

Package ‘chipseq’

October 12, 2021

Title chipseq: A package for analyzing chipseq data

Version 1.42.0

Author Deepayan Sarkar, Robert Gentleman, Michael Lawrence, Zizhen Yao

Description Tools for helping process short read data for chipseq experiments

Depends R (>= 2.10), methods, BiocGenerics (>= 0.1.0), S4Vectors (>= 0.17.25), IRanges (>= 2.13.12), GenomicRanges (>= 1.31.8), ShortRead

Imports methods, stats, lattice, BiocGenerics, IRanges, GenomicRanges, ShortRead

Suggests BSgenome, GenomicFeatures, TxDb.Mmusculus.UCSC.mm9.knownGene

Maintainer Bioconductor Package Maintainer
<maintainer@bioconductor.org>

License Artistic-2.0

LazyLoad yes

biocViews ChIPSeq, Sequencing, Coverage, QualityControl, DataImport

git_url <https://git.bioconductor.org/packages/chipseq>

git_branch RELEASE_3_13

git_last_commit 735c9dd

git_last_commit_date 2021-05-19

Date/Publication 2021-10-12

R topics documented:

chipseqFilter	2
coverageplot	3
cstest	3
diffPeakSummary	4
estimate.mean.fraglen	5
islandDepthPlot	7

laneSubsample	8
peakCutoff	8
peakSummary-methods	9
subsetSummary	10
Index	12

chipseqFilter	<i>Filtering ChIP-seq reads</i>
---------------	---------------------------------

Description

Convenience for creating an [SRFilter](#) object appropriate for ChIP-seq data. Typically, the result is passed to [readAligned](#) when loading reads.

Usage

```
chipseqFilter(exclude = "[_MXY]", uniqueness = c("location", "sequence", "location*sequence", "none"),
```

Arguments

- exclude A regular expression for excluding chromosomes by name. Just like the parameter to [bsapply](#).
- uniqueness The criteria used to determine whether a read is unique. A read may be unique if it maps to a unique location, has a unique sequence or both. Specifying none avoids this test entirely.
- hasStrand Whether to require that the read is mapped to a strand, which usually translates to whether the read was mapped at all.

Value

An SRFilter object

Author(s)

M. Lawrence

Examples

```
sp <- SolexaPath(system.file("extdata", package="ShortRead"))

filter <- chipseqFilter()
aln <- readAligned(sp, "s_2_export.txt", filter=filter)
## allow mapping to the same location (but only if sequence is different)
filter <- chipseqFilter(uniqueness = "sequence")
aln <- readAligned(sp, "s_2_export.txt", filter=filter)
## allow sex chromosomes
filter <- chipseqFilter(exclude = "[M_]")
aln <- readAligned(sp, "s_2_export.txt", filter=filter)
```

coverageplot	<i>Plot coverage on a small interval.</i>
--------------	---

Description

A function that plots one or two coverage vectors over a relatively small interval in the genome.

Usage

```
coverageplot(peaks1, peaks2 = NULL, i = 1,  
             xlab = "Position", ylab = "Coverage",  
             opposite = TRUE, ...)
```

Arguments

peaks1, peaks2	A set of peaks as described by ranges over a coverage vector.
i	Which peak to use.
xlab, ylab	Axis labels.
opposite	Logical specifying whether the two peaks should be plotted on opposite sides (appropriate for positive and negative strand peaks).
...	extra arguments.

Author(s)

Deepayan Sarkar

Examples

```
cov <- Rle(c(1:10, seq(10, 1, -2), seq(1,5,2), 4:1), rep(1:2, 11))  
peaks <- slice(cov, 3)  
peaks.cov <- Views(cov, ranges(peaks))  
peaks.cov.rev <- rev(peaks.cov)  
coverageplot(peaks.cov, peaks.cov.rev, ylab = "Example")
```

cstest	<i>A test ChIP-Seq dataset</i>
--------	--------------------------------

Description

A small subset of a ChIP-Seq dataset downloaded from the Short-Read Archive.

Usage

```
data(cstest)
```

Format

The dataset is on object of class `GRangesList` with read alignments from three chromosomes in two lanes representing CTCF and GFP pull-down in mouse.

Source

Short Read Archive, GEO accession number GSM288351 <http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSM288351>

References

Chen X., Xu H., Yuan P., Fang F., Huss M., Vega V.B., Wong E., Orlov Y.L., Zhang W., Jiang J., Loh Y.H., Yeo H.C., Yeo Z.X., Narang V., Govindarajan K.R., Leong B., Shahab A.S., Ruan Y., Bourque G., Sung W.K., Clarke N.D., Wei C.L., Ng H.H. (2008), "Integration of External Signaling Pathways with the Core Transcriptional Network in Embryonic Stem Cells". *Cell*, 133:1106-1117.

Examples

```
data(cstest)
names(cstest)
cstest$gfp
```

diffPeakSummary	<i>A function to identify and produce summary statistics for differentially expressed peaks.</i>
-----------------	--

Description

Given two sets of peaks, this function combines them and summarizes the individual coverage vectors under the combined peak set.

Usage

```
diffPeakSummary(ranges1, ranges2,
                 viewSummary = list(sums = viewSums, maxs = viewMaxs))
```

Arguments

<code>ranges1</code>	First set of peaks (typically an <code>RleViewsList</code>).
<code>ranges2</code>	Second set of peaks (typically an <code>RleViewsList</code>).
<code>viewSummary</code>	A list of the per peak summary functions.

Value

A data.frame with one row for each peak in the combined data. The chromosome, start and stop nucleotide positions (+ strand) are given as are the summary statistics requested.

Author(s)

D. Sarkar

Examples

```
data(cstest)
library(BSgenome.Mmusculus.UCSC.mm9)
seqlevels(cstest) <- seqlevels(Mmusculus)
seqlengths(cstest) <- seqlengths(Mmusculus)
## find peaks
findPeaks <- function(reads) {
  reads.ext <- resize(reads, width = 200)
  slice(coverage(reads.ext), lower = 8)
}
peakSummary <- diffPeakSummary(findPeaks(cstest$gfp), findPeaks(cstest$ctcf))
```

`estimate.mean.fraglen` *Estimate summaries of the distribution of fragment lengths in a short-read experiment. The methods are designed for ChIP-Seq experiments and may not work well in data without peaks.*

Description

`estimate.mean.fraglen` implements three methods for estimating mean fragment length. The other functions are related helper functions implementing various methods, but may be useful by themselves for diagnostic purposes. Many of these operations are potentially slow.

`sparse.density` is intended to be similar to [density](#), but returns the results in a run-length encoded form. This is useful when long stretches of the range of the data have zero density.

Usage

```
estimate.mean.fraglen(x, method = c("SISSR", "coverage", "correlation"),
  ...)

basesCovered(x, shift = seq(5, 300, 5), seqLen = 100, verbose = FALSE)

densityCorr(x, shift = seq(0, 500, 5), center = FALSE,
  width = seqLen * 2L, seqLen = 100L, maxDist = 500L, ...)

sparse.density(x, width = 50, kernel = "epanechnikov",
  from = start(rix)[1] - 10L,
  to = end(rix)[length(rix)] + 10L)
```

Arguments

x	For estimate.mean.fraglen, typically an AlignedRead or a GRanges object. For basesCovered and densityCorr, a list with elements "+" and "-" representing locations of reads aligned to positive and negative strands (the values should be integers denoting the location where the first sequenced base matched.) densityCorr has also come to support GRanges input directly. For sparse.density, a numeric or integer vector for which density is to be computed.
method	Character string giving method to be used. method = "SISSR" implements the method described in Jothi et al (see References below). method = "correlation" implements the method described in Kharchenko et al (see References below), where the idea is to compute the density of tag start positions separately for each strand, and then determine the amount of shift that maximizes the correlation between these two densities. method = "coverage" computes the optimal shift for which the number of bases covered by any read is minimized.
shift	Integer vector giving amount of shifts to be tried when optimizing. The current algorithm simply evaluates all supplied values and reports the one giving minimum coverage or maximum correlation.
seqLen	For the "coverage" method, the assumed length of each read for computing the coverage. Typically the read length. This is added to the shift estimated by "coverage" and "correlation" to come up with the actual fragment length.
verbose	Logical specifying whether progress information should be printed during execution.
center	For the "correlation" method, whether the calculations should incorporate centering by the mean density. The default is not to do so; as the density is zero over most of the genome, this slightly improves efficiency at negligible loss in accuracy.
width	half-bandwidth used in the computation. This needs to be specified as an integer, data-driven rules are not supported.
kernel	A character string giving the density kernel.
from, to	specifies range over which the density is to be computed.
maxDist	If distance to nearest neighbor is more than this, the position is discarded. This removes isolated points, which are not very informative.
...	Extra arguments, passed on as appropriate to other functions.

Details

For the correlation method, the range over which densities are computed only cover the range of reads; that is, the beginning and end of chromosomes are excluded.

Value

estimate.mean.fraglen gives an estimate of the mean fragment length.

basesCovered and densityCorr give a vector of the corresponding objective function evaluated at the supplied values of shift.

sparse.density returns an object of class "Rle".

Author(s)

Deepayan Sarkar, Michael Lawrence

References

R. Jothi, S. Cuddapah, A. Barski, K. Cui, and K. Zhao. Genome-wide identification of in vivo protein-DNA binding sites from ChIP-Seq data. *Nucleic Acids Research*, 36:5221–31, 2008.

P. V. Kharchenko, M. Y. Tolstorukov, and P. J. Park. Design and analysis of ChIP experiments for DNA-binding proteins. *Nature Biotechnology*, 26:1351–1359, 2008.

Examples

```
data(cstest)
estimate.mean.fraglen(cstest[["ctcf"]], method = "coverage")
```

islandDepthPlot	<i>Plot island depth distribution</i>
-----------------	---------------------------------------

Description

Plots the distribution of island depths using points for the observed islands and a line for the Poisson estimate of the noise. Useful for choosing a depth corresponding to a desired FDR.

Usage

```
islandDepthPlot(x, maxDepth = 20L)
```

Arguments

x	A coverage object, e.g., RleList .
maxDepth	The maximum depth to plot (there are usually some outliers).

Author(s)

D. Sarkar, M. Lawrence

See Also

[peakCutoff](#) for calculating a cutoff value for an FDR.

Examples

```
data(cstest)
cov <- coverage(resize(cstest$ctcf, width=200))
islandDepthPlot(cov)
```

laneSubsample	<i>Subsample short read alignment locations</i>
---------------	---

Description

Subsamples data from multiple lanes on a per-chromosome basis.

Usage

```
laneSubsample(lane1, lane2, fudge = 0.05)
```

Arguments

lane1, lane2	Two lanes of data, each of class "GRanges".
fudge	A numeric fudge factor. For each chromosome, if the difference in the sizes relative to the size of the first dataset is less than fudge, no subsampling is done.

Value

laneSubsample returns a list similar to its input, but with the larger dataset subsampled to be similar to the smaller one.

Author(s)

D. Sarkar

Examples

```
data(cstest)
## subsample to compare lanes
cstest.sub <- laneSubsample(cstest[[1]], cstest[[2]])
unlist(cstest.sub)
```

peakCutoff	<i>Calculate a peak cutoff</i>
------------	--------------------------------

Description

Calculates a peak cutoff value given an FDR, assuming a Poisson noise distribution estimated from the frequency of singleton and doubleton islands.

Usage

```
peakCutoff(cov, fdr.cutoff = 0.001, k = 2:20)
```

Arguments

cov	The coverage object, e.g., an RleList object.
fdr.cutoff	The maximum-allowed FDR for calculating the cutoff.
k	The coverage levels at which to estimate an FDR value. The maximal value that is less than fdr.cutoff is chosen for calculating the cutoff. Usually best left to the default.

Value

A numeric value to use for calling peaks

Author(s)

D. Sarkar and M. Lawrence

See Also

[islandDepthPlot](#) for the graphical equivalent; the vignette for a bit more explanation.

Examples

```
data(cstest)
cov <- coverage(resize(cstest$ctcf, width=200))
peakCutoff(cov)
```

peakSummary-methods *Summarizing peak sets*

Description

Summarizes a set of peaks into a [GRanges](#) object with columns of statistics like the peak maxima and integrals (sums).

Usage

```
peakSummary(x, ...)
```

Arguments

x	An object containing peaks, usually a RleViewsList .
...	Arguments to pass to methods

Value

A [GRanges](#) object of the peaks, with columns named max, maxpos (position of the maximum, centered), and sum.

See Also

[view-summarization-methods](#) in the IRanges package for view summarization methods like viewMaxs and viewSums.

subsetSummary	<i>Compute summaries for cumulative subsets of a short-read data set.</i>
---------------	---

Description

THIS FUNCTION IS DEFUNCT!

Divides a short-read dataset into several subsets, and computes various summaries cumulatively. The goal is to study the characteristics of the data as a function of sample size.

Usage

```
subsetSummary(x, chr, nstep, props = seq(0.1, 1, 0.1),
  chromlens = seqlengths(x), fg.cutoff = 6, seqLen = 200,
  fdr.cutoff = 0.001, use.fdr = FALSE, resample = TRUE,
  islands = TRUE, verbose = getOption("verbose"))
```

Arguments

x	A "GRanges" object representing alignment locations at the sample level.
chr	The chromosome for which the summaries are to be obtained. Must specify a valid element of x
nstep	The number of maps in each increment for the full dataset (not per-chromosome). This will be translated to a per-chromosome number proportionally.
props	Alternatively, an increasing sequence of proportions determining the size of each subset. Overrides nstep.
chromlens	A named vector of per-chromosome lengths, typically the result of seqlengths .
fg.cutoff	The coverage depth above which a region would be considered foreground.
seqLen	The number of bases to which to extend each read before computing coverage.
resample	Logical; whether to randomly reorder the reads before dividing them up into subsets. Useful to remove potential order effects (for example, if data from two lanes were combined to produce x).
fdr.cutoff	The maximum false discovery rate for a region that is considered to be foreground.
use.fdr	Whether to use the FDR detected peaks when calling foreground and background.
islands	Logical. If TRUE, the whole island would be considered foreground if the maximum depth equals or exceeds fg.cutoff. If FALSE, only the region above the cutoff would be considered foreground.
verbose	logical controlling whether progress information will be shown during computation (which is potentially long-running).

Value

A data frame with various per-subset summaries.

Note

This function should be considered preliminary, in that it might change significantly or simply be removed in a subsequent version. If you like it the way it is, please notify the maintainer.

Author(s)

Deepayan Sarkar, Michael Lawrence

Index

- * **datasets**
 - cstest, [3](#)
- * **hplot**
 - coverageplot, [3](#)
- * **manip**
 - laneSubsample, [8](#)
- * **methods**
 - peakSummary-methods, [9](#)
- * **univar**
 - estimate.mean.fraglen, [5](#)
 - subsetSummary, [10](#)
- * **utilities**
 - laneSubsample, [8](#)

AlignedRead, [6](#)

basesCovered (estimate.mean.fraglen), [5](#)

bsapply, [2](#)

chipseqFilter, [2](#)

coverageplot, [3](#)

cstest, [3](#)

density, [5](#)

densityCorr (estimate.mean.fraglen), [5](#)

densityCorr, GenomicRanges
(estimate.mean.fraglen), [5](#)

densityCorr, list
(estimate.mean.fraglen), [5](#)

diffPeakSummary, [4](#)

diffPeakSummary, RleViewsList, RleViewsList-method
(diffPeakSummary), [4](#)

estimate.mean.fraglen, [5](#)

estimate.mean.fraglen, AlignedRead-method
(estimate.mean.fraglen), [5](#)

estimate.mean.fraglen, GRanges-method
(estimate.mean.fraglen), [5](#)

GRanges, [6](#), [9](#)

islandDepthPlot, [7](#), [9](#)

laneSubsample, [8](#)

peakCutoff, [7](#), [8](#)

peakSummary (peakSummary-methods), [9](#)

peakSummary, RleViews-method
(peakSummary-methods), [9](#)

peakSummary, RleViewsList-method
(peakSummary-methods), [9](#)

peakSummary-methods, [9](#)

readAligned, [2](#)

RleList, [7](#), [9](#)

RleViewsList, [4](#), [9](#)

seqlengths, [10](#)

sparse.density (estimate.mean.fraglen),
[5](#)

SRFilter, [2](#)

subsetSummary, [10](#)

view-summarization-methods, [10](#)