# Class 11

Benjie Miao (PID: A69026849)

# Lab 11: Protein Structure Prediction with AlphaFold

First, we used Colab to predict the structure of the given HIV sequence. We store it into the HIVPrDimer\_23119 folder.

```
# Change this for YOUR results dir name
  results_dir <- "HIVPrDimer_23119/"</pre>
  # File names for all PDB models
  pdb_files <- list.files(path=results_dir,</pre>
                           pattern="*.pdb",
                           full.names = TRUE)
  # Print our PDB file names
  basename(pdb files)
[1] "HIVPrDimer_23119_unrelaxed_rank_001_alphafold2_multimer_v3_model_1_seed_000.pdb"
[2] "HIVPrDimer_23119_unrelaxed_rank_002_alphafold2_multimer_v3_model_5_seed_000.pdb"
[3] "HIVPrDimer_23119_unrelaxed_rank_003_alphafold2_multimer_v3_model_4_seed_000.pdb"
[4] "HIVPrDimer_23119_unrelaxed_rank_004_alphafold2_multimer_v3_model_2_seed_000.pdb"
[5] "HIVPrDimer_23119_unrelaxed_rank_005_alphafold2_multimer_v3_model_3_seed_000.pdb"
  library(bio3d)
Warning: package 'bio3d' was built under R version 4.3.2
  # Read all data from Models
  # and superpose/fit coords
  pdbs <- pdbaln(pdb_files, fit=TRUE, exefile="msa")</pre>
```

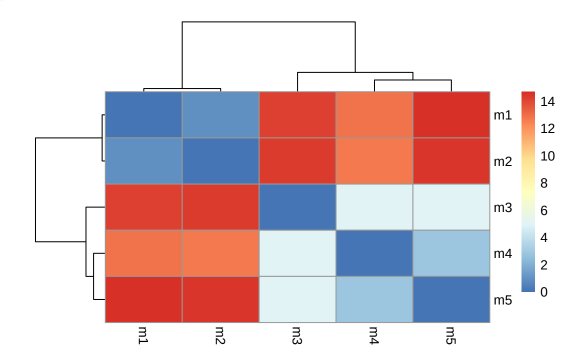
### Reading PDB files: HIVPrDimer\_23119/HIVPrDimer\_23119\_unrelaxed\_rank\_001\_alphafold2\_multimer\_v3\_model\_1\_seed\_000 HIVPrDimer\_23119/HIVPrDimer\_23119\_unrelaxed\_rank\_002\_alphafold2\_multimer\_v3\_model\_5\_seed\_000 HIVPrDimer\_23119/HIVPrDimer\_23119\_unrelaxed\_rank\_003\_alphafold2\_multimer\_v3\_model\_4\_seed\_000 HIVPrDimer\_23119/HIVPrDimer\_23119\_unrelaxed\_rank\_004\_alphafold2\_multimer\_v3\_model\_2\_seed\_000 HIVPrDimer\_23119/HIVPrDimer\_23119\_unrelaxed\_rank\_005\_alphafold2\_multimer\_v3\_model\_3\_seed\_000 Extracting sequences name: HIVPrDimer 23119/HIVPrDimer 23119 unrelaxed rank 001 alphafold2 multimer pdb/seq: 1 pdb/seq: 2 name: HIVPrDimer\_23119/HIVPrDimer\_23119\_unrelaxed\_rank\_002\_alphafold2\_multimer\_ pdb/seq: 3 name: HIVPrDimer\_23119/HIVPrDimer\_23119\_unrelaxed\_rank\_003\_alphafold2\_multimer\_ pdb/seq: 4 name: HIVPrDimer\_23119/HIVPrDimer\_23119\_unrelaxed\_rank\_004\_alphafold2\_multimer\_ pdb/seq: 5 name: HIVPrDimer\_23119/HIVPrDimer\_23119\_unrelaxed\_rank\_005\_alphafold2\_multimer\_ pdbs 50 PQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGI [Truncated\_Name:1]HIVPrDimer [Truncated\_Name:2]HIVPrDimer PQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGI [Truncated\_Name:3]HIVPrDimer PQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGI [Truncated\_Name:4]HIVPrDimer PQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGI [Truncated\_Name:5]HIVPrDimer PQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGI \*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\* 1 50 100 [Truncated\_Name:1]HIVPrDimer GGFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNFP [Truncated\_Name:2]HIVPrDimer GGFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNFP [Truncated\_Name:3]HIVPrDimer GGFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNFP [Truncated\_Name:4]HIVPrDimer GGFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNFP [Truncated\_Name:5]HIVPrDimer GGFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNFP \*\*\*\*\*\*\*\*\*\*\*\*\*\* 51 100

[Truncated\_Name:1]HIVPrDimer [Truncated\_Name:2]HIVPrDimer [Truncated\_Name:3]HIVPrDimer [Truncated\_Name:4]HIVPrDimer QITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGIG QITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGIG QITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGIG QITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGIG

```
[Truncated_Name:5]HIVPrDimer
                             QITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGIG
                             **************
                           101
                                                                            150
                           151
                                                                          198
[Truncated_Name:1]HIVPrDimer
                             GFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNF
[Truncated Name:2]HIVPrDimer
                             GFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNF
[Truncated_Name:3]HIVPrDimer
                             GFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNF
[Truncated_Name:4]HIVPrDimer
                             GFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNF
[Truncated_Name:5]HIVPrDimer
                             GFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNF
                             *************
                           151
                                                                          198
Call:
 pdbaln(files = pdb_files, fit = TRUE, exefile = "msa")
Class:
 pdbs, fasta
Alignment dimensions:
 5 sequence rows; 198 position columns (198 non-gap, 0 gap)
+ attr: xyz, resno, b, chain, id, ali, resid, sse, call
  rd <- rmsd(pdbs, fit=T)</pre>
Warning in rmsd(pdbs, fit = T): No indices provided, using the 198 non NA positions
  range(rd)
[1] 0.000 14.689
  library(pheatmap)
```

Warning: package 'pheatmap' was built under R version 4.3.2

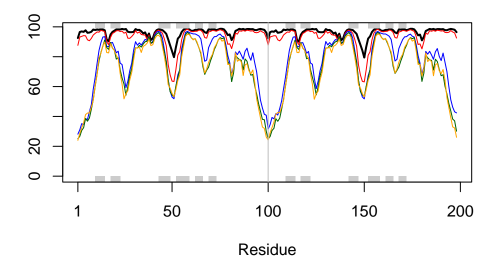
```
colnames(rd) <- paste0("m",1:5)
rownames(rd) <- paste0("m",1:5)
pheatmap(rd)</pre>
```



```
# Read a reference PDB structure
pdb <- read.pdb("1hsg")</pre>
```

Note: Accessing on-line PDB file

```
plotb3(pdbs$b[1,], typ="1", lwd=2, sse=pdb)
points(pdbs$b[2,], typ="1", col="red")
points(pdbs$b[3,], typ="1", col="blue")
points(pdbs$b[4,], typ="1", col="darkgreen")
points(pdbs$b[5,], typ="1", col="orange")
abline(v=100, col="gray")
```



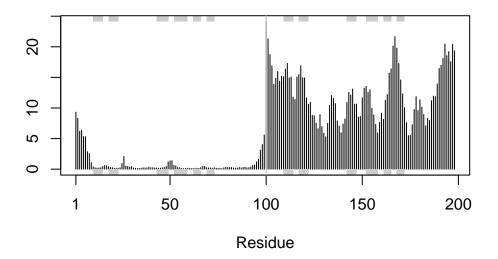
### core <- core.find(pdbs)</pre>

```
core size 197 of 198
                      vol = 6154.839
core size 196 of 198
                      vol = 5399.676
                      vol = 5074.795
core size 195 of 198
core size 194 of 198
                      vol = 4802.518
core size 193 of 198
                      vol = 4520.256
core size 192 of 198
                      vol = 4305.362
core size 191 of 198
                      vol = 4089.792
core size 190 of 198
                      vol = 3886.145
core size 189 of 198
                      vol = 3758.321
core size 188 of 198
                      vol = 3620.18
                      vol = 3496.698
core size 187 of 198
core size 186 of 198
                      vol = 3389.985
core size 185 of 198
                      vol = 3320.114
                      vol = 3258.683
core size 184 of 198
core size 183 of 198
                      vol = 3208.591
core size 182 of 198
                      vol = 3156.736
core size 181 of 198
                      vol = 3141.668
core size 180 of 198
                      vol = 3136.574
core size 179 of 198
                      vol = 3155.52
core size 178 of 198
                     vol = 3185.362
```

```
core size 177 of 198 vol = 3204.487
core size 176 of 198
                      vol = 3211.978
core size 175 of 198
                      vol = 3234.993
core size 174 of 198
                      vol = 3244.062
core size 173 of 198
                      vol = 3237.845
core size 172 of 198
                      vol = 3218.77
core size 171 of 198
                      vol = 3180.743
core size 170 of 198
                      vol = 3130.369
core size 169 of 198
                      vol = 3067.881
core size 168 of 198
                      vol = 2989.546
core size 167 of 198
                      vol = 2928.272
core size 166 of 198
                      vol = 2851.193
core size 165 of 198
                      vol = 2780.877
core size 164 of 198
                      vol = 2708.433
core size 163 of 198
                      vol = 2636.516
core size 162 of 198
                      vol = 2563.25
core size 161 of 198
                      vol = 2478.024
                      vol = 2404.793
core size 160 of 198
core size 159 of 198
                      vol = 2330.997
core size 158 of 198
                      vol = 2250.477
core size 157 of 198
                      vol = 2159.432
core size 156 of 198
                      vol = 2070.759
core size 155 of 198
                      vol = 1983.579
core size 154 of 198
                      vol = 1917.913
core size 153 of 198
                      vol = 1842.556
core size 152 of 198
                      vol = 1775.398
core size 151 of 198
                      vol = 1695.133
core size 150 of 198
                      vol = 1632.173
core size 149 of 198
                      vol = 1570.391
core size 148 of 198
                      vol = 1497.238
core size 147 of 198
                      vol = 1434.802
core size 146 of 198
                      vol = 1367.706
core size 145 of 198
                      vol = 1302.596
core size 144 of 198
                      vol = 1251.985
core size 143 of 198
                      vol = 1207.976
core size 142 of 198
                      vol = 1167.112
core size 141 of 198
                      vol = 1118.27
core size 140 of 198
                      vol = 1081.664
core size 139 of 198
                      vol = 1029.75
core size 138 of 198
                      vol = 981.766
core size 137 of 198
                      vol = 944.446
core size 136 of 198
                      vol = 899.224
core size 135 of 198 vol = 859.402
```

```
core size 134 of 198
                     vol = 814.694
core size 133 of 198
                      vol = 771.862
core size 132 of 198
                      vol = 733.807
core size 131 of 198
                      vol = 702.053
core size 130 of 198
                      vol = 658.757
core size 129 of 198
                      vol = 622.574
core size 128 of 198
                      vol = 578.29
core size 127 of 198
                      vol = 543.07
core size 126 of 198
                      vol = 510.934
core size 125 of 198
                      vol = 481.595
core size 124 of 198
                      vol = 464.672
core size 123 of 198
                      vol = 451.721
core size 122 of 198
                      vol = 430.417
core size 121 of 198
                      vol = 409.141
core size 120 of 198
                      vol = 378.942
core size 119 of 198
                      vol = 348.325
core size 118 of 198
                      vol = 324.738
core size 117 of 198
                      vol = 312.394
core size 116 of 198
                      vol = 300.89
core size 115 of 198
                      vol = 279.976
core size 114 of 198
                      vol = 263.434
core size 113 of 198
                      vol = 250.263
core size 112 of 198
                      vol = 229.592
core size 111 of 198
                      vol = 209.929
core size 110 of 198
                      vol = 196.379
core size 109 of 198
                      vol = 180.628
core size 108 of 198
                      vol = 167.088
core size 107 of 198
                      vol = 155.875
core size 106 of 198
                      vol = 142.595
core size 105 of 198
                      vol = 128.924
core size 104 of 198
                      vol = 114.054
core size 103 of 198
                      vol = 100.936
core size 102 of 198
                      vol = 90.431
core size 101 of 198
                      vol = 81.972
core size 100 of 198
                      vol = 74.017
core size 99 of 198
                     vol = 66.855
core size 98 of 198
                     vol = 59.525
core size 97 of 198
                     vol = 52.263
core size 96 of 198
                     vol = 43.699
core size 95 of 198
                     vol = 35.813
core size 94 of 198
                     vol = 28.888
core size 93 of 198
                     vol = 20.692
core size 92 of 198 vol = 14.975
```

```
core size 91 \text{ of } 198 \text{ vol} = 9.146
 core size 90 of 198 vol = 5.232
 core size 89 of 198 vol = 3.53
 core size 88 of 198 vol = 2.657
core size 87 of 198 vol = 1.998
core size 86 \text{ of } 198 \text{ vol} = 1.333
core size 85 of 198 vol = 1.141
core size 84 of 198 vol = 1.012
core size 83 of 198 vol = 0.891
core size 82 of 198 vol = 0.749
core size 81 of 198 vol = 0.618
core size 80 of 198 vol = 0.538
 core size 79 of 198 vol = 0.479
FINISHED: Min vol (0.5) reached
  core.inds <- print(core, vol=0.5)</pre>
# 80 positions (cumulative volume <= 0.5 Angstrom^3)
  start end length
         25
1
     10
                 16
2
     27
         48
                 22
3
     53 94
                 42
  xyz <- pdbfit(pdbs, core.inds, outpath="corefit_structures")</pre>
  rf <- rmsf(xyz)</pre>
  plotb3(rf, sse=pdb)
  abline(v=100, col="gray", ylab="RMSF")
```



## Predict alignment error

```
library(jsonlite)
  # Listing of all PAE JSON files
  pae_files <- list.files(path=results_dir,</pre>
                            pattern=".*model.*\\.json",
                            full.names = TRUE)
  pae1 <- read_json(pae_files[1],simplifyVector = TRUE)</pre>
  pae5 <- read_json(pae_files[5],simplifyVector = TRUE)</pre>
  attributes(pae1)
$names
[1] "plddt"
              "max_pae" "pae"
                                    "ptm"
                                               "iptm"
  # Per-residue pLDDT scores
  # same as B-factor of PDB..
  head(pae1$plddt)
```

```
[1] 92.50 96.56 96.94 96.62 97.69 96.00
```

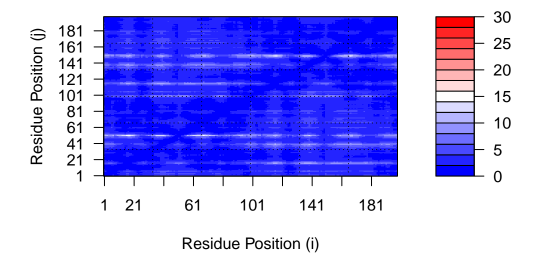
```
pae1$max_pae
```

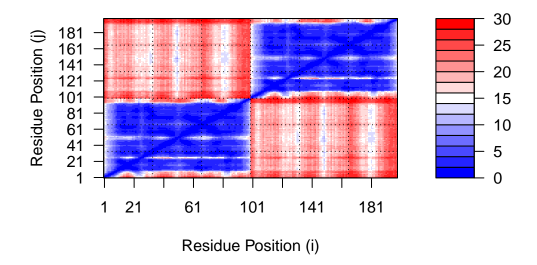
### [1] 15.54688

```
pae5$max_pae
```

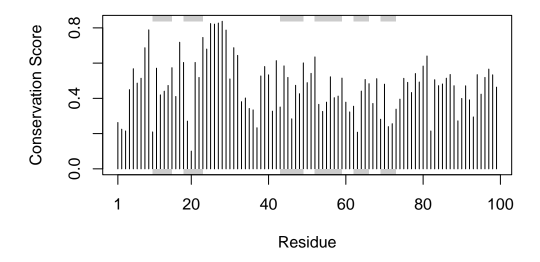
### [1] 29.29688

The first entry is a better alignment.





### Residue conservation from alignment file



```
con <- consensus(aln, cutoff = 0.9)
con$seq</pre>
```

```
m1.pdb <- read.pdb(pdb_files[1])
occ <- vec2resno(c(sim[1:99], sim[1:99]), m1.pdb$atom$resno)
write.pdb(m1.pdb, o=occ, file="m1_conserv.pdb")</pre>
```

Then we can use  ${\tt Mol*}$  to visualize the pdb file.