The Pviz User Guide

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1 Introduction

Pviz is an R package inspired by and depending on Gviz. It introduces new types of track and extends the existing ones in order to deal with amino-acid based data.

This package keeps most of the mechanics of Gviz, notably the use of DisplayParameters and the same plotting function: plotTracks. Therefore, the user is invited to refer to Gviz help pages and vignette for more information and examples.

As with any R package, it should first be loaded in the session

> library(Pviz)

2 Gviz tracks

Pviz extends and uses the most common classes of Gviz to make them easier to use with amino acid data. We removed the requirement for a genome and a chromosome when creating these tracks. Moreover, they support the functions defined in Pviz.

2.1 ATrack

ATrack extends Gviz's AnnotationTrack and behave the same way. However, it does not require to specify a chromosome and a genome. Please refer to Gviz documentation for more details about AnnotationTrack and the available DisplayParameters.

```
> at <- ATrack(start = c(250, 480), end = c(320, 520), name = "Annotations")
> plotTracks(at, from = 1, to = 600)
```





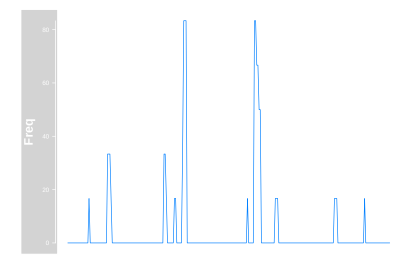


2.2 DTrack

Naturally DTrack extends Givz's DataTrack. Here again, please refer to Gviz documentation for details on how to use DataTrack.

Some example data are available in this package. Expression frequency for peptide in hxb2 envelope.

```
> data(PvizVignette)
> dt <- DTrack(data = freqEx, start = posEx, width = 15, name = "Freq")
> plotTracks(dt, from = 1, to = 850, type = "l")
```



3 Pviz new track types

Pviz introduce some new track types to deal with amino-acid based data. The new tracks look can be modified using the DisplayParameters and will most of the time offer the same options as the ones available for Gviz tracks.

3.1 ProteinAxisTrack

This track acts as a replacement for the <code>GenomeAxisTrack</code>. I comes with the same coloration, transparency and other customization options but loses the DNA representation for a simple segment.

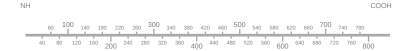
```
> pat <- ProteinAxisTrack()
> plotTracks(pat, from = 1, to = 850)
```



Just like in GenomeAxisTrack, it is possible to use littleTicks to get a more precise scale. Moreover, because Pviz, has been made to deal with

peptides and protein, the option addNc can display indicators for N-term and C-term ends on the axis.

> pat <- ProteinAxisTrack(addNC = TRUE, littleTicks = TRUE)
> plotTracks(pat, from = 1, to = 850)



3.2 SequenceTrack

This new track simply displays a selected sequence. It can takes both AAstring or regular string and thus, can be used for both AA and DNA sequences.

Note that the first amino acid of the sequence should correspond to the first position of any other element you choose to display at the same time.

The previously loaded dataset also contains an AAstring object for the sequence of the envelope of hxb2.

> seqEx

857-letter "AAString" instance seq: MRVKEKYQHLWRWGWRWGTMLLGMLMICSATEKLWV...VAEGTDRVIEVVQGACRAIRHIPRRIRQGLERILL*

> st <- SequenceTrack(sequence = seqEx, name = "env")
> plotTracks(trackList = c(pat, st), from = 1, to = 50)





MRVKEKYQHLWRWGWRWGTMLLGMLMICSATEKLWVTVYYGVPWWKEATT

Note that if the plotting ranges becomes too wide, the letters will stack and it may lead to an unreadable output. The default parameters usually allows to plot up to 100-150 characters depending on your graphic window. The user can modify the character size via the DisplayParameters system when creating the track with the cex argument. Here is an example of bad plotting.

```
> plotTracks(trackList = c(pat, st), cex = 0.5)
```

```
NH COOH
42 71 100 128 157 185 214 242 271 300 228 357 385 414 442 471 500 528 557 585 614 642 571 700 728 757 785 814

57 85 114 142 171 200 228 257 285 314 342 371 400 428 457 465 514 542 571 600 628 657 665 714 742 771 800 828
```

Althought the character expansion has been set to less than 1. The ranges are still too wide for a correct display.

3.3 ProbeTrack

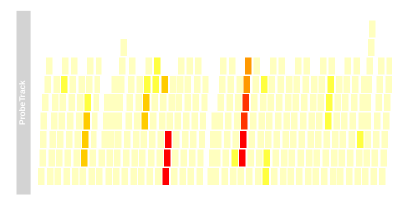
This track is designed to display peptide microarray data. It draws each peptide relative to its position in the sequence and enclose them in rectangles colored depending on their intensity. To create this track, the sequence of the peptides, their intensity or frequency and their starting position have to be passed as arguments.

```
> pt <- ProbeTrack(pepEx, freqEx, posEx)
> plotTracks(pt, from = 460, to = 530)
```



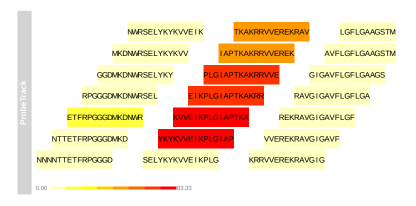
Unlike in SequenceTrack, the size of the characters in each peptide sequence depends on the plotting ranges (the user can still choose to change the size manually) and if the ranges become too wide, the characters will appear as dots or completely disappear instead of stacking on top of each other. While it loses the sequence information, it might be relevant to locate regions where peptides have high intensity/frequency.

> plotTracks(pt)



For a more explicit display, a legend has been implemented for this track and can be called during track creation or in the plotting function. The legend displays the scale of intensities.

> plotTracks(pt, from = 460, to = 530, legend = TRUE)



4 Example of plot

Naturally, the interest of Pviz, just like its parent Gviz is the display of multiple tracks at once. Here is an example of what Pviz can render, using the tracks previously created.

```
> pt <- ProbeTrack(pepEx, freqEx, posEx, cex = 0.8)
> plotTracks(trackList = c(pat, st, at, pt, dt), from = 460, to = 530,
+ type = "1", legend = TRUE)
```

