

Package ‘Pviz’

February 21, 2025

Type Package

Title Peptide Annotation and Data Visualization using Gviz

Version 1.41.0

Author Renan Sauteraud, Mike Jiang, Raphael Gottardo

Maintainer Renan Sauteraud <rsautera@fhcrc.org>

Description Pviz adapts the Gviz package for protein sequences and data.

License Artistic-2.0

Depends R(>= 3.0.0), Gviz(>= 1.7.10)

Imports biovizBase, Biostrings, GenomicRanges, IRanges, data.table,
methods

Suggests knitr, pepDat

biocViews Visualization, Proteomics, Microarray

VignetteBuilder knitr

git_url <https://git.bioconductor.org/packages/Pviz>

git_branch devel

git_last_commit 048901b

git_last_commit_date 2024-10-29

Repository Bioconductor 3.21

Date/Publication 2025-02-20

Contents

ATrack	2
CladeTrack	3
DTrack	4
plot_clade	4
plot_inter	5
ProbeTrack	6
ProteinAxisTrack	7
ProteinSequenceTrack	8

Index	10
--------------	-----------

ATrack

ATrack class

Description

This class contains Gviz's AnnotationTrack and adds default values to the genome and chromosome slot

Usage

```
ATrack(range = NULL, start = NULL, end = NULL, width = NULL, group, id,
        stacking = "squish", name = "ATrack", fun, selectFun, ...)
```

Arguments

range, start, end, width, group, id, stacking, name, fun, selectFun, ...
Arguments to be passed to AnnotationTrack.

Author(s)

Renan Sauteraud

See Also

[AnnotationTrack](#), [GdObject](#)

Examples

```
# Object construction
aTrack <- ATrack(start = c(20, 60), end = c(40, 100), name = "random.anno",
id=c("small", "big"))
#Stacking example
a2Track <- ATrack(start = c(20, 30), end = c(40, 100), name = "stacking=dense",
id = c("small", "big"), stacking = "dense", fill=c("black", "orange"))
a3Track <- ATrack(start = c(20, 30), end = c(40, 100), name = "no stacking",
id = c("small", "big"), fill = c("black", "orange"))
#Plotting
plotTracks(trackList = c(aTrack, a2Track, a3Track), showFeatureId = TRUE)
```

`CladeTrack`*CladeTrack*

Description

This track can be used to display the result of pepStat analysis for a single clade. It contains DTrack.

Usage

```
CladeTrack(restab, clade, name = clade, ...)
```

Arguments

<code>restab</code>	A data.frame. The result of a peptide microarray analysis, as returned by pepStat's <code>restab</code> function.
<code>clade</code>	A character. The clade to plot.
<code>name</code>	A character. The name of the track, used in the title panel when plotting. By default, the <code>clade</code> name.
<code>...</code>	Additional argument to be passed to <code>DataTrack</code> . They will be treated as display parameters.

Slots

`clade` A character. The clade to display.

Author(s)

Renan Sauteraud

See Also

DTrack

Examples

```
if(require(pepDat)){
  data(restab)
  ct <- CladeTrack(restab, clade = "M", type = "1", legend = TRUE)
  plotTracks(ct)
}
```

DTrack

DTrack class

Description

This class contains Gviz's DataTrack and adds default values to the genome and chromosome slot

Usage

```
DTrack(range = NULL, start = NULL, end = NULL, width = NULL, data,
        name = "DTrack", ...)
```

Arguments

range, start, end, width, data, name, ...
 Arguments to be passed to DataTrack.

Details

Refer to DataTrack for details regarding the constructor.

Author(s)

Renan Sauteraud

See Also

[DataTrack](#), [GdObject](#)

Examples

```
dTrack <- DTrack(start=seq(1,1000, len=100), width=10, data=matrix(runif(400),
  nrow=4), name="random data")
```

plot_clade

Plot frequency of response for a single clade.

Description

Plot an axis and the frequency of response of a single selected clade.

Usage

```
plot_clade(restab, clade, sequence = NULL, from = 0,
           to = max(restab$position), ...)
```

Arguments

restab	A data.frame. The result of a peptide microarray analysis, as returned by pepStat's restab function.
clade	A character. The clade to plot.
sequence	An optional character or AAString. The sequence of the ProteinSequenceTrack to plot. It should be the sequence of the reference genome used in the peptideSet that generated the restab.
from	A numeric, the start coordinate of the plot.
to	A numeric, the end coordinate of the plot.
...	Additional arguments to be passed to plotTracks.

Author(s)

Renan Sauteraud

See Also

restab, plot_inter, plotTracks

Examples

```
if(require(pepDat)){
  data(restab)
  plot_clade(restab, clade = c("A", "M"))
}
```

plot_inter

Plot frequency of response for each group

Description

Plot an axis and the frequency of response of each group, averaged by peptides at each position.

Usage

```
plot_inter(restab, sequence = NULL, from = 0, to = max(restab$position),
  ...)
```

Arguments

restab	A data.frame. The result of a peptide microarray analysis, as returned by pepStat's restab function.
sequence	A character or an AAString. If not NULL, the sequence of the ProteinSequenceTrack to plot. It should be the sequence of the reference genome used in the peptideSet that generated the restab.
from	A numeric, the start coordinate of the plot.
to	A numeric, the end coordinate of the plot.
...	Additional arguments to be passed to plotTracks.

Author(s)

Renan Sauteraud

See Also

restab, plot_clade, plotTracks

Examples

```
if(require(pepDat)){
  data(restab_aggregate)
  plot_inter(restab_aggregate)
}
```

 ProbeTrack

ProbeTrack

Description

This track can be used to display the frequency of antibody binding for each probe on an arrayas predicted by pepStat's function makeCalls.

Usage

```
ProbeTrack(sequence, intensity, probeStart, restab = NULL, group = NULL,
  name = "ProbeTrack", ...)
```

Arguments

sequence	A character vector. The sequence of peptides to display.
intensity	A numeric vector. The frequency of binding or the baseline corrected intensity for the peptides.
probeStart	A numeric vector. The start position of the peptides.
name	A character. The name of the track used in the title panel when plotting
restab	A data.frame containing all the above parameters, as outputted by pepStat's restab function.
group	A character. The group to display on the ProbeTrak. This is only required when restab is not NULL. See details section for more information.
...	Arguments to be passed to DataTrack.

Details

The vectors for the arguments sequence, freq and probeStart should be of the same length. If restab is provided, the three previous arguments will be ignored and group must be specified. group must be a valid column name in restab, data.frame.

Slots

sequence A character vector. The probes sequence.

probeStart A numeric vector. The start position of the probes.

intensity A numeric vector. The frequency of response of each probe. Or the baseline corrected intensity of the signal.

Author(s)

Renan Sauteraud

See Also

[GdObject](#)

restab

Examples

```
if(require(pepDat)){
  data(restab)
  pt <- ProbeTrack(sequence = restab$peptide,
                   intensity = restab$group2,
                   probeStart = restab$start)
  plotTracks(pt)
  plotTracks(pt, from = 460, to = 560, legend=TRUE)
}
```

ProteinAxisTrack

ProteinAxisTrack

Description

A track to display an axis for protein or peptide sequences

Usage

```
ProteinAxisTrack(range = NULL, name = "Axis", addNC = FALSE, id = NULL,
  ...)
```

Arguments

range, name, id, ...

Arguments to be passed to GenomeAxisTrack.

addNC A logical. If TRUE, display the Amino-terminal and Carboxyl-terminal ends on the axis.

Author(s)

Renan Sauteraud

See Also

[GenomeAxisTrack](#)

Examples

```
# Object construction
paxTrack <- ProteinAxisTrack()
pax2 <- ProteinAxisTrack(addNC=TRUE)
pax3 <- ProteinAxisTrack(littleTicks=TRUE)
# Plotting
plotTracks(c(paxTrack,pax2,pax3), from=1, to=100)
```

ProteinSequenceTrack *ProteinSequenceTrack*

Description

A track to display peptides and protein sequences.

Usage

```
ProteinSequenceTrack(sequence = NULL, name = "Sequence", ...)
```

Arguments

sequence	A character or AAString of length one. The sequence to display.
name	A character. The name of the track used in the title panel when plotting
...	Additional items which will all be interpreted as display parameters.

Author(s)

Renan Sauteraud

See Also

[SequenceTrack](#), [DisplayPars](#)

Examples

```
if(require(pepDat)){
  data(pep_hxb2)
  hxb2_seq <- metadata(pep_hxb2)$sequence
  st<-ProteinSequenceTrack(sequence=hxb2_seq, name="env")

  # Plotting amino acids
  plotTracks(st, to = 20)

  # When the range becomes wider, only coloured squares are displayed
```



```
plotTracks(st, to = 100)  
  
# When overplotting, a single line will mark the ProteinSequenceTrack  
plotTracks(st)  
}
```

Index

AnnotationTrack, [2](#)
ATrack, [2](#)
ATrack-class (ATrack), [2](#)

CladeTrack, [3](#)
CladeTrack-class (CladeTrack), [3](#)

DataTrack, [4](#)
DisplayPars, [8](#)
DTrack, [4](#)
DTrack-class (DTrack), [4](#)

GdObject, [2](#), [4](#), [7](#)
GenomeAxisTrack, [8](#)

plot_clade, [4](#)
plot_inter, [5](#)
plotTracks, [5](#), [6](#)
ProbeTrack, [6](#)
ProbeTrack-class (ProbeTrack), [6](#)
ProteinAxisTrack, [7](#)
ProteinAxisTrack-class
 (ProteinAxisTrack), [7](#)
ProteinSequenceTrack, [8](#)
ProteinSequenceTrack-class
 (ProteinSequenceTrack), [8](#)

SequenceTrack, [8](#)