# Class 11

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### A quick look at the PDB

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

```
##
                              X.ray
                                      NMR
                                             EM Multiple.methods Neutron Other
                                                                                  Total
## Protein (only)
                             142303 11804 5999
                                                                        70
                                                                              32 160385
                                                              177
## Protein/Oligosaccharide
                               8414
                                       31 979
                                                                        0
                                                                                   9429
                                                                5
                                      274 1986
                                                                                   9754
## Protein/NA
                               7491
                                                                3
                                                                        0
                                                                               0
                                                                         2
## Nucleic acid (only)
                               2368
                                     1372
                                                                8
                                                                                   3811
                                             60
                                                                               1
## Other
                                149
                                       31
                                              3
                                                                                    183
                                                                0
                                                                         0
                                                                               0
## Oligosaccharide (only)
                                 11
                                        6
                                              0
                                                                1
                                                                         0
                                                                               4
                                                                                      22
```

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

```
totalxray <- sum(db$X.ray)
totalem <- sum(db$EM)
total.sums <- sum(db$Total)

perc <- ( (totalxray + totalem)/total.sums ) * 100
perc</pre>
```

## [1] 92.47157

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

```
propProtein <- (160385 / total.sums)
propProtein</pre>
```

## [1] 0.8736328

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?

There are 591 HIV-protease structures. It is a very important structure.

## Visualizing HIV-1 protease structure using VMD

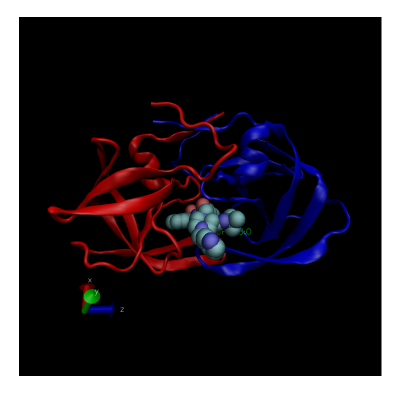
Q4: Water molecules normally have 3 atoms. Why do we see just one atom per water molecule in this structure?

We only see the oxygen atoms only because the hydrogen atoms are too small to see.

Q5: There is a conserved water molecule in the binding site. Can you identify this water molecule? What residue number does this water molecule have (see note below)?

HOH308:0

## VMD Structure visualization image



### Introduction to Bio3D in R

Load bio3d package

library(bio3d)

Read the pdb file

pdb <- read.pdb("1hsg") # accessing the online 1hsg file</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
```

To extract the sequence

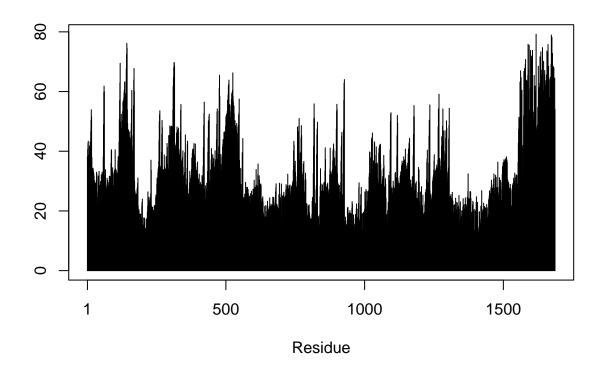
#### aa123(pdbseq(pdb))

```
[1] "PRO" "GLN" "ILE" "THR" "LEU" "TRP" "GLN" "ARG" "PRO" "LEU" "VAL" "THR"
##
    [13] "ILE" "LYS" "ILE" "GLY" "GLY" "GLN" "LEU" "LYS" "GLU" "ALA" "LEU" "LEU"
    [25] "ASP" "THR" "GLY" "ALA" "ASP" "ASP" "THR" "VAL" "LEU" "GLU" "GLU" "MET"
    [37] "SER" "LEU" "PRO" "GLY" "ARG" "TRP" "LYS" "PRO" "LYS" "MET" "ILE" "GLY"
    [49] "GLY" "ILE" "GLY" "GLY" "PHE" "ILE" "LYS" "VAL" "ARG" "GLN" "TYR" "ASP"
##
    [61] "GLN" "ILE" "LEU" "ILE" "GLU" "ILE" "CYS" "GLY" "HIS" "LYS" "ALA" "ILE"
##
    [73] "GLY" "THR" "VAL" "LEU" "VAL" "GLY" "PRO" "THR" "PRO" "VAL" "ASN" "ILE"
##
   [85] "TLE" "GLY" "ARG" "ASN" "LEU" "LEU" "THR" "GLN" "TLE" "GLY" "CYS" "THR"
##
   [97] "LEU" "ASN" "PHE" "PRO" "GLN" "ILE" "THR" "LEU" "TRP" "GLN" "ARG" "PRO"
  [109] "LEU" "VAL" "THR" "ILE" "LYS" "ILE" "GLY" "GLY" "GLN" "LEU" "LYS" "GLU"
  [121] "ALA" "LEU" "LEU" "ASP" "THR" "GLY" "ALA" "ASP" "ASP" "THR" "VAL" "LEU"
  [133] "GLU" "GLU" "MET" "SER" "LEU" "PRO" "GLY" "ARG" "TRP" "LYS" "PRO" "LYS"
  [145] "MET" "ILE" "GLY" "GLY" "ILE" "GLY" "GLY" "PHE" "ILE" "LYS" "VAL" "ARG"
## [157] "GLN" "TYR" "ASP" "GLN" "ILE" "LEU" "ILE" "GLU" "ILE" "CYS" "GLY" "HIS"
## [169] "LYS" "ALA" "ILE" "GLY" "THR" "VAL" "LEU" "VAL" "GLY" "PRO" "THR" "PRO"
## [181] "VAL" "ASN" "ILE" "ILE" "GLY" "ARG" "ASN" "LEU" "LEU" "THR" "GLN" "ILE"
## [193] "GLY" "CYS" "THR" "LEU" "ASN" "PHE"
```

Plot of B-factor

```
plot.bio3d(pdb$atom$b, sse=pdb)
```

```
## Warning in plotb3(...): Length of input 'sse' does not equal the length of input
## 'x'; Ignoring 'sse'
```



Q7: How many amino acid residues are there in this pdb object?

There are 198 amino acid residues.

Q8: Name one of the two non-protein residues?

A non-protein residue is HOH, or water.

Q9: How many protein chains are in this structure?

There are 2 protein chains in this structure.

Find the attributes of the object

#### attributes(pdb)

```
## $names
## [1] "atom" "xyz" "seqres" "helix" "sheet" "calpha" "remark" "call"
##
## $class
## [1] "pdb" "sse"
```

Look at ATOM part of the pdb file

#### head(pdb\$atom)

```
type eleno elety alt resid chain resno insert
                                                          Х
                                                                        z o
                                                                                b
                                                                  У
## 1 ATOM
                    N <NA>
                             PRO
                                      Α
                                            1
                                                <NA> 29.361 39.686 5.862 1 38.10
              1
## 2 ATOM
              2
                             PRO
                                      Α
                                                <NA> 30.307 38.663 5.319 1 40.62
                   CA <NA>
                                            1
## 3 ATOM
                    C <NA>
                             PRO
                                                <NA> 29.760 38.071 4.022 1 42.64
              3
                                      Α
                                            1
## 4 ATOM
              4
                    O <NA>
                             PRO
                                      Α
                                            1
                                                <NA> 28.600 38.302 3.676 1 43.40
                                              <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
## 1 <NA>
               N
                   <NA>
## 2
      <NA>
               С
                   <NA>
## 3
     <NA>
               С
                   <NA>
## 4
      <NA>
               0
                   <NA>
## 5
      <NA>
               С
                   <NA>
## 6 <NA>
               C
                   <NA>
```

The atom data in R

#### head(pdb\$atom)

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