

Package ‘CNViz’

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Type Package

Title Copy Number Visualization

Version 1.19.0

Description CNViz takes probe, gene, and segment-level log2 copy number ratios and launches a Shiny app to visualize your sample's copy number profile. You can also integrate loss of heterozygosity (LOH) and single nucleotide variant (SNV) data.

Depends R (>= 4.0), shiny (>= 1.5.0)

Imports dplyr, stats, utils, grDevices, plotly, karyoploteR,
CopyNumberPlots, GenomicRanges, magrittr, DT, scales, graphics

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Encoding UTF-8

biocViews Visualization, CopyNumberVariation, Sequencing, DNASEq

RoxygenNote 7.1.1

Suggests rmarkdown, knitr

VignetteBuilder knitr

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all_tcga2018_data	<i>Data from 2018 TCGA studies from cBioPortal</i>
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Description

A dataset containing the study name and aggregated gene level copy number data

Usage

all_tcga2018_data

Format

A data frame with 14944 rows and 6 variables:

- hugoGeneSymbol** hugo gene symbol
- Gain** proportion of cohort with gain in this gene
- Amplification** proportion of cohort with amplification in this gene
- ShallowDeletion** proportion of cohort with shallow deletion in this gene
- DeepDeletion** proportion of cohort with deep deletion in this gene
- study_name** cancer type and sample size

Source

<https://github.com/waldronlab/cBioPortalData> See data-raw folder.

cbio_studies	<i>Names of 2018 TCGA studies from cBioPortal</i>
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Description

A dataset containing the names and studyIds of the 2018 TCGA studies from cBioPortal.

Usage

```
cbio_studies
```

Format

A data frame with 32 rows and 2 variables:

Cancer Name of diagnosis and sample size

studyId studyId that can be used in the cBioPortalData R package

Source

<https://github.com/waldronlab/cBioPortalData> See data-raw folder.

cytoband_data	<i>Genomic locations of cytoband labels</i>
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Description

A dataset containing the chr, start and end position for cytobands according to hg38.

Usage

```
cytoband_data
```

Format

A data frame with 863 rows and 6 variables:

chrom chromosome

chromStart start position

chromEnd end position

name cytoband name

gieStain color

color HEX color

Source

<https://genome.ucsc.edu/cgi-bin/hgTables>

gene_data	<i>Gene data for vignette example</i>
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Description

A dataset containing simulated gene data as sample input for launchCNViz

Usage

```
data(gene_data)
```

Format

A dataframe with 112 rows and 6 variables

chr chromosome

start start location

end end location

gene gene name

log2 log2 copy number ratio

weight weight given to log2 value

loh loss of heterozygosity

Source

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launchCNViz	<i>Launches CNViz, a shiny app to visualize your sample's copy number data.</i>
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Description

CNViz launches a shiny application to visualize your sample's copy number data. At least one of probe_data, gene_data, or segment_data must be supplied; sample_name, variant_data and meta_data are all optional. The more inputs supplied, the more informative the application will be. See the CNViz vignette for more information. Use the hg38 reference genome. CNViz only displays a single sample's data.

Usage

```
launchCNViz(  
  sample_name = "sample",  
  probe_data = data.frame(),  
  gene_data = data.frame(),  
  segment_data = data.frame(),  
  variant_data = data.frame(),  
  meta_data = data.frame()  
)
```

Arguments

sample_name	A string with the ID/name of your sample.
probe_data	A dataframe or GRanges object containing probe-level data. If a dataframe, column names must include chr, gene, start, end, log2. chr/seqnames column should be formatted as 'chr1' through 'chrX', 'chrY'. start, end and log2 should be numeric. If a GRanges object, gene and log2 are metadata columns. Optional column/metadata: weight, where weight is numeric.
gene_data	A dataframe or GRanges object containing gene-level data - one row per gene. If a dataframe, column names must include chr, gene, start, end, log2. chr/seqnames column should be formatted as 'chr1' through 'chrX', 'chrY'. start, end and log2 should be numeric. If a GRanges object, gene and log2 are metadata columns. Optional columns/metadata: weight, loh; where weight is numeric and loh values are TRUE or FALSE.
segment_data	A dataframe or GRanges object containing segment-level data. If a dataframe, column names must include chr, start, end, log2. chr column should be formatted as 'chr1' through 'chrX', 'chrY'. start, end and log2 should be numeric. If a GRanges object, log2 is a metadata column. Optional column/metadata: loh; where loh values are TRUE or FALSE.
variant_data	A dataframe or VRanges object containing SNVs and short indels and columns of your choosing. If a dataframe, the only required columns are gene and mutation_id. Optional column: start; where start indicates the starting position of the mutation. If a VRanges object, make sure gene is one of the metadata columns, so it can be tied to the gene or probe data; a mutation_id column can also be included, otherwise it will be constructed. Additional columns might include depth, allelic_fraction, ref, alt.
meta_data	A dataframe containing your sample's metadata - columns of your choosing. Optional column: ploidy; ploidy will be rounded to the nearest whole number. Additional columns might include purity. This dataframe should only have one row.

Value

a Shiny application

Examples

```

probes <- data.frame(chr = c("chr1", "chr1", "chr4", "chr4", "chrX"),
  gene = c("NOTCH2", "NOTCH2", "KIT", "TET2", "BTK"),
  start = c(119922221, 119967406, 54732072, 105243553, 101360541),
  end = c(119922461, 119967646, 54732192, 105243793, 101360781),
  log2 = c(-0.0832403, -0.0578757, 0.2131540, -0.3189430, -0.7876670),
  weight = c(0.684114, 0.681546, 0.606129, 0.682368, 0.405772))
segments <- data.frame(chr = c("chr1", "chr1", "chr4", "chr4", "chrX"),
  start = c(1050069, 124932724, 1942322, 51743951, 1198732),
  end = c(122026459, 246947668, 49712061, 188110779, 37098762),
  log2 = c(1, 1, 1, 1, 0.5849625), loh = c(FALSE, FALSE, FALSE, TRUE, TRUE))
meta <- data.frame(purity = c(.5),
  ploidy = c(2), sex = c("Female"))

launchCNViz(sample_name = "sample123", probe_data = probes,
  segment_data = segments, meta_data = meta)

```

meta_data

Metadata for vignette example

Description

A dataset containing simulated metadata as sample input for launchCNViz

Usage

```
data(meta_data)
```

Format

A dataframe with 1 rows and 2 variables

purity sample purity

ploidy tumor ploidy

Source

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probe_data	<i>Probe data for vignette example</i>
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Description

A dataset containing simulated probe data as sample input for launchCNViz

Usage

```
data(probe_data)
```

Format

A data frame with 2006 rows and 6 variables:

chr chromosome

start start location

end end location

gene gene name

log2 log2 copy number ratio

weight weight given to log2 value

Source

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segment_data	<i>Segment data for vignette example</i>
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Description

A dataset containing simulated segment data as sample input for launchCNViz

Usage

```
data(segment_data)
```

Format

A dataframe with 101 rows and 5 variables

chr chromosome

start start location

end end location

log2 log2 copy number ratio

loh loss of heterozygosity

Source

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variant_data	<i>Variant data for vignette example</i>
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Description

A dataset containing simulated SNV and indel data as sample input for launchCNViz

Usage

```
data(variant_data)
```

Format

A dataframe with 119 rows and 4 variables

- gene** gene name
- mutation_id** string with information about snv
- depth** read depth
- start** starting location

Source

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