# class11:alphafold

## Tin

Here we read the results from alpha fold and try to interpret all the models and quality score metrics:

```
library(bio3d)
pth <- "hivdimer_23119/"
pdb.files <- list.files(path = pth, full.names = TRUE, pattern = ".pdb")</pre>
```

Align and supperpose all these models

```
file.exists(pdb.files)
```

[1] TRUE TRUE TRUE TRUE TRUE

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

#### Reading PDB files:

```
hivdimer_23119//hivdimer_23119_unrelaxed_rank_001_alphafold2_multimer_v3_model_2_seed_000.pd hivdimer_23119//hivdimer_23119_unrelaxed_rank_002_alphafold2_multimer_v3_model_5_seed_000.pd hivdimer_23119//hivdimer_23119_unrelaxed_rank_003_alphafold2_multimer_v3_model_4_seed_000.pd hivdimer_23119//hivdimer_23119_unrelaxed_rank_004_alphafold2_multimer_v3_model_1_seed_000.pd hivdimer_23119//hivdimer_23119_unrelaxed_rank_005_alphafold2_multimer_v3_model_3_seed_000.pd .....
```

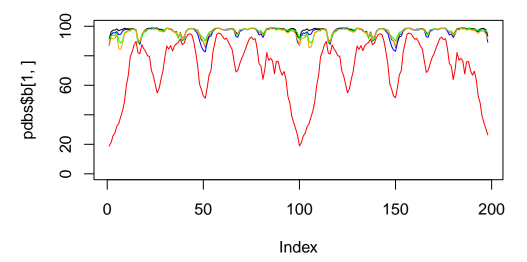
#### Extracting sequences

```
pdb/seq: 1 name: hivdimer_23119//hivdimer_23119_unrelaxed_rank_001_alphafold2_multimer_v3_rank_nose. 2 name: hivdimer_23119//hivdimer_23119_unrelaxed_rank_002_alphafold2_multimer_v3_rank_nose. 3 name: hivdimer_23119//hivdimer_23119_unrelaxed_rank_003_alphafold2_multimer_v3_rank_nose. 4 name: hivdimer_23119//hivdimer_23119_unrelaxed_rank_004_alphafold2_multimer_v3_rank_nose. 5 name: hivdimer_23119//hivdimer_23119_unrelaxed_rank_005_alphafold2_multimer_v3_rank_nose. 5 name: hivdimer_23119//hivdimer_23119_unrelaxed_rank_nose. 6 name: hivdimer_
```

```
library(bio3dview)
#view.pdbs(pdbs)
```

\$b is where the confidence score is stored for the model.

```
plot(pdbs$b[1,], typ ="1", ylim=c(0,100))
lines(pdbs$b[2,], typ="1", col="blue")
lines(pdbs$b[3,], typ="1", col="green")
lines(pdbs$b[4,], typ="1", col="orange")
lines(pdbs$b[5,], typ="1", col="red")
```



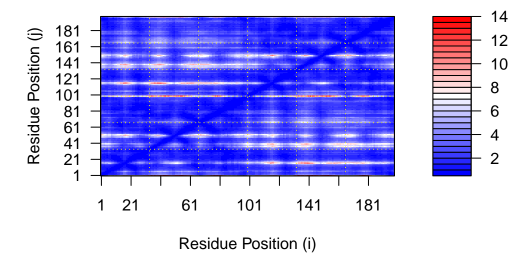
#Predicted alignment error for domains

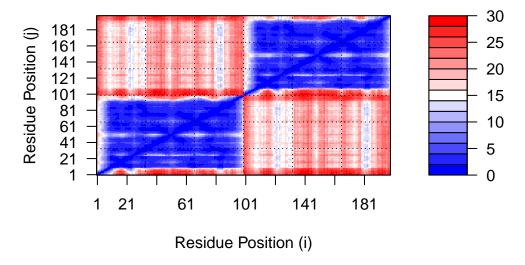
```
pae1 <- read_json(pae_files[1],simplifyVector = TRUE)
pae5 <- read_json(pae_files[5],simplifyVector = TRUE)
attributes(pae1)</pre>
```

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

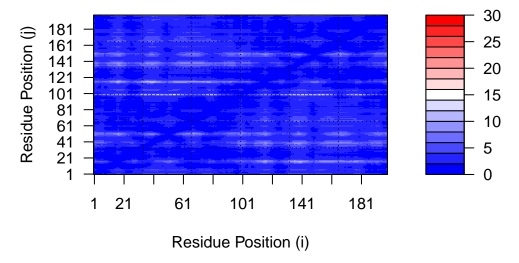
```
# Per-residue pLDDT scores
# same as B-factor of PDB..
head(pae1$plddt)
```

### [1] 91.44 96.06 97.38 97.38 98.19 96.94









# Score Residue Conservation from aligment file

AlphaFold returns it's large alignment file used for analysis. Here we read this file and score conservation per position

[1] "hivdimer\_23119//hivdimer\_23119.a3m"

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

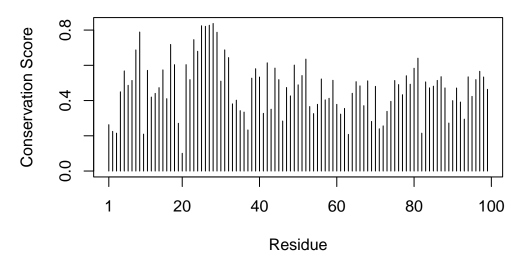
```
[1] " ** Duplicated sequence id's: 101 **"
[2] " ** Duplicated sequence id's: 101 **"
```

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

[1] 5378 132

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

```
plotb3(sim[1:99],
     ylab="Conservation Score")
```



Find the consensus sequence at a very high cut-off to find invarient residues

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

- [127] "-" "-" "-" "-" "-" "-"