# The Pviz User Guide

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# 1 Example of plot

```
> library(HIV.db)
> library(Pviz)
> alignObj <- readAlign()</pre>
> refScale <- alignObj[[1]]</pre>
> refSeq <- alignObj[[2]]</pre>
> HIV_db <- loadFeatures(ref = "env")
> envBase <- getFeature(HIV_db)</pre>
> envStart = getHXB2Coordinates(envBase)[1, ][1]
> envEnd = getHXB2Coordinates(envBase)[1, ][2]
> proteins <- getFeature(HIV_db, category = "protein", start = envStart,
      end = envEnd, frame = getFrame(envBase))
> antis <- getEpitope(envBase, name = c("VRC01"))</pre>
> helix <- getChildren(envBase, category = c("helix"))</pre>
> rpext <- ProteinAxisTrack(littleTicks = TRUE)
> rpref <- ProteinAxisTrack(refScale = refScale, adNC = TRUE)
> sTrack <- SequenceTrack(refSeq)
> data(pepMicroarrayEx)
> p1Track <- ProbeTrack(pepMicroarrayEx$probeSeq, pepMicroarrayEx$probeFreq,</p>
      pepMicroarrayEx$probePos, protein = "gp120", name = "sequence(B)")
> a2Track <- ATrack(id = proteins@values@unlistData@listData[["name"]],</pre>
      start = start(proteins), end = end(proteins), genome = "hxb2",
```

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```
name = "Protein", protein = "gp120", fill = "navyblue", size = 1)
> a3Track <- ATrack(id = helix@values@unlistData@listData[["name"]],</pre>
      start = start(helix), end = end(helix), genome = "hxb2",
     name = "Helix", fill = "orange", protein = "gp120")
> a6Track <- ATrack(id = antis@values@unlistData@listData[["name"]],</pre>
      start = start(antis), end = end(antis), genome = "hxb2",
      name = "Epitopes", fill = "gray", protein = "gp120")
> data(pepExprEx)
> library(IRanges)
> d6Track <- DTrack(range = IRanges(start = pepExprEx$dPos, width = 1),</pre>
      groups = rownames(pepExprEx$dExpr), data = pepExprEx$dExpr,
     genome = "hxb2", protein = "gp120", col = c("orange", "gray"),
      cex = 1
> plotTracks(trackList = c(rpext, rpref, sTrack, a2Track, a3Track,
      a6Track, p1Track, d6Track), from = 1, to = 150, type = c("p",
      "smooth"), stacking = "dense", legend = TRUE, showFeatureId = TRUE)
```

## 2 Use extended coordinates system

The extended coordinate system is based on a multiple alignment. It is a scale from 0 to the length of the alignment (i.e. reference sequence + gaps).

The refScale is a scale used to translate coordinates between the extended system and the normal system (which is based uniquely on the reference sequence length).

### 2.1 Data formating

In the following examples, hxb2 is the reference sequence and is aligned with different subtypes.

First, get the refScale:

```
> alignObj <- readAlign(filename = system.file("extdata/alignment.fasta",
+    package = "Pviz"))
> refScale <- alignObj[[1]]
> refSeq <- alignObj[[2]]</pre>
```

Now, the refScale can be used to translate coordinates into the extended system.

Example of loading HIV\_db using extended coordinates.

```
> library(HIV.db)
> HIV_db <- loadFeatures(ref = "env", refScale = refScale)
> envBase <- getFeature(HIV_db)
   The positions in envBase are the ones observed in the alignment.
   To translate all the positions in pep_hxb2, use convertDB().
> data(pep_hxb2)
> ndb <- convertDB(db = pep_hxb2)
\lceil \lceil 1 \rceil \rceil
                              "C"
[1] "CRF01" "B"
                      "D"
                                       "CRF02" "M"
                                                        " A "
[[2]]
[1] "MRVKETQMNWPNLWK----WGTLILGLVIICSASDNLWVTVYYGVPVWRDADTTLFCASDAKAHETEVHNVWATHACVPT
[2] "MRVKGIRKNYQHLWR----WGTMLLGMLMICSAAEQLWVTVYYGVPVWKEATTTLFCASDAKAYDTEVHNVWATHACVPT
[3] "MRVRGIERNYQHLWR----WGTMLLGMLMICSVAEQLWVTVYYGVPVWKEATTTLFCASDAKSYKTEAHNIWATHACVPT
[4] "MRVRGILRNWQQWWI----WGILGFWMLMICNVVGNLWVTVYYGVPVWKEAKTTLFCASDAKAYEKEVHNVWATHACVPT
[5] "MRVMGIQKNYPLLWR----WGTIIFWIMIICNA-ENLWVTVYYGVPVWRDAETTLFCASDAKAYDTEVHNVWATHACVPT
```

[6] "MRVMGIQRNWQHLWR----WGTLILGMLIICSAAENLWVTVYYGVPVWKEATTTLFCASDAKAYETEVHNVWATHACVPT "MRVMGIQRNCQHLWR----WGTMILGMIIICSAAENLWVTVYYGVPVWKDAETTLFCASDAKAYETEVHNVWATHACVPT

```
user system elapsed 2.856 0.000 2.861
```

ndb is pep\_hxb2 with updated ranges, aligned and trimmed columns.

To convert coordinates into the extended system, use coord2ext() with the refScale defined earlier:

```
> start <- coord2ext(c(200, 450), ref = refScale)
> start
[1] 223 474
```

#### 2.2 Plotting with both coordinates system

All objects used to create tracks should be in extended coordinate system.

```
> sTrack <- SequenceTrack(refSeq)
> pax1 <- ProteinAxisTrack()
> pax2 <- ProteinAxisTrack(refScale = refScale, col.gap = "blue")
> proteins <- getFeature(HIV_db, category = "protein", start = envStart,</pre>
```

```
+ end = envEnd, frame = getFrame(envBase))
> aTrack <- ATrack(id = proteins@values@unlistData@listData$name,
+ start = start(proteins), end = end(proteins))
> aTrack <- ATrack(id = helix@values@unlistData@listData$name,
+ start = start(helix), end = end(helix), fill = "red")
> plotTracks(c(pax1, pax2, sTrack, aTrack), from = 1, to = 160)
```

The first axis track displays the extended scale, while the second displays the reference coordinates, it also shows the gaps in the reference sequence with respect of the alignment.