** is a logical Rle with the positions of the chromosome that passed a cutoff of 0.

**References**


**See Also**

`loadCoverage`, `genomeInfo`
### genomeDataRaw

**Genome samples processed data**

**Description**

10kb region from chr21 processed for 31 RNA-seq samples described in `genomeInfo`. The TopHat BAM files are included in the package and this is the output of `loadCoverage` applied to it with `cutoff=NULL`. For more information check the example of `loadCoverage`.

**Format**

A list with two components.

- **coverage** is a DataFrame object where each column represents a sample.
- **position** is NULL because no bases were filtered.

**References**


**See Also**

`loadCoverage`, `genomeInfo`

### genomeFstats

**F-statistics for the example data**

**Description**

Calculated F-statistics for a 10kb region from chr21 processed for 31 RNA-seq samples described in `genomeInfo`. For more information check the example of `calculateStats`.

**Format**

A numeric Rle of length 1434 with the calculated F-statistics as exemplified in `calculateStats`.

**See Also**

`calculateStats`
**genomeInfo**

**Genome samples information**

**Description**

Information for the 31 samples downloaded from the Short Read Archive from studies comparing CEU and YRI populations. This data is used to specify the adjustment variables in `calculateStats`. The data is sorted according to the BAM files identifiers. Reads were 36bp long.

**Format**

A data.frame with 5 columns:

- **run**: The short name used to identify the sample BAM file.
- **library.layout**: Whether it was a single-end library or a paired-end library.
- **hapmap.id**: The HapMap identifier of the person sequenced. Note that some were sequenced more than once.
- **gender**: Whether the person sequence is a female or a male.
- **pop**: The population the person belongs to.

**Details**

The samples are from:

- **10** unrelated females from the YRI population.
- **5** unrelated females from the CEU population.
- **5** unrelated males (unrelated to the females too) from the CEU population.

**References**


**See Also**

`genomeData`, `calculateStats`
Candidate DERs for example data

Description

Candidate Differentially Expressed Regions (DERs) for the example data. For more information check `calculatePvalues`.

Format

A list with four components.

- `regions`  a GRanges object with the candidate DERs.
- `nullStats`  a numeric Rle with the mean F-statistics for the null DERs found from the permutations.
- `nullWidths`  an integer Rle with the width of each null candidate DER.
- `nullPermutation`  an integer Rle with the permutation number for each candidate DER. It identifies which permutation cycle created the null candidate DER.

See Also

- `calculatePvalues`

Genomic State for Hsapiens.UCSC.hg19.knownGene

Description

Pre-computed genomic state for Hsapiens UCSC hg19 knownGene annotation built using `makeGenomicState` for TxDb.Hsapiens.UCSC.hg19.knownGene version 2.14.0. The object has been subset for chr21 only.

Format

A GRangesList with two components.

- `fullGenome`  classifies each region as either being exon, intron or intergenic.
- `codingGenome`  classifies the regions as being promoter, exon, intro, 5UTR, 3UTR or intergenic.

See Also

- `makeGenomicState`
getRegionCoverage

Extract coverage information for a set of regions

Description

This function extracts the raw coverage information calculated by fullCoverage at each base for a set of regions found with calculatePvalues. It can further calculate the mean coverage per sample for each region.

Usage

getRegionCoverage(fullCov = NULL, regions, totalMapped = NULL, targetSize = 8e+07, files = NULL, ...)

Arguments

fullCov A list where each element is the result from loadCoverage used with returnCoverage = TRUE. Can be generated using fullCoverage. Alternatively, specify files to extract the coverage information from the regions of interest. This can be helpful if you do not wish to store fullCov for memory reasons.

regions The $regions output from calculatePvalues. It is important that the seqlengths information is provided.

totalMapped The total number of reads mapped for each sample. Providing this data adjusts the coverage to reads in targetSize library. By default, to reads per 80 million reads.

targetSize The target library size to adjust the coverage to. Used only when totalMapped is specified.

files A character vector with the full path to the sample BAM files (or BigWig files). The names are used for the column names of the DataFrame. Check rawFiles for constructing files. files can also be a BamFileList object created with BamFileList or a BigWigFileList object created with BigWigFileList.

... Arguments passed to other methods and/or advanced arguments.

Details

When fullCov is the output of loadCoverage with cutoff non-NULL, getRegionCoverage assumes that the regions come from the same data. Meaning that filterData was not used again. This ensures that the regions are a subset of the data available in fullCov.

If fullCov is NULL and files is specified, this function will attempt to read the coverage from the files. Note that if you used 'totalMapped' and 'targetSize' before, you will have to specify them again to get the same results.

See also advancedArg with fun='loadCoverage' for other details.

You should use at most one core per chromosome.
loadCoverage

Load the coverage information from a group of BAM files

Description

For a group of samples this function reads the coverage information for a specific chromosome directly from the BAM files. It then merges them into a DataFrame and removes the bases that do not pass the cutoff.

Usage

loadCoverage(files, chr, cutoff = NULL, filter = "one", chrlen = NULL, output = NULL, bai = NULL, ...)

Arguments

files A character vector with the full path to the sample BAM files (or BigWig files). The names are used for the column names of the DataFrame. Check rawFiles for constructing files. files can also be a BamFileList, BamFile, BigWigFileList, or BigWigFile object.
loadCoverage

chr
Chromosome to read. Should be in the format matching the one used in the raw data.

cutoff
This argument is passed to filterData.

filter
Has to be either 'one' (default) or 'mean'. In the first case, at least one sample has to have coverage above cutoff. In the second case, the mean coverage has to be greater than cutoff.

chrlen
The chromosome length in base pairs. If it’s NULL, the chromosome length is extracted from the BAM files.

output
If NULL then no output is saved in disk. If auto then an automatic name is constructed using UCSC names (chrXCovInfo.Rdata for example). If another character is specified, then that name is used for #’ the output file.

bai
The full path to the BAM index files. If NULL it is assumed that the BAM index files are in the same location as the BAM files and that they have the .bai extension. Ignored if files is a BamFileList object or if inputType==’BigWig’.

...arguments passed to other methods and/or advanced arguments.

Details
The ... argument can be used to control which alignments to consider when reading from BAM files. See scanBamFlag.

Parallelization for loading the data in chunks is used only used when tilewidth is specified. You may use up to one core per tile.

If you set the advanced argument dropNd = TRUE, bases with CIGAR string "D" (deletion from reference) will be excluded from the base-level coverage calculation.

If you are working with data from an organism different from 'Homo sapiens' specify so by setting the global 'species' and 'chrsStyle' options. For example: options(species = 'arabidopsis_thaliana') options(chrsStyle = 'NCBI')

Value
A list with two components.

coverage is a DataFrame object where each column represents a sample. The number of rows depends on the number of base pairs that passed the cutoff and the information stored is the coverage at that given base.

position is a logical Rle with the positions of the chromosome that passed the cutoff.

Author(s)
Leonardo Collado-Torres, Andrew Jaffe

Examples

datadir <- system.file('extdata', 'genomeData', package='derfinder')
files <- rawFiles(datadir = datadir, samplepatt = '*accepted_hits.bam$',
                  fileterm = NULL)
## Shorten the column names
names(files) <- gsub('_accepted_hits.bam', '', names(files))

## Read and filter the data, only for 2 files
dataSmall <- loadCoverage(files = files[1:2], chr = 'chr21', cutoff = 0)

## Not run:
## Export to BigWig files
createBw(list('chr21' = dataSmall))

## Load data from BigWig files
dataSmall.bw <- loadCoverage(c(ERR009101 = 'ERR009101.bw', ERR009102 = 'ERR009102.bw'), chr = 'chr21')

## Compare them
mapply(function(x, y) { x - y }, dataSmall$coverage, dataSmall.bw$coverage)

## Note that the only difference is the type of Rle (integer vs numeric) used
## to store the data.

## End(Not run)

makeGenomicState <- function(txdb, chrs = c(1:22, "X", "Y"), ...)

## Arguments

txdb A TxDb object.
chrs The names of the chromosomes to use as denoted in the txdb object. Check isActiveSeq.
...
Arguments passed to other methods and/or advanced arguments.

## Value

A GRangesList object with two elements: fullGenome and codingGenome. Both have metadata information for the type of region (theRegion), transcript IDs (tx_id), transcript name (tx_name), and gene ID (gene_id). fullGenome classifies each region as either being exon, intron or intergenic. codingGenome classifies the regions as being promoter, exon, intro, 5UTR, 3UTR or intergenic.
Author(s)

Andrew Jaffe, Leonardo Collado-Torres

See Also

TxDb

Examples

```r
## Load the example data base from the GenomicFeatures vignette
library('GenomicFeatures')
samplefile <- system.file('extdata', 'hg19_knownGene_sample.sqlite',
                          package='GenomicFeatures')
txdb <- loadDb(samplefile)

## Generate genomic state object, only for chr6
sampleGenomicState <- makeGenomicState(txdb, chrs='chr6')
## Not run:
## Create the GenomicState object for Hsapiens.UCSC.hg19.knownGene
library('TxDb.Hsapiens.UCSC.hg19.knownGene')
txdb <- TxDb.Hsapiens.UCSC.hg19.knownGene

## Creating this GenomicState object takes around 8 min for all chrs and
## around 30 secs for chr21
GenomicState.Hsapiens.UCSC.hg19.knownGene.chr21 <-
  makeGenomicState(txdb=txdb, chrs='chr21')

## For convinience, this object is already included in derfinder
library('testthat')
expect_that(GenomicState.Hsapiens.UCSC.hg19.knownGene.chr21,
            is_equivalent_to(genomicState))

## Hsapiens ENSEMBL GRCh37
library('GenomicFeatures')
## Can take several minutes and speed will depend on your internet speed
xx <- makeTxDbPackageFromBiomart(version = '0.99', maintainer = 'Your Name',
                                 author='Your Name')
txdb <- loadDb(file.path('TxDb.Hsapiens.BioMart.ensemble.GRCh37.p11', 'inst',
                          'extdata', 'TxDb.Hsapiens.BioMart.ensemble.GRCh37.p11.sqlite'))

## Creating this GenomicState object takes around 13 min
GenomicState.Hsapiens.ensemble.GRCh37.p11 <- makeGenomicState(txdb=txdb,
                                                              chrs=c(1:22, 'X', 'Y'))

## Save for later use
save(GenomicState.Hsapiens.ensemble.GRCh37.p11,
     file='GenomicState.Hsapiens.ensemble.GRCh37.p11.Rdata')

## End(Not run)
```
makeModels

Build model matrices for differential expression

Description

Builds the model matrices for testing for differential expression by comparing a model with a grouping factor versus one without it. It adjusts for the confounders specified and the median coverage of each sample. The resulting models can be used in calculateStats.

Usage

makeModels(sampleDepths, testvars, adjustvars = NULL, testIntercept = FALSE)

Arguments

sampleDepths  Per sample library size adjustments calculated with sampleDepth.
testvars  A vector or matrix specifying the variables to test. For example, a factor with the group memberships when testing for differences across groups. It's length should match the number of columns used from coverageInfo$coverage.
adjustvars  Optional matrix of adjustment variables (e.g. measured confounders, output from SVA, etc.) to use in fitting linear models to each nucleotide. These variables have to be specified by sample and the number of rows must match the number of columns used. It will also work if it is a vector of the correct length.
testIntercept  If TRUE then testvars is ignored and mod0 will contain the column medians and any adjusting variables specified, but no intercept.

Value

A list with two components.

- mod  The alternative model matrix.
- mod0  The null model matrix.

Author(s)

Leonardo Collado-Torres

See Also

sampleDepth, calculateStats
mergeResults

Examples

## Collapse the coverage information
collapsedFull <- collapseFullCoverage(list(genomeData$coverage),
    verbose=TRUE)

## Calculate library size adjustments
sampleDepths <- sampleDepth(collapsedFull, probs=c(0.5), nonzero=TRUE,
    verbose=TRUE)

## Build the models
models <- makeModels(sampleDepths, testvars=group, adjustvars=adjustvars)

mergeResults

Merge results from different chromosomes

Description

This function merges the results from running `analyzeChr` on several chromosomes and assigns genomic states using `annotateRegions`. It re-calculates the p-values and q-values using the pooled areas from the null regions from all chromosomes. Once the results have been merged, `derfinderReport::generateReport` can be used to generate an HTML report of the results. The `derfinderReport` package is available at https://github.com/lcolladotor/derfinderReport.

Usage

mergeResults(chrs = c(1:22, "X", "Y"), prefix = ".",
    significantCut = c(0.05, 0.1), genomicState, minoverlap = 20,
    mergePrep = FALSE, ...)

Arguments

chrs The chromosomes of the files to be merged.

prefix The main data directory path, which can be useful if `analyzeChr` is used for several parameters and the results are saved in different directories.

significantCut A vector of length two specifying the cutoffs used to determine significance. The first element is used to determine significance for the p-values and the second element is used for the q-values just like in `calculatePvalues`.

genomicState A GRanges object created with `makeGenomicState`. It can be either the genomicState$fullGenome or genomicState$codingGenome component.

minoverlap Determines the minimum overlap needed when annotating regions with `annotateRegions`.

mergePrep If TRUE the output from `preprocessCoverage` is merged.

Arguments passed to other methods and/or advanced arguments.
Details
If you want to calculate the FWER, supply optionsStats which is produced by `analyzeChr`.

Value
Seven Rdata files.

- **fullFstats.Rdata**: Full F-statistics from all chromosomes in a list of Rle objects.
- **fullTime.Rdata**: Timing information from all chromosomes.
- **fullNullSummary.Rdata**: A DataFrame with the null region information: statistic, width, chromosome and permutation identifier. It's ordered by the statistics.
- **fullRegions.Rdata**: GRanges object with regions found and with full annotation from `matchGenes`. Note that the column `strand` from `matchGenes` is renamed to `annoStrand` to comply with GRanges specifications.
- **fullCoveragePrep.Rdata**: A list with the pre-processed coverage data from all chromosomes.
- **fullAnnotatedRegions.Rdata**: A list as constructed in `annotateRegions` with the assigned genomic states.
- **optionsMerge.Rdata**: A list with the options used when merging the results. Used in `derfinderReport::generateReport`.

Author(s)
Leonardo Collado-Torres

See Also
`analyzeChr`, `calculatePvalues`, `annotateRegions`

Examples
### The output will be saved in the 'generateReport-example' directory
dir.create('generateReport-example', showWarnings = FALSE, recursive = TRUE)

### For convenience, the derfinder output has been pre-computed
file.copy(system.file(file.path('extdata', 'chr21'), package='derfinder', mustWork=TRUE), 'generateReport-example', recursive=TRUE)

### Merge the results from the different chromosomes. In this case, there's only one: chr21
mergeResults(chrs='21', prefix='generateReport-example', genomicState=genomicState$fullGenome)

### Not run:
### You can then explore the wallclock time spent on each step
load(file.path('generateReport-example', 'fullRegions.Rdata'))

### Process the time info
time <- lapply(fullTime, function(x) data.frame(diff(x)))
time <- do.call(rbind, time)
colnames(time) <- 'sec'
time$sec <- as.integer(round(time$sec))
time$min <- time$sec / 60
time$chr <- paste0('chr', gsub('\..*', '', rownames(time)))
time$step <- gsub('.\..*', '', rownames(time))
rownames(time) <- seq_len(nrow(time))

## Make plot
library('ggplot2')
ggplot(time, aes(x=step, y=min, colour=chr)) + geom_point() +
  labs(title='Wallclock time by step') +
  scale_colour_discrete(limits=chrs) +
  scale_x_discrete(limits=names(fullTime[[1]][-1])) + ylab('Time (min)') +
  xlab('Step')

## End(Not run)

---

**preprocessCoverage**

Transform and split the data

---

**Description**

This function takes the coverage data from `loadCoverage`, scales the data, does the log2 transformation, and splits it into appropriate chunks for using `calculateStats`.

**Usage**

```r
preprocessCoverage(coverageInfo, groupInfo = NULL, cutoff = 5,
colsubset = NULL, lowMemDir = NULL, ...)
```

**Arguments**

- `coverageInfo` A list containing a DataFrame with the coverage data and a logical Rle with the positions that passed the cutoff. This object is generated using `loadCoverage`.
- `groupInfo` A factor specifying the group membership of each sample. If NULL no group mean coverages are calculated. If the factor has more than one level, the first one will be used to calculate the log2 fold change in `calculatePvalues`.
- `cutoff` The base-pair level cutoff to use. It's behavior is controlled by `filter`.
- `colsubset` Optional vector of column indices of `coverageInfo$coverage` that denote samples you wish to include in analysis.
- `lowMemDir` If specified, each chunk is saved into a separate Rdata file under `lowMemDir` and later loaded in `fstats.apply` when running `calculateStats` and `calculatePvalues`. Using this option helps reduce the memory load as each fork in `bplapply` loads only the data needed for the chunk processing. The downside is a bit longer computation time due to input/output.
- `...` Arguments passed to other methods and/or advanced arguments.
### Details

If chunksize is NULL, then mc.cores is used to determine the chunksize. This is useful if you want to split the data so each core gets the same amount of data (up to rounding).


If lowMemDir is specified then $coverageProcessed is NULL and $mclapplyIndex is a vector with the chunk identifiers.

### Value

A list with five components.

- **coverageProcessed**: contains the processed coverage information in a DataFrame object. Each column represents a sample and the coverage information is scaled and log2 transformed. Note that if colsubset is not NULL the number of columns will be less than those in coverageInfo$coverage. The total number of rows depends on the number of base pairs that passed the cutoff and the information stored is the coverage at that given base. Further note that filterData is re-applied if colsubset is not NULL and could thus lead to fewer rows compared to coverageInfo$coverage.

- **mclapplyIndex**: is a list of logical Rle objects. They contain the partitioning information according to chunksize.

- **position**: is a logical Rle with the positions of the chromosome that passed the cutoff.

- **meanCoverage**: is a numeric Rle with the mean coverage at each filtered base.

- **groupMeans**: is a list of Rle objects containing the mean coverage at each filtered base calculated by group. This list has length 0 if groupInfo=NULL.

### Author(s)

Leonardo Collado-Torres

### See Also

- filterData, loadCoverage, calculateStats

### Examples

```r
## Split the data and transform appropriately before using calculateStats()
dataReady <- preprocessCoverage(genomeData, cutoff = 0, scalefac = 32,
                               chunksize = 1e3, colsubset = NULL, verbose = TRUE)
names(dataReady)
dataReady
```
**rawFiles**  
*Construct full paths to a group of raw input files*

**Description**

For a group of samples this function creates the list of paths to the raw input files which can then be used in `loadCoverage`. The raw input files are either BAM files or BigWig files.

**Usage**

```r
rawFiles(datadir = NULL, sampledirs = NULL, samplepatt = NULL,  
          fileterm = "accepted_hits.bam")
```

**Arguments**

- `datadir` The main directory where each of the `sampledirs` is a sub-directory of `datadir`.
- `sampledirs` A character vector with the names of the sample directories. If `datadir` is `NULL` it is then assumed that `sampledirs` specifies the full path to each sample.
- `samplepatt` If specified and `sampledirs` is set to `NULL`, then the directories matching this pattern in `datadir` (set to `.` if it’s set to `NULL`) are used as the sample directories.
- `fileterm` Name of the BAM or BigWig file used in each sample. By default it is set to `accepted_hits.bam` since that is the automatic name generated when aligning with TopHat. If `NULL` it is then ignored when reading the rawfiles. This can be useful if all the raw files are stored in a single directory.

**Details**

This function can also be used to identify a set of BigWig files.

**Value**

A vector with the full paths to the raw files and sample names stored as the vector names.

**Author(s)**

Leonardo Collado-Torres

**See Also**

`loadCoverage`

**Examples**

```r
## Get list of BAM files included in derfinder
datadir <- system.file('exdata', 'genomeData', package='derfinder')
files <- rawFiles(datadir=datadir, samplepatt='*accepted_hits.bam*',  
                 fileterm=NULL)
files
```
regionMatrix

Identify regions data by a coverage filter and get a count matrix

Description

Given a set of un-filtered coverage data (see fullCoverage), create candidate regions by applying a
cutoff on the coverage values, and obtain a count matrix where the number of rows corresponds to
the number of candidate regions and the number of columns corresponds to the number of samples.
The values are the mean coverage for a given sample for a given region.

Usage

regionMatrix(fullCov, cutoff = 5, filter = "mean", L, runFilter = TRUE,
returnBP = TRUE, ...)

Arguments

fullCov  A list where each element is the result from loadCoverage used with returnCoverage = TRUE.
Can be generated using fullCoverage. If runFilter = FALSE, then returnMean = TRUE
must have been used.
cutoff  The base-pair level cutoff to use. It’s behavior is controlled by filter.
filter  Has to be either 'one' (default) or 'mean'. In the first case, at least one sample
has to have coverage above cutoff. In the second case, the mean coverage has
to be greater than cutoff.
L  The width of the reads used.
runFilter  This controls whether to run filterData or not. If set to FALSE then returnMean = TRUE
must have been used to create each element of fullCov.
returnBP  If TRUE, returns $bpCoverage explained below.
...  Arguments passed to other methods and/or advanced arguments.

Details

This function uses several other derfinder-package functions. Inspect the code if interested.
You should use at most one core per chromosome.

Value

A list with one entry per chromosome. Then per chromosome, a list with three components.

regions  A set of regions based on the coverage filter cutoff as returned by findRegions.
bpCoverage  A list with one element per region. Each element is a matrix with numbers of rows
equal to the number of base pairs in the region and number of columns equal to the number of
samples. It contains the base-level coverage information for the regions. Only returned when
returnBP = TRUE.
coverageMatrix  A matrix with the mean coverage by sample for each candidate region.
**sampleDepth**

**Calculate adjustments for library size**

**Description**

For a given data set calculate the per-sample coverage adjustments. Hector Corrada’s group proposed calculating the sum of the coverage for genes below a given sample quantile. In this function, we calculate the sample quantiles of interest by sample, and then the sum of the coverage for bases below or equal to quantiles of interest. The resulting values are transformed \( \log_2(x + \text{scalefac}) \) to avoid very large numbers that could potentially affect the stability of the F-statistics calculation. The sample coverage adjustments are then used in `makeModels` for constructing the null and alternative models.

**Usage**

```r
sampleDepth(collapsedFull, probs = c(0.5, 1), scalefac = 32, ...)
```
sampleDepth

Arguments

- **collapsedFull**: The full coverage data collapsed by sample as produced by `collapseFullCoverage`.
- **probs**: Number(s) between 0 and 1 representing the quantile(s) of interest. For example, 0.5 is the median.
- **scalefac**: Number added to the sample coverage adjustments before the log2 transformation.
- ... Arguments passed to other methods and/or advanced arguments.

Value

A matrix (vector of length(probs) == 1) with the library size depth adjustments per sample to be used in `makeModels`. The number of rows corresponds to the number of quantiles used for the sample adjustments.

Author(s)

Leonardo Collado-Torres

References


See Also

collapseFullCoverage, makeModels

Examples

```r
## Collapse the coverage information
collapsedFull <- collapseFullCoverage(list(genomeData$coverage),
  verbose=TRUE)

## Calculate library size adjustments
sampleDepths <- sampleDepth(collapsedFull, probs=c(0.5, 1), verbose=TRUE)
sampleDepths
```
Index

*Topic datasets
  genomedata, 23
genomedataraw, 24
genomefstats, 24
genomeinfo, 25
genomeRegions, 26
genomicstate, 26

*Topic package
  derfinderMpackage, 2

advanced_arg (advancedArg), 3
advancedArg, 3, 14, 27
analyze_chr (analyzeChr), 4
analyzeChr, 4, 33, 34
annotate_regions (annotateRegions), 6
annotateRegions, 6, 33, 34
annotateTranscripts, 4, 5

BamFile, 28
BamFileList, 14, 22, 27, 28
BigWigFile, 28
BigWigFileList, 14, 22, 27, 28
bplapply, 5, 33

calculate_pvalues (calculatePvalues), 7
calculate_stats (calculateStats), 10
calculatePvalues, 2, 4–7, 7, 20, 21, 26–28, 33–35
calculateStats, 2, 4, 5, 7, 10, 10, 20, 24, 25, 32, 35, 36
coerce_gr (coerceGR), 11
coerceGR, 11, 15, 16
collapse_full_coverage (collapseFullCoverage), 12
collapseFullCoverage, 12, 40
countOverlaps, 6
coverage_to_exon (coverageToExon), 13
coverageToExon, 13
create_bw (createBw), 15
create_bw_sample (createBwSample), 16
createBw, 15
createBwSample, 15, 16
derfinder-package, 2, 38
derfinder-package, 2
export, 15, 16
extended_map_seqlevels
  (extendedMapSeqlevels), 17
extendedMapSeqlevels, 17
filter_data (filterData), 18
filterData, 12, 18, 19, 22, 27, 29, 36, 38
find_regions (findRegions), 20
findOverlaps, 6
findRegions, 8, 20, 38
fstats.apply, 5, 8, 35
full_coverage (fullCoverage), 22
fullCoverage, 11–16, 22, 27, 28, 38
genomedata, 23, 25
genomedataraw, 24
genomefstats, 24
genomeinfo, 23, 24, 25
genomeRegions, 26
 genomicstate, 26
get_region_coverage
  (getRegionCoverage), 27
getRegionCoverage, 13, 14, 27, 27
GRanges, 11, 12, 15, 16
GRangesList, 15

isActiveSeq, 30

load_coverage (loadCoverage), 28
loadCoverage, 2, 4, 11–13, 15, 16, 18–20,
  22–24, 27, 28, 35–38

make_genomic_state (makeGenomicState), 30
make_models (makeModels), 32
makeGenomicState, 6, 7, 13, 26, 30, 33
makeModels, 2, 4, 5, 7, 10, 32, 39, 40
mapSeqlevels, 17
matchGenes, 2, 4, 5, 34
mclapply, 36
merge_results (mergeResults), 33
mergeResults, 33

preprocess_coverage
  (preprocessCoverage), 35
preprocessCoverage, 2, 4, 5, 7, 8, 10, 18–20,
  33, 35

qf, 4
quantile, 4
qvalue, 8

raw_files (rawFiles), 37
rawFiles, 2, 14, 22, 27, 28, 37
region_matrix (regionMatrix), 38
regionFinder, 20, 21
regionMatrix, 38

sample_depth (sampleDepth), 39
sampleDepth, 12, 13, 32, 39
scanBamFlag, 29
SnowParam, 3

TxDb, 30, 31
TxDb.Hsapiens.UCSC.hg19.knownGene, 5