Class 10: Structural Bioinformatics (pt1)

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1: Introduction to the RCSB Protein Data Bank (PDB)

The main repository of biomolecular structure data is called the PDB found at https://www.rcsb.org/

Let's see what this database contains. I went to PDB > Analyze > PDB Statistics > By Exp Method and Molecular Type.

```
pdbstats <- read.csv("Data Export Summary.csv")
pdbstats</pre>
```

	Molecular.Type	X.ray	EM	NMR	Multiple.methods	Neutron	Other
1	Protein (only)	169,563	16,774	12,578	208	81	32
2	Protein/Oligosaccharide	9,939	2,839	34	8	2	0
3	Protein/NA	8,801	5,062	286	7	0	0
4	Nucleic acid (only)	2,890	151	1,521	14	3	1
5	Other	170	10	33	0	0	0
6	Oligosaccharide (only)	11	0	6	1	0	4
	Total						

```
1 199,236
2 12,822
3 14,156
4 4,580
5 213
6 22
```

pdbstats\$X.ray

```
[1] "169,563" "9,939" "8,801" "2,890" "170" "11"
```

The numbers are in quotes, and they are , because the comma caused them to be characters. We cannot do calculation with character, so we need to convert them into numbers.

I can fix this by replacing "," for nothing "" with the sub() function

```
x <- pdbstats$X.ray
sum(as.numeric(sub(",", "", x)))</pre>
```

[1] 191374

Or I can use the **readr** package and the **read_csv()** function

```
library("readr")
pdbstats <- read_csv("Data Export Summary.csv")

Rows: 6 Columns: 8
-- Column specification ------
Delimiter: ","
chr (1): Molecular Type
dbl (3): Multiple methods, Neutron, Other
num (4): X-ray, EM, NMR, Total

i Use `spec()` to retrieve the full column specification for this data.
i Specify the column types or set `show_col_types = FALSE` to quiet this message.</pre>
```

pdbstats

#	A tibble: 6 x 8								
	`Molecular Type`	`X-ray`	EM	NMR	`Multiple	methods`	${\tt Neutron}$	Other	Total
	<chr></chr>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>		<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>
1	Protein (only)	169563	16774	12578		208	81	32	199236
2	Protein/Oligosacc~	9939	2839	34		8	2	0	12822
3	Protein/NA	8801	5062	286		7	0	0	14156
4	Nucleic acid (onl~	2890	151	1521		14	3	1	4580
5	Other	170	10	33		0	0	0	213
6	Oligosaccharide (~	11	0	6		1	0	4	22

I want to clean the column names so they are all lower case and don't have spaces in them.

colnames(pdbstats)

```
[1] "Molecular Type" "X-ray" "EM" "NMR"
[5] "Multiple methods" "Neutron" "Other" "Total"
```

library("janitor")

Attaching package: 'janitor'

The following objects are masked from 'package:stats':

chisq.test, fisher.test

```
df <- clean_names(pdbstats)
df</pre>
```

```
# A tibble: 6 x 8
 molecular_type
                          x_ray
                                    em
                                         nmr multiple_methods neutron other
                                                                               total
  <chr>
                          <dbl> <dbl> <dbl>
                                                         <dbl>
                                                                  <dbl> <dbl>
                                                                                <dbl>
1 Protein (only)
                         169563 16774 12578
                                                           208
                                                                     81
                                                                           32 199236
2 Protein/Oligosacchar~
                           9939
                                 2839
                                          34
                                                             8
                                                                      2
                                                                            0
                                                                               12822
                                                             7
3 Protein/NA
                           8801
                                 5062
                                         286
                                                                      0
                                                                            0
                                                                                14156
4 Nucleic acid (only)
                           2890
                                   151
                                        1521
                                                            14
                                                                      3
                                                                            1
                                                                                 4580
5 Other
                                    10
                                                             0
                                                                            0
                            170
                                          33
                                                                      0
                                                                                  213
6 Oligosaccharide (onl~
                             11
                                     0
                                           6
                                                              1
                                                                      0
                                                                                   22
```

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

Total number of X-ray structures

```
sum(df$x_ray)
```

[1] 191374

Total number of structures

```
sum(df$total)
```

[1] 231029

Percent of X-ray structures.

```
sum(df$x_ray)/sum(df$total) *100
```

[1] 82.83549

Percent of EM structures

```
sum(df\$em) / sum(df\$total) *100
```

[1] 10.75017

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

Percent of protein structure

```
sum(df$total[1:3]) / sum(df$total) *100
```

[1] 97.91585

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?

24,695

2. Visualizing the HIV-1 protease structure using Mol*

The main Mol* homepage at https://molstar.org/viewer/ We can input our own PDB files or just give it a PDB databse accession code (4 letter PDB code)



Figure 1: molecular view of 1 HSG

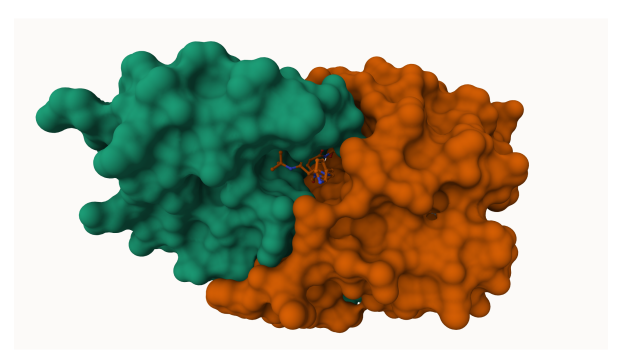


Figure 2: Molecular surface representation of the polymer

The important role of water

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

Because there are a lot of water molecules around the protein. If we show all the atoms for every water molecular, they would cover up the polymer and ligand.

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?

It has residue number 308.

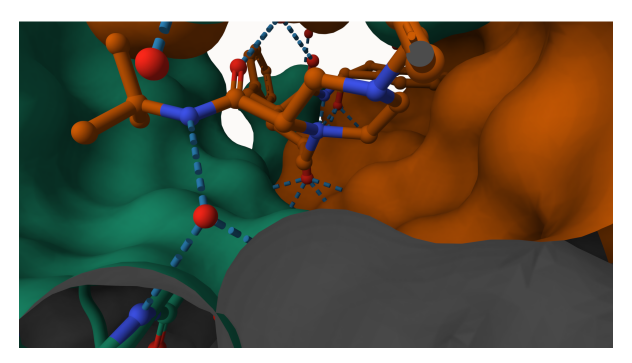


Figure 3: The conserved water

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.

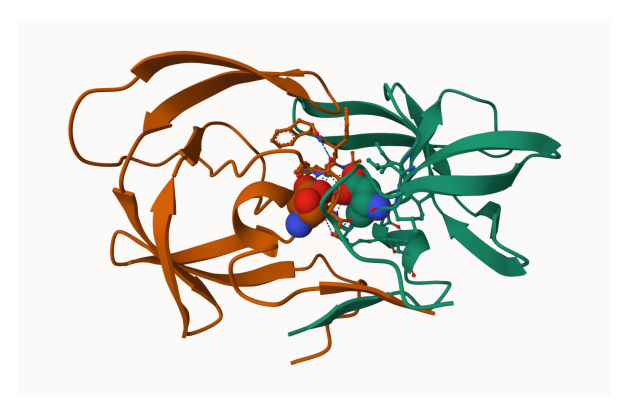


Figure 4: The important ASP25 amino acids in chain A and B

3. Introduction to Bio3D in R

We can use the **bio3d** package for structural bioinformatics to read PDB data into R

```
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
     Q7: How many amino acid residues are there in this pdb object?
length(pdbseq(pdb))
[1] 198
     Q8: Name one of the two non-protein residues?
MK1
     Q9: How many protein chains are in this structure?
2 chains, A and B
Looking at the pdb project
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
                                                       X
                                                               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
                                         1
                                             <NA> 30.307 38.663 5.319 1 40.62
                                   Α
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
5 ATOM
                          PRO
                                             <NA> 30.508 37.541 6.342 1 37.87
                CB <NA>
                                   Α
                                             <NA> 29.296 37.591 7.162 1 38.40
6 ATOM
           6
                CG <NA>
                          PRO
  segid elesy charge
  <NA>
            N
                <NA>
1
2
  <NA>
            С
                <NA>
3 <NA>
            С
                <NA>
4 <NA>
            0
                <NA>
5 <NA>
            С
                <NA>
            С
                <NA>
6 <NA>
```

Let's try a new function not yet in the bio3d package. It requires the **r3dmol** and **shiny** package that we need to install with **install.packages("r3dmol")** and **install.packages("shiny")**.

```
library(r3dmol)
source("https://tinyurl.com/viewpdb")
#view.pdb(pdb, backgroundColor = "lightblue")
```

4. Predicting functional dynamics

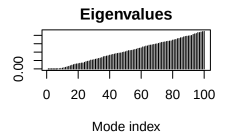
We can use the nma() function in bio3d to predict the large-scale functional motions of biomolecules.

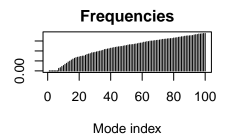
```
adk <- read.pdb("6s36")

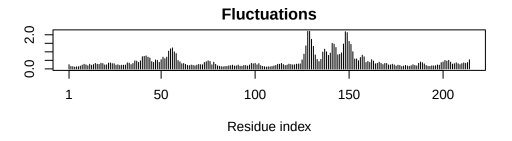
Note: Accessing on-line PDB file
   PDB has ALT records, taking A only, rm.alt=TRUE</pre>
```

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:
      \tt MRIILLGAPGAGKGTQAQFIMEKYGIPQISTGDMLRAAVKSGSELGKQAKDIMDAGKLVT
      DELVIALVKERIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFDVPDELIVDKI
      VGRRVHAPSGRVYHVKFNPPKVEGKDDVTGEELTTRKDDQEETVRKRLVEYHQMTAPLIG
      YYSKEAEAGNTKYAKVDGTKPVAEVRADLEKILG
+ attr: atom, xyz, seqres, helix, sheet,
        calpha, remark, call
m <- nma(adk)
 Building Hessian...
                            Done in 0.023 seconds.
 Diagonalizing Hessian...
                            Done in 0.467 seconds.
```







Write out a trajectory of the predicted molecular motion:

Load this file into Mol*