The pepDat User Guide

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1 Introduction

pepDat is an R package that stores sample files, data for vignettes and peptide collections. It is intended to be used in conjunction with other packages for peptide analysis and visualisation: pepStat anf Pviz. As with any R package, it should first be loaded in the session

library(pepDat)

2 Peptide collections

2.1 Information

For each peptide, the following collections display information about the position relative to the reference sequence, the alignment, the trimmed alignment, the zSums for the physicochemical properties as well as the clades where they can be found.

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2.2 Structure

The datasets in this package are GRanges objects. For more information about the class, its accessors and setters, please refer to GenomicRanges documentation.

2.3 pep_hxb2

This collection is based on the alignment of the reference HIV sequence hxb2 and seven subtypes (clades) A, B, C, D, M, CRF01 and CRF02.

2.4 pep_hxb2JPT

This collection adds a few more clades to the previous one: CM244, CON_01_AE, LAI_A04321 and MN_DD328842.

2.5 pep_mac239

This collection is for SIV, with the clades mac239 and E660.

2.6 pep_m239smE543

This collection is for SIV, with the clades mac239 and E543.

3 Loading the collections

The peptide collections can be loaded like any other R dataset.

```
data(pep_hxb2)
head(pep_hxb2)
## GRanges with 6 ranges and 10 metadata columns:
##
                      segnames
                                  ranges strand |
                                                              names
##
                         <Rle> <IRanges>
                                          <Rle>
                                                       <character>
                                  [1, 16]
                                               * | MRVKETQMNWPNLWK
##
     MRVKETQMNWPNLWK
                         gp160
##
     MRVMGIQKNYPLLWR
                         gp160
                                 [1, 16]
                                               * | MRVMGIQKNYPLLWR
                         gp160
     MRVMGIQRNCQHLWR
                                 [1, 16]
                                               * | MRVMGIQRNCQHLWR
##
                                 [1, 16]
##
     MRVKGIRKNYQHLWR
                         gp160
                                               * | MRVKGIRKNYQHLWR
##
     MRVRGILRNWQQWWI
                         gp160
                                  [1, 16]
                                               * | MRVRGILRNWQQWWI
     MRVRGIERNYQHLWR
                                               * | MRVRGIERNYQHLWR
                                  [1, 16]
##
                         gp160
##
                                  aligned
                                                    trimmed
                                                                 seqNb
##
                                 <factor>
                                                   <factor> <integer>
     MRVKETQMNWPNLWK MRVKETQMNWPNL----WK MRVKenWPNL----WK
##
                                                                     1
##
     MRVMGIQKNYPLLWR MRVMGIQKNYPLL----WR MRVMgnYPLL----WR
                                                                     1
     MRVMGIQRNCQHLWR MRVMGIQRNCQHL----WR MRVMgnCQHL----WR
                                                                     1
##
     MRVKGIRKNYQHLWR MRVKGIRKNYQHL----WR MRVKgnYQHL----WR
##
                                                                     1
     MRVRGILRNWQQWWI MRVRGILRNWQQW----WI MRVRgnWQQW----WI
##
                                                                     1
     MRVRGIERNYQHLWR MRVRGIERNYQHL----WR MRVRgnYQHL----WR
##
                                                                     1
```

##		clade	z1	z2	z3	z4
##		<character></character>	<numeric></numeric>	<numeric></numeric>	<numeric></numeric>	<numeric></numeric>
##	MRVKETQMNWPNLWK	CRF01	-3.14	9.87	-8.14	6.72
##	MRVMGIQKNYPLLWR	CRF02	-13.12	3.19	-11.96	7.09
##	MRVMGIQRNCQHLWR	A	1.65	4.17	-11.22	10.31
##	MRVKGIRKNYQHLWR	В	3.95	9.78	-18.51	12.55
##	MRVRGILRNWQQWWI	C	-11.42	10.26	-16.19	10.33
##	MRVRGIERNYQHLWR	D	6.00	10.76	-17.14	8.52
##		z5				
##		<numeric></numeric>				
##	MRVKETQMNWPNLWK	1.36				
##	MRVMGIQKNYPLLWR	0.76				
##	MRVMGIQRNCQHLWR	-2.99				
##	MRVKGIRKNYQHLWR	-0.87				
##	MRVRGILRNWQQWWI	-2.37				
##	MRVRGIERNYQHLWR	-1.91				
##						
##	seqlengths:					
##	gp160					
##	NA					

4 Create collections

While the package comes with datasets for HIV and SIV. It is possible to create new collections for different organisms or proteins using the create_db function in pepStat. Please refer to pepStat's user guide and ?create_db for more information.