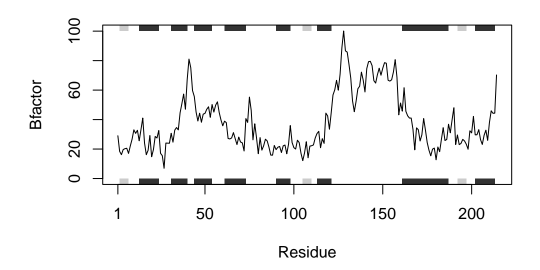
HW Q6

Kevin (A16482696)

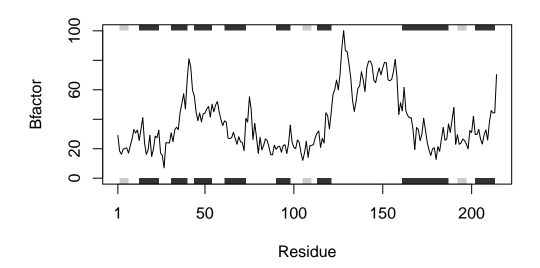
```
df <- data.frame(a=1:10, b=seq(200,400,length=10),c=11:20,d=NA)
dfa <- (dfa - min(dfa)) / (max(dfa) - min(dfa))
df$b <- (df$b - min(df$a)) / (max(df$b) - min(df$b))
dfc <- (dfc - min(dfc)) / (max(dfc) - min(dfc))
df$d <- (df$d - min(df$d)) / (max(df$a) - min(df$d))
library(bio3d)
s1 \leftarrow read.pdb("4AKE") # 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")</pre>
s2.chainA <- trim.pdb(s2, chain="A", elety="CA")</pre>
s3.chainA <- trim.pdb(s1, chain="A", elety="CA")</pre>
s1.b <- s1.chainA$atom$b</pre>
s2.b <- s2.chainA$atom$b
s3.b <- s3.chainA$atom$b
```



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



plotb3(s3.b, sse=s3.chainA, typ="1", ylab="Bfactor")



```
# create function to read general proteins
proteinclusters <- function(proteinstructures) {

# combine the proteins as vectors into matrices and combine those matricies
allproteins <- do.call(rbind,proteinstructures)

# measure the distance in those matricies
proteinmatrix <- dist(allproteins)

# use hierachical clustering to cluster the combined proteins
hc <- hclust(proteinmatrix)

#plot it
plot(hc)
}

# assign proteins to name
proteinstructures <- list(s1.b, s2.b, s3.b)
# perform the function
proteinclusters(proteinstructures)</pre>
```

Cluster Dendrogram



proteinmatrix hclust (*, "complete")