Lab 09

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PDB database composition statistics

```
pdbstats <- read.csv("Data Export Summary.csv", row.names = 1)</pre>
```

Q1:The percentage of structures solved by X-ray and electron microscopy is:

```
pdbstats[,] <- lapply(pdbstats, function(x) as.numeric(gsub(",", "", x)))
total = sum(pdbstats$Total)
percentage = round(c(sum(pdbstats$X.ray),sum(pdbstats$EM)) / total *100, 2)</pre>
```

Based on the answer, there are 85.52% from X-ray structures and 7.48% from electron microscopy.

Q2: The proportion of protein structures in PDB:

```
percentage = round(sum(pdbstats[1,7]) / total *100, 2)
```

Based on the answer, there are 86.81% of proteins (only) in PDB.

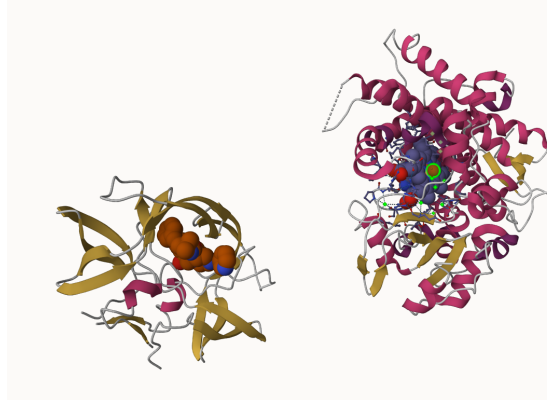
Q3: Based on the result from PDB website, there are 4929 structures of HIV protease structures in current PDB.

Visualizing the HIV-1 protease

Q4: The water molecules are being simplified for a better visualization of the protein itself.

Q5: The critical conserved water molecule has the residue number of 509, which is in the middle of Photoporphyrin IX.

Q6:indinavir or larger ligands and substranst might entering the binding site through the gap



between the two proteins.

Introduction to Bio3D

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

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)</pre>
```

```
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
Q7: Number of amino acid residues on this PDB is 198.
Q8: The non-protein residues are HOH and MK1.
Q9: The number of protein chains in this structure is 2.
  attributes(pdb)
$names
[1] "atom"
            "xyz"
                      "seqres" "helix" "sheet" "calpha" "remark" "call"
$class
[1] "pdb" "sse"
  head(pdb$atom)
 type eleno elety alt resid chain resno insert
                                                                   z o
1 ATOM
                                 A 1 <NA> 29.361 39.686 5.862 1 38.10
          1
                N < NA >
                          PRO
2 ATOM
          2
               CA <NA>
                         PRO
                                 Α
                                       1 <NA> 30.307 38.663 5.319 1 40.62
3 ATOM
               C <NA>
                         PRO
                                       1 <NA> 29.760 38.071 4.022 1 42.64
          3
                                 Α
4 ATOM
          4
                O <NA>
                         PRO
                                 Α
                                       1 <NA> 28.600 38.302 3.676 1 43.40
                                        1 <NA> 30.508 37.541 6.342 1 37.87
5 ATOM
               CB <NA>
                          PRO
          5
                                 Α
6 ATOM
               CG <NA>
                          PRO
                                            <NA> 29.296 37.591 7.162 1 38.40
 segid elesy charge
1 <NA>
                <NA>
           N
```

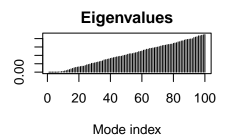
```
2 <NA> C <NA>
3 <NA> C <NA>
4 <NA> O <NA>
5 <NA> C <NA>
C <NA>
```

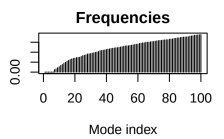
Predicting functional motions of a single structure

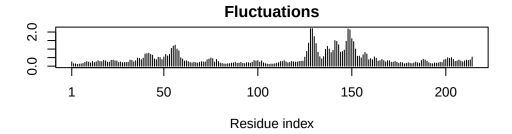
```
adk <- read.pdb("6s36")
 Note: Accessing on-line PDB file
  PDB has ALT records, taking A only, rm.alt=TRUE
  adk
Call: read.pdb(file = "6s36")
  Total Models#: 1
    Total Atoms#: 1898, XYZs#: 5694 Chains#: 1 (values: A)
    Protein Atoms#: 1654 (residues/Calpha atoms#: 214)
    Nucleic acid Atoms#: 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
  m <-nma(adk)
```

Building Hessian... Done in 0.013 seconds. Diagonalizing Hessian... Done in 0.276 seconds.

plot(m)







mktrj(m, file="adk_m7.pdb")

Comparative structure analysis of Adenylate Kinase

#install.packages("bio3d")
#install.packages("devtools")
#install.packages("BiocManager")

#BiocManager::install("msa")
#devtools::install_bitbucket("Grantlab/bio3d-view")

Q10: package msa is found only on BioConductor not CRAN.

Q11: the bio3d-view is not found in either BioConductor or CRAN.

Q12: It would be True that funcitons can be used to install packages from GitHub and BitBucket.

```
library(bio3d)
  aa <- get.seq("1ake_A")</pre>
Warning in get.seq("lake_A"): Removing existing file: seqs.fasta
Fetching... Please wait. Done.
  aa
                                                                            60
             MRIILLGAPGAGKGTQAQFIMEKYGIPQISTGDMLRAAVKSGSELGKQAKDIMDAGKLVT
pdb | 1AKE | A
            61
                                                                            120
             DELVIALVKERIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFDVPDELIVDRI
pdb | 1AKE | A
                                                                            120
           121
                                                                            180
pdb|1AKE|A
             VGRRVHAPSGRVYHVKFNPPKVEGKDDVTGEELTTRKDDQEETVRKRLVEYHQMTAPLIG
           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
Q13: The sequence in total is 214 aa long.
```

```
# Blast or hmmer search
  #b <- blast.pdb(aa)</pre>
  # Plot a summary of search results
  #hits <- plot(b)</pre>
  # List out some 'top hits'
  #head(hits$pdb.id)
  hits <- NULL
  hits$pdb.id <- c('1AKE_A','6S36_A','6RZE_A','3HPR_A','1E4V_A','5EJE_A','1E4Y_A','3X2S_A','
  # 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$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/6S36.pdb.gz exists. Skipping download
Warning in get.pdb(hits$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

1		
	I	0%
 =====	I	8%
 ==========	1	15%
 ===================================	I	23%
 ===================================	I	31%
 ===================================	I	38%
ı ====================================	1	46%
' ======= !	I	54%
' ======== !		62%
ı ====================================	1	69%
 	I	77%
 ===================================	1	85%
l .		

```
92%
  ______
  |-----| 100%
  # Align releated PDBs
  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
            name: pdbs/split_chain/1AKE_A.pdb
pdb/seq: 1
  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
pdb/seq: 3
            name: pdbs/split_chain/6RZE_A.pdb
  PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 4
            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: 5

```
pdb/seq: 6
             name: pdbs/split_chain/5EJE_A.pdb
   PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 7
             name: pdbs/split_chain/1E4Y_A.pdb
pdb/seq: 8
             name: pdbs/split_chain/3X2S_A.pdb
             name: pdbs/split_chain/6HAP_A.pdb
pdb/seq: 9
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
              name: pdbs/split_chain/3GMT_A.pdb
pdb/seq: 12
              name: pdbs/split_chain/4PZL_A.pdb
pdb/seq: 13
  # Vector containing PDB codes for figure axis
  ids <- basename.pdb(pdbs$id)</pre>
  # Draw schematic alignment
  #plot(pdbs, labels=ids) # Commented due to margin errors
  anno <- pdb.annotate(ids)</pre>
  unique(anno$source)
```

- [1] "Escherichia coli"
- [2] "Escherichia coli K-12"
- [3] "Escherichia coli 0139:H28 str. E24377A"
- [4] "Escherichia coli str. K-12 substr. MDS42"
- [5] "Photobacterium profundum"
- [6] "Burkholderia pseudomallei 1710b"
- [7] "Francisella tularensis subsp. tularensis SCHU S4"

anno

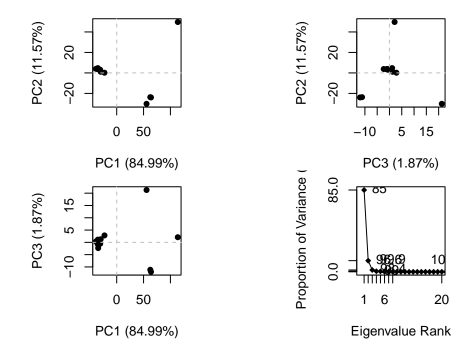
	structureId	chainId	macromoleculeType	chainLength	experimentalTechnique
1AKE_A	1AKE	A	Protein	214	X-ray
6S36_A	6S36	A	Protein	214	X-ray
6RZE_A	6RZE	A	Protein	214	X-ray
3HPR_A	3HPR	A	Protein	214	X-ray
1E4V_A	1E4V	A	Protein	214	X-ray
5EJE_A	5EJE	A	Protein	214	X-ray
1E4Y_A	1E4Y	Α	Protein	214	X-ray
$3X2S_A$	3X2S	A	Protein	214	X-ray

```
6HAP_A
              6HAP
                                                        214
                                                                             X-ray
                          Α
                                       Protein
6HAM_A
              6HAM
                          Α
                                       Protein
                                                        214
                                                                             X-ray
4K46_A
              4K46
                                                        214
                                                                             X-ray
                          Α
                                       Protein
3GMT_A
              3GMT
                                                        230
                                                                             X-ray
                          Α
                                       Protein
4PZL A
              4PZL
                          Α
                                       Protein
                                                        242
                                                                             X-ray
       resolution
                         scopDomain
                                                                              pfam
1AKE A
             2.00 Adenylate kinase Adenylate kinase, active site lid (ADK lid)
6S36_A
             1.60
                               <NA> Adenylate kinase, active site lid (ADK_lid)
6RZE A
             1.69
                               <NA> Adenylate kinase, active site lid (ADK lid)
                                <NA> Adenylate kinase, active site lid (ADK_lid)
3HPR_A
             2.00
             1.85 Adenylate kinase Adenylate kinase, active site lid (ADK_lid)
1E4V_A
             1.90
                                <NA> Adenylate kinase, active site lid (ADK_lid)
5EJE_A
                   Adenylate kinase Adenylate kinase, active site lid (ADK_lid)
1E4Y_A
             1.85
             2.80
                                <NA> Adenylate kinase, active site lid (ADK_lid)
3X2S_A
             2.70
                                <NA> Adenylate kinase, active site lid (ADK_lid)
6HAP_A
6HAM_A
             2.55
                               <NA> Adenylate kinase, active site lid (ADK_lid)
4K46_A
             2.01
                               <NA> Adenylate kinase, active site lid (ADK_lid)
3GMT_A
             2.10
                               <NA> Adenylate kinase, active site lid (ADK_lid)
             2.10
                               <NA> Adenylate kinase, active site lid (ADK_lid)
4PZL_A
                ligandId
1AKE A
                     AP5
6S36 A CL (3), NA, MG (2)
6RZE_A
          NA (3),CL (2)
3HPR_A
                     AP5
1E4V_A
                     AP5
5EJE_A
                  AP5,CO
                     AP5
1E4Y_A
3X2S_A
         JPY (2), AP5, MG
6HAP_A
                     AP5
6HAM_A
                     AP5
4K46_A
            ADP, AMP, PO4
3GMT_A
                S04 (2)
4PZL_A
             CA, FMT, GOL
                                                                                 ligandName
                                                          BIS (ADENOSINE) -5'-PENTAPHOSPHATE
1AKE A
6S36 A
                                            CHLORIDE ION (3), SODIUM ION, MAGNESIUM ION (2)
                                                           SODIUM ION (3), CHLORIDE ION (2)
6RZE A
3HPR A
                                                          BIS (ADENOSINE) -5'-PENTAPHOSPHATE
1E4V_A
                                                          BIS (ADENOSINE) -5'-PENTAPHOSPHATE
5EJE A
                                         BIS(ADENOSINE)-5'-PENTAPHOSPHATE, COBALT (II) ION
                                                          BIS (ADENOSINE) -5'-PENTAPHOSPHATE
1E4Y_A
3X2S A N-(pyren-1-ylmethyl)acetamide (2),BIS(ADENOSINE)-5'-PENTAPHOSPHATE,MAGNESIUM ION
6HAP_A
                                                          BIS (ADENOSINE) -5'-PENTAPHOSPHATE
```

```
6HAM_A
                                                        BIS (ADENOSINE) -5'-PENTAPHOSPHATE
4K46_A
                         ADENOSINE-5'-DIPHOSPHATE, ADENOSINE MONOPHOSPHATE, PHOSPHATE ION
3GMT_A
                                                                          SULFATE ION (2)
4PZL_A
                                                        CALCIUM ION, FORMIC ACID, GLYCEROL
                                                  source
1AKE A
                                        Escherichia coli
6S36 A
                                        Escherichia coli
6RZE_A
                                        Escherichia coli
3HPR_A
                                  Escherichia coli K-12
1E4V_A
                                        Escherichia coli
                 Escherichia coli 0139:H28 str. E24377A
5EJE_A
1E4Y_A
                                        Escherichia coli
               Escherichia coli str. K-12 substr. MDS42
3X2S_A
                 Escherichia coli 0139:H28 str. E24377A
6HAP_A
6HAM_A
                                   Escherichia coli K-12
4K46_A
                               Photobacterium profundum
3GMT_A
                        Burkholderia pseudomallei 1710b
4PZL_A Francisella tularensis subsp. tularensis SCHU S4
1AKE A STRUCTURE OF THE COMPLEX BETWEEN ADENYLATE KINASE FROM ESCHERICHIA COLI AND THE INHIB
6S36 A
6RZE A
3HPR_A
1E4V_A
5EJE_A
                                                                                          Crys
1E4Y_A
3X2S_A
6HAP_A
6HAM_A
4K46_A
3GMT_A
4PZL_A
                                                                                      The crys
                                                      citation rObserved
                                                                            rFree
1AKE_A
                       Muller, C.W., et al. J Mol Biol (1992)
                                                                 0.19600
                                                                               NA
                        Rogne, P., et al. Biochemistry (2019)
6S36 A
                                                                 0.16320 0.23560
                        Rogne, P., et al. Biochemistry (2019)
6RZE A
                                                                 0.18650 0.23500
       Schrank, T.P., et al. Proc Natl Acad Sci U S A (2009)
3HPR A
                                                                 0.21000 0.24320
1E4V A
                         Muller, C.W., et al. Proteins (1993)
                                                                 0.19600
5EJE_A Kovermann, M., et al. Proc Natl Acad Sci U S A (2017)
                                                                 0.18890 0.23580
1E4Y_A
                         Muller, C.W., et al. Proteins (1993)
                                                                 0.17800
3X2S_A
                      Fujii, A., et al. Bioconjug Chem (2015)
                                                                 0.20700 0.25600
                     Kantaev, R., et al. J Phys Chem B (2018)
6HAP_A
                                                                 0.22630 0.27760
6HAM_A
                     Kantaev, R., et al. J Phys Chem B (2018)
                                                                 0.20511 0.24325
```

```
4K46_A
                          Cho, Y.-J., et al. To be published
                                                                  0.17000 0.22290
3GMT_A Buchko, G.W., et al. Biochem Biophys Res Commun (2010)
                                                                  0.23800 0.29500
                             Tan, K., et al. To be published
4PZL_A
                                                                  0.19360 0.23680
         rWork spaceGroup
1AKE_A 0.19600
               P 21 2 21
6S36_A 0.15940
                  C 1 2 1
6RZE_A 0.18190
                  C 1 2 1
3HPR_A 0.20620
                P 21 21 2
1E4V_A 0.19600
                P 21 2 21
5EJE_A 0.18630
                P 21 2 21
1E4Y_A 0.17800
                 P 1 21 1
3X2S_A 0.20700 P 21 21 21
6HAP_A 0.22370
                  I 2 2 2
6HAM_A 0.20311
                     P 43
4K46_A 0.16730 P 21 21 21
                 P 1 21 1
3GMT_A 0.23500
4PZL_A 0.19130
                     P 32
```

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>
```

