## Package 'RepViz'

October 14, 2021

Type Package

Title Replicate oriented Visualization of a genomic region

```
Version 1.8.0
Author Thomas Faux, Kalle Rytkönen, Asta Laiho, Laura L. Elo
Maintainer Thomas Faux, Asta Laiho <faux.thomas1@gmail.com> <asta.laiho@utu.fi>
Description RepViz enables the view of a genomic region in a simple and
     efficient way. RepViz allows simultaneous viewing of both
     intra- and intergroup variation in sequencing counts of the
     studied conditions, as well as their comparison to the output
     features (e.g. identified peaks) from user selected data analysis
     methods. The RepViz tool is primarily designed for chromatin data
     such as ChIP-seq and ATAC-seq, but can also be used with other
     sequencing data such as RNA-seq, or combinations of different
     types of genomic data.
License GPL-3
Encoding UTF-8
RoxygenNote 6.1.1
VignetteBuilder knitr
Depends R (>= 3.5.1), GenomicRanges (>= 1.30.0), Rsamtools (>=
     1.34.1), IRanges (>= 2.14.0), biomaRt (>= 2.36.0), S4Vectors
     (>= 0.18.0), graphics, grDevices, utils
Suggests knitr, testthat
biocViews WorkflowStep, Visualization, Sequencing, ChIPSeq, ATACSeq,
     Software, Coverage, Genomic Variation
git_url https://git.bioconductor.org/packages/RepViz
git_branch RELEASE_3_13
git_last_commit 700aac6
git_last_commit_date 2021-05-19
Date/Publication 2021-10-14
```

2 RepViz

### **R** topics documented:

RepViz		
Index		3
RepViz	Plot a genomic region	

#### **Description**

Plot a genomic region

#### Usage

```
RepViz(region, genome = c("hg19", "hg38", "mm10"), BAM = NULL,
BED = NULL, avgTrack = TRUE, geneTrack = TRUE, max = NULL,
verbose = TRUE)
```

#### **Arguments**

```
region
                   a GRange object with chr, start, end
                   a character vector 'hg19', 'hg38' or 'mm10'
genome
BAM
                   a path to the BAM related csv input file
BED
                   a path to the BED related csv input file
                   a logical indicating if the average track should be included or not
avgTrack
geneTrack
                   a logical indicating if the gene track should be included or not
                   a numerical vector containing the yaxis maximum value of each BAM track
max
                   a logical indicating whether the progress of the plotting is shown
verbose
```

#### Value

displays the region specified by the user

#### **Examples**

```
region <- GRanges('chr12:110938000-110940000')
setwd(tempdir())
#Copying the files to the user working directory
file.copy(from = list.files(system.file('extdata', package = 'RepViz'), full.names = TRUE),
    to = tempdir())
#Generate the visualization of the given region
RepViz::RepViz(region = region,
    genome = 'hg19',
    BAM = 'BAM_input.csv',
    BED = 'BED_input.csv',
    avgTrack = TRUE,
    geneTrack = TRUE)</pre>
```

# **Index**

RepViz, 2