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alphafold analysis

AUTHOR shivani

```
library(bio3d)
```

PDB File names of models

Align and superpose

```
pdbs <- pdbaln(files, fit=TRUE, exefile="msa")</pre>
Reading PDB files:
hiv monomer 94b5b//hiv monomer 94b5b unrelaxed rank 001 alphafold2 ptm model 5 seed 000.p
hiv_monomer_94b5b//hiv_monomer_94b5b_unrelaxed_rank_002_alphafold2_ptm_model_1_seed_000.p
hiv_monomer_94b5b//hiv_monomer_94b5b_unrelaxed_rank_003_alphafold2_ptm_model_4_seed_000.p
db
hiv_monomer_94b5b//hiv_monomer_94b5b_unrelaxed_rank_004_alphafold2_ptm_model_3_seed_000.p
hiv monomer 94b5b//hiv monomer 94b5b unrelaxed rank 005 alphafold2 ptm model 2 seed 000.p
db
. . . . .
Extracting sequences
pdb/seq: 1
             name:
hiv_monomer_94b5b//hiv_monomer_94b5b_unrelaxed_rank_001_alphafold2_ptm_model_5_seed_000.p
db
pdb/seq: 2
hiv_monomer_94b5b//hiv_monomer_94b5b_unrelaxed_rank_002_alphafold2_ptm_model_1_seed_000.p
db
pdb/seq: 3
             name:
hiv monomer 94b5b//hiv monomer 94b5b unrelaxed rank 003 alphafold2 ptm model 4 seed 000.p
db
pdb/seq: 4
             name:
hiv monomer 94b5b//hiv monomer 94b5b unrelaxed rank 004 alphafold2 ptm model 3 seed 000.p
db
```

RMSD analysis

pdb/seq: 5

db

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hiv_monomer_94b5b//hiv_monomer_94b5b_unrelaxed_rank_005_alphafold2_ptm_model_2_seed_000.p

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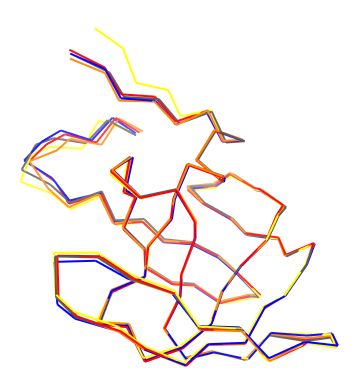
```
rd <- rmsd(pdbs)</pre>
```

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

```
mean(rd)
```

[1] 0.54368

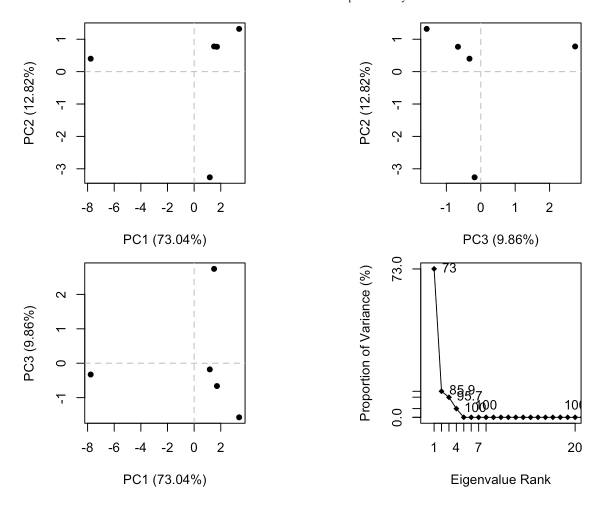
```
source("http://tinyurl.com/newviewngl")
library(NGLVieweR)
view.pdbs(pdbs)
```



#PCA

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

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Residue conservaation from alignment file

[1] "hiv_monomer_94b5b//hiv_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

score residue conservation:

```
sim <- conserv(aln)
```

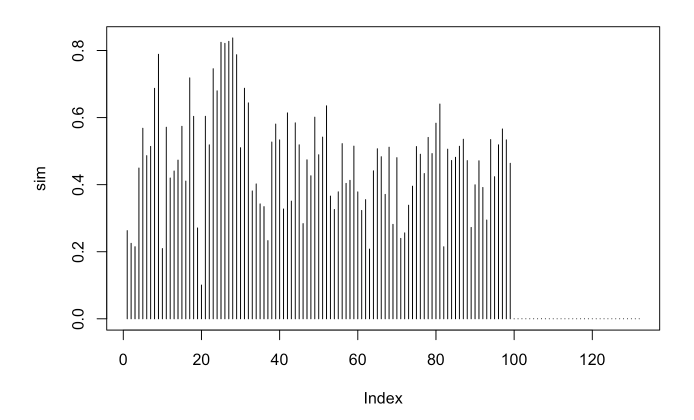
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```
con <- consensus(aln, cutoff = 0.9)
con$seq</pre>
```

plot the conservation along

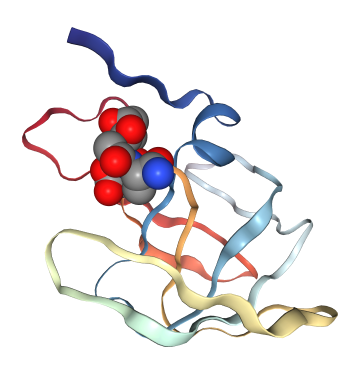
```
plot(sim, typ="h")
```



conserved positions in structure:

```
pdb <- read.pdb(files[1])
view.pdb(pdb, backgroundColor="pink", highlight=atom.select(pdb, resno=25:28), highlight.</pre>
```

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Instead of HIV Dimer analysis, Barry let us analyze our own novel protein sequence in Alphafold