class10-structural-bioinformatics

Aaron (PID A17544470)

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The PDB database

The main repository of biomolecular structure is called the PDB found at :http://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

The commas in the numbers is causing data to be converted to character data instead of numeric.

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

This can be fixed 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.

```
##install.packages("readr")
library("readr")
pdbstats <- read_csv("Data Export Summary.csv")</pre>
```

```
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.

pdbstats

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

Total number of X-ray structures

sum(df\$x_ray)

[1] 191374

Total number of structures

sum(df\$total)

[1] 231029

Percentage of X-ray structures

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

[1] 82.83549

Percentage of EM structures

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

[1] 10.75017

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

Percentage of protein structures

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

[1] 86.23852

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?

2. Using Mol*

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

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

We see just one atom per water molecule in this structure because both hydrogens are involved in hydrogen bonding and show as sticks in a ball and stick representation.

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, this water molecule is HOH308



Figure 1: Molecular view of 1HSG $\,$

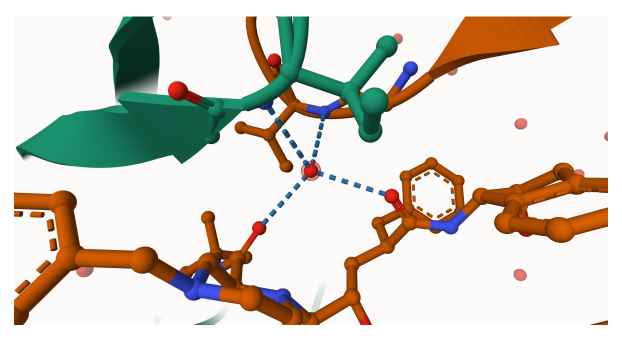
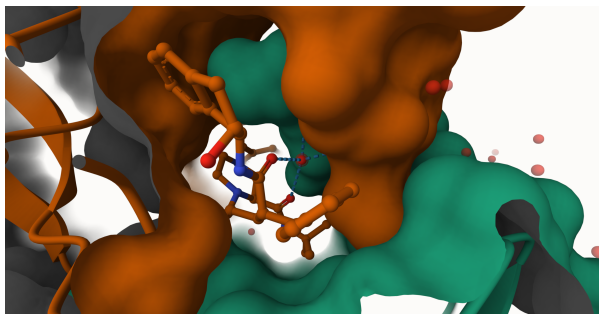


Figure 2: Molecular view of an important water



> 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.

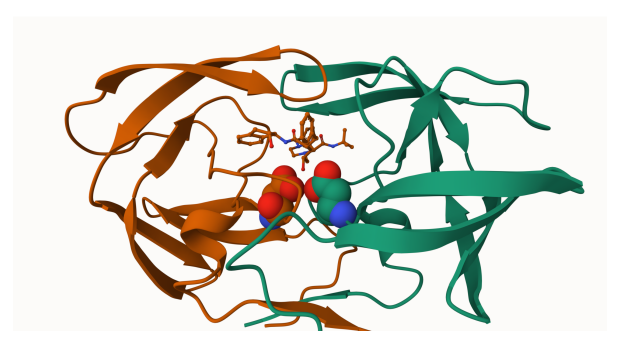


Figure 3: Molecular view of aspartic acid

3. Introduction to Bio3D in R

We can use the **biod3d** package for structural bioinformatics to read PDB 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 object in more detail
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
                          PRO
                                            <NA> 29.361 39.686 5.862 1 38.10
           1
                 N < NA >
                                  Α
                                        1
2 ATOM
           2
                CA <NA>
                          PRO
                                        1
                                            <NA> 30.307 38.663 5.319 1 40.62
                                  Α
3 ATOM
           3
                C <NA>
                          PRO
                                        1
                                            <NA> 29.760 38.071 4.022 1 42.64
                                  Α
                                        1 <NA> 28.600 38.302 3.676 1 43.40
4 ATOM
           4
                 O <NA>
                          PRO
                                  Α
5 ATOM
           5
                          PRO
                                        1
                                            <NA> 30.508 37.541 6.342 1 37.87
                CB <NA>
                                  Α
6 ATOM
           6
                CG <NA>
                          PRO
                                        1
                                            <NA> 29.296 37.591 7.162 1 38.40
  segid elesy charge
  <NA>
                <NA>
           N
1
2
  <NA>
           C
                <NA>
3 <NA>
           C
                <NA>
4 <NA>
               <NA>
           0
           С
5 <NA>
                <NA>
  <NA>
           C
                <NA>
```

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

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

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

adk</pre>
```

```
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:

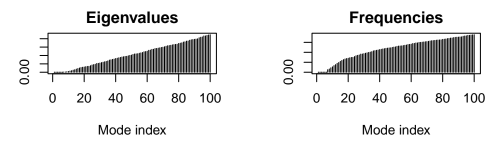
MRIILLGAPGAGKGTQAQFIMEKYGIPQISTGDMLRAAVKSGSELGKQAKDIMDAGKLVT DELVIALVKERIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFDVPDELIVDKI VGRRVHAPSGRVYHVKFNPPKVEGKDDVTGEELTTRKDDQEETVRKRLVEYHQMTAPLIG YYSKEAEAGNTKYAKVDGTKPVAEVRADLEKILG

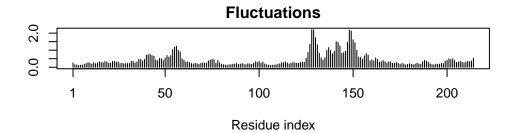
+ attr: atom, xyz, seqres, helix, sheet, calpha, remark, call

m <- nma(adk)

Building Hessian... Done in 0.012 seconds. Diagonalizing Hessian... Done in 0.262 seconds.

plot(m)





Write out a trajectory of the predicted molecular motion:

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