

Class 12_Structural Bioinformatics II

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Comparative analysis of protein structures

Using the bio3d package.

```
library(bio3d)

pdb <- read.pdb("1hel")
```

```
## Note: Accessing on-line PDB file
```

```
pdb
```

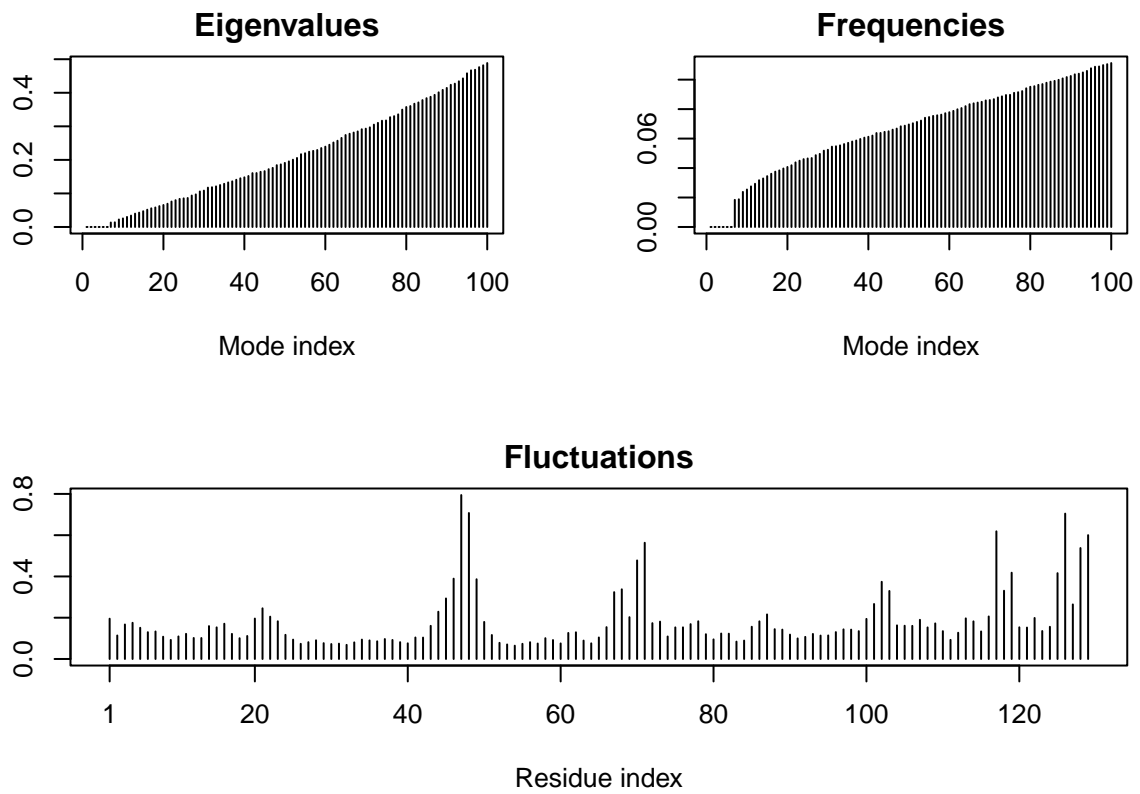
```
##
## Call: read.pdb(file = "1hel")
##
## Total Models#: 1
## Total Atoms#: 1186, XYZs#: 3558 Chains#: 1 (values: A)
##
## Protein Atoms#: 1001 (residues/Calpha atoms#: 129)
## Nucleic acid Atoms#: 0 (residues/phosphate atoms#: 0)
##
## Non-protein/nucleic Atoms#: 185 (residues: 185)
## Non-protein/nucleic resid values: [ HOH (185) ]
##
## Protein sequence:
## KVFGRCELAAAMKRHGLDNYRGYSLGNWVCAAKFESNFNTQATNRNTDGSTDYGILQINS
## RWWCNDGRTPGSRNLCNIPCSALLSSDITASVNCAKKIVSDGNGMNAWVAWRNRCKGTDV
## QAWIRGCRL
##
## + attr: atom, xyz, seqres, helix, sheet,
## calpha, remark, call
```

Let's use a bioinformatics method called NMA (Normal Mode Analysis) to predict the dynamics (flexibility) of this enzyme.

```
modes <- nma(pdb)
```

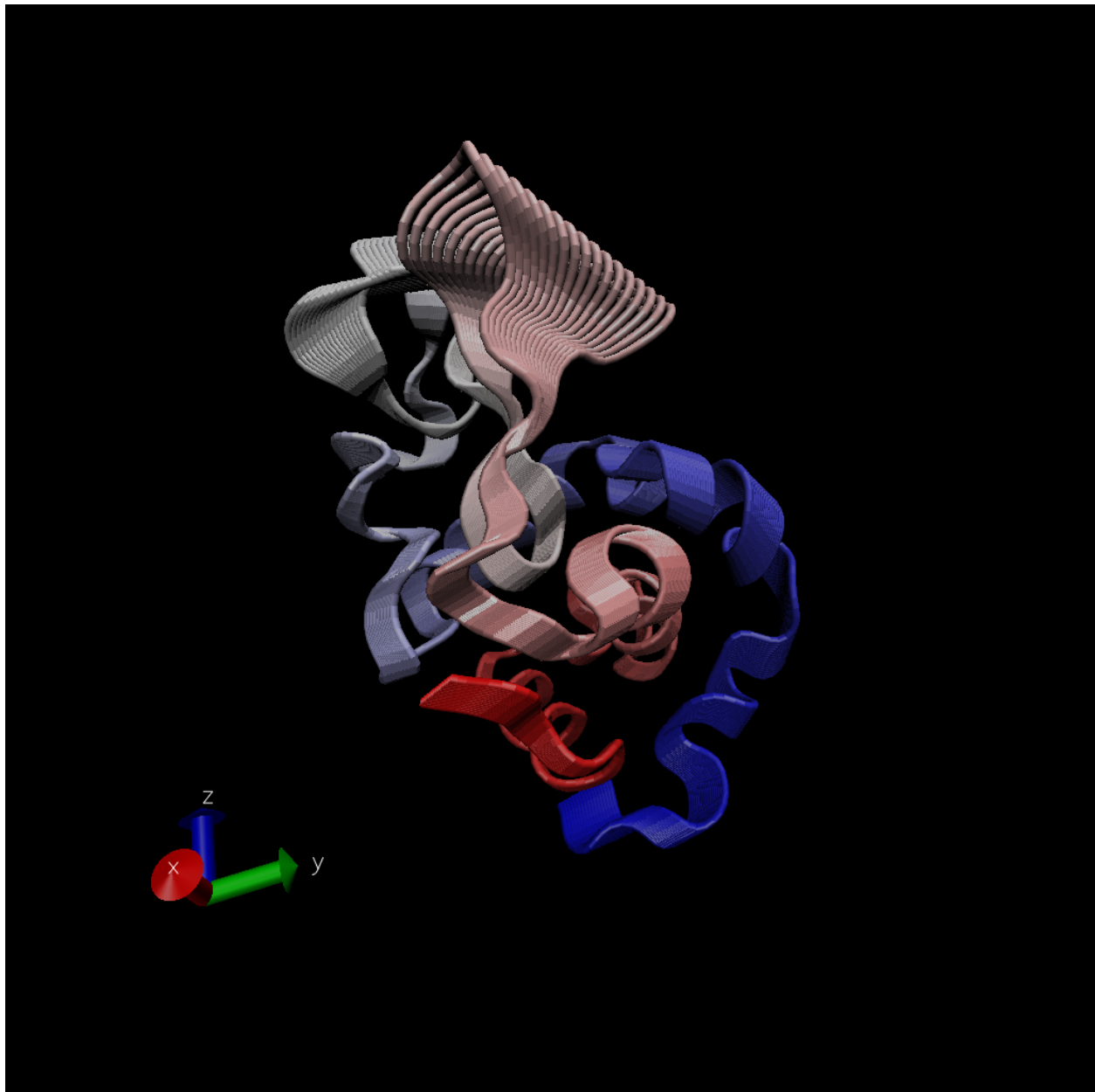
```
## Building Hessian... Done in 0.031 seconds.
## Diagonalizing Hessian... Done in 0.093 seconds.
```

```
plot.nma(modes)
```



Make a “move” of its predicted motion. We often call this a “trajectory”.

```
mktrj(modes, file = "nma.pdb")
```



Analysis of ADK

```
library(bio3d)  
aa <- get.seq("1ake_A")
```

```
## Warning in get.seq("1ake_A"): Removing existing file: seqs.fasta
```

```
## Fetching... Please wait. Done.
```

```
aa
```

```
##          1          .          .          .          .          .          60
## pdb|1AKE|A  MRIILLGAPGAGKGTQAQFIMEKYGIPQISTGDMLRAAVKSGSELGKQAKDIMDAGKLV
##          1          .          .          .          .          .          60
##
##          61          .          .          .          .          .          120
## pdb|1AKE|A  DELVIALVKERIAQEDCRNGFLLDGFPR TIPQADAMKEAGINVDYVLEFDVPDELIVDRI
##          61          .          .          .          .          .          120
##
##          121         .          .          .          .          .          180
## pdb|1AKE|A  VGRRVHAPSGRVYHVKNPPKVEGKDDVTGEELTTRKDDQEETVRKRLVEYHQMTAPLIG
##          121         .          .          .          .          .          180
##
##          181         .          .          .          214
## pdb|1AKE|A  YYSKEAEAGNTKYAKVDGTPVAEVRADLEKILG
##          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 BLAST from BLAST
# blast <- blast.pdb(aa)
```

```
# hits <- plot(blast)
```

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

```
# Download related PDB files
# files <- get.pdb(hits$ pdb.id, path="pdb", split=TRUE, gzip=TRUE)
```

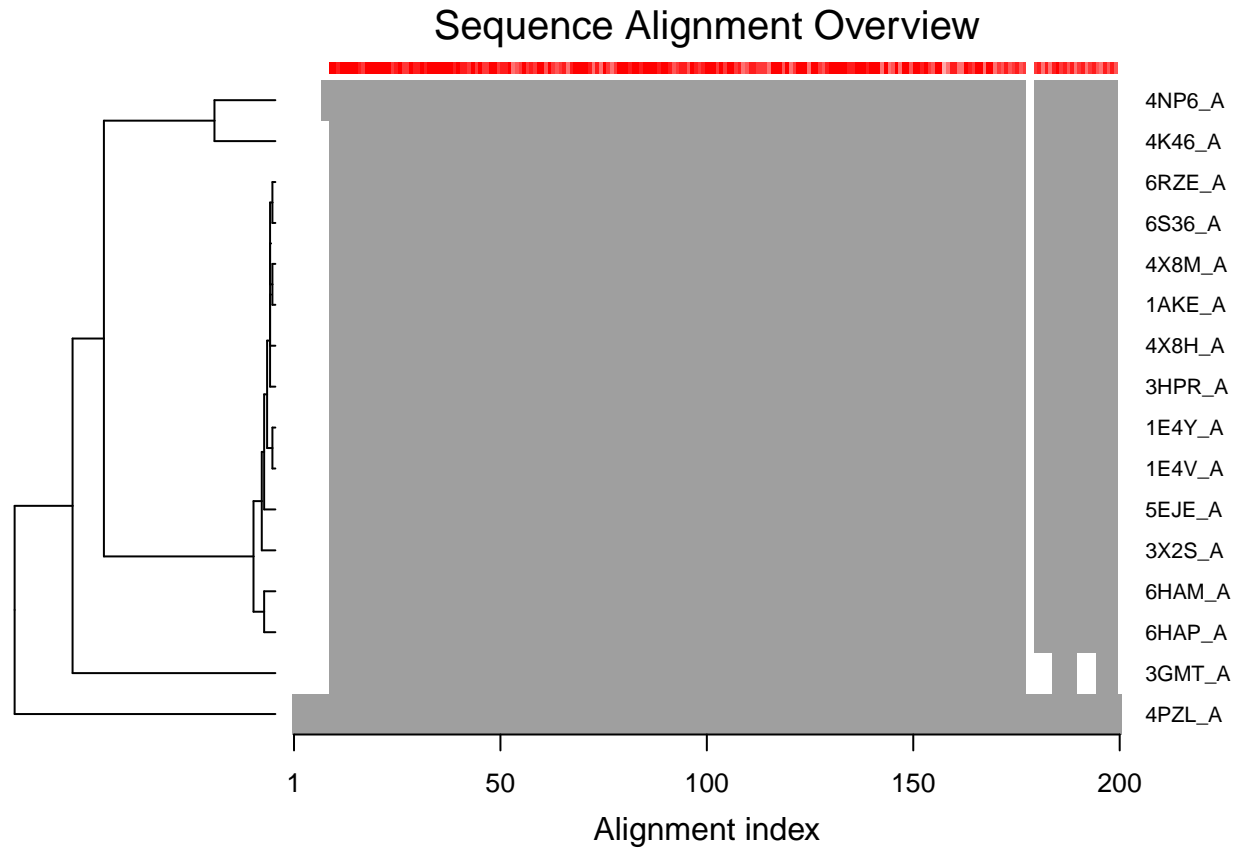
Multiple structure alignment

```
# pdbs <- pdbaln(files, fit = TRUE)
```

```
# Align related PDBs
# pdbs <- pdbaln(files, fit = TRUE)#, exefile="msa")
# save(pdbs, blast, file="mydata.RData")
load(file="mydata.RData")
```

```
# Vector containing PDB codes for figure axis
ids <- basename.pdb(pdbs$id)
```

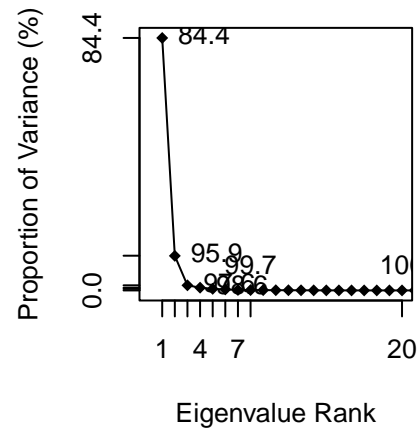
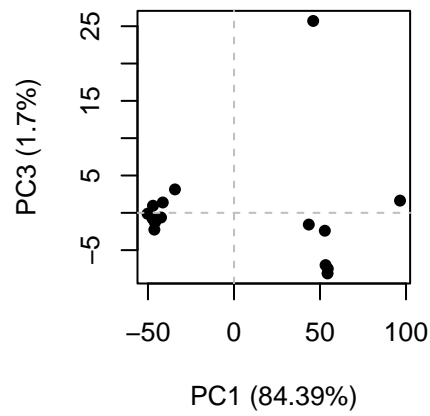
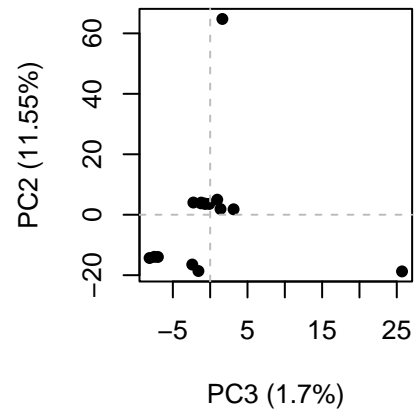
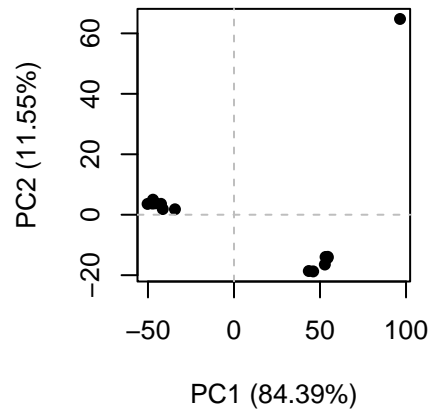
```
# Draw schematic alignment
plot(pdbs, labels=ids)
```



PCA

We will use the `bio3d::pca()` function which is designed for protein structure data.

```
# Perform PCA  
pc.xray <- pca(pdbbs)  
plot(pc.xray)
```



Make a trajectory visualization of the motion captured by the first Principal Component.

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

