# **R** documentation

of all in 'man/'

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annoCuration, genomicCompartments-method Curate Gene Annotations

## **Description**

This method curates gene annotations by performing several filtering steps on the provided annotation data. It filters out non-canonical chromosomes, retains known coding and non-coding genes, keeps genes with a length greater than 2kb, removes duplicate annotations with the same coordinates, extends the transcription start site (TSS) by 1kb upstream, and removes overlapping annotations. The curated annotations are then set as the curated annotation data of the 'genomic-Compartments' object.

#### Usage

```
annoCuration(theobject, geneannovec, nbcpu = 8)
```

## **Arguments**

theobject An object of class 'genomicCompartments'.

geneannovec A character vector containing the paths to gene annotation files.

Number of CPU cores to use for parallel processing (default is 8).

#### Value

Returns the updated 'genomicCompartments' object with curated gene annotations.

## **Examples**

```
## Not run:
# Create a genomicCompartments object
gc_obj <- genomeCompart(peakspathvec, geneannovec)
# Perform annotation curation
gc_obj <- annoCuration(gc_obj, geneannovec)
## End(Not run)</pre>
```

bivalentPromoters, genomicCompartments-method

\*\*Identify Bivalent Promoters\*\*

## **Description**

This method identifies bivalent promoters by creating a merged GRanges object from peaks of H3K4me3 and H3K27me3 histone marks at TSS-/+1Kb.

#### Usage

bivalentPromoters(theobject, peakspathvec)

## **Arguments**

theobject An object of class 'genomicCompartments'.

peakspathvec A named vector specifying paths to the peak files of H3K4me3 and H3K27me3.

#### Value

Returns the modified 'genomicCompartments' object with bivalent promoters assigned.

## **Examples**

```
## Not run:
# Create a genomicCompartments object
gc_obj <- genomeCompart(peakspathvec, geneannovec)
# Identify bivalent promoters
bivalentPromoters(gc_obj, peakspathvec = c(H3K4me3 = "path/to/H3K4me3.gff",
H3K27me3 = "path/to/H3K27me3.gff"))
## End(Not run)</pre>
```

 ${\it boxplotGlcnacLevels, genomicCompartments-method} \\ {\it Generate Boxplot of Glcnac Levels}$ 

# Description

This method generates boxplots of Glcnac levels across the different genomic compartments stored in the given genomicCompartments object.

## Usage

## **Arguments**

the object An object of class 'genomic Compartments'.

bwpath Character string specifying the path to the BigWig file containing Glcnac levels.

outputfolder Character string specifying the output folder path for saving results.

includerepeats Logical indicating whether to include repeats in the analysis.

plotviolin Logical indicating whether to plot to current device (default is FALSE).

#### Value

Save the boxplot to outputfolder.

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## **Examples**

buildGR

Build GRanges Object from File

## **Description**

This function reads a file containing genomic coordinates and constructs a 'GRanges' object.

#### Usage

```
buildGR(currentpath)
```

## **Arguments**

currentpath

A character string representing the GFF file path to read. The file should contain tabular data with the following columns:

- V1 Chromosome names
- V4 Start positions
- V5 End positions
- V9 Feature names
- V7 Strand information

## Value

A 'GRanges' object with the specified genomic coordinates and feature names.

```
## Not run:
# Example file path
path <- "path/to/genomic/coordinates/file.gff"
# Build GRanges object
gr <- buildGR(path)
## End(Not run)</pre>
```

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buildGRNoReading

Build GRanges Object from Data Frame

## **Description**

This function constructs a 'GRanges' object from a data frame.

## Usage

```
buildGRNoReading(fi)
```

#### **Arguments**

fi

A data frame with columns representing genomic coordinates and other relevant information in GFF format. The expected columns are:

- V1 Chromosome names
- V4 Start positions
- V5 End positions
- V9 Feature names
- V7 Strand information

#### Value

A 'GRanges' object with the specified genomic coordinates and feature names.

## **Examples**

```
## Not run:
# Example data frame
df <- data.frame(V1 = "chr1", V4 = 1000, V5 = 2000, V9 = "gene1", V7 = "+")
# Build GRanges object
gr <- buildGRNoReading(df)
## End(Not run)</pre>
```

buildIntervalsObject

Build Genomic Intervals Object

# Description

This function constructs a 'genomeCompart' object, defines various genomic features, and optionally includes repeat annotations.

# Usage

```
buildIntervalsObject(peakspathvec, geneannos, includerepeats = FALSE,
  repeatsannovec = NA)
```

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#### **Arguments**

peakspathvec	A named vector of file paths to peak files in GFF format for various ChIP-seq and ATAC-seq data. Expected names include: "H3K27ac", "Ser5P", "Ser2P", "H3K4me3", "H3K27me3", "ATACSeq", "Suz12", "RING1B", "H3K9me3".
geneannos	A list of gene annotation files to be used for defining non-redundant gene sets and other features.
includerepeats	A logical value indicating whether to include repeat annotations (LINE, LTR, SINE). Default set to FALSE.
repeatsannovec	If include repeats is TRUE, a vector of paths to the LINE, LTR, and SINE coordinates. Default is $\ensuremath{NA}.$
enhancers	Logical indicating if active and poised enhancers should be computed. Default is FALSE.

#### **Details**

This function performs the following steps:

- Builds the initial 'genomeCompart' object.
- Defines a non-redundant set of gencode genes by removing overlapping genes, genes closer than 1kb to each other, and genes shorter than 2kb.
- Identifies active promoters using H3K27ac at TSS±1Kb.
- Identifies promoters with Ser5P (initiation) at TSS±1Kb.
- Identifies elongating genes by overlapping Ser2P from TSS+1kb to TES.
- Defines termination sites at TES+1kb.
- Identifies bivalent promoters using H3K4me3 and H3K27me3 at TSS±1kb.
- Defines active enhancers using H3K27ac, H3K4me1, and ATAC-seq, ensuring no overlap with gene annotations.
- Defines poised enhancers using H3K27me3, H3K4me1, and Suz12, ensuring no overlap with gene annotations.
- Defines polycomb domains using Suz12 and RING1B overlaps.
- Defines heterochromatin using H3K9me3.
- Optionally includes repeat annotations (SINE, LINE, LTR) if 'includerepeats' is TRUE.

#### Value

An updated 'genomeCompart' object with defined genomic features and, optionally, repeat annotations.

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## End(Not run)

checkParams

Validate Parameters given as input to different functions of the package

#### **Description**

This function validates various parameters required by several functions to ensure they meet the expected criteria.

#### Usage

#### **Arguments**

peakspathqueryvec

A vector of file paths to peak files in GFF format.

glcnacbwvec A vector of file paths to glc bigwig files.

querynamevec A vector of names corresponding to the peak files.

geneannovec A vector of file paths to gene annotation files, expected to be in the order of

gencode, refGene, and refseq.

peakspathcategoriesvec

A vector of names for each peak file, expected to contain strings identifying

categories such as H3K27ac, H3K4me1, etc.

repeatsannovec A vector of file paths to repeat annotation files in GFF format. The file names

are expected to contain the substrings LINE, LTR, and SINE.

countstable A vector containing the path to a count table file.

includerepeats A logical value indicating whether to include repeat annotations.

countsannotype A string specifying the annotation type for the counts table, either "ensembl" or

"entrez".

## **Details**

This function performs the following validation checks:

- Ensures there are no more than two files in peakspathqueryvec.
- Checks that the number of peak files matches the number of bigwig files.
- Verifies that the number of query names matches the number of peak files.
- Validates that gene annotation files are ordered as gencode, refGene, and refseq.
- Confirms that category peak files contain the expected strings for categories like H3K27ac, H3K4me1, etc.
- If repeat annotations are included, checks that the repeat files contain the expected substrings LINE, LTR, and SINE.
- Ensures only one counts table is provided.
- Validates that the counts table annotation type is either "ensembl" or "entrez".

#### Value

None. The function stops with an error message if any validation check fails.

## **Examples**

```
## Not run:
peakspathqueryvec <- c("path/to/peak1.gff", "path/to/peak2.gff")</pre>
glcnacbwvec <- c("path/to/bw1", "path/to/bw2")</pre>
querynamevec <- c("query1", "query2")</pre>
geneannovec <- c("gencode", "refGene", "refseq")</pre>
peak spath categories vec <- c("H3K27ac_path", "H3K4me1_path", "H3K27me3_path", "H3K27me3
 "H3K4me3_path", "Suz12_path", "RING1B_path", "H3K9me3_path", "Ser5P_path",
 "Ser2P_path", "ATAC_path")
repeatsannovec <- c("LINE_path", "LTR_path", "SINE_path")</pre>
countstable <- c("path/to/counts_table")</pre>
includerepeats <- TRUE
countsannotype <- "ensembl"</pre>
checkParams(peakspathqueryvec, glcnacbwvec, querynamevec, geneannovec,
peakspathcategoriesvec, repeatsannovec, countstable, includerepeats,
countsannotype)
## End(Not run)
```

complexUpsetDiagram, list-method

Generate Complex Upset Diagram

# Description

This method generates a plot showing the upset diagram of the overlaps of GlcNAc peaks with different genomic compartments and the corresponding average binding values for each peak.

#### Usage

```
complexUpsetDiagram(theobject, includerepeats, outfold)
```

# Arguments

plotglclevels

the object A list of objects of class 'genomic Compartments'.

outfold Character string specifying the output folder path for saving the upset diagram

and boxplot.

includerepeats Logical indicating whether to include repeats in the analysis. Default is FALSE. includeenhancers

Logical indicating whether to include enhancers in the analysis. Default is

FALSE.

Logical indicating if the violin plot of the values of the glc peaks should be

plotted on top. Default is FALSE.

minsize Minimum number of overlap for a category to be indicated on the plot. Default

is 10.

#### Value

No return value. The function generates and saves a complex upset plot in the specified output folder.

## **Examples**

```
## Not run:
# Create a list of genomicCompartments objects
gc_list <- list(gc_obj1, gc_obj2, gc_obj3)
# Generate complex upset diagram
complexUpsetDiagram(gc_list, includerepeats = FALSE,
  outfold = "path/to/output_folder")
## End(Not run)</pre>
```

```
enhancer, {\tt genomicCompartments-method} \\ {\it Identify\ Enhancers}
```

## **Description**

This method identifies active or poisedd enhancers based on the overlap of histone marks and ATAC-seq peaks.

## Usage

```
enhancer(theobject, peakspath1, peakspath2, peakspath3, label1, label2,
label3, statelabel)
```

## **Arguments**

theobject	An object of class 'genomicCompartments'.
peakspath1	Path to the peaks file of the first histone mark.
peakspath2	Path to the peaks file of the second histone mark.
peakspath3	Path to the peaks file of the third histone mark or ATAC-seq.
label1	Label for the first histone mark.
label2	Label for the second histone mark.
label3	Label for the third histone mark or ATAC-seq.
statelabel	Label indicating the state of enhancers ('active' or 'poised').

## **Details**

Active enhancers are defined as the overlap of H3K27ac/H3K4me1/ATAC-seq. Poised enhancers are defined as the overlap of H3K27me3/H3K4me1/PRC2. They should not overlap with the combination of UCSC refGene, NCBI RefSeq, and GENCODE VM25.

#### Value

Returns the modified 'genomicCompartments' object with enhancers assigned.

#### **Examples**

```
## Not run:
# Create a genomicCompartments object
gc_obj <- genomeCompart(peakspathvec, geneannovec)
# Identify active enhancers
enhancer(gc_obj, peakspath1 = "path/to/H3K27ac.gff",
    peakspath2 = "path/to/H3K4me1.gff",
    peakspath3 = "path/to/ATAC-seq.gff",
label1 = "H3K27ac", label2 = "H3K4me1", label3 = "ATAC-seq",
statelabel = "active")
## End(Not run)</pre>
```

extractCompCoordWithPeak,genomicCompartments-method

Extract Compartment Coordinates with Peaks

## **Description**

This method performs overlap analysis between reference peaks and each compartment defined in the 'genomicCompartments' object. It extracts compartments containing peaks and outputs their coordinates in GFF format.

# Usage

```
extractCompCoordWithPeak(theobject, outfold, includerep)
```

#### **Arguments**

theobject An object of class 'genomicCompartments'.

outfold Path to the folder where output files will be written.

includerep Logical indicating whether to include repeat regions in the analysis. Default is

FALSE.

includeenhancers

Logical indicating whether to include enhancers in the analysis. Default is

FALSE.

## Value

Returns the modified 'genomicCompartments' object with compartments containing peaks.

```
## Not run:
# Create a genomicCompartments object
gc_obj <- genomeCompart(peakspathvec, geneannovec)
# Extract compartment coordinates with peaks
extractCompCoordWithPeak(gc_obj, outfold = "output_folder")
## End(Not run)</pre>
```

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filterChromAndStrand Filter Chromosomes and Strand Information in GRanges

## **Description**

This function filters out unwanted chromosomes and standardizes strand information in a 'GRanges' object.

## Usage

filterChromAndStrand(currentgr)

## **Arguments**

currentgr

A 'GRanges' object containing genomic ranges to be filtered.

#### **Details**

This function performs the following operations:

- If the strand information contains neither "+" and "-" strands, all strand information is set to "+".
- Chromosomes with names containing "Un", "random", or "chrM" are removed.
- The sequence levels of the resulting 'GRanges' object are updated to include only the remaining chromosomes.

## Value

A filtered 'GRanges' object with unwanted chromosomes removed and strand information standardized.

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genomeCompart

Create Genomic Compartments Object

#### **Description**

This function is the constructor of the genomicCompartments object. It is called by the function buildIntervalsObject.

## Usage

```
genomeCompart(peakspathvec, geneannovec)
```

#### **Arguments**

peakspathvec Character vector containing the path to the peaks file.

geneannovec Character vector containing the paths to gene annotation files.

#### **Details**

peakspathvec is made of the GFF file containing glc peaks and of the GFF files containing the ChIP-seq and ATAC-seq peak coordinates. The geneannovec contains path to the gencode, refgene, and refseq gff files.

#### Value

Returns an object of class 'genomicCompartments' containing the processed peaks and gene annotations.

#### See Also

buildIntervalsObject

genomicCompartments-class

Genomic Compartments Class

#### **Description**

The 'genomicCompartments' class is designed to store various genomic data, including reference peaks, gene annotations, and different types of genomic regions such as promoters, enhancers, and chromatin domains.

#### Slots

referencePeaks A 'GRanges' object containing reference peaks (required).

geneAnnotations A 'GRangesList' object containing gene annotations (required).

curatedAnno A 'GRanges' object for curated annotations.

activeProm A 'GRanges' object for active promoters defined as K27ac in TSS-/+1Kb.

activeEnh A 'GRanges' object for active enhancers defined as H3K27ac/H3K4me1/ATAC-seq. They should not overlap with the combination of UCSC refGene, NCBI RefSeq, and GEN-CODE VM25.

poisedEnh A 'GRanges' object for poised enhancers defined as H3K27me3/H3K4me1/PRC2. They should not overlap with the combination of UCSC refGene, NCBI RefSeq, and GENCODE VM25. Note: use a non redundant set of genes i.e. no overlap, remove genes closer to 1kb from one another, remove genes that are too short i.e. length < 2kb.

PcGDomain A 'GRanges' object for PcG domains defined as Suz12 and RING1B overlap.

heteroChrom A 'GRanges' object for heterochromatin regions (H3K9me3)

bivalentProm A 'GRanges' object for bivalent promoters defined as the overlap of H3K4me3/H3K27me3.

initiation A 'GRanges' object for initiation regions defined as TSS-1/+1kb overlapping with Ser5P peaks.

elongation A 'GRanges' object for elongation regions defined as TSS+1kb to TES overlapping Ser2P.

termination A 'GRanges' object for termination regions defined as TES+1kb

SINE A 'GRanges' object for SINE elements.

LINE A 'GRanges' object for LINE elements.

LTR A 'GRanges' object for LTR elements.

regionsMatrix A matrix representing various regions.

glcPeakVals A list of glc peaks.

 ${\tt activePromWithPeaks}\ A\ {\tt `GRanges'\ object\ for\ active\ promoters\ with\ glc\ peaks}.$ 

activeEnhWithPeaks A 'GRanges' object for active enhancers with glc peaks.

 ${\tt poisedEnhWithPeaks}\ A\ `GRanges'\ object\ for\ poised\ enhancers\ with\ glc\ peaks.$ 

PcGDomainWithPeaks A 'GRanges' object for PcG domains with glc peaks.

heteroChromWithPeaks A 'GRanges' object for heterochromatin regions with glc peaks.

bivalentPromWithPeaks A 'GRanges' object for bivalent promoters with glc peaks.

initiationWithPeaks A 'GRanges' object for initiation regions with glc peaks.

elongationWithPeaks A 'GRanges' object for elongation regions with glc peaks.

terminationWithPeaks A 'GRanges' object for termination regions with glc peaks.

#### Constructor

new("genomicCompartments", referencePeaks = grglc, geneAnnotations = grlist, ...)

#### Accessors

```
In the following snippets, x is a genomicCompartments object.
getRefPeaks(x): Get the reference glc peaks
getGenesFullAnno(x): Get the full gene annotations
getCuratedAnno(x): Get the curated annotations
getActiveProm(x): Get the active promoters
getActiveEnh(x): Get the active enhancers
getPoisedEnh(x): Get the poised enhancers
getPcGDomain(x): Get the PcG domains
getHeteroChrom(x): Get the heterochromatin regions
getBivalentProm(x): Get the bivalent promoters
getInitiation(x): Get the initiation regions
getElongation(x): Get the elongation regions
getTermination(x): Get the termination regions
getSINE(x): Get the SINE regions
getLINE(x): Get the LINE regions
getLTR(x): Get the LTR regions
getRegionsMatrix(x): Get the regions matrix
getGlcPeakVal(x): Get the glc peak values
getActivePromWithPeaks(x): Get the active promoters with glc peaks
getActiveEnhWithPeaks(x): Get the active enhancers with glc peaks
getPoisedEnhWithPeaks(x): Get the poised enhancers with glc peaks
getPcGDomainWithPeaks(x): Get the PcG domains with glc peaks
getHeteroChromWithPeaks(x): Get the heterochromatin regions with glc peaks
getBivalentPromWithPeaks(x): Get the bivalent promoters with glc peaks
getInitiationWithPeaks(x): Get the initiation regions with glc peaks
getElongationWithPeaks(x): Get the elongation regions with glc peaks
getTerminationWithPeaks(x): Get the termination regions with glc peaks
```

#### **Subsetting**

```
In the following snippets, x is a genomicCompartments object.
setCurratedAnno(x, value): Set the curated annotations
setActivePromoters(x, value): Set the active promoters
setInitiationSer5PProm(x, value): Set the initiation Ser5P promoters
setElongationSer2P(x, value): Set the elongation Ser2P regions
setTermination(x, value): Set the termination regions
setBivalentProm(x, value): Set the bivalent promoters
setActiveEnhancer(x, value): Set the active enhancers
setPoisedEnhancer(x, value): Set the poised enhancers
setPolycombsDomain(x, value): Set the PcG domains
setHeterochromatin(x, value): Set the heterochromatin regions
setSINE(x, value): Set the SINE regions
setLINE(x, value): Set the LINE regions
setLTR(x, value): Set the LTR regions
setRegionsMatrix(x, value): Set the regions matrix
setGlcPeakVal(x, value): Set the glc peak values
```

setActivePromWithPeaks(x, value): Set the active promoters with glc peaks setActiveEnhWithPeaks(x, value): Set the active enhancers with glc peaks setPoisedEnhWithPeaks(x, value): Set the poised enhancers with glc peaks setPcGDomainWithPeaks(x, value): Set the PcG domains with glc peaks setHeteroChromWithPeaks(x, value): Set the heterochromatin regions with glc peaks setBivalentPromWithPeaks(x, value): Set the bivalent promoters with glc peaks setInitiationWithPeaks(x, value): Set the initiation regions with glc peaks setElongationWithPeaks(x, value): Set the elongation regions with glc peaks setTerminationWithPeaks(x, value): Set the termination regions with glc peaks

## **Description**

This method creates a matrix for generating an UpSet plot, indicating overlaps between glc peaks and genomic compartments.

## Usage

```
matrixForUpset(theobject, includerepeats, glcpeakvalues)
```

#### **Arguments**

```
theobject An object of class 'genomicCompartments'.

includerepeats Logical indicating whether to include repeats in the analysis.

includeenhancers

Logical indicating whether to include enhancers in the analysis.

glcpeakvalues Numeric vector of glc peak values.
```

#### Value

Returns the modified 'genomicCompartments' object with the regions matrix assigned.

```
## Not run:
# Create a genomicCompartments object
gc_obj <- genomeCompart(peakspathvec, geneannovec)
# Create matrix for UpSet plot
matrixForUpset(gc_obj, includerepeats = FALSE, glcpeakvalues = c(1, 2, 3))
## End(Not run)</pre>
```

outputGlcPeaksCoordPerCompartment,genomicCompartments-method

Output Glc Peaks Coordinates per Compartment

## **Description**

This method calculates the overlap of glc peaks with each compartment defined in the 'genomic-Compartments' object.

## Usage

#### **Arguments**

theobject An object of class 'genomicCompartments'.

outputfolder Path to the folder where output files will be written.

glcpeakspath Path to the GFF file containing glc peaks coordinates.

includerepeats Logical indicating whether to include repeat regions in the analysis.

includeenhancers

Logical indicating whether to include enhancers in the analysis.

#### Value

Returns nothing explicitly; Write GFF and BED files of the coordinates of the glc peaks overlapping with a specific compartment.

```
## Not run:
# Parameters
peakspathquery <- "/path/to/glc.gff"</pre>
peakspathcategoriesvec <- c(H3K27ac = "/path/to/H3K27ac.gff",</pre>
       H3K4me1 = "/path/to/H3K4me1.gff", H3K27me3 = "/path/to/H3K27me3.gff",
       H3K4me3 = "/path/to/H3K4me3.gff", Suz12 = NA, RING1B = NA,
       H3K9me3 = "/path/to/H3K9me3.gff", Ser5P = "/path/to/Ser5P.gff",
       Ser2P = "/path/to/Ser2P.gff", ATACSeq = "/path/to/ATACseq.gff")
peakspathvec <- c(peakspathquery, peakspathcategoriesvec)</pre>
geneannovec <- c(gencode = "/path/to/gencode.gff",</pre>
       refgene = "/path/to/refGeneUCSC.gff",
       refseq = "/path/to/refseq.gff")
# Create a genomicCompartments object
gc_obj <- genomeCompart(peakspathvec, geneannovec)</pre>
# Output glc peaks coordinates per compartment
outputGlcPeaksCoordPerCompartment(gc_obj, outputfolder, peakspathquery,
includerepeats = FALSE, includeenhancers = TRUE)
## End(Not run)
```

```
overlapOnGenes, genomicCompartments-method 
Overlap Peaks on Genes
```

## **Description**

This method performs overlap analysis between peaks and curated annotations at specified genomic regions (TSS or TES). It assigns overlapping peaks to specific compartments based on the provided labels.

## Usage

```
overlapOnGenes(theobject, regionlabel, peaklabel = NA, peakpath = NA)
```

## **Arguments**

theobject An object of class 'genomicCompartments'.

regionlabel Character specifying the genomic region label ("TSS", "GBTES", or "TES").

peaklabel Character specifying the peak label ("H3K27ac", "Ser5P", or "Ser2P").

Path to the file containing peaks data. NA if regionlabel is "TES".

## Value

Returns the modified 'genomicCompartments' object with assigned compartments.

#### **Examples**

```
## Not run:
# Create a genomicCompartments object
gc_obj <- genomeCompart(peakspathvec, geneannovec)
# Perform overlap analysis on TSS with H3K27ac peaks
overlapOnGenes(gc_obj, regionlabel = "TSS", peaklabel = "H3K27ac",
peakpath = "path/to/peaks.gff")
## End(Not run)</pre>
```

```
polycombsDomains, genomicCompartments-method

*Identify Polycomb Domains*
```

# Description

This method identifies Polycomb domains based on the overlap of two marks.

# Usage

```
polycombsDomains(theobject, peakspath1, peakspath2, label1, label2,
domainname)
```

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## **Arguments**

theobject An object of class 'genomicCompartments'.

peakspath1 Path to the peaks file of the first polycomb.

peakspath2 Path to the peaks file of the second polycomb.

label1 Label for the first polycomb.label2 Label for the second polycomb.

domainname Name or label for the identified Polycomb domains.

#### Value

Returns the modified 'genomicCompartments' object with Polycomb domains assigned.

#### **Examples**

```
## Not run:
# Create a genomicCompartments object
gc_obj <- genomeCompart(peakspathvec, geneannovec)
# Identify Polycomb domains
polycombsDomains(gc_obj, peakspath1 = "path/to/SUZ12.gff",
    peakspath2 = "path/to/RING1B.gff",
    label1 = "SUZ12", label2 = "RING1B", domainname = "PolycombDomains")
## End(Not run)</pre>
```

retrieveconversiontab Retrieve Gene Conversion Table

#### **Description**

This function retrieves a conversion table of gene identifiers from Ensembl BioMart.

## Usage

```
retrieveconversiontab(biomartstr, datasetstr, hoststr, alternativemirroropt)
```

# **Arguments**

biomartstr A string specifying the BioMart database to use (e.g., "ENSEMBL\_MART\_ENSEMBL").

datasetstr A string specifying the dataset to use within the BioMart database (e.g., "hsapi-

ens\_gene\_ensembl").

hoststr A string specifying the host URL for BioMart (e.g., "http://www.ensembl.org").

 $alternative \verb|mirroropt|$ 

A string specifying an alternative mirror option for BioMart, if needed.

## Details

The function filters the retrieved data to include only rows with non-empty and non-NA gene IDs.

#### Value

A data frame containing the conversion table with columns for RefSeq, Entrez, and Ensembl gene identifiers.

# **Examples**

```
## Not run:
biomartstr <- "ENSEMBL_MART_ENSEMBL"</pre>
datasetstr <- "hsapiens_gene_ensembl"</pre>
hoststr <- "http://www.ensembl.org"</pre>
alternativemirroropt <- "useast"</pre>
geneconvert <- retrieveconversiontab(biomartstr, datasetstr, hoststr,</pre>
alternativemirroropt)
head(geneconvert)
## End(Not run)
```

```
retrieveGlcPeakVal,genomicCompartments-method
                        Retrieve Glc Peak Values
```

#### **Description**

This method retrieves glc peak values associated with genomic compartments.

#### Usage

```
retrieveGlcPeakVal(theobject, includerepeats, bwpath)
```

## **Arguments**

theobject An object of class 'genomicCompartments'.

Character string specifying the path to the bigWig file containing glc peak valbwpath

includerepeats Logical indicating whether to include repeats in the analysis. Default is FALSE.

includeenhancers

Logical indicating whether to include enhancers in the analysis. Default is FALSE.

## Value

Returns the modified 'genomicCompartments' object with glc peak values assigned.

```
## Not run:
# Create a genomicCompartments object
gc_obj <- genomeCompart(peakspathvec, geneannovec)</pre>
# Retrieve glc peak values
gc_obj <- retrieveGlcPeakVal(gc_obj, includerepeats = FALSE,</pre>
bwpath = "path/to/bigWig/files")
```

```
## End(Not run)
```

```
{\it set Heterochromatin \verb|<---|, genomic Compartments-method} \\ Allocates\ heterochromatic\ region\ coordinates
```

# Description

This method stores the heterochromatin region coordinates in an object of class 'genomicCompartments'.

## Usage

```
setHeterochromatin(obj) <- value</pre>
```

#### **Arguments**

theobject An object of class 'genomicCompartments'.

value GRanges object of heterochromatic region coordinates.

#### Value

The updated object of class 'genomicCompartments' with the new heterochromatin regions.

# **Examples**

```
## Not run:
# Create a genomicCompartments object
gc_obj <- new("genomicCompartments")
# Define new heterochromatin regions
new_heterochromatin <- GRanges(seqnames = "chr1",
ranges = IRanges(start = 1, end = 1000000))
# Set the heterochromatin regions
setHeterochromatin(gc_obj) <- new_heterochromatin
## End(Not run)</pre>
```

```
setLINE<-,genomicCompartments-method

*Allocate LINE coordinates*
```

## **Description**

This method stores the LINE coordinates in an object of class 'genomicCompartments'.

#### Usage

```
setLINE(obj) <- value</pre>
```

## **Arguments**

the object An object of class 'genomic Compartments'.

value GRanges object of LINE region coordinates.

#### Value

The updated object of class 'genomicCompartments' with the new LINE regions.

## **Examples**

```
## Not run:
# Create a genomicCompartments object
gc_obj <- new("genomicCompartments")
# Define new LINE regions
new_LINE <- GRanges(seqnames = "chr1",
ranges = IRanges(start = 1, end = 1000000))
# Set the LINE regions
setLINE(gc_obj) <- new_LINE
## End(Not run)</pre>
```

```
setLTR<-,genomicCompartments-method

*Allocate LTR coordinates*
```

# Description

This method stores the heterochromatin region coordinates in an object of class 'genomicCompartments'.

## Usage

```
setLTR(obj) <- value</pre>
```

## **Arguments**

the object An object of class 'genomic Compartments'.

value GRanges object of LTR region coordinates.

# Value

The updated object of class 'genomicCompartments' with the new LTR regions.

```
## Not run:
# Create a genomicCompartments object
gc_obj <- new("genomicCompartments")
# Define new LTR regions
new_LTR <- GRanges(seqnames = "chr1",
ranges = IRanges(start = 1, end = 1000000))</pre>
```

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```
# Set the LTR regions
setLTR(gc_obj) <- new_LINE
## End(Not run)</pre>
```

```
setSINE<-, genomicCompartments-method

*Allocates SINE coordinates*
```

## **Description**

This method stores the SINE coordinates in an object of class 'genomicCompartments'.

## Usage

```
setSINE(obj) <- value</pre>
```

#### **Arguments**

the object An object of class 'genomic Compartments'.

value GRanges object of SINE region coordinates.

#### Value

The updated object of class 'genomicCompartments' with the new SINE regions.

## **Examples**

```
## Not run:
# Create a genomicCompartments object
gc_obj <- new("genomicCompartments")
# Define new SINE regions
new_SINE <- GRanges(seqnames = "chr1",
ranges = IRanges(start = 1, end = 1000000))
# Set the SINE regions
setSINE(gc_obj) <- new_SINE
## End(Not run)</pre>
```

tryGetBM

Attempt to Retrieve Information from Ensembl Using biomaRt

# Description

This function attempts to retrieve information about genes from Ensembl using the 'biomaRt' package. It tries up to 5 times before giving up and raising an error. The function supports specifying attributes, values, and filters for the query.

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#### Usage

```
tryGetBM(attributes, ensembl, values = NULL, filters = NULL)
```

#### **Arguments**

attributes Character vector specifying the attributes to retrieve.

ensembl 'Mart' object representing the connection to the Ensembl database.

values Optional. List or vector of values for the filter.

filters Optional. Character vector specifying the filters to apply.

#### Value

Returns a data frame with the retrieved information if the query is successful.

#### **Examples**

```
## Not run:
ensembl <- tryUseMart(biomart = "ensembl", dataset = "hsapiens_gene_ensembl",
host = "https://www.ensembl.org")
attributes <- c("ensembl_gene_id", "external_gene_name", "chromosome_name",
    "start_position", "end_position")
gene_info <- tryGetBM(attributes = attributes, ensembl = ensembl)
## End(Not run)</pre>
```

tryUseMart

Attempt to Connect to Ensembl Using biomaRt

#### **Description**

This function attempts to establish a connection to the Ensembl database using #' the 'biomaRt' package. It tries up to 5 times before giving up and raising an error. The function supports using an alternative mirror if specified.

## Usage

```
tryUseMart(biomart = "ensembl", dataset, host, alternativemirror)
```

#### **Arguments**

biomart Character string specifying the BioMart database to use. Default is "ensembl".

dataset Character string specifying the dataset to use within the BioMart database.

host Character string specifying the host URL to connect to.

alternativemirror

Logical value indicating whether to use an alternative mirror. Default is 'FALSE'.

#### Value

Returns a 'Mart' object if the connection is successful.

## **Examples**

```
## Not run:
ensembl <- tryUseMart(biomart = "ensembl", dataset = "hsapiens_gene_ensembl",
host = "https://www.ensembl.org")
## End(Not run)</pre>
```

```
\label{local_compartments} up set \textit{Diagram}, \textit{genomicCompartments-method} \\ \textit{Generate UpSet Diagram}
```

# Description

This method generates an UpSet diagram based on the overlap matrix of glc peaks and genomic compartments.

#### Usage

```
upsetDiagram(theobject, outputfolder, includerepeats, plotupset = FALSE)
```

## **Arguments**

theobject An object of class 'genomicCompartments'.

outputfolder Character string specifying the output folder where the UpSet diagram will be saved.

includerepeats Logical indicating whether to include repeats in the analysis.

plotupset Logical indicating whether to plot on device the upset plot ('TRUE') or not

('FALSE').

## Value

Does not return anything but saves the plot to outpufolder.

```
## Not run:
# Create a genomicCompartments object
gc_obj <- genomeCompart(peakspathvec, geneannovec)
# Generate UpSet diagram
upsetDiagram(gc_obj, outputfolder = "results/upset_plots",
includerepeats = FALSE, plotupset = TRUE)
## End(Not run)</pre>
```

 ${\it violinplotExpression, genomicCompartments-method} \\ {\it Generate\ Violin\ Plot\ for\ Gene\ Expression}$ 

# **Description**

This method generates violin plots for gene expression across genomic compartments.

# Usage

```
violinplotExpression(theobject, outfold, includerep, rnacounts, refseqpath,
  geneidtab, countsannotype, plotviolin = FALSE)
```

## **Arguments**

theobject	An object of class 'genomicCompartments'.
outfold	Character string specifying the output folder path for saving results.
includerep	Logical indicating whether to include repeats in the analysis.
rnacounts	Character string specifying the path to the RNA-seq count table.
refseqpath	Character string specifying the path to the original RefSeq annotation file.
geneidtab	Table containing genes ID.
countsannotype	Character string specifying the type of counts (e.g., entrez or not) used for expression values.
plotviolin	Logical indicating whether to plot on current device (default is FALSE).

# Value

Save the violin plot to outfold.

```
## Not run:
# Create a genomicCompartments object
gc_obj <- genomeCompart(peakspathvec, geneannovec)
# Generate violin plots for gene expression
violinplotExpression(gc_obj, outfold = "path/to/output_folder",
includerep = TRUE, rnacounts = "path/to/rna_counts",
refseqpath = "path/to/refseq_annotation",
geneidtab = geneidtab, countsannotype = "entrez", plotviolin = TRUE)
## End(Not run)</pre>
```

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