Class 11 Lab

Karleen Guerrero (A16791042)

```
library(readr)
pdbdb <- read_csv("~/Downloads/pdb_stats (1).csv")

Rows: 6 Columns: 8
-- Column specification ------
Delimiter: ","
chr (1): Molecular Type
dbl (3): Multiple methods, Neutron, Other
num (4): X-ray, EM, NMR, Total

i Use `spec()` to retrieve the full column specification for this data.
i Specify the column types or set `show_col_types = FALSE` to quiet this message.

Q1: What percentage of structures in the PDB are solved by X-Ray and Electron
Microscopy.</pre>
```

<pre>pdbdb <- read.csv('pdb_stats.csv',</pre>	row.names = 1)
pdbdb	

	X.ray	EM	NMR	Multiple.methods	Neutron	Other
Protein (only)	167,192	15,572	12,529	208	77	32
Protein/Oligosaccharide	9,639	2,635	34	8	2	0
Protein/NA	8,730	4,697	286	7	0	0
Nucleic acid (only)	2,869	137	1,507	14	3	1
Other	170	10	33	0	0	0
Oligosaccharide (only)	11	0	6	1	0	4
	Total					
Protein (only)	195,610					
Protein/Oligosaccharide	12,318					
Protein/NA	13,720					
Nucleic acid (only)	4,531					

```
Other 213
Oligosaccharide (only) 22
```

I need to remove the comma and conert to numeric to do math:

```
as.numeric( sub(",", "", pdbdb$Total) )

[1] 195610 12318 13720 4531 213 22

#as.numeric(pdbdb$Total)
```

I could turn this into a function to fix the whole table or any future table I read like this:

```
x <- pdbdb$Total
as.numeric( sub(",", "", x) )

[1] 195610 12318 13720 4531 213 22

comma2numeric <- function(x) {
   as.numeric( sub(",", "", x) )</pre>
```

Test it

```
comma2numeric(pdbdb$X.ray)
```

[1] 167192 9639 8730 2869 170 11

```
apply(pdbdb, 2, comma2numeric)
```

	X.ray	EM	NMR	Multiple.methods	Neutron	Other	Total
[1,]	167192	15572	12529	208	77	32	195610
[2,]	9639	2635	34	8	2	0	12318
[3,]	8730	4697	286	7	0	0	13720
[4,]	2869	137	1507	14	3	1	4531
[5,]	170	10	33	0	0	0	213
[6,]	11	0	6	1	0	4	22

Or try a different read/import function

```
library(readr)
pdbdb <- read_csv("pdb_stats.csv")</pre>
Rows: 6 Columns: 8
-- Column specification -----
Delimiter: ","
chr (1): Molecular Type
dbl (3): Multiple methods, Neutron, Other
num (4): X-ray, EM, NMR, Total
i Use `spec()` to retrieve the full column specification for this data.
i Specify the column types or set `show_col_types = FALSE` to quiet this message.
sum(pdbdb$Total)
[1] 226414
     Q1: What percentage of structures in the PDB are solved by X-Ray and Electron
     Microscopy.
sum(pdbdb$'X-ray')/sum(pdbdb$Total) * 100
[1] 83.30359
sum(pdbdb$EM)/sum(pdbdb$Total)* 100
[1] 10.18091
```

Q2: What proportion of structures in the PDB are protein?

Mol*

 Mol^* (pronounced "molstar") is a new web-based molecular viewar that we will need to learn the basics of here.

https://molstar.org/viewer/

We will use PDB code: 1HSG

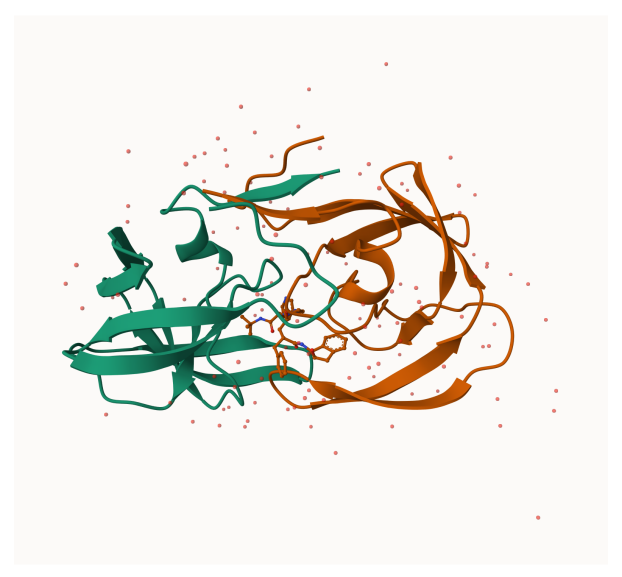


Figure 1: A first image from molstar

Some more custom images:

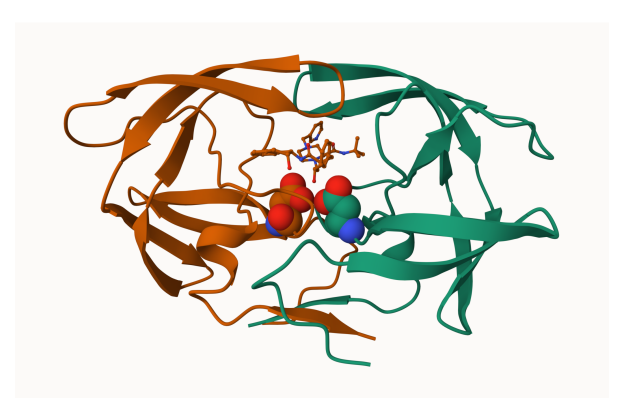


Figure 2: The all important catalytic ASP25 amino acid

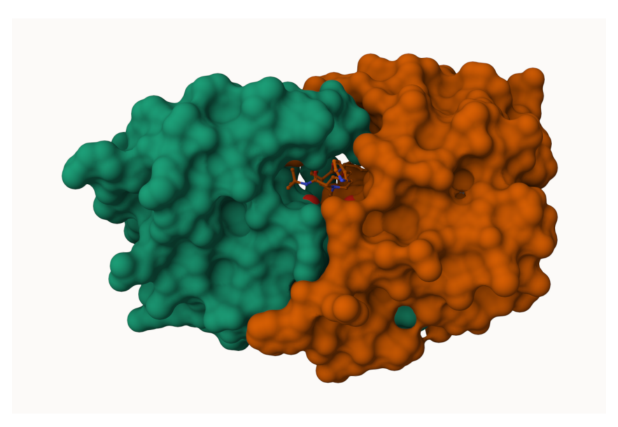


Figure 3: Surafce display showing Merk compound in the peptide binding pocket

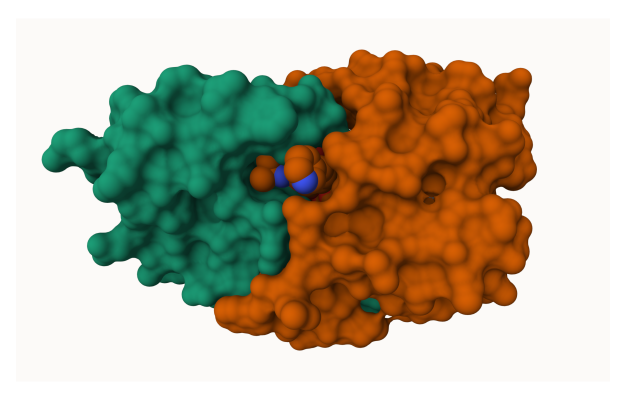


Figure 4: Ligand in pocket

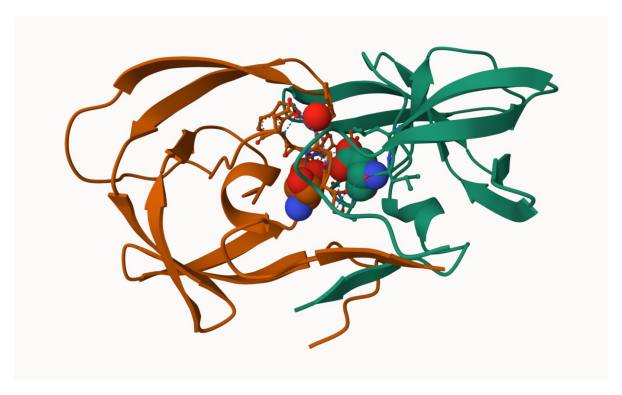


Figure 5: Water molecule

Q3: Type HIV in the PDB website search box on the home page and determine how many HIV-1 protease structures are in the current PDB?

5 structures

The Bio3D pachage

The bio3d package allows us to do all sorts of structural bioinformatics work in R. Let's start with how it can read these PDB files:

```
library(bio3d)

pdb <- read.pdb("1hsg")</pre>
```

Note: Accessing on-line PDB file

pdb

```
Call: read.pdb(file = "1hsg")
   Total Models#: 1
     Total Atoms#: 1686, XYZs#: 5058 Chains#: 2 (values: A B)
     Protein Atoms#: 1514 (residues/Calpha atoms#: 198)
     Nucleic acid Atoms#: 0 (residues/phosphate atoms#: 0)
     Non-protein/nucleic Atoms#: 172 (residues: 128)
     Non-protein/nucleic resid values: [ HOH (127), MK1 (1) ]
   Protein sequence:
      PQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGIGGFIKVRQYD
      QILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNFPQITLWQRPLVTIKIGGQLKE
      ALLDTGADDTVLEEMSLPGRWKPKMIGGIGGFIKVRQYDQILIEICGHKAIGTVLVGPTP
      VNIIGRNLLTQIGCTLNF
+ attr: atom, xyz, seqres, helix, sheet,
        calpha, remark, call
attributes(pdb)
$names
[1] "atom"
                     "segres" "helix" "sheet" "calpha" "remark" "call"
             "xyz"
$class
[1] "pdb" "sse"
head(pdb$atom)
  type eleno elety alt resid chain resno insert
                                                            у
1 ATOM
          1
                N < NA >
                         PRO
                                 Α
                                       1
                                           <NA> 29.361 39.686 5.862 1 38.10
2 ATOM
                                       1
               CA <NA>
                         PRO
                                 Α
                                           <NA> 30.307 38.663 5.319 1 40.62
3 ATOM
          3
               C <NA>
                         PRO
                                Α
                                      1 <NA> 29.760 38.071 4.022 1 42.64
4 ATOM
          4
                O <NA>
                         PRO
                                       1 <NA> 28.600 38.302 3.676 1 43.40
                                 Α
5 ATOM
          5
               CB <NA>
                         PRO
                                      1 <NA> 30.508 37.541 6.342 1 37.87
                                 Α
                         PRO
                                 Α
                                     1 <NA> 29.296 37.591 7.162 1 38.40
6 ATOM
          6
               CG <NA>
  segid elesy charge
1 <NA>
           N
               <NA>
2 <NA>
           С
               <NA>
```

```
3 <NA> C <NA>
4 <NA> O <NA>
5 <NA> C <NA>
C <NA>
```

pdbseq(pdb)

```
3
                    5
                         6
                             7
                                  8
                                       9
                                          10
                                              11
                                                   12
                                                        13
                                                             14
                                                                 15
                                                                      16
                                                                          17
                                                                               18
"P" "Q" "I" "T" "L" "W" "Q" "R" "P" "L" "V" "T" "I" "K"
                                                               "I"
                                                                     "G"
                                                                          "G"
                                                                              "0"
         23
                            27
                                      29
              24
                   25
                       26
                                 28
                                          30
                                              31
                                                   32
                                                        33
                                                             34
                                                                 35
                                                                      36
                                                                          37
                                                                               38
                                                                     "M"
         "L" "L"
                 "D"
                      "T"
                           "G"
                                " A "
                                    "D"
                                         "D"
                                              "T"
                                                  "V"
                                                      "L"
                                                           "E"
                                                                "E"
                                                                          "S"
                                                                                   ייקיי
     42
                                                   52
          43
              44
                   45
                        46
                            47
                                 48
                                      49
                                          50
                                               51
                                                        53
                                                             54
                                                                 55
                                                                      56
                                                                          57
                                                                               58
"R" "W"
        "K"
             "P" "K" "M"
                           "I"
                                    "G" "I"
                                              "G" "G" "F"
                               "G"
                                                            "I"
                                                                "K"
                                                                     "V"
                                                                          "R"
                                                                              "0"
                                                                                   "Y"
                                                                                       "D"
              64
                   65
                       66
                            67
                                 68
                                      69
                                          70
                                               71
                                                   72
                                                        73
                                                             74
                                                                 75
                                                                      76
                                                                          77
                       "I"
                                                  "I"
                                                      "G"
                                                            "T"
                                                                "V"
             "I" "E"
                           "C"
                                "G"
                                    "H"
                                         "K" "A"
     82
          83
              84
                   85
                        86
                            87
                                 88
                                      89
                                          90
                                               91
                                                   92
                                                        93
                                                             94
                                                                 95
                                                                      96
                                                                          97
                 "I"
                                         "L"
                                              "T"
                                                  "Q"
                                                            "G"
                                                                "C"
                                                                                       "P"
         "N"
             "I"
                       "G"
                           "R"
                                "N"
                                    "L"
                                                       "I"
                                                                     "T"
                                                                          "L"
                                                                              "N"
      3
                    6
                         7
                                  9
                             8
                                      10
                                          11
                                               12
                                                   13
                                                        14
                                                             15
                                                                 16
                                                                      17
                                                                          18
                                                                               19
                                                                                    20
                                                                                         21
"O" "I"
        "T"
                 ''W''
                      "Q"
                           "R"
                                "P"
                                              "T"
             "L"
                                    "L"
                                         "V"
                                                  "I"
                                                       "K"
                                                            "I"
                                                                "G"
                                                                     "G"
                                                                          "0"
                                                                              "L"
                                                                                   "K"
                                                                                       "E"
                            28
                                 29
                                      30
                                          31
                                               32
                                                   33
                                                             35
                                                                 36
                                                                      37
                                                                          38
         "L"
             "D" "T"
                       "G"
                                "D"
                                    "D"
                                         "T"
                                              "V"
                                                  "L"
                                                      "E"
                                                            "E"
                                                                "M"
                                                                     "S"
                                                                          "L"
                                                                              "P"
                                                                                   "G"
                                                                                       "R"
                           "A"
          44
              45
                   46
                        47
                            48
                                 49
                                      50
                                          51
                                               52
                                                   53
                                                        54
                                                             55
                                                                 56
                                                                      57
                                                                          58
                                                                               59
                                                                                        61
"W" "K"
        "P" "K" "M" "I" "G"
                                              "G" "F"
                                "G"
                                    "I"
                                         "G"
                                                       "I"
                                                            "K"
                                                                "V"
                                                                     "R"
                                                                          "0"
                                                                              "Y"
                                                                                       "Q"
                                 69
                                      70
                                          71
                                               72
                                                   73
                                                        74
                                                             75
                                                                 76
         64
              65
                   66
                       67
                            68
                                                                      77
                                                                          78
                                                                               79
"T" "L" "I" "E" "I" "C" "G"
                                        "A"
                                             " T "
                                                  "G"
                                                       "T"
                                                                          "G"
                                                                              ייקיי
                                "H"
                                    "K"
                                 89
                                      90
                                          91
                                              92
                                                   93
                                                        94
                                                             95
                                                                 96
                                                                          98
         84
              85
                  86
                       87
                            88
"V" "N" "I" "I" "G" "R" "N" "L" "L" "T" "Q" "I" "G" "C" "T" "L" "N" "F"
```

Q7: How many amino acid residues are there in this pdb object?

sum(pdb\$calpha)

[1] 198

Q8: Name one of the two non-protein residues?

HOH and MK1

Q9: How many protein chains are in this structure?

2

```
unique(pdb$atom$chain)
```

```
[1] "A" "B"
```

m <- nma(adk)

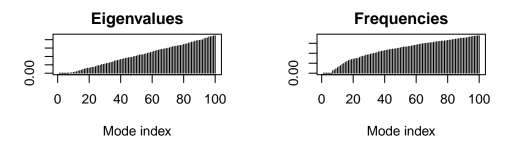
Predicting functional mortions of a single structure

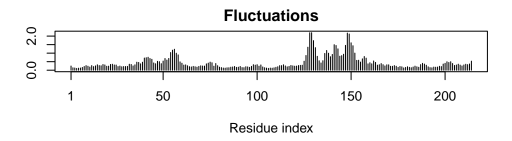
Lets do bioinformatics rediction of functional motions- i.e the movements that one of these molecules needs to make to do its stuff.

```
adk <- read.pdb("6s36")
  Note: Accessing on-line PDB file
   PDB has ALT records, taking A only, rm.alt=TRUE
adk
 Call: read.pdb(file = "6s36")
   Total Models#: 1
     Total Atoms#: 1898, XYZs#: 5694 Chains#: 1 (values: A)
     Protein Atoms#: 1654 (residues/Calpha atoms#: 214)
     Nucleic acid Atoms#: 0 (residues/phosphate atoms#: 0)
     Non-protein/nucleic Atoms#: 244 (residues: 244)
     Non-protein/nucleic resid values: [ CL (3), HOH (238), MG (2), NA (1) ]
   Protein sequence:
      MRIILLGAPGAGKGTQAQFIMEKYGIPQISTGDMLRAAVKSGSELGKQAKDIMDAGKLVT
      DELVIALVKERIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFDVPDELIVDKI
      VGRRVHAPSGRVYHVKFNPPKVEGKDDVTGEELTTRKDDQEETVRKRLVEYHQMTAPLIG
      YYSKEAEAGNTKYAKVDGTKPVAEVRADLEKILG
+ attr: atom, xyz, seqres, helix, sheet,
        calpha, remark, call
# Perform flexiblity prediction
```

Building Hessian... Done in 0.031 seconds. Diagonalizing Hessian... Done in 0.314 seconds.

plot(m)





Comparative Analysis of Protein Structures

library(bio3d)

Here we will find and analyze all ADK structures in the PDB database.

We will start with a single database accession id:"1ake_A"

```
id <- "1ake_A"
aa <-get.seq(id)</pre>
```

Warning in get.seq(id): Removing existing file: seqs.fasta

Fetching... Please wait. Done.

I ran the following

install.packages("BiocManager") BiocManager::install("msa")

Q10. Which of the packages above is found only on BioConductor and not CRAN?

The 'msa' package is from BioConductor and not CRAN

Q11. Which of the above packages is not found on BioConductor or CRAN?:

The "bio3d-view" package is not found on BioConductor or CRAN

Q12. True or False? Functions from the devtools package can be used to install packages from GitHub and BitBucket?

TRUE

Q13. How many amino acids are in this sequence, i.e. how long is this sequence?

There are 214 Amino Acids in the sequence.

```
attributes(aa)
```

#hits <- plot(b)</pre>

```
$names
[1] "id" "ali" "call"

$class
[1] "fasta"

ncol(aa$ali)

[1] 214

#b <- blast.pdb(aa)

#attributes(b)

#plot(b$hit.tbl)</pre>
```

```
#hits
```

```
#hits$pdb.id
```

Pre-calculated results:

```
hits <- NULL
hits$pdb.id <- c('1AKE_A','6S36_A','6RZE_A','3HPR_A','1E4V_A','5EJE_A','1E4Y_A','3X2S_A','6H.
# Download related PDB files
files <- get.pdb(hits$pdb.id, path="pdbs", split=TRUE, gzip=TRUE)</pre>
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/1AKE.pdb.gz exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/6S36.pdb.gz exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/6RZE.pdb.gz exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/3HPR.pdb.gz exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/1E4V.pdb.gz exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/5EJE.pdb.gz exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/1E4Y.pdb.gz exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/3X2S.pdb.gz exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/6HAP.pdb.gz exists. Skipping download
```

Warning in get.pdb(hits\$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE): pdbs/6HAM.pdb.gz exists. Skipping download

Warning in get.pdb(hits\$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE): pdbs/4K46.pdb.gz exists. Skipping download

Warning in get.pdb(hits\$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE): pdbs/3GMT.pdb.gz exists. Skipping download

Warning in get.pdb(hits\$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE): pdbs/4PZL.pdb.gz exists. Skipping download

 	l	0%
 ===== 		8%
 =========		15%
 ===================================		23%
 ===================================	l	31%
 ===================================	l	38%
 	l	46%
 	l	54%
 =======	l	62%
 	l	69%
 	l	77%
 	l	85%
 		92%
 ===================================		100%

Next we will use the odbaln() function to alighn and also optionally fit (i.e superpose) the identified PDB structures

```
pdbs <- pdbaln(files, fit = TRUE, exefile="msa")</pre>
Reading PDB files:
pdbs/split_chain/1AKE_A.pdb
pdbs/split_chain/6S36_A.pdb
pdbs/split_chain/6RZE_A.pdb
pdbs/split_chain/3HPR_A.pdb
pdbs/split_chain/1E4V_A.pdb
pdbs/split_chain/5EJE_A.pdb
pdbs/split_chain/1E4Y_A.pdb
pdbs/split_chain/3X2S_A.pdb
pdbs/split_chain/6HAP_A.pdb
pdbs/split_chain/6HAM_A.pdb
pdbs/split_chain/4K46_A.pdb
pdbs/split_chain/3GMT_A.pdb
pdbs/split_chain/4PZL_A.pdb
   PDB has ALT records, taking A only, rm.alt=TRUE
     PDB has ALT records, taking A only, rm.alt=TRUE
       PDB has ALT records, taking A only, rm.alt=TRUE
    PDB has ALT records, taking A only, rm.alt=TRUE
Extracting sequences
pdb/seq: 1
             name: pdbs/split_chain/1AKE_A.pdb
   PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 2
             name: pdbs/split_chain/6S36_A.pdb
   PDB has ALT records, taking A only, rm.alt=TRUE
             name: pdbs/split_chain/6RZE_A.pdb
pdb/seq: 3
   PDB has ALT records, taking A only, rm.alt=TRUE
             name: pdbs/split_chain/3HPR_A.pdb
pdb/seq: 4
```

PDB has ALT records, taking A only, rm.alt=TRUE

PDB has ALT records, taking A only, rm.alt=TRUE

name: pdbs/split_chain/1E4V_A.pdb

name: pdbs/split_chain/5EJE_A.pdb

pdb/seq: 5

pdb/seq: 6

Align releated PDBs

```
pdb/seq: 7
             name: pdbs/split_chain/1E4Y_A.pdb
pdb/seq: 8
             name: pdbs/split_chain/3X2S_A.pdb
pdb/seq: 9
             name: pdbs/split_chain/6HAP_A.pdb
pdb/seq: 10
              name: pdbs/split_chain/6HAM_A.pdb
   PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 11
              name: pdbs/split_chain/4K46_A.pdb
   PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 12
              name: pdbs/split_chain/3GMT_A.pdb
pdb/seq: 13
              name: pdbs/split chain/4PZL A.pdb
```

pdbs

[Truncated_Name:1]1AKE_A.pdb
[Truncated_Name:2]6S36_A.pdb
[Truncated_Name:3]6RZE_A.pdb
[Truncated_Name:4]3HPR_A.pdb
[Truncated_Name:5]1E4V_A.pdb
[Truncated_Name:6]5EJE_A.pdb
[Truncated_Name:7]1E4Y_A.pdb
[Truncated_Name:8]3X2S_A.pdb
[Truncated_Name:9]6HAP_A.pdb
[Truncated_Name:10]6HAM_A.pdb
[Truncated_Name:11]4K46_A.pdb
[Truncated_Name:12]3GMT_A.pdb
[Truncated_Name:13]4PZL_A.pdb

40 ----MRIILLGAPGAGKGTQAQFIMEKYGIPQIS -----MRIILLGAPGAGKGTQAQFIMEKYGIPQIS -----MRIILLGAPGAGKGTQAQFIMEKYGIPQIS -----MRIILLGAPGAGKGTQAQFIMEKYGIPQIS -----MRIILLGAPVAGKGTQAQFIMEKYGIPQIS ----MRIILLGAPGAGKGTQAQFIMEKYGIPQIS -----MRIILLGALVAGKGTQAQFIMEKYGIPQIS ----MRIILLGAPGAGKGTQAQFIMEKYGIPQIS ----MRIILLGAPGAGKGTQAQFIMEKYGIPQIS -----MRIILLGAPGAGKGTQAQFIMEKYGIPQIS -----MRIILLGAPGAGKGTQAQFIMAKFGIPQIS ----MRLILLGAPGAGKGTQANFIKEKFGIPQIS TENLYFQSNAMRIILLGAPGAGKGTQAKIIEQKYNIAHIS **^**** *****

40

[Truncated_Name:1]1AKE_A.pdb
[Truncated_Name:2]6S36_A.pdb
[Truncated_Name:3]6RZE_A.pdb
[Truncated_Name:4]3HPR_A.pdb
[Truncated_Name:5]1E4V_A.pdb
[Truncated_Name:6]5EJE_A.pdb
[Truncated_Name:7]1E4Y_A.pdb
[Truncated_Name:8]3X2S_A.pdb
[Truncated_Name:9]6HAP_A.pdb
[Truncated_Name:10]6HAM_A.pdb
[Truncated_Name:11]4K46_A.pdb
[Truncated_Name:12]3GMT_A.pdb

TGDMLRAAVKSGSELGKQAKDIMDAGKLVTDELVIALVKE
TGDMLRAAVKSGSELGKQAKDIMDAGKLVTDEIIIALVKE
TGDMLRAAIKSGSELGKQAKDIMDAGKLVTDEIIIALVKE
TGDMLRAAIKSGSELGKQAKDIMDAGKLVTDEIIIALVKE

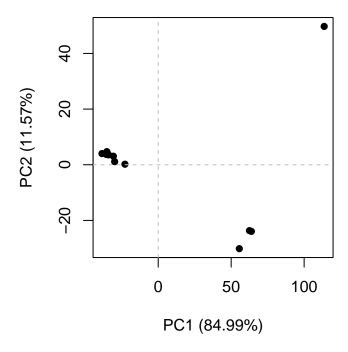
1

[Truncated_Name:13]4PZL_A.pdb	.pdb TGDMIRETIKSGSALGQELKKVLDAGELVSDEFIIKIVKD						
	****^*	^* *^	**	* ^*	* ** *	^^ ^*·	^^
	41						80
	81				•		120
[Truncated_Name:1]1AKE_A.pdb	RIAQEDO	CRNGFLL	DGFPR	ΓΙΡQΑΙ	DAMKEAGI	IVDYVLE:	FD
[Truncated_Name:2]6S36_A.pdb	RIAQEDO	CRNGFLL	DGFPR	ΓΙΡQΑΙ	DAMKEAGI	IVDYVLE	FD
[Truncated_Name:3]6RZE_A.pdb	RIAQEDO	CRNGFLL	DGFPR	ΓΙΡQΑΙ	DAMKEAGI	IVDYVLE	FD
[Truncated_Name:4]3HPR_A.pdb	RIAQEDO	CRNGFLL	DGFPR	ΓΙΡQΑΙ	DAMKEAGI	IVDYVLE	FD
[Truncated_Name:5]1E4V_A.pdb	RIAQEDO	CRNGFLL	DGFPR	ΓΙΡQΑΙ	DAMKEAGI	IVDYVLE	FD
[Truncated_Name:6]5EJE_A.pdb	RIAQEDO	CRNGFLL	DGFPR	ΓΙΡQΑΙ	DAMKEAGI	IVDYVLE	FD
[Truncated_Name:7]1E4Y_A.pdb	RIAQEDO	CRNGFLL	DGFPR	ΓΙΡQΑΙ	DAMKEAGI	IVDYVLE!	FD
[Truncated_Name:8]3X2S_A.pdb	RIAQEDS	RNGFLL	DGFPR	ΓΙΡQΑΙ	DAMKEAGI	IVDYVLE!	FD
[Truncated_Name:9]6HAP_A.pdb	RICQEDS	RNGFLL	DGFPR	ΓΙΡQΑΙ	DAMKEAGI	IVDYVLE!	FD
[Truncated_Name:10]6HAM_A.pdb	RICQEDS	RNGFLL	DGFPRT	ΓΙΡQΑΙ	DAMKEAGI	IVDYVLE:	FD
[Truncated_Name:11]4K46_A.pdb	RIAQDDO	CAKGFLL	DGFPRT	ΓΙΡQΑΙ	OGLKEVGV	/VDYVIE	FD
[Truncated_Name:12]3GMT_A.pdb	RLKEADO	CANGYLF	DGFPRT	ΓΙΑQΑΙ	DAMKEAGV <i>I</i>	AIDYVLE:	ID
[Truncated_Name:13]4PZL_A.pdb					QELDKLGVI		
1	*^ *		** ***	-	· -	^**^^*	
	81						120
	-	•			-		
	121						160
[Truncated_Name:1]1AKE_A.pdb	VPDELIV	DRIVGR	RVHAPS	SGRVYF	IVKFNPPKV	/EGKDDV	TG
[Truncated_Name:2]6S36_A.pdb	VPDELIV	/DKIVGR	RVHAPS	SGRVYF	HVKFNPPKV	/EGKDDV	TG
[Truncated_Name:3]6RZE_A.pdb	VPDELIV	DAIVGR	RVHAPS	SGRVYF	HVKFNPPKV	/EGKDDV	TG
[Truncated_Name:4]3HPR_A.pdb	VPDELIV	DRIVGR	RVHAPS	SGRVYF		/EGKDDG	TG
[Truncated_Name:5]1E4V_A.pdb	VPDELIV	DRIVGR	RVHAPS	SGRVYF	HVKFNPPKV	JEGKDDV'	TG
[Truncated_Name:6]5EJE_A.pdb	VPDELIV	DRIVGR	RVHAPS	SGRVYF		/EGKDDV	TG
[Truncated_Name:7]1E4Y_A.pdb	VPDELIV	DRIVGR	RVHAPS	SGRVYF	IVKFNPPKV	/EGKDDV	TG
[Truncated_Name:8]3X2S_A.pdb					IVKFNPPKV		
[Truncated_Name:9]6HAP_A.pdb	VPDELIV	DRIVGR	RVHAPS	SGRVYF	IVKFNPPKV	/EGKDDV	TG
[Truncated_Name:10]6HAM_A.pdb	VPDELTV	DR.TVGR.	RVHAPS	SGRVYF		/FGKDDV	TG
[Truncated Name:11]4K46_A.pdb					INVYNPPKV	. —	
[Truncated_Name:12]3GMT_A.pdb					IVKFNPPKV		
[Truncated_Name:13]4PZL_A.pdb					TKFNPPKV		
[II uncaved_Name: 10] II ZL_A.pub	* ^^	·^ ^ **			* ^****		
	121	444	4 4 4		, ,,,,,,,	e dedede	160
	121	•		•	•		100
	161						200
[Truncated_Name:1]1AKE_A.pdb		מחחדדים	VBKBI	YEAHUM	MTAPLIGYY	/SKFAFA	
[Truncated_Name: 2] 6S36_A.pdb					TAPLIGY:		
[Truncated_Name:3]6RZE_A.pdb					TTAPLIGY:		
[Truncated_Name:3] ORZE_A.pdb [Truncated_Name:4] 3HPR_A.pdb				-			
[11 dicated_wame:4] SHPK_A.pdb	CCLIIK	NUMERI	N UVUT/	ᄓᄱᄓ	TAPLIGY!	OVENEV	ΔI/I

```
[Truncated_Name:5]1E4V_A.pdb
                                EELTTRKDDQEETVRKRLVEYHQMTAPLIGYYSKEAEAGN
[Truncated_Name:6]5EJE_A.pdb
                                EELTTRKDDQEECVRKRLVEYHQMTAPLIGYYSKEAEAGN
[Truncated_Name:7]1E4Y_A.pdb
                                EELTTRKDDQEETVRKRLVEYHQMTAPLIGYYSKEAEAGN
[Truncated_Name:8]3X2S_A.pdb
                                EELTTRKDDQEETVRKRLCEYHQMTAPLIGYYSKEAEAGN
[Truncated Name:9]6HAP A.pdb
                                EELTTRKDDQEETVRKRLVEYHQMTAPLIGYYSKEAEAGN
[Truncated Name: 10] 6HAM A.pdb
                                EELTTRKDDQEETVRKRLVEYHQMTAPLIGYYSKEAEAGN
[Truncated Name:11]4K46 A.pdb
                                EDLVIREDDKEETVLARLGVYHNQTAPLIAYYGKEAEAGN
[Truncated_Name:12]3GMT_A.pdb
                                EPLVQRDDDKEETVKKRLDVYEAQTKPLITYYGDWARRGA
[Truncated_Name:13]4PZL_A.pdb
                                EPLITRTDDNEDTVKQRLSVYHAQTAKLIDFYRNFSSTNT
                                     * ** *^ *
                                                **
                              161
                                                                        200
                              201
                                                           227
[Truncated_Name:1]1AKE_A.pdb
                                T--KYAKVDGTKPVAEVRADLEKILG-
[Truncated_Name:2]6S36_A.pdb
                                T--KYAKVDGTKPVAEVRADLEKILG-
[Truncated_Name:3]6RZE_A.pdb
                                T--KYAKVDGTKPVAEVRADLEKILG-
[Truncated_Name:4]3HPR_A.pdb
                                T--KYAKVDGTKPVAEVRADLEKILG-
                                T--KYAKVDGTKPVAEVRADLEKILG-
[Truncated_Name:5]1E4V_A.pdb
[Truncated_Name:6]5EJE_A.pdb
                                T--KYAKVDGTKPVAEVRADLEKILG-
[Truncated Name:7]1E4Y A.pdb
                                T--KYAKVDGTKPVAEVRADLEKILG-
[Truncated Name:8]3X2S A.pdb
                                T--KYAKVDGTKPVAEVRADLEKILG-
[Truncated Name:9]6HAP A.pdb
                                T--KYAKVDGTKPVCEVRADLEKILG-
[Truncated_Name:10]6HAM_A.pdb
                                T--KYAKVDGTKPVCEVRADLEKILG-
[Truncated Name:11]4K46 A.pdb
                                T--QYLKFDGTKAVAEVSAELEKALA-
[Truncated_Name:12]3GMT_A.pdb
                                E----YRKISG-
[Truncated_Name: 13] 4PZL_A.pdb
                                KIPKYIKINGDQAVEKVSQDIFDQLNK
                              201
                                                           227
Call:
  pdbaln(files = files, fit = TRUE, exefile = "msa")
Class:
 pdbs, fasta
Alignment dimensions:
  13 sequence rows; 227 position columns (204 non-gap, 23 gap)
+ attr: xyz, resno, b, chain, id, ali, resid, sse, call
```

pc.xray <-pca(pdbs)</pre>

```
plot(pc.xray, pc.axes =c(1,2))
```

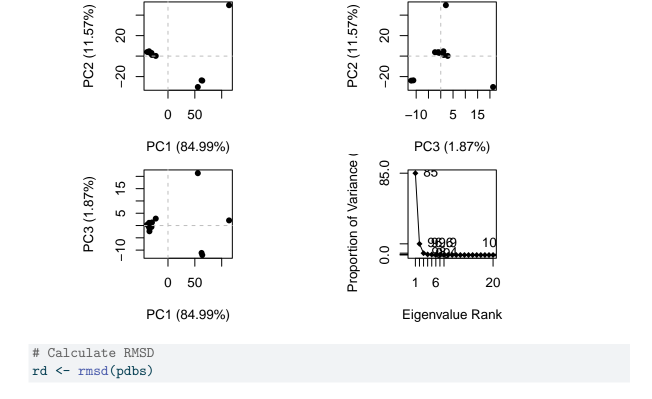


```
uniprot <- 248838887
pdb <- 195610
pdb/uniprot * 100
```

[1] 0.0786091

Principal Component Analysis

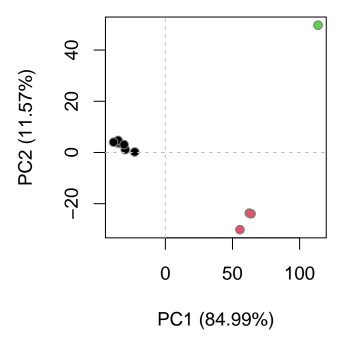
```
# Perform PCA
pc.xray <- pca(pdbs)
plot(pc.xray)</pre>
```



Warning in rmsd(pdbs): No indices provided, using the 204 non NA positions

```
# Structure-based clustering
hc.rd <- hclust(dist(rd))
grps.rd <- cutree(hc.rd, k=3)

plot(pc.xray, 1:2, col="grey50", bg=grps.rd, pch=21, cex=1)</pre>
```



To visualize the major structural variations in the ensemble the function mktrj() can be used to generate a trajectory PDB file by interpolating along a give PC (eigenvector):

```
pc1 <- mktrj(pc.xray, pc=1, file="pc_1.pdb")</pre>
```

Q14. What do you note about this plot? Are the black and colored lines similar or different? Where do you think they differ most and why?

```
# NMA of all structures
modes <- nma(pdbs)</pre>
```

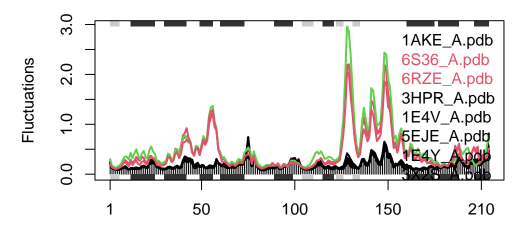
Details of Scheduled Calculation:

- ... 13 input structures
- ... storing 606 eigenvectors for each structure
- ... dimension of x\$U.subspace: (612x606x13)
- ... coordinate superposition prior to NM calculation
- ... aligned eigenvectors (gap containing positions removed)
- estimated memory usage of final 'eNMA' object: 36.9 Mb

0%

plot(modes, pdbs, col=grps.rd)

Extracting SSE from pdbs\$sse attribute



Residue number (reference PDB: 1AKE_A)