GenomeGraphs
November 11, 2009

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AnnotationTrack-class

Class "AnnotationTrack"

Description
A generic object to store annotation

Objects from the Class
Objects can be created by calls of the form \texttt{new("AnnotationTrack", \ldots)}.

Slots
\begin{itemize}
  \item \texttt{chr}: Object of class "numeric"
  \item \texttt{strand}: Object of class "numeric"
  \item \texttt{regions}: Object of class "dfOrNULL"
  \item \texttt{dp}: Object of class "DisplayPars"
\end{itemize}

Extends
Class "gdObject ", directly.

Methods
\begin{itemize}
  \item \texttt{drawGD} signature(gdObject = "AnnotationTrack"):
  \item \texttt{getPlotId} signature(obj = "AnnotationTrack"):
  \item \texttt{initialize} signature(.Object = "AnnotationTrack"):
\end{itemize}
BaseTrack-class

Author(s)
James Bullard

Examples
showClass("AnnotationTrack")

Class "BaseTrack" represents base specific data

Description
Represents specific data, e.g. how many times was every base sequenced

Objects from the Class
Objects can be created by calls of the form new("BaseTrack", ...).

Slots
- **base**: Object of class "numeric". Is a vector of base positions
- **value**: Object of class "numeric". Is a vector of corresponding values for every base
- **strand**: Object of class "character" represents that DNA strand
- **dp**: Object of class DisplayPars to control various features of how the track is displayed.

Extends
Class "gdObject", directly.

Methods
- **show** signature(object = "BaseTrack"): ...

Author(s)
Steffen Durinck

References
http://www.stat.berkeley.edu/~steffen/

See Also
objects to See Also as gdPlot

Examples
if (interactive()) {
  data("exampleData", package="GenomeGraphs")
  ga <- new("GenomeAxis")
  bt <- new("BaseTrack", base = yeastCons1[,1], value = yeastCons1[,2],
            dp = DisplayPars(color = "darkblue"))
  gdPlot(list(ga, bt))
}

DisplayPars-class

The DisplayPars class is used to specify graphical parameters to gdObjects.

Description

The DisplayPars functions analogously to par and gp. Generally the class is instantiated using the DisplayPars function rather than directly.

Examples

#

## DisplayPars-class

Class "DisplayPars" is used to specify graphical parameters to gdObjects.

Description

The DisplayPars functions analogously to par and gp. Generally the class is instantiated using the DisplayPars function rather than directly.

Objects from the Class

Objects can be created by calls of the form DisplayPars(...) rather than calls to: new("DisplayPars", ...). by calling the DisplayPars function directly in the constructor the gdObjects are guaranteed to have the appropriate defaults.

Slots

pars: Object of class "environment" Generally this slot is not accessed directly.

Methods

getPar signature(obj = "DisplayPars"): gets a graphical parameter by name

initialize signature(.Object = "DisplayPars"): This constructor should not be called directly.

setPar signature(obj = "DisplayPars"): sets a graphical parameter - see the example below. Often it is easier to set the graphical parameter from within the gdObject.

show signature(object = "DisplayPars"): prints current graphical parameters

Warning

The DisplayPars class should not be manipulated directly. The preferred method for interacting with the class can be seen in the example below.

Author(s)

James Bullard
Examples

```r
showClass("DisplayPars")

if (interactive()) {
  minbase = 10000
  maxbase = 15000
  mart <- useMart("ensembl", dataset = "scerevisiae_gene_ensembl")
  genesplus <- new("GeneRegion", start = minbase, end = maxbase, biomart = mart,
                   strand = "+", chromosome = "I", dp = DisplayPars(size = 2))
  ## plot it.
  gdPlot(list(genesplus, new("Title", title = "genes")), minbase, maxbase)

  ## to obtain a list of the current graphical parameters:
  print(genesplus@dp)

  ## to set a parameter:
  setPar(genesplus, "protein_coding", "pink")
  gdPlot(list(genesplus, new("Title", title = "genes")), minbase, maxbase)
}
```

DisplayPars

`DisplayPars` constructs objects of type `DisplayPars` which are used to effect the display of `gdObject`

Description

`DisplayPars` takes any number of named arguments which will be used by the `drawGD` method of the `gdObject`. These arguments are analagous to both `par` and `gp` of the traditional and grid graphics systems respectively. Different functions support different graphical parameters - thus it is necessary to consult the documentation of the particular `gdObject` to determine which `DisplayPars` will be used.

Usage

`DisplayPars(...)`

Arguments

```
... name value pairs
```

Details

It is not recommended to call `new("DisplayPars", ...)` directly; rather this function `DisplayPars()` should be called instead. If a `gdObject` has already been instantiated then the appropriate method for changing graphical parameters is: `setPar`.

Value

Returns an object of type `DisplayPars`, generally this will be called during a call to the `new` function for a particular `gdObject`. 
Author(s)

James Bullard

Examples

```r
minbase = 10000
gmaxbase = 15000
mart <- useMart("ensembl", dataset = "scerevisiae_gene_ensembl")
genesplus <- new("GeneRegion", start = minbase, end = maxbase, biomart = mart,
    strand = "+", chromosome = "I", dp = DisplayPars(color = "red"))
gaxis <- new("GenomeAxis", add53 = TRUE, add35 = TRUE)
genesminus <- new("GeneRegion", start = minbase, end = maxbase, biomart = mart,
    strand = "-", chromosome = "I", dp = DisplayPars(color = "purple", size = 2))
title <- new("Title", title = "genes in a region")
gdPlot(list(genesplus, gaxis, genesminus, title), minbase, maxbase)
```

drawGD

Generic called on each gdObject to do the plotting.

Description

This generic gets called on each of the gdObjects (obviously the generic is implemented by a method for each object) and thus if a user wishes to implement new gdObjects they need to have access to this generic.

Usage

```r
drawGD(gdObject, minBase, maxBase, vpPosition, ...)
```

Arguments

- `gdObject`
- `minBase`
- `maxBase`
- `vpPosition`
- `...` Ignored
Class "ExonArray" representing probe level exon array data from Affymetrix

Description

Represents probe level exon array data from Affymetrix. Makes it possible to visualize alternative splicing as measured by the Affymetrix exon array platform and relate it to known transcript isoforms annotated by Ensembl.

Objects from the Class

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

Slots

intensity: Object of class "matrix", array data matrix containing probes as the rows and samples as the columns

probeStart: Object of class "numeric" vector with the start positions of the probes

probeEnd: Object of class "numeric" vector with the end positions of the probes

probeId: Object of class "character" vector containing the probeset identifiers

nProbes: Object of class "numeric" vector defining how many probes there are for each exon/probeset

size: Object of class "numeric" specifying the size of the ExonArray plot in the final plot

displayProbesets: Object of class "logical" used to indicate if probe set names should be plotted or not

probesetSize: Object of class "numeric" font size of the probeset identifiers to be plotted

color: Object of class "character" vector of colors or one color that will be used to draw the intensity values

mapColor: Object of class "character" specifying the color of the lines that map the probeset positions to the Ensembl gene annotation

lwd: Object of class "numeric" vector of line widths to be used to plot the intensity data.

lty: Object of class "character" vector of line types to be used to plot the intensity data.

Extends

Class "gdObject", directly.

Methods

show signature(object = "ExonArray"): ...

Author(s)

Steffen Durinck

References

http://www.stat.berkeley.edu/~steffen/
See Also

objects to See Also as `gdPlot`

Examples

```r
if(interactive()){
  data("unrData", package="GenomeGraphs")
  library(biomaRt)
  mart = useMart("ensembl", dataset="hsapiens_gene_ensembl")

  title = new("Title", title = "ENSG000000009307", dp = DisplayPars(color = "darkslategray"))
  exonmapcol = rep("khaki", length(unrNProbes))
  exonmapcol[28]="darkred"
  probeSetCol = rep("grey", length(unrNProbes))
  probeSetCol[27:28]="darkslategray"
  probeSetLwd = rep(1, length(unrNProbes))
  probeSetLwd[27:28]=3

  exon = new("ExonArray", intensity = unrData, probeStart = unrPositions[,3], probeEnd=unrPositions[,4])
  exon2 = new("ExonArray", intensity = unrData, probeStart = unrPositions[,3], probeEnd=unrPositions[,4],
               mapColor = exonmapcol, probeSetColor = probeSetCol, probeSetLwd = probeSetLwd, displayProbesets=FALSE)
  affyModel = new("GeneModel", exonStart = unrPositions[,3], exonEnd = unrPositions[,4])
  gene = new("Gene", id = "ENSG000000009307", biomart = mart)
  transcript = new("Transcript", id = "ENSG000000009307", biomart = mart)
  legend = new("Legend", legend = c("affyModel","gene"), dp = DisplayPars(color= c("darkgreen","orange")))

  gdPlot(list(title,exonarray1 = exon2,exonarray2= exon, AffymetrixModel= affyModel, gene, legend),
         minBase = min(exon@probeStart), maxBase=max(exon@probeEnd))
}
```

---

**exonProbePos**

*Contains dummy exon probe positions*

### Description

Contains dummy exon probe positions

### Examples

```r
#
```

---

**gdObject-class**

*Class "gdObject" is the parent class of all of the objects in the system.*

### Description

The gdObject is the superclass of all the classes in the system and provides some basic functionality for displaying and managing graphical parameters.

### Objects from the Class

Objects can be created by calls of the form `new("gdObject", ...)` Generally, this class is meant to be subclassed and not created directly.
Slots

dp: Object of class "DisplayPars"

Methods

ggetCex signature(obj = "gdObject"): ...
ggetColor signature(obj = "gdObject"): ...
ggetLty signature(obj = "gdObject"): ...
ggetLwd signature(obj = "gdObject"): ...
ggetPar signature(obj = "gdObject"): ...
ggetPch signature(obj = "gdObject"): ...
ggetPointSize signature(obj = "gdObject"): ...
ggetSize signature(obj = "gdObject"): ...
ginitialize signature(.Object = "gdObject"): ...
gsetPar signature(obj = "gdObject"): ...
gshowDisplayOptions signature(obj = "gdObject"): ...
gshowDisplayOptions signature(obj = "character"): ...

Author(s)

James Bullard

Examples

showClass("gdObject")

gdPlot

gdPlot is the main plotting function of the GenomeGraphs package

Description

gdPlot is the main plotting function of the GenomeGraphs package. A collection of objects are given as a list and these will then be plotted in the order given.

Usage

gdPlot(gdObjects, minBase = NA, maxBase = NA, overlays = NULL,
        labelColor = "black", labelCex = 1, labelRot = 90)

Arguments

gdObjects: This is either a list of gdObjects which will be plotted from top to bottom or a single gdObjects to be plotted.

minBase: minBase defines the minimum base that will be plotted, if ommitted a minimum is determined from the objects in gdObjects if possible.

maxBase: maxBase defines the maximum base that will be plotted, if ommitted a minimum is determined from the objects in gdObjects if possible.
Overview overlays defines a set of regions to overlay on the plot. This argument is either a list or a single Overlay object.

**labelColor** Draw the labels with the given colors.

**labelCex** Character expansion factor.

**labelRot** Rotate the track labels labelRot degrees.

**Author(s)**

Steffen Durinck and James Bullard

**References**

http://www.stat.berkeley.edu/~steffen/

**Examples**

data("exampleData", package="GenomeGraphs")

minbase = min(probestart)
maxbase = max(probestart)

mart = useMart("ensembl", dataset="hsapiens_gene_ensembl")

genesplus = new("GeneRegion", start = minbase, end = maxbase, strand = "+", chromosome = "3")
genesmin = new("GeneRegion", start = minbase, end = maxbase, strand = "-", chromosome = "3")

seg <- new("Segmentation", segments = segments, 
          segmentStart = segStart, segmentEnd = segEnd, dp = DisplayPars(color = "dodgerblue2", lwd=2,lty = "dashed"))

cop <- new("GenericArray", intensity = cn, probeStart = probestart, 
           segmentation = seg, dp = DisplayPars(size=3, color = "seagreen", type="dot"))

ideog = new("Ideogram", chromosome = "3")

expres = new("GenericArray", intensity = intensity, probeStart = exonProbePos, dp = DisplayPars(
           genomeAxis = new("GenomeAxis", add53 = TRUE, add35=TRUE)

gdPlot(list(ideog,expres,cop,genesplus,genomeAxis,genesmin), minBase = minbase, maxBase = maxbase)

---

**geneBiomart**

*AnnotationTrack objects from biomaRt*

**Description**

Convenience function to construct an AnnotationTrack object from biomaRt.

**Usage**

geneBiomart(id, biomart, type = "ensembl_gene_id", dp = NULL)
Gene-class

Arguments

- **id**: ~Describe id here~~
- **biomart**: ~Describe biomart here~~
- **type**: ~Describe type here~~
- **dp**: ~Describe dp here~~

Value

An AnnotationTrack object

Author(s)

James Bullard

---

**Gene-class**

Class "Gene" represents the Ensembl Gene level annotation

Description

Class "Gene" represents the Ensembl Gene level annotation. Upon creation of an object of this class, intron and exon boundaries are retrieved from Ensembl

Objects from the Class

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

Slots

- **id**: Object of class "character", representing a unique identifier for the gene or a vector of identifiers for genes that are located near each other (or at least on the same chromosome)
- **type**: Object of class "character", representing the type of identifier used, e.g. hgnc_symbol, entrezgene and ensembl_gene_id, check the listFilters function of the biomaRt package for more identifier options
- **size**: Object of class "numeric", specifies the size of the plot
- **color**: Object of class "character", specifies the color of the exons
- **biomart**: Object of class "Mart", contains the link to the Ensembl database and should be created using the useMart function from the biomaRt package
- **ens**: Object of class "data.frame", contains the output from the Ensembl query, users don’t need to give a value to this

Methods

- **initialize** signature(.Object = "Gene"): ...
- **drawGD** signature(.Object = "Gene"): ...
- **show** signature(object = "Gene"): ...

Author(s)

Jim Bullard and Steffen Durinck
GeneModel-class

Class "GeneModel", represents a custom gene model

Description

This class represents a custom gene model defined by exon boundaries. An example of this class could be an Affymetrix gene model used to create the Affy Exon array

Objects from the Class

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

Slots

- **exonStart**: Object of class "numeric", vector containing the start positions of the exons that are to be drawn
- **exonEnd**: Object of class "numeric", vector containing the end positions of the exons that are to be drawn
- **chromosome**: Object of class "numeric", chromosome name
- **dp**: Object of class "DisplayPars", color of the exons and size of the exon model in the final plot

Methods

No methods defined with class "GeneModel" in the signature.

Author(s)

Steffen Durinck

References

http://www.stat.berkeley.edu/~steffen/

See Also

objects to See Also as `gdPlot`
Examples

data("unrData", package="GenomeGraphs")
affyModel = new("GeneModel", exonStart = unrPositions[,3], exonEnd = unrPositions[,4])
gdPlot(list(affyModel), minBase = min(unrPositions[,3]), maxBase=max(unrPositions[,4]))

---

geneRegionBiomart  **Construct an AnnotationTrack object from biomaRt.**

Description

This function constructs an AnnotationTrack object from Biomart. It is a convenience function.

Usage

geneRegionBiomart(chr, start, end, strand, biomart, dp = NULL, chrFunction = function(x) x, strandFunction = function(x) x)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>chr</td>
<td>An integer</td>
</tr>
<tr>
<td>start</td>
<td>The start location</td>
</tr>
<tr>
<td>end</td>
<td>The end location</td>
</tr>
<tr>
<td>strand</td>
<td>An integer -1, 0, 1</td>
</tr>
<tr>
<td>biomart</td>
<td>A mart</td>
</tr>
<tr>
<td>dp</td>
<td>A DisplayPars object</td>
</tr>
<tr>
<td>chrFunction</td>
<td>A function which takes chr and converts it into the correct representation for biomaRt. For instance yeast likes to have chromosomes as roman numerals so you can use as.roman here.</td>
</tr>
<tr>
<td>strandFunction</td>
<td>Analagous to chrFunction, but for strand. The internal representation of strand is -1,0,1 whereas biomaRt has different species dependent representations.</td>
</tr>
</tbody>
</table>

Value

An AnnotationTrack object.

Author(s)

James Bullard
GeneRegion-class  
Class "GeneRegion", representing gene structures in a defined genomic region

Description
Given a start and end position and a chromosome name, all gene structures in this region will be retrieved from Ensembl upon creation of the object.

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

Slots
- **start**: Object of class "numeric", start position
- **end**: Object of class "numeric", end position
- **chromosome**: Object of class "character", chromosome name
- **strand**: Object of class "character", represents the strand from which the gene structures should be retrieved. Value is either + or -
- **size**: Object of class "numeric", represents the size of the GeneRegion in the final plot
- **color**: Object of class "character", represents the color to be used to plot the exons
- **biomart**: Object of class "Mart", containing the link to the Ensembl database. This should be created by the `useMart` function from the `biomaRt` package
- **ens**: Object of class "data.frame", output of the `biomaRt` query, should not be used by users

Methods
- **drawGD** signature(.Object = "GeneRegion"):...
- **initialize** signature(.Object = "GeneRegion"):...
- **show** signature(object = "GeneRegion"):...

Author(s)
Steffen Durinck

References
http://www.stat.berkeley.edu/~steffen/

See Also
objects to See Also as `gdPlot`
Examples

```r
if(interactive()){
  mart = useMart("ensembl", dataset="hsapiens_gene_ensembl")
  plusStrand = new("GeneRegion", chromosome = "17", start = 30450000, end = 30550000, strand = "+")
  genomeAxis = new("GenomeAxis", add53=TRUE)
  gdPlot(list(genomeAxis, plusStrand), minBase = 30450000, maxBase = 30550000)
}
```

GenericArray-class

Class "GenericArray", representing array data

Description

The Generic Array class is a class that can be used to create plots from array data such as microarrays and arrayCGH platforms. It can represent the data as line plots or dot plots and segments can be included as well.

Objects from the Class

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

Slots

- **intensity**: Object of class "matrix", matrix containing the intensities of expression or cgh data. Rows should be probes, columns samples
- **probeStart**: Object of class "numeric", start position of the probes
- **probeEnd**: Object of class "numeric", end position of the probes if available
- **type**: Object of class "character", has two values: line and point. If line is selected then a line will be plotted for each sample. If point is selected, points will be plotted.
- **segments**: Object of class "list", if segments are available they have to be represented as a list, each position in the list should correspond to each sample as defined by the columns of the intensity matrix
- **segmentStart**: Object of class "list", containing the start positions of the segments
- **segmentEnd**: Object of class "list", containing the end positions of the segments
- **color**: Object of class "character", represents the color to be used to plot the intensity matrix, can be a vector of colors for multiple samples
- **lty**: Object of class "character", if line is selected as type, lty specifies which line type should be used. Can be a vector of line types for multiple samples
- **pch**: Object of class "numeric", if point is selected as type then pch represents which symbol should be used to plot the points
- **pointSize**: Object of class "numeric", specifies the point size if point is selected as type
- **lwd**: Object of class "numeric", specifies the line width if line is selected as type. Can be a vector of line widths if multiple samples are present
- **size**: Object of class "numeric" representing the size of the Array plot in the final plot
- **segmentColor**: Object of class "character", If segments are to be plotted, this will define the color of the segment lines
Methods

```r
show signature(object = "GenericArray");
```

Author(s)

Steffen Durinck

References

http://www.stat.berkeley.edu/~steffen/

See Also

objects to See Also as `gdPlot`

Examples

```r
if(interactive()){  
data("exampleData", package="GenomeGraphs")  

minbase <- 180292097  
maxbase <- 180492096  
ideog <- new("Ideogram", chromosome = "3")  
expres <- new("GenericArray", intensity = intensity, probeStart = exonProbePos,  
    dp = DisplayPars(color="darkred", type="point"))  
gdPlot(list(ideog, expres), minBase = minbase, maxBase = maxbase)
}
```

GenomeAxis-class  
Class "GenomeAxis", representing a genomic coordinate axis

Description

Represents a genomic coordinate axis

Objects from the Class

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

Slots

- **add53**: Object of class "logical", indicating if 5 to 3 prime direction needs to be plotted
- **add35**: Object of class "logical", indicating if 3 to 5 prime direction needs to be plotted
- **dp**: Object of class "DisplayPars", containing the display parameters such as size of the plot and color
- **littleTicks**: Object of class "logical", indicating if the genome axis should be dense for improved locating of regions of interest.

Methods

No methods defined with class "GenomeAxis" in the signature.
getPar

Author(s)

Steffen Durinck

References

http://www.stat.berkeley.edu/~steffen/

See Also

objects to See Also as gdPlot

Examples

if(interactive()){
mart = useMart("ensembl", dataset="hsapiens_gene_ensembl")
genomeAxis = new("GenomeAxis", add53=TRUE)
plusStrand = new("GeneRegion", chromosome = "17", start = 30450000, end = 30550000, strand = +)
gdPlot(list(genomeAxis, plusStrand), minBase = 30450000, maxBase = 30550000)
}

getPar

Retrieves a display parameter from an object.

Description

Retrieves a display parameter from an object.

Usage

getPar(obj, name, ...)

Arguments

obj A gdObject or DisplayPars object.
name Name of parameter to return.
... Ignored

Examples

a <- new("GenomeAxis")
getPar(a, "size")
HighlightRegion-class

Class "HighlightRegion" is used to highlight vertical blocks of genomic regions.

Description

HighlightRegion is used to highlight a genomic region of interest. The class offers the ability to highlight or block out regions of interest.

Objects from the Class

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

Slots

- **start**: Object of class "numeric" genomic start position.
- **end**: Object of class "numeric" genomic end position.
- **region**: Object of class "numericOrNull" start and end number of the tracks to be covered by the region. These start from the first track (top = 1) to the last track: `length(gdObjects)` in the call to `gdObject`
- **coords**: Object of class "character" can be either "genomic" or "absolute", if the coordinates are "absolute" then one can plot things using the coordinate space defined by: lower-left (0,0) upper-right (1,1). In this case, start = x0, end = x1 and then region = (y0, y1). See the examples for more details.
- **dp**: Object of class "DisplayPars" specifys the various display parameters.

Extends

Class "gdObject", directly.

Methods

No methods defined with class "HighlightRegion" in the signature.

Author(s)

James Bullard

Examples

```r
if (interactive()) {
  data("exampleData", package="GenomeGraphs")

  ga <- new("GenomeAxis")
  grF <- new("GeneRegion", start = 10000, end = 20000, chromosome = "I", strand = "+", biomart = yeastMart)
  grR <- new("GeneRegion", start = 10000, end = 20000, chromosome = "I", strand = "-", biomart = yeastMart)
  bt <- new("BaseTrack", base = yeastCons1[,1], value = yeastCons1[,2])
  hr1 <- new("HighlightRegion", start = 11000, end = 13000, 
              dp = DisplayPars(alpha = 1, color = "red", lty = "dashed", lwd = 3))
  hr2 <- new("HighlightRegion", start = 15900, end = 16500)
}
Ideogram-class

Class "Ideogram", represent an Ideogram

Description

An ideogram is a representation of a chromosome containing the banding pattern. Note that currently ideograms are only available for hsapiens.

Objects from the Class

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

Slots

- **chromosome**: Object of class "character", representing the chromosome that needs to be drawn. E.g. 3 if chromosome 3 needs to be drawn or Y for Y chromosome.
- **dp**: Object of class "DisplayPars", can be used to specify the size (default 1) of the ideogram in the final plot and to specify the highlighting color

Methods

No methods defined with class "Ideogram" in the signature.

Author(s)

Steffen Durinck

References

http://www.stat.berkeley.edu/~steffen/

See Also

- objects to See Also as `gdPlot`

Examples

```r
if(interactive()){
  data("exampleData", package="GenomeGraphs")

  minbase <- 180292097
  maxbase <- 180492096
  ideog <- new("Ideogram", chromosome = "3")
  expres <- new("GenericArray", intensity = intensity, probeStart = exonProbePos,
                dp = DisplayPars(color="darkred", type="point"))
  gdPlot(list(ideog, expres), minBase = minbase, maxBase = maxbase)
}
```
ideogramTab  
Contains info to plot ideograms

Description
Contains info to plot ideograms

Format
The format is: chr "ideogramTab"

Source
NCBI

Examples

data(ideogramTab)
## maybe str(ideogramTab) ; plot(ideogramTab) ...

intensity  
Contains dummy intensity data

Description
Contains dummy intensity data

Examples
#

Legend-class  
Class "Legend", represents a legend to add to a plot

Description
This class represents a legend

Objects from the Class
Objects can be created by calls of the form new("Legend", ...).

Slots

legend: Object of class "character", vector with names of the items in the legend
dp: Object of class "DisplayPars" size of the legend (size), the size of the font (cex) and the colors (color) of the legend
**makeAnnotationTrack**

**Methods**

No methods defined with class "Legend" in the signature.

**Author(s)**

Steffen Durinck

**References**

http://www.stat.berkeley.edu/~steffen/

**See Also**

objects to See Also as gdPlot

**Examples**

```
showClass("Legend")
```

```
makeAnnotationTrack regions = NULL, chr = NULL, strand = NULL, start = NULL, end = NULL, feature = NULL, group = NULL, ID = NULL, dp = NULL
```

**Description**

Convenience function for constructing objects of class AnnotationTrack.

**Usage**

```
makeAnnotationTrack(regions = NULL, chr = NULL, strand = NULL, start = NULL, end = NULL, feature = NULL, group = NULL, ID = NULL, dp = NULL)
```

**Arguments**

- `regions` A dataframe with columns start, end, feature, group, ID. start and end delineate the boundaries of the boxes feature can be used to color the boxes. Group denotes linking so generally exons from a gene form a group. Finally, ID can be used to plot names on boxes.
- `chr` The chromosome of the regions (can be ignored)
- `strand` The strand of the regions (can be ignored)
- `start` If regions is missing then we construct a dataframe from the remaining parameters.
- `end` Construct regions with this vector
- `feature` Construct regions with this feature vector or scalar
- `group` Defines a grouping
- `ID` Defines an ID for each annotation bit
- `dp` DisplayPars, in this case we can create a mapping between feature and color. So let's say in the feature column you have: gene, transcript, gene, then in the dp you can say gene = 'blue' and transcript = 'green'
makeBaseTrack

Value

Returns an object of class AnnotationTrack

Examples

```r
a <- makeAnnotationTrack(start = c(10, 15, 25), end = c(12, 19, 31),
                         group = c(1,1,2), feature = c("gene", "gene", "tf"),
                         ID = paste("id", 1:3, sep = ""), dp = DisplayPars(gene = 'blue'))
gdPlot(a, minBase = 0, maxBase = 40)
```

makeBaseTrack

Creates an object of class BaseTrack

Description

Creates an object of class BaseTrack, which can represent many datasets containing a base given by a vector of positions and a corresponding vector with values for these base positions

Usage

```r
makeBaseTrack(base, value, strand, segmentation, dp = NULL)
```

Arguments

- **base**: Numeric vector of base positions
- **value**: Numeric vector with values for these base positions
- **strand**: Character either + or - representing the strand
- **segmentation**: Object of class Segmentation, used when segments are needed to be drawn
- **dp**: Object of class DisplayPars representing the display parameters of the plot

Value

Object of class BaseTrack

Author(s)

Jim Bullard and Steffen Durinck

References

~put references to the literature/web site here ~

See Also

DisplayPars, gdPlot
Examples

```r
##---- Should be DIRECTLY executable !! ----
##-- ==> Define data, use random, 
##-- or do help(data=index) for the standard data sets.

## The function is currently defined as
function (base, value, strand, segmentation, dp = NULL)
{
  pt <- getClass("BaseTrack")@prototype
  if (is.null(dp))
    dp <- pt@dp
  if (missing(strand))
    strand <- pt@strand
  if (missing(segmentation))
    segmentation <- pt@segmentation
  if (missing(base))
    stop("Need base argument to know the base positions to plot the data on the genome")
  if (missing(value))
    stop("Need value argument")
  new("BaseTrack", base = base, value = value, strand = strand, 
       dp = dp, segmentation = segmentation)
}
```

**makeExonArray**

Creates and object of class ExonArray

Description

Creates an object of class ExonArray, representing exon array microarray data

Usage

```r
makeExonArray(intensity, probeStart, probeEnd, probeId, nProbes, displayProbesets)
```

Arguments

- **intensity**: Matrix of intensities, probes in the rows, samples in the columns
- **probeStart**: Vector of probe start positions
- **probeEnd**: Vector of probe end positions (optional)
- **probeId**: Character vector containing the probe identifiers
- **nProbes**: Vector indicating how many probes are in each probeset
- **displayProbesets**: Logical indicating if the probeset identifier should be displayed or not
- **dp**: Object of class DisplayPars to set the display parameters

Value

Object of ExonArray class

Author(s)

Steffen Durinck and Jim Bullard
makeGeneModel

Creates an object of class GeneModel

Description

Creates an object of class GeneModel representing a custom annotation or gene model.

Usage

makeGeneModel(start, end, chromosome, dp = NULL)

Arguments

start Vector of start positions for exons
end Vector of end positions for exons
chromosome chromosome name
dp Display parameters represented as an object of class DisplayPars
Value

Object of class GeneModel

Author(s)

Steffen Durinck and Jim Bullard

References

~put references to the literature/web site here ~

See Also

DisplayPars

Examples

```r
##---- Should be DIRECTLY executable !! ----
##-- ==> Define data, use random,
##-- or do help(data=index) for the standard data sets.

## The function is currently defined as
## function (start, end, chromosome, dp = NULL)
## {
##   if (is.null(dp))
##     dp <- getClass("GeneModel")@prototype@dp
##   new("GeneModel", exonStart = start, exonEnd = end, dp = dp)
## }
```

makeGene

Creates an object of class Gene

Description

Creates an object of class Gene. This represents a gene structure as annotated in Ensembl.

Usage

`makeGene(id, type, biomart, dp = NULL)`

Arguments

- `id`: An identifier used to specify of which gene the intron-exon structure should be retrieved.
- `type`: The type of identifiers used, examples are `ensembl_gene_id`, `hgnc_symbol`, `entrezgene`. See listAttributes function of the `biomaRt` package for more info.
- `biomart`: Mart object, created by the `useMart` function of `biomaRt`.
- `dp`: object of class `DisplayPars`, determines the display of features on the plot.

Value

An object of class Gene
Author(s)

Steffen Durinck and Jim Bullard

References

~put references to the literature/web site here ~

See Also

gdPlot

Examples

```r
##---- Should be DIRECTLY executable !! ----
##-- ==> Define data, use random,
##-- or do help(data=index) for the standard data sets.

## The function is currently defined as
## function (id, type, biomart, dp = NULL)
## {
##  if (missing(id))
##    stop("Need to specify a gene identifier for creating a Gene")
##  pt <- getClass("Gene")@prototype
##  if (is.null(dp))
##    dp <- pt@dp
##  if (missing(type))
##    type <- pt@type
##  new("Gene", id = id, type = type, biomart = biomart, dp = dp)
## }
```

---

**makeGeneRegion**

Creates an object of class Gene containing the intron-exon structures of genes

Description

Creates an object of class Gene containing the intron-exon structures of genes. Given a start and end position, strand and chromosome, all the intron-exon structures of all genes laying in this region will be retrieved.

Usage

```r
makeGeneRegion(start, end, chromosome, strand, biomart, dp = NULL)
```

Arguments

- **start**: Start position on chromosome
- **end**: End position on chromosome
- **chromosome**: Chromosome name
- **strand**: Strand either + or -
- **biomart**: Mart object, created by the useMart function of biomaRt
- **dp**: Object of class DisplayPars, determines the display of features on the plot
makeGenericArray

Value

An object of class Gene

Author(s)

Steffen Durinck and Jim Bullard

References

~put references to the literature/web site here ~

See Also

gdPlot

Examples

```r
## Should be DIRECTLY executable !! ----
##-- ==> Define data, use random,  
##-- or do help(data=index) for the standard data sets.
## The function is currently defined as
## function (start, end, chromosome, strand, biomart, dp = NULL) 
{
  if (missing(start))
    stop("Need to specify a start for creating a GeneRegion")
  pt <- getClass("GeneRegion")@prototype
  if (is.null(dp))
    dp <- pt@dp
  if (is.numeric(chromosome))
    chromosome <- as.character(chromosome)
  new("GeneRegion", start = start, end = end, chromosome = chromosome,
       strand = strand, biomart = biomart, dp = dp)
}
```

makeGenericArray  Creates an object of class GenericArray

Description

Creates an object of class Generic Array representing microarray data. This could be gene expression, array CGH, etc.

Usage

```r
makeGenericArray(intensity, probeStart, probeEnd, segmentation, dp = NULL)
```
makeGenomeAxis

Arguments

- **intensity**: Matrix of intensities, probes in the rows, samples in the columns
- **probeStart**: Vector of start positions for the probes
- **probeEnd**: Vector of end positions for probes (optional)
- **segmentation**: Object of class Segmentation, needs to be added if segments should be plotted as well
- **dp**: Object of class DisplayPars which handles the display parameters for plotting

Value

Object of class GenericArray

Author(s)

Jim Bullard and Steffen Durinck

References

BMC bioinformatics 2009

See Also

gdPlot

Examples

```r
function (intensity, probeStart, probeEnd, segmentation, dp = NULL)
{
  pt <- getClass("GenericArray")@prototype
  if (is.null(dp))
    dp <- pt@dp
  if (missing(probeEnd))
    probeEnd <- pt@probeEnd
  if (missing(segmentation))
    segmentation <- pt@segmentation
  if (missing(probeStart))
    stop("Need probeStart argument to know where to plot the data on the genome")
  new("GenericArray", intensity = intensity, probeStart = probeStart,
      probeEnd = probeEnd, dp = dp, segmentation = segmentation)
}
```

makeGenomeAxis Creates an object of class GenomeAxis

Description

Creates an object of class GenomeAxis, representing a genome coordinate axis.

Usage

```r
makeGenomeAxis(add53 = FALSE, add35 = FALSE, littleTicks = FALSE, dp = NULL)
```
makeIdeogram

Arguments

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>add53</td>
<td>Add a 5 to 3 prime label</td>
</tr>
<tr>
<td>add35</td>
<td>Add a 3 to 5 prime label</td>
</tr>
<tr>
<td>littleTicks</td>
<td>Add smaller ticks between larger ticks</td>
</tr>
<tr>
<td>dp</td>
<td>Set the display parameters see DisplayPars</td>
</tr>
</tbody>
</table>

Value

Object of class GenomeAxis

Author(s)

Jim Bullard and Steffen Durinck

References

~put references to the literature/web site here ~

See Also

DisplayPars, gdPlot

Examples

```r
##---- Should be DIRECTLY executable !! ----
##-- ==> Define data, use random,
##-- or do help(data=index) for the standard data sets.

## The function is currently defined as
function (add53 = FALSE, add35 = FALSE, littleTicks = FALSE, dp = NULL)
{
  if (is.null(dp))
    dp <- getClass("GenomeAxis")@prototype@dp
  new("GenomeAxis", add53 = add53, add35 = add35, littleTicks = littleTicks, dp = dp)
}
```

makeIdeogram

*Creates object of class Ideogram*

Description

Creates object of class Ideogram

Usage

```r
makeIdeogram(chromosome, dp = NULL)
```

Arguments

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>chromosome</td>
<td>Chromosome to represent (currently human only)</td>
</tr>
<tr>
<td>dp</td>
<td>Display parameters such as color and size</td>
</tr>
</tbody>
</table>
makeLegend

Creates an object of class Legend

Description

Creates an object of class Legend which can be used to plot a legend

Usage

makeLegend(text, fill, cex)

Arguments

text Vector of characters representing the legend
fill Vector of colors to fill the legend boxes
cex Font size of the legend
Value

Object of class Legend

Author(s)

Jim Bullard and Steffen Durinck

References

~put references to the literature/web site here ~

See Also

See Also as gdPlot

Examples

```r
##---- Should be DIRECTLY executable !! ----
##-- ==> Define data, use random,
##-- or do help(data=index) for the standard data sets.

## The function is currently defined as
function (text, fill, cex)
{
  dp <- getClass("Legend")@prototype@dp
  if (!missing(cex))
    setPar(dp, "cex", cex)
  if (!missing(fill))
    setPar(dp, "color", fill)
  new("Legend", legend = text, dp = dp)
}
```

makeRectangleOverlay

Create a rectangular overlay

Description

Construct rectangular overlays.

Usage

makeRectangleOverlay(start, end, region = NULL, coords = c("genomic", "absolute")

Arguments

- `start`: Start position in coords coordinates
- `end`: End position in coords coordinates
- `region`: Which tracks to span, or the y (vertical range)
- `coords`: Which coordinate system to use, if absolute then the range is from 0,1 and region become the y coordinates
- `dp`: The display parameters
The rectangular overlay can be used to plot overlays in either genomic or absolute coordinates. If coordinates are absolute then the region argument becomes the y arguments.

An object of class RectangleOverlay

Examples

data("exampleData", package = "GenomeGraphs")
cop <- makeGenericArray(intensity = cn, probeStart = probestart,
                        dp = DisplayPars(size=3, color = "seagreen", type="dot"))
gdPlot(list(makeGenomeAxis(), cop), overlays =
       makeRectangleOverlay(start = 180350000, end = 180350000 + 1e5, dp = DisplayPars(alpha = .3)))

makeSegmentation

Create objects of class segmentation

Construct objects of class segmentation

makeSegmentation(start, end, value, dp = NULL)

Either a list or a vector. If it is a list then it is a list of vectors of start position (this is the way it is represented in the segmentation class) If it is a vector it is a vector of start positions.

Same as start, but the corresponding end positions.

The y value of the segmentation, ie. segments(start[i], value[i], end[i], value[i])

The Display parameters.

An object of class Segmentation

Examples

data("exampleData", package="GenomeGraphs")
seg <- makeSegmentation(segStart, segEnd, segments,
                        dp = DisplayPars(color = "black", lwd=2, lty = "solid"))
cop <- makeGenericArray(intensity = cn, probeStart = probestart,
                        segmentation = seg, dp = DisplayPars(size=3, color = "seagreen"),
gdPlot(cop)
makeTextOverlay  

Create objects of class TextOverlay

Description
Create objects of class TextOverlay

Usage

makeTextOverlay(text, xpos, ypos, region = NULL, coords = c("genomic", "absolute"), dp = NULL)

Arguments

text  The text to plot
xpos  The xposition of the text
ypos  The yposition of the text
region  
coords  
dp  The display parameters

Value

Returns class of TextOverlay

Examples

data("exampleData", package="GenomeGraphs")
seg <- makeSegmentation(segStart, segEnd, segments,
  dp = DisplayPars(color = "black", lwd=2,lty = "solid")
) cop <- makeGenericArray(intensity = cn, probeStart = probestart,
  segmentation = seg, dp = DisplayPars(size=3, color = "seagreen",
  gdPlot(cop, overlay = makeTextOverlay("Overlay Text", xpos = .5, ypos = .5, coords = "absolute"))

makeTitle  

Creates an object of class Title

Description
Creates an object of class Title which can be used to add a title to the plot

Usage

makeTitle(text, cex, color, size)

Arguments

text  The text that will make up the title

  cex  Font size of the title

color  Font color of the title

  size  Size of the viewport in which the title resides
**makeTranscript**

*Creates an object of class Transcript*

**Description**

Creates an object of class Transcript. This represents all known transcript structures in Ensembl.

**Usage**

```r
makeTranscript(id, type, biomart, dp = NULL)
```

**Arguments**

- **id**
  - An identifier used to specify of which gene/transcript the transcript structures should be retrieved

- **type**
  - The type of identifiers used, examples are ensembl_gene_id, hgnc_symbol, entrezgene. See listAttributes function of thebiomaRt package for more info

- **biomart**
  - Mart object, created by the useMart function of biomaRt

- **dp**
  - object of class DisplayPars, determines the display of features on the plot
Value
An object of class Transcript

Author(s)
Steffen Durinck and Jim Bullard

References
~put references to the literature/web site here ~

See Also
gdPlot

Examples
```r
## Should be DIRECTLY executable !! ----
##-- ==> Define data, use random,  
##-- or do help(data=index) for the standard data sets.

## The function is currently defined as
## function (id, type, biomart, dp = NULL)
## {
##   if (missing(id))
##     stop("Need to specify a gene identifier for creating a Transcript")
##   pt <- getClass("Transcript")@prototype
##   if (is.null(dp))
##     dp <- pt@dp
##   if (missing(type))
##     type = pt@type
##   new("Transcript", id = id, type = type, biomart = biomart,
##       dp = dp)
## }
```

---

**MappedRead-class**  
**Represents mapped reads**

### Description
Represents mapped reads

### Slots
- **start**: Object of class "numeric", containing start position of the reads
- **end**: Object of class "numeric", containing end position of the reads
- **strand**: Object of class "numeric", containing strand to which the reads map
- **chromosome**: Object of class "numeric", containing chromosome to which the reads map

### Methods
- **show** signature(object = "MappedRead"): ...
Author(s)
Steffen Durinck

References
http://www.stat.berkeley.edu/~steffen/

Examples
```r
## maybe str(MappedRead) ; plot(MappedRead) ...
```

---

Overlay-class  

Class "Overlay"

---

Description
Superclass of overlay objects.

Objects from the Class
Objects from this class are generally not created.

Slots
- `dp`: Object of class "DisplayPars"

Extends
Class "gdObject", directly.

Methods
No methods defined with class "Overlay" in the signature.

Examples
```r
showClass("Overlay")
```

---

probestart  
Contains dummy expression array probe start positions

---

Description
Contains dummy expression array probe start positions

Examples
```r
#
```
RectangleOverlay-class

Class "RectangleOverlay"

Description
Rectangular Overlay

Objects from the Class
Objects can be created by calls of the form makeRectangleOverlay (makeRectangleOverlay).

Slots
start: Object of class "numeric" ~
end: Object of class "numeric" ~
region: Object of class "numericOrNull" ~
coords: Object of class "character" ~
dp: Object of class "DisplayPars" ~

Extends
Class "Overlay", directly. Class "gdObject", by class "Overlay", distance 2.

Methods
drawOverlay signature(obj = "RectangleOverlay"):

Examples
showClass("RectangleOverlay")

segEnd
Contains dummy copy number segmentation end positions

Description
Contains dummy copy number segmentation end positions

Examples
#

Segmentable-class  

Class "Segmentable"

Description

The Segmentable class is used as an interface to determine whether or not a segmentation should be plotted.

Objects from the Class

This object is mostly intended to be extended.

Slots

- **segmentation**: Object of class "SegmentationOrNULL"

Methods

- **getSegmentation** signature(obj = "Segmentable"): returns the segmentation, this is essentially the method which is implemented by subclasses.

See Also

- Segmentation

Examples

- `showClass("Segmentable")`

Segmentation-class  

Class "Segmentation" is used to specify segmentations to any class that extends Segmentable (GenericArray, BaseTrack)

Description

A Segmentation object provides line segments to various gdObjects

Objects from the Class

Objects can be created by calls of the form `new("Segmentation", segments = list(1), segmentStart = list(1000), segmentEnd = list(1010)).`

Slots

- **segments**: Object of class "list"
- **segmentStart**: Object of class "list"
- **segmentEnd**: Object of class "list"
- **dp**: Object of class "DisplayPars"
Extends

Class "gdObject", directly.

Methods

getSegmentEnd signature(obj = "Segmentation"):

getSegmentStart signature(obj = "Segmentation"):

getSegments signature(obj = "Segmentation"):

Author(s)

James Bullard

Examples

showClass("Segmentation")

segments

Contains dummy copy number segment data

Description

Contains dummy copy number segment data

Examples

#

segStart

Contains dummy copy number segmentation start positions data

Description

Contains dummy copy number segmentation start positions

Examples

#
**seqDataEx**

This is an example data set from chromosome 4 of yeast from various publicly available datasets.

**Description**

This was a small dataset constructed from publicly available datasets. Please see references for details.

**Usage**

```r
data(seqDataEx)
```

**Format**

```r
data("seqDataEx", package = "GenomeGraphs") names(seqDataEx)
```

**References**

Ugrappa Nagalakshmi et. al. The transcriptional landscape of the yeast genome defined by RNA sequencing. Science, 2008


Adam Siepel, et. al. Evolutionarily conserved elements in vertebrate, insect, worm, and yeast genomes. Genome Res, 2005

**Examples**

```r
data(seqDataEx)
```

---

**setPar**

*Sets a display parameter*

**Description**

Sets a display parameter

**Usage**

```r
setPar(obj, name, val, ...)
```

**Arguments**

- `obj` An object, usually a gdObject.
- `name` Name of display parameter to set.
- `val` Value of display parameter.
- `...` Ignored
Examples

```r
a <- new("GenomeAxis")
setPar(a, "size", 100)
gdPlot(a, minBase = 10, maxBase = 10000)
```

showDisplayOptions  Print standard display options. DisplayPars for an object or a class

Description

Prints the available display options for a class or name of a class.

Usage

```r
showDisplayOptions(obj, ...)
```

Arguments

- `obj` Either an object of subclass gdObject or a character naming a class
- `...` ~Describe ... here~~

Value

Returns a DisplayPars object which is generally printed to the screen.

Examples

```r
showDisplayOptions("GenericArray")
```

TextOverlay-class  Class "TextOverlay"

Description

Textual overlay classes

Objects from the Class

Objects can be created by calls of the form `makeTextOverlay`

Slots

- `text`: Object of class "character"
- `xpos`: Object of class "numeric"
- `ypos`: Object of class "numeric"
- `region`: Object of class "numericOrNull"
- `coords`: Object of class "character"
- `dp`: Object of class "DisplayPars"
Title-class

Extends

Class "Overlay", directly. Class "gdObject", by class "Overlay", distance 2.

Methods

drawOverlay signature(obj = "TextOverlay"): ...

Examples

showClass("TextOverlay")

Title-class

Class "Title" representing the title of a plot

Description

Represent the title of a plot

Objects from the Class

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

Slots

- **title**: Object of class "character" which will be used as title
- **dp**: Object of class "DisplayPars" specifying the size and color of the title in the final plot

Methods

No methods defined with class "Title" in the signature.

Author(s)

Steffen Durinck

References

http://www.stat.berkeley.edu/~steffen/

See Also

objects to See Also as gdPlot

Examples

showClass("Title")
Transcript-class  

Represent known transcript isoforms as annotated by Ensembl

Description

Represent known transcript isoforms as annotated by Ensembl

Objects from the Class

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

Slots

- **id**: Object of class "character", represents the gene identifier that should be used to retrieve the transcript level annotation
- **type**: Object of class "character", represents the type of identifiers used to specify the gene e.g. hgnc_symbol, entrezgene and ensembl_gene_id
- **size**: Object of class "numeric", represents the size of the plot of this object
- **transcriptSize**: Object of class "numeric", represents the size of the transcripts in the plot
- **numOfTranscripts**: Object of class "numeric", should not be used by users
- **color**: Object of class "character", color of the exons
- **biomart**: Object of class "Mart", containing the links to the Ensembl database. This object should be created with the useMart function of the biomaRt package
- **ens**: Object of class "data.frame", should not be used by the users. Contains the output from the biomaRt query

Methods

- **drawGD** signature(.Object = "Transcript"):
- **initialize** signature(.Object = "Transcript"):
- **show** signature(object = "Transcript"):

Author(s)

Steffen Durinck

References

http://www.stat.berkeley.edu/~steffen/

See Also

objects to See Also as `gdPlot`
TranscriptRegion-class

Class "TranscriptRegion", representing a genomic region with transcripts

Description

Upon creation of this object, transcripts present in a specified region will be retrieved from Ensembl

Objects from the Class

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

Slots

- **start**: Object of class "numeric", the start base of the genomic region
- **end**: Object of class "numeric", the end base of the genomic region
- **chromosome**: Object of class "character", the chromosome
- **size**: Object of class "numeric", the size of the plot for this object
- **biomart**: Object of class "Mart", contains link to Ensembl and should be created using the useMart function of the biomaRt package
- **ens**: Object of class "data.frame", users should not specify this, it contains the output of the query to Ensembl

Methods

- **show** signature(object = "TranscriptRegion"): ...

Author(s)

Steffen Durinck

References

http://www.stat.berkeley.edu/~steffen/

See Also

objects to See Also as gdPlot

Examples

showClass("TranscriptRegion")
unrData  
**Contains exon array data**

**Description**
Contains exon array data from the publically available dataset on human tissue panels, given by Affymetrix. The data was contributed to the package by Elizabeth Purdom.

**Examples**
```
#
```

unrNProbes  
**Contains exon array data**

**Description**
Contains the number of probes per exon array probeset id from the publically available dataset on human tissue panels, given by Affymetrix. The data was contributed to the package by Elizabeth Purdom.

**Examples**
```
#
```

unrPositions  
**Contains probe start and end positions of exon array probes**

**Description**
Contains probe start and end positions from the publically available dataset on human tissue panels, given by Affymetrix. The data was contributed to the package by Elizabeth Purdom.

**Examples**
```
#
```

yeastCons1  
**Contains dummy yeast conservation data**

**Description**
Contains dummy yeast base conservation data.

**Examples**
```
#
```
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