# lab6\_supp\_Q6

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### Can you improve this analysis code?

x <- read.pdb(pdb)

library(bio3d) s1 <- read.pdb("4AKE") # kinase with drug s2 <- read.pdb("1AKE") # kinase no drug s3 <- read.pdb("1E4Y") # kinase with drug 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.chainAatomb s2.b <- s2.chainAatomb s3.b <- s3.chainAatomb plotb3(s1.b, sse=s1.chainA, typ="l", ylab="Bfactor") plotb3(s2.b, sse=s2.chainA, typ="l", ylab="Bfactor") plotb3(s3.b, sse=s3.chainA, typ="l", ylab="Bfactor")

Q6. How would you generalize the original code above to work with any set of input protein structures?

Lets make a function that encompasses all the work of the code above in a single function.

First we need to open the Bio3d package using the library function.

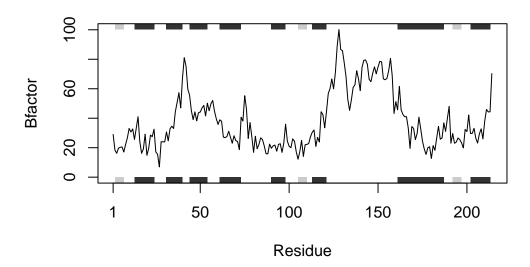
```
library(bio3d)
```

Now we will make a function encompassing the three requirments for a function: -name -input argument -function body

```
pdbs <- c("4AKE", "1AKE", "1E4Y" )
bfactor_analysis <- function(pdbs){
    #use lapply to process PDB file. lapply will apply the function to each element in our inposence of the second pdb function for the second pdb function for the pdb files and parse their contents into the second pdb function for the pdb files and parse their contents into the pdb files and parse the pdb fi
```

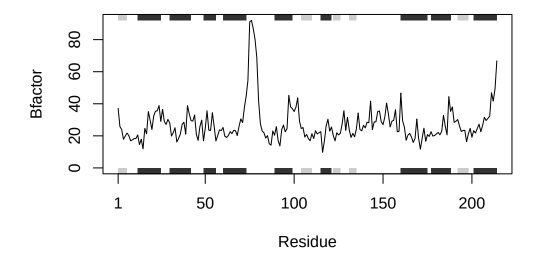
Note: Accessing on-line PDB file

## **B-factors for 4AKE**

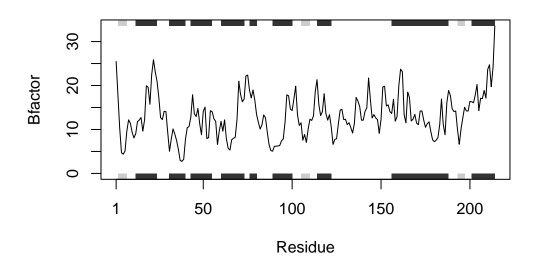


Note: Accessing on-line PDB file PDB has ALT records, taking A only, rm.alt=TRUE

# **B-factors for 1AKE**



## **B-factors for 1E4Y**



\$`4AKE`

\$`4AKE`\$structure

Call: read.pdb(file = pdb)

Total Models#: 1

Total Atoms#: 3459, XYZs#: 10377 Chains#: 2 (values: A B)

Protein Atoms#: 3312 (residues/Calpha atoms#: 428)
Nucleic acid Atoms#: 0 (residues/phosphate atoms#: 0)

Non-protein/nucleic Atoms#: 147 (residues: 147) Non-protein/nucleic resid values: [ HOH (147) ]

#### Protein sequence:

MRIILLGAPGAGKGTQAQFIMEKYGIPQISTGDMLRAAVKSGSELGKQAKDIMDAGKLVT DELVIALVKERIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFDVPDELIVDRI VGRRVHAPSGRVYHVKFNPPKVEGKDDVTGEELTTRKDDQEETVRKRLVEYHQMTAPLIG YYSKEAEAGNTKYAKVDGTKPVAEVRADLEKILGMRIILLGAPGA...<cut>...KILG + attr: atom, xyz, seqres, helix, sheet, calpha, remark, call

#### \$`4AKE`\$chain

Call: trim.pdb(pdb = x, chain = "A", elety = "CA")

Total Models#: 1

Total Atoms#: 214, XYZs#: 642 Chains#: 1 (values: A)

Protein Atoms#: 214 (residues/Calpha atoms#: 214) Nucleic acid Atoms#: 0 (residues/phosphate atoms#: 0)

Non-protein/nucleic Atoms#: 0 (residues: 0)
Non-protein/nucleic resid values: [ none ]

#### Protein sequence:

MRIILLGAPGAGKGTQAQFIMEKYGIPQISTGDMLRAAVKSGSELGKQAKDIMDAGKLVT DELVIALVKERIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFDVPDELIVDRI VGRRVHAPSGRVYHVKFNPPKVEGKDDVTGEELTTRKDDQEETVRKRLVEYHQMTAPLIG YYSKEAEAGNTKYAKVDGTKPVAEVRADLEKILG

+ attr: atom, helix, sheet, seqres, xyz, calpha, call

#### \$`4AKE`\$bfactor

[1] 29.02 18.44 16.20 19.67 20.26 20.55 17.05 22.13 26.71 33.05 [11] 30.66 32.73 25.61 33.19 41.03 24.09 16.18 19.14 29.19 14.79 23.94 [21] 19.63 28.54 27.49 32.56 17.13 15.50 6.98 24.07 24.00 [31] 30.70 24.70 32.84 34.60 33.01 44.60 50.74 57.32 47.04 67.13 [41] 81.04 75.20 59.68 55.63 45.12 39.04 44.31 38.21 43.70 44.19 [51] 47.00 48.67 41.54 50.22 45.07 49.77 52.04 44.82 39.75 35.79 [61] 38.92 37.93 27.18 26.86 27.53 31.16 27.08 23.03 28.12 24.78 [71] 24.22 18.69 40.67 38.08 55.26 46.29 26.25 37.14 27.50 16.86 [81] 27.76 19.27 22.22 26.70 25.52 21.22 15.90 15.84 22.44 19.61 22.19 22.73 16.80 23.25 35.95 24.42 20.96 [91] 21.23 21.79 17.64 [101] 20.00 25.99 24.39 17.19 12.16 17.35 24.97 14.08 22.01 22.26 [111] 22.78 27.47 30.49 32.02 20.90 27.03 23.84 44.37 42.47 33.48 [121] 44.56 56.67 60.18 66.62 59.95 70.81 88.63 100.11 86.60 85.80 [131] 77.48 68.13 52.66 45.34 52.43 60.90 62.64 72.19 66.75 58.73 [141] 74.57 79.29 79.53 76.58 66.40 64.76 70.48 74.84 70.11 74.82 [151] 78.61 78.24 66.70 66.10 67.01 72.28 80.64 68.54 43.23 51.24 [161] 45.72 61.60 45.61 42.57 41.03 41.02 33.34 19.48 34.38 33.11

```
[171] 25.48 29.68 40.71 32.91 24.41 19.20 15.43 19.93 20.66 12.72
[181] 21.40 18.21 26.68 34.50 25.77 26.52 36.85 31.05 39.84 48.03
[191] 23.04 29.57 23.00 23.80 26.59 25.49 23.25 19.89 32.37 30.97
[201] 42.16 29.64 29.69 33.15 26.38 23.17 29.35 32.80 25.92 38.01
[211] 45.95 44.26 44.35 70.26
$ 1AKE 1
$`1AKE`$structure
Call: read.pdb(file = pdb)
  Total Models#: 1
    Total Atoms#: 3804, XYZs#: 11412 Chains#: 2 (values: A B)
    Protein Atoms#: 3312 (residues/Calpha atoms#: 428)
    Nucleic acid Atoms#: 0 (residues/phosphate atoms#: 0)
    Non-protein/nucleic Atoms#: 492 (residues: 380)
    Non-protein/nucleic resid values: [ AP5 (2), HOH (378) ]
  Protein sequence:
     MRIILLGAPGAGKGTQAQFIMEKYGIPQISTGDMLRAAVKSGSELGKQAKDIMDAGKLVT
     DELVIALVKERIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFDVPDELIVDRI
     VGRRVHAPSGRVYHVKFNPPKVEGKDDVTGEELTTRKDDQEETVRKRLVEYHQMTAPLIG
     YYSKEAEAGNTKYAKVDGTKPVAEVRADLEKILGMRIILLGAPGA...<br/>cut>...KILG
+ attr: atom, xyz, seqres, helix, sheet,
       calpha, remark, call
$`1AKE`$chain
Call: trim.pdb(pdb = x, chain = "A", elety = "CA")
  Total Models#: 1
    Total Atoms#: 214, XYZs#: 642 Chains#: 1 (values: A)
    Protein Atoms#: 214 (residues/Calpha atoms#: 214)
    Nucleic acid Atoms#: 0 (residues/phosphate atoms#: 0)
    Non-protein/nucleic Atoms#: 0 (residues: 0)
    Non-protein/nucleic resid values: [ none ]
```

#### Protein sequence:

MRIILLGAPGAGKGTQAQFIMEKYGIPQISTGDMLRAAVKSGSELGKQAKDIMDAGKLVT DELVIALVKERIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFDVPDELIVDRI VGRRVHAPSGRVYHVKFNPPKVEGKDDVTGEELTTRKDDQEETVRKRLVEYHQMTAPLIG YYSKEAEAGNTKYAKVDGTKPVAEVRADLEKILG

+ attr: atom, helix, sheet, seqres, xyz, calpha, call \$`1AKE`\$bfactor [1] 37.14 25.76 23.90 17.83 19.86 21.75 20.21 16.92 17.47 18.35 18.31 20.57 [13] 14.56 17.87 11.87 24.63 21.29 35.13 29.68 23.96 32.34 35.34 35.64 38.91 [25] 29.00 36.55 28.83 27.15 30.28 28.13 19.90 21.95 25.07 16.15 18.35 21.19 [37] 27.13 28.55 21.10 38.88 33.63 29.51 29.21 33.01 20.92 17.17 25.84 29.80 [49] 16.89 24.66 35.62 23.52 23.37 34.41 25.96 16.79 20.20 23.72 23.29 25.23 [61] 19.81 19.00 20.21 22.62 21.40 23.47 23.20 20.21 25.90 30.58 28.25 37.60 [73] 44.66 54.46 91.10 92.02 86.85 80.21 68.72 42.01 27.69 23.06 21.98 18.60 [85] 20.17 15.06 14.20 23.07 20.36 25.76 17.02 13.71 23.88 26.72 22.58 24.51 [97] 45.23 38.07 36.97 35.17 37.83 43.69 29.14 24.56 25.20 19.27 20.88 18.27 [109] 16.96 21.38 18.33 23.18 21.15 21.97 22.63 9.74 16.71 26.18 30.39 22.95 [121] 25.51 20.28 16.86 21.94 20.59 21.64 27.42 35.72 23.47 31.57 23.71 19.01 [133] 21.52 19.40 24.32 34.28 23.96 23.14 26.60 24.94 28.49 28.18 41.64 23.85 [145] 28.67 28.76 35.16 35.46 28.74 26.99 31.74 40.41 33.73 25.57 29.13 29.74 [157] 36.32 22.58 22.82 46.67 29.44 25.40 17.27 20.38 21.55 19.19 15.89 18.37 [169] 30.51 18.47 11.70 18.45 24.75 16.63 20.80 19.62 22.56 19.87 20.22 21.16 [181] 22.13 20.66 22.82 32.86 26.04 20.60 44.44 35.28 38.03 28.46 29.10 30.19 [193] 26.17 22.71 23.39 23.44 16.27 21.26 24.67 19.12 23.26 21.75 24.59 27.26 [205] 22.63 26.40 31.60 29.57 30.90 32.29 46.86 41.73 49.31 66.76 \$`1E4Y` \$`1E4Y`\$structure Call: read.pdb(file = pdb) Total Models#: 1 Total Atoms#: 3428, XYZs#: 10284 Chains#: 2 (values: A B) Protein Atoms#: 3314 (residues/Calpha atoms#: 428) Nucleic acid Atoms#: 0 (residues/phosphate atoms#: 0) Non-protein/nucleic Atoms#: 114 (residues: 2)

Non-protein/nucleic resid values: [ AP5 (2) ]

#### Protein sequence:

MRIILLGALVAGKGTQAQFIMEKYGIPQISTGDMLRAAVKSGSELGKQAKDIMDAGKLVT DELVIALVKERIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFDVPDELIVDRI VGRRVHAPSGRVYHVKFNPPKVEGKDDVTGEELTTRKDDQEETVRKRLVEYHQMTAPLIG YYSKEAEAGNTKYAKVDGTKPVAEVRADLEKILGMRIILLGALVA...<cut>...KILG

+ attr: atom, xyz, seqres, helix, sheet, calpha, remark, call

#### \$`1E4Y`\$chain

Call: trim.pdb(pdb = x, chain = "A", elety = "CA")

Total Models#: 1

Total Atoms#: 214, XYZs#: 642 Chains#: 1 (values: A)

Protein Atoms#: 214 (residues/Calpha atoms#: 214)
Nucleic acid Atoms#: 0 (residues/phosphate atoms#: 0)

Non-protein/nucleic Atoms#: 0 (residues: 0)
Non-protein/nucleic resid values: [ none ]

#### Protein sequence:

MRIILLGALVAGKGTQAQFIMEKYGIPQISTGDMLRAAVKSGSELGKQAKDIMDAGKLVT DELVIALVKERIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFDVPDELIVDRI VGRRVHAPSGRVYHVKFNPPKVEGKDDVTGEELTTRKDDQEETVRKRLVEYHQMTAPLIG YYSKEAEAGNTKYAKVDGTKPVAEVRADLEKILG

+ attr: atom, helix, sheet, seqres, xyz, calpha, call

#### \$`1E4Y`\$bfactor

[1] 25.46 17.86 10.28 4.73 4.36 5.10 9.59 12.19 11.41 9.39 8.08 9.01 [13] 11.77 12.15 12.72 9.62 12.18 19.95 19.59 15.73 22.51 25.87 23.08 20.97 [25] 17.28 12.69 12.24 14.14 14.05 9.38 5.03 7.78 10.13 8.96 7.50 5.48 [37] 2.97 2.73 3.23 7.81 10.40 10.67 12.79 17.90 13.56 12.94 14.78 11.31 [49] 8.79 14.13 15.10 7.92 8.15 14.28 14.04 12.42 11.84 6.57 9.59 11.84 [61] 9.61 12.18 7.89 5.74 5.31 7.67 7.99 8.24 12.34 20.98 17.93 16.30 [73] 16.94 22.19 22.36 18.96 17.18 18.99 16.65 13.39 11.61 10.10 11.03 13.31 [85] 12.66 9.44 6.60 5.20 5.06 6.16 6.20 6.24 6.34 7.39 7.86 11.66 [97] 17.87 17.67 14.63 14.30 16.98 19.84 13.36 10.93 11.52 7.56 8.85 7.07 [109] 10.08 12.34 12.05 13.10 18.63 21.34 15.73 13.16 14.04 18.13 13.59 12.12

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[121] 13.37 10.57 6.60 7.73 7.91 11.31 14.38 14.60 12.25 12.33 11.10 11.53 [133] 10.44 9.18 11.36 17.28 16.45 15.21 12.11 12.12 14.10 14.94 21.72 16.82 [145] 12.61 13.40 12.64 12.24 9.13 12.31 19.68 19.83 15.34 15.61 14.07 13.64 [157] 16.87 11.89 12.92 19.93 23.72 23.13 13.35 11.51 18.51 17.24 11.92 12.36 [169] 13.42 11.45 11.09 14.19 14.22 12.15 10.49 11.29 11.74 9.53 7.65 7.21 [181] 7.56 8.14 11.07 16.93 11.12 8.79 16.03 18.87 17.72 14.72 14.08 14.21 [193] 9.99 6.63 10.11 12.64 15.06 14.21 14.20 16.39 16.31 16.07 17.83 20.24 [205] 14.28 17.10 17.00 18.88 17.13 23.68 24.72 19.74 24.12 33.57
```