Package 'regionReport'

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Type Package

Title Generate HTML or PDF reports for a set of genomic regions or DESeq2/edgeR results

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Description Generate HTML or PDF reports to explore a set of regions such as the results from annotation-agnostic expression analysis of RNA-seq data at base-pair resolution performed by derfinder. You can also create reports for DESeq2 or edgeR results.

License Artistic-2.0

LazyData true

URL https://github.com/leekgroup/regionReport

BugReports https://support.bioconductor.org/t/regionReport/

VignetteBuilder knitr

```
biocViews DifferentialExpression, Sequencing, RNASeq, Software,
Visualization, Transcription, Coverage, ReportWriting,
DifferentialMethylation, DifferentialPeakCalling,
ImmunoOncology, QualityControl
```

Depends R(>= 3.2)

Imports BiocStyle (>= 2.5.19), derfinder (>= 1.25.3), DEFormats, DESeq2, GenomeInfoDb, GenomicRanges, knitr (>= 1.6), knitrBootstrap (>= 0.9.0), methods, RefManageR, rmarkdown (>= 0.9.5), S4Vectors, SummarizedExperiment, utils

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RoxygenNote 7.1.1

Encoding UTF-8

Roxygen list(markdown = TRUE)

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derfinderReport	Generate a	HTML/PDF	report	exploring	the	basic	results	from
	derfinder							

Description

This function generates a HTML report exploring the basic results from single base-level approach derfinder analysis results (<www.bioconductor.org/packages/derfinder>). The HTML report itself is generated using rmarkdown (http://rmarkdown.rstudio.com/). It works best after using merg-eResults.

Usage

```
derfinderReport(
   prefix,
   outdir = "basicExploration",
   output = "basicExploration",
   project = prefix,
   browse = interactive(),
   nBestRegions = 100,
   makeBestClusters = TRUE,
   nBestClusters = 2,
   fullCov = NULL,
   hg19 = TRUE,
   p.ideos = NULL,
```

derfinderReport

```
txdb = NULL,
device = "png",
significantVar = "qvalue",
customCode = NULL,
template = NULL,
theme = NULL,
digits = 2,
...
```

```
)
```

Arguments

prefix	The main data directory path where mergeResults was run. It should be the same as mergeResults(prefix).
outdir	The name of output directory relative to prefix.
output	The name of output HTML file (without the html extension).
project	The title of the project.
browse	If TRUE the HTML report is opened in your browser once it's completed.
nBestRegions makeBestCluster	The number of region plots to make, ordered by area.
	If TRUE, plotCluster is used on the nBestClusters regions by area. Note that these plots take some time to make.
nBestClusters	The number of region cluster plots to make by taking the nBestClusters re- gions ranked by area of the cluster.
fullCov	A list where each element is the result from loadCoverage used with cutoff=NULL. Can be generated using fullCoverage.
hg19	If TRUE then the reference is assumed to be hg19 and chromosome lengths as well as the default transcription database (TxDb.Hsapiens.UCSC.hg19.knownGene) will be used.
p.ideos	A list where each element is the result of plotIdeogram. If it's NULL and hg19=TRUE then they are created for the hg19 human reference.
txdb	Specify the transcription database to use for making the plots for the top regions by area. If NULL and hg19=TRUE then TxDb.Hsapiens.UCSC.hg19.knownGene is used.
device	The graphical device used when knitting. See more at http://yihui.name/knitr/options (dev argument).
significantVar	A character variable specifying whether to use the p-values, the FDR adjusted p-values or the FWER adjusted p-values to determine significance. Has to be either 'pvalue', 'qvalue' or 'fwer'.
customCode	An absolute path to a child R Markdown file with code to be evaluated before the reproducibility section. Its useful for users who want to customize the report by adding conclusions derived from the data and/or further quality checks and plots.
template	Template file to use for the report. If not provided, will use the default file found in basicExploration/basicExploration.Rmd within the package source.

theme	A ggplot2 theme to use for the plots made with ggplot2.
digits	The number of digits to round to in the interactive table of the top nBestRegions. Note that p-values and adjusted p-values won't be rounded.
	Arguments passed to other methods and/or advanced arguments. Advanced ar- guments:
	chrsStyle The naming style of the chromosomes. By default, UCSC. See se- qlevelsStyle.
	species Species name. See extendedMapSeqlevels for more information.
	currentStyle Current naming style used. See extendedMapSeqlevels for more information.
	fullRegions Part of the output of mergeResults. Specify it only if you have already loaded it in memory.
	fullNullSummary Part of the output of mergeResults. Specify it only if you have already loaded it in memory.
	fullAnnotatedRegions Part of the output of mergeResults. Specify it only if you have already loaded it in memory.
	optionsStats Part of the output of analyzeChr. Specify it only if you have already loaded it in memory.
	optionsMerge Part of the output of mergeResults. Specify it only if you have already loaded it in memory.
	overviewParams A two element list with base_size and areaRel that control the text size for the genomic overview plots.
	<pre>output_format Either html_document, pdf_document or knitrBootstrap::bootstrap_document unless you modify the YAML template.</pre>
	clean Logical, whether to clean the results or not. Passed to render.
	Passed to extendedMapSeqlevels.

Details

Set output_format to 'knitrBootstrap::bootstrap_document' or 'pdf_document' if you want a HTML report styled by knitrBootstrap or a PDF report respectively. If using knitrBootstrap, we recommend the version available only via GitHub at https://github.com/jimhester/knitrBootstrap which has nicer features than the current version available via CRAN. You can also set the output_format to 'html_document' for a HTML report styled by rmarkdown. The default is set to 'BiocStyle::html_document'.

If you modify the YAML front matter of template, you can use other values for output_format.

The HTML report styled with knitrBootstrap can be smaller in size than the 'html_document' report.

Value

An HTML report with a basic exploration of the derfinder results. See the example output at http: //leekgroup.github.io/regionReport/reference/derfinderReport-example/basicExploration/ basicExploration.html.

derfinderReport

Author(s)

Leonardo Collado-Torres

See Also

mergeResults, analyzeChr, fullCoverage

```
## Load derfinder
library("derfinder")
```

```
## The output will be saved in the 'derfinderReport-example' directory
dir.create("derfinderReport-example", showWarnings = FALSE, recursive = TRUE)
```

```
## For convenience, the derfinder output has been pre-computed
file.copy(system.file(file.path("extdata", "chr21"),
    package = "derfinder",
    mustWork = TRUE
), "derfinderReport-example", recursive = TRUE)
## Not run:
## If you prefer, you can generate the output from derfinder
initialPath <- getwd()
setwd(file.path(initialPath, "derfinderReport-example"))
## Collapse the coverage information
collapsedFull <- collapseFullCoverage(list(genomeData$coverage),
    verbose = TRUE
)
```

```
## Calculate library size adjustments
sampleDepths <- sampleDepth(collapsedFull,
    probs = c(0.5), nonzero = TRUE,
    verbose = TRUE
)</pre>
```

```
## Build the models
group <- genomeInfo$pop
adjustvars <- data.frame(genomeInfo$gender)
models <- makeModels(sampleDepths, testvars = group, adjustvars = adjustvars)
## Analyze chromosome 21
analyzeChr(
    chr = "21", coverageInfo = genomeData, models = models,
    cutoffFstat = 1, cutoffType = "manual", seeds = 20140330, groupInfo = group,
    mc.cores = 1, writeOutput = TRUE, returnOutput = FALSE
)</pre>
```

```
## Change the directory back to the original one
setwd(initialPath)
```

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

```
## Merge the results from the different chromosomes. In this case, there's
## only one: chr21
mergeResults(
    chrs = "21", prefix = "derfinderReport-example",
    genomicState = genomicState$fullGenome
)
## Load the options used for calculating the statistics
load(file.path("derfinderReport-example", "chr21", "optionsStats.Rdata"))
## Generate the HTML report
report <- derfinderReport(</pre>
    prefix = "derfinderReport-example", browse = FALSE,
    nBestRegions = 15, makeBestClusters = TRUE,
    fullCov = list("21" = genomeDataRaw$coverage), optionsStats = optionsStats
)
if (interactive()) {
    ## Browse the report
    browseURL(report)
}
## See the example output at
## http://leekgroup.github.io/regionReport/reference/derfinderReport-example/basicExploration/basicExploration
## Not run:
## Note that you can run the example using:
example("derfinderReport", "regionReport", ask = FALSE)
## End(Not run)
```

DESeq2Report

Generate a HTML/PDF report exploring DESeq2 results

Description

This function generates a HTML report with exploratory data analysis plots for DESeq2 results created with DESeq. Other output formats are possible such as PDF but lose the interactivity. Users can easily append to the report by providing a R Markdown file to customCode, or can customize the entire template by providing an R Markdown file to template.

Usage

```
DESeq2Report(
   dds,
   project = "",
   intgroup,
   colors = NULL,
```

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DESeq2Report

```
res = NULL,
nBest = 500,
nBestFeatures = 20,
customCode = NULL,
outdir = "DESeq2Exploration",
output = "DESeq2Exploration",
browse = interactive(),
device = "png",
template = NULL,
searchURL = "http://www.ncbi.nlm.nih.gov/gene/?term=",
theme = NULL,
digits = 2,
....)
```

Arguments

dds	A DESeqDataSet object with the results from running DESeq.
project	The title of the project.
intgroup	interesting groups: a character vector of names in colData(x) to use for group- ing. This parameter is passed to functions such as plotPCA.
colors	vector of colors used in heatmap. If NULL, then a a default set of colors will be used. This argument is passed to pheatmap.
res	A DESeqResults object. If NULL, then results will be used on dds with default parameters.
nBest	The number of features to include in the interactive table. Features are ordered by their adjusted p-values.
nBestFeatures	The number of best features to make plots of their counts. We recommend a small number, say 20.
customCode	An absolute path to a child R Markdown file with code to be evaluated before the reproducibility section. Its useful for users who want to customize the report by adding conclusions derived from the data and/or further quality checks and plots.
outdir	The name of output directory.
output	The name of output HTML file (without the html extension).
browse	If TRUE the HTML report is opened in your browser once it's completed.
device	The graphical device used when knitting. See more at http://yihui.name/ knitr/options (dev argument).
template	Template file to use for the report. If not provided, will use the default file found in DESeq2Exploration/DESeq2Exploration.Rmd within the package source.
searchURL	A url used for searching the name of the features in the web. By default http://www.ncbi.nlm.nih.gov/ is used which is the recommended option when features are genes. It's only used when the output is a HTML file.
theme	A ggplot2 theme to use for the plots made with ggplot2.

digits	The number of digits to round to in the interactive table of the top nBestFeatures. Note that p-values and adjusted p-values won't be rounded.
	Arguments passed to other methods and/or advanced arguments. Advanced ar- guments:
	software The name of the package used for performing the differential expression analysis. Either DESeq2 or edgeR.
	dge A DGEList object. NULL by default and only used by edgeReport.
	theCall The function call. NULL by default and only used by edgeReport.
	<pre>output_format Either html_document, pdf_document or knitrBootstrap::bootstrap_document unless you modify the YAML template.</pre>
	clean Logical, whether to clean the results or not. Passed to render.

Details

Set output_format to 'knitrBootstrap::bootstrap_document' or 'pdf_document' if you want a HTML report styled by knitrBootstrap or a PDF report respectively. If using knitrBootstrap, we recommend the version available only via GitHub at https://github.com/jimhester/knitrBootstrap which has nicer features than the current version available via CRAN. You can also set the output_format to 'html_document' for a HTML report styled by rmarkdown. The default is set to 'BiocStyle::html_document'.

If you modify the YAML front matter of template, you can use other values for output_format.

The HTML report styled with knitrBootstrap can be smaller in size than the 'html_document' report.

Value

An HTML report with a basic exploration for the given set of DESeq2 results. See an example at http://leekgroup.github.io/regionReport/reference/DESeq2Report-example/DESeq2Exploration.
html.

Author(s)

Leonardo Collado-Torres

```
## Load example data from the pasilla package as done in the DESeq2 vignette
## at
## <https://bioconductor.org/packages/release/bioc/vignettes/DESeq2/inst/doc/DESeq2.html#count-matrix-input>.
library("pasilla")
pasCts <- system.file("extdata",
    "pasilla_gene_counts.tsv",
    package = "pasilla", mustWork = TRUE
)
pasAnno <- system.file("extdata",
    "pasilla_sample_annotation.csv",
    package = "pasilla", mustWork = TRUE
)
cts <- as.matrix(read.csv(pasCts, sep = "\t", row.names = "gene_id"))</pre>
```

edgeReport

```
coldata <- read.csv(pasAnno, row.names = 1)</pre>
coldata <- coldata[, c("condition", "type")]</pre>
coldata$condition <- factor(coldata$condition)</pre>
coldata$type <- factor(coldata$type)</pre>
rownames(coldata) <- sub("fb", "", rownames(coldata))</pre>
cts <- cts[, rownames(coldata)]</pre>
## Create DESeqDataSet object from the pasilla package
library("DESeq2")
dds <- DESeqDataSetFromMatrix(</pre>
    countData = cts,
    colData = coldata,
    design = ~condition
)
dds <- DESeq(dds)
## The output will be saved in the 'DESeq2Report-example' directory
dir.create("DESeq2Report-example", showWarnings = FALSE, recursive = TRUE)
## Generate the HTML report
report <- DESeq2Report(dds, "DESeq2-example", c("condition", "type"),</pre>
    outdir = "DESeq2Report-example"
)
if (interactive()) {
    ## Browse the report
    browseURL(report)
}
## See the example output at
## http://leekgroup.github.io/regionReport/reference/DESeq2Report-example/DESeq2Exploration.html
## Not run:
## Note that you can run the example using:
example("DESeq2Report", "regionReport", ask = FALSE)
## End(Not run)
```

edgeReport

Generate a HTML/PDF report exploring edgeR results

Description

This function generates a HTML report with exploratory data analysis plots for edgeR results created. Other output formats are possible such as PDF reports but they lose the interactivity. Users can easily append to the report by providing a R Markdown file to customCode, or can customize the entire template by providing an R Markdown file to template.

Usage

```
edgeReport(
  dge,
  object,
 project = "",
  intgroup,
  colors = NULL,
  pAdjustMethod = "BH",
  alpha = 0.1,
  independentFiltering = FALSE,
  filter,
  theta,
  filterFun,
  nBest = 500,
 nBestFeatures = 20,
 customCode = NULL,
 outdir = "edgeRexploration",
  output = "edgeRexploration",
  browse = interactive(),
  device = "png",
  template = NULL,
  searchURL = "http://www.ncbi.nlm.nih.gov/gene/?term=",
  theme = NULL,
 digits = 2,
  . . .
)
```

Arguments

dge	A DGEList object.
object	A DGEExact or DGELRT object that contains p-values stored in object\$table\$PValue
project	The title of the project.
intgroup	interesting groups: a character vector of names in colData(x) to use for group- ing. This parameter is passed to functions such as plotPCA.
colors	vector of colors used in heatmap. If NULL, then a a default set of colors will be used. This argument is passed to pheatmap.
pAdjustMethod	the method to use for adjusting p-values, see p.adjust. This argument will be passed to results.
alpha	the significance cutoff used for optimizing the independent filtering (by default 0.1). If the adjusted p-value cutoff (FDR) will be a value other than 0.1, alpha should be set to that value. This argument will be passed to results.
independentFilt	tering
	logical, whether independent filtering should be applied automatically. By de- fault it's set to FALSE in contrast with the default used in results to match edgeR's behavior.

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edgeReport

filter	the vector of filter statistics over which the independent filtering will be opti- mized. By default the logCPM will be used if independentFiltering is set to TRUE. It can also be a length 1 character vector specifying one of the column names of object\$table.
theta	the quantiles at which to assess the number of rejections from independent fil- tering. This argument is passed results.
filterFun	an optional custom function as described in results.
nBest	The number of features to include in the interactive table. Features are ordered by their adjusted p-values.
nBestFeatures	The number of best features to make plots of their counts. We recommend a small number, say 20.
customCode	An absolute path to a child R Markdown file with code to be evaluated before the reproducibility section. Its useful for users who want to customize the report by adding conclusions derived from the data and/or further quality checks and plots.
outdir	The name of output directory.
output	The name of output HTML file (without the html extension).
browse	If TRUE the HTML report is opened in your browser once it's completed.
device	The graphical device used when knitting. See more at http://yihui.name/knitr/options (dev argument).
template	Template file to use for the report. If not provided, will use the default file found in DESeq2Exploration/DESeq2Exploration.Rmd within the package source.
searchURL	A url used for searching the name of the features in the web. By default http://www.ncbi.nlm.nih.gov/ is used which is the recommended option when features are genes. It's only used when the output is a HTML file.
theme	A ggplot2 theme to use for the plots made with ggplot2.
digits	The number of digits to round to in the interactive table of the top nBestFeatures. Note that p-values and adjusted p-values won't be rounded.
•••	Arguments passed to other methods and/or advanced arguments. Advanced ar- guments:
	software The name of the package used for performing the differential expression analysis. Either DESeq2 or edgeR.
	dge A DGEList object. NULL by default and only used by edgeReport.
	theCall The function call. NULL by default and only used by edgeReport.
	<pre>output_format Either html_document, pdf_document or knitrBootstrap::bootstrap_document unless you modify the YAML template.</pre>
	clean Logical, whether to clean the results or not. Passed to render.

Details

Set output_format to 'knitrBootstrap::bootstrap_document' or 'pdf_document' if you want a HTML report styled by knitrBootstrap or a PDF report respectively. If using knitrBootstrap, we recommend the version available only via GitHub at https://github.com/jimhester/knitrBootstrap which has nicer features than the current version available via CRAN.

If you modify the YAML front matter of template, you can use other values for output_format.

This report is similar to the one created by DESeq2Report with two additional plots exclusive for edgeR results. We designed the reports to be very similar intentionally and use the Bioconductor package DEFormats to achieve this goal.

Value

An HTML report with a basic exploration for the given set of edgeR results. See the example report at http://leekgroup.github.io/regionReport/reference/edgeReport-example/edgeRexploration.html.

Author(s)

Leonardo Collado-Torres

```
## Create example data using DEFormats
library("DEFormats")
set.seed(20160407)
counts <- simulateRnaSeqData()</pre>
group <- rep(c("A", "B"), each = 3)</pre>
## Create DGEList object
library("edgeR")
dge <- DGEList(counts, group = group)</pre>
## Perform DE analysis with edgeR
design <- model.matrix(~group)</pre>
dge <- estimateDisp(dge, design)</pre>
fit <- glmFit(dge, design)</pre>
lrt <- glmLRT(fit, coef = 2)</pre>
## The output will be saved in the 'edgeReport-example' directory
dir.create("edgeReport-example", showWarnings = FALSE, recursive = TRUE)
## Generate the HTML report
report <- edgeReport(dge, lrt,</pre>
    project = "edgeR-example", intgroup = "group",
    outdir = "edgeReport-example"
)
if (interactive()) {
    ## Browse the report
    browseURL(report)
}
## See the example report at
## http://leekgroup.github.io/regionReport/reference/edgeReport-example/edgeRexploration.html
## Not run:
## Note that you can run the example using:
example("edgeReport", "regionReport", ask = FALSE)
```

End(Not run)

renderReport

Generate a HTML/PDF report exploring a set of genomic regions

Description

This function generates a HTML report with quality checks, genome location exploration, and an interactive table with the results. Other output formats are possible such as PDF but lose the interactivity. Users can easily append to the report by providing a R Markdown file to customCode, or can customize the entire template by providing an R Markdown file to template.

Usage

```
renderReport(
  regions,
  project = "",
  pvalueVars = c(`P-values` = "pval"),
  densityVars = NULL,
  significantVar = mcols(regions)$pval <= 0.05,</pre>
  annotation = NULL,
  nBestRegions = 500,
  customCode = NULL,
  outdir = "regionExploration",
  output = "regionExploration",
  browse = interactive(),
  txdb = NULL,
  device = "png",
 densityTemplates = list(Pvalue = templatePvalueDensity, Common = templateDensity,
   Manhattan = templateManhattan),
  template = NULL,
  theme = NULL,
 digits = 2,
  . . .
)
templatePvalueDensity
templateDensity
templateManhattan
templatePvalueHistogram
templateHistogram
```

Arguments

regions	The set of genomic regions of interest as a GRanges object. All sequence lengths must be provided.
project	The title of the project.
pvalueVars	The names of the variables with values between 0 and 1 to plot density values by chromosome and a table for commonly used cutoffs. Most commonly used to explore p-value distributions. If a named character vector is provided, the names are used in the plot titles.
densityVars	The names of variables to use for making density plots by chromosome. Com- monly used to explore scores and other variables given by region. If a named character vector is provided, the names are used in the plot titles.
significantVar	A logical variable differentiating statistically significant regions from the rest. When provided, both types of regions are compared against each other to see differences in width, location, etc.
annotation	The output from matchGenes used on regions. Note that this can take time for a large set of regions so it's better to pre-compute this information and save it.
nBestRegions	The number of regions to include in the interactive table.
customCode	An absolute path to a child R Markdown file with code to be evaluated before the reproducibility section. Its useful for users who want to customize the report by adding conclusions derived from the data and/or further quality checks and plots.
outdir	The name of output directory.
output	The name of output HTML file (without the html extension).
browse	If TRUE the HTML report is opened in your browser once it's completed.
txdb	Specify the transcription database to use for identifying the closest genes via matchGenes. If NULL it will use TxDb.Hsapiens.UCSC.hg19.knownGene by default.
device	The graphical device used when knitting. See more at http://yihui.name/knitr/options (dev argument).
densityTemplate	25
	A list of length 3 with templates for the p-value density plots (variables from pvalueVars), the continuous variables density plots (variables from densityVars), and Manhattan plots for the p-value variables (pvalueVars). These templates are processed by whisker.render. Check the default templates for more information. The densityTemplates argument is available for those users interested in customizing these plots. For example, to show histograms instead of density plots use templatePvalueHistogram and templateHistogram instead of templatePvalueDensity and templateDensity respectively.
template	Template file to use for the report. If not provided, will use the default file found in regionExploration/regionExploration.Rmd within the package source.
theme	A ggplot2 theme to use for the plots made with ggplot2.
digits	The number of digits to round to in the interactive table of the top nBestRegions. Note that p-values and adjusted p-values won't be rounded.

. . .

Arguments passed to other methods and/or advanced arguments. Advanced arguments: overviewParams A two element list with base_size and areaRe1 that control

- the text size for the genomic overview plots.
- output_format Either html_document, pdf_document or knitrBootstrap::bootstrap_document unless you modify the YAML template.
- clean Logical, whether to clean the results or not. Passed to render.

Format

An object of class character of length 1.

Details

Set output_format to 'knitrBootstrap::bootstrap_document' or 'pdf_document' if you want a HTML report styled by knitrBootstrap or a PDF report respectively. If using knitrBootstrap, we recommend the version available only via GitHub at https://github.com/jimhester/knitrBootstrap which has nicer features than the current version available via CRAN. You can also set the output_format to 'html_document' for a HTML report styled by rmarkdown. The default is set to 'BiocStyle::html_document'.

If you modify the YAML front matter of template, you can use other values for output_format.

The HTML report styled with knitrBootstrap can be smaller in size than the 'html_document' report.

Value

An HTML report with a basic exploration for the given set of genomic regions. See the example report at http://leekgroup.github.io/regionReport/reference/renderReport-example/regionExploration.html.

Author(s)

Leonardo Collado-Torres

```
## Load derfinder for an example set of regions
library("derfinder")
regions <- genomeRegions$regions
## Assign chr length
```

```
library("GenomicRanges")
seqlengths(regions) <- c("chr21" = 48129895)</pre>
```

```
## The output will be saved in the 'renderReport-example' directory
dir.create("renderReport-example", showWarnings = FALSE, recursive = TRUE)
## Generate the HTML report
report <- renderReport(regions, "Example run",</pre>
   pvalueVars = c(
       "Q-values" = "qvalues", "P-values" = "pvalues"
   ), densityVars = c(
       "Area" = "area", "Mean coverage" = "meanCoverage"
   ),
   significantVar = regions$qvalues <= 0.05, nBestRegions = 20,</pre>
   outdir = "renderReport-example"
)
if (interactive()) {
    ## Browse the report
   browseURL(report)
}
## See the example report at
## http://leekgroup.github.io/regionReport/reference/renderReport-example/regionExploration.html
## Check the default templates. For users interested in customizing these
## plots.
## For p-value variables:
cat(regionReport::templatePvalueDensity)
## For continous variables:
cat(regionReport::templateDensity)
## For Manhattan plots
cat(regionReport::templateManhattan)
*****
## bumphunter example mentioned in the vignette ##
****
## Load bumphunter
library("bumphunter")
## Create data from the vignette
pos <- list(</pre>
   pos1 = seq(1, 1000, 35),
   pos2 = seq(2001, 3000, 35),
   pos3 = seq(1, 1000, 50)
)
chr <- rep(paste0("chr", c(1, 1, 2)), times = sapply(pos, length))</pre>
pos <- unlist(pos, use.names = FALSE)</pre>
## Find clusters
```

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```
## Build simulated bumps
Indexes <- split(seq_along(cl), cl)</pre>
beta1 <- rep(0, length(pos))</pre>
for (i in seq(along = Indexes)) {
    ind <- Indexes[[i]]</pre>
    x <- pos[ind]</pre>
    z \le scale(x, median(x), max(x) / 12)
    beta1[ind] <- i * (-1)^(i + 1) * pmax(1 - abs(z)^3, 0)^3 ## multiply by i to vary size
}
## Build data
beta0 <- 3 * sin(2 * pi * pos / 720)
X <- cbind(rep(1, 20), rep(c(0, 1), each = 10))
set.seed(23852577)
error <- matrix(rnorm(20 * length(beta1), 0, 1), ncol = 20)</pre>
y <- t(X[, 1]) %x% beta0 + t(X[, 2]) %x% beta1 + error</pre>
## Perform bumphunting
tab <- bumphunter(y, X, chr, pos, cl, cutoff = .5)</pre>
## Explore data
lapply(tab, head)
library("GenomicRanges")
## Build GRanges with sequence lengths
regions <- GRanges(</pre>
    seqnames = tab$table$chr,
    IRanges(start = tab$table$start, end = tab$table$end),
    strand = "*", value = tab$table$value, area = tab$table$area,
    cluster = tab$table$cluster, L = tab$table$L, clusterL = tab$table$clusterL
)
## Assign chr lengths
seqlengths(regions) <- seqlengths(</pre>
    getChromInfoFromUCSC("hg19", as.Seqinfo = TRUE)
)[
    names(seqlengths(regions))
]
## Explore the regions
regions
## Now create the report
report <- renderReport(regions, "Example bumphunter",</pre>
    pvalueVars = NULL,
    densityVars = c(
        "Area" = "area", "Value" = "value",
        "Cluster Length" = "clusterL"
    ), significantVar = NULL,
    output = "bumphunter-example", outdir = "bumphunter-example",
    device = "png"
)
```

See the example report at
http://leekgroup.github.io/regionReport/reference/bumphunter-example/bumphunter-example.html

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