Package ‘idiogram’

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Author Karl J. Dykema <karl.dykema@vai.org>
Maintainer Karl J. Dykema <karl.dykema@vai.org>
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Description

This function will take the name of a data package and build a chromLocation object representing that data set. It has also been modified to allow further breakup of the chromLocs.

Usage

\[
\text{buildChromLocation.2(dataPkg, major=NULL)}
\]

Arguments

dataPkg The name of the data package to be used
major name of major breakpoint by which to divide chromosomes, "arms", "bands", and "mb" currently work.

Details

The requested data set must be available in the user's .libPaths(), and the function will throw an error if this is not the case.

If the data package is present, the necessary information will be extracted from the data package and a chromLocation object will be created.

If "major" is set to "arms", the the chromLocs object is populated with data from the chromosome arms; "1p", "1q", "2p", etc... Rat and Human chromosomes follow this pattern, so data packages from both species should work with this function.

If "major" is set to "bands", the chromosomes are divided up based upon which band they fall into.

If "major" is set to "mb", chromosomes are split into 3000+ megabase segments. Note, this creates a very large chromLocation object.

Note, "major" can contain multiple breakpoint names, eg. major=c("arms","bands")

If the "major" argument is used, it stores a list of the extra chromosome names. chromLoc@chromLocs$armList - (or bandList, mbList)

Value

A chromLocation object representing the specified data set.

Author(s)

Main author: Jeff Gentry with minor additions by: Karl Dykema
Examples

```r
## A bit of a hack to not have a package dependency on hgu95av2
## but need to fiddle w/ the warn level to not fail the example anyways.
curWarn <- getOption("warn")
options(warn=0)
on.exit(options(warn=curWarn), add=TRUE)
if (require(hgu95av2.db) & require(idiogram)) {
data(Hs.cytoband)
z <- buildChromLocation.2("hgu95av2.db", major="arms")
} else print("This example requires the hgu95av2.db data package")
```

---

**cytoband-class**

*Class* "cytoband"

---

**Description**

Cytogenetic banding information

**Objects from the Class**

Objects can be created by calls of the form `new("cytoband", ...)`.

**Slots**

- `stain`: typical staining designation from classical cytogenetics
- `band`: character string representing the band name/number
- `start`: position in basepairs for the start of a given band
- `end`: position in basepairs for the end of a given band
- `length`: length in basepairs for a given band

**Author(s)**

Karl Dykema <karl.dykema@vai.org>

**See Also**

See Also as `Hs.cytoband`

**Examples**

```r
## None
```
**Hs.cytoband**  
*Cytogenetic Banding information*

**Description**

Cytogenetic banding information for Homo sapiens (Hs), Mus musculus (Mm) and Rattus norvegicus (Rn)

**Usage**

```r
data(Hs.cytoband)
```

**Format**

Hs.cytoband is an environment containing 24 objects of class "cytoband" as defined by the idiogram package. Rn.cytoband and Mm.cytoband have also been included for the rat and mouse genomes.

**Source**

UCSC Genome Browser http://genome.ucsc.edu/

**Examples**

```r
data(Hs.cytoband)
cyto <- get("1", env=Hs.cytoband)
bands <- matrix(cyto@end-cyto@start, ncol=1)
barplot(bands, col="white")
```

---

**idiograb**  
*idiograb*

**Description**

idiograb reads the position of the graphics pointer when the (first) mouse button is pressed.  
idiogram also reads a second position of the graphics pointer after another mouse button press.  
The two points selected are used to define a diagonal line from which a bounding box will be constructed. It then retrieves the gene identifiers of the points that lie within the bounding box.

**Usage**

```r
idiograb(idio, show.box = TRUE, brush = NULL, ...)
```
idual

Arguments

- `idiogram` point coordinates and corresponding gene identifiers from an idiogram function call
- `show.box` boolean. if TRUE, a box is drawn showing the selected region
- `brush` a color to highlight the points within the selected region
- `...` additional plotting parameters passed to `points` to modify the points within the selected region

Details

Coordinates can be passed in a plotting structure (a list with `x`, `y`, and `labels` components). Typically this is generated from a call to `idiogram`.

The points selected are used to define the top-left and bottom-right locations or the bottom-left and top-right locations for bounding box. These locations can be selected in any order. A character vector of the labels of all the points that lie within the selected region is returned.

Value

A character vector of gene identifiers

Author(s)

Karl Dykema <karl.dykema@vai.org>

See Also

idiogram

Examples

data(idiogramExample)
ip <- idiogram(colo.eset[,1],ucsf.chr,chr="1")
if(interactive()) idiogram(ip,brush="red")

idiogram

Plotting of Genomic Data

Description

Function for plotting genomic data along with corresponding cytogenetic banding information

Usage

`idiogram(data, genome, chr=NULL, organism=NULL, method=c("plot","matplot","image"), margin=c("ticks","idiogram"), grid.col=c("red","grey"), grid.lty=c(1,2), widths=c(1,2), relative=FALSE, dlim=NULL, main=NA, xlab=NA, ylab=NA, cex.axis=.7, na.color = par("bg"), cen.color="red", mb=FALSE, ...)`
Arguments

- **data**: a vector or matrix of numeric data to plot. The names/rownames attribute needs to contain corresponding gene identifiers.
- **genome**: a chromLocation object associated with the specified data set. See below for details.
- **chr**: which chromosome to plot
- **organism**: if NULL, determination of the host organism will be retrieved from the organism slot of the chromLocation object. Otherwise "h", "r", or "m" can be used to specify human, rat, or mouse chromosome information.
- **method**: plotting method
- **margin**: type of banding information to display in the plot margin
- **grid.col**: a two element vector specifying the centromere and band grid colors.
- **grid.lty**: a two element vector specifying the centromere and band grid line type.
- **widths**: a two element vector specifying the relative width of the margin idiogram two the adjacent graph. This option is currently ignored.
- **relative**: If relative is TRUE, the vertical height of the plot is scaled relative to the size of largest chromosome.
- **dlim**: a two element vector specifying the minimum and maximum values for x. Data in x that exceed the min and max limits will be set to the min/max limit values before plotting.
- **main**: an overall title for the plot. Defaults to the chromosome name.
- **xlab**: a title for the x axis.
- **ylab**: a title for the y axis.
- **cex.axis**: the magnification to be used for axis annotation relative to the current.
- **na.color**: color to be used for NA values, defaults to: par("bg")
- **cen.color**: color to be used for the centromere when margin="idiogram", defaults to: "red"
- **mb**: if TRUE, subset data to include just those row names starting with the value of the argument chr. Otherwise, use chromLoc as outlined below.
- **...**: additional graphical parameters can be given as arguments.

Details

This function displays cytogenetic banding information in the plot margin and calls a secondary plotting function to display associated data at the same relative position. Cytogenetic data for human, mouse, and rat genomes are currently included.

The data is arranged by associating gene identifiers to genomic location using a chromLoc annotation object built using the buildChromLocation function from the annotation package. As such, a vector of data is to be plotted, the names attribute of the vector needs to contain the gene identifiers. Likewise if a matrix of data is to be plotted, the rownames attribute of the matrix needs to contain the gene identifiers.

To date, plot can be called for vector data, while matplot and image can be called for matrix data. Most additional plotting arguments can be passed down via .... However, the idiogram function
idiogram plots the axis independently. Currently, only the cex.axis, col.axis, and font.axis parameters are intercepted from ... and redirected to the specialized axis call. Other parameters that effect the axis should be set via par.

The function midiogram is a simple wrapper around idiogram to plot all the chromosomes from a particular organism using sensible default values. The "m" refers to plotting multiple idiograms.

Author(s)
Kyle Furge <kyle.furge@vai.org> and Karl Dykema <karl.dykema@vai.org>

See Also
buildChromLocation, Hs.cytoband, idiograb

Examples

library(idiogram)

## NOTE: This requires an annotation package to work.
## In this example packages "hu6800.db" and "golubEsets" are used.
## They can be downloaded from http://www.bioconductor.org
## "hu6800.db" is under MetaData, "golubEsets" is under Experimental Data.

if(require(hu6800.db) && require(golubEsets)) {
  library(golubEsets)
data(Golub_Train)

  hu.chr <- buildChromLocation("hu6800")
ex <- assayData(Golub_Train)$exprs[,1]

  ## make sure the names() attribute is set correctly
gN <- names(ex)
gN[1:10]

  idiogram(ex,hu.chr,chr="1")

  colors <- rep("black",times=length(ex))
  colors[ex > 10000] <- "red"
  pts[ex > 10000] <- 2
  idiogram(ex,hu.chr,chr="1",col=colors,pch=pts,font.axis=2,cex.axis=1)
  abline(v=0,col="darkgreen")

  ## An example of the dlim option. It is most useful for making
  ## consistant multi-panel plots
  colors <- rep("black",times=length(ex))
  colors[ex > 10000] <- "red"
  colors[ex < 0] <- "blue"
idiogramExample
data included for idiogram package examples

Description
colo.eset and ucsf.chr are included for use in example plots. vai.chr is a chromLocation object included for examples in the package 'reb'. Please see the aCGH Bioconductor package for more information.

Usage
data(idiogramExample)
library(idiogram)
data(Hs.cytoband)
data(idiogramExample)
idiogram(colo.eset, ucsf.chr, chr="1")
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