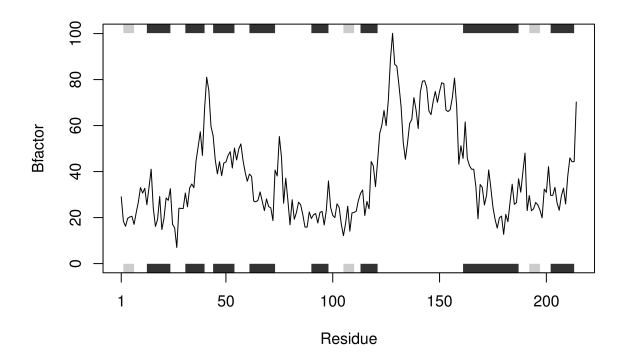
HW6

Yu Zhang 2019/10/18

```
library(bio3d)
prostr <- function(x, chain_name = "A", elety_name = "CA", yl = "Bfactor", atom_col = "b"){
    s <- read.pdb(x)
    s.chain <- trim.pdb(s, chain = chain_name, elety = elety_name)
    s.atom <- s.chain$atom[,atom_col]
    plotb3(s.atom, sse=s.chain, typ="l", ylab = yl)
}
prostr("4AKE")</pre>
```

Note: Accessing on-line PDB file



#Inputs: PDB ID (e.g., "4AKE")
#Optional Inputs: we can also specify which chain or atom or ylabel or elemet within dataframe atom we
#Process: the function first reads the pdb file and choose chainA, atom CA, and b, then makes a plot
#Output: the plot for the specified protein with x axis representing residues and y axis representing B