# Package 'coMET'

## November 5, 2014

Type Package

**Title** coMET: visualisation of regional epigenome-wide association scan (EWAS) results and DNA comethylation patterns.

**Version** 0.99.4

Date 2014-11-05

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**Description** Visualisation of EWAS results in a genomic region. In addition to phenotype-association P-values, coMET also generates plots of comethylation patterns and provides a series of annotation tracks.

**Depends** R (>= 3.1.0), grid, biomaRt, Gviz, rtracklayer, GenomicRanges

Suggests knitr, RUnit, BiocGenerics, BiocStyle

Imports colortools, hash, grDevices, gridExtra, ggbio, ggplot2, trackViewer

**License** GPL (>= 2)

URL http://epigen.kcl.ac.uk/comet

biocViews

Software, DifferentialMethylation, Visualization, Sequencing, Genetics, FunctionalGenomics

VignetteBuilder knitr

NeedsCompilation no

Repository Bioconductor

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## R topics documented:

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## Description

coMET is an R package for visualising EWAS results in a genomic region. Along with phenotype-association plots, coMET also generates plots of co-methylation patterns and provides a series of annotation tracks.

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#### **Details**

Package: coMET
Type: Package
Version: 0.99.4
Date: 2014-11-05
License: GPL (>=2)

coMET is an R package that can generate regional plots of EWAS results, DNA co-methylation patterns, and genomic information. A coMET figure includes 3 panels with a plot of P-values from EWAS, customized annotation tracks, and a triangle heatmap plot which demonstrates the correlation structure of DNA methylation at the CpG sites in the genomic region. Plots are created as PDF or EPS files.

#### Author(s)

Tiphaine C. Martin, Idil Yet, Pei-Chien Tsai, Jordana T. Bell Maintainer: Tiphaine Martin <tiphaine.martin@kcl.ac.uk>

Website: http://www.epigen.kcl.ac.uk/comet

#### References

Martin, T.C, Erte, I, Tsai, P-C, Bell, J.T., coMET: an R plotting package to visualize regional plots of epigenome-wide association scan results, QG14, 2014.

```
extdata <- system.file("extdata", package="coMET",mustWork=TRUE)</pre>
configfile <- file.path(extdata, "config_cyp1b1_zoom_4comet.txt")
myinfofile <- file.path(extdata, "cyp1b1_infofile.txt")</pre>
myexpressfile <- file.path(extdata, "cyp1b1_infofile_exprGene_region.txt")</pre>
mycorrelation <- file.path(extdata, "cyp1b1_res37_rawMatrix.txt")</pre>
chrom <- "chr2"
start <- 38290160
end <- 38303219
gen <- "hg19"
if(interactive()){
  genetrack <-genesENSEMBL(gen,chrom,start,end,showId=FALSE)</pre>
  snptrack <- snpBiomart(chrom, start, end,</pre>
                          dataset="hsapiens_snp_som", showId=FALSE)
  strutrack <- structureBiomart(chrom, start, end,</pre>
                                   strand, dataset="hsapiens_structvar_som")
  clinVariant<-ClinVarMainTrack(gen,chrom,start,end)</pre>
  clinCNV<-ClinVarCnvTrack(gen,chrom,start,end)</pre>
  gwastrack <-GWASTrack(gen,chrom,start,end)</pre>
  geneRtrack <-GeneReviewsTrack(gen,chrom,start,end)</pre>
```

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```
listgviz <- list(genetrack,snptrack,strutrack,clinVariant,</pre>
                 clinCNV,gwastrack,geneRtrack)
comet(config.file=configfile, MYDATA.FILE=myinfofile, CORMATRIX.FILE=mycorrelation,
      MYDATA.LARGE.FILE=myexpressfile, TRACKS.GVIZ=listgviz,
      VERBOSE=FALSE, PRINT.IMAGE=FALSE,DISP.PVALUEPLOT=FALSE)
} else {
 data(geneENSEMBLtrack)
 data(snpBiomarttrack)
 data(ISCAtrack)
 data(strucBiomarttrack)
 data(ClinVarCnvTrack)
 data(clinVarMaintrack)
 data(GWASTrack)
 data(GeneReviewTrack)
 listgviz <- list(genetrack,snptrack,strutrack,clinVariant,</pre>
                 clinCNV,gwastrack,geneRtrack)
comet(config.file=configfile, MYDATA.FILE=myinfofile, CORMATRIX.FILE=mycorrelation,
      MYDATA.LARGE.FILE=myexpressfile, TRACKS.GVIZ=listgviz,
      VERBOSE=FALSE, PRINT.IMAGE=FALSE,DISP.PVALUEPLOT=FALSE)
}
```

chromatinHMMAll

Creating multiple chromHMM tracks from the UCSC genome browser

#### **Description**

Create multiple chromHMM Broad tracks by connecting to the UCSC genome browser using the GViz bioconductor package

#### Usage

#### **Arguments**

gen	the name of the genome
chr	the chromosome of interest
start	the first position in region of interest (the smallest value)
end	the last position in region of interest (the biggest value)
mySession	the object session from the function browserSession of rtracklayer
track.name	the name of the track, for example : Broad ChromHMM
pattern	the pattern of the track to visualise
table.name	the name of the table from the track

chromatinHMMOne 5

#### Value

list of AnnotationTrack objects of GViz

#### Author(s)

Tiphaine Martin

#### **Examples**

```
library("Gviz")
  gen <- "hg19"
  chr <- "chr2"
  start <- 38290160
  end <- 38313219
if(interactive()){
  BROWSER.SESSION="UCSC"
  mySession <- browserSession(BROWSER.SESSION)</pre>
  genome(mySession) <- gen</pre>
  track.name="Broad ChromHMM"
  tablestrack<-tableNames(ucscTableQuery(mySession, track=track.name))</pre>
  table.name<-tablestrack[1]
  PATTERN.REGULATION<-"GM12878"
 chromhmmPattern<-chromatinHMMAll(gen,chr,start,end,mySession,track.name,PATTERN.REGULATION)</pre>
  plotTracks(chromhmmPattern, from = start, to =end)
  chromhmmNoPattern<-chromatinHMMAll(gen,chr,start,end,mySession,track.name)</pre>
  plotTracks(chromhmmNoPattern, from = start, to =end)
} else {
  data(chromhmmPattern)
  plotTracks(chromhmmPattern, from = start, to =end)
  data(chromhmmNoPattern)
  plotTracks(chromhmmNoPattern, from = start, to =end)
}
```

 $\hbox{chromatin} \hbox{HMMOne}$ 

Creating one chromHMM track from the UCSC genome browser

## Description

Create one track of only one type of chromHMM Broad element from the UCSC genome browser using the Gviz bioconductor package

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#### Usage

## Arguments

gen	the name of the genome
chr	the chromosome of interest
start	the first position in region of interest (the smallest value)
end	the last position in region of interest (the biggest value)
mySession	the object session from the function browserSession of rtracklayer
track.name	the name of the track( Broad ChromHMM )
table.name	the name of the table from the track

#### Value

An AnnotationTrack object of Gviz

#### Author(s)

Tiphaine Martin

```
library("Gviz")
  gen <- "hg19"
  chr <- "chr2"
  start <- 38290160
  end <- 38303219
if(interactive()) {
  BROWSER.SESSION="UCSC"
  mySession <- browserSession(BROWSER.SESSION)</pre>
  genome(mySession) <- gen</pre>
  {\tt track.name="Broad ChromHMM"}
  tablestrack<-tableNames(ucscTableQuery(mySession, track=track.name))</pre>
  table.name<-tablestrack[1]</pre>
  chromhmmtrack one <-chromatin HMMOne (gen, chr, start, end, my Session, track.name, table.name)\\
  plotTracks(chromhmmtrackone, from = start, to =end)
}else {
  data(chromhmmtrackone)
  plotTracks(chromhmmtrackone, from = start, to =end)
}
```

chrUCSC2ENSEMBL 7

chrUCSC2ENSEMBL	Removing "chr" to the chromosome number from UCSC to transform
	it to ENSEMBL chromosome format

## Description

Removing "chr" at the beginning of the chromosome number

#### Usage

```
chrUCSC2ENSEMBL(chr)
```

#### **Arguments**

chr

the chromosome number in UCSC format

#### Value

the number of chromosome at ENSEMBL format

#### Author(s)

Tiphaine Martin

#### **Examples**

```
chr<-"chr7"
chrUCSC2ENSEMBL(chr)</pre>
```

ClinVarCnvTrack

Create one track of the genomic positions of variants from the ClinVar database (CNV only)

## Description

Create one track of the genomic positions of variants from the ClinVar database (CNV only, Variants excluded) using the Gviz bioconductor package

## Usage

```
ClinVarCnvTrack(gen, chr, start, end, showId = FALSE)
```

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#### **Arguments**

gen	the name of the genome
chr	the chromosome of interest
start	the first position in region of interest (the smallest value)
end	the last position in region of interest (the biggest value)

showId Show the ID of the genetic elements

#### Value

An UcscTrack object of Gviz

#### Author(s)

Tiphaine Martin

#### **Examples**

```
library("Gviz")
chrom <- "chr2"
start <- 38290160
end <- 38303219
gen <- "hg19"
if(interactive()){
   clinCNV<-ClinVarCnvTrack(gen,chrom,start,end)
   plotTracks(clinCNV, from = start, to =end)
}else {
   data(ClinVarCnvTrack)
   plotTracks(clinCNV, from = start, to =end)
}</pre>
```

ClinVarMainTrack

Create one track of the genomic positions of variants from the ClinVar database (variants only)

#### **Description**

Create one track of the genomic positions of variants from the ClinVar database (Variants only, CNV excluded) using the Gviz bioconductor package

## Usage

```
ClinVarMainTrack(gen, chr, start, end, showId=FALSE)
```

#### **Arguments**

gen	the name of the genome
chr	the chromosome of interest
start	the first position in region of interest (the smallest value)
end	the last position in region of interest (the biggest value)
showId	Show the ID of the genetic elements

#### Value

An UcscTrack object of Gviz

#### Author(s)

Tiphaine Martin

#### **Examples**

```
library("Gviz")
gen <- "hg19"
chrom <- "chr2"
start <- 38290160
end <- 38303219

if(interactive()) {
   clinVariant<-ClinVarMainTrack(gen,chrom,start,end)
   plotTracks(clinVariant, from = start, to =end)
}else{
   data(clinVarMaintrack)
   plotTracks(clinVariant, from = start, to =end)
}</pre>
```

comet

Visualize EWAS results in a genomic region of interest

#### **Description**

coMET is an R-based package to visualize EWAS (epigenome-wide association scans) results in a genomic region of interest. The main feature of coMET is to plot the the significance level of EWAS results in the selected region, along with correlation in DNA methylation values between CpG sites in the region. The coMET package generates plots of phenotype-association, co-methylation patterns, and a series of annotation tracks.

#### Usage

```
comet(MYDATA.FILE = NULL, MYDATA.FORMAT = "SITE", MYDATA.LARGE.FILE = NULL,
   MYDATA.LARGE.FORMAT = "SITE", CORMATRIX.FILE = NULL, CORMATRIX.METHOD = "spearman",
   CORMATRIX.FORMAT = "RAW", CORMATRIX.COLOR.SCHEME = "bluewhitered", MYDATA.REF = NULL,
    START = NULL, END = NULL, ZOOM = FALSE, LAB.Y = "log", PVAL.THRESHOLD = 1e-05,
   DISP.PVAL.THRESHOLD = 1, DISP.ASSOCIATION = FALSE, DISP.ASSOCIATION.LARGE = FALSE,
     DISP.REGION = FALSE, DISP.REGION.LARGE = FALSE, SYMBOLS = "circle-fill",
     SYMBOLS.LARGE = NA, SAMPLE.LABELS = NULL, SAMPLE.LABELS.LARGE = NULL,
   USE.COLORS = TRUE, DISP.COLOR.REF = TRUE, COLOR.LIST = NULL, COLOR.LIST.LARGE = NULL,
     DISP.MYDATA = TRUE, BIOFEAT.USER.FILE = NULL, BIOFEAT.USER.TYPE = NULL,
   BIOFEAT.USER.TYPE.PLOT = NULL, GENOME = "hg19", DATASET.GENE = "hsapiens_gene_ensembl",
     TRACKS.GVIZ = NULL, TRACKS.GGBIO = NULL, TRACKS.TRACKVIEWER = NULL,
     DISP.MYDATA.NAMES = TRUE, DISP.COLOR.BAR = TRUE, DISP.PHYS.DIST = TRUE,
     DISP.LEGEND = TRUE, DISP.MARKER.LINES = TRUE, DISP.CORMATRIXMAP = TRUE,
     DISP.PVALUEPLOT =TRUE, DISP.TYPE = "symbol", DISP.MULT.LAB.X = FALSE,
     DISP.CONNECTING.LINES = TRUE, PALETTE.FILE = NULL, IMAGE.TITLE = NULL,
    IMAGE.NAME = "coMET", IMAGE.TYPE = NULL, IMAGE.SIZE = 3.5, FONT.FACTOR = NULL,
     SYMBOL.FACTOR = NULL, PRINT.IMAGE = TRUE, CONNECTING.LINES.FACTOR = 1.5,
     CONNECTING.LINES.ADJ = 0.01, CONNECTING.LINES.VERT.ADJ = -1,
     CONNECTING.LINES.FLEX = 0, config.file = NULL, VERBOSE = FALSE)
```

#### **Arguments**

MYDATA.FILE Name of the info file describing the coMET parameters

MYDATA. FORMAT Format of the input data in MYDATA.FILE. There are 4 different options: SITE, REGION, SITE\_ASSOC, REGION\_ASSOC

MYDATA.LARGE.FILE

Name of additional info files describing the coMET parameters. File names should be comma-separated.

MYDATA.LARGE.FORMAT

Format of additional data to be visualised in the p-value plot. File names should be comma-separated.

CORMATRIX.FILE Name of the raw data file or the pre-computed correlation matrix file. CORMATRIX.METHOD

Options for calculating the correlation matrix: spearman, pearson and kendall

CORMATRIX.FORMAT

Format of the input CORMATRIX.FILE. There are two options: raw file (RAW if CpG sites are by column and samples by row or RAW\_REV if CpG site are by row and samples by column) and pre-computed correlation matrix (CORMATRIX)

CORMATRIX.COLOR.SCHEME

Color scheme options: heat, bluewhitered, cm, topo, gray, bluetored

MYDATA.REF The name of the reference CpG-site listed in MYDATA.FILE

START The first nucleotide position to be visualised END the last nucleotide position to be visualised

ZOOM Default=False

LAB.Y Scale of the y-axis. Options: log or ln

PVAL.THRESHOLD Significance threshold to be displayed as a red dashed line

DISP.PVAL.THRESHOLD

Display only the findings that pass the value put in DISP.PVAL.THRESHOLD

DISP.ASSOCIATION

This logical option works only if MYDATA.FILE contains the effect direction (MYDATA.FORMAT=SITE ASSOC or REGION ASSOC). The value can be TRUE or FALSE: if FALSE (default), for each point of data in the p-value plot, the color of symbol is the color of co-methylation pattern between the point and the reference site; if TRUE, the effect direction is shown. If the association is positive, the color is the one defined with the option COLOR.LIST. On the other hand, if the association is negative, the color is the opposed color.

DISP.ASSOCIATION.LARGE

This logical option works only if MYDATA.LARGE.FILE contains the effect direction (MYDATA.LARGE.FORMAT=SITE\_ASSOC or REGION\_ASSOC). The value can be TRUE or FALSE: if FALSE (default), for each point of data in the p-value plot, the color of symbol is the color of co-methylation pattern between the point and the reference site; if TRUE, the effect direction is shown. If the association is positive, the color is the one defined with the option COLOR.LIST.LARGE. On the other hand, if the association is negative, the color is the opposed color.

DISP.REGION

This logical option works only if MYDATA.FILE contains regions (MYDATA.FORMAT=REGION or REGION\_ASSOC). The value can be TRUE or FALSE (default). If TRUE, the genomic element will be shown by a continuous line with the color of the element, in addition to the symbol at the center of the region. If FALSE, only the symbol is shown.

DISP.REGION.LARGE

This logical option works only if MYDATA.LARGE.FILE contains regions (MY-DATA.LARGE.FORMAT=REGION or REGION\_ASSOC). The value can be TRUE or FALSE (default). If TRUE, the genomic element will be shown by a continuous line with the color of the element, in addition to the symbol at the center of the region. If FALSE, only the symbol is shown.

**SYMBOLS** 

The symbol shown in the p-value plot. Options: circle, square, diamond, triangle. Symbols can be filled by appending -fill, e.g. square-fill. Example: circle, diamond-fill, triangle

SYMBOLS.LARGE

The symbol to visualise the data defined in MYDATA.LARGE.FILE. Options: circle, square, diamond, triangle; Symbols can either be filled or not filled by appending -fill e.s., square-fill. Example: circle,diamond-fill,triangle

SAMPLE.LABELS

Labels for the sample described in MYDATA.FILE to include in the legend SAMPLE.LABELS.LARGE

> Labels for the sample described in MYDATA.LARGE.FILE to include in the legend

USE.COLORS

Use the colors defined or use the grey color scheme

DISP. COLOR. REF Logical option TRUE or FALSE (TRUE default). if TRUE, the connection line related to the reference probe is in purple, if FALSE if the connection line related to the reference probe stay black.

COLOR.LIST List of colors for displaying the P-value symbols related to the data in MY-DATA.FILE

COLOR.LIST.LARGE

List of colors for displaying the P-value symbols related to the data in MY-DATA.LARGE.FILE

DISP.MYDATA logical option TRUE or FALSE. TRUE (default). If TRUE, the P-value plot is shown; if FALSE the plot will be defined by GViz

BIOFEAT.USER.FILE

Name of data file to visualise in the tracks. File names should be commaseparated.

BIOFEAT.USER.TYPE

Track type, where multiple tracks can be shown (comma-separated): DataTrack, AnnotationTrack, GeneRegionTrack.

BIOFEAT.USER.TYPE.PLOT

Format of the plot if the data are shown with the Gviz's function called Data-Track (comma-separated)

GENOME The human genome reference file. e.g. "hg19" for Human genome 19 (NCBI 37), "grch37" (GRCh37), "grch38" (GRCh38)

DATASET.GENE The gene names from ENSEMBL. e.g. hsapiens\_gene

TRACKS.GVIZ list of tracks created by Gviz.

TRACKS.GGBIO list of tracks created by ggbio.
TRACKS.TRACKVIEWER

list of tracks created by track viewer.

DISP.MYDATA.NAMES

logical option TRUE or FALSE. If True (default), the names of the CpG sites are displayed.

DISP. COLOR. BAR Color legend for the correlation matrix (range -1 to 1). Default: blue-white-red

DISP.PHYS.DIST logical option (TRUE or FALSE). TRUE (default). Display the bp distance on the plots

DISP.LEGEND logical option TRUE or FALSE. TRUE (default) Display the sample labels and corresponding symbols on the lower right side

DISP.MARKER.LINES

logical option TRUE or FALSE. TRUE (default), if FALSE the red line for PVAL.THRESHOLD is not shown

DISP.CORMATRIXMAP

logical option TRUE or FALSE. TRUE (default), if FALSE correlation matrix is not shown

DISP.PVALUEPLOT

logical option (TRUE or FALSE). TRUE (default), if FALSE the pvalue plot is not shown

DISP.TYPE Default: symbol

DISP.MULT.LAB.X

logical option TRUE or FALSE. FALSE (default). Display evenly spaced X-axis labels; up to 5 labels are shown.

#### DISP.CONNECTING.LINES

logical option TRUE or FALSE. TRUE (default) displays connecting lines between p-value plot and correlation matrix

PALETTE. FILE File that contains color scheme for the heatmap. Colors are hexidecimal HTML

color codes; one color per line; if you do not want to use this option, use the

color defined by the option CORMATRIX.COLOR.SCHEME

IMAGE.TITLE Title of the plot

IMAGE.NAME Path and Name of the plot file without extension

IMAGE.TYPE Options: pdf or eps

IMAGE.SIZE Default: 3.5 inches. Possible sizes: 3.5 or 7
FONT.FACTOR Font size of the sample labels. Range: 0-1

SYMBOL. FACTOR Size of the symbols. Range: 0-1

PRINT. IMAGE Print image in file or not.

CONNECTING.LINES.FACTOR

Length of the connecting lines. Range: 0-2

CONNECTING.LINES.ADJ

Position of the connecting lines horizontally. Negative values shift the connecting lines to the left and positive values shift the lines to the right. Range: (-1;1) option -1 means no connecting lines.

CONNECTING.LINES.VERT.ADJ

Position of the connecting lines vertically. Can be used to vertically adjust the position of the connecting lines in relation to the CpG-site names. Negative value shift the connecting lines down. Range: (-0.5 - 0), option -1 mean the default value related to the plot size (-0.5 for 3.5 plot size; -0.7 for 7.5 plot size)

CONNECTING.LINES.FLEX

Adjusts the spread of the connecting lines. Range: 0-2

config.file Configuration file that contains the values of these options instead of defining

these by command line

VERBOSE logical option TRUE or FALSE. TRUE (default). If TRUE, shows comments.

#### Value

Create a plot

#### Author(s)

Tiphaine Martin

```
extdata <- system.file("extdata", package="coMET",mustWork=TRUE)
configfile <- file.path(extdata, "config_cyp1b1_zoom_4comet.txt")
myinfofile <- file.path(extdata, "cyp1b1_infofile.txt")
myexpressfile <- file.path(extdata, "cyp1b1_infofile_exprGene_region.txt")
mycorrelation <- file.path(extdata, "cyp1b1_res37_rawMatrix.txt")</pre>
```

```
chrom <- "chr2"
start <- 38290160
end <- 38303219
gen <- "hg19"
if(interactive()){
 cat("interactive")
 genetrack <-genesENSEMBL(gen,chrom,start,end,showId=FALSE)</pre>
 snptrack <- snpBiomart(chrom, start, end,</pre>
                        dataset="hsapiens_snp_som", showId=FALSE)
 strutrack <- structureBiomart(chrom, start, end,</pre>
                               strand, dataset="hsapiens_structvar_som")
 clinVariant<-ClinVarMainTrack(gen,chrom,start,end)</pre>
 clinCNV<-ClinVarCnvTrack(gen,chrom,start,end)</pre>
 gwastrack <-GWASTrack(gen,chrom,start,end)</pre>
 geneRtrack <-GeneReviewsTrack(gen,chrom,start,end)</pre>
 listgviz <- list(genetrack,snptrack,strutrack,clinVariant,</pre>
                 clinCNV,gwastrack,geneRtrack)
comet(config.file=configfile, MYDATA.FILE=myinfofile, CORMATRIX.FILE=mycorrelation,
      MYDATA.LARGE.FILE=myexpressfile, TRACKS.GVIZ=listgviz,
      VERBOSE=FALSE, PRINT.IMAGE=FALSE, DISP.PVALUEPLOT=FALSE)
} else {
 cat("Non interactive")
 data(geneENSEMBLtrack)
 data(snpBiomarttrack)
 data(ISCAtrack)
 data(strucBiomarttrack)
 data(ClinVarCnvTrack)
 data(clinVarMaintrack)
 data(GWASTrack)
 data(GeneReviewTrack)
 listgviz <- list(genetrack,snptrack,strutrack,clinVariant,</pre>
                 clinCNV,gwastrack,geneRtrack)
comet(config.file=configfile, MYDATA.FILE=myinfofile, CORMATRIX.FILE=mycorrelation,
      MYDATA.LARGE.FILE=myexpressfile, TRACKS.GVIZ=listgviz,
      VERBOSE=FALSE, PRINT.IMAGE=FALSE, DISP.PVALUEPLOT=FALSE)
}
```

comet.web

Visualize EWAS results in a genomic region of interest with predefined annotation tracks

#### **Description**

coMET is an R-based package to visualize EWAS (epigenome-wide association scans) results in a genomic region of interest. The main feature of coMET is to plot the the significance level of EWAS

results in the selected region, along with correlation in DNA methylation values between CpG sites in the region. The coMET package generates plots of phenotype-association, co-methylation patterns, and a series of annotation tracks.

#### Usage

```
comet.web(MYDATA.FILE = NULL, MYDATA.FORMAT = c("SITE", "REGION", "SITE_ASSO", "REGION_ASSO"),
          MYDATA.LARGE.FILE = NULL,
         MYDATA.LARGE.FORMAT = c("SITE", "REGION", "SITE_ASSO", "REGION_ASSO"),
      CORMATRIX.FILE = NULL, CORMATRIX.METHOD = c("spearman", "pearson", "kendall"),
          CORMATRIX.FORMAT = c("CORMATRIX", "RAW", "RAW_REV"),
          CORMATRIX.COLOR.SCHEME = "heat", MYDATA.REF = NULL,
          GENOME="hg19", START = NULL, END = NULL, ZOOM = FALSE, LAB.Y = "log",
          PVAL.THRESHOLD = 1e-07, DISP.PVAL.THRESHOLD = 1,
          DISP.ASSOCIATION = FALSE, DISP.ASSOCIATION.LARGE = FALSE,
        DISP.REGION = FALSE, DISP.REGION.LARGE = FALSE, SYMBOLS = "circle-fill",
          SYMBOLS.LARGE = NA, SAMPLE.LABELS = NULL, SAMPLE.LABELS.LARGE = NULL,
          USE.COLORS = TRUE, DISP.COLOR.REF = TRUE, COLOR.LIST = NULL,
          COLOR.LIST.LARGE = NULL, BIOFEAT.USER.FILE = NULL,
          BIOFEAT.USER.TYPE = c("GeneRegion", "Annotation", "Data"),
          BIOFEAT.USER.TYPE.PLOT = NULL, LIST.TRACKS = "geneENSEMBL,CGI,
          ChromHMM, DNAse, RegENSEMBL, SNP", PATTERN. REGULATION = "GM12878",
         IMAGE.TITLE = NULL, IMAGE.NAME = "coMET", IMAGE.TYPE = c("pdf", "eps"),
       IMAGE.SIZE = 3.5, PRINT.IMAGE = FALSE, config.file = NULL, VERBOSE = FALSE)
```

#### **Arguments**

MYDATA.FILE Name of the info file describing the coMET parameters

MYDATA. FORMAT Format of the input data in MYDATA.FILE. There are 4 different options: SITE, REGION, SITE ASSOC, REGION ASSOC

MYDATA.LARGE.FILE

Name of additional info files describing the coMET parameters. File names should be comma-separated.

MYDATA.LARGE.FORMAT

Format of additional data to be visualised in the p-value plot. File names should be comma-separated.

CORMATRIX.FILE Name of the raw data file or the pre-computed correlation matrix file.

CORMATRIX.METHOD

Options for calculating the correlation matrix: spearman, pearson and kendall

CORMATRIX.FORMAT

Format of the input CORMATRIX.FILE. There are three options: raw file (RAW if CpG sites are by column and samples by row or RAW\_REV if CpG site are by row and samples by column) and pre-computed correlation matrix (CORMATRIX)

CORMATRIX.COLOR.SCHEME

Color scheme options: heat, bluewhitered, cm, topo, gray, bluetored

MYDATA.REF The name of the reference CpG-site listed in MYDATA.FILE

GENOME The human genome reference file. e.g. "hg19" for Human genome 19 (NCBI

37), "grch37" (GRCh37), "grch38" (GRCh38)

START The first nucleotide position to be visualised END the last nucleotide position to be visualised

ZOOM logical option TRUE or FALSE. FALSE (default)

LAB. Y Scale of the y-axis. Options: log or ln

PVAL.THRESHOLD Significance threshold to be displayed as a red dashed line

DISP.PVAL.THRESHOLD

Display only the findings that pass the value put in DISP.PVAL.THRESHOLD

#### DISP.ASSOCIATION

This logical option works only if MYDATA.FILE contains the effect direction (MYDATA.FORMAT=SITE\_ASSOC or REGION\_ASSOC). The value can be TRUE or FALSE: if FALSE (default), for each point of data in the p-value plot, the color of symbol is the color of co-methylation pattern between the point and the reference site; if TRUE, the effect direction is shown. If the association is positive, the color is the one defined with the option COLOR.LIST. On the other hand, if the association is negative, the color is the opposed color.

#### DISP.ASSOCIATION.LARGE

This logical option works only if MYDATA.LARGE.FILE contains the effect direction (MYDATA.LARGE.FORMA=SITE\_ASSOC or REGION\_ASSOC). The value can be TRUE or FALSE: if FALSE (default), for each point of data in the p-value plot, the color of symbol is the color of co-methylation pattern between the point and the reference site; if TRUE, the effect direction is shown. If the association is positive, the color is the one defined with the option COLOR.LIST.LARGE. On the other hand, if the association is negative, the color is the opposed color.

DISP.REGION

This logical option works only if MYDATA.FILE contains regions (MYDATA.FORMAT=REGION or REGION\_ASSOC). The value can be TRUE or FALSE (default). If TRUE, the genomic element will be shown by a continuous line with the color of the element, in addition to the symbol at the center of the region. If FALSE, only the symbol is shown.

#### DISP.REGION.LARGE

This logical option works only if MYDATA.LARGE.FILE contains regions (MYDATA.LARGE.FORMAT=REGION or REGION\_ASSOC). The value can be TRUE or FALSE (default). If TRUE, the genomic element will be shown by a continuous line with the color of the element, in addition to the symbol at the center of the region. If FALSE, only the symbol is shown.

SYMBOLS The symbol shown in the p-value plot. Options: circle, square, diamond, tri-

angle. Symbols can be filled by appending -fill, e.g. square-fill. Example:

circle, diamond-fill, triangle

SYMBOLS.LARGE The symbol to visualise the data defined in MYDATA.LARGE.FILE. Options:

circle, square, diamond, triangle; Symbols can either be filled or not filled by

appending -fill e.s., square-fill. Example: circle,diamond-fill,triangle

SAMPLE.LABELS Labels for the sample described in MYDATA.FILE to include in the legend

#### SAMPLE.LABELS.LARGE

Labels for the sample described in MYDATA.LARGE.FILE to include in the legend

USE. COLORS Use the colors defined or use the grey color scheme

DISP. COLOR. REF Logical option TRUE or FALSE (TRUE default). if TRUE, the connection line

related to the reference probe is in purple, if FALSE if the connection line related

to the reference probe stay black.

COLOR.LIST List of colors for displaying the P-value symbols related to the data in MY-

DATA.FILE

List of colors for displaying the P-value symbols related to the data in MY-DATA.LARGE.FILE

BIOFEAT.USER.FILE

COLOR, LIST, LARGE

Name of data file to visualise in the tracks. File names should be commaseparated.

BIOFEAT.USER.TYPE

Track type, where multiple tracks can be shown (comma-separated): DataTrack, AnnotationTrack, GeneRegionTrack.

BIOFEAT.USER.TYPE.PLOT

Format of the plot if the data are shown with the Gviz's function called Data-

Track (comma-separated)

List of annotation tracks to visualise. Options include geneENSEMBL, CGI, ChromHMM, DNAse, RegENSEMBL, SNP, transcriptENSEMBL, SNPstoma, SNPstru, SNPstrustoma, ISCA, COSMIC, GAD, ClinVar, GeneReviews, GWAS,

ClinVarCNV, GCcontent, genesUCSC, xenogenesUCSC.

PATTERN.REGULATION

LIST.TRACKS

The cell/tissue or the list of cells/tissues to visualise in the regulation region defined by Broad ChromHMM

IMAGE.TITLE Title of the plot

IMAGE.NAME path and Name of the plot file without extension

IMAGE.TYPE Options: pdf or eps

IMAGE.SIZE Default: 3.5 inches. Possible sizes: 3.5 or 7

PRINT.IMAGE Print image in file or not.

config.file Configuration file that contains the values of these options instead of defining

these by command line

VERBOSE logical option TRUE or FALSE. TRUE (default). If TRUE, shows comments.

#### Author(s)

Tiphaine Martin

```
extdata <- system.file("extdata", package="coMET",mustWork=TRUE)
configfile <- file.path(extdata, "config_cyp1b1_zoom_4webserver.txt")</pre>
```

18 CoreillCNVTrack

CoreillCNVTrack Create one track of the genomic positions of CNV in chromosomal

aberration and inherited disorders from the NIGMS Human Genetic

Cell Repository data

#### Description

Create one track of the genomic positions of copy-number variants (CNVs) in chromosomal aberration and inherited disorder cell lines from the NIGMS Human Genetic Cell Repository using the Gviz bioconductor package

#### Usage

```
CoreillCNVTrack(gen, chr, start, end, showId=FALSE)
```

#### **Arguments**

gen the name of the genome chr the chromosome of interest

start the first position in the region of interest (the smallest value) end the last position in the region of interest (the largest value)

showId Show the ID of the genetic elements

#### Value

An UcscTrack object of Gviz

#### Author(s)

Tiphaine Martin

```
library("Gviz")
gen <- "hg19"
chrom <- "chr2"
start <- 38290160
end <- 38303219
if(interactive()){</pre>
```

COSMICRawTrack 19

```
coreilVariant<-CoreillCNVTrack(gen,chrom,start,end)
plotTracks(coreilVariant, from = start, to =end)
} else {
   data(coreilVarianttrack)
   plotTracks(coreilVariant, from = start, to =end)
}</pre>
```

COSMICRawTrack

Get the genomic positions of variants of COSMIC

## Description

Get the genomic positions of variants of COSMIC, the "Catalogue Of Somatic Mutations In Cancer"

#### Usage

```
COSMICRawTrack(gen, chr, start, end, showId=FALSE)
```

## Arguments

gen	the name of genome
chr	the chromosome of interest
start	the first position of region of interest (the smallest value)
end	the last position of region of interest (the biggest value)
showId	Show Id of genetic elements

#### Author(s)

Tiphaine Martin

```
##---- Should be DIRECTLY executable !! ----
##-- ==> Define data, use random,
##--or do help(data=index) for the standard data sets.
library("Gviz")
chrom <- "chr2"
start <- 38290160
end <- 38303219
gen <- "hg19"
cosmicVariant<-COSMICRawTrack(gen,chrom,start,end)
plotTracks(cosmicVariant)</pre>
```

20 COSMICTrack

COSMICTrack	Create one track of the genomic positions of variants from CO	SMIC
	great	

#### **Description**

Create one track of the genomic positions of variants from COSMIC, the "Catalogue Of Somatic Mutations In Cancer" using the Gviz bioconductor package

#### Usage

```
COSMICTrack(gen, chr, start, end, showId=FALSE)
```

#### **Arguments**

gen	the name of the genome
chr	the chromosome of interest
start	the first position in the region of interest (the smallest value)
end	the last position in the region of interest (the largest value)

Show the ID of the genetic elements

#### Value

showId

An UcscTrack object of Gviz

#### Author(s)

Tiphaine Martin

```
library("Gviz")
chrom <- "chr2"
start <- 38290160
end <- 38303219
gen <- "hg19"
if(interactive()){
  cosmicVariant<-COSMICTrack(gen,chrom,start,end)
  plotTracks(cosmicVariant, from = start, to =end)
}else {
  data(cosmicVarianttrack)
  plotTracks(cosmicVariant, from = start, to =end)
}</pre>
```

cpgIslandsUCSC 21

cpg	$T \sim 1$	~~~	_ I I	CCC
CITO	1 5 1	and	S 1 1	1 71

create track CpG Island from UCSC

## Description

create track CpG Island from UCSC using the Gviz bioconductor package

#### Usage

```
cpgIslandsUCSC(gen, chr, start, end)
```

## Arguments

gen	the name of the genome
chr	the chromosome of interest
start	the first position in the region of interest (the smallest value)
end	the last position in the region of interest (the largest value)

#### Value

An UcscTrack object of Gviz

## Author(s)

Tiphaine Martin

```
library("Gviz")
chrom <- "chr2"
start <- 38290160
end <- 38303219
gen <- "hg19"

if(interactive()) {
  cpgIstrack<-cpgIslandsUCSC(gen, chrom, start, end)
  plotTracks(cpgIstrack, from = start, to =end)
}else {
  data(cpgIslandtrack)
  plotTracks(cpgIstrack, from = start, to =end)
}</pre>
```

22 DNAseUCSC

DNAseUCSC	Creation of an UCSC's DNase clusters track	

#### **Description**

Creation of DNase cluster track from a connection to UCSC genome browser in using the GViz bioconductor package

#### Usage

```
DNAseUCSC(gen, chr, start, end, mySession, track.name = "DNase Clusters", table.name = NULL)
```

#### **Arguments**

gen	the name of the genome
chr	the chromosome of interest
start	the first position in the region of interest (the smallest value)
end	the last position in the region of interest (the largest value)
mySession	the object session from the function browserSession of rtracklayer
track.name	the name of the track DNAseUCSC. "DNase Clusters" (default)
table.name	the name of the table from the track

#### Value

An AnnotationTrack object of Gviz

#### Author(s)

Tiphaine Martin

```
library("Gviz")
gen <- "hg19"
chr <- "chr7"
start <- 38290160
end <- 38303219
if(interactive()){
   BROWSER.SESSION="UCSC"
   mySession <- browserSession(BROWSER.SESSION)
   genome(mySession) <- gen
   track.name="Broad ChromHMM"
   tablestrack<-tableNames(ucscTableQuery(mySession, track=track.name))
   table.name<-tableStrack[1]
   dnasetrack<-DNAseUCSC(gen,chr,start,end,mySession)
   plotTracks(dnasetrack, from = start, to =end)
}else {</pre>
```

GADTrack 23

```
data(dnasetrack)
plotTracks(dnasetrack, from = start, to =end)
}
```

**GADTrack** 

Create one track of the genomic positions of variants from the Genetic Association Database (GAD)

#### Description

Create one track of the genomic positions of variants from the Genetic Association Database (GAD) (archive of human genetic association studies of complex diseases and disorders) using the Gviz bioconductor package

#### Usage

```
GADTrack(gen, chr, start, end, showId=FALSE)
```

#### **Arguments**

gen	the name of the genome
chr	the chromosome of interest
start	the first position in the region of interest (the smallest value)
end	the last position in the region of interest (the largest value)

showId Show the ID of the genetic elements

#### Value

An UcscTrack object of Gviz

#### Author(s)

Tiphaine Martin

```
library("Gviz")
gen2 <- "hg19"
chrom2 <- "chr2"
start2 <- 38290160
end2 <- 38303219

if(interactive()) {
   gadtrack<-GADTrack(gen=gen2 ,chr=chrom2 ,start=start2 ,end=end2)
   plotTracks(gadtrack, from = start2, to =end2)
} else {
   data(gadtrack)</pre>
```

24 gcContent

```
plotTracks(gadtrack, from = start2, to =end2)
}
```

gcContent

Create one track of GC content from UCSC

#### **Description**

Create a track of GC content from UCSC using the Gviz bioconductor package

#### Usage

```
gcContent(gen, chr, start, end)
```

## Arguments

gen the name of the genome chr the chromosome of interest

start the first position in the region of interest (the smallest value) end the last position in the region of interest (the largest value)

#### Value

A UcscTrack object of Gviz

#### Author(s)

Tiphaine Martin

```
library("Gviz")
gen <- "hg19"
chr <- "chr7"
start <- 38290160
end <- 38303219

if(interactive()){
   gctrack<-gcContent(gen,chr,start,end)
   plotTracks(gctrack,from= start, to=end)
} else {
   data(gctrack)
   plotTracks(gctrack,from= start, to=end)
}</pre>
```

GeneReviewsTrack 25

GeneReviewsTrack	Create one track of the genomic positions of variants from GeneReviews
	views

## Description

Create one track of the genomic positions of variants from GeneReviews using the Gviz bioconductor package

## Usage

```
GeneReviewsTrack(gen, chr, start, end, showId=FALSE)
```

#### **Arguments**

gen	the name of the genome
chr	the chromosome of interest
start	the first position in the region of interest (the smallest value)
end	the last position in the region of interest (the largest value)
showId	Show the ID of the genetic elements

#### Value

An UcscTrack object of Gviz

#### Author(s)

Tiphaine Martin

```
library("Gviz")
gen <- "hg19"
chr <- "chr2"
start <- 38290160
end <- 38303219
if(interactive()){
  geneRtrack <-GeneReviewsTrack(gen,chrom,start,end,showId=TRUE)
  plotTracks(geneRtrack, from = start, to = end)
} else {
  data(GeneReviewTrack)
  plotTracks(geneRtrack, from = start, to = end)
}</pre>
```

26 genesENSEMBL

genesENSEMBL	Create one track of the genes in the genomic regions of interest from EMSEMBL
	EMSEMBL

## Description

Create one track of the genes in the genomic regions of interest from EMSEMBL using the Gviz bioconductor package

#### Usage

```
genesENSEMBL(gen, chr, start, end, showId=FALSE)
```

## Arguments

gen	the name of the genome
chr	the chromosome of interest
start	the first position in the region of interest (the smallest value)
end	the last position in the region of interest (the largest value)
showId	Show the ID of the genetic elements

#### Value

A BiomartGeneRegionTrack object of Gviz

#### Author(s)

Tiphaine Martin

```
library("Gviz")
gen <- "hg19"
chrom <- "chr2"
start <- 38290160
end <- 38303219
if(interactive()) {
  genetrack <-genesENSEMBL(gen,chrom,start,end,showId=FALSE)
  plotTracks(genetrack, from = start, to =end)
} else {
  data(geneENSEMBLtrack)
  plotTracks(genetrack, from = start, to =end)
}</pre>
```

genesNameENSEMBL 27

•	Obtain the genes names in the genomic regions of interest from EN-SEMBL
---	---

## Description

Obtain the genes names in the genomic regions of interest from ENSEMBL

## Usage

```
genesNameENSEMBL(gen, chr, start, end, dataset)
```

## Arguments

gen	the name of the genome
chr	the chromosome of interest
start	the first position in the region of interest (the smallest value)
end	the last position in the region of interest (the largest value)
dataset	Name of the database to select genes

## Author(s)

Tiphaine Martin

```
library("Gviz")
gen <- "hg19"
chr <- "chr7"
start <- 38290160
end <- 38303219

if(interactive()){
   dataset<- "hsapiens_gene_ensemb1"
   geneNameEnsemb1<- genesNameENSEMBL(gen,chr,start,end,dataset)
   geneNameEnsemb1
} else {
   data(geneNameEnsemb1)
   geneNameEnsemb1
}</pre>
```

28 GWASTrack

GWASTrack Create one track of the genomic positions of variants from the GWAS catalog	GWASTrack	
---	-----------	--

## Description

Create one track of the genomic positions of variants from the NHGRI Catalog of Published Genome-Wide Association Studies using the Gviz bioconductor package

#### Usage

```
GWASTrack(gen, chr, start, end, showId=FALSE)
```

#### **Arguments**

gen	the name of the genome
chr	the chromosome of interest
start	the first position in the region of interest (the smallest value)
end	the last position in the region of interest (the largest value)
showId	Show the ID of the genetic elements

#### Value

An UcscTrack object of Gviz

#### Author(s)

Tiphaine Martin

```
library("Gviz")
gen <- "hg19"
chr <- "chr2"
start <- 37949607
end <- 37965207

if(interactive()) {
   gwastrack <-GWASTrack(gen,chrom,start,end)
   plotTracks(gwastrack, from = start, to =end)
} else {
   data(GWASTrack)
   plotTracks(gwastrack, from = start, to =end)
}</pre>
```

HistoneAll 29

HistoneAll	Create multiple tracks of histone modifications from the UCSC genome browser

## Description

Create multiple tracks of histone modifications from the UCSC genome browser (ENCODE/Broad) using the Gviz bioconductor package

## Usage

#### **Arguments**

n	the name of the genome	
r	the chromosome of interest	
art	the first position in the region of interest (the smallest value)	
d	the last position in the region of interest (the largest value)	
Session	the object session from the function browserSession of rtracklayer	r
ttern	The cell type	
ack.name	the name of the track, for example: "Broad Histone"	
ble.name	the name of the table from the track	
d Session ttern ack.name	the first position in the region of interest (the smallest value) the last position in the region of interest (the largest value) the object session from the function browserSession of rtrackl The cell type the name of the track, for example: "Broad Histone"	laye

#### Value

A list of AnnotationTrack object of Gviz

#### Author(s)

Tiphaine Martin

```
library("Gviz")
gen <- "hg19"
chr <- "chr2"
start <- 38290160
end <- 38313219

if(interactive()){
   BROWSER.SESSION="UCSC"
   mySession <- browserSession(BROWSER.SESSION)
   genome(mySession) <- gen
   pattern1 <- "GM12878"</pre>
```

30 HistoneOne

```
histonalltrack<-HistoneAll(gen,chr,start,end,mySession, pattern=pattern1,track.name="Broad Histone")
plotTracks(histonalltrack, from = start, to =end)
} else {
   data(histonalltrack)
   plotTracks(histonalltrack, from = start, to =end)
}</pre>
```

HistoneOne

Create one track of one histone modification profile from the UCSC genome browser

## Description

Create one track of one histone modification profile from the UCSC genome browser (ENCODE/Broad) using the Gviz bioconductor package

## Usage

#### **Arguments**

gen	the name of the genome
chr	the chromosome of interest
start	the first position in the region of interest (the smallest value)
end	the last position in the region of interest (the largest value)
mySession	the object session from the function browserSession of rtracklayer
track.name	the name of the track, for example: "Broad Histone"
table.name	the name of the table from the track

#### Value

An AnnotationTrack object of Gviz

#### Author(s)

Tiphaine Martin

ISCATrack 31

#### **Examples**

```
library("Gviz")
gen <- "hg19"
chr <- "chr2"
start <- 38290160
end <- 38303219

if(interactive()) {
   BROWSER.SESSION="UCSC"
   mySession <- browserSession(BROWSER.SESSION)
   genome(mySession) <- gen
   histoneonetrack<-HistoneOne(gen,chr,start,end,mySession)
   plotTracks(histoneonetrack, from = start, to =end)
} else {
   data(histoneonetrack)
   plotTracks(histoneonetrack, from = start, to =end)
}</pre>
```

**ISCATrack** 

Create one track of the genomic positions of variants from ISCA

## Description

Create one track of the genomic positions of variants from International Standards for Cytogenomic Arrays (ISCA) Consortium using the Gviz bioconductor package

#### Usage

```
ISCATrack(gen, chr, start, end, mySession, table.name, showId=FALSE)
```

## Arguments

gen	the name of the genome
chr	the chromosome of interest
start	the first position in the region of interest (the smallest value)
end	the last position in the region of interest (the largest value)
mySession	the object session from the function browserSession of rtracklayer
table.name	A table of ISCAT classifications: iscaBenign, iscaCuratedBenign, iscaCuratedPathogenic, iscaLikelyBenign, iscaLikelyPathogenic, iscaPathGainCum, iscaPathLossCum, iscaPathogenic, iscaUncertain
showId	Show the ID of the genetic elements

#### Value

An UcscTrack object of Gviz

32 knownGenesUCSC

#### Author(s)

Tiphaine Martin

## **Examples**

```
library("Gviz")
gen <- "hg19"
chr <- "chr2"
start <- 38292433
end <- 38305492

if(interactive()){
   BROWSER.SESSION="UCSC"
   mySession <- browserSession(BROWSER.SESSION)
   genome(mySession) <- gen
   iscatrack <-ISCATrack(gen,chrom,start,end,mySession, table="iscaPathogenic")
   plotTracks(iscatrack, from = start, to =end)
} else {
   data(ISCAtrack)
   plotTracks(iscatrack, from = start, to =end)
}</pre>
```

knownGenesUCSC

Create a track of known genes from the UCSC genome browser

#### **Description**

Create a track of known genes from the UCSC genome browser using the Gviz bioconductor package

#### Usage

```
knownGenesUCSC(gen, chr, start, end)
```

#### **Arguments**

gen	the name of the genome
chr	the chromosome of interest
start	the first position in the region of interest (the smallest value)

end the last position in the region of interest (the largest value)

#### Value

An UcscTrack object of Gviz

refGenesUCSC 33

#### Author(s)

Tiphaine Martin

#### **Examples**

```
library("Gviz")
gen <- "hg19"
chr <- "chr7"
start <- 38290160
end <- 38303219

if(interactive()) {
   genesUcsctrack<-knownGenesUCSC(gen,chr,start,end)
   plotTracks(genesUcsctrack, from = start, to =end)
}else {
   data(genesUcsctrack)
   plotTracks(genesUcsctrack, from = start, to =end)
}</pre>
```

refGenesUCSC

Create track reference Genes from UCSC

#### **Description**

Create track reference Genes from UCSC

## Usage

```
refGenesUCSC(gen, chr, start, end, showId=FALSE)
```

#### **Arguments**

gen	the name of genome
chr	the chromosome of interest
start	the first position of region of interest (the smallest value)
end	the last position of region of interest (the biggest value)
showId	Show Id of genetic elements

#### Author(s)

Tiphaine Martin

```
##--- Should be DIRECTLY executable !! ----
##-- ==> Define data, use random,
##--or do help(data=index) for the standard data sets.
```

34 regulationBiomart

regulationBiomart

Create a regulation track from ENSEMBL

## Description

Create a 'Regulation' track from ENSEMBL using the Gviz bioconductor package

#### Usage

```
regulationBiomart(gen, chr, start, end)
```

#### **Arguments**

gen	the name of the genome
chr	the chromosome of interest
start	the first position in the region of interest (the smallest value)
end	the last position in the region of interest (the largest value)

#### Value

An AnnotationTrack object of Gviz

#### Author(s)

Tiphaine Martin

```
library("Gviz")
gen <- "hg19"
chr <- "chr7"
start <- 38290160
end <- 38303219

if(interactive()){
   regulationENSEMBLtrack<-regulationBiomart(gen,chr,start,end)
   plotTracks(regulationENSEMBLtrack, from = start, to =end)
} else {
   data(regulationENSEMBLtrack)
   plotTracks(regulationENSEMBLtrack, from = start, to =end)
}</pre>
```

RepeatMaskerTrack 35

RepeatMaskerTrack	Create one track of the genomic positions of regions from Repeat- MaskerTrack
	MaskerTrack

## Description

Create one track of the genomic positions of regions from RepeatMaskerTrack using the Gviz bioconductor package

## Usage

```
RepeatMaskerTrack(gen, chr, start, end, showId=FALSE)
```

#### **Arguments**

gen	the name of the genome
chr	the chromosome of interest
start	the first position in the region of interest (the smallest value)
end	the last position in the region of interest (the largest value)
showId	Show the ID of the genetic elements

#### Value

An UcscTrack object of Gviz

#### Author(s)

Tiphaine Martin

```
library("Gviz")
gen <- "hg19"
chr <- "chr2"
start <- 38290160
end <- 38303219
if(interactive()){
  rmtrack <-RepeatMaskerTrack(gen,chrom,start,end)
  plotTracks(geneRtrack, from = start, to = end)
} else {
  data(RepeatMaskerTrack)
  plotTracks(rmtrack, from = start, to = end)
}</pre>
```

36 snpBiomart

snpBiomart	Create a short variation track from ENSEMBL
	v

## Description

Create a 'Short Variation' track from ENSEMBL using the Gviz bioconductor package

## Usage

```
snpBiomart(chr, start, end, dataset, showId=FALSE, title = NULL)
```

## Arguments

chr	the chromosome of interest
start	the first position in the region of interest (the smallest value)
end	the last position in the region of interest (the largest value)
dataset	The name of the database. Example "hsapiens_snp_som"
showId	Show the the ID of element or not
title	The name of the annotation track

#### Value

An AnnotationTrack object of Gviz

#### Author(s)

Tiphaine Martin

snpLocationsUCSC 37

snpLocationsUCSC	Create a SNP track from UCSC
------------------	------------------------------

## Description

Create a SNP track from UCSC using the Gviz bioconductor package

## Usage

```
snpLocationsUCSC(gen, chr, start, end, track)
```

## Arguments

gen	the name of the genome
chr	the chromosome of interest
start	the first position in the region of interest (the smallest value)
end	the last position in the region of interest (the largest value)
track	The name of the database. Example "snp138"

#### Value

An UcscTrack object of Gviz

## Author(s)

Tiphaine Martin

```
library("Gviz")
gen <- "hg19"
chr <- "chr7"
start <- 38290160
end <- 38303219

if(interactive()) {
    snpUCSCtrack<-snpLocationsUCSC(gen,chr,start,end,"snp138")
    plotTracks(snpUCSCtrack, from = start, to =end)
} else {
    data(snpUCSCtrack)
    plotTracks(snpUCSCtrack, from = start, to =end)
}</pre>
```

38 structureBiomart

structureBiomart	Create a structural variation track from ENSEMBL	

#### **Description**

Create a 'Structural Variation' track from ENSEMBL using the Gviz bioconductor package

#### Usage

```
structureBiomart(chr, start, end, strand, dataset, showId=FALSE, title = NULL)
```

#### Arguments

chr	the chromosome of interest
start	the first position in the region of interest (the smallest value)
end	the last position in the region of interest (the largest value)
strand	the strand to extract structure data for
dataset	The name of the database. Example "hsapiens_structvar_som"
showId	Show the the ID of the element
title	The name of the annotation track

#### Value

An AnnotationTrack object of Gviz

## Author(s)

Tiphaine Martin

transcriptENSEMBL 39

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Create a track of transcripts from ENSEMBL

#### **Description**

Create a track to visualize different transcripts from ENSEMBL using the Gviz bioconductor package

#### Usage

```
transcriptENSEMBL(gen, chr, start, end, showId = FALSE)
```

## Arguments

gen	the name of the genome
chr	the chromosome of interest
start	the first position in the region of interest (the smallest value)
end	the last position in the region of interest (the largest value)
showId	Show the ID of the genetic elements

#### Value

A BiomartGeneRegionTrack object of Gviz

## Author(s)

Tiphaine Martin

```
library("Gviz")
gen <- "hg19"
chr <- "chr7"
start <- 38290160
end <- 38303219

if(interactive()){
   transENSMBLtrack<-transcriptENSEMBL(gen,chr,start,end)
   plotTracks(transENSMBLtrack, from = start, to =end)
} else {
   data(transENSMBLtrack)
   plotTracks(transENSMBLtrack, from = start, to =end)
}</pre>
```

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