class9: Structural Bioinformatics (pt1)

Ari_Fon (PID: A15390446)

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#The PDB database The PDB is the main repository for 3D structure data of biomolecules.

Here we explore it's composition. We obtained most recent stats from http://www.rcsb.org/stats/summary

```
tbl <- read.csv("Data Export Summary.csv", row.names = 1)
```

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

```
tot.method <- colSums(tbl)
round(tot.method/tot.method["Total"] * 100, digits = 3)</pre>
```

##	X.ray	NMR	EM Multi	ple.methods
##	87.197	7.284	5.354	0.106
##	Neutron	Other	Total	
##	0.039	0.020	100.000	

X.Ray is 87.197% and EM is 5.354%

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

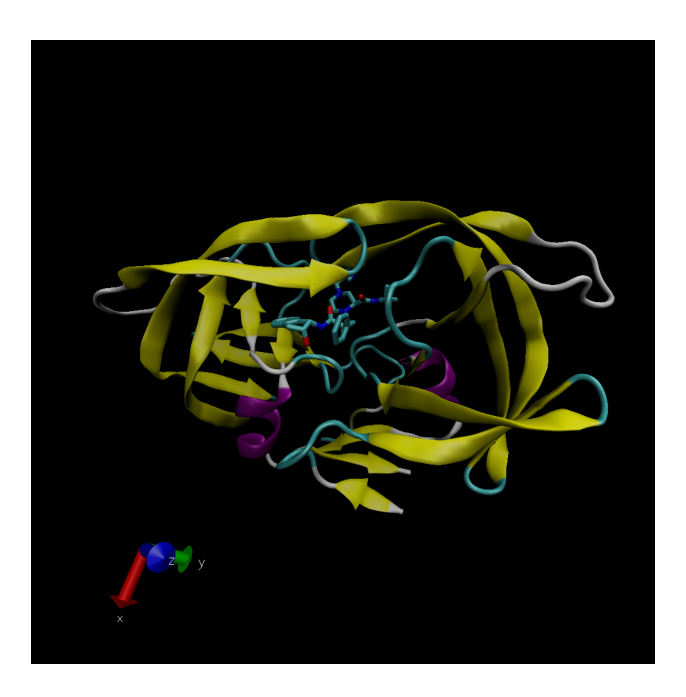
```
ans <- tbl$Total[1]/sum(tbl$Total)*100
round(ans, 3)</pre>
```

[1] 87.27

The answer to this question is 87.27 % of total structures

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?

Here is a VMD generated image of HIV-protease, PBD code: 1hsg



Bio3D package for structural bioinformatics

We will load the bio3d package

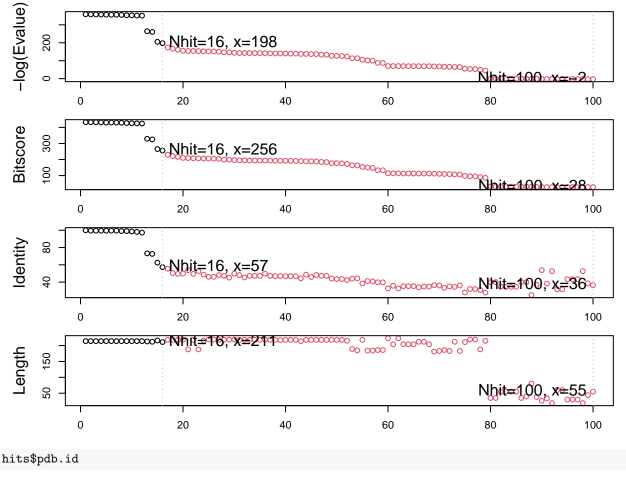
```
library(bio3d)
pdb <- read.pdb("1hsg")

## Note: Accessing on-line PDB file
pdb</pre>
```

```
##
   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
head(pdb$atom)
    type eleno elety alt resid chain resno insert
                                                        Х
                                                               у
                   N <NA>
                            PRO
                                              <NA> 29.361 39.686 5.862 1 38.10
## 1 ATOM
             1
                                    Α
                                          1
                            PRO
## 2 ATOM
             2
                  CA <NA>
                                          1 <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
                                         1 <NA> 28.600 38.302 3.676 1 43.40
                   O <NA>
                            PRO
## 4 ATOM
             4
                                    Α
                                   A 1 <NA> 30.508 37.541 6.342 1 37.87
## 5 ATOM
             5
                  CB <NA>
                            PRO
             6
                  CG <NA>
                            PRO
                                   A 1 <NA> 29.296 37.591 7.162 1 38.40
## 6 ATOM
    segid elesy charge
##
## 1 <NA>
             N
                 <NA>
## 2 <NA>
              C
                  <NA>
              C <NA>
## 3 <NA>
## 4 <NA>
              O <NA>
## 5
     <NA>
              С
                  <NA>
              С
                  <NA>
## 6 <NA>
Extract the sequence for ADK:
aa <- get.seq("1ake_A")</pre>
## Warning in get.seq("lake A"): Removing existing file: seqs.fasta
## Fetching... Please wait. Done.
##
                                                                          60
              MRIILLGAPGAGKGTQAQFIMEKYGIPQISTGDMLRAAVKSGSELGKQAKDIMDAGKLVT
## pdb|1AKE|A
##
```

##

```
## pdb|1AKE|A DELVIALVKERIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFDVPDELIVDRI
              61
##
## pdb|1AKE|A VGRRVHAPSGRVYHVKFNPPKVEGKDDVTGEELTTRKDDQEETVRKRLVEYHQMTAPLIG
                  . . .
##
             181
## pdb|1AKE|A YYSKEAEAGNTKYAKVDGTKPVAEVRADLEKILG
##
## Call:
   read.fasta(file = outfile)
##
##
## Class:
##
   fasta
##
## Alignment dimensions:
## 1 sequence rows; 214 position columns (214 non-gap, 0 gap)
## + attr: id, ali, call
blast <- blast.pdb(aa)</pre>
## Searching ... please wait (updates every 5 seconds) RID = OS1ZC3T401R
## Reporting 100 hits
hits <- plot(blast)</pre>
##
    * Possible cutoff values: 197 -3
##
              Yielding Nhits: 16 100
##
   * Chosen cutoff value of: 197
##
##
             Yielding Nhits: 16
```

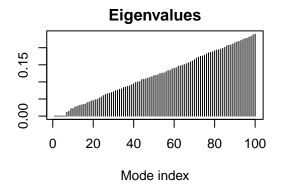


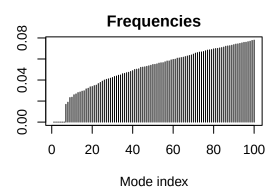
```
## [1] "1AKE_A" "4X8M_A" "6S36_A" "6RZE_A" "4X8H_A" "3HPR_A" "1E4V_A" "5EJE_A" 
## [9] "1E4Y_A" "3X2S_A" "6HAP_A" "6HAM_A" "4K46_A" "4NP6_A" "3GMT_A" "4PZL_A"
```

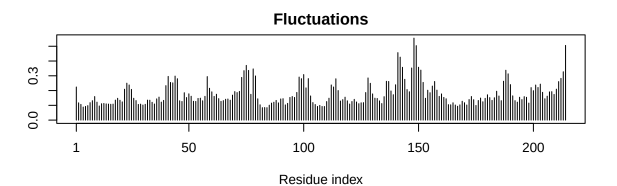
Normal mode analysis (NMA)

```
pdb <- read.pdb("1ake")</pre>
     Note: Accessing on-line PDB file
##
##
      PDB has ALT records, taking A only, rm.alt=TRUE
pdb
##
##
    Call: read.pdb(file = "1ake")
##
##
      Total Models#: 1
##
        Total Atoms#: 3804, XYZs#: 11412 Chains#: 2 (values: A B)
##
##
        Protein Atoms#: 3312 (residues/Calpha atoms#: 428)
        Nucleic acid Atoms#: 0 (residues/phosphate atoms#: 0)
##
```

```
##
##
        Non-protein/nucleic Atoms#: 492 (residues: 380)
##
        Non-protein/nucleic resid values: [ AP5 (2), HOH (378) ]
##
##
      Protein sequence:
##
         MRIILLGAPGAGKGTQAQFIMEKYGIPQISTGDMLRAAVKSGSELGKQAKDIMDAGKLVT
##
         DELVIALVKERIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFDVPDELIVDRI
         VGRRVHAPSGRVYHVKFNPPKVEGKDDVTGEELTTRKDDQEETVRKRLVEYHQMTAPLIG
##
##
         YYSKEAEAGNTKYAKVDGTKPVAEVRADLEKILGMRIILLGAPGA...<cut>...KILG
##
## + attr: atom, xyz, seqres, helix, sheet,
           calpha, remark, call
##
Trim to chain A only.
chain <- trim.pdb(pdb, chain= "A")</pre>
chain
##
##
    Call: trim.pdb(pdb = pdb, chain = "A")
##
##
      Total Models#: 1
        Total Atoms#: 1954, XYZs#: 5862 Chains#: 1 (values: A)
##
##
##
        Protein Atoms#: 1656 (residues/Calpha atoms#: 214)
##
        Nucleic acid Atoms#: 0 (residues/phosphate atoms#: 0)
##
##
        Non-protein/nucleic Atoms#: 298 (residues: 242)
##
        Non-protein/nucleic resid values: [ AP5 (1), HOH (241) ]
##
##
      Protein sequence:
##
         \tt MRIILLGAPGAGKGTQAQFIMEKYGIPQISTGDMLRAAVKSGSELGKQAKDIMDAGKLVT
##
         DELVIALVKERIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFDVPDELIVDRI
##
         VGRRVHAPSGRVYHVKFNPPKVEGKDDVTGEELTTRKDDQEETVRKRLVEYHQMTAPLIG
         YYSKEAEAGNTKYAKVDGTKPVAEVRADLEKILG
##
## + attr: atom, helix, sheet, seqres, xyz,
##
           calpha, call
Run a bioinformatics method to predict the flexibility and "functional motions" of this protein chain.
modes <- nma(chain)</pre>
   Building Hessian...
                             Done in 0.014 seconds.
                                 Done in 0.291 seconds.
## Diagonalizing Hessian...
plot(modes)
```







```
m7 <- mktrj.nma(modes, mode=7, file="mode_7.pdb")

pdb <- read.pdb("1ake")

## Note: Accessing on-line PDB file

## Warning in get.pdb(file, path = tempdir(), verbose = FALSE): /var/folders/66/

## q1kfbssd0fq0rdf6rp3vpn7c0000gn/T//Rtmpd7zg9k/1ake.pdb exists. Skipping download

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

chain <- trim.pdb(pdb, chain="A")

modes <-nma(chain)

## Building Hessian... Done in 0.013 seconds.

## Diagonalizing Hessian... Done in 0.285 seconds.

mktrj.nma(modes, mode=7, file="mode_7.pdb")
```

