Package 'BioCartaImage'

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Type Package

Title BioCarta Pathway Images

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biocViews Software, Pathways, BioCarta, Visualization

Description The core functionality of the package is to provide coordinates of genes on the BioCarta pathway images and to provide methods to add self-defined graphics to the genes of interest.

VignetteBuilder knitr

URL https://github.com/jokergoo/BioCartaImage

BugReports https://github.com/jokergoo/BioCartaImage/issues

License MIT + file LICENSE NeedsCompilation no RoxygenNote 7.2.3 Encoding UTF-8 Roxygen list(markdown = TRUE) git_url https://git.bioconductor.org/packages/BioCartaImage git_branch devel git_last_commit_f747f75 git_last_commit_date 2025-04-15 Repository Bioconductor 3.22 Date/Publication 2025-07-18 Author Zuguang Gu [aut, cre] (ORCID: <https://orcid.org/0000-0002-7395-8709>) Maintainer Zuguang Gu <z.gu@dkfz.de>

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BioCartaImage-package The BioCartaImage package

Description

BioCarta is a valuable source of biological pathways which not only provides well manually curated pathways, but also remarkable and intuitive pathway images. One useful features of pathway analysis which is to highlight genes of interest on the pathway images is lost. Since the original source of BioCarta (biocarte.com) is lost from the internet, we digged out the data from the internet archive and formatted it into a package.

Details

The core functionality of this package is to highlight certain genes on the pathway image. The **BioCartaImage** package wraps the pathway image as well as gene locations into a graphic object

A simple use is as follows:

```
library(BioCartaImage)
library(grid)
grid.newpage()
grid.biocarta("h_RELAPathway", color = c("1387" = "yellow"))
```

where "h_RELAPathway" is a BioCarta pathway ID, "1387" (in the EntreZ ID type) is the gene to be highlighted. grid.biocarta() is a low-level grid graphical function which adds the pathway graphic to a certain position in the plot.

More advanced use is first to create a graphic object (a grob), later to add more complex graphics to it:

```
grid.newpage()
grob = biocartaGrob("h_RELAPathway")
grob2 = mark_gene(grob, "1387", function(x, y) {
    pos = pos_by_polygon(x, y)
    pushViewport(viewport(x = pos[1] - 10, y = pos[2],
        width = unit(4, "cm"), height = unit(4, "cm"),
        default.units = "native", just = "right"))
```

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all_pathways

```
grid.rect(gp = gpar(fill = "red"))
grid.text("add whatever\nyou want here")
popViewport()
}, capture = TRUE)
grid.draw(grob2)
```

Here biocartaGrob() creates a grob for the pathway image and mark_gene() adds more graphics which are defined by the self-defined function.

For more details, please go to the vignette of this package.

all_pathways All BioCarta pathways

Description

All BioCarta pathways

Usage

all_pathways()

Details

The original BioCarta website (biocarta.com) is retired, but the full list of pathways can be found from archived websites such as https://web.archive.org/web/20170122225118/https://cgap.nci.nih.gov/Pathways/BioCarta_Pathways or https://www.gsea-msigdb.org/gsea/msigdb/human/genesets.jsp?collection=CP:BIOCARTA.

Value

A vector of pathway IDs (the primary pathway IDs on BioCarta).

Examples

all_pathways()

BIOCARTA_PATHWAYS Pre-computed data objects

Description

Pre-computed data objects

Usage

BIOCARTA_PATHWAYS

PATHWAY2BC

PATHWAY2ENTREZ

PATHWAY2MSIGDB

BC2ENTREZ

Format

An object of class list of length 314.

An object of class data.frame with 4428 rows and 2 columns.

An object of class data.frame with 5196 rows and 2 columns.

An object of class data. frame with 292 rows and 2 columns.

An object of class data.frame with 1739 rows and 2 columns.

Details

BIOCARTA_PATHWAYS, PATHWAY2BC, PATHWAY2ENTREZ and BC2ENTREZ are collected from web.archive.org (https://web.archive.org/web/20170122225118/https://cgap.nci.nih.gov/Pathways/BioCarta_ Pathways). PATHWAY2MSIGDB is collected from MSigDB database (https://www.gsea-msigdb. org/gsea/msigdb/human/genesets.jsp?collection=CP:BIOCARTA). The script for generating these datasets can be found at:

system.file("script", "process.R", package = "BioCartaImage")

Value

- BIOCARTA_PATHWAYS: A list of pathway objects. The pathway object is explained in get_pathway().
- PATHWAY2BC: A two-column data frame of pathway IDs and BC IDs.
- PATHWAY2ENTREZ: A two-column data frame of pathway IDs and gene Entrez IDs.
- PATHWAY2MSIGDB: A two-column data frame of pathway IDs and MSigDB IDs.
- BC2ENTREZ: A two-column data frame of BC IDs and gene EntreZ IDs.

The nodes in the original BioCarta pathways are proteins and some of them do not have one-toone mapping to genes, such as protein families or complex. Here BC_ID is the primary ID of proteins/single nodes in BioCarta Pathways and this package provides mapping to gene EntreZ IDs.

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genes_in_pathway Genes in a pathway

Description

Genes in a pathway

Usage

genes_in_pathway(pathway)

Arguments

pathway A BioCarta pathway ID, a MSigDB ID or a biocarta_pathway object. All valid BioCarta pathway IDs are in all_pathways(). The list of MSigDB IDs for BioCarta pathways can be found at https://www.gsea-msigdb.org/gsea/ msigdb/human/genesets.jsp?collection=CP:BIOCARTA.

Value

A character vector of Entrez IDs.

Examples

genes_in_pathway("h_RELAPathway")

get_pathway

Get a single pathway

Description

Get a single pathway

Usage

get_pathway(pathway_id)

Arguments

pathway_idA BioCarta pathway ID. All valid BioCarta pathway IDs are in all_pathways().To make it more convenient to use, the value can also be a MSigDB pathway IDin the BioCarta catalogue. The format should look like: "BIOCARTA_RELA_PATHWAY".

A biocarta_pathway object. The object is a simple list and contains the following elements:

- id: The pathway ID.
- name: The pathway name.
- bc: The nodes in the original BioCarta pathways are proteins and some of them do not have one-to-one mapping to genes, such as protein families or complex. Here bc contains the primary IDs of proteins/single nodes in the pathway. The mapping to genes can be obtained by genes_in_pathway().
- shape: The shape of the corresponding protein/node in the pathway image.
- coords: It is a list of integer vectors, which contains coordinates of the corresponding shapes, in the unit of pixels. This information is retrieved from the HTML source code (in the <area> tag), so the the coordinates start from the top left of the image. The format of the coordinate vectors is c(x1, y1, x2, y2, ...).
- image_file: The file name of the pathway image.

The bc, shape and coords elements have the same length and in the same order.

See Also

The BioCarta pathways on MSigDB: https://www.gsea-msigdb.org/gsea/msigdb/human/genesets.jsp?collection=CP:BIOCARTA.

Examples

```
get_pathway("h_RELAPathway")
get_pathway("BIOCARTA_RELA_PATHWAY")
```

get_pathway_image Download the pathway image

Description

Download the pathway image

Usage

get_pathway_image(pathway)

```
image_dimension(pathway)
```

Arguments

pathway A BioCarta pathway ID, a MSigDB ID or a biocarta_pathway object. All valid BioCarta pathway IDs are in all_pathways(). The list of MSigDB IDs for BioCarta pathways can be found at https://www.gsea-msigdb.org/gsea/ msigdb/human/genesets.jsp?collection=CP:BIOCARTA.

Details

The images are downloaded from https://data.broadinstitute.org/gsea-msigdb/msigdb/ biocarta/human/.

grid.biocarta

Value

get_pathway_image() returns a raster object. image_dimension() returns an integer vector of the height and width of the image.

Examples

```
img = get_pathway_image("h_RELAPathway")
class(img)
# you can directly plot the raster object
plot(img)
```

image_dimension("h_RELAPathway")

grid.biocarta Draw a BioCarta pathway

Description

Draw a BioCarta pathway

Usage

```
grid.biocarta(
  pathway,
  color = NULL,
  x = unit(0.5, "npc"),
  y = unit(0.5, "npc"),
  width = NULL,
  height = NULL,
  just = "centre",
  default.units = "npc",
  name = NULL
)
biocartaGrob(
  pathway,
  color = NULL,
  x = unit(0.5, "npc"),
  y = unit(0.5, "npc"),
  width = NULL,
  height = NULL,
  just = "centre",
  default.units = "npc",
  name = NULL
)
```

Arguments

```
pathway
```

A BioCarta pathway ID, a MSigDB ID or a biocarta_pathway object. All valid BioCarta pathway IDs are in all_pathways(). The list of MSigDB IDs for BioCarta pathways can be found at https://www.gsea-msigdb.org/gsea/msigdb/human/genesets.jsp?collection=CP:BIOCARTA.

color	A named vector where names should correspond to Entrez IDs.
х	A numeric vector or unit object specifying x-location.
У	A numeric vector or unit object specifying y-location.
width	A numeric vector or unit object specifying width.
height	A numeric vector or unit object specifying width.
just	The same as in grid::viewport().
default.units	The same as in grid::viewport().
name	The same as in grid::viewport().

Details

The graphics object contains a pathway image and genes highlighted on the image.

The aspect ratio of the image is kept. If one of width and height is set, the other dimension is calculated by the aspect ratio. If both of width and height is set or inherit from parent viewport, the width and height are automatically adjust to let one dimension completely fill the viewport.

Value

biocartaGrob() returns a gTree object.

Examples

```
library(grid)
grid.newpage()
grid.biocarta("h_RELAPathway")
grob = biocartaGrob("h_RELAPathway")
```

Description

Internal functions for drawing the pathway grob

Usage

```
## S3 method for class 'biocarta_pathway_grob'
makeContext(x)
```

S3 method for class 'biocarta_pathway_grob'
grobWidth(x)

S3 method for class 'biocarta_pathway_grob'
grobHeight(x)

Arguments

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A grob returned by biocartaGrob().

mark_gene

Value

makeContext() returns a grob object.

grobWidth() returns a unit object.

grobHeight() returns a unit object.

mark_gene

Mark a gene on the pathway image

Description

Mark a gene on the pathway image

Usage

mark_gene(grob, entrez_id, fun, min_area = 0, capture = FALSE)

Arguments

grob	A grob returned by biocartaGrob().
entrez_id	A single Entrez ID.
fun	A self-defined function to add graphics to the selected gene.
min_area	Multiple polygons may be used for one single gene in the image. It can be used to select the largest polygon. The unit for calculating the area is the pixel in the image (or more properly, square pixels).
capture	It is suggested to let fun() directly return grob/gTree objects. But you can also directly use functions such as grid.points() or grid.lines() in fun((). In this case, capture must be set to TRUE to capture these graphics.

Details

fun() should be applied to each gene. It is possible an Entrez gene is mapped to multiple nodes in the image, so more precisely, fun() is applied to every node that contains the input gene.

fun() only accepts two arguments, x and y which are two vectors of xy-coordinates that define the polygon. The helper function pos_by_polygon() can be used to get positions around the polygon.

There are two ways to use fun(). First, fun() directly returns a grob. It can be a simple grob, such as by grid::pointsGrob() or complex grob by grid::gTree() and grid::gList(). Second, fun() can directly include plotting functions such as grid::grid.points(), in this case, capture argument must be set to TRUE to capture these graphics.

Value

If capture = FALSE, it must return a grob where new graphics are already added.

Examples

```
library(grid)
grid.newpage()
grob = biocartaGrob("h_RELAPathway")
# gene 1387 is a gene in the pathway
grob2 = mark_gene(grob, "1387", function(x, y) {
pos = pos_by_polygon(x, y)
pointsGrob(pos[1], pos[2], default.units = "native", pch = 16,
 gp = gpar(col = "yellow"))
})
grid.draw(grob2)
grid.newpage()
grob3 = mark_gene(grob, "1387", function(x, y) {
pos = pos_by_polygon(x, y)
grid.points(pos[1], pos[2], default.units = "native", pch = 16,
 gp = gpar(col = "yellow"))
}, capture = TRUE)
grid.draw(grob3)
grid.newpage()
grob4 = mark_gene(grob, "1387", function(x, y) {
pos = pos_by_polygon(x, y)
pushViewport(viewport(x = pos[1] - 10, y = pos[2],
 width = unit(4, "cm"), height = unit(4, "cm"),
 default.units = "native", just = "right"))
 grid.rect(gp = gpar(fill = "red"))
popViewport()
}, capture = TRUE)
grid.draw(grob4)
```

pos_by_polygon Position around a polygon

Description

Position around a polygon

Usage

```
pos_by_polygon(
    x,
    y,
    where = c("left", "right", "top", "bottom", "topleft", "topright", "bottomleft",
        "bottomright")
)
```

Arguments

х	x-coordinate of a polygon.
у	y-coordinate of a polygon.
where	Which side of the polygon? It should take value in c("left", "right", "top",
	"bottom", "topleft", "topright", "bottomleft", "bottomright").

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print.biocarta_pathway

Value

A numeric scalar of length two, which is the xy-coordinate of the point.

Examples

```
x = c(235, 235, 237, 241, 246, 248, 250, 250, 250, 253,
256, 260, 264, 263, 261, 257, 252, 247, 241, 237, 235)
y = c(418, 409, 402, 397, 394, 395, 396, 404, 411, 416, 417,
416, 415, 422, 429, 434, 437, 436, 432, 426, 418)
pos_by_polygon(x, y, "left")
pos_by_polygon(x, y, "bottomleft")
```

```
print.biocarta_pathway
```

Print the biocarta_pathway object

Description

Print the biocarta_pathway object

Usage

```
## S3 method for class 'biocarta_pathway'
print(x, ...)
```

Arguments

х	A biocarta_pathway object.
	Other arguments.

Details

It prints two numbers:

- The number of nodes without removing duplicated ones.
- The number of unique genes that are mapped to the pathway.

Value

Nothing.

Examples

```
p = get_pathway("h_RELAPathway")
p
```

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