Class 10

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```
string <- c("10", "100", 1, "1,000")
  as.numeric(string)+1
Warning: NAs introduced by coercion
[1] 11 101
              2 NA
     Q. Write a function to fix this non numeric table
  x <- string
  as.numeric(gsub(",","", x))
[1]
      10 100
                  1 1000
  data.file <- "Data Export Summary.csv"</pre>
  stats <- read.csv(data.file, row.names=1)</pre>
  head(stats)
                                            NMR Multiple.methods Neutron Other
                           X.ray
                                      EM
                         158,844 11,759 12,296
Protein (only)
                                                              197
                                                                        73
                                                                              32
Protein/Oligosaccharide
                           9,260
                                                                8
                                                                         1
                                                                               0
                                  2,054
                                             34
                                                                7
                                                                         0
                                                                               0
Protein/NA
                           8,307
                                   3,667
                                            284
Nucleic acid (only)
                           2,730
                                     113
                                         1,467
                                                               13
                                                                         3
                                                                               1
                                                                         0
Other
                             164
                                                                               0
Oligosaccharide (only)
                              11
                           Total
Protein (only)
                         183,201
Protein/Oligosaccharide 11,357
```

```
Protein/NA 12,265
Nucleic acid (only) 4,327
Other 205
Oligosaccharide (only) 22
```

```
rm.comma <- function(x) {
   as.numeric(gsub(",", "", x))
}

statsn <- apply(stats, 2, rm.comma)
head(statsn)</pre>
```

	X.ray	EM	NMR	${\tt Multiple.methods}$	Neutron	Other	Total
[1,]	158844	11759	12296	197	73	32	183201
[2,]	9260	2054	34	8	1	0	11357
[3,]	8307	3667	284	7	0	0	12265
[4,]	2730	113	1467	13	3	1	4327
[5,]	164	9	32	0	0	0	205
[6,]	11	0	6	1	0	4	22

rownames(statsn) <- rownames(stats)
statsn</pre>

	X.ray	EM	NMR	Multiple.methods	Neutron	Other
Protein (only)	158844	11759	12296	197	73	32
Protein/Oligosaccharide	9260	2054	34	8	1	0
Protein/NA	8307	3667	284	7	0	0
Nucleic acid (only)	2730	113	1467	13	3	1
Other	164	9	32	0	0	0
Oligosaccharide (only)	11	0	6	1	0	4
	Total					
Protein (only)	183201					
Protein/Oligosaccharide	11357					
Protein/NA	12265					
Nucleic acid (only)	4327					
Other	205					
Oligosaccharide (only)	22					

of all of the protein sequences on pdb(251,600,768), only 183201 have been experimentally determined

```
round(183201/251600768*100,2)
```

[1] 0.07

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

```
(sum(statsn[,1]) + sum(statsn[,2]))/sum(statsn[,7])*100
```

[1] 93.15962

```
totals <- apply(statsn, 2, sum)
round(totals/totals["Total"]*100,2)</pre>
```

X.ray	EM	NMR	Multiple.methods
84.83	8.33	6.68	0.11
Neutron	Other	Total	
0.04	0.02	100.00	

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

```
sum(statsn[1:3,7])/sum(statsn[,7])
```

[1] 0.9784556

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?

I found 211,377 examples.

#pdb format



Q4: Water molecules normally have 3 atoms. Why do we see just one atom per water molecule in this structure?

That is because the hydrogen is hidden, you can only see the oxygen. This is because the resolution is 2 angstrom, Hydrogen is only 1 A.

Q5: There is a critical "conserved" water molecule in the binding site. Can you identify this water molecule? What residue number does this water molecule have

Yes, HOH308

Q6: Generate and save a figure clearly showing the two distinct chains of HIV-protease along with the ligand. You might also consider showing the catalytic residues ASP 25 in each chain and the critical water (we recommend "Ball & Stick" for these side-chains). Add this figure to your Quarto document. Discussion Topic: Can you think of a way in which indinavir, or even larger ligands and substrates, could enter the binding site?

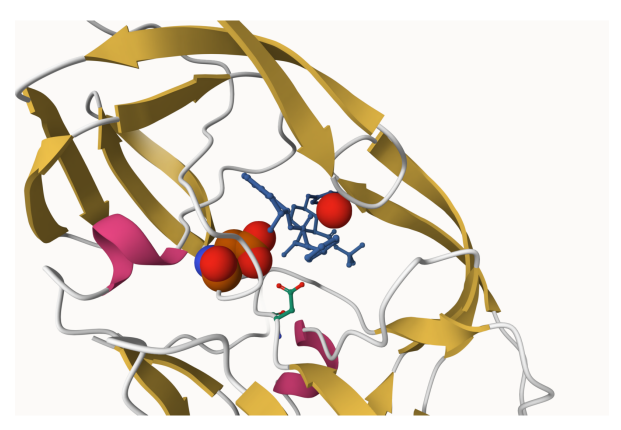


Figure 1: A lovely image

```
#3. Introduction to Bio3D in R
  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
     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
```

```
attributes(pdb)
$names
[1] "atom" "xyz"
                   "seqres" "helix" "sheet" "calpha" "remark" "call"
$class
[1] "pdb" "sse"
  head(pdb$atom$resid)
[1] "PRO" "PRO" "PRO" "PRO" "PRO" "PRO"
#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
```

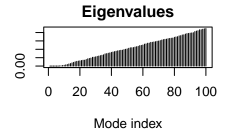
YYSKEAEAGNTKYAKVDGTKPVAEVRADLEKILG

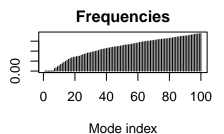
+ attr: atom, xyz, seqres, helix, sheet, calpha, remark, call

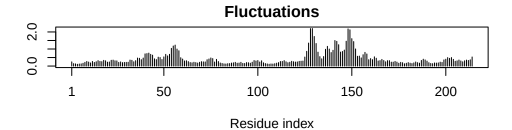
```
# Perform flexiblity prediction
m <- nma(adk)</pre>
```

Building Hessian... Done in 0.015 seconds. Diagonalizing Hessian... Done in 0.272 seconds.

plot(m)







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

- Q10. Which of the packages above is found only on BioConductor and not CRAN? $_{\rm msa}$
 - Q11. Which of the above packages is not found on BioConductor or CRAN?:

bio3d-view

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

TRUE

Search and retrieve ADK structures

```
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
                                                                       60
                                                                       120
pdb|1AKE|A
            DELVIALVKERIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFDVPDELIVDRI
                                                                       120
           121
                                                                       180
pdb|1AKE|A VGRRVHAPSGRVYHVKFNPPKVEGKDDVTGEELTTRKDDQEETVRKRLVEYHQMTAPLIG
          121
                                                                       180
          181
                                             214
pdb|1AKE|A
            YYSKEAEAGNTKYAKVDGTKPVAEVRADLEKILG
          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. How many amino acids are in this sequence, i.e. how long is this sequence?
```

Search and retrieve ADK structures

```
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

į	1	0%
 ====	1	8%
 =======	1	15%
 ===================================	I	23%
 	ı	31%
 	1	38%
 	·	46%
į	ı	
		54%
 	I	62%
	I	69%
	1	77%

```
85%
                                                                         92%
              #Align and superpose structures
  # 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 has ALT records, taking A only, rm.alt=TRUE
            name: pdbs/split_chain/6S36_A.pdb
pdb/seq: 2
   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
             name: pdbs/split chain/5EJE A.pdb
pdb/seq: 6
   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
pdb/seq: 9
             name: pdbs/split_chain/6HAP_A.pdb
pdb/seq: 10
              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: 11
   PDB has ALT records, taking A only, rm.alt=TRUE
              name: pdbs/split_chain/3GMT_A.pdb
pdb/seq: 12
pdb/seq: 13
              name: pdbs/split_chain/4PZL_A.pdb
  # Vector containing PDB codes for figure axis
  ids <- basename.pdb(pdbs$id)</pre>
  # Draw schematic alignment
  #plot(pdbs, labels=ids)
#Annotate collected PDB structures
  anno <- pdb.annotate(ids)</pre>
  unique(anno$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
                                       Protein
                                                        214
                                                                              X-ray
                          Α
3HPR_A
               3HPR
                                                        214
                          Α
                                       Protein
                                                                              X-ray
1E4V_A
               1E4V
                          Α
                                       Protein
                                                        214
                                                                              X-ray
5EJE_A
                                                        214
               5EJE
                          Α
                                       Protein
                                                                              X-ray
1E4Y A
               1E4Y
                          Α
                                       Protein
                                                        214
                                                                              X-ray
3X2S A
               3X2S
                          Α
                                       Protein
                                                        214
                                                                              X-ray
6HAP A
               6HAP
                          Α
                                       Protein
                                                        214
                                                                              X-ray
6HAM A
               6HAM
                           Α
                                       Protein
                                                        214
                                                                              X-ray
4K46_A
               4K46
                                                        214
                          Α
                                       Protein
                                                                              X-ray
                                                        230
3GMT_A
               3GMT
                          Α
                                       Protein
                                                                              X-ray
               4PZL
                                                        242
4PZL_A
                          Α
                                       Protein
                                                                              X-ray
                                                                      ligandId
       resolution
                         scopDomain
                                                        pfam
1AKE_A
             2.00 Adenylate kinase Adenylate kinase (ADK)
                                                                            AP5
                                <NA> Adenylate kinase (ADK) CL (3),NA,MG (2)
             1.60
6S36_A
6RZE_A
             1.69
                                <NA> Adenylate kinase (ADK)
                                                                 NA (3),CL (2)
3HPR_A
             2.00
                                <NA> Adenylate kinase (ADK)
                                                                            AP5
1E4V_A
             1.85 Adenylate kinase Adenylate kinase (ADK)
                                                                            AP5
5EJE_A
             1.90
                                <NA> Adenylate kinase (ADK)
                                                                         AP5,CO
1E4Y_A
             1.85 Adenylate kinase Adenylate kinase (ADK)
                                                                            AP5
3X2S A
             2.80
                                <NA> Adenylate kinase (ADK)
                                                                JPY (2), AP5, MG
6HAP A
             2.70
                                <NA> Adenylate kinase (ADK)
                                                                            AP5
             2.55
                                <NA> Adenylate kinase (ADK)
6HAM A
                                                                            AP5
4K46_A
             2.01
                                <NA> Adenylate kinase (ADK)
                                                                   ADP, AMP, PO4
3GMT_A
             2.10
                                <NA> Adenylate kinase (ADK)
                                                                        S04 (2)
                                <NA> Adenylate kinase (ADK)
4PZL_A
             2.10
                                                                    CA, FMT, GOL
                                                                                  ligandName
1AKE_A
                                                           BIS (ADENOSINE) -5'-PENTAPHOSPHATE
                                             CHLORIDE ION (3), SODIUM ION, MAGNESIUM ION (2)
6S36_A
6RZE_A
                                                            SODIUM ION (3), CHLORIDE ION (2)
3HPR_A
                                                           BIS (ADENOSINE) -5'-PENTAPHOSPHATE
1E4V_A
                                                           BIS (ADENOSINE) -5'-PENTAPHOSPHATE
                                         BIS(ADENOSINE)-5'-PENTAPHOSPHATE, COBALT (II) ION
5EJE_A
1E4Y_A
                                                           BIS (ADENOSINE) -5'-PENTAPHOSPHATE
3X2S_A N-(pyren-1-ylmethyl)acetamide (2),BIS(ADENOSINE)-5'-PENTAPHOSPHATE,MAGNESIUM ION
                                                           BIS (ADENOSINE) -5'-PENTAPHOSPHATE
6HAP A
6HAM A
                                                           BIS (ADENOSINE) -5'-PENTAPHOSPHATE
                          ADENOSINE-5'-DIPHOSPHATE, ADENOSINE MONOPHOSPHATE, PHOSPHATE ION
4K46 A
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
5EJE_A
                 Escherichia coli 0139:H28 str. E24377A
1E4Y_A
                                       Escherichia coli
3X2S A
               Escherichia coli str. K-12 substr. MDS42
6HAP_A
                 Escherichia coli 0139:H28 str. E24377A
6HAM A
                                  Escherichia coli K-12
4K46_A
                               Photobacterium profundum
                        Burkholderia pseudomallei 1710b
3GMT_A
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
                       Muller, C.W., et al. J Mol Biol (1992)
1AKE_A
                                                                 0.19600
                                                                               NA
6S36_A
                        Rogne, P., et al. Biochemistry (2019)
                                                                 0.16320 0.23560
6RZE_A
                        Rogne, P., et al. Biochemistry (2019)
                                                                 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
                                                                              NA
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
6HAP_A
                     Kantaev, R., et al. J Phys Chem B (2018)
                                                                 0.22630 0.27760
                     Kantaev, R., et al. J Phys Chem B (2018)
6HAM A
                                                                 0.20511 0.24325
                          Cho, Y.-J., et al. To be published
4K46 A
                                                                 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
                                                                 0.19360 0.23680
4PZL A
         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 3GMT_A 0.23500 P 1 21 1 4PZL_A 0.19130 P 32
```

#Principal component analysis

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

