Class 6 HW - Write a Function

Jessica Le (PID: A17321021)

Can you improve this code analysis?

```
library(bio3d)
s1 <- read.pdb("4AKE") # kinase with drug

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

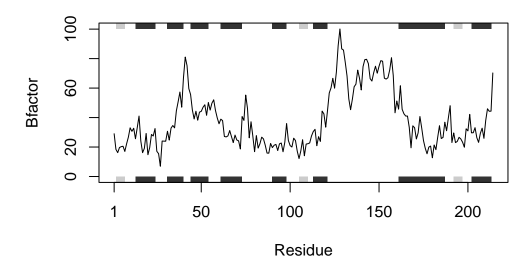
s3 <- read.pdb("1E4Y") # kinase with drug

Note: Accessing on-line PDB file
s1.chainA <- trim.pdb(s1, chain="A", elety="CA")
s2.chainA <- trim.pdb(s2, chain="A", elety="CA")
s3.chainA <- trim.pdb(s1, chain="A", elety="CA")
s1.b <- s1.chainA$atom$b
s2.b <- s2.chainA$atom$b
s2.b <- s2.chainA$atom$b
s3.b <- s3.chainA$atom$b
plotb3(s1.b, sse=s1.chainA, typ="l", ylab="Bfactor")</pre>
```



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





The generated function takes the input of a protein code from the Protein Data Bank to generate an output of a plot of the beta factor of the atoms in chain A of the specified protein. It will work with any set of input protein structures if plot_pdb is called.

```
library(bio3d)
plot_pdb <- function(pdb_code) {
   s <- read.pdb(pdb_code)
   s.chainA <- trim.pdb(s, chain="A", elety="CA")
   s.b <- s.chainA$atom$b
   plotb3(s.b, sse=s.chainA, typ="l", ylab="Bfactor")
}</pre>
```

This is a breakdown of the generated function.

```
#Call installed bio3d database
library(bio3d)

#Takes the input of a pdb code and creates pdb file that is read using `read.pdb`
plot_pdb <- function(pdb_code) {</pre>
```

```
#Reduces the PDB structures to a smaller subset of atoms using function `trim.pdb` and the is
s.chainA <- trim.pdb(s, chain="A", elety="CA")

#Selecting beta factor from atoms and the input is "s.chainA"
s.b <- s.chainA$atom$b

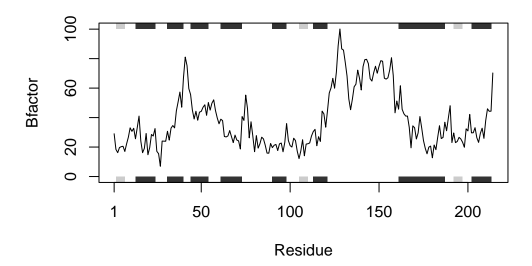
#Plotting the beta factor vs. residues for the specified atom where the input is "s.b"
plotb3(s.b, sse=s.chainA, typ="l", ylab="Bfactor")
}</pre>
```

Running the generated function by calling plot_pdb(), the output plots for the three specified proteins are.

```
sapply(c("4AKE", "1AKE", "1E4Y"), FUN=plot_pdb)
```

Note: Accessing on-line PDB file

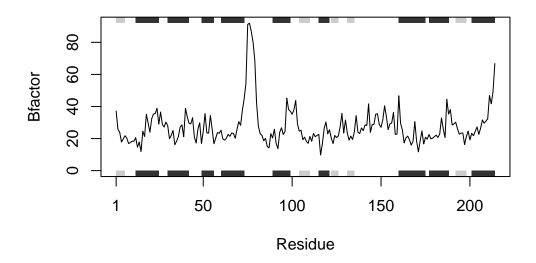
Warning in get.pdb(file, path = tempdir(), verbose = FALSE): /var/folders/s8/d27zs9y55db6qp39x9h69v9r0000gn/T//RtmpfCHHjJ/4AKE.pdb exists. Skipping download



Note: Accessing on-line PDB file

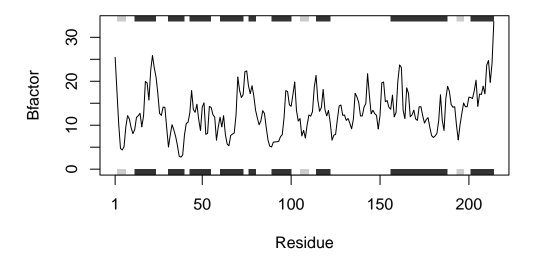
Warning in get.pdb(file, path = tempdir(), verbose = FALSE):
/var/folders/s8/d27zs9y55db6qp39x9h69v9r0000gn/T//RtmpfCHHjJ/1AKE.pdb exists.
Skipping download

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



Note: Accessing on-line PDB file

Warning in get.pdb(file, path = tempdir(), verbose = FALSE): /var/folders/s8/d27zs9y55db6qp39x9h69v9r0000gn/T//RtmpfCHHjJ/1E4Y.pdb exists. Skipping download



\$`4AKE` NULL

\$`1AKE`

NULL

\$`1E4Y`

NULL

"