# Package 'ChIPQC'

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```
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     TxDb.Mmusculus.UCSC.mm10.knownGene,
     TxDb.Mmusculus.UCSC.mm9.knownGene,
     TxDb.Rnorvegicus.UCSC.rn4.ensGene,
     TxDb.Celegans.UCSC.ce6.ensGene,
     TxDb.Dmelanogaster.UCSC.dm3.ensGene
Suggests BiocStyle
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     ChIPQC_IF.R plots.r dbaplots.R utilities.R report.r
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ChIPQC-package

ChIPQC - Quality metrics for ChIPseq data

#### **Description**

ChIPQC analyzes aligned reads (in .bam format) for ChIP-seq samples and their associated controls, computing a variety of quality control metrics and statistics, and providing reporting and plotting functions to enable assessment of experimental data for further analysis.

### **Details**

Package: ChIPQC
Type: Package
Version: 0.1

Date: 2014-03-01 License: GPL3

ChIPQC primarily uses two object classes: ChIPQCsample, which encapsulates the information about individual samples, and ChIPQCexperiment, which encapsulates information about larger ChIP-seq experiments (consisting of a number of samples). The primary entry point is the constructor function ChIPQC, which takes a description of an entire experiment, constructs objects for all the samples, and computes the quality metrics.

### Author(s)

Tom Carroll and Rory Stark

Maintainers: Tom Carroll <tc.infomatics@gmail.com> and Rory Stark <rory.stark@cruk.cam.ac.uk>

### References

Frontiers?

#### See Also

ChIPQC is designed to work closely with the DiffBind package, which provides functionality for analyzing ChIP-seq experiments, including performing differential binding analysis to identify significantly differentially bound peaks.

average peak signal-methods

Retrieve average peak profiles

### **Description**

Retrieve the average peak profile for a sample or set of samples.

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#### Methods

signature(object = "ChIPQCexperiment") Retrieve a matrix of the average peak profiles for all of the samples in an ChIP-seq experiment. Each column represents a sample, and each row a base pair position, centered on peak summits.

signature(object = "list") Retrieve a matrix of the average peak profiles for all of the samples in a list of ChIPQC samples. Each column represents a sample, and each row a base pair position, centered on peak summits.

signature(object = "ChIPQCsample") Retrieve a vector representing the average peak profile for a sample. Each column represents a basepair position, centered on the peak summits.

#### Author(s)

Thomas Carroll and Rory Stark

#### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

### **Examples**

```
data(example_QCexperiment)
CTCFprofile = averagepeaksignal(QCsample(exampleExp,1))
length(CTCFprofile)
plot(CTCFprofile,type='l',ylab="mean pileup")
allprofiles = averagepeaksignal(exampleExp)
dim(allprofiles)
for(i in 1:ncol(allprofiles)) lines(allprofiles[,i],col=i)
```

ChIPQC

Construct a ChIPQCexperiment object

### Description

Constructs a new ChIPQCexperiment object.

### Usage

### Arguments

experiment

A specification of the ChIP-seq experiment to evaluate. This can either be a dataframe, a filename for a .csv file, or a DBA-object object as defined in the DiffBind package. Columns names in sample sheet may include:

- SampleID: Identifier string for sample
- Tissue: Identifier string for tissue type
- Factor: Identifier string for factor

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- Condition: Identifier string for condition
- Treatment: Identifier string for treatment
- Replicate: Replicate number of sample
- bamReads: file path for bam file containing aligned reads for ChIP sample
- bamControl: file path for bam file containing aligned reads for control sample
- ControlID: Identifier string for control sample
- Peaks: path for file containing peaks for sample. Format determined by PeakCaller field or caller parameter
- PeakCaller: Identifier string for peak caller used. If Peaks is not a bed file, this will determine how the Peaks file is parsed. If missing, will use default peak caller specified in caller parameter. Possible values:
  - "raw": text file file; peak score is in fourth column
  - "bed": .bed file; peak score is in fifth column
  - "narrow": default peak.format: narrowPeaks file
  - "macs": MACS .xls file
  - "swembl": SWEMBL .peaks file
  - "bayes": bayesPeak file
  - "fp4": FindPeaks v4
- PeakFormat: string indicating format for peak files; see PeakCaller and dba.peakset
- ScoreCol: column in peak files that contains peak scores
- LowerBetter: logical indicating that lower scores signify better peaks

See the documentation for the sampleSheet parameter of dba for details.

annotation

Either a character string indicating the genome and version to use as a genomic annotation, or a previously defined annotation (obtained using QCannotation on a previously defined ChIPQCexperiment object.) May be left unspecified, in which case no genomic feature analysis is performed. The following annotation specifiers are supported:

- "hg19": Human, version 19
- "hg18": Human, version 18
- "mm10": Mouse, version 10
- "mm9": Mouse, version 19
- "rn4": Rat, version 4
- "ce6": C. Elgans, version 6
- "dm3": D. Melanogaster, version 3

Alternatively, you can construct your own annotation; see the package vignette for more information.

chromosomes

Specification of which chromosomes to use for computing QC statistics. If missing, the first chromosome which has a peak is checked. If NULL, all chromosomes will be checked (which may be time-consuming). This can be a character string (e.g. "chr18") or a vector or list of character strings. If it is an integer or vector of integers, the chromosomes will be checked based on the order that they are listed in a peak set.

samples

list of ChIPCsample objects. If present, the sample objects will be taken directly from this list instead of being computed using the ChIPQCsample constructor.

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consensus If consensus is a GRanges object, all samples will use this peakset when com-

puting peak-based metrics. If consensus=TRUE, a consensus peakset will be generated and used for all samples, derived by merging overlapping peaks in all provided peaksets, keeping any peaks that overlap in at least two samples To avoid this behavior, set consensus=FALSE; this will result in only supplied peaksets being used for calculation of peak-based metrics (and no peak-based metric being computed for samples with no peakset specified, such as controls).

bCount if TRUE, the peak scores for all samples will be based on read counts using

dba. count using a consensus peakset. If consenus is missing, any samples (such as controls) that are not already associated with a peakset will be associated with the consensus peakset (if consensus is not missing, all samples will be associated with the consensus peakset). Note that the re-counting process

may be time-consuming.

mapQCth An integer representing a mapping quality score threshold. Only reads with

mapping quality scores above this threshold will be used for some statistics.

blacklist A GRanges object or filename specifying a bed file containing genomic regions

that should be excluded from the analysis. If missing and the annotation is "hg19", a default blacklist, blacklist\_hg19 derived from the UCSC list, will be used. No blacklist is used if this is set to NULL, or is left missing and the

annotation is not "hg19".

profileWin An integer indicating the width, in base pairs, of the window to be used for peak

profiles. Peaks will be centered on their summits, and include half the window

size upstream and half downstream of this point.

fragmentLength An integer indicating the expected fragment length of the libraries. Optional, as

this value will be computed for each library.

shifts A vector of values to try when computing optimal shift sizes.

... additional parameters passed to dba. count if bCount=TRUE.

### Details

ChIPQC first constructs a new DBA-object object if one is not provided. Next it computes the annotation if one is not provided. The main loop constructs new ChIPQCsample objects for each sample (and unique control sample).

#### Value

A ChIPQCexperiment object.

#### Note

ChIPQC uses bplapply from the BiocParallel package to build the ChIPQCsample object in parallel, if supported. Control of the parallelization can be effected using BiocParallel functions, such as register.

### Author(s)

Thomas Carroll and Rory Stark

### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample, DiffBind

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### **Examples**

```
## Not run: exampleExp = ChIPQC(samples,annotation="hg19")
data(example_QCexperiment)
exampleExp

## Not run: tamoxifen = ChIPQC(samples, ,annotation="hg19", consensus=TRUE, bCounts=T)
data(tamoxifen_QC)
tamoxifen
```

ChIPQC-data

Example data sets for ChIPQC package, each containing a ChIPQC experiment object, as well as a pre-compiled blacklist for hg19.

### **Description**

The tamoxifen\_QC example data set contains a ChIPQCexperiment object named tamoxifen. This data set, based on Ross-Innes et al (2012), includes 11 ER ChIP libraries, and their input controls, divided into tamoxifen responsive and tamoxifen resistant sample groups. Only data for chromosome 18 (chr18) are included.

The example\_QCexperiment data set contains a ChIPQCexperiment object named exampleExp. This data set, derived from ENCODE data, includes 6 ChIP libraries. Only data for chromosome 22 (chr22) are included.

blacklist\_hg19 data set includes a GRanges object named blacklist\_hg19 containing blacklisted regions for the human genome, derived from the UCSC blacklist.

### Usage

```
data(tamoxifen_QC)
data(example_QCexperiment)
data(blacklist_hg19)
```

### **Format**

tamoxifen\_QC: A single ChIPQCexperiment object named tamoxifen is loaded. This object is used for the ChIPQC-package examples and vignette. This object can also be used with the DiffBind package (see related data objects tamoxifen).

example\_QCexperiment: A single ChIPQCexperiment object named exampleExp is loaded. This object is used for the ChIPQC-package examples and vignette.

blacklist\_hg19: A single GRanges object named blacklist.hg19 that is used by default when processing hg19 data sets.

#### Source

Ross-Innes, C. S., Stark, R., Teschendorff, A. E., Holmes, K. A., Ali, H. R., Dunning, M. J., Brown, G. D., Gojis, O., Ellis, I. O., Green, A. R., Ali, S., Chin, S.-F., Palmieri, C., Caldas, C., and Carroll, J. S. (2012). Differential oestrogen receptor binding is associated with clinical outcome in breast cancer. Nature 481, 389-393.

#### **Examples**

```
data(tamoxifen_QC)
tamoxifen
plotRegi(tamoxifen)

data(example_QCexperiment)
exampleExp

data(blacklist_hg19)
blacklist.hg19
```

ChIPQCexperiment-class

ChIPQCexperiment instances

### Description

The ChIPQCexperiment class is built around a list of ChIPQCsample objects, each representing a ChIP or control sample in a ChIP-seq experiment. These objects are created using the ChIPQC function.

#### **Slots**

```
.Data: Object of class "list": internal
Samples: Object of class "list": List of ChIPQCsample objects.

DBA: Object of class "DBA": DBA-object object (from package DiffBind)
annotation: Object of class "list": annotation data
```

### Extends

Class "list".

### Methods

```
QCmetrics signature(object = "ChIPQCexperiment"): see QCmetrics.

QCsample signature(object = "ChIPQCexperiment"): see QCsample.

QCcontrol signature(object = "ChIPQCexperiment"): see QCcontrol.

QCannotation signature(object = "ChIPQCexperiment"): see QCannotation.

QCdba signature(object = "ChIPQCexperiment"): see QCdba.

averagepeaksignal signature(object = "ChIPQCexperiment"): see averagepeaksignal.

coveragehistogram signature(object = "ChIPQCexperiment"): see coveragehistogram.

crosscoverage signature(object = "ChIPQCexperiment"): see crosscoverage.

flagtagcounts signature(object = "ChIPQCexperiment"): see flagtagcounts.

fragmentlength signature(object = "ChIPQCexperiment"): see fragmentlength.

FragmentLengthCrossCoverage signature(object = "ChIPQCexperiment"): see FragmentLengthCrossCoverage

frip signature(object = "ChIPQCexperiment"): see frip.
```

```
mapped signature(object = "ChIPQCexperiment"): see mapped.
reads signature(object = "ChIPQCexperiment"): see reads.
duplicates signature(object = "ChIPQCexperiment"): see duplicates.
duplicateRate signature(object = "ChIPQCexperiment"): see duplicateRate.
Normalisedaveragepeaksignal signature(object = "ChIPQCexperiment"): see Normalisedaveragepeaksignal.
peaks signature(object = "ChIPQCexperiment"):see peaks.
readlength signature(object = "ChIPQCexperiment"): see readlength.
ReadLengthCrossCoverage signature(object = "ChIPQCexperiment"): see ReadLengthCrossCoverage.
RelativeCrossCoverage signature(object = "ChIPQCexperiment"):see RelativeCrossCoverage.
ribl signature(object = "ChIPQCexperiment"): see ribl.
rip signature(object = "ChIPQCexperiment"): see rip.
show signature(object = "ChIPQCexperiment"): see show.
ssd signature(object = "ChIPQCexperiment"): see ssd.
regi signature(object = "ChIPQCexperiment"): see regi.
plotCC signature(object = "ChIPQCexperiment"): see plotCC.
plotCoverageHist signature(object = "ChIPQCexperiment"): see plotCoverageHist.
plotFribl signature(object = "ChIPQCexperiment"): see plotFribl.
plotPeakProfile signature(object = "ChIPQCexperiment"): see plotPeakProfile.
plotRap signature(object = "ChIPQCexperiment"): see plotRap.
plotRegi signature(object = "ChIPQCexperiment"): see plotRegi.
plotCorHeatmap signature(object = "ChIPQCexperiment"): see plotCorHeatmap.
plotPrincomp signature(object = "ChIPQCexperiment"): see plotPrincomp.
ChIPQCreport signature(object = "ChIPQCexperiment"): see ChIPQCreport.
```

#### Author(s)

Thomas Carroll and Rory Stark

#### See Also

ChIPQC-package, ChIPQCsample, DiffBind

```
## Not run: exampleExp = ChIPQC(samples)
data(example_QCexperiment)
exampleExp

## Not run: tamoxifen = ChIPQC(samples, consensus=TRUE, bCounts=T)
data(tamoxifen_QC)
tamoxifen
```

ChIPQCreport-methods Generate a summary QC report

### **Description**

Generate an HTML summary QC report.

#### Methods

signature(experiment = "ChIPQCexperiment", facet=TRUE, reportName="ChIPQC", reportFolder="ChIPQCre Generates an summary QC report for the experiment in HTML format.

experiment ChIPQCexperiment object

facet logical indicating whether or not to facet using experimental metadata.

reportName filename of main report file (.html).

reportFolder directory name where plot graphics will be saved

facetBy metadata fields to use for faceting colourBy metadata field to color by

signature(experiment = "list", facet=TRUE, reportName="ChIPQC", reportFolder="ChIPQCreport", facetE Generates an summary QC report for a list of ChIPQCsample objects in HTML format.

experiment list object

facet logical indicating whether or not to facet using experimental metadata.

reportName filename of main report file (.html).

reportFolder directory name where plot graphics will be saved

facetBy metadata fields to use for faceting

colourBy metadata field to color by

signature(sample = "ChIPQCsample", reportName="ChIPQC", reportFolder="ChIPQCreport",)
Generate a summary QC report for a sample in HTML format.

sample ChIPQCsample object

reportName filename of main report file (.html).

reportFolder directory name where plot graphics will be saved

### Note

ChIPQCreport uses Nozzle.R2 for generating HTML.

### Author(s)

Thomas Carroll and Rory Stark

### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

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#### **Examples**

```
data(example_QCexperiment)
ChIPQCreport(exampleExp,facetBy=c("Tissue","Factor"))
#report in ChIPQCreport/Example.html

data(tamoxifen_QC)
ChIPQCreport(tamoxifen,facetBy="Tissue",colourBy="Condition")
#report in ChIPQCreport/ChIPQC.html

ChIPQCreport(tamoxifen,facetBy=c("Tissue","Condition"))
#report in ChIPQCreport/ChIPQC.html
```

ChIPQCsample-class

Class "ChIPQCsample"

### **Description**

Object containing quality metrics computed for a ChIP-seq (or associated control) sample.

### **Objects from the Class**

Objects can be created using the ChIPQCsample function.

### **Constructor Function**

ChIPQCsample(reads, peaks, annotation, chromosomes = NULL, mapQCth = 15, blacklist, profileWin = 400, fragmentLength = 125, shifts = 1:300, runCrossCor = FALSE, verboseT=FALSE)

- · readscharacter string filename of .bam file
- peaksGRanges object or character string filename of peaks. If present, peak-based metrics will be computed.
- annotation Either a character string indicating the genome and version to use as a genomic
  annotation, or a previously defined annotation (obtained using QCannotation on a previously
  defined ChIPQCexperiment object.) May be left unspecified, in which case no genomic feature analysis is performed. The following annotation specifiers are supported:

```
"hg20"
           Human, version 20
"hg19"
           Human, version 19
"hg18"
           Human, version 18
"mm10"
          Mouse, version 10
"mm9"
          Mouse, version 19
"rn4"
           Rat, version 4
"ce6"
           C. Elgans, version 6
"dm3"
          D. Melanogaster, version 3
```

Alternatively, you can construct your own annotation; see the package Vignette for more information.

• chromosomes Specification of which chromosomes to use for computing QC statistics. If missing, the first chromosome which has a peak is checked. If NULL, all chromosomes will be checked (which may be time-consuming). This can be a character string (e.g. "chr18") or a vector or list of character strings. If it is an integer or vector of integers, the chromosomes will be checked based on the order that they are listed in a peak set.

- mapQCth An integer representing a mapping quality score threshold. Only reads with mapping quality scores above this threshold will be used for some statistics.
- blacklist A GRanges object or filename specifying a bed file containing genomic regions that should be excluded from the analysis. If missing and the annotation is "hg19", a default blacklist, blacklist\_hg19 derived from the UCSC list, will be used. No blacklist is used if this is set to NULL, or is left missing and the annotation is not "hg19".
- profileWin An integer indicating the width, in base pairs, of the window to be used for peak profiles. Peaks will be centered on their summits, and include half the window size upstream and half downstream of this point.
- fragmentLength An integer indicating the expected fragment length of the libraries. Optional, as this value will be computed.
- shifts A vector of values to try when computing optimal shift sizes.
- runCrossCor Compute cross-correlation in addition to cross-coverage. This will take more compute time, and is currently not used in the final report.
- verboseT TRUE or FALSE, specifying whether to report progress. Default is TRUE. When set to FALSE ChIPQC does not report any progress until complete.

#### **Slots**

AveragePeakSignal: Object of class "list" CrossCoverage: Object of class "numeric" CrossCorrelation: Object of class "numeric" SSD: Object of class "numeric" SSDBL: Object of class "numeric" CountsInPeaks: Object of class "numeric" CountsInBlackList: Object of class "numeric" CountsInFeatures: Object of class "list" PropInFeatures: Object of class "list" CoverageHistogram: Object of class "numeric" FlagAndTagCounts: Object of class "numeric" readlength: Object of class "numeric" segnames: Object of class "Rle" ranges: Object of class "IRanges" strand: Object of class "Rle" elementMetadata: Object of class "DataFrame" seginfo: Object of class "Seginfo" metadata: Object of class "list"

### **Extends**

Class "GRanges"

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#### Methods

```
averagepeaksignal signature(object = "ChIPQCsample"): see averagepeaksignal.
coveragehistogram signature(object = "ChIPQCsample"): see coveragehistogram.
crosscoverage signature(object = "ChIPOCsample"): see crosscoverage.
flagtagcounts signature(object = "ChIPQCsample"): see flagtagcounts.
fragmentlength signature(object = "ChIPQCsample"): see fragmentlength.
FragmentLengthCrossCoverage signature(object = "ChIPQCsample"): see FragmentLengthCrossCoverage.
frip signature(object = "ChIPQCsample"): see frip.
mapped signature(object = "ChIPQCsample"): see mapped.
reads signature(object = "ChIPQCsample"): see reads.
duplicates signature(object = "ChIPQCsample"): see duplicates.
duplicateRate signature(object = "ChIPQCsample"): see duplicateRate.
Normalisedaveragepeaksignal signature(object = "ChIPQCsample"): see Normalisedaveragepeaksignal.
peaks signature(object = "ChIPQCsample"):see peaks.
readlength signature(object = "ChIPQCsample"): see readlength.
ReadLengthCrossCoverage signature(object = "ChIPQCsample"): see ReadLengthCrossCoverage.
RelativeCrossCoverage signature(object = "ChIPQCsample"):see RelativeCrossCoverage.
ribl signature(object = "ChIPQCsample"): see ribl.
rip signature(object = "ChIPQCsample"): see rip.
show signature(object = "ChIPQCsample"): see show.
ssd signature(object = "ChIPQCsample"): see ssd.
regi signature(object = "ChIPQCsample"): see regi.
plotCC signature(object = "ChIPQCsample"): see plotCC.
plotCoverageHist signature(object = "ChIPQCsample"): see plotCoverageHist.
plotFribl signature(object = "ChIPQCsample"): see plotFribl.
plotPeakProfile signature(object = "ChIPQCsample"): see plotPeakProfile.
plotRap signature(object = "ChIPQCsample"): see plotRap.
plotRegi signature(object = "ChIPQCsample"): see plotRegi.
```

### Author(s)

Thomas Carroll and Rory Stark

#### References

Carroll TS, Liang Z, Salama R, Stark R and Santiago Id (in press). Impact of artefact removal on ChIP quality metrics in ChIP-seq and ChIP-exo data. Frontiers in Genetics.

### See Also

```
ChIPQC-package, ChIPQCsample
```

#### **Examples**

coveragehistogram-methods

Retrieve histogram data representing densities of coverage pileups

### **Description**

Retrieve histogram data representing densities of coverage pileups.

### Methods

- signature(object = "ChIPQCexperiment") Retrieve a matrix of coverage histogram data for all samples in a ChIP-seq experiment. Each column represents a sample, and each row a pileup height, with the value representing the number of basepair positions that report this pileup height.
- signature(object = "list") Retrieve a matrix of coverage histogram data for all ChIPQCsamples in a list. Each column represents a sample, and each row a pileup height, with the value representing the number of basepair positions that report this pileup height.
- signature(object = "ChIPQCsample") Retrieve a vector representing coverage histogram data for a sample. Values represent the number of base pairs positions that report the pileup value. The value in position 1 of the vector the contains the number of examined basepair positions that are overlapped by exactly zero reads, while position 2 shows the number of basepair positions overlapped by exactly one read, etc.

### Author(s)

Thomas Carroll and Rory Stark

### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

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crosscoverage-methods Retrieve the cross coverage values for a range of shift sizes

### Description

Retrieves the cross-coverage values for a range of shift sizes.

#### Methods

signature(object = "ChIPQCexperiment") Retrieve a matrix of cross-coverage data for all samples in an ChIP-seq experiment. Each column represents a sample, and each row a shift size, with the value representing the cross-coverage using that size read.

signature(object = "list") Retrieve a matrix of cross-coverage data for all samples in a list of ChIPQCsample objects. Each column represents a sample, and each row a shift size, with the value representing the cross-coverage using that size read.

signature(object = "ChIPQCsample") Retrieve a vector of cross-coverage data for a sample. Each position in the vector corresponds to a shift size, with the value representing the cross-coverage using that size read.

#### Author(s)

Thomas Carroll and Rory Stark

#### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

### **Examples**

```
data(example_QCexperiment)
CTCFcoverage = crosscoverage(QCsample(exampleExp,1))
length(CTCFcoverage)
plot(CTCFcoverage,type='1',
    ylab="Cross-coverage",
    xlab="Fragment length")

allcoverages = crosscoverage(exampleExp)
dim(allcoverages)
for(i in 1:ncol(allcoverages)) lines(allcoverages[,i],col=i)
```

duplicateRate-methods Retrieve duplication rates

### Description

Retrieve duplication rates.

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#### Methods

signature(object = "ChIPQCexperiment", bFiltered) Retrieve a vector of the duplication rates for each sample in an experiment. A read is considered duplicated if another read maps to the same genomic location; the duplication rate is the number of duplicated reads divided by the total number of reads for a sample. If bFiltered=TRUE (or is missing), only reads that pass the mapping quality filter for each sample are included. if bFiltered=FALSE, all reads for each sample will be included.

signature(object = "list", bFiltered) Retrieve a vector of the duplication rates for each sample in a list of ChIPQCsample objects. A read is considered duplicated if another read maps to the same genomic location; the duplication rate is the number of duplicated reads divided by the total number of reads for a sample. If bFiltered=TRUE (or is missing), only reads that pass the mapping quality filter for each sample are included. if bFiltered=FALSE, all reads for each sample will be included.

signature(object = "ChIPQCsample", bFiltered) Retrieve the duplication rate for a sample. A read is considered duplicated if another read maps to the same genomic location; the duplication rate is the number of duplicated reads divided by the total number of reads for the sample. If bFiltered=TRUE (or is missing), only reads that pass the mapping quality filter for the sample are included. if bFiltered=FALSE, all reads for the sample will be included.

#### Author(s)

Thomas Carroll and Rory Stark

#### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

### **Examples**

```
data(example_QCexperiment)
duplicateRate(exampleExp)
duplicateRate(QCsample(exampleExp,1))
```

duplicates-methods

Retrieve numbers of duplicate reads.

### **Description**

Retrieve the numbers of duplicate reads.

### Methods

signature(object = "ChIPQCexperiment", bFiltered) Retrieve a vector of the numbers of duplicate reads for each sample in an experiment. A read is considered duplicated if another read maps to the same genomic location. If bFiltered=TRUE (or is missing), this will be the number of duplicates that pass the mapping quality filter for each sample. if bFiltered=FALSE, it will be the total number of duplicates for each sample.

signature(object = "list", bFiltered) Retrieve a vector of the numbers of duplicate reads for each sample in a list of ChIPQCsample objects. A read is considered duplicated if another read maps to the same genomic location. If bFiltered=TRUE (or is missing), this will be the number of duplicates that pass the mapping quality filter for each sample. if bFiltered=FALSE, it will be the total number of duplicates for each sample.

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signature(object = "ChIPQCsample", bFiltered) Retrieve the number of duplicates for a sample. A read is considered duplicated if another read maps to the same genomic location. If bFiltered=TRUE (or is missing), this will be the number of duplicates that pass the mapping quality filter. if bFiltered=FALSE, it will be the total number of duplicates for the sample.

### Author(s)

Thomas Carroll and Rory Stark

#### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

### **Examples**

```
data(example_QCexperiment)
duplicates(exampleExp)
duplicates(QCsample(exampleExp,1))
```

flagtagcounts-methods Retrieve numbers of reads that pass various filters

### **Description**

Retrieve numbers of reads that pass various filters

### Methods

signature(object = "ChIPQCexperiment") Retrieve a matrix of counts passing various filters for all the samples in an experiment. Each column represents the counts in a sample.

signature(object = "list") Retrieve a matrix of counts passing various filters for all the samples in a list of ChIPQCsample objects. Each column represents the counts in a sample.

signature(object = "ChIPQCsample") Retrieve a vector of counts passing various filters. The
values are:

UnMapped Number of reads that are not mapped (aligned)
Mapped Number of reads that are mapped (aligned)

Duplicates Number of reads that align to exactly the same place as another read

MapQPass Number of reads with a mapping quality score greater than or equal to the specified threshold

MapQPassandDup Number of reads that are mapped (aligned) and not duplicates

#### Author(s)

Thomas Carroll and Rory Stark

### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

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#### **Examples**

```
data(example_QCexperiment)
flagtagcounts(exampleExp)
flagtagcounts(QCsample(exampleExp,1))
```

fragmentlength-methods

Retrieve the estimated fragment length

### **Description**

Retrieve the estimated fragment length.

#### Methods

signature(object = "ChIPQCexperiment") Retrieve a vector of estimated fragments sizes, one
for each sample in the experiment.

signature(object = "list") Retrieve a vector of estimated fragments sizes, one for each sample in a list of ChIPQCsample objects.

signature(object = "ChIPQCsample", width) Retrieve the estimated fragment length for a sample. If width is missing, the readlength derived from the bam file is used as the read length.

### Author(s)

Thomas Carroll and Rory Stark

#### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

### **Examples**

```
data(example_QCexperiment)
fragmentlength(exampleExp)
fragmentlength(QCsample(exampleExp,1))
```

frip-methods

Retrieve fraction of reads in peaks

### **Description**

Retrieve the fraction of reads in peaks

#### Methods

signature(object = "ChIPQCexperiment") Retrieve a vector of values representing the proportion of reads that overlap peaks for each sample in an experiment.

signature(object = "list") Retrieve a vector of values representing the proportion of reads that overlap peaks for each sample in a list of ChIPQCsample objects.

signature(object = "ChIPQCsample") Retrieve a value representing the proportion of reads that overlap the peaks for a sample.

mapped-methods 19

### Author(s)

Thomas Carroll and Rory Stark

### See Also

```
ChIPQC-package, ChIPQCexperiment, ChIPQCsample
```

### **Examples**

```
data(example_QCexperiment)
frip(exampleExp)
frip(QCsample(exampleExp,1))
```

mapped-methods

Retrieve numbers of mapped reads

### **Description**

Retrieve the numbers of mapped reads.

### Methods

```
signature(object = "ChIPQCexperiment") Retrieve a vector of the numbers of mapped (aligned)
reads for each sample in an experiment.
```

signature(object = "list") Retrieve a vector of the numbers of mapped (aligned) reads for each sample in a list of ChIPQCsample objects.

signature(object = "ChIPQCsample") Retrieve the number of mapped (aligned) reads in a sample.

### Author(s)

Thomas Carroll and Rory Stark

### See Also

```
ChIPQC-package, ChIPQCexperiment, ChIPQCsample
```

```
data(example_QCexperiment)
mapped(exampleExp)
mapped(QCsample(exampleExp,1))
```

20 peaks-methods

Normalisedaveragepeaksignal-methods

Retrieve normalised average peak profiles

### **Description**

Retrieve normalised average peak profiles

#### Methods

signature(object = "ChIPQCexperiment") Retrieve a matrix of normalised average peak signal data for all samples in a ChIP-seq experiment. Each column represents a sample, and each row a base pair position, centered on peak summits.

signature(object = "list") Retrieve a matrix of normalised average peak signal data for all samples in a list of ChIPQCsample objects. Each column represents a sample, and each row a base pair position, centered on peak summits.

signature(object = "ChIPQCsample") Retrieve a vector representing the normalised average peak profile for a sample. Each column represents a basepair position, centered on the peak summits.

### Author(s)

Thomas Carroll and Rory Stark

### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

### **Examples**

```
data(example_QCexperiment)
CTCFprofile = Normalisedaveragepeaksignal(QCsample(exampleExp,1))
length(CTCFprofile)
plot(CTCFprofile, type='l',ylab="normalised mean pileup")
allprofiles = Normalisedaveragepeaksignal(exampleExp)
dim(allprofiles)
for(i in 1:ncol(allprofiles)) lines(allprofiles[,i],col=i)
```

peaks-methods

Retrieve peaks

### Description

Retrieve peaks.

plotCC-methods 21

#### Methods

signature(object = "ChIPQCexperiment") Retrieve a GRangesList of the peaks associated with all the samples in an experiment.

signature(object = "list") Retrieve a GRangesList of the peaks associated with all the samples in a list of ChIPQCsample objects.

signature(object = "ChIPQCsample") Retrieve a GRanges object containing the peaks associated with a sample.

### Author(s)

Thomas Carroll and Rory Stark

### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

### **Examples**

```
data(example_QCexperiment)
expPeaks = peaks(exampleExp)
length(expPeaks)
peaks(QCsample(exampleExp,1))
```

plotCC-methods

Generate Cross-Coverage plots

### **Description**

Generate Cross-Coverage plots.

### Methods

signature(object = "ChIPQCexperiment", method) Generate cross-coverage plots for all the samples in an experiment.

signature(object = "list", method) Generate cross-coverage plots for list of samples in an
 experiment.

signature(object = "ChIPQCsample", methods) Generate cross-coverage plots for a sample.
Supported methods include:

"Coverage" [default] Coverage plot

### Note

plotCC uses ggplot2 for plotting, and returns a ggplot2 plot dataframe.

### Author(s)

Thomas Carroll and Rory Stark

#### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

### Examples

```
data(example_QCexperiment)
plotCC(exampleExp)
plotCC(exampleExp,excludedBox=TRUE)
plotCC(QCsample(exampleExp,1))
plotCC(QCsample(exampleExp)[1:4])
```

plotCorHeatmap-methods

Generate Correlation Heatmap for ChIP samples

### **Description**

Generates correlation heatmap for ChIP samples.

### Methods

signature(object = "ChIPQCexperiment", attributes, ...) Generate correlation heatmap, including clustering dendrogram, for all the samples in an experiment that are associated with a peakset.

attributes character string, or vector of character strings, representing metadata field names, for use in labeling additional parameters passed to dba.plotHeatmap

### Note

plotCorHeatmap uses dba.plotHeatmap for plotting.

### Author(s)

Rory Stark and Thomas Carroll

### See Also

ChIPQC-package, ChIPQCexperiment, dba.plotHeatmap

```
data(tamoxifen_QC)
plotCorHeatmap(tamoxifen,attributes=c("Tissue","Condition","Replicate"))
```

plotCoverageHist-methods

Generate coverage histogram plot

### **Description**

Generate coverage histogram plot.

#### Methods

signature(object = "ChIPQCexperiment") Generate coverage histogram plots for all the samples in an experiment.

signature(object = "list") Generate coverage histogram plots for all the samples in a list of ChIPQCsamples.

signature(object = "ChIPQCsample") Generate coverage histogram plots for a sample.

#### Note

Uses ggplot2 for plotting, and returns a ggplot2 plot dataframe.

### Author(s)

Thomas Carroll and Rory Stark

### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

### Examples

```
data(example_QCexperiment)
plotCoverageHist(exampleExp,facetBy=c("Tissue","Factor"))
plotCoverageHist(QCsample(exampleExp,1))
```

plotFribl-methods

Generate fraction of reads in blacklist plot

### Description

Generate fraction of reads in blacklist plot.

### Methods

```
signature(object = "ChIPQCexperiment", type="barstacked", facet=T, facetBy=c("Tissue", "Factor"), AsF Generate fraction of reads in blacklist plots for all the samples in an experiment.
```

```
signature(object = "list", type="barstacked",facet=T, facetBy=c("Sample"),AsPercent=TRUE) Generate fraction of reads in blacklist plots for all the samples in a list of ChIPQCsample objects..
```

signature(object = "ChIPQCsample", type="barstacked", AsPercent=TRUE) Generate fraction of reads in blacklist plots for a sample.

24 plotFrip-methods

#### Note

plotFribl uses ggplot2 for plotting, and returns a ggplot2 plot dataframe.

#### Author(s)

Thomas Carroll and Rory Stark

#### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

### **Examples**

```
data(example_QCexperiment)
plotFribl(exampleExp)
plotFribl(QCsample(exampleExp,1))
```

plotFrip-methods

Generate fraction of reads in peaks plot

### **Description**

Generate fraction of reads in peaks plot.

#### Methods

```
signature(object = "ChIPQCexperiment", type="barstacked", facet=T, facetBy=c("Tissue", "Factor"), AsF
Generate fraction of reads in peaks plots for all the samples in an experiment.
```

```
signature(object = "list", type="barstacked",facet=T, facetBy=c("Sample"),AsPercent=TRUE)
Generate fraction of reads in peaks plots for all the samples in a list of ChIPQCsample objects.
```

signature(object = "ChIPQCsample", type="barstacked", facet=T, facetBy=c("Tissue", "Factor"), AsPerce
Generate fraction of reads in peaks plots for a sample.

### Note

plotFrip usesggplot2 for plotting, and returns a ggplot2 plot dataframe.

### Author(s)

Thomas Carroll and Rory Stark

### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

```
data(example_QCexperiment)
plotFrip(exampleExp)
plotFrip(QCsample(exampleExp,1))
```

```
plotPeakProfile-methods
```

Generate peak profile plots

### Description

Generate peak profile plots.

### Methods

```
signature(object = "ChIPQCexperiment", method) Generate peak profile plots for all the samples in an experiment.
```

signature(object = "list", method) Generate peak profile plots for all the samples in a list of ChIPQCsample objects..

signature(object = "ChIPQCsample", method) Generate peak profile plots for a sample.

### Note

plotPeakProfile uses ggplot2 for plotting, and returns a ggplot2 plot dataframe.

### Author(s)

Thomas Carroll and Rory Stark

### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

### **Examples**

```
data(example_QCexperiment)
plotCoverageHist(exampleExp,facetBy=c("Tissue","Factor"))
plotCoverageHist(QCsample(exampleExp,1))
data(tamoxifen_QC)
plotCoverageHist(tamoxifen,facetBy=c("Tissue","Condition"))
```

plotPrincomp-methods Generate Princi

Generate Principal Components Analysis plot for ChIP samples

### **Description**

Generate principal components analysis plot for ChIP samples.

26 plotRap-methods

#### Methods

```
signature(object = "ChIPQCexperiment", attributes, ...) Generate principal components analysis plot, for all the samples in an experiment that are associated with a peakset.
```

attributes character string, or vector of character strings, representing metadata field names, for use grouping samples by additional parameters passed to dba.plotPCA

### Note

plotPrincomp uses dba.plotPCA for plotting.

#### Author(s)

Rory Stark and Thomas Carroll

#### See Also

ChIPQC-package, ChIPQCexperiment, dba.plotPCA

### **Examples**

```
data(tamoxifen_QC)
plotPrincomp(tamoxifen,attributes=c("Condition"))
```

plotRap-methods

Generate reads in peaks plot

### Description

Generate reads in peaks plot.

### Methods

```
signature(object = "ChIPQCexperiment", type="barstacked", facet=T, facetBy=c("Tissue", "Factor"))
    Generate reads in peaks plots for all the samples in an experiment.
signature(object = "list", type="barstacked", facet=T, facetBy=c("Sample")) Generate
    reads in peaks plots for all the samples in a list of ChIPQCsample objects.
signature(object = "ChIPQCexperiment", type="barstacked", facet=T, facetBy=c("Tissue", "Factor"))
    Generate reads in peaks plots for all the samples in an experiment.
signature(object = "ChIPQCsample", type="barstacked", facet=T, facetBy=c("Tissue", "Factor"))
    Generate reads in peaks plots for a sample.
```

### Note

plotRap uses ggplot2 for plotting, and returns a ggplot2 plot dataframe.

### Author(s)

Thomas Carroll and Rory Stark

plotRegi-methods 27

### See Also

```
ChIPQC-package, ChIPQCexperiment, ChIPQCsample
```

### **Examples**

```
data(example_QCexperiment)
plotRap(exampleExp, facetBy=c("Tissue", "Factor"))
plotRap(QCsample(exampleExp,1))
```

plotRegi-methods

Generate relative enrichment of genomic features plot

### **Description**

Generate relative enrichment of genomic features plot.

#### Methods

```
signature(object = "ChIPQCexperiment", facet=T, facetBy=c("Tissue", "Factor")) Generate
relative enrichment of genomic features plots for all the samples in an experiment.
```

signature(object = "list", facet=T, facetBy=c("Sample")) Generate relative enrichment of
genomic features plots for all the samples in a list of ChIPQCsample objects.

signature(object = "ChIPQCsample") Generate relative enrichment of genomic features plots
for a sample.

### Note

plotRegi uses ggplot2 for plotting, and returns a ggplot2 plot dataframe.

#### Author(s)

Thomas Carroll and Rory Stark

### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

```
data(example_QCexperiment)
plotRegi(exampleExp,facetBy=c("Tissue","Factor"))
plotRegi(QCsample(exampleExp,1))
data(tamoxifen_QC)
plotRegi(tamoxifen,facetBy=c("Tissue","Condition"))
```

28 QCannotation-methods

plotSSD-methods

Generate SSD plot

### **Description**

Generate SSD metric plot. If blacklists supplied, will generate SSD prior and post blacklisting

### Methods

```
signature(object = "ChIPQCexperiment", facet=T, facetBy=c("Tissue", "Factor")) Generate SSD metric plot for all samples in experiment. If blacklists supplied, will generate SSD prior and post blacklisting
```

signature(object = "list", facet=T, facetBy=c("Tissue", "Factor")) Generate SSD metric plot for list of samples in experiment. If blacklists supplied, will generate SSD prior and post blacklisting

signature(object = "ChIPQCsample") Generate SSD metric plot for single sample. If blacklists supplied, will generate SSD prior and post blacklisting

### Note

plotSSD uses ggplot2 for plotting, and returns a ggplot2 plot gg object.

### Author(s)

Thomas Carroll and Rory Stark

### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

### **Examples**

```
data(example_QCexperiment)
plotSSD(exampleExp, facetBy=c("Tissue", "Factor"))
plotSSD(QCsample(exampleExp,1))
plotSSD(QCsample(exampleExp)[1:3])
data(tamoxifen_QC)
plotSSD(tamoxifen, facetBy=c("Tissue", "Condition"))
```

QCannotation-methods

Retrieve an annotation description, or a processed annotation

### Description

Retrieve an annotation description, or a processed annotation, from a ChIPQCexperiment object.

QCcontrol-methods 29

#### Methods

signature(object = "ChIPQCexperiment", bRetrieve=FALSE) Retrieve the annotation. If bRetrieve=FALSE (default), the character string describing the annotation is returned (currently only "hg19" is supported). If bRetrieve=TRUE, a processed annotation is returned (in the form of a list). This can be used in subsequent calls to ChIPQC and/or ChIPQCsample for efficiency purposes.

### Author(s)

Rory Stark and Thomas Carroll

### See Also

ChIPQC-package, ChIPQCsample, ChIPQCexperiment

### **Examples**

```
data(example_QCexperiment)
QCannotation(exampleExp)
```

QCcontrol-methods

Retrieve control objects associated with a sample

### **Description**

Retrieve the ChIPQCsample objects representing controls, or a specific ChIPQCsample representing the control for a specific sample, from a ChIPQCexperiment object.

### **Methods:**

signature(object = "ChIPQCexperiment", sampleID) Get the control sample (as a ChIPQCsample object) associated with a ChIP sample, or, if sampleID is missing, a list of all samples used as controls. sampleID is a character string or an integer.

### Author(s)

Rory Stark and Thomas Carroll

### See Also

ChIPQC-package, ChIPQCsample, ChIPQCexperiment

```
data(tamoxifen_QC)
controls = QCcontrol(tamoxifen)
length(controls)
names(controls)
controls[[1]]
bt474control = QCcontrol(tamoxifen,"BT4741")
bt474control
```

30 QCmetadata-methods

QCdba-methods

Retrieve the DBA-object object associated with an experiment

### **Description**

Retrieve the DBA-object object associated with a ChIPQCexperiment object.

#### **Methods:**

signature(object = "ChIPQCexperiment") Retrieves the DBA-object object associated with a ChIPQCexperiment. This object can be used with DiffBind functions to further analyse a ChIP-seq experiment, including performing a differential binding analysis.

### Author(s)

Rory Stark and Thomas Carroll

#### See Also

ChIPQC-package, ChIPQCexperiment, DiffBind, dba

### **Examples**

```
data(tamoxifen_QC)
tamoxifenDBA = QCdba(tamoxifen)
## library(DiffBind)
## tamoxifenDBA
```

QCmetadata-methods

Retrieve metadata associated with an experiment

### **Description**

Retrieve metadata for a ChIPQCexperiment object.

### **Methods:**

signature(object = "ChIPQCexperiment") Retrieve a data frame containing metadata for all the samples in a ChIP-seq experiment represented by a ChIPQCexperiment object.

signature(object = "list") Retrieve a data frame containing metadata for all the samples in a list of ChIPQCsample objects.

### Author(s)

Rory Stark and Thomas Carroll

### See Also

ChIPQC-package, ChIPQCsample, ChIPQCexperiment

QCmetrics-methods 31

### **Examples**

```
data(tamoxifen_QC)
meta = QCmetadata(tamoxifen)
meta
```

QCmetrics-methods

Retrieve consolidated set of QC metrics

### **Description**

Retrieves a consolidated set of QC metrics.

### Methods

```
signature(object = "ChIPQCexperiment") Retrieves a matrix of QC metrics for all the samples in an experiment, with a column of values for each sample.
```

signature(object = "list") Retrieves a matrix of QC metrics for all the samples in a list of ChIPQCsample objects, with a column of values for each sample.

signature(object = "ChIPQCsample") Retrieves a vector of QC metrics for a sample.

### Author(s)

Rory Stark and Thomas Carroll

### See Also

ChIPQC-package, ChIPQCsample, ChIPQCexperiment

### **Examples**

```
data(example_QCexperiment)
QCmetrics(exampleExp)

data(tamoxifen_QC)
QCmetrics(tamoxifen)
QCmetrics(QCsample(tamoxifen,1))
```

QCsample-methods

Retrieve sample objects associated with an experiment

### Description

Retrieves a list of ChIPQCsample objects, or one specific ChIPQCsample object, from a ChIPQCexperiment object.

32 readlength-methods

#### Methods

signature(object = "ChIPQCexperiment", sampleID) If sampleID is missing or equal to 0, the full list of ChIPQCsample objects is returned. If sampleID is an integer n, the ChIPQCsample object corresponding to the nth sample is returned. If sampleID is a character string, the ChIPQCsample object corresponding to the sample with that ID is returned.

#### Author(s)

Rory Stark and Thomas Carroll

#### See Also

ChIPQC-package, ChIPQCsample, ChIPQCexperiment

### **Examples**

```
data(example_QCexperiment)
samples = QCsample(exampleExp)
length(samples)
names(samples)
samples$CTCF_1
```

readlength-methods

Retrieve read length values

### **Description**

Retrieve read length values.

### Methods

signature(object = "ChIPQCexperiment") Retrieve a vector of read length values, one for
each sample in an experiment.

signature(object = "list") Retrieve a vector of read length values, one for each sample in a
list of ChIPQCsample objects.

signature(object = "ChIPQCsample") Retrieve the read length value for a sample.

### Author(s)

Thomas Carroll and Rory Stark

### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

```
data(example_QCexperiment)
readlength(exampleExp)
readlength(QCsample(exampleExp,1))
```

ReadLength Fragment Length Cross Coverage-methods

Retrieve the cross coverage values when extending reads to the optimal fragment length.

### **Description**

Retrieve the cross coverage values when extending reads to the optimal fragment length.

#### **Methods**

- signature(object = "ChIPQCexperiment") Retrieve a vector of cross-coverage values for all samples in a ChIP-seq experiment, when all reads are shifted by the optimal fragment length (the maximum cross-coverage value).
- signature(object = "list") Retrieve a vector of cross-coverage values for all samples in a list of ChIPQCsample objects, when all reads are shifted by the optimal fragment length (the maximum cross-coverage value).
- signature(object = "ChIPQCsample") Retrieve the cross-coverage value for a sample, when all reads are shifted by the optimal fragment length (the maximum cross-coverage value).

### Author(s)

Thomas Carroll and Rory Stark

#### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

### **Examples**

```
data(example_QCexperiment)
FragmentLengthCrossCoverage(exampleExp)
FragmentLengthCrossCoverage(QCsample(exampleExp,1))
```

 ${\tt ReadLengthReadLengthCrossCoverage-methods}$ 

 $Retrieve\ the\ cross\ coverage\ values\ without\ extending\ reads$ 

### Description

Retrieve the cross coverage values without extending reads.

### Methods

- signature(object = "ChIPQCexperiment") Retrieve a vector of cross-coverage values for all samples in a ChIP-seq experiment, with no shift.
- signature(object = "list") Retrieve a vector of cross-coverage values for all samples in a list of ChIPQCsample objects, with no shift.
- signature(object = "ChIPQCsample") Retrieve the cross-coverage value for a sample, with no shift.

34 reads-methods

#### Author(s)

Thomas Carroll and Rory Stark

#### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

### **Examples**

```
data(example_QCexperiment)
ReadLengthCrossCoverage(exampleExp)
ReadLengthCrossCoverage(QCsample(exampleExp,1))
```

reads-methods

Retrieve numbers of reads

### **Description**

Retrieve the numbers of reads.

#### Methods

- signature(object = "ChIPQCexperiment", bFiltered) Retrieve a vector of the numbers of reads for each sample in an experiment. If bFiltered=TRUE (or is missing), this will be the number of reads that pass the mapping quality filter for each sample. if bFiltered=FALSE, it will be the total number of reads for each sample.
- signature(object = "list", bFiltered) Retrieve a vector of the numbers of reads for each sample in a list of ChIPQCsample objects. If bFiltered=TRUE (or is missing), this will be the number of reads that pass the mapping quality filter for each sample. if bFiltered=FALSE, it will be the total number of reads for each sample.
- signature(object = "ChIPQCsample", bFiltered) Retrieve the number of reads for a sample. If bFiltered=TRUE (or is missing), this will be the number of reads that pass the mapping quality filter. if bFiltered=FALSE, it will be the total number of reads for the sample.

### Author(s)

Thomas Carroll and Rory Stark

### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

```
data(example_QCexperiment)
reads(exampleExp)
reads(QCsample(exampleExp,1))
```

regi-methods 35

|--|

### **Description**

Retrieve genomic profile information in terms of relative enrichment over background genomic distribution.

#### Methods

signature(object = "ChIPQCexperiment") Retrieve a matrix of relative enrichment values for a variety of genomic features. Each column represents the enrichment values for one sample in the experiment.

signature(object = "list") Retrieve a matrix of relative enrichment values for a variety of genomic features. Each column represents the enrichment values for one sample in a list of ChIPQCsample objects.

signature(object = "ChIPQCsample") Retrieve a vector of relative enrichment values for a variety of genomic features for a sample. Relative enrichment is computed as the proportion of reads overlapping a genomic feature type compared to the overall proportion of base pairs in the genome comprising those features. Genomic features include:

3 UTRs 5 UTRs 5 UTRs

Intronic (non-coding) portions of gene bodies

Transcripts Transcribed regions, including exons

Promoters 500 500 bp regions immediately upstream of annotated TSSs

Promoters2000to500 2500bp regions from 2000bp immediately upstream of annotated TSSs to 500bp downstream 2000bp regions from 20000bp immediately upstream of annotated TSSs to 2000bp downstream 2000bp downstream

#### Author(s)

Thomas Carroll and Rory Stark

### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

```
data(example_QCexperiment)
genomicprofile = regi(exampleExp)
heatmap(genomicprofile)
regi(QCsample(exampleExp,1))
```

36 ribl-methods

RelativeCrossCoverage-methods

Retrieve the relative cross coverage values for a range of shift sizes

### **Description**

Retrieve the relative cross-coverage values for a range of shift sizes

### Methods

- signature(object = "ChIPQCexperiment") Retrieve a vector of relative cross-coverage values for all samples in a ChIP-seq experiment, computed based on the maximal value (when extending the reads to the optimal fragment length) versus the cross-coverage values using non-extended reads.
- signature(object = "list") Retrieve a vector of relative cross-coverage values for all samples in a list of ChIPQCsample objects, computed based on the maximal value (when extending the reads to the optimal fragment length) versus the cross-coverage values using non-extended reads.
- signature(object = "ChIPQCsample") Retrieve the relative cross-coverage value for a sample, computed based on the maximal value (when extending the reads to the optimal fragment length) versus the cross-coverage values using non-extended reads.

#### Author(s)

Thomas Carroll and Rory Stark

#### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

### **Examples**

```
data(example_QCexperiment)
RelativeCrossCoverage(exampleExp)
RelativeCrossCoverage(QCsample(exampleExp,1))
```

ribl-methods

Retrieve numbers of reads overlapping blacklisted regions

### **Description**

Retrieve the numbers of reads overlapping blacklisted regions.

#### Methods

- signature(object = "ChIPQCexperiment") Retrieve a vector of the numbers of reads overlapping blacklisted regions for each sample in an experiment.
- signature(object = "list") Retrieve a vector of the numbers of reads overlapping blacklisted regions for each sample in a list of ChIPQCsample objects.
- signature(object = "ChIPQCsample") Retrieve the number of reads overlapping blacklisted regions in a sample.

rip-methods 37

### Author(s)

Thomas Carroll and Rory Stark

### See Also

```
ChIPQC-package, ChIPQCexperiment, ChIPQCsample
```

### **Examples**

```
data(example_QCexperiment)
ribl(exampleExp)
ribl(QCsample(exampleExp,1))
```

rip-methods

Retrieve numbers of reads overlapping peaks

### **Description**

Retrieve the numbers of reads overlapping peaks.

### Methods

signature(object = "ChIPQCexperiment") Retrieve a vector of the numbers of reads overlapping peaks for each sample in an experiment.

signature(object = "list") Retrieve a vector of the numbers of reads overlapping peaks for each sample in a list of ChIPQCsample objects.

signature(object = "ChIPQCsample") Retrieve the number of reads overlapping peaks in a sample.

### Author(s)

Thomas Carroll and Rory Stark

### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

```
data(example_QCexperiment)
rip(exampleExp)
rip(QCsample(exampleExp,1))
```

38 ssd-methods

ssd-methods

Retrieve SSD (squared sum of deviations) values of peak coverage

### **Description**

Retrieve SSD (squared sum of deviations) values of peak coverage density.

### Methods

signature(object = "ChIPQCexperiment") Retrieve a vector of SSD values, one for each sample in an experiment.

signature(object = "list") Retrieve a vector of SSD values, one for each sample in a list of ChIPQCsample objects.

signature(object = "ChIPQCsample") Retrieve the SSD (squared sum of deviations) for a sample, computed from the standard deviation of the coveragehistogram.

### Note

uses the SSD calculation from the chipseq package.

### Author(s)

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### See Also

ChIPQC-package, ChIPQCexperiment, ChIPQCsample

```
data(example_QCexperiment)
ssd(exampleExp)
ssd(QCsample(exampleExp,1))
```

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