Lab 09

AUTHOR
Nichelle Camden

1. Introduction to the RCSB Protein Data Bank (PDB)

Q1: What percentage of structures in the PDB are solved by X-Ray and Electron Microscopy.

169,916 out of 196,979 total, about 86%.

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

most of them! 171,351 out of 196,979 total, about 87%

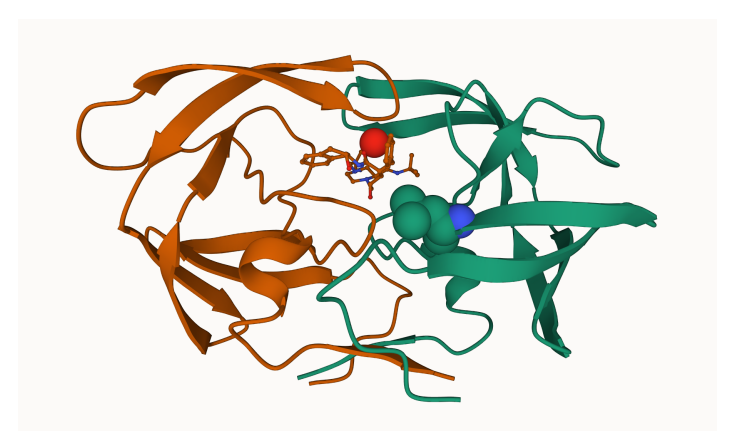
#Q3: Type HIV in the PDB website search box on the home page and determine how many HIV-1 protease structures are in the current PDB?

22,183

2. Visualizing the HIV-1 protease structure

Viewing PDB sturctures with Mol*

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Reading and working with structyres in R

The bio3d package for structural bioinformatics has a lot of features for reading and working with biomolecular sequences and structures.

```
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:

LATTEMODE EATTURED FROM LAND LAFFILISE ADMINISTRATION TO A DATE OF THE PROPERTY OF THE PROPERT

QILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNFPQITLWQRPLVTIKIGGQLKE ALLDTGADDTVLEEMSLPGRWKPKMIGGIGGFIKVRQYDQILIEICGHKAIGTVLVGPTP VNIIGRNLLTQIGCTLNF

+ attr: atom, xyz, segres, helix, sheet,

```
calpha, remark, call
```

```
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

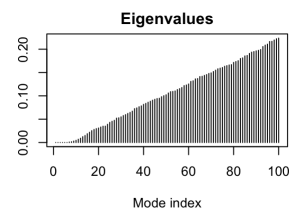
Normal mode analysis (NMA) is a bioinformatics method for predicting functional motions. It will show us the parts of the protein that are "flexible" (i.e. the most dynamic).

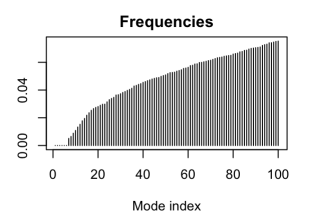
```
m <- nma(adk)
```

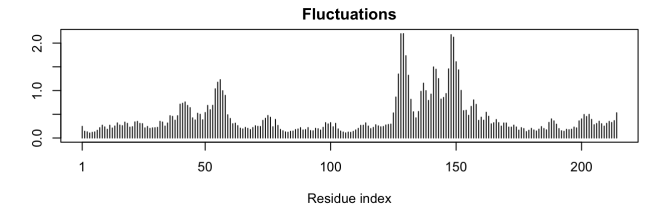
Building Hessian... Done in 0.084 seconds. Diagonalizing Hessian... Done in 0.634 seconds.

plot(m)

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Make a "movie" of this thing moving.

```
mktrj(m, file="adk_nma.pdb")
```

Comparative analysis of all ADK structures

First, we get the sequence of ADK and use this to search the PDB database.

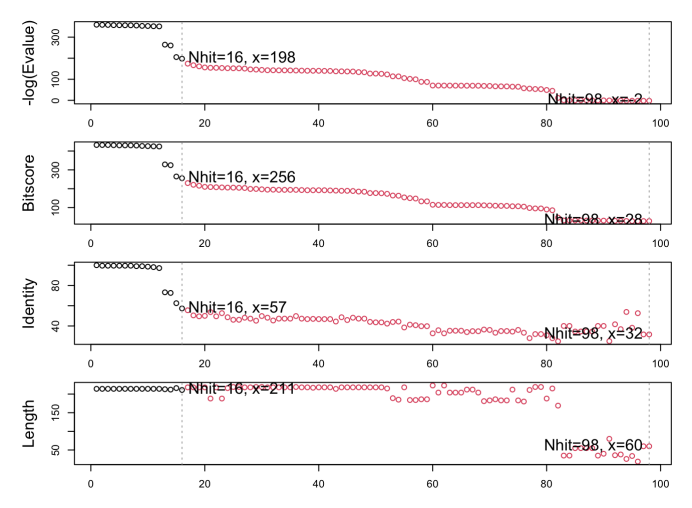
```
aa <- get.seq("lake_a")</pre>
```

Warning in get.seq("1ake_a"): Removing existing file: seqs.fasta

Fetching... Please wait. Done.

```
120
             61
            121
                                                                              180
              VGRRVHAPSGRVYHVKFNPPKVEGKDDVTGEELTTRKDDQEETVRKRLVEYH0MTAPLIG
pdb | 1AKE | A
            121
                                                                              180
            181
                                                  214
              YYSKEAEAGNTKYAKVDGTKPVAEVRADLEKILG
pdb | 1AKE | A
            181
                                                  214
Call:
  read.fasta(file = outfile)
Class:
  fasta
Alignment dimensions:
  1 sequence rows; 214 position columns (214 non-gap, 0 gap)
+ attr: id, ali, call
Run a blast search of the PDB database using this sequence
 blast <- blast.pdb(aa)</pre>
 Searching ... please wait (updates every 5 seconds) RID = P02CFPWU016
 Reporting 98 hits
 #blast
examine the results
 hits <- plot(blast)</pre>
  * Possible cutoff values:
                                 197 -3
             Yielding Nhits:
                                 16 98
  * Chosen cutoff value of:
                                 197
             Yielding Nhits:
                                 16
```

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download all the ADK structures in the PDB database

```
hits$pdb.id
```

```
[1] "1AKE_A" "4X8M_A" "6S36_A" "6RZE_A" "4X8H_A" "3HPR_A" "1E4V_A" "5EJE_A"
```

[9] "1E4Y A" "3X2S A" "6HAP A" "6HAM A" "4K46 A" "4NP6 A" "3GMT A" "4PZL A"

#pdb.annotate(hits\$pdb.id)

```
# 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/ 4X8M.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

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10/31/22, 10:51 AM

Lab 09 Warning in get.pdb(hits\$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE): pdbs/ 4X8H.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 4NP6.pdb.gz exists. Skipping download 3GMT.pdb.gz exists. Skipping download

Warning in get.pdb(hits\$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE): pdbs/

Warning in get.pdb(hits\$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE): pdbs/

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

0% 6% 12% 19% ========== 25% _____ 31% _____ 38% _____

 ===================================		44%
====================================		50%
 ===================================		56%
 =========		62%
 ===================================	ı	69%
 ===================================	ı	75%
 ===================================	ı	81%
 ===================================		88%
 ===================================	ı	94%
 ===================================		100%

Viewing all these structures looks like a mess. We need to try something else...

We will align and superimpose these structures.

```
library("BiocManager")
pdbs <- pdbaln(files, fit = TRUE, exefile="msa")</pre>
```

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```
Reading PDB files:
pdbs/split_chain/1AKE_A.pdb
pdbs/split_chain/4X8M_A.pdb
pdbs/split_chain/6S36_A.pdb
pdbs/split_chain/6RZE_A.pdb
pdbs/split_chain/4X8H_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/4NP6_A.pdb
pdbs/split_chain/3GMT_A.pdb
pdbs/split_chain/4PZL_A.pdb
```

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

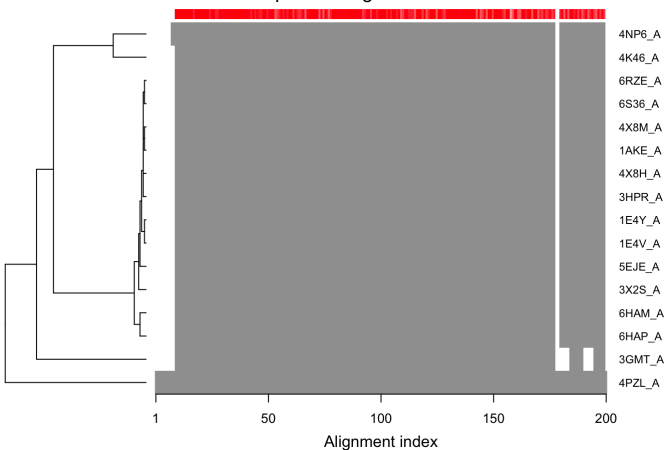
- .. PDB has ALT records, taking A only, rm.alt=TRUE
- . PDB has ALT records, taking A only, rm.alt=TRUE
- .. PDB has ALT records, taking A only, rm.alt=TRUE
- .. PDB has ALT records, taking A only, rm.alt=TRUE

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```
PDB has ALT records, taking A only, rm.alt=TRUE
   PDB has ALT records, taking A only, rm.alt=TRUE
Extracting sequences
             name: pdbs/split_chain/1AKE_A.pdb
pdb/seq: 1
   PDB has ALT records, taking A only, rm.alt=TRUE
             name: pdbs/split_chain/4X8M_A.pdb
pdb/seq: 2
pdb/seq: 3
             name: pdbs/split_chain/6S36_A.pdb
   PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 4
             name: pdbs/split_chain/6RZE_A.pdb
   PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 5
             name: pdbs/split_chain/4X8H_A.pdb
pdb/seq: 6
             name: pdbs/split_chain/3HPR_A.pdb
  PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 7
             name: pdbs/split_chain/1E4V_A.pdb
pdb/seq: 8
             name: pdbs/split_chain/5EJE_A.pdb
   PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 9
             name: pdbs/split_chain/1E4Y_A.pdb
pdb/seq: 10
              name: pdbs/split_chain/3X2S_A.pdb
pdb/seq: 11
              name: pdbs/split_chain/6HAP_A.pdb
pdb/seq: 12
              name: pdbs/split_chain/6HAM_A.pdb
   PDB has ALT records, taking A only, rm.alt=TRUE
              name: pdbs/split chain/4K46 A.pdb
pdb/seq: 13
   PDB has ALT records, taking A only, rm.alt=TRUE
              name: pdbs/split chain/4NP6 A.pdb
pdb/seq: 14
pdb/seq: 15
              name: pdbs/split_chain/3GMT_A.pdb
pdb/seq: 16
              name: pdbs/split_chain/4PZL_A.pdb
#pdbs
# Vector containing PDB codes for figure axis
ids <- basename.pdb(pdbs$id)</pre>
# Draw schematic alignment
plot(pdbs, labels=ids)
```

```
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```

Sequence Alignment Overview



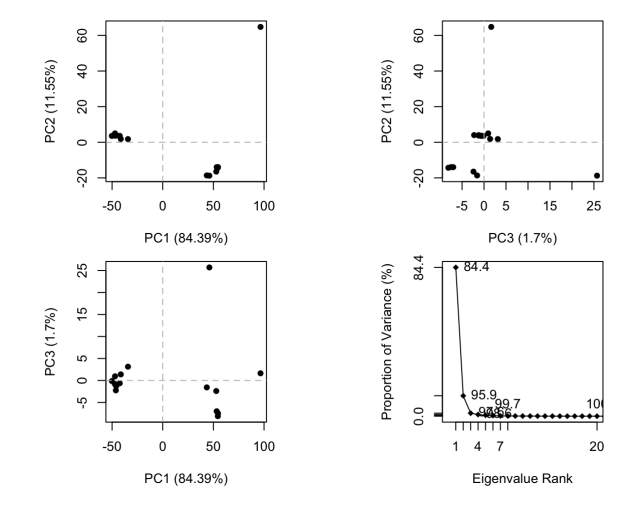
PCA to the rescue...

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

and plot my results

```
plot(pc.xray)
```

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```
# Calculate RMSD
rd <- rmsd(pdbs)</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)</pre>
```

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

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