Class 11: Structural Bioinformatics (pt2)

Lilia Jimenez (PID:A16262599)

Alphafold has changed the game for protein structure prediction and allows anyone with sufficient bioninfromatics skills to predict the structure if virtually any protein.

we ran alphafold via googlecolab at: https://github.com/sokrypton/ColabFold

in particular we used their alphafold2_mmseqs2 version that uses mmseqs2 rather than hmmmer for sequence search

The main output include a set of PDB structure files along with matching JSON format files that tell us how good the resulting models might be lets start by loading these structures up in Mol*

- [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)

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

```
pdb/seq: 1 name: hivprdimer_23119/hivprdimer_23119_unrelaxed_rank_001_alphafold2_multimer_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_
```

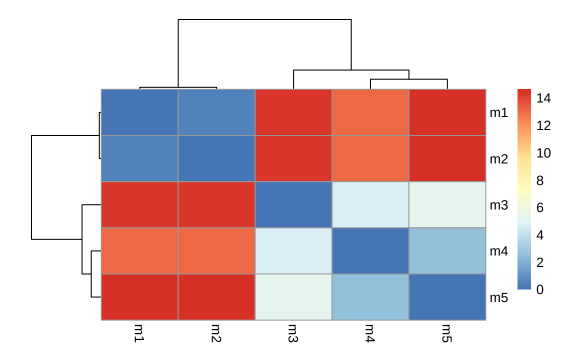
core<-core.find(pdbs)</pre>

```
core size 197 of 198 vol = 4578.346
core size 196 of 198 vol = 3931.108
core size 195 of 198 vol = 3709.733
core size 194 of 198 vol = 3496.019
core size 193 of 198 vol = 3302.432
core size 192 of 198 vol = 3146.474
core size 191 of 198 vol = 3048.964
core size 190 of 198 vol = 2970.354
core size 189 of 198 vol = 2893.012
core size 188 of 198 vol = 2831.825
core size 187 \text{ of } 198 \text{ vol} = 2774.506
core size 186 \text{ of } 198 \text{ vol} = 2728.043
core size 185 \text{ of } 198 \text{ vol} = 2704.946
core size 184 of 198 vol = 2701.981
core size 183 \text{ of } 198 \text{ vol} = 2715.909
core size 182 of 198 vol = 2809.853
core size 181 of 198 vol = 2888.95
core size 180 of 198 vol = 2967.282
core size 179 of 198 vol = 3036.256
core size 178 of 198
                      vol = 3066.287
core size 177 of 198
                      vol = 3096.833
core size 176 of 198
                      vol = 3056.414
core size 175 of 198 vol = 3014.768
```

```
core size 174 of 198 vol = 2975.013
core size 173 of 198
                      vol = 2898.051
core size 172 of 198
                      vol = 2810.173
core size 171 of 198
                      vol = 2747.532
core size 170 of 198
                      vol = 2684.434
core size 169 of 198
                      vol = 2620.353
core size 168 of 198
                      vol = 2550.877
core size 167 of 198
                      vol = 2492.582
core size 166 of 198
                      vol = 2422.978
                      vol = 2358.916
core size 165 of 198
                      vol = 2298.292
core size 164 of 198
core size 163 of 198
                      vol = 2235.918
core size 162 of 198
                      vol = 2171.02
core size 161 of 198
                      vol = 2093.559
core size 160 of 198
                      vol = 2029.144
core size 159 of 198
                      vol = 1950.957
core size 158 of 198
                      vol = 1881.015
                      vol = 1801.506
core size 157 of 198
core size 156 of 198
                      vol = 1728.892
core size 155 of 198
                      vol = 1660.037
core size 154 of 198
                      vol = 1586.149
core size 153 of 198
                      vol = 1532.718
core size 152 of 198
                      vol = 1460.186
core size 151 of 198
                      vol = 1399.251
core size 150 of 198
                      vol = 1333.908
core size 149 of 198
                      vol = 1271.747
                      vol = 1219.496
core size 148 of 198
core size 147 of 198
                      vol = 1176.003
core size 146 of 198
                      vol = 1138.478
core size 145 of 198
                      vol = 1102.124
core size 144 of 198
                      vol = 1049.642
core size 143 of 198
                      vol = 1014.063
core size 142 of 198
                      vol = 970.575
core size 141 of 198
                      vol = 929.178
core size 140 of 198
                      vol = 889.104
core size 139 of 198
                      vol = 846.668
core size 138 of 198
                      vol = 805.8
core size 137 of 198
                      vol = 775.034
                      vol = 743.09
core size 136 of 198
core size 135 of 198
                      vol = 715.695
core size 134 of 198
                      vol = 689.788
core size 133 of 198
                      vol = 660.329
core size 132 of 198 vol = 630.966
```

```
core size 131 of 198 vol = 597.207
core size 130 of 198
                      vol = 566.989
core size 129 of 198
                      vol = 532.89
core size 128 of 198
                      vol = 496.208
core size 127 of 198
                      vol = 463.183
core size 126 of 198
                      vol = 431.893
core size 125 of 198
                      vol = 408.864
core size 124 of 198
                      vol = 376.61
core size 123 of 198
                      vol = 362.377
                      vol = 353.633
core size 122 of 198
core size 121 of 198
                      vol = 331.501
core size 120 of 198
                      vol = 312.518
core size 119 of 198
                      vol = 286.715
core size 118 of 198
                      vol = 262.336
core size 117 of 198
                      vol = 245.109
core size 116 of 198
                      vol = 228.342
core size 115 of 198
                      vol = 210.366
                      vol = 197.519
core size 114 of 198
core size 113 of 198
                      vol = 179.392
core size 112 of 198
                      vol = 161.891
core size 111 of 198
                      vol = 148.359
core size 110 of 198
                      vol = 134.477
core size 109 of 198
                      vol = 121.261
core size 108 of 198
                      vol = 109.516
core size 107 of 198
                      vol = 103.031
core size 106 of 198
                      vol = 96.443
core size 105 of 198
                      vol = 88.455
core size 104 of 198
                      vol = 81.816
core size 103 of 198
                      vol = 74.88
core size 102 of 198
                      vol = 68.386
core size 101 of 198
                      vol = 65.937
core size 100 of 198
                      vol = 62.345
core size 99 of 198 vol = 58.836
                     vol = 52.868
core size 98 of 198
core size 97 of 198
                     vol = 47.796
core size 96 of 198
                     vol = 41.292
core size 95 of 198
                     vol = 33.831
core size 94 of 198
                     vol = 24.912
core size 93 of 198
                     vol = 18.912
core size 92 of 198
                     vol = 12.7
core size 91 of 198
                     vol = 7.35
core size 90 of 198
                     vol = 4.922
core size 89 of 198 vol = 3.421
```

```
core size 88 of 198 vol = 2.553
 core size 87 of 198 vol = 1.917
 core size 86 of 198 vol = 1.513
 core size 85 of 198 vol = 1.201
 core size 84 of 198 vol = 1.046
 core size 83 of 198 vol = 0.922
 core size 82 of 198 vol = 0.755
 core size 81 of 198 vol = 0.668
 core size 80 of 198 vol = 0.596
 core size 79 of 198 vol = 0.549
 core size 78 of 198 vol = 0.493
 FINISHED: Min vol (0.5) reached
  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 14.631
  library(pheatmap)
  colnames(rd) <- paste0("m",1:5)</pre>
  rownames(rd) <- paste0("m",1:5)</pre>
  pheatmap(rd)
```



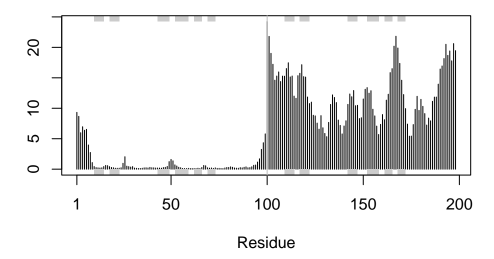
read a reference pdb structure fule

```
pdb <- read.pdb("1hsg")</pre>
```

Note: Accessing on-line PDB file

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

```
core.inds <- print(core, vol=0.5)</pre>
# 79 positions (cumulative volume <= 0.5 Angstrom^3)</pre>
  start end length
1
     10
          25
                  16
2
                  21
     28
          48
     53
                  42
         94
  xyz <- pdbfit(pdbs, core.inds, outpath="corefit_structures")</pre>
  rf <- rmsf(xyz)</pre>
  plotb3(rf, sse=pdb)
  abline(v=100, col="gray", ylab="RMSF")
```



If the predicted model has more than one domain, each domain may have high confidence, yet the relative positions of the domains may not. The estimated reliability of relative domain positions is in graphs of predicted aligned error (PAE) which are included in the downloadable zip file and analyzed in R above.

##Predicted Alignment Error for domains

```
head(pae1$plddt)
```

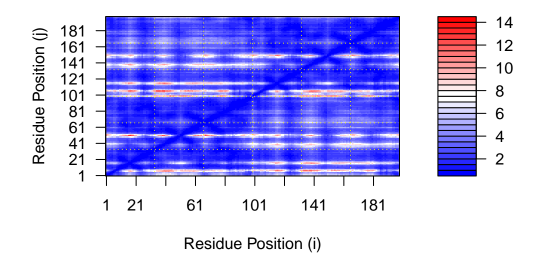
[1] 87.81 92.00 91.81 91.88 94.25 88.00

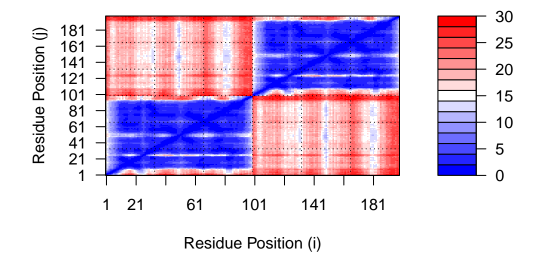
```
pae1$max_pae
```

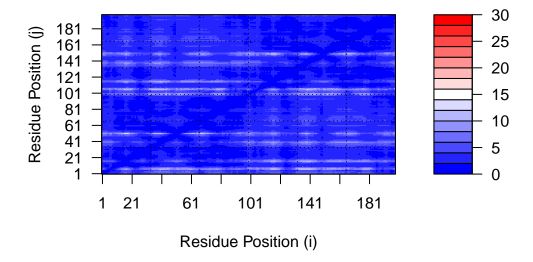
[1] 14.09375

 $pae5$max_pae$

[1] 29.29688







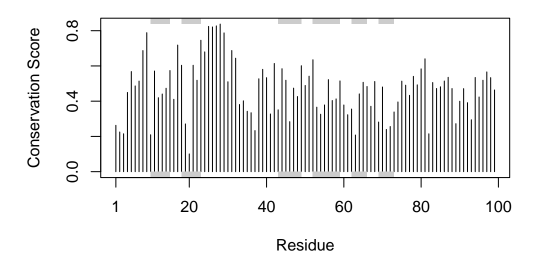
##Residue conservation from alignment file

[1] "hivprdimer_23119/hivprdimer_23119.a3m"

```
aln <- read.fasta(aln_file[1], to.upper = TRUE)
[1] " ** Duplicated sequence id's: 101 **"
[2] " ** Duplicated sequence id's: 101 **"

dim(aln$ali)</pre>
```

[1] 5378 132



```
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>
```