

Class 11: Structural Bioinformatics pt 2

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AlphaFold Data Base (AFDB)

The EBI maintains the largest database of AlphaFold structure prediction models at:
<https://alphafold.ebi.ac.uk>

From last class (before Halloween) we saw that the PDB had 244,290 (Oct 2025)

The total number of protein sequences in UniProtKB is 199,579,901

Key Point: This is a tiny fraction of sequence space that has structural coverage
(0.12%)

```
244290 / 199579901 * 100
```

```
[1] 0.1224021
```

AFDB is attempting to address this gap...

There are two “Quality Scores” from AlphaFold. One for residues (i.e. each amino acid) called **pLDDT** score. The other **PAE** score measures the confidence in the relative position of two residues (i.e. a score for every pair of residues).

Generating your own structure predictions

Image of all 5 models

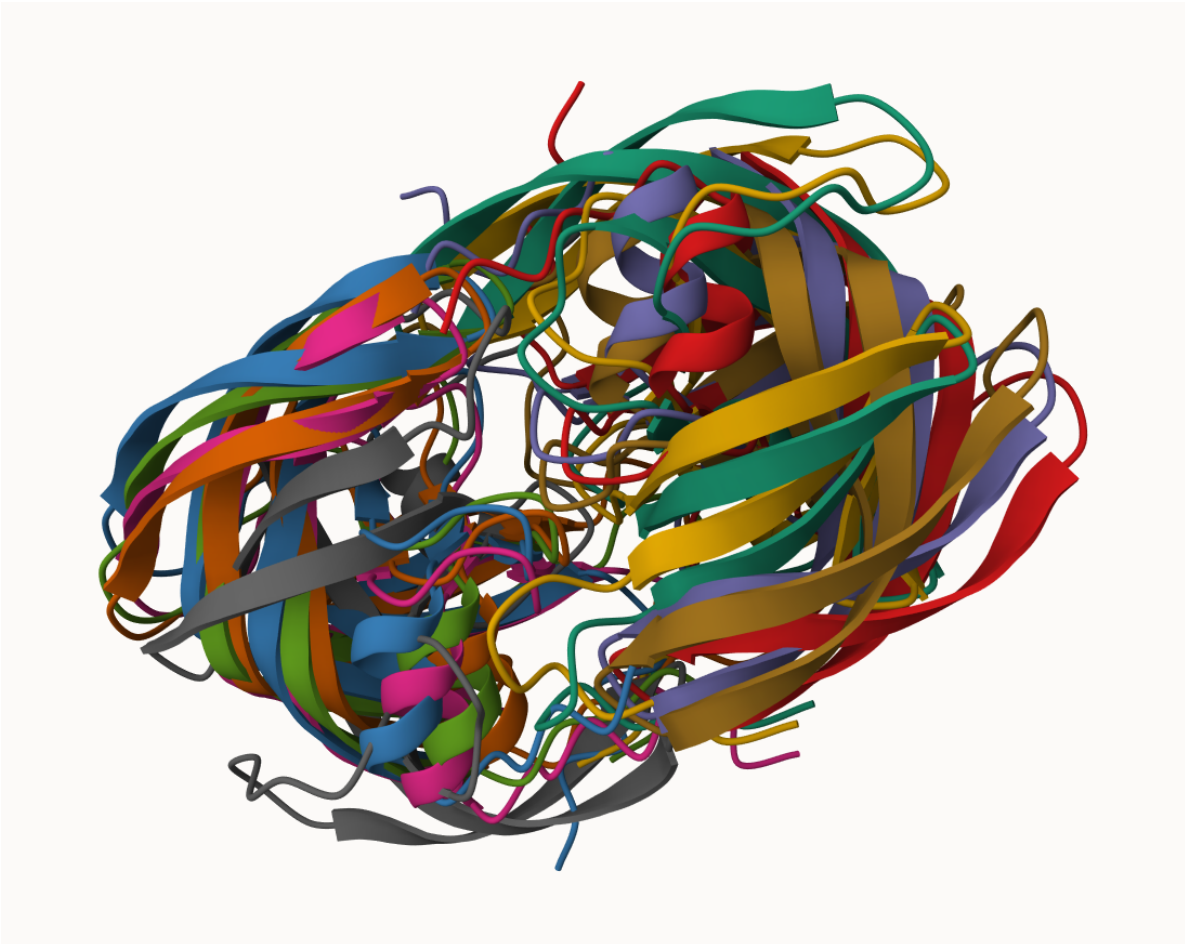


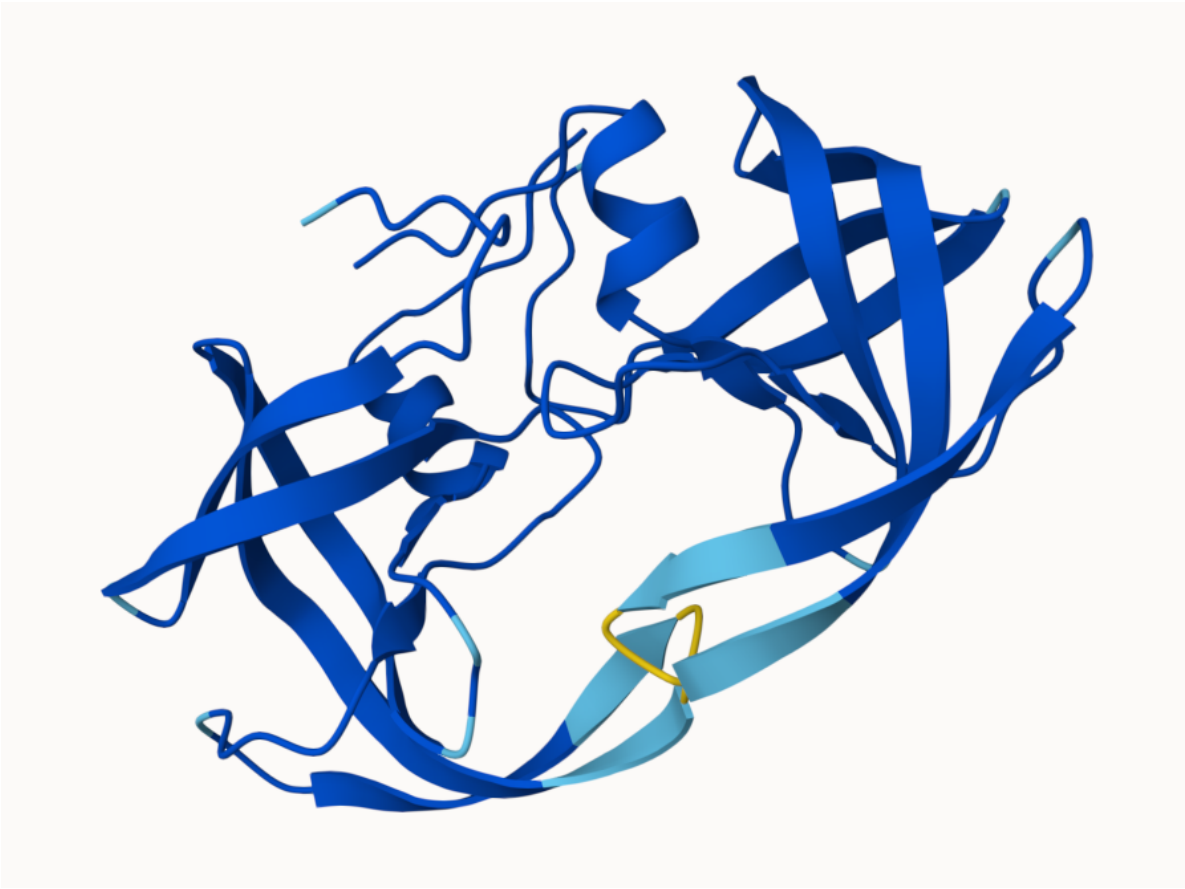
Image of the 1st models



pLDDT score of 1st model



pLDDT score of 4th model



Custom analysis of resulting models in R

Read key result file into R. The first thing I need to know is what my results directory/folder is called (i.e. its name is different for every AlphaFold run/job)

```
results_dir <- "HIVPR_dimer_23119/"

# File names for all PDB models
pdb_files <- list.files(path=results_dir,
                         pattern="*.pdb",
                         full.names = TRUE)

# Print our PDB file names
basename(pdb_files)
```

```
[1] "HIVPR_dimer_23119_unrelaxed_rank_001_alphaFold2_multimer_v3_model_4_seed_000.pdb"
```

```
[2] "HIVPR_dimer_23119_unrelaxed_rank_002_alphaFold2_multimer_v3_model_1_seed_000.pdb"
[3] "HIVPR_dimer_23119_unrelaxed_rank_003_alphaFold2_multimer_v3_model_5_seed_000.pdb"
[4] "HIVPR_dimer_23119_unrelaxed_rank_004_alphaFold2_multimer_v3_model_2_seed_000.pdb"
[5] "HIVPR_dimer_23119_unrelaxed_rank_005_alphaFold2_multimer_v3_model_3_seed_000.pdb"
```

```
library(bio3d)

m1 <- read.pdb(pdb_files[1])
m1
```

Call: read.pdb(file = pdb_files[1])

Total Models#: 1

Total Atoms#: 1514, XYZs#: 4542 Chains#: 2 (values: A B)

Protein Atoms#: 1514 (residues/Calpha atoms#: 198)

Nucleic acid Atoms#: 0 (residues/phosphate atoms#: 0)

Non-protein/nucleic Atoms#: 0 (residues: 0)

Non-protein/nucleic resid values: [none]

Protein sequence:

```
PQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWPKMIGGIGGFVKVRQYD  
QILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNFPQITLWQRPLVTIKIGGQLKE  
ALLDTGADDTVLEEMSLPGRWPKMIGGIGGFVKVRQYDQILIEICGHKAIGTVLVGPTP  
VNIIGRNLLTQIGCTLNF
```

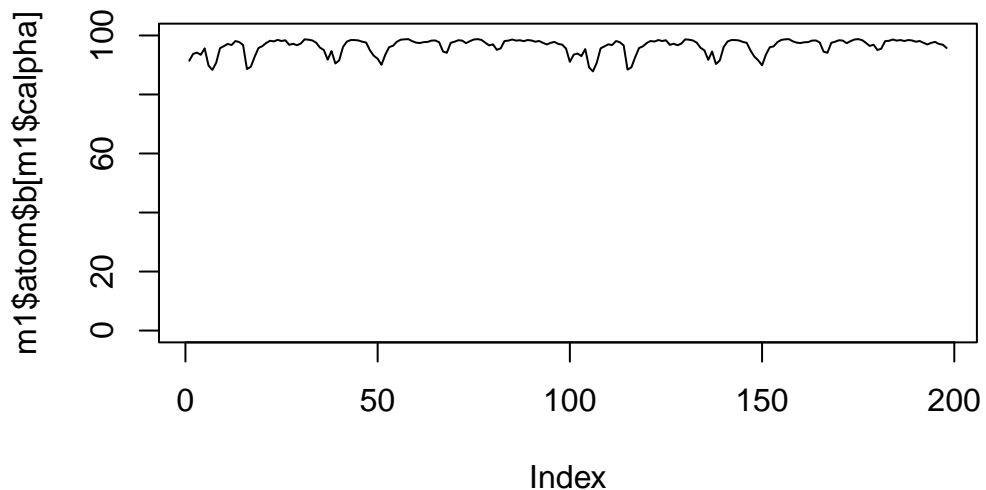
+ attr: atom, xyz, calpha, call

```
head(m1$atom)
```

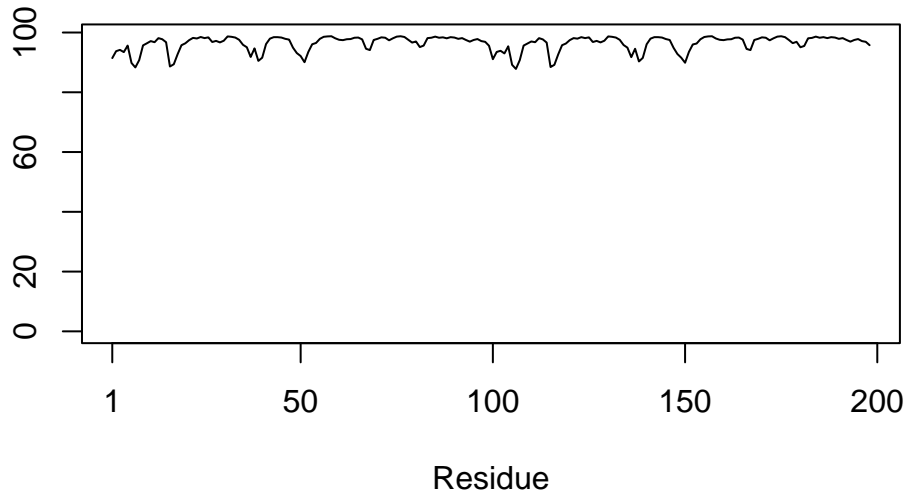
	type	eleno	elety	alt	resid	chain	resno	insert	x	y	z	o	b
1	ATOM	1	N	<NA>	PRO	A	1	<NA>	-16.656	5.527	4.973	1	91.44
2	ATOM	2	CA	<NA>	PRO	A	1	<NA>	-17.000	4.836	3.725	1	91.44
3	ATOM	3	C	<NA>	PRO	A	1	<NA>	-16.375	3.453	3.623	1	91.44
4	ATOM	4	CB	<NA>	PRO	A	1	<NA>	-16.453	5.773	2.643	1	91.44
5	ATOM	5	O	<NA>	PRO	A	1	<NA>	-15.445	3.137	4.375	1	91.44
6	ATOM	6	CG	<NA>	PRO	A	1	<NA>	-15.336	6.512	3.307	1	91.44
	segid	elesy	charge										
1	<NA>	N	<NA>										

```
2 <NA>      C  <NA>
3 <NA>      C  <NA>
4 <NA>      C  <NA>
5 <NA>      0  <NA>
6 <NA>      C  <NA>
```

```
plot( m1$atom$b[m1$calpha], typ="l", ylim=c(0,100))
```



```
plot.bio3d(m1$atom$b[m1$calpha], type="l")
```



Residue conservation from alignment file

Find the large AlphaFold alignment file

```
aln_file <- list.files(path=results_dir,
                        pattern=".a3m$",
                        full.names = TRUE)
aln_file
```

```
[1] "HIVPR_dimer_23119//HIVPR_dimer_23119.a3m"
```

Read this into R

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

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

How many sequences are in this alignment

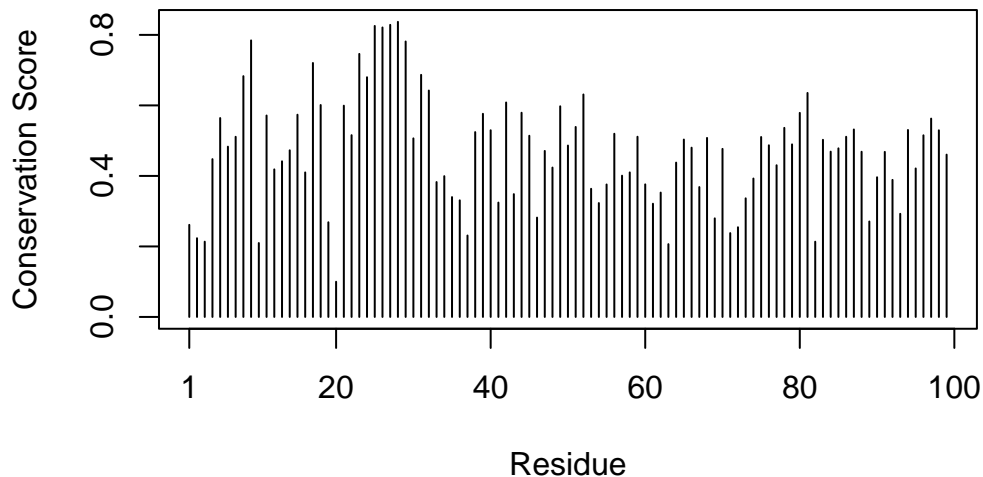
```
dim(aln$ali)
```

```
[1] 5397 132
```

We can score residue conservation in the alignment with the conserv() function.

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

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



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

```
[1] "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-"  
[19] "-" "-" "-" "-" "-" "D" "T" "G" "A" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-"  
[37] "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-"  
[55] "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-"  
[73] "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-"  
[91] "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-"  
[109] "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-" "-"  
[127] "-" "-" "-" "-" "-" "-"
```