class10

Wanning Cui

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

```
read.csv("Data Export Summary.csv")
```

```
Molecular.Type
                                       EM
                                              NMR Multiple.methods Neutron Other
                             X.ray
           Protein (only) 158,844 11,759 12,296
                                                                197
                                                                         73
                                                                               32
2 Protein/Oligosaccharide
                             9,260
                                   2,054
                                                                  8
                                                                          1
                                                                                0
3
               Protein/NA
                             8,307
                                   3,667
                                                                  7
                                                                          0
                                                                                0
                                              284
                                      113 1,467
4
      Nucleic acid (only)
                                                                 13
                                                                          3
                             2,730
                                                                                1
5
                     Other
                               164
                                        9
                                               32
                                                                  0
                                                                          0
                                                                                0
  Oligosaccharide (only)
                                11
                                        0
                                                6
                                                                  1
                                                                          0
                                                                                4
```

Total 1 183,201

2 11,357

3 12,265

4 4,327

5 205

6 22

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

	X.ray	EM	NMR	Multiple.methods	Neutron	Other
Protein (only)	158,844	11,759	12,296	197	73	32
Protein/Oligosaccharide	9,260	2,054	34	8	1	0
Protein/NA	8,307	3,667	284	7	0	0
Nucleic acid (only)	2,730	113	1,467	13	3	1
Other	164	9	32	0	0	0
Oligosaccharide (only)	11	0	6	1	0	4

```
Total
Protein (only)
                         183,201
Protein/Oligosaccharide 11,357
Protein/NA
                          12,265
Nucleic acid (only)
                           4,327
Other
                             205
Oligosaccharide (only)
                              22
  string <- c("10","100","1", "1,000")
  numeric_vector <- as.numeric(gsub(",", "", string)) + 1</pre>
  x <- string
  as.numeric(gsub(",", "", x))
Г1]
      10 100
                 1 1000
  rm.comma <- function(x){</pre>
    as.numeric(gsub(",", "", x))
  }
  pdbstats <- apply(stats,2,rm.comma)</pre>
  rownames(pdbstats) <- rownames(stats)</pre>
  pdbstats
                                         NMR Multiple.methods Neutron Other
                          X.ray
                                   EM
                                                           197
                                                                    73
                                                                          32
Protein (only)
                         158844 11759 12296
Protein/Oligosaccharide
                           9260 2054
                                          34
                                                             8
                                                                     1
                                                                           0
                                                             7
                                                                     0
                                                                           0
Protein/NA
                           8307 3667
                                         284
Nucleic acid (only)
                           2730
                                  113 1467
                                                            13
                                                                     3
                                                                           1
                            164
                                          32
                                                             0
                                                                     0
                                                                           0
Other
                                    9
                                                                     0
                                                                           4
Oligosaccharide (only)
                             11
                                    0
                                           6
                                                             1
                          Total
Protein (only)
                         183201
Protein/Oligosaccharide 11357
Protein/NA
                          12265
Nucleic acid (only)
                           4327
Other
                            205
Oligosaccharide (only)
                             22
```

totals <- apply(pdbstats,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	

Q. Write a function to fix this non-numeric table

We can use the gsub() function.

Will add the rownames from the original wee table...

The percentage is 93.16

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

The main repository of structural data is the PDB. Let's examine what it contains. At the time of writing, there are 183,201 protein structures In UniProt, there are 251600,768 protein sequences.

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

[1] 0.07

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?

7434

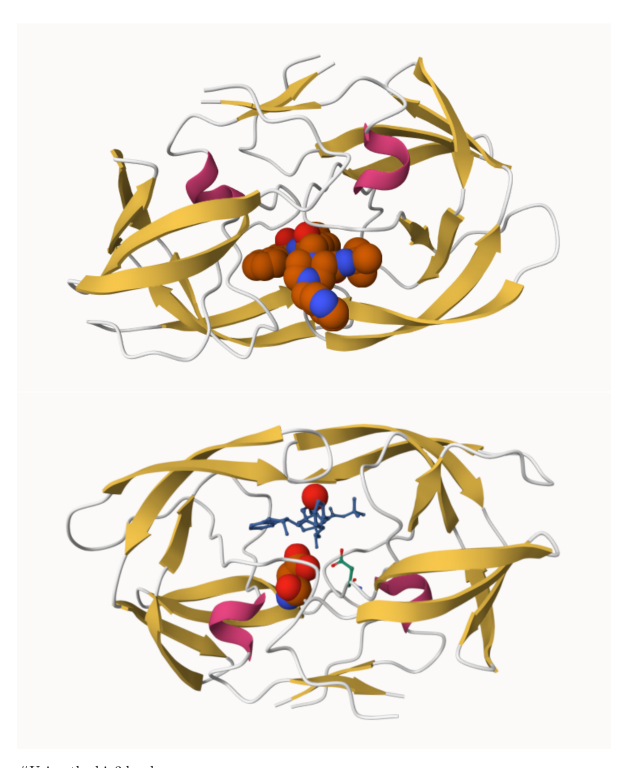
#2 Visualizing structure >Q4: Water molecules normally have 3 atoms. Why do we see just one atom per water molecule in this structure?

There are no hydrogen atoms observed in the structure. It's smaller than the 2A resolution.

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

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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.



#Using the bio3d pakcage

```
library(bio3d)
Warning: package 'bio3d' was built under R version 4.3.2
  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
  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
                                                       Х
                                                              у
1 ATOM
           1
                 N < NA >
                          PRO
                                  Α
                                        1
                                             <NA> 29.361 39.686 5.862 1 38.10
2 ATOM
           2
                CA <NA>
                          PRO
                                             <NA> 30.307 38.663 5.319 1 40.62
                                  Α
3 ATOM
                 C <NA>
                                        1 <NA> 29.760 38.071 4.022 1 42.64
                          PRO
                                        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
5 ATOM
           5
                CB <NA>
                          PRO
                                  Α
6 ATOM
                CG <NA>
                          PRO
                                            <NA> 29.296 37.591 7.162 1 38.40
  segid elesy charge
  <NA>
           N
                <NA>
1
  <NA>
            С
                <NA>
3
  <NA>
           С
                <NA>
  <NA>
            O <NA>
  <NA>
           C
                <NA>
  <NA>
           C
                <NA>
```

head(pdb\$atom\$resid)

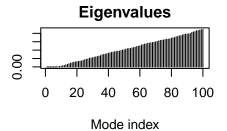
```
[1] "PRO" "PRO" "PRO" "PRO" "PRO" "PRO"
```

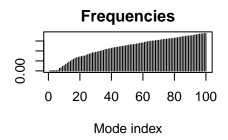
aa321(pdb\$atom\$resid[pdb\$calpha])

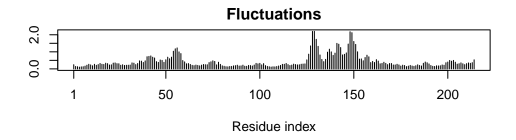
#Predicting functional motions of a single structure

Run a Normal Mode Analysis (NMA) - a bioinformatics method to predict functional motions.

```
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
  modes <- nma(adk)</pre>
Building Hessian...
                            Done in 0.03 seconds.
Diagonalizing Hessian...
                           Done in 0.28 seconds.
  plot(modes)
```







mktrj(modes,pdb=adk,file="adk_m7.pdb")