class10 : Structural Bioinformatics pt 1

The PDB database

First let's see what is in the PDB database, the main repository of protein structures.

Downloaded composition stats from: https://www.rcsb.org/stats/summary

For context: Release 2023_2024 of 13-Sep-2023 of UniProtKB/TrEMBL contains 251600768 sequence entries.

https://tinyurl.com/statspdb

```
stats <- read.csv("PDBstats.csv", row.names = 1)
stats</pre>
```

	X.ray	EM	NMR	Multiple.methods	Neutron	Other
Protein (only)	158,844	11,759	12,296	197	73	32
Protein/Oligosaccharide	9,260	2,054	34	8	1	0
Protein/NA	8,307	3,667	284	7	0	0
Nucleic acid (only)	2,730	113	1,467	13	3	1
Other	164	9	32	0	0	0
Oligosaccharide (only)	11	0	6	1	0	4
	Total					
Protein (only)	183,201					
Protein/Oligosaccharide	11,357					
Protein/NA	12,265					
Nucleic acid (only)	4,327					
Other	205					
Oligosaccharide (only)	22					

There is a problem here due to the commas in the numbers. This causes R to treat them as characters.

```
x <- stats$X.ray
  X
[1] "158,844" "9,260"
                          "8,307"
                                    "2,730"
                                               "164"
                                                          "11"
  as.numeric(gsub(",","",x))
[1] 158844
              9260
                             2730
                                     164
                     8307
                                              11
  rm.comma <- function(x) {as.numeric(gsub(",","",x))}</pre>
  rm.comma(stats$EM)
[1] 11759 2054 3667
                          113
                                  9
                                         0
I can use apply() to fix the whole table...
  pdbstats <- apply(stats, 2, rm.comma)</pre>
  rownames(pdbstats) <- rownames(stats)</pre>
  head(pdbstats)
                                          NMR Multiple.methods Neutron Other
                           X.ray
                                    EM
Protein (only)
                                                            197
                                                                      73
                          158844 11759 12296
Protein/Oligosaccharide
                            9260 2054
                                           34
                                                              8
                                                                       1
                                                                             0
Protein/NA
                            8307
                                  3667
                                          284
                                                              7
                                                                       0
                                                                             0
Nucleic acid (only)
                                                             13
                                                                       3
                            2730
                                   113 1467
                                                                             1
Other
                             164
                                     9
                                           32
                                                              0
                                                                       0
                                                                             0
Oligosaccharide (only)
                              11
                                     0
                                            6
                                                              1
                                                                             4
                           Total
Protein (only)
                          183201
Protein/Oligosaccharide
                           11357
Protein/NA
                           12265
Nucleic acid (only)
                            4327
                             205
Other
Oligosaccharide (only)
                              22
```

totals <- apply(pdbstats,2,sum) totals/totals["Total"]</pre>

X.ray	EM	NMR	${\tt Multiple.methods}$
0.8483231383	0.0832730146	0.0667953467	0.0010691797
Neutron	Other	Total	
0.0003642780	0.0001750427	1.0000000000	

Q1: What percentage of structures in the PDB are solved by X-Ray and Electron Microscopy?

```
round(totals/totals["Total"]*100,2)
```

X.ray	EM	NMR	${\tt Multiple.methods}$
84.83	8.33	6.68	0.11
Neutron	Other	Total	
0.04	0.02	100.00	

Q2: What proportion of structures in the PDB are protein?

```
round(pdbstats[1,"Total"] / sum(pdbstats[, "Total"])*100, 2)
```

[1] 86.67

Q3: Type HIV in the PDB website search box on the home page and determine how many HIV-1 protease structures are in the current PDB?

Skip for time

Protein structures in PDB as a fraction of UniProt sequences. See: https://www.uniprot.org/help/release-statistics

```
round((pdbstats[1,"Total"]/251600768)*100,2)
```

[1] 0.07

Here is a lovely figure of HIP-Pr with the catalytic ASP residues, the MK1 compund and the all important water 308



Q4: Water molecules normally have 3 atoms. Why do we see just one atom per water molecule in this structure?

The resolution can't see the small hydrogen atoms because the size is so small.

Q5: There is a critical "conserved" water molecule in the binding site. Can you identify this water molecule? What residue number does this water molecule have?

HOH 308

Q6: Generate and save a figure clearly showing the two distinct chains of HIV-protease along with the ligand. You might also consider showing the catalytic residues ASP 25 in each chain and the critical water (we recommend "Ball & Stick" for these side-chains). Add this figure to your Quarto document.



The bio3d package for structural bioinformatics

library(bio3d)

```
pdb <- read.pdb("1hsg")</pre>
  Note: Accessing on-line PDB file
  pdb
 Call:
        read.pdb(file = "1hsg")
   Total Models#: 1
     Total Atoms#: 1686, XYZs#: 5058 Chains#: 2 (values: A B)
     Protein Atoms#: 1514 (residues/Calpha atoms#: 198)
     Nucleic acid Atoms#: 0 (residues/phosphate atoms#: 0)
     Non-protein/nucleic Atoms#: 172 (residues: 128)
     Non-protein/nucleic resid values: [ HOH (127), MK1 (1) ]
   Protein sequence:
      PQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGIGGFIKVRQYD
      QILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNFPQITLWQRPLVTIKIGGQLKE
      ALLDTGADDTVLEEMSLPGRWKPKMIGGIGGFIKVRQYDQILIEICGHKAIGTVLVGPTP
      VNIIGRNLLTQIGCTLNF
+ attr: atom, xyz, seqres, helix, sheet,
        calpha, remark, call
     Q7: How many amino acid residues are there in this pdb object?
198
     Q8: Name one of the two non-protein residues?
HOH
     Q9: How many protein chains are in this structure?
2 chains
  attributes(pdb)
```

```
$names
[1] "atom"
             "xyz"
                       "seqres" "helix" "sheet" "calpha" "remark" "call"
$class
[1] "pdb" "sse"
  head(pdb$atom)
 type eleno elety alt resid chain resno insert
                                                         Х
                                                                 у
                                                                       z o
                                              <NA> 29.361 39.686 5.862 1 38.10
1 ATOM
           1
                 N < NA >
                           PRO
                                    Α
                                          1
2 ATOM
           2
                CA <NA>
                           PRO
                                    Α
                                          1
                                              <NA> 30.307 38.663 5.319 1 40.62
3 ATOM
           3
                 C <NA>
                           PRO
                                              <NA> 29.760 38.071 4.022 1 42.64
                                    Α
                                          1
4 ATOM
           4
                           PRO
                                          1 <NA> 28.600 38.302 3.676 1 43.40
                 O <NA>
                                    Α
           5
                                          1 <NA> 30.508 37.541 6.342 1 37.87
5 ATOM
                CB <NA>
                           PRO
                                    Α
6 ATOM
           6
                CG <NA>
                           PRO
                                    Α
                                              <NA> 29.296 37.591 7.162 1 38.40
  segid elesy charge
1
  <NA>
            N
                <NA>
2
  <NA>
            С
                <NA>
3
  <NA>
            С
                <NA>
  <NA>
            0
                <NA>
            С
  <NA>
                <NA>
            С
                 <NA>
   <NA>
#Predicting functional motions of a single structure Let's finish today with a bioinformatics
calculation to perdict the functional motions of a PB structure
  adk <- read.pdb("6s36")
 Note: Accessing on-line PDB file
  PDB has ALT records, taking A only, rm.alt=TRUE
  adk
        read.pdb(file = "6s36")
   Total Models#: 1
     Total Atoms#: 1898, XYZs#: 5694 Chains#: 1
```

Protein Atoms#: 1654 (residues/Calpha atoms#: 214) Nucleic acid Atoms#: 0 (residues/phosphate atoms#: 0)

Non-protein/nucleic Atoms#: 244 (residues: 244)

Non-protein/nucleic resid values: [CL (3), HOH (238), MG (2), NA (1)]

Protein sequence:

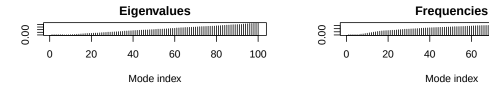
MRIILLGAPGAGKGTQAQFIMEKYGIPQISTGDMLRAAVKSGSELGKQAKDIMDAGKLVT DELVIALVKERIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFDVPDELIVDKI VGRRVHAPSGRVYHVKFNPPKVEGKDDVTGEELTTRKDDQEETVRKRLVEYHQMTAPLIG YYSKEAEAGNTKYAKVDGTKPVAEVRADLEKILG

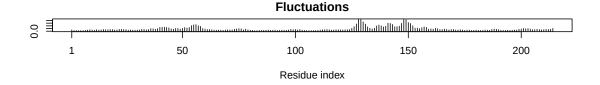
+ attr: atom, xyz, segres, helix, sheet, calpha, remark, call

m <- nma(adk)

Building Hessian... Done in 0.08 seconds. Diagonalizing Hessian... Done in 0.616 seconds.

plot(m)





60

80

100

```
mktrj(m,file="adk_m7.pdb")
mktrj(m, file="adk_m7.pdb")
```

#4. Comparative structure analysis of Adenylate Kinase

We need some packages for today's class. These include 'bio3d' and 'msa'.

We 'msa' package is from BioConductor. These packages focus on genomics type work and are managed by the 'BiocManager' package.

Install 'install.packages("BiocManager")' and then 'BiocManager::install("msa")' all entered in the R "brain" console.

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

True

bio3d-view

121

121

pdb|1AKE|A

```
library(bio3d)
  aa <- get.seq("1ake_A")</pre>
Warning in get.seq("lake_A"): Removing existing file: seqs.fasta
Fetching... Please wait. Done.
  aa
              MRIILLGAPGAGKGTQAQFIMEKYGIPQISTGDMLRAAVKSGSELGKQAKDIMDAGKLVT
pdb | 1AKE | A
             61
pdb | 1AKE | A
              DELVIALVKERIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFDVPDELIVDRI
```

60

60

120

120

180

180

VGRRVHAPSGRVYHVKFNPPKVEGKDDVTGEELTTRKDDQEETVRKRLVEYHQMTAPLIG

```
181
                                                 214
             YYSKEAEAGNTKYAKVDGTKPVAEVRADLEKILG
pdb|1AKE|A
           181
                                                 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
  # Blast or hmmer search
  #b <- blast.pdb(aa)</pre>
  # Plot a summary of search results
  #hits <- plot(b)</pre>
  # List out some 'top hits'
  #head(hits$pdb.id)
  hits <- NULL
  hits$pdb.id <- c('1AKE_A','6S36_A','6RZE_A','3HPR_A','1E4V_A','5EJE_A','1E4Y_A','3X2S_A','
Now we can download all these structures for further analysis with the 'get.pdb()' function.
  # 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.gz exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/6S36.pdb.gz exists. Skipping download
```

```
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/6RZE.pdb.gz exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/3HPR.pdb.gz exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/1E4V.pdb.gz exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/5EJE.pdb.gz exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/1E4Y.pdb.gz exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/3X2S.pdb.gz exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/6HAP.pdb.gz exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/6HAM.pdb.gz exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/4K46.pdb.gz exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/3GMT.pdb.gz exists. Skipping download
Warning in get.pdb(hits$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE):
pdbs/4PZL.pdb.gz exists. Skipping download
                                                                             0%
                                                                             8%
```

No we have all these related structures we can Align and Supperpose...

```
# 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/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
- .. 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 pdb/seq: 2 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 pdb/seq: 6 name: pdbs/split_chain/5EJE_A.pdb 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 name: pdbs/split_chain/6HAM_A.pdb pdb/seq: 10 PDB has ALT records, taking A only, rm.alt=TRUE pdb/seq: 11 name: pdbs/split_chain/4K46_A.pdb

PDB has ALT records, taking A only, rm.alt=TRUE

name: pdbs/split_chain/3GMT_A.pdb

name: pdbs/split_chain/4PZL_A.pdb

pdbs

pdb/seq: 12

pdb/seq: 13

[Truncated_Name: 6] 5EJE_A.pdb -----MRIILLGAPGAGKGTQAQFIMEKYGIPQIS [Truncated_Name:7]1E4Y_A.pdb -----MRIILLGALVAGKGTQAQFIMEKYGIPQIS [Truncated_Name:8]3X2S_A.pdb -----MRIILLGAPGAGKGTQAQFIMEKYGIPQIS [Truncated_Name:9]6HAP_A.pdb ----MRIILLGAPGAGKGTQAQFIMEKYGIPQIS [Truncated Name:10]6HAM A.pdb ----MRIILLGAPGAGKGTQAQFIMEKYGIPQIS [Truncated Name:11]4K46 A.pdb -----MRIILLGAPGAGKGTQAQFIMAKFGIPQIS [Truncated Name:12]3GMT A.pdb ----MRLILLGAPGAGKGTQANFIKEKFGIPQIS [Truncated_Name:13]4PZL_A.pdb TENLYFQSNAMRIILLGAPGAGKGTQAKIIEQKYNIAHIS **^**** ****** * 1 40 41 80 [Truncated_Name:1]1AKE_A.pdb TGDMLRAAVKSGSELGKQAKDIMDAGKLVTDELVIALVKE [Truncated_Name:2]6S36_A.pdb TGDMLRAAVKSGSELGKQAKDIMDAGKLVTDELVIALVKE [Truncated_Name:3]6RZE_A.pdb TGDMLRAAVKSGSELGKQAKDIMDAGKLVTDELVIALVKE [Truncated_Name: 4] 3HPR_A.pdb TGDMLRAAVKSGSELGKQAKDIMDAGKLVTDELVIALVKE [Truncated_Name:5]1E4V_A.pdb TGDMLRAAVKSGSELGKQAKDIMDAGKLVTDELVIALVKE [Truncated_Name: 6] 5EJE_A.pdb TGDMLRAAVKSGSELGKQAKDIMDACKLVTDELVIALVKE [Truncated_Name:7]1E4Y_A.pdb TGDMLRAAVKSGSELGKQAKDIMDAGKLVTDELVIALVKE [Truncated Name:8]3X2S A.pdb TGDMLRAAVKSGSELGKQAKDIMDCGKLVTDELVIALVKE [Truncated Name:9]6HAP A.pdb TGDMLRAAVKSGSELGKQAKDIMDAGKLVTDELVIALVRE [Truncated Name:10]6HAM A.pdb TGDMLRAAIKSGSELGKQAKDIMDAGKLVTDEIIIALVKE [Truncated_Name:11]4K46_A.pdb TGDMLRAAIKAGTELGKQAKSVIDAGQLVSDDIILGLVKE [Truncated Name:12]3GMT A.pdb TGDMLRAAVKAGTPLGVEAKTYMDEGKLVPDSLIIGLVKE [Truncated_Name:13]4PZL_A.pdb TGDMIRETIKSGSALGQELKKVLDAGELVSDEFIIKIVKD 41 80 81 [Truncated_Name:1]1AKE_A.pdb RIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFD [Truncated_Name:2]6S36_A.pdb RIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFD [Truncated_Name:3]6RZE_A.pdb RIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFD [Truncated_Name: 4] 3HPR_A.pdb RIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFD ${\tt RIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFD}$ [Truncated_Name:5]1E4V_A.pdb [Truncated Name:6]5EJE A.pdb RIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFD [Truncated Name:7]1E4Y A.pdb RIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFD [Truncated Name:8]3X2S A.pdb RIAQEDSRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFD [Truncated_Name: 9] 6HAP_A.pdb RICQEDSRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFD [Truncated_Name:10]6HAM_A.pdb RICQEDSRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFD [Truncated_Name:11]4K46_A.pdb RIAQDDCAKGFLLDGFPRTIPQADGLKEVGVVVDYVIEFD

14

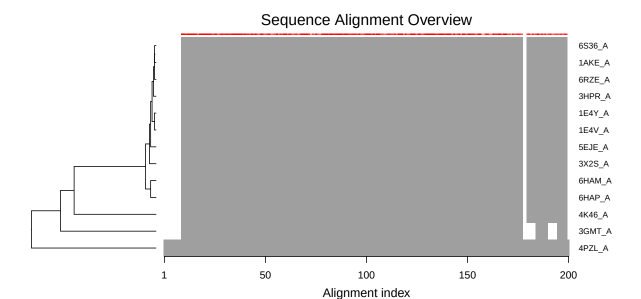
RLKEADCANGYLFDGFPRTIAQADAMKEAGVAIDYVLEID

[Truncated_Name:12]3GMT_A.pdb

[Truncated_Name: 13] 4PZL_A.pdb

	81
	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 VPDELIVDAIVGRRVHAPSGRVYHVKFNPPKVEGKDDVTG VPDELIVDAIVGRRVHAPSGRVYHVKFNPPKVEGKDDVTG VPDELIVDRIVGRRVHAPSGRVYHVKFNPPKVEGKDDVTG VPDELIVDRIVGRRVHAPSGRVYHVKFNPPKVEGKDDVTG VPDELIVDRIVGRRVHAPSGRVYHVKFNPPKVEGKDDVTG VPDELIVDRIVGRRVHAPSGRVYHVKFNPPKVEGKDDVTG VPDELIVDRIVGRRVHAPSGRVYHVKFNPPKVEGKDDVTG VPDELIVDRIVGRRVHAPSGRVYHVKFNPPKVEGKDDVTG VPDELIVDRIVGRRVHAPSGRVYHVKFNPPKVEGKDDVTG VPDELIVDRIVGRRVHAPSGRVYHVKFNPPKVEGKDDVTG VADSVIVERMAGRRAHLASGRTYHNVYNPPKVEGKDDVTG VPFSEIIERMSGRRTHPASGRTYHVKFNPPKVEGKDDVTG VADNLLIERITGRRIHPASGRTYHTKFNPPKVADKDDVTG
	121
[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	EELTTRKDDQEETVRKRLVEYHQMTAPLIGYYSKEAEAGN EELTTRKDDQEETVRKRLVEYHQMTAPLIGYYSKEAEAGN EELTTRKDDQEETVRKRLVEYHQMTAPLIGYYSKEAEAGN EELTTRKDDQEETVRKRLVEYHQMTAPLIGYYSKEAEAGN EELTTRKDDQEETVRKRLVEYHQMTAPLIGYYSKEAEAGN EELTTRKDDQEETVRKRLVEYHQMTAPLIGYYSKEAEAGN EELTTRKDDQEETVRKRLVEYHQMTAPLIGYYSKEAEAGN EELTTRKDDQEETVRKRLVEYHQMTAPLIGYYSKEAEAGN EELTTRKDDQEETVRKRLCEYHQMTAPLIGYYSKEAEAGN EELTTRKDDQEETVRKRLVEYHQMTAPLIGYYSKEAEAGN EELTTRKDDQEETVRKRLVEYHQMTAPLIGYYSKEAEAGN EELTTRKDDQEETVRKRLVEYHQMTAPLIGYYSKEAEAGN EDLVIREDDKEETVLARLGVYHNQTAPLIAYYGKEAEAGN EPLVQRDDDKEETVKKRLDVYEAQTKPLITYYGDWARRGA EPLITRTDDNEDTVKQRLSVYHAQTAKLIDFYRNFSSTNT * * * * * * * * * * * * * * * * * * *
[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	201

```
[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
  # Vector containing PDB codes for figure axis
  ids <- basename.pdb(pdbs$id)</pre>
  # Draw schematic alignment
  plot(pdbs, labels=ids)
```



Now I can search the PDB database for related sequences:

Side-note: Let's annotate these structures (in other words find out what they are, what species they are from, stuff about the experiment they were solved in etc.)

For this we can use the 'pdb.annotate()'

```
#anno <-pdb.annotate(hits$pdb.id)
anno <- pdb.annotate(ids)
unique(anno$source)</pre>
```

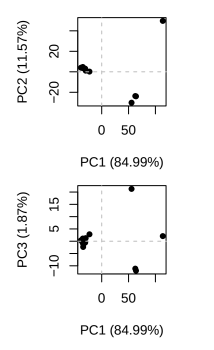
- [1] "Escherichia coli"
- [2] "Escherichia coli K-12"
- [3] "Escherichia coli 0139:H28 str. E24377A"
- [4] "Escherichia coli str. K-12 substr. MDS42"
- [5] "Photobacterium profundum"
- [6] "Burkholderia pseudomallei 1710b"
- [7] "Francisella tularensis subsp. tularensis SCHU S4"

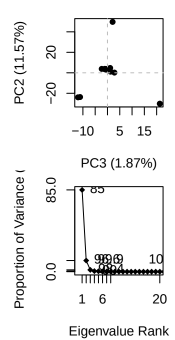
```
#attributes(anno)
head(anno)
```

structureId chainId macromoleculeType chainLength experimentalTechnique

```
1AKE_A
              1AKE
                                                       214
                                                                           X-ray
                         Α
                                      Protein
6S36_A
              6S36
                         Α
                                      Protein
                                                       214
                                                                           X-ray
6RZE_A
              6RZE
                                                       214
                         Α
                                      Protein
                                                                           X-ray
3HPR_A
              3HPR
                                                       214
                         Α
                                      Protein
                                                                           X-ray
1E4V A
              1E4V
                         Α
                                      Protein
                                                       214
                                                                           X-ray
              5EJE
5EJE A
                         Α
                                      Protein
                                                       214
                                                                           X-ray
       resolution
                        scopDomain
                                                                            pfam
1AKE_A
             2.00 Adenylate kinase Adenylate kinase, active site lid (ADK_lid)
                               <NA> Adenylate kinase, active site lid (ADK_lid)
6S36_A
             1.60
                               <NA> Adenylate kinase, active site lid (ADK_lid)
6RZE_A
             1.69
                               <NA> Adenylate kinase, active site lid (ADK_lid)
3HPR_A
             2.00
             1.85 Adenylate kinase Adenylate kinase, active site lid (ADK lid)
1E4V_A
                               <NA> Adenylate kinase, active site lid (ADK_lid)
5EJE_A
             1.90
               ligandId
                                                                ligandName
                                         BIS (ADENOSINE) -5'-PENTAPHOSPHATE
1AKE_A
                    AP5
6S36_A CL (3),NA,MG (2)
                           CHLORIDE ION (3), SODIUM ION, MAGNESIUM ION (2)
6RZE_A
          NA (3),CL (2)
                                          SODIUM ION (3), CHLORIDE ION (2)
3HPR_A
                    AP5
                                         BIS (ADENOSINE) -5'-PENTAPHOSPHATE
1E4V_A
                    AP5
                                         BIS (ADENOSINE) -5'-PENTAPHOSPHATE
5EJE A
                 AP5, CO BIS (ADENOSINE) -5'-PENTAPHOSPHATE, COBALT (II) ION
                                        source
1AKE A
                              Escherichia coli
6S36_A
                             Escherichia coli
6RZE_A
                             Escherichia coli
3HPR_A
                        Escherichia coli K-12
1E4V_A
                              Escherichia coli
5EJE_A Escherichia coli 0139:H28 str. E24377A
1AKE A STRUCTURE OF THE COMPLEX BETWEEN ADENYLATE KINASE FROM ESCHERICHIA COLI AND THE INHIB
6S36_A
6RZE_A
3HPR_A
1E4V_A
5EJE_A
                                                                                           Crys
                                                      citation rObserved rFree
1AKE A
                      Muller, C.W., et al. J Mol Biol (1992)
                                                                  0.1960
                       Rogne, P., et al. Biochemistry (2019)
6S36 A
                                                                  0.1632 0.2356
6RZE A
                       Rogne, P., et al. Biochemistry (2019)
                                                                  0.1865 0.2350
3HPR_A Schrank, T.P., et al. Proc Natl Acad Sci U S A (2009)
                                                                  0.2100 0.2432
                        Muller, C.W., et al. Proteins (1993)
1E4V A
                                                                  0.1960
5EJE_A Kovermann, M., et al. Proc Natl Acad Sci U S A (2017)
                                                                  0.1889 0.2358
        rWork spaceGroup
1AKE_A 0.1960 P 21 2 21
```

```
# Perform PCA
pc.xray <- pca(pdbs)
plot(pc.xray)</pre>
```



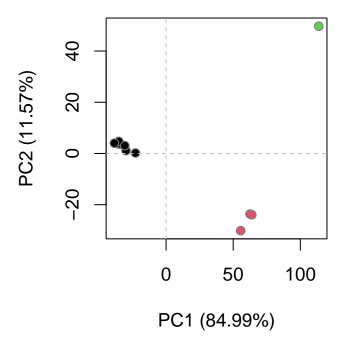


```
# Calculate RMSD
rd <- rmsd(pdbs)</pre>
```

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

```
# Structure-based clustering
hc.rd <- hclust(dist(rd))
grps.rd <- cutree(hc.rd, k=3)

plot(pc.xray, 1:2, col="grey50", bg=grps.rd, pch=21, cex=1)</pre>
```



Class 11

- $[1] \ "hivpr_dimer_23119_unrelaxed_rank_001_alphafold2_multimer_v3_model_1_seed_000.pdb" \ and \ an approximate the content of the content$
- [2] "hivpr_dimer_23119_unrelaxed_rank_002_alphafold2_multimer_v3_model_5_seed_000.pdb"
- [3] "hivpr_dimer_23119_unrelaxed_rank_003_alphafold2_multimer_v3_model_4_seed_000.pdb"
- [4] "hivpr_dimer_23119_unrelaxed_rank_004_alphafold2_multimer_v3_model_2_seed_000.pdb"
- [5] "hivpr_dimer_23119_unrelaxed_rank_005_alphafold2_multimer_v3_model_3_seed_000.pdb"

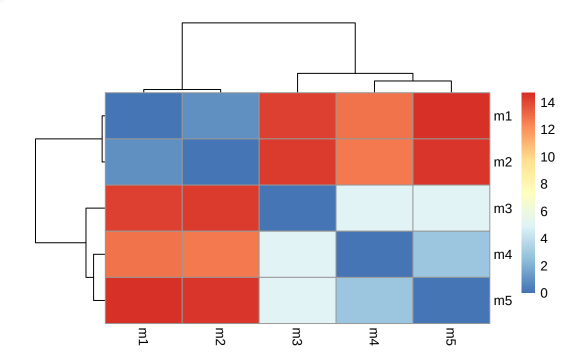
```
library(bio3d)
     # Read all data from Models
     # and superpose/fit coords
     pdbs <- pdbaln(pdb_files, fit=TRUE, exefile="msa")</pre>
Reading PDB files:
hivpr_dimer_23119/hivpr_dimer_23119_unrelaxed_rank_001_alphafold2_multimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer_v3_model_1_seed_001_nultimer
hivpr_dimer_23119/hivpr_dimer_23119_unrelaxed_rank_003_alphafold2_multimer_v3_model_4_seed_0
hivpr_dimer_23119/hivpr_dimer_23119_unrelaxed_rank_005_alphafold2_multimer_v3_model_3_seed_0
Extracting sequences
                            name: hivpr_dimer_23119/hivpr_dimer_23119_unrelaxed_rank_001_alphafold2_multime
pdb/seq: 1
                            name: hivpr_dimer_23119/hivpr_dimer_23119_unrelaxed_rank_002_alphafold2_multimer_
pdb/seq: 2
pdb/seq: 3
                            name: hivpr_dimer_23119/hivpr_dimer_23119_unrelaxed_rank_003_alphafold2_multime:
pdb/seq: 4
                            name: hivpr_dimer_23119/hivpr_dimer_23119_unrelaxed_rank_004_alphafold2_multimer
pdb/seq: 5
                            name: hivpr_dimer_23119/hivpr_dimer_23119_unrelaxed_rank_005_alphafold2_multimer
     pdbs
 [Truncated_Name:1]hivpr_dime
                                                                   PQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGI
[Truncated_Name:2]hivpr_dime
                                                                   PQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGI
[Truncated_Name:3]hivpr_dime
                                                                   PQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGI
[Truncated_Name:4]hivpr_dime
                                                                   PQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGI
 [Truncated_Name:5]hivpr_dime
                                                                   PQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGI
                                                                   **************
                                                                                                                                                                            50
                                                                 51
                                                                                                                                                                            100
```

[Truncated_Name:1]hivpr_dime [Truncated_Name:2]hivpr_dime [Truncated_Name:3]hivpr_dime [Truncated_Name:4]hivpr_dime [Truncated_Name:5]hivpr_dime GGFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNFP GGFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNFP GGFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNFP GGFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNFP GGFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNFP

```
*****************
                            51
                                                                            100
                           101
                                                                            150
[Truncated Name:1]hivpr dime
                             QITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGIG
[Truncated Name:2]hivpr dime
                             QITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGIG
[Truncated Name:3] hivpr dime
                             QITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGIG
[Truncated_Name:4]hivpr_dime
                             QITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGIG
[Truncated_Name:5]hivpr_dime
                             QITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGIG
                             **************
                           101
                                                                            150
                           151
                                                                          198
[Truncated_Name:1]hivpr_dime
                             GFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNF
[Truncated_Name:2]hivpr_dime
                             GFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNF
[Truncated_Name:3]hivpr_dime
                             GFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNF
[Truncated_Name:4]hivpr_dime
                             GFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNF
[Truncated_Name:5]hivpr_dime
                             GFIKVRQYDQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNF
                             **************
                           151
                                                                          198
Call:
 pdbaln(files = pdb_files, fit = TRUE, exefile = "msa")
Class:
 pdbs, fasta
Alignment dimensions:
 5 sequence rows; 198 position columns (198 non-gap, 0 gap)
+ attr: xyz, resno, b, chain, id, ali, resid, sse, call
  rd <- rmsd(pdbs, fit=T)</pre>
Warning in rmsd(pdbs, fit = T): No indices provided, using the 198 non NA positions
  range(rd)
[1] 0.000 14.689
```

```
library(pheatmap)

colnames(rd) <- paste0("m",1:5)
rownames(rd) <- paste0("m",1:5)
pheatmap(rd)</pre>
```



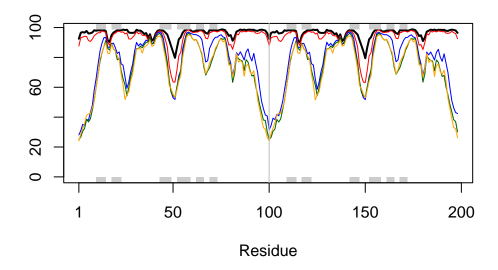
```
# Read a reference PDB structure
pdb <- read.pdb("1hsg")</pre>
```

Note: Accessing on-line PDB file

Warning in get.pdb(file, path = tempdir(), verbose = FALSE):
/var/folders/13/j10nl4n15ls4kl7cbgcgxp_c0000gn/T//RtmpcuXv0R/1hsg.pdb exists.
Skipping download

```
plotb3(pdbs$b[1,], typ="l", lwd=2, sse=pdb)
points(pdbs$b[2,], typ="l", col="red")
points(pdbs$b[3,], typ="l", col="blue")
points(pdbs$b[4,], typ="l", col="darkgreen")
```

```
points(pdbs$b[5,], typ="l", col="orange")
abline(v=100, col="gray")
```



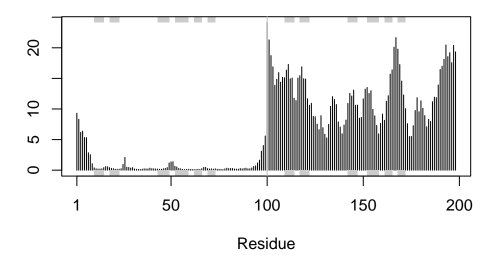
core <- core.find(pdbs)</pre>

```
core size 197 of 198
                      vol = 6154.839
core size 196 of 198
                      vol = 5399.676
core size 195 of 198
                      vol = 5074.795
core size 194 of 198
                      vol = 4802.518
core size 193 of 198
                      vol = 4520.256
core size 192 of 198
                      vol = 4305.362
                      vol = 4089.792
core size 191 of 198
core size 190 of 198
                      vol = 3886.145
core size 189 of 198
                      vol = 3758.321
core size 188 of 198
                      vol = 3620.18
core size 187 of 198
                      vol = 3496.698
core size 186 of 198
                      vol = 3389.985
core size 185 of 198
                      vol = 3320.114
core size 184 of 198
                      vol = 3258.683
core size 183 of 198
                      vol = 3208.591
core size 182 of 198 vol = 3156.736
```

```
core size 181 of 198 vol = 3141.668
core size 180 of 198
                      vol = 3136.574
core size 179 of 198
                      vol = 3155.52
core size 178 of 198
                      vol = 3185.362
core size 177 of 198
                      vol = 3204.487
core size 176 of 198
                      vol = 3211.978
core size 175 of 198
                      vol = 3234.993
core size 174 of 198
                      vol = 3244.062
core size 173 of 198
                      vol = 3237.845
core size 172 of 198
                      vol = 3218.77
core size 171 of 198
                      vol = 3180.743
core size 170 of 198
                      vol = 3130.369
core size 169 of 198
                      vol = 3067.881
core size 168 of 198
                      vol = 2989.546
core size 167 of 198
                      vol = 2928.272
core size 166 of 198
                      vol = 2851.193
core size 165 of 198
                      vol = 2780.877
                      vol = 2708.433
core size 164 of 198
core size 163 of 198
                      vol = 2636.516
core size 162 of 198
                      vol = 2563.25
core size 161 of 198
                      vol = 2478.024
core size 160 of 198
                      vol = 2404.793
core size 159 of 198
                      vol = 2330.997
core size 158 of 198
                      vol = 2250.477
core size 157 of 198
                      vol = 2159.432
core size 156 of 198
                      vol = 2070.759
core size 155 of 198
                      vol = 1983.579
core size 154 of 198
                      vol = 1917.913
core size 153 of 198
                      vol = 1842.556
core size 152 of 198
                      vol = 1775.398
core size 151 of 198
                      vol = 1695.133
core size 150 of 198
                      vol = 1632.173
core size 149 of 198
                      vol = 1570.391
core size 148 of 198
                      vol = 1497.238
core size 147 of 198
                      vol = 1434.802
core size 146 of 198
                      vol = 1367.706
core size 145 of 198
                      vol = 1302.596
core size 144 of 198
                      vol = 1251.985
core size 143 of 198
                      vol = 1207.976
core size 142 of 198
                      vol = 1167.112
core size 141 of 198
                      vol = 1118.27
core size 140 of 198
                      vol = 1081.664
core size 139 of 198 vol = 1029.75
```

```
core size 138 of 198
                     vol = 981.766
core size 137 of 198
                      vol = 944.446
core size 136 of 198
                      vol = 899.224
core size 135 of 198
                      vol = 859.402
core size 134 of 198
                      vol = 814.694
core size 133 of 198
                      vol = 771.862
core size 132 of 198
                      vol = 733.807
core size 131 of 198
                      vol = 702.053
core size 130 of 198
                      vol = 658.757
                      vol = 622.574
core size 129 of 198
core size 128 of 198
                      vol = 578.29
core size 127 of 198
                      vol = 543.07
core size 126 of 198
                      vol = 510.934
core size 125 of 198
                      vol = 481.595
core size 124 of 198
                      vol = 464.672
core size 123 of 198
                      vol = 451.721
core size 122 of 198
                      vol = 430.417
                      vol = 409.141
core size 121 of 198
core size 120 of 198
                      vol = 378.942
core size 119 of 198
                      vol = 348.325
core size 118 of 198
                      vol = 324.738
core size 117 of 198
                      vol = 312.394
core size 116 of 198
                      vol = 300.89
core size 115 of 198
                      vol = 279.976
core size 114 of 198
                      vol = 263.434
core size 113 of 198
                      vol = 250.263
core size 112 of 198
                      vol = 229.592
core size 111 of 198
                      vol = 209.929
core size 110 of 198
                      vol = 196.379
core size 109 of 198
                      vol = 180.628
core size 108 of 198
                      vol = 167.088
core size 107 of 198
                      vol = 155.875
core size 106 of 198
                      vol = 142.595
core size 105 of 198
                      vol = 128.924
core size 104 of 198
                      vol = 114.054
core size 103 of 198
                      vol = 100.936
core size 102 of 198
                      vol = 90.431
core size 101 of 198
                      vol = 81.972
core size 100 of 198
                     vol = 74.017
core size 99 of 198
                     vol = 66.855
core size 98 of 198
                     vol = 59.525
core size 97 of 198
                     vol = 52.263
core size 96 of 198 vol = 43.699
```

```
core size 95 of 198 vol = 35.813
 core size 94 of 198 vol = 28.888
 core size 93 of 198 vol = 20.692
 core size 92 of 198 vol = 14.975
 core size 91 of 198 vol = 9.146
 core size 90 of 198 \text{ vol} = 5.232
 core size 89 of 198 vol = 3.53
 core size 88 of 198 vol = 2.657
 core size 87 of 198 vol = 1.998
 core size 86 of 198 vol = 1.333
 core size 85 of 198 vol = 1.141
 core size 84 of 198 vol = 1.012
 core size 83 of 198 vol = 0.891
 core size 82 of 198 vol = 0.749
 core size 81 of 198 vol = 0.618
 core size 80 of 198 vol = 0.538
 core size 79 of 198 vol = 0.479
FINISHED: Min vol (0.5) reached
  core.inds <- print(core, vol=0.5)</pre>
# 80 positions (cumulative volume <= 0.5 Angstrom^3)
  start end length
         25
     10
                16
2
     27
        48
                22
3
    53 94
                42
  xyz <- pdbfit(pdbs, core.inds, outpath="corefit_structures")</pre>
  rf <- rmsf(xyz)
  plotb3(rf, sse=pdb)
  abline(v=100, col="gray", ylab="RMSF")
```

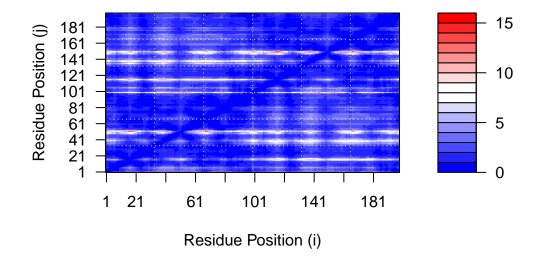


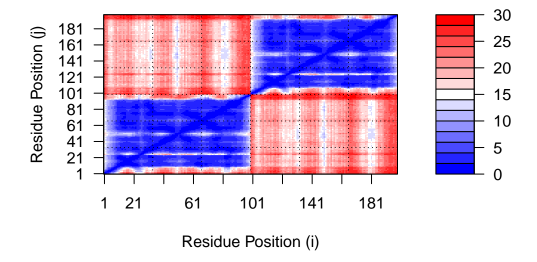
```
library(jsonlite)
  # Listing of all PAE JSON files
  pae_files <- list.files(path=results_dir,</pre>
                            pattern=".*model.*\\.json",
                            full.names = TRUE)
  pae1 <- read_json(pae_files[1],simplifyVector = TRUE)</pre>
  pae5 <- read_json(pae_files[5],simplifyVector = TRUE)</pre>
  attributes(pae1)
$names
[1] "plddt"
              "max_pae" "pae"
                                    "ptm"
                                               "iptm"
  # Per-residue pLDDT scores
  # same as B-factor of PDB..
  head(pae1$plddt)
```

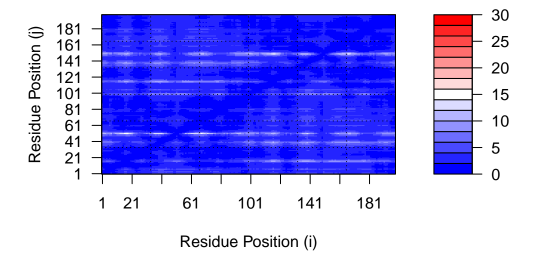
[1] 92.50 96.56 96.94 96.62 97.69 96.00

```
pae1$max_pae
```

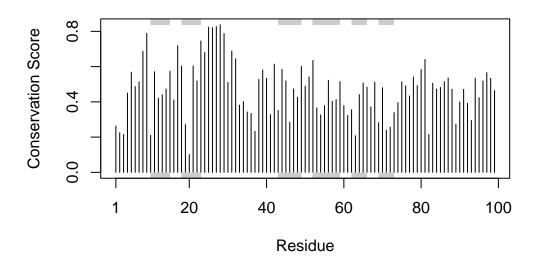
[1] 15.54688







[1] 5378 132



[127] "-" "-" "-" "-" "-"

```
m1.pdb <- read.pdb(pdb_files[1])
occ <- vec2resno(c(sim[1:99], sim[1:99]), m1.pdb$atom$resno)
write.pdb(m1.pdb, o=occ, file="m1_conserv.pdb")</pre>
```

sessionInfo()

R version 4.3.1 (2023-06-16)
Platform: x86_64-apple-darwin20 (64-bit)
Running under: macOS Big Sur 11.7.10

Matrix products: default
BLAS: /Library/Frameworks/R.framework/Versions/4.3-x86_64/Resources/lib/libRblas.0.dylib
LAPACK: /Library/Frameworks/R.framework/Versions/4.3-x86_64/Resources/lib/libRlapack.dylib;

locale:

[1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8

time zone: America/Los_Angeles

tzcode source: internal

attached base packages:

[1] stats graphics grDevices utils datasets methods base

other attached packages:

[1] jsonlite_1.8.7 pheatmap_1.0.12 bio3d_2.4-4

loaded via a namespace (and not attached):

[1]	crayon_1.5.2	httr_1.4.7	cli_3.6.1
[4]	knitr_1.45	rlang_1.1.2	xfun_0.41
[7]	glue_1.6.2	S4Vectors_0.40.1	colorspace_2.1-0
[10]	RCurl_1.98-1.13	Biostrings_2.70.1	htmltools_0.5.7
[13]	stats4_4.3.1	scales_1.2.1	rmarkdown_2.25
[16]	grid_4.3.1	munsell_0.5.0	evaluate_0.23
[19]	bitops_1.0-7	fastmap_1.1.1	lifecycle_1.0.4
[22]	yam1_2.3.7	IRanges_2.36.0	<pre>GenomeInfoDb_1.38.1</pre>
[25]	compiler_4.3.1	RColorBrewer_1.1-3	Rcpp_1.0.11
[28]	XVector_0.42.0	rstudioapi_0.15.0	digest_0.6.33
[31]	R6_2.5.1	<pre>GenomeInfoDbData_1.2.11</pre>	curl_5.1.0
[34]	parallel_4.3.1	gtable_0.3.4	tools_4.3.1
[37]	zlibbioc_1.48.0	$msa_1.34.0$	BiocGenerics_0.48.1

