

# Lecture 6 Lab Homework

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```
#Load bio3d package into RStudio
library(bio3d)

#Create function that can be applied to any protein in PDB
#input for the function is protein identifier name
#output for function is a line plot for protein atomic chain data
protein_plot <- function(x){

  #read pdb file of input protein from protein database
  protein_file <- read.pdb(x)

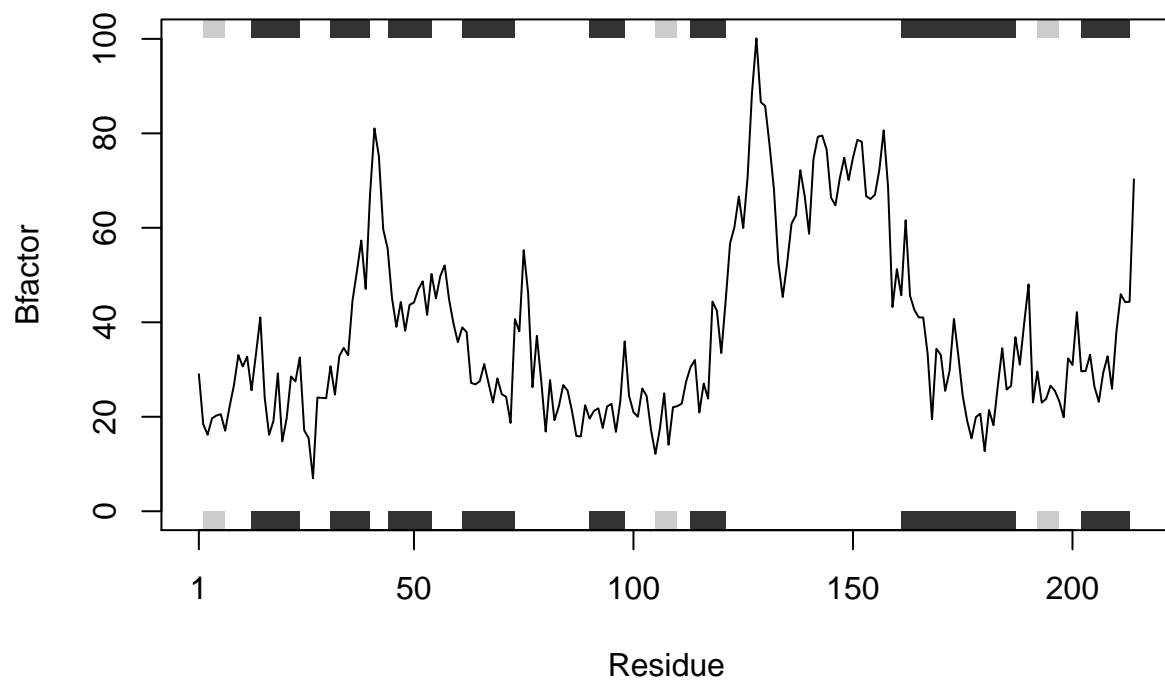
  #produce a smaller pdb file with subset of specified atom chain
  subset_chain <- trim.pdb(protein_file, chain="A", elety="CA")

  #Assign specified atom values in subset chain to variable
  atom_values <- subset_chain$atom$b

  #Plot the specified atom values in line graph
  plotb3(atom_values, sse=subset_chain, typ="l", ylab="Bfactor")
}

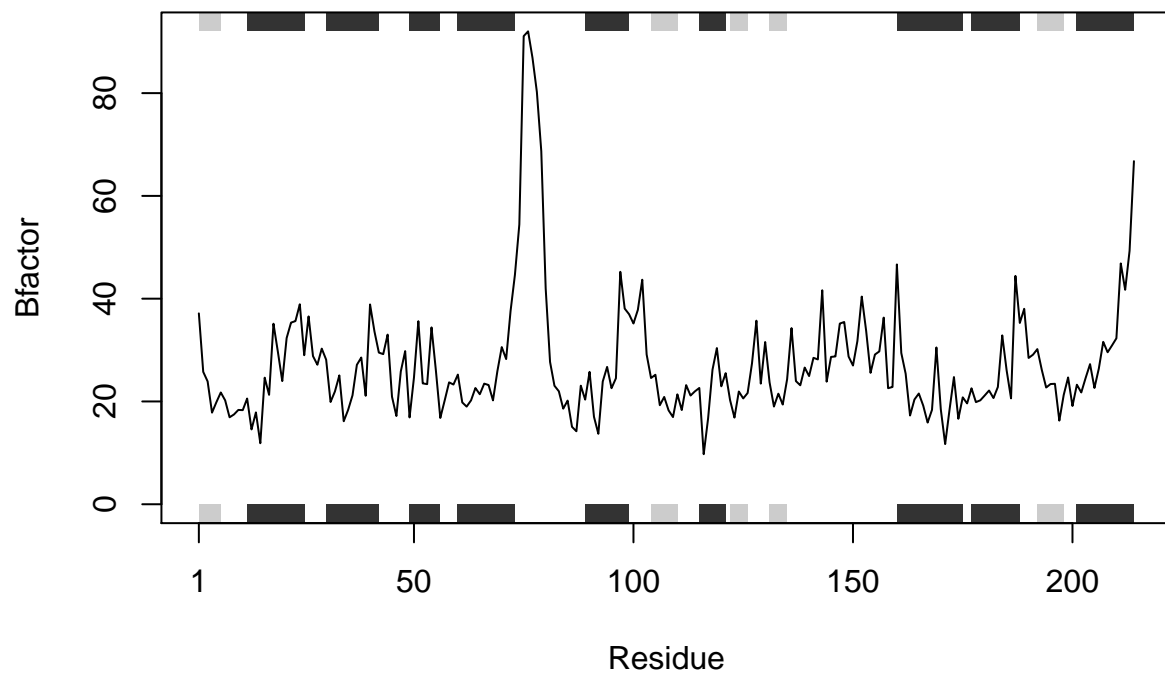
#examples for protein_plot() function
protein_plot("4AKE") #kinase with drug
```

## Note: Accessing on-line PDB file



```
protein_plot("1AKE") #kinase with no drug
```

```
## Note: Accessing on-line PDB file
## PDB has ALT records, taking A only, rm.alt=TRUE
```



```
protein_plot("1E4Y") #kinase with drug
```

```
## Note: Accessing on-line PDB file
```

