Class 6 Homework

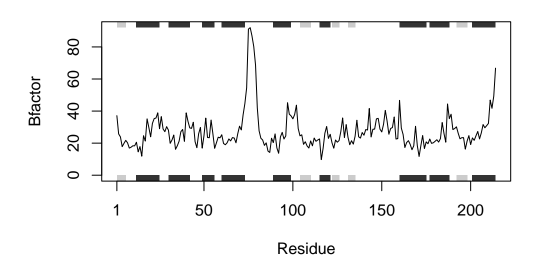
Mike Simonyan

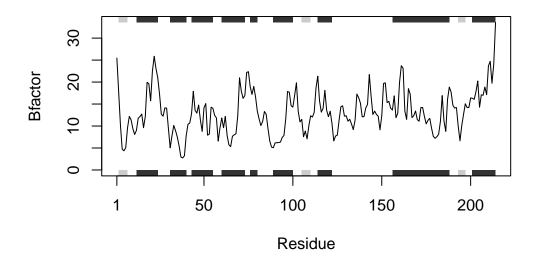
4/26/23

```
# Can you improve this analysis code?
library(bio3d)
s1 <- read.pdb("4AKE") # kinase with drug</pre>
Note: Accessing on-line PDB file
s2 <- read.pdb("1AKE") # kinase no drug
Note: Accessing on-line PDB file
PDB has ALT records, taking A only, rm.alt=TRUE
s3 <- read.pdb("1E4Y") # kinase with drug
Note: Accessing on-line PDB file
s1.chainA <- trim.pdb(s1, chain="A", elety="CA")</pre>
s2.chainA <- trim.pdb(s2, chain="A", elety="CA")</pre>
s3.chainA <- trim.pdb(s3, chain="A", elety="CA")</pre>
s1.b <- s1.chainA$atom$b</pre>
s2.b <- s2.chainA$atom$b</pre>
s3.b <- s3.chainA$atom$b
plotb3(s1.b, sse=s1.chainA, typ="l", ylab="Bfactor")
```



plotb3(s2.b, sse=s2.chainA, typ="1", ylab="Bfactor")





We will start by using read.pdb to read the protein.

```
protein_x <- read.pdb("1E4Y")</pre>
```

Note: Accessing on-line PDB file

Warning in get.pdb(file, path = tempdir(), verbose = FALSE): /var/folders/g7/6kj1lwl51s7_mw596z6h96y40000gn/T//Rtmp06Uyuv/1E4Y.pdb exists. Skipping download

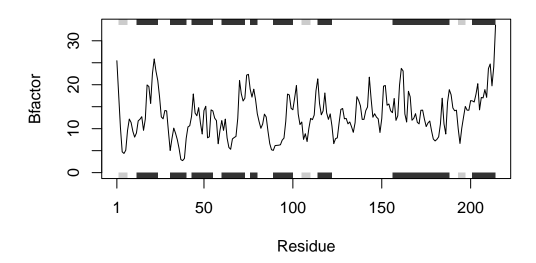
We will then use trim.pdb with chain ("A") and elety ("CA").

```
protein_x.chainA <- trim.pdb(protein_x, chain="A", elety="CA")</pre>
```

In order to for chain A to be specific to an atom, we must call this function.

```
protein_x.b <- protein_x.chainA$atom$b</pre>
```

Finally, we will plot a line graph (typ="l")



```
protein_x.chainA <- trim.pdb(protein_x, chain="A", elety="CA")
protein_x.b <- protein_x.chainA$atom$b
plotb3(protein_x.b, sse=protein_x.chainA, typ="l", ylab="Bfactor")
}

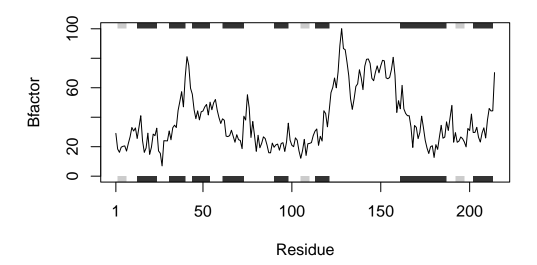
# test to see if the functoin can be generalized to work with any set of input protein str
plot_protein_x("4AKE")</pre>
```

Note: Accessing on-line PDB file

we are turning this into a function

plot_protein_x <- function(x){
 protein_x <- read.pdb(x)</pre>

Warning in get.pdb(file, path = tempdir(), verbose = FALSE): /var/folders/g7/6kj1lwl51s7_mw596z6h96y40000gn/T//Rtmp06Uyuv/4AKE.pdb exists. Skipping download

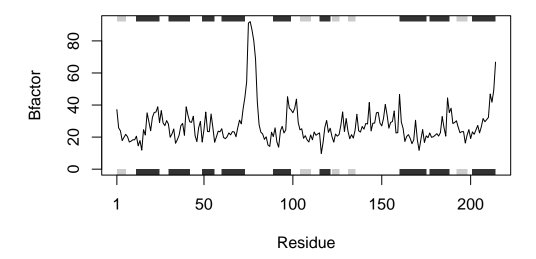


plot_protein_x("1AKE")

Note: Accessing on-line PDB file

Warning in get.pdb(file, path = tempdir(), verbose = FALSE): /var/folders/g7/6kj1lwl51s7_mw596z6h96y40000gn/T//Rtmp06Uyuv/1AKE.pdb exists. Skipping download

PDB has ALT records, taking A only, rm.alt=TRUE



plot_protein_x("1E4Y")

Note: Accessing on-line PDB file

Warning in get.pdb(file, path = tempdir(), verbose = FALSE):
/var/folders/g7/6kj1lwl51s7_mw596z6h96y40000gn/T//Rtmp06Uyuv/1E4Y.pdb exists.
Skipping download

