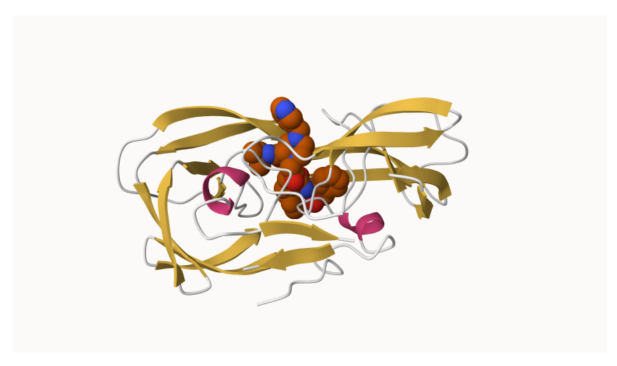
Lab 10 (Structural Bioinformatics)

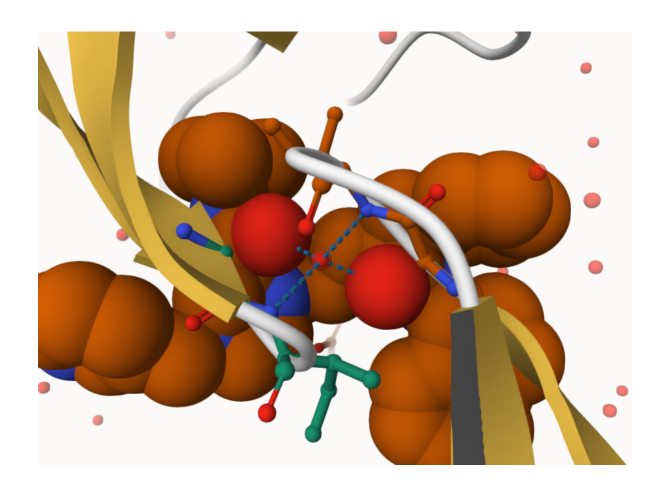
Jazz Zhang (A16149005)

```
PDB_summary <- read.csv("Data Export Summary.csv", row.names = 1)</pre>
  rm.comma <- function(x){</pre>
    as.numeric(gsub(",", "", x))
  PDB_stats <- apply(PDB_summary, 2, rm.comma)</pre>
  rownames(PDB_stats) <- rownames(PDB_summary)</pre>
  sum(PDB_stats[,"X.ray"])/sum(PDB_stats[,"Total"])
[1] 0.8483231
  sum(PDB_stats[,"EM"])/sum(PDB_stats[,"Total"])
[1] 0.08327301
Q1. 84.83% by X ray; 8.33% by EM
  PDB_stats["Protein (only)","Total"]/sum(PDB_stats[,"Total"])
[1] 0.8667026
Q2. 86.67\%
Q3. 211,377
```



 $\mathbf{Q4}.$ Hydrogen's not displayed because the atom size is smaller than $\mathbf{2A}$ (resolution of the structure)

Q5. 308





library(bio3d)

```
Warning: package 'bio3d' was built under R version 4.3.1
```

```
pdb <- read.pdb("1hsg")

Note: Accessing on-line PDB file

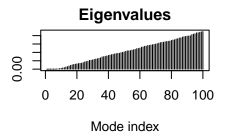
pdb

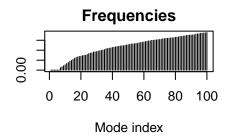
Call: read.pdb(file = "1hsg")

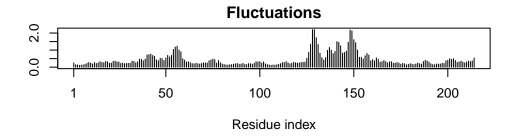
Total Models#: 1</pre>
```

```
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. 198
Q8. HOH; MK1
Q9. 2
  attributes(pdb)
$names
[1] "atom"
                     "segres" "helix" "sheet" "calpha" "remark" "call"
            "xyz"
$class
[1] "pdb" "sse"
  head(pdb$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
                                       1 <NA> 28.600 38.302 3.676 1 43.40
4 ATOM
                         PRO
          4
               O <NA>
                                 Α
5 ATOM
               CB <NA>
                         PRO
                                 Α
                                      1 <NA> 30.508 37.541 6.342 1 37.87
6 ATOM
               CG <NA>
                         PRO
                                 Α
                                      1 <NA> 29.296 37.591 7.162 1 38.40
 segid elesy charge
```

```
1 <NA>
         N <NA>
2 <NA>
           C <NA>
           C <NA>
3 <NA>
4 <NA>
           O <NA>
           C <NA>
5 <NA>
6 <NA>
           C <NA>
  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
  m <- nma(adk)
Building Hessian...
                           Done in 0.01 seconds.
Diagonalizing Hessian... Done in 0.25 seconds.
```







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