class10 structural bioinformatics p1

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the PDB database

Here we examine the size and composition of the main database of biomolecular structures - the PDB.

Get a CSV file from the PDB database and read it into R.

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

	X.ray	EM	NMR	Multiple.methods	Neutron	Other
Protein (only)	161,663	12,592	12,337	200	74	32
Protein/Oligosaccharide	9,348	2,167	34	8	2	0
Protein/NA	8,404	3,924	286	7	0	0
Nucleic acid (only)	2,758	125	1,477	14	3	1
Other	164	9	33	0	0	0
Oligosaccharide (only)	11	0	6	1	0	4
	Total					
Protein (only)	186,898					
Protein/Oligosaccharide	11,559					
Protein/NA	12,621					
Nucleic acid (only)	4,378					
Other	206					
Oligosaccharide (only)	22					

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

My pbdstats dataframe has numbers with commas in them. This may cause us problems.

```
sum(as.numeric(pdbstats$X.ray, pdbstats$EM))
```

Warning: NAs introduced by coercion

[1] NA

We found a function called $\mathtt{gsub}()$. I can use this to make a function that I can use for every column in the table.

```
commasum <- function(x) {</pre>
  sum(as.numeric(gsub(",", "", x)))
}
totals <- apply(pdbstats, 2, commasum)</pre>
totals
         X.ray
                               EM
                                                NMR Multiple.methods
        182348
                           18817
                                              14173
                                                                   230
       Neutron
                                              Total
                           Other
            79
                                             215684
                               37
round(totals/totals["Total"]*100, 2)
         X.ray
                               EM
                                                NMR Multiple.methods
         84.54
                            8.72
                                               6.57
                                                                 0.11
       Neutron
                           Other
                                              Total
          0.04
                            0.02
                                             100.00
```

85% were solved by x-ray and 9% by EM.

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

```
commasum(pdbstats["Protein (only)", "Total"])/totals["Total"]*100

Total
86.65362
87%
```

• Q. What fraction of Uniprot structures are represented in the PDB? 0.09%

```
215684/249751891*100
```

[1] 0.08635931

visualizing protein structure

We will learn the basics of Mol* (mol-star) homepage: https://molstar.org/viewer/ We will play with PDB 1HSG:

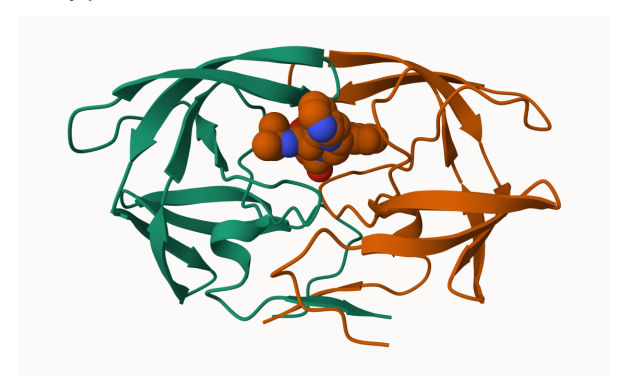


Figure 1: HIV-Pr with a bound inhibitor

back to R and working with PDB structures

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

Note: Accessing on-line PDB file
hiv

Call: read.pdb(file = "1hsg")</pre>
```

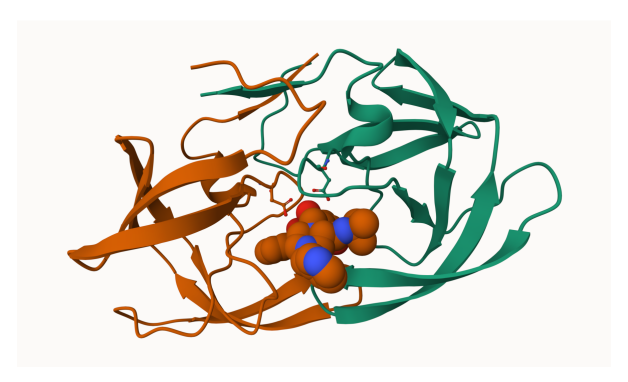


Figure 2: with ASP25 residues represented as ball-and-stick figures

```
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:

PQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGIGGFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNFPQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGIGGFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNF

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

head(hiv\$atom)

```
type eleno elety alt resid chain resno insert
                                                         X
                                                                 У
                                                                       z o
1 ATOM
           1
                  N <NA>
                           PRO
                                               <NA> 29.361 39.686 5.862 1 38.10
                                    Α
                                          1
2 ATOM
           2
                           PRO
                                               <NA> 30.307 38.663 5.319 1 40.62
                 CA <NA>
                                    Α
                                          1
3 ATOM
                 C <NA>
                                               <NA> 29.760 38.071 4.022 1 42.64
           3
                           PRO
                                    Α
                                          1
4 ATOM
                  O <NA>
                           PRO
                                               <NA> 28.600 38.302 3.676 1 43.40
                                          1
                                               <NA> 30.508 37.541 6.342 1 37.87
5 ATOM
           5
                 CB <NA>
                           PRO
                                    Α
                                          1
6 ATOM
           6
                 CG <NA>
                           PRO
                                          1
                                               <NA> 29.296 37.591 7.162 1 38.40
                                    Α
  segid elesy charge
  <NA>
                 <NA>
1
            N
2
  <NA>
            C
                 <NA>
3
  <NA>
            С
                 <NA>
  <NA>
            0
                 <NA>
   <NA>
            C
                 <NA>
            C
   <NA>
                 <NA>
```

pdbseq(hiv)

```
5
                     6
                       7
                            8
                                9 10 11 12 13 14 15 16 17 18
"P" "Q" "I" "T" "L" "W" "Q" "R" "P" "L" "V" "T" "I" "K" "I" "G" "G" "G" "L" "K"
21 22 23 24 25 26 27 28 29 30 31 32 33 34 35
                                                         36 37
                                                                  38
"E" "A" "L" "L" "D" "T" "G" "A" "D" "D" "T" "V" "L" "E" "E" "M" "S" "L" "P" "G"
41 42 43 44 45
                  46 47 48
                              49 50 51 52 53 54 55 56 57 58
"R" "W" "K" "P" "K" "M" "I" "G" "G" "I" "G" "G" "F" "I" "K" "V" "R" "O" "Y" "D"
                                      71
        63
           64
               65
                    66
                       67
                           68
                               69
                                   70
                                          72
                                              73
                                                  74
                                                      75
                                                          76
                                                              77
"Q" "I" "L" "I" "E" "I" "C" "G" "H" "K" "A" "I" "G" "T" "V" "L" "V" "G" "P" "T"
        83 84 85
                   86 87
                           88
                               89 90 91 92 93
                                                  94 95
                                                          96 97
"P" "V" "N" "T" "T" "G" "R" "N" "I," "I," "T" "G" "T" "G" "C" "T" "I," "N"
                               10 11
                                       12 13 14
                                                      16
                 6
                     7
                        8
                             9
                                                  15
                                                          17
                                                               18
                                                                  19
ייטיי יידיי יידיי
                                                                     "K" "E"
                   27
                                      32 33
    23
        24
            25
               26
                       28
                           29
                               30 31
                                              34
                                                   35
                                                       36
                                                          37
                                                               38
                                                                  39
"A" "L" "L" "D" "T" "G" "A" "D" "D" "T" "V" "L" "E" "E" "M" "S" "L" "P"
                                                                     "G" "R"
            45
               46
                    47
                       48
                           49
                               50 51
                                      52 53
                                              54
                                                   55
                                                      56
                                                           57
                                                               58
"W" "K" "P" "K" "M" "I" "G" "G" "I" "G" "G" "F" "I" "K" "V" "R" "Q" "Y"
                                                                     "ם" "ם"
62 63
       64 65 66 67
                       68 69
                               70 71 72
                                          73
                                              74
                                                  75 76
                                                          77
                                                              78
                                                                  79
                                                                      80
                                                                          81
"I" "L" "I" "E" "I" "C" "G" "H" "K" "A" "I" "G" "T" "V" "L" "V" "G" "P"
82 83 84 85 86 87 88 89 90 91 92 93 94 95 96 97
                                                              98
"V" "N" "I" "I" "G" "R" "N" "L" "L" "T" "Q" "I" "G" "C" "T" "L" "N" "F"
```

here we will do a normal mode analysis (NMA) to predict functional motions of a kinase protein

```
adk <- read.pdb("6s36")
 Note: Accessing on-line PDB file
  PDB has ALT records, taking A only, rm.alt=TRUE
  adk
       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
  modes <- nma(adk)
Building Hessian...
                           Done in 0.046 seconds.
Diagonalizing Hessian...
                           Done in 0.526 seconds.
  plot(modes)
```