Alphafold Analysis

Sung Lien A16628474

Here we analyze our AlphaFold structure prediction models. The input directory/folder comes from the ColabFold folder:

```
# Change this for YOUR results dir name
results_dir <- "hivpr_monomer_94b5b/"</pre>
```

```
[1] "hivpr_monomer_94b5b_unrelaxed_rank_001_alphafold2_ptm_model_5_seed_000.pdb"
```

- [3] "hivpr_monomer_94b5b_unrelaxed_rank_003_alphafold2_ptm_model_4_seed_000.pdb"
- [4] "hivpr_monomer_94b5b_unrelaxed_rank_004_alphafold2_ptm_model_3_seed_000.pdb"
- [5] "hivpr_monomer_94b5b_unrelaxed_rank_005_alphafold2_ptm_model_2_seed_000.pdb"

I will use the Bio3D package for analysis

```
library(bio3d)
```

Align and superpose

```
pdbs <- pdbaln(pdb_files, fit=TRUE, exefile="msa")</pre>
```

^{[2] &}quot;hivpr_monomer_94b5b_unrelaxed_rank_002_alphafold2_ptm_model_1_seed_000.pdb"

```
Reading PDB files:
hivpr_monomer_94b5b//hivpr_monomer_94b5b_unrelaxed_rank_001_alphafold2_ptm_model_5_seed_000.
hivpr_monomer_94b5b//hivpr_monomer_94b5b_unrelaxed_rank_002_alphafold2_ptm_model_1_seed_000.
hivpr_monomer_94b5b//hivpr_monomer_94b5b_unrelaxed_rank_003_alphafold2_ptm_model_4_seed_000.
hivpr_monomer_94b5b//hivpr_monomer_94b5b_unrelaxed_rank_004_alphafold2_ptm_model_3_seed_000.
hivpr_monomer_94b5b//hivpr_monomer_94b5b_unrelaxed_rank_005_alphafold2_ptm_model_2_seed_000.
Extracting sequences
            name: hivpr monomer 94b5b//hivpr monomer 94b5b unrelaxed rank 001 alphafold2 pt
pdb/seq: 1
            name: hivpr_monomer_94b5b//hivpr_monomer_94b5b_unrelaxed_rank_002_alphafold2_ptm
pdb/seq: 2
pdb/seq: 3
            name: hivpr_monomer_94b5b//hivpr_monomer_94b5b_unrelaxed_rank_003_alphafold2_ptm
            name: hivpr_monomer_94b5b//hivpr_monomer_94b5b_unrelaxed_rank_004_alphafold2_ptm
pdb/seq: 4
pdb/seq: 5
            name: hivpr_monomer_94b5b//hivpr_monomer_94b5b_unrelaxed_rank_005_alphafold2_ptm
pdbs
                                                                              50
[Truncated_Name:1]hivpr_mono
                              PQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGI
[Truncated_Name:2]hivpr_mono
                              PQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGI
[Truncated_Name:3]hivpr_mono
                              PQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGI
                              PQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGI
[Truncated_Name:4]hivpr_mono
[Truncated_Name:5]hivpr_mono
                              PQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGI
                              **************
                              1
                                                                              50
                                                                             99
                             51
[Truncated_Name:1]hivpr_mono
                              GGFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNF
[Truncated_Name:2]hivpr_mono
                              GGFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNF
[Truncated_Name:3]hivpr_mono
                              GGFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNF
[Truncated_Name:4]hivpr_mono
                              GGFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNF
[Truncated_Name:5]hivpr_mono
                              GGFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNF
                              **************
                             51
                                                                             99
```

Call:

pdbaln(files = pdb_files, fit = TRUE, exefile = "msa")

Class:

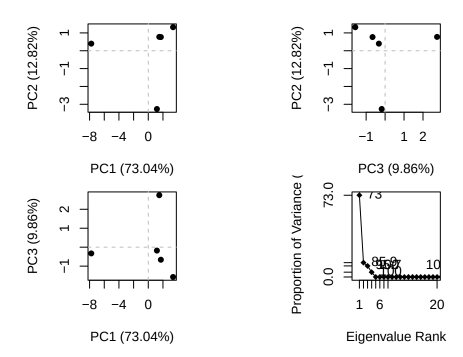
pdbs, fasta

Alignment dimensions:

5 sequence rows; 99 position columns (99 non-gap, 0 gap)

+ attr: xyz, resno, b, chain, id, ali, resid, sse, call

```
pc <- pca(pdbs)
plot(pc)</pre>
```



RMSD analysis

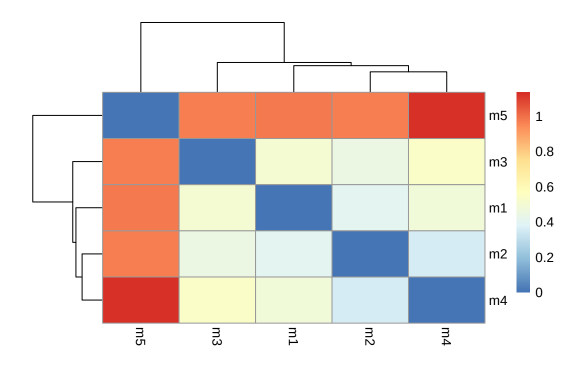
RMSD is a common measure od structural distance used in structural biology.

```
rd <- rmsd(pdbs, fit=T)
```

Warning in rmsd(pdbs, fit = T): No indices provided, using the 99 non NA positions

```
library(pheatmap)

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

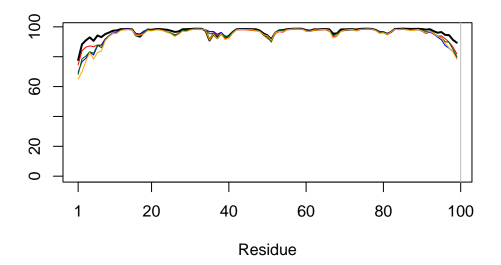


Read a reference PDB structure

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

Note: Accessing on-line PDB file

```
plotb3(pdbs$b[1,], typ="l", lwd=2, sse=pdb$sse[1:length(pdbs$b[1,])])
points(pdbs$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 <- core.find(pdbs)</pre>

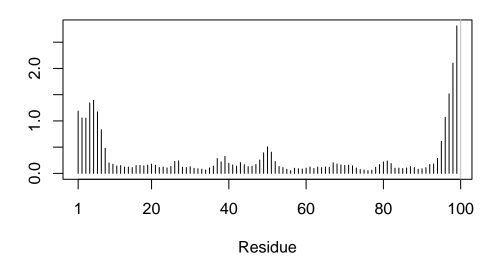
```
core size 98 of 99
                  vol = 3.178
core size 97 of 99
                   vol = 2.565
core size 96 of 99
                   vol = 2.035
                   vol = 1.636
core size 95 of 99
core size 94 of 99
                   vol = 1.281
core size 93 of 99
                   vol = 0.945
                   vol = 0.653
core size 92 of 99
core size 91 of 99 vol = 0.481
FINISHED: Min vol (0.5) reached
```

core.inds <- print(core, vol=0.5)</pre>

```
xyz <- pdbfit(pdbs, core.inds, outpath="corefit_structures")

rf <- rmsf(xyz)

plotb3(rf, sse=pdb$sse[1:length(pdbs$b[1,])])
abline(v=100, col="gray", ylab="RMSF")</pre>
```



#Predicted Alignment Error for domains

```
$names
[1] "plddt" "max_pae" "pae" "ptm"
```

attributes(pae1)

```
# Per-residue pLDDT scores
# same as B-factor of PDB..
head(pae1$plddt)
```

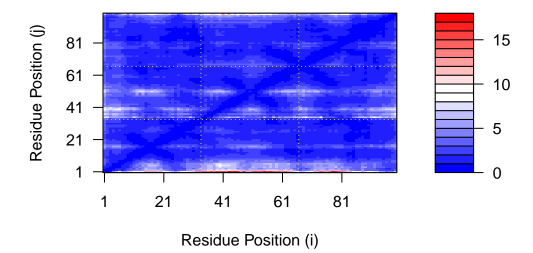
[1] 77.81 88.31 90.94 92.94 90.62 94.19

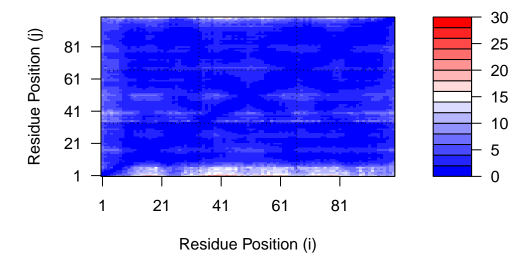
```
pae1$max_pae
```

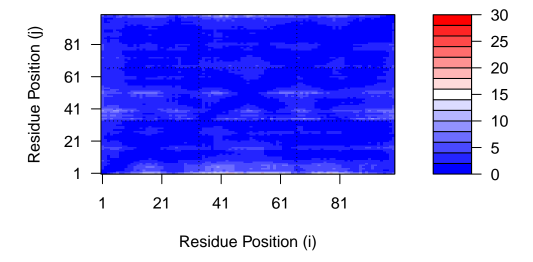
[1] 17.59375

```
pae5$max_pae
```

[1] 20.67188







#Residue conservation from alignment file

[1] "hivpr_monomer_94b5b//hivpr_monomer_94b5b.a3m"

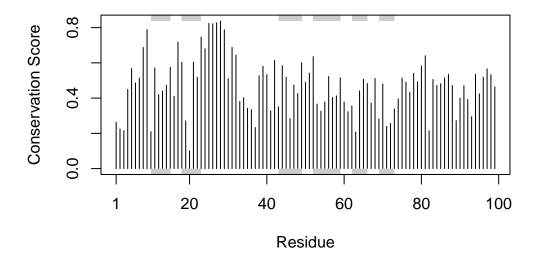
```
aln <- read.fasta(aln_file[1], to.upper = TRUE)</pre>
```

[1] " ** Duplicated sequence id's: 101 **"

```
dim(aln$ali)
```

[1] 5378 132

```
sim <- conserv(aln)</pre>
```



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