Bimm 143 Class 12: Structural Bioinformatics Part 2

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Introduction to Bio3D in R

```
# load the library
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
aa123(pdbseq(pdb))
##
     [1] "PRO" "GLN" "ILE" "THR" "LEU" "TRP" "GLN" "ARG" "PRO" "LEU" "VAL" "THR"
    [13] "ILE" "LYS" "ILE" "GLY" "GLY" "GLN" "LEU" "LYS" "GLU" "ALA" "LEU" "LEU"
    [25] "ASP" "THR" "GLY" "ALA" "ASP" "ASP" "THR" "VAL" "LEU" "GLU" "GLU" "MET"
##
    [37] "SER" "LEU" "PRO" "GLY" "ARG" "TRP" "LYS" "PRO" "LYS" "MET" "ILE" "GLY"
    [49] "GLY" "ILE" "GLY" "GLY" "PHE" "ILE" "LYS" "VAL" "ARG" "GLN" "TYR" "ASP"
##
    [61] "GLN" "ILE" "LEU" "ILE" "GLU" "ILE" "CYS" "GLY" "HIS" "LYS" "ALA" "ILE"
    [73] "GLY" "THR" "VAL" "LEU" "VAL" "GLY" "PRO" "THR" "PRO" "VAL" "ASN" "ILE"
    [85] "ILE" "GLY" "ARG" "ASN" "LEU" "LEU" "THR" "GLN" "ILE" "GLY" "CYS" "THR"
   [97] "LEU" "ASN" "PHE" "PRO" "GLN" "ILE" "THR" "LEU" "TRP" "GLN" "ARG" "PRO"
  [109] "LEU" "VAL" "THR" "ILE" "LYS" "ILE" "GLY" "GLY" "GLN" "LEU" "LYS" "GLU"
## [121] "ALA" "LEU" "LEU" "ASP" "THR" "GLY" "ALA" "ASP" "ASP" "THR" "VAL" "LEU"
```

```
## [133] "GLU" "GLU" "MET" "SER" "LEU" "PRO" "GLY" "ARG" "TRP" "LYS" "PRO" "LYS"
  [145] "MET" "ILE" "GLY" "GLY" "ILE" "GLY" "PHE" "ILE" "LYS" "VAL" "ARG"
## [157] "GLN" "TYR" "ASP" "GLN" "ILE" "LEU" "ILE" "GLU" "ILE" "CYS" "GLY" "HIS"
## [169] "LYS" "ALA" "ILE" "GLY" "THR" "VAL" "LEU" "VAL" "GLY" "PRO" "THR" "PRO"
## [181] "VAL" "ASN" "ILE" "ILE" "GLY" "ARG" "ASN" "LEU" "LEU" "THR" "GLN" "ILE"
## [193] "GLY" "CYS" "THR" "LEU" "ASN" "PHE"
# ATOM records
head(pdb$atom)
     type eleno elety alt resid chain resno insert
                    N < NA >
                             PRO
                                    Α
                                                <NA> 29.361 39.686 5.862 1 38.10
              1
                   CA <NA>
## 2 ATOM
              2
                             PRO
                                     Α
                                                <NA> 30.307 38.663 5.319 1 40.62
## 3 ATOM
                    C <NA>
                             PRO
                                     Α
                                            1 <NA> 29.760 38.071 4.022 1 42.64
                                            1 <NA> 28.600 38.302 3.676 1 43.40
## 4 ATOM
                    O <NA>
                             PRO
                                            1 <NA> 30.508 37.541 6.342 1 37.87
                   CB <NA>
## 5 ATOM
              5
                             PRO
                                     Α
## 6 ATOM
                   CG <NA>
                             PRO
                                            1 <NA> 29.296 37.591 7.162 1 38.40
     segid elesy charge
##
## 1
     <NA>
                   <NA>
## 2
      <NA>
               С
                   <NA>
## 3
     <NA>
                   <NA>
## 4
     <NA>
               0
                   <NA>
## 5
      <NA>
                   <NA>
## 6 <NA>
                   <NA>
     Q7: How many amino acid residues are there in this pdb object?
198
     Q8: Name one of the two non-protein residues?
HOH
     Q9: How many protein chains are in this structure?
2
```

Comparative Analysis of Protein Structures

```
library(bio3d)
pdb <- read.pdb("1hel")</pre>
##
     Note: Accessing on-line PDB file
pdb
##
    Call: read.pdb(file = "1hel")
##
##
##
      Total Models#: 1
##
        Total Atoms#: 1186, XYZs#: 3558 Chains#: 1 (values: A)
##
##
        Protein Atoms#: 1001 (residues/Calpha atoms#: 129)
##
        Nucleic acid Atoms#: 0 (residues/phosphate atoms#: 0)
##
##
        Non-protein/nucleic Atoms#: 185 (residues: 185)
##
        Non-protein/nucleic resid values: [ HOH (185) ]
```

```
##
## Protein sequence:
## KVFGRCELAAAMKRHGLDNYRGYSLGNWVCAAKFESNFNTQATNRNTDGSTDYGILQINS
## RWWCNDGRTPGSRNLCNIPCSALLSSDITASVNCAKKIVSDGNGMNAWVAWRNRCKGTDV
## QAWIRGCRL
##
## + attr: atom, xyz, seqres, helix, sheet,
## calpha, remark, call
```

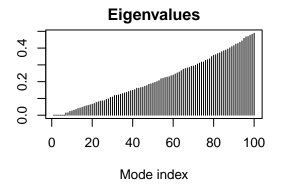
Let's use bioinformatics methods called NMA (Normal Mode Analysis) to predict the dynamics (flexibility) of the enzyme

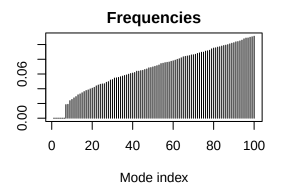
```
modes <- nma(pdb)

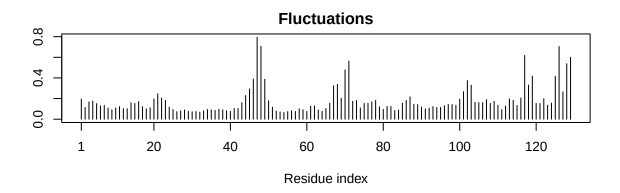
## Building Hessian... Done in 0.037 seconds.

## Diagonalizing Hessian... Done in 2.209 seconds.

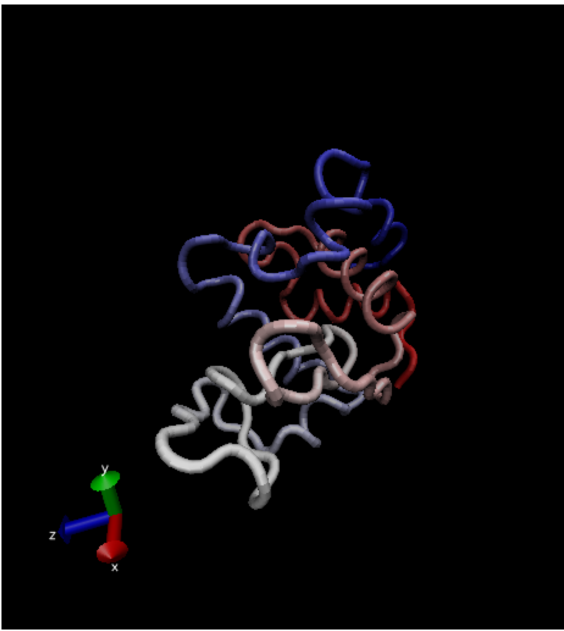
plot(modes)
```







Make a "move" of its predicted motion. We often call this a "trajectory" mktrj(modes, file="nma.pdb")



Upload image of nma.pdb # Analysis of ADK

Q10. Which of the packages above is found only on BioConductor and not CRAN? "msa" $\,$

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

The packages that are downloaded from GitHub or Bitbucket are not on Bioconductor or CRAN. In this case it is "Grantlab/bio3d-view" from bitbucket

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

TRUE

Search and Retreive ADK structures

```
aa <- get.seq("1ake_A")</pre>
## Warning in get.seq("lake_A"): Removing existing file: seqs.fasta
## Fetching... Please wait. Done.
    Q13. How many amino acids are in this sequence, i.e. how long is this sequence?
214
# List out some 'top hits'
hits <- NULL
hits$pdb.id <- c('1AKE_A','4X8M_A','6S36_A','6RZE_A','4X8H_A','3HPR_A','1E4V_A','5EJE_A','1E4Y_A','3X2S
# Download releated PDB files
files <- get.pdb(hits$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
## 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$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE): pdbs/
## 4K46.pdb.gz exists. Skipping download
## Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE): pdbs/
## 4NP6.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
```

Multiple Structure Alignment # Align releated PDBs pdbs <- pdbaln(files, fit = TRUE, exefile="./muscle")</pre> ## 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 ## ## ## 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/4X8M_A.pdb name: pdbs/split_chain/6S36_A.pdb pdb/seq: 3 ## PDB has ALT records, taking A only, rm.alt=TRUE name: pdbs/split_chain/6RZE_A.pdb pdb/seq: 4 ## 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 name: pdbs/split_chain/1E4V_A.pdb ## pdb/seq: 7 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

pdb/seq: 13

pdb/seq: 12

name: pdbs/split_chain/6HAP_A.pdb

name: pdbs/split_chain/6HAM_A.pdb

name: pdbs/split_chain/4K46_A.pdb

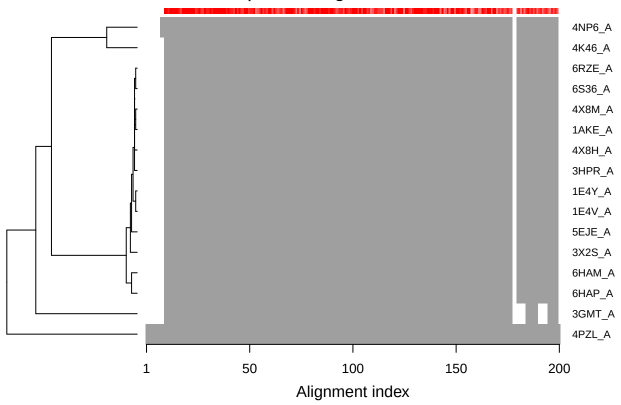
PDB has ALT records, taking A only, rm.alt=TRUE

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

```
## pdb/seq: 14    name: pdbs/split_chain/4NP6_A.pdb
## pdb/seq: 15    name: pdbs/split_chain/3GMT_A.pdb
## pdb/seq: 16    name: pdbs/split_chain/4PZL_A.pdb

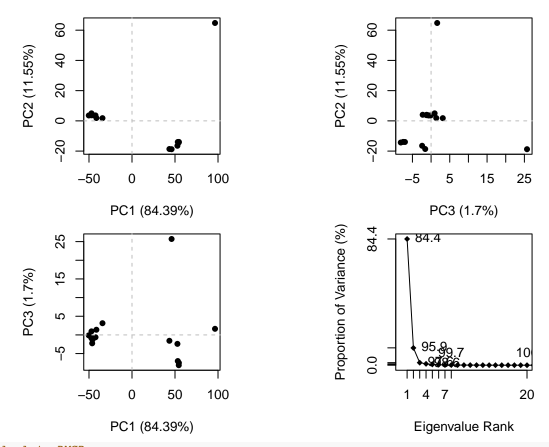
# Vector containing PDB codes for figure axis
ids <- basename.pdb(pdbs$id)
# Draw schematic alignment
plot(pdbs, labels=ids)</pre>
```

Sequence Alignment Overview



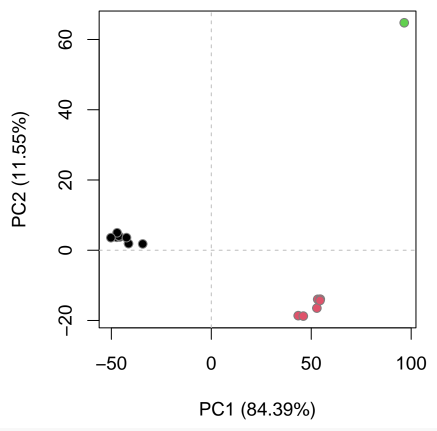
Use bio3d pca() function

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

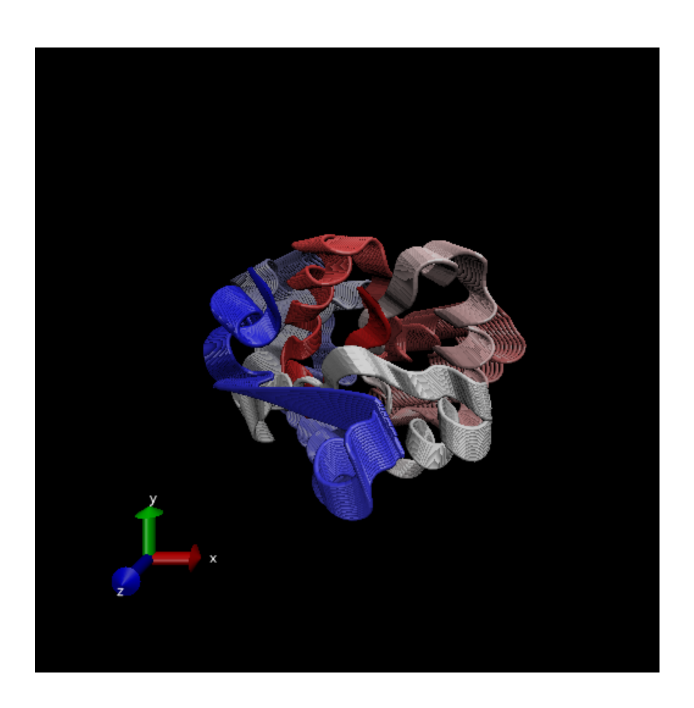


```
# Calculate RMSD
rd <- rmsd(pdbs)</pre>
```

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



Visualize first principal component
pc1 <- mktrj(pc.xray, pc=1, file="pc_1.pdb")</pre>



Find a gene VMD file:CATF845.fwd CATF

