class 10

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Comparative structure analysis of Adenylate Kinase

adenelate kinase (ADK) is an important drug target and we would love to know how it works.

We will perform PCA on the complete collection of ADK structures in the protein data-bank (PDB) to reveal detailed features and mechanistic principles of these essential shape changing transitions.

First install the packages needed > Q10. Which of the packages above is found only on BioConductor and not CRAN?

msa

Q11. Which of the above packages is not found on BioConductor or CRAN?:

Grantlab/bio3d-view

Q12. True or False? Functions from the devtools package can be used to install packages from GitHub and BitBucket?

TRUE.

Search and retrieve ADK structures

Fetch the query sequence for chain A of the PDB ID 1AKE (example ADK sequence). We will use this to find all ADK structures in the PDB.

```
library(bio3d)
aa <- get.seq("1ake_A")</pre>
```

Warning in get.seq("lake_A"): Removing existing file: seqs.fasta

Fetching... Please wait. Done.

aa

```
60
             MRIILLGAPGAGKGTQAQFIMEKYGIPQISTGDMLRAAVKSGSELGKQAKDIMDAGKLVT
pdb|1AKE|A
            61
                                                                            120
             DELVIALVKERIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFDVPDELIVDRI
           121
                                                                            180
pdb|1AKE|A
             VGRRVHAPSGRVYHVKFNPPKVEGKDDVTGEELTTRKDDQEETVRKRLVEYHQMTAPLIG
           121
                                                                            180
           181
                                                214
pdb|1AKE|A
             YYSKEAEAGNTKYAKVDGTKPVAEVRADLEKILG
                                                214
Call:
  read.fasta(file = outfile)
Class:
  fasta
Alignment dimensions:
  1 sequence rows; 214 position columns (214 non-gap, 0 gap)
+ attr: id, ali, call
     Q13. How many amino acids are in this sequence, i.e. how long is this sequence?
214 amino acids.
Now we can use this sequence as a query to BLAST search the PDB to find similar sequences
```

Blast or hmmer search

and structures.

```
# Plot a summary of search results
#hits <- plot(b)</pre>
```

Let's see what's in hits object.

```
#hits$pdb.id
```

Because we do not want to rerun the code above everytime, we just manually spell the hits vector

```
hits <- NULL
  hits$pdb.id <- c('1AKE_A','6S36_A','6RZE_A','3HPR_A','1E4V_A','5EJE_A','1E4Y_A','3X2S_A','
  # Download releated PDB files
  files <- get.pdb(hits$pdb.id, path="pdbs", split=TRUE, gzip=TRUE)</pre>
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/1AKE.pdb exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/6S36.pdb exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/6RZE.pdb exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/3HPR.pdb exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/1E4V.pdb exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/5EJE.pdb exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/1E4Y.pdb exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/3X2S.pdb exists. Skipping download
```

Warning in get.pdb(hits\$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE): pdbs/6HAP.pdb exists. Skipping download

Warning in get.pdb(hits\$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE): pdbs/6HAM.pdb exists. Skipping download

Warning in get.pdb(hits\$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE): pdbs/4K46.pdb exists. Skipping download

Warning in get.pdb(hits\$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE): pdbs/3GMT.pdb exists. Skipping download

Warning in get.pdb(hits\$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE): pdbs/4PZL.pdb exists. Skipping download

1		
	I	0%
 =====	I	8%
 ==========	1	15%
 ===================================	I	23%
 ===================================	I	31%
 ===================================	I	38%
ı ====================================	1	46%
' ======= !	I	54%
' ======== !		62%
ı ====================================	1	69%
 	I	77%
 ===================================	1	85%
l .		

Align and superpose structures

```
Align and superpose these structures which are all over place.
  # Align releated PDBs
  pdbs <- pdbaln(files, fit = TRUE, exefile="msa")</pre>
Reading PDB files:
pdbs/split_chain/1AKE_A.pdb
pdbs/split_chain/6S36_A.pdb
pdbs/split_chain/6RZE_A.pdb
pdbs/split_chain/3HPR_A.pdb
pdbs/split_chain/1E4V_A.pdb
pdbs/split_chain/5EJE_A.pdb
pdbs/split_chain/1E4Y_A.pdb
pdbs/split_chain/3X2S_A.pdb
pdbs/split_chain/6HAP_A.pdb
pdbs/split_chain/6HAM_A.pdb
pdbs/split_chain/4K46_A.pdb
pdbs/split_chain/3GMT_A.pdb
pdbs/split_chain/4PZL_A.pdb
   PDB has ALT records, taking A only, rm.alt=TRUE
     PDB has ALT records, taking A only, rm.alt=TRUE
       PDB has ALT records, taking A only, rm.alt=TRUE
    PDB has ALT records, taking A only, rm.alt=TRUE
. . .
Extracting sequences
             name: pdbs/split_chain/1AKE_A.pdb
pdb/seq: 1
   PDB has ALT records, taking A only, rm.alt=TRUE
             name: pdbs/split_chain/6S36_A.pdb
   PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 3
             name: pdbs/split_chain/6RZE_A.pdb
```

PDB has ALT records, taking A only, rm.alt=TRUE pdb/seq: 4 name: pdbs/split_chain/3HPR_A.pdb PDB has ALT records, taking A only, rm.alt=TRUE pdb/seq: 5 name: pdbs/split_chain/1E4V_A.pdb name: pdbs/split chain/5EJE A.pdb pdb/seq: 6 PDB has ALT records, taking A only, rm.alt=TRUE pdb/seq: 7 name: pdbs/split chain/1E4Y A.pdb pdb/seq: 8 name: pdbs/split_chain/3X2S_A.pdb pdb/seq: 9 name: pdbs/split_chain/6HAP_A.pdb pdb/seq: 10 name: pdbs/split_chain/6HAM_A.pdb PDB has ALT records, taking A only, rm.alt=TRUE name: pdbs/split_chain/4K46_A.pdb pdb/seq: 11 PDB has ALT records, taking A only, rm.alt=TRUE pdb/seq: 12 name: pdbs/split_chain/3GMT_A.pdb pdb/seq: 13 name: pdbs/split_chain/4PZL_A.pdb

Let's have a look at our pdbs object.

pdbs

[Truncated_Name:1]1AKE_A.pdb [Truncated_Name:2]6S36_A.pdb [Truncated_Name:3]6RZE_A.pdb [Truncated_Name:4]3HPR_A.pdb

40 ----MRIILLGAPGAGKGTQAQFIMEKYGIPQIS -----MRIILLGAPGAGKGTQAQFIMEKYGIPQIS -----MRIILLGAPGAGKGTQAQFIMEKYGIPQIS ----MRIILLGAPGAGKGTQAQFIMEKYGIPQIS ----MRIILLGAPVAGKGTQAQFIMEKYGIPQIS ----MRIILLGAPGAGKGTQAQFIMEKYGIPQIS -----MRIILLGALVAGKGTQAQFIMEKYGIPQIS ----MRIILLGAPGAGKGTQAQFIMEKYGIPQIS -----MRIILLGAPGAGKGTQAQFIMEKYGIPQIS -----MRIILLGAPGAGKGTQAQFIMEKYGIPQIS ----MRIILLGAPGAGKGTQAQFIMAKFGIPQIS -----MRLILLGAPGAGKGTQANFIKEKFGIPQIS TENLYFQSNAMRIILLGAPGAGKGTQAKIIEQKYNIAHIS 1 40

41 80
TGDMLRAAVKSGSELGKQAKDIMDAGKLVTDELVIALVKE
TGDMLRAAVKSGSELGKQAKDIMDAGKLVTDELVIALVKE
TGDMLRAAVKSGSELGKQAKDIMDAGKLVTDELVIALVKE
TGDMLRAAVKSGSELGKQAKDIMDAGKLVTDELVIALVKE

[Truncated_Name:5] 1E4V_A.pdb [Truncated_Name:6] 5EJE_A.pdb [Truncated_Name:7] 1E4Y_A.pdb [Truncated_Name:8] 3X2S_A.pdb [Truncated_Name:9] 6HAP_A.pdb [Truncated_Name:10] 6HAM_A.pdb [Truncated_Name:11] 4K46_A.pdb [Truncated_Name:12] 3GMT_A.pdb [Truncated_Name:13] 4PZL_A.pdb TGDMLRAAVKSGSELGKQAKDIMDAGKLVTDELVIALVKE
TGDMLRAAVKSGSELGKQAKDIMDACKLVTDELVIALVKE
TGDMLRAAVKSGSELGKQAKDIMDAGKLVTDELVIALVKE
TGDMLRAAVKSGSELGKQAKDIMDCGKLVTDELVIALVKE
TGDMLRAAVKSGSELGKQAKDIMDAGKLVTDELVIALVRE
TGDMLRAAIKSGSELGKQAKDIMDAGKLVTDEIIIALVKE
TGDMLRAAIKAGTELGKQAKSVIDAGQLVSDDIILGLVKE
TGDMLRAAVKAGTPLGVEAKTYMDEGKLVPDSLIIGLVKE
TGDMIRETIKSGSALGQELKKVLDAGELVSDEFIIKIVKD

[Truncated_Name:1]1AKE_A.pdb
[Truncated_Name:2]6S36_A.pdb
[Truncated_Name:3]6RZE_A.pdb
[Truncated_Name:4]3HPR_A.pdb
[Truncated_Name:5]1E4V_A.pdb
[Truncated_Name:6]5EJE_A.pdb
[Truncated_Name:7]1E4Y_A.pdb
[Truncated_Name:8]3X2S_A.pdb
[Truncated_Name:9]6HAP_A.pdb
[Truncated_Name:10]6HAM_A.pdb
[Truncated_Name:11]4K46_A.pdb
[Truncated_Name:12]3GMT_A.pdb
[Truncated_Name:12]3GMT_A.pdb

RIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFD
RIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFD
RIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFD
RIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFD
RIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFD
RIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFD
RIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFD
RIAQEDSRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFD
RICQEDSRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFD
RICQEDSRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFD
RICQEDSRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFD
RIAQDDCAKGFLLDGFPRTIPQADGLKEVGVVVDYVIEFD
RIKEADCANGYLFDGFPRTIAQADAMKEAGVAIDYVLEID
RISKNDCNNGFLLDGVPRTIPQAQELDKLGVNIDYIVEVD

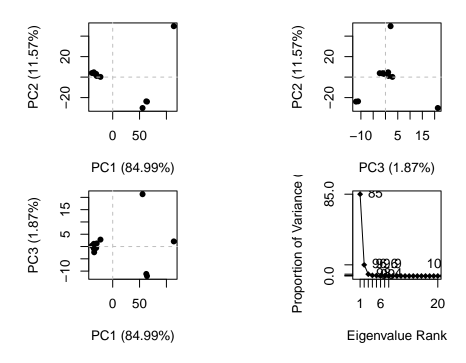
121 . . . 160

[Truncated_Name:1]1AKE_A.pdb
[Truncated_Name:2]6S36_A.pdb
[Truncated_Name:3]6RZE_A.pdb
[Truncated_Name:4]3HPR_A.pdb
[Truncated_Name:5]1E4V_A.pdb
[Truncated_Name:6]5EJE_A.pdb
[Truncated_Name:7]1E4Y_A.pdb
[Truncated_Name:8]3X2S_A.pdb
[Truncated_Name:9]6HAP_A.pdb
[Truncated_Name:10]6HAM_A.pdb
[Truncated_Name:11]4K46_A.pdb
[Truncated_Name:12]3GMT_A.pdb
[Truncated_Name:13]4PZL_A.pdb

VPDELIVDRIVGRRVHAPSGRVYHVKFNPPKVEGKDDVTG
VPDELIVDKIVGRRVHAPSGRVYHVKFNPPKVEGKDDVTG
VPDELIVDAIVGRRVHAPSGRVYHVKFNPPKVEGKDDVTG
VPDELIVDRIVGRRVHAPSGRVYHVKFNPPKVEGKDDVTG
VPDELIVDRIVGRRVHAPSGRVYHVKFNPPKVEGKDDVTG
VPDELIVDRIVGRRVHAPSGRVYHVKFNPPKVEGKDDVTG
VPDELIVDRIVGRRVHAPSGRVYHVKFNPPKVEGKDDVTG
VPDELIVDRIVGRRVHAPSGRVYHVKFNPPKVEGKDDVTG
VPDELIVDRIVGRRVHAPSGRVYHVKFNPPKVEGKDDVTG
VPDELIVDRIVGRRVHAPSGRVYHVKFNPPKVEGKDDVTG
VPDELIVDRIVGRRVHAPSGRVYHVKFNPPKVEGKDDVTG
VADSVIVERMAGRRAHLASGRTYHNVYNPPKVEGKDDVTG
VADNLLIERITGRRIHPASGRTYHVKFNPPKVEGKDDVTG

```
121
                                                                        160
                              161
                                                                        200
[Truncated Name:1]1AKE A.pdb
                                EELTTRKDDQEETVRKRLVEYHQMTAPLIGYYSKEAEAGN
[Truncated Name:2]6S36 A.pdb
                                EELTTRKDDQEETVRKRLVEYHQMTAPLIGYYSKEAEAGN
[Truncated Name:3]6RZE A.pdb
                                EELTTRKDDQEETVRKRLVEYHQMTAPLIGYYSKEAEAGN
[Truncated_Name: 4] 3HPR_A.pdb
                                EELTTRKDDQEETVRKRLVEYHQMTAPLIGYYSKEAEAGN
[Truncated_Name:5]1E4V_A.pdb
                                EELTTRKDDQEETVRKRLVEYHQMTAPLIGYYSKEAEAGN
[Truncated_Name: 6] 5EJE_A.pdb
                                EELTTRKDDQEECVRKRLVEYHQMTAPLIGYYSKEAEAGN
[Truncated_Name:7]1E4Y_A.pdb
                                EELTTRKDDQEETVRKRLVEYHQMTAPLIGYYSKEAEAGN
[Truncated_Name:8]3X2S_A.pdb
                                EELTTRKDDQEETVRKRLCEYHQMTAPLIGYYSKEAEAGN
[Truncated_Name: 9] 6HAP_A.pdb
                                EELTTRKDDQEETVRKRLVEYHQMTAPLIGYYSKEAEAGN
[Truncated_Name: 10] 6HAM_A.pdb
                                EELTTRKDDQEETVRKRLVEYHQMTAPLIGYYSKEAEAGN
[Truncated_Name:11]4K46_A.pdb
                                EDLVIREDDKEETVLARLGVYHNQTAPLIAYYGKEAEAGN
[Truncated_Name:12]3GMT_A.pdb
                                EPLVQRDDDKEETVKKRLDVYEAQTKPLITYYGDWARRGA
[Truncated_Name:13]4PZL_A.pdb
                                EPLITRTDDNEDTVKQRLSVYHAQTAKLIDFYRNFSSTNT
                              161
                                                                        200
                              201
                                                           227
[Truncated Name:1]1AKE A.pdb
                                T--KYAKVDGTKPVAEVRADLEKILG-
[Truncated_Name:2]6S36_A.pdb
                                T--KYAKVDGTKPVAEVRADLEKILG-
[Truncated Name:3]6RZE A.pdb
                                T--KYAKVDGTKPVAEVRADLEKILG-
[Truncated_Name: 4] 3HPR_A.pdb
                                T--KYAKVDGTKPVAEVRADLEKILG-
[Truncated_Name:5]1E4V_A.pdb
                                T--KYAKVDGTKPVAEVRADLEKILG-
[Truncated_Name: 6] 5EJE_A.pdb
                                T--KYAKVDGTKPVAEVRADLEKILG-
[Truncated_Name:7]1E4Y_A.pdb
                                T--KYAKVDGTKPVAEVRADLEKILG-
[Truncated_Name:8]3X2S_A.pdb
                                T--KYAKVDGTKPVAEVRADLEKILG-
[Truncated_Name:9]6HAP_A.pdb
                                T--KYAKVDGTKPVCEVRADLEKILG-
[Truncated_Name:10]6HAM_A.pdb
                                T--KYAKVDGTKPVCEVRADLEKILG-
[Truncated_Name:11]4K46_A.pdb
                                T--QYLKFDGTKAVAEVSAELEKALA-
[Truncated_Name:12]3GMT_A.pdb
                                E----YRKISG-
[Truncated_Name:13]4PZL_A.pdb
                                KIPKYIKINGDQAVEKVSQDIFDQLNK
                              201
                                                           227
Call:
 pdbaln(files = files, fit = TRUE, exefile = "msa")
Class:
 pdbs, fasta
```

```
Alignment dimensions:
  13 sequence rows; 227 position columns (204 non-gap, 23 gap)
+ attr: xyz, resno, b, chain, id, ali, resid, sse, call
Use PCA to analyze the aligned and superposed structure
  # Perform PCA
  pc.xray <- pca(pdbs)</pre>
  pc.xray
Call:
  pca.pdbs(pdbs = pdbs)
Class:
  pca
Number of eigenvalues:
  612
        Eigenvalue Variance Cumulative
   PC 1
          2824.299
                     84.993
                                84.993
   PC 2
           384.613
                    11.574
                                96.568
   PC 3
            62.077
                     1.868
                                98.436
   PC 4
            19.614
                      0.590
                                99.026
   PC 5
            14.644
                      0.441
                                99.467
   PC 6
             5.228
                      0.157
                                99.624
   (Obtained from 13 conformers with 612 xyz input values).
+ attr: L, U, z, au, sdev, mean, call
  plot(pc.xray)
```



Color the clusters

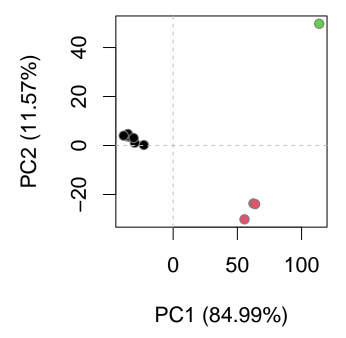
rmsd() will calculate all pairwise root-mean-square deviation values of the structural ensemble distance between different structures . This facilitates clustering analysis based on the pairwise structural deviation:

rmsd(pdbs)

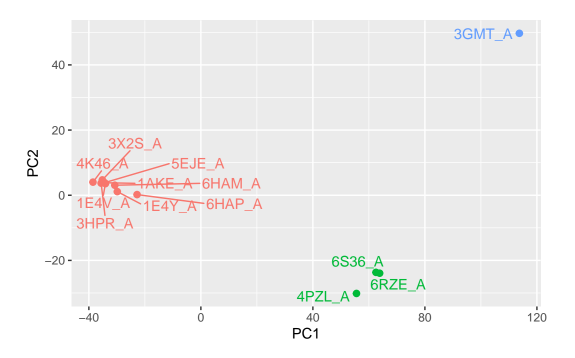
Warning in rmsd(pdbs): No indices provided, using the 204 non NA positions

```
1AKE_A 6S36_A 6RZE_A 3HPR_A 1E4V_A 5EJE_A 1E4Y_A 3X2S_A 6HAP_A 6HAM_A
1AKE_A
        0.000
                7.097
                        7.200
                               0.311
                                       0.251
                                              0.427
                                                      0.941
                                                              0.621
                                                                     1.355
                                                                             0.851
6S36_A
        7.097
                0.000
                        0.434
                               7.184
                                       7.077
                                              7.111
                                                      6.785
                                                              7.195
                                                                     6.346
                                                                             6.859
        7.200
                               7.289
                                              7.212
                                                      6.882
                                                              7.290
6RZE_A
                0.434
                       0.000
                                       7.178
                                                                     6.441
                                                                             6.959
3HPR_A
        0.311
                7.184
                       7.289
                               0.000
                                       0.382
                                              0.507
                                                      1.002
                                                              0.643
                                                                     1.426
                                                                             0.909
1E4V_A
        0.251
                7.077
                       7.178
                               0.382
                                              0.475
                                                              0.636
                                       0.000
                                                      0.971
                                                                     1.377
                                                                             0.851
5EJE_A
        0.427
                7.111
                       7.212
                               0.507
                                       0.475
                                              0.000
                                                      1.093
                                                              0.702
                                                                     1.526
                                                                             0.989
1E4Y_A
        0.941
                6.785
                        6.882
                               1.002
                                       0.971
                                              1.093
                                                      0.000
                                                              0.961
                                                                     1.067
                                                                             0.748
3X2S_A
        0.621
                7.195
                       7.290
                               0.643
                                       0.636
                                              0.702
                                                      0.961
                                                              0.000
                                                                     1.448
                                                                             0.899
6HAP_A
        1.355
                6.346
                        6.441
                               1.426
                                       1.377
                                              1.526
                                                      1.067
                                                              1.448
                                                                     0.000
                                                                             1.162
6HAM_A
        0.851
                6.859
                       6.959
                               0.909
                                       0.851
                                              0.989
                                                      0.748
                                                              0.899
                                                                     1.162
                                                                             0.000
```

```
4K46_A 1.013 7.420 7.518 0.951 1.039 1.059 1.182 0.870 1.787 1.019
3GMT_A 10.863 6.336 6.308 10.950 10.852 10.882 10.632 10.902 10.207 10.646
4PZL_A 6.917 2.381 2.457 6.987 6.896 6.953 6.562 6.967 6.110 6.684
      4K46_A 3GMT_A 4PZL_A
1AKE A 1.013 10.863 6.917
6S36_A 7.420 6.336 2.381
6RZE A 7.518 6.308 2.457
3HPR_A 0.951 10.950 6.987
1E4V A 1.039 10.852 6.896
5EJE_A 1.059 10.882 6.953
1E4Y_A 1.182 10.632 6.562
3X2S_A 0.870 10.902 6.967
6HAP_A 1.787 10.207 6.110
6HAM_A 1.019 10.646 6.684
4K46_A 0.000 11.156 7.199
3GMT_A 11.156 0.000 7.047
4PZL_A 7.199 7.047 0.000
  # Calculate RMSD
  rd <- rmsd(pdbs)
Warning in rmsd(pdbs): No indices provided, using the 204 non NA positions
  # Structure-based clustering
  hc.rd <- hclust(dist(rd))</pre>
  grps.rd <- cutree(hc.rd, k=3)</pre>
  grps.rd
1AKE_A 6S36_A 6RZE_A 3HPR_A 1E4V_A 5EJE_A 1E4Y_A 3X2S_A 6HAP_A 6HAM_A 4K46_A
    1
                 2
                    1 1
                                 1 1 1 1
3GMT_A 4PZL_A
    3
  #colored by groups from `cutree`
  plot(pc.xray, 1:2, col="grey50", bg=grps.rd, pch=21, cex=1.1)
```



plot using ggplot



Optional further visualization

we can make a wee movie ("trajectory") of the major difference of ADK.

```
# Visualize first principal component
pc1 <- mktrj(pc.xray, pc=1, file="pc_1.pdb")</pre>
```