

# Class 9: Structural Bioinformatics

Wade Ingersoll (PID: A69038080)

## Table of contents

The PDB database . . . . .	1
Visualizing structure data . . . . .	7
Bio3D package for structural bioinformatics . . . . .	9
Prediction of functional motions . . . . .	13
Play with 3D viewing in R . . . . .	15
4. Comparative structure analysis of Adenylate Kinase . . . . .	16
Overview . . . . .	16
Setup . . . . .	16
Search and retrieve ADK structures . . . . .	17
Align and superpose structures . . . . .	20
Optional: Viewing our superposed structures . . . . .	23
Principal component analysis . . . . .	28
5. Optional further visualization . . . . .	30

## The PDB database

The main repository for biomolecular structure data is the Protein Data Bank (PDB)  
<https://www.rcsb.org/>

Let's have a quick look at the composition of this database:

```
# Read in data file
stats <- read.csv("Data Export Summary.csv")

# View it
stats
```

	Molecular.Type	X.ray	EM	NMR	Integrative	Multiple.methods
1	Protein (only)	176,378	20,438	12,709	342	221
2	Protein/Oligosaccharide	10,284	3,396	34	8	11
3	Protein/NA	9,007	5,931	287	24	7
4	Nucleic acid (only)	3,077	200	1,554	2	15
5	Other	174	13	33	3	0
6	Oligosaccharide (only)	11	0	6	0	1
	Neutron	Other	Total			
1	83	32	210,203			
2	1	0	13,734			
3	0	0	15,256			
4	3	1	4,852			
5	0	0	223			
6	0	4	22			

We need to remove commas:

```
as.numeric(sub(","," ", stats$X.ray))
```

```
[1] 176378 10284 9007 3077 174 11
```

However, this is not efficient so we will use a different function from the **readr** package.

```
library(readr)

stats <- read_csv("Data Export Summary.csv")
```

```
Rows: 6 Columns: 9
-- Column specification -----
Delimiter: ","
chr (1): Molecular Type
dbl (4): Integrative, Multiple methods, Neutron, Other
num (4): X-ray, EM, NMR, Total

i Use `spec()` to retrieve the full column specification for this data.
i Specify the column types or set `show_col_types = FALSE` to quiet this message.
```

```
stats
```

```

# A tibble: 6 x 9
`Molecular Type` `X-ray`    EM    NMR Integrative `Multiple methods` Neutron
<chr>            <dbl>    <dbl> <dbl>        <dbl>            <dbl>    <dbl>
1 Protein (only) 176378 20438 12709        342            221     83
2 Protein/Oligosacch~ 10284  3396   34          8             11      1
3 Protein/NA       9007   5931   287         24             7      0
4 Nucleic acid (only) 3077   200    1554        2             15      3
5 Other             174    13     33          3             0      0
6 Oligosaccharide (o~ 11     0      6           0             1      0
# i 2 more variables: Other <dbl>, Total <dbl>

```

Now we can answer Q1 (Percent X-Ray and EM)

```
n.total <- sum(stats$Total)
n.xray <- sum(stats$`X-ray`)
n.em <- sum(stats$EM)

round( n.xray/n.total * 100, 2 )
```

```
[1] 81.43
```

```
round( n.em/n.total * 100, 2 )
```

```
[1] 12.27
```

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

**Answer:** X-Ray = 81.43% and EM = 12.27% (see code above for calculations)

Now let's answer Q2

```
n.total <- sum(stats$Total)
protein <- stats$Total[1]

round( protein/n.total * 100, 2 )
```

```
[1] 86.05
```

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

**Answer:** 86.05% (see code above for calculations)

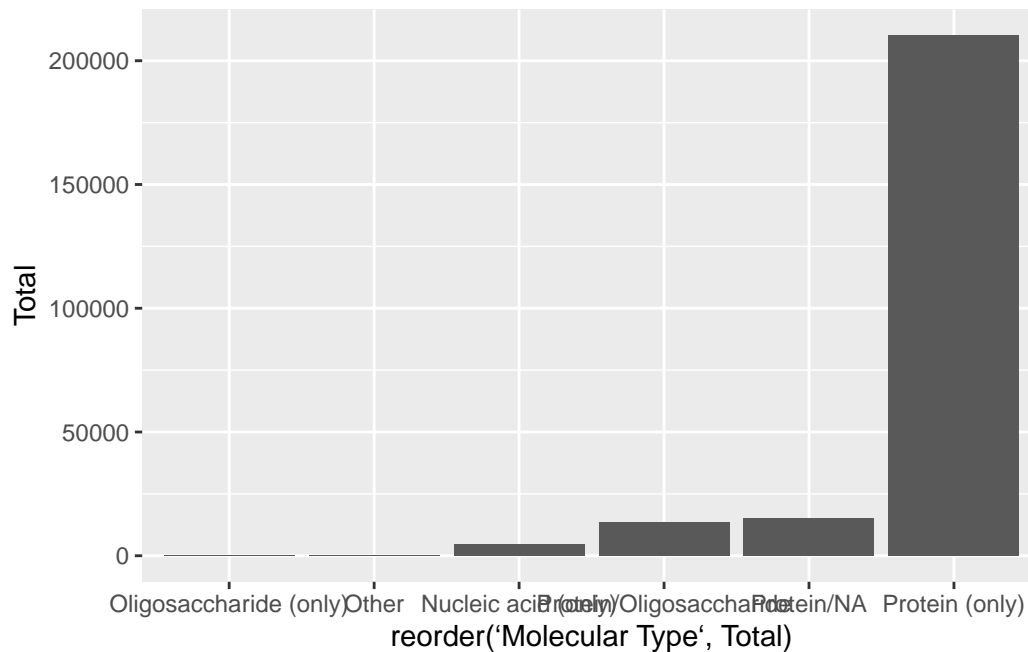
Now let's answer Q3

**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? (Make a barplot overview)

**Answer:** 1,150

```
library(ggplot2)

ggplot(stats) +
  aes(reorder(`Molecular Type`, Total), Total) +
  geom_col()
```



## Visualizing structure data

The Mol\* viewer embedded in many bioinformatics websites. The homepage is <https://molstar.org/> (use pdb accession number 1hsg)

I can insert any figure or image file using markdown format

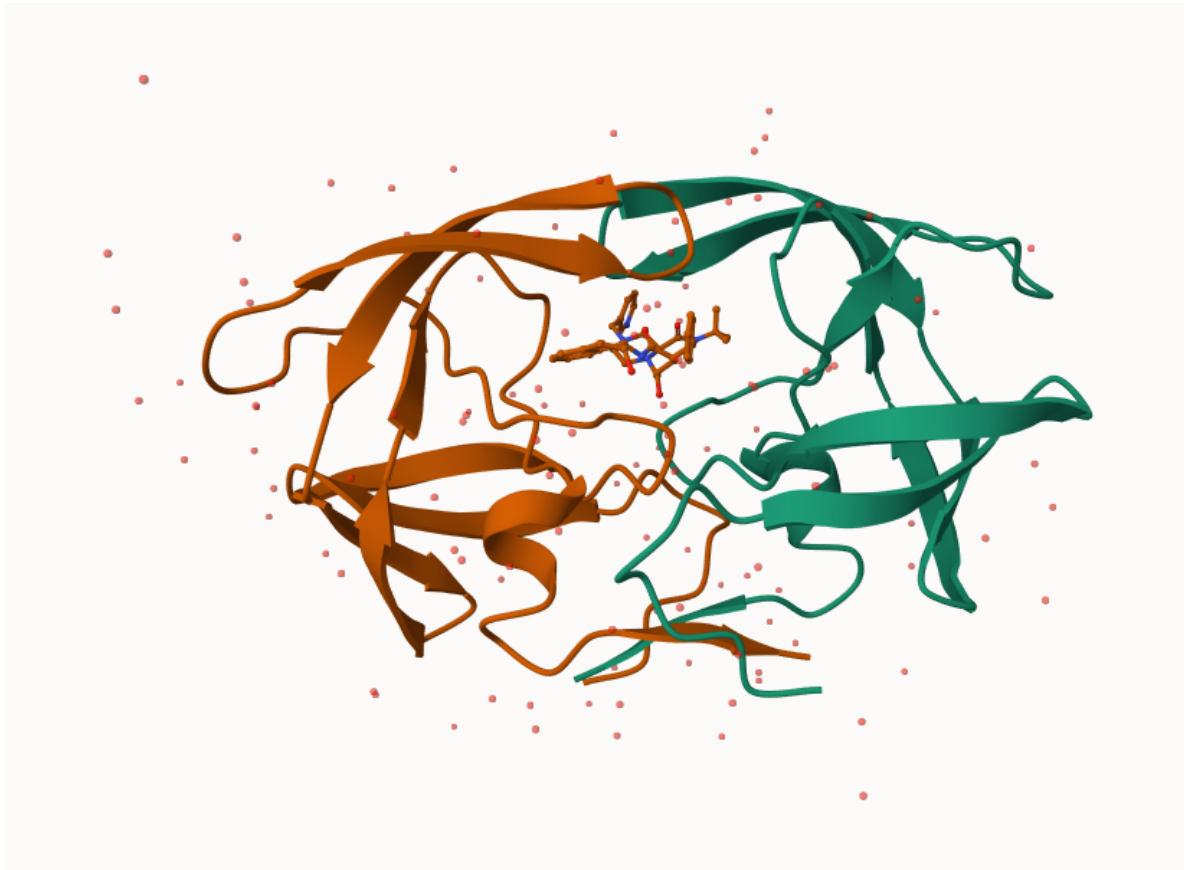


Figure 1: The HIV-Pr dimer with bound inhibitor



Figure 2: The catalytic ASP25 and active-site water

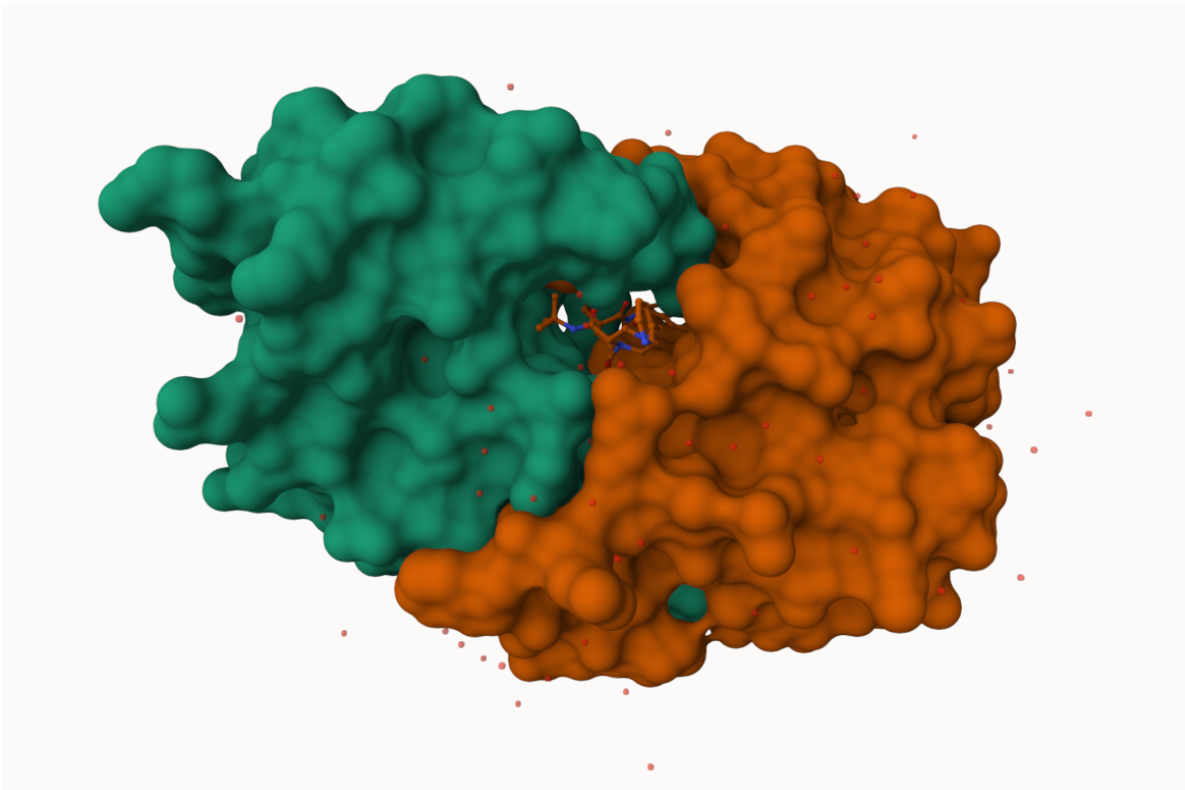


Figure 3: molecular surface

### Bio3D package for structural bioinformatics

We can use the bio3d package to read and analyze biomolecular data in R:

```
library(bio3d)  
  
hiv <- read.pdb("1hsg")
```

Note: Accessing on-line PDB file

```
hiv
```

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:
```

```
PQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWPKMIGGIGGFIKVRQYD  
QILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNFPQITLWQRPLVTIKIGGQLKE  
ALLDTGADDTVLEEMSLPGRWPKMIGGIGGFIKVRQYDQILIEICGHKAIGTVLVGPTP  
VNIIGRNLLTQIGCTLNF
```

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

```
head(hiv$atom)
```

	type	eleno	elety	alt	resid	chain	resno	insert	x	y	z	o	b
1	ATOM	1	N	<NA>	PRO	A	1	<NA>	29.361	39.686	5.862	1	38.10
2	ATOM	2	CA	<NA>	PRO	A	1	<NA>	30.307	38.663	5.319	1	40.62
3	ATOM	3	C	<NA>	PRO	A	1	<NA>	29.760	38.071	4.022	1	42.64
4	ATOM	4	O	<NA>	PRO	A	1	<NA>	28.600	38.302	3.676	1	43.40
5	ATOM	5	CB	<NA>	PRO	A	1	<NA>	30.508	37.541	6.342	1	37.87
6	ATOM	6	CG	<NA>	PRO	A	1	<NA>	29.296	37.591	7.162	1	38.40
	segid	elesy	charge										
1	<NA>	N	<NA>										
2	<NA>	C	<NA>										
3	<NA>	C	<NA>										
4	<NA>	O	<NA>										
5	<NA>	C	<NA>										
6	<NA>	C	<NA>										

```
Let's get the sequence
```

```
pdbseq( hiv )
```

1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20
"P"	"Q"	"I"	"T"	"L"	"W"	"Q"	"R"	"P"	"L"	"V"	"T"	"I"	"K"	"I"	"G"	"G"	"Q"	"L"	"K"
21	22	23	24	25	26	27	28	29	30	31	32	33	34	35	36	37	38	39	40

```

"E" "A" "L" "L" "D" "T" "G" "A" "D" "D" "T" "V" "L" "E" "E" "M" "S" "L" "P" "G"
41 42 43 44 45 46 47 48 49 50 51 52 53 54 55 56 57 58 59 60
"R" "W" "K" "P" "K" "M" "I" "G" "G" "I" "G" "F" "I" "K" "V" "R" "Q" "Y" "D"
61 62 63 64 65 66 67 68 69 70 71 72 73 74 75 76 77 78 79 80
"Q" "I" "L" "I" "E" "I" "C" "G" "H" "K" "A" "I" "G" "T" "V" "L" "V" "G" "P" "T"
81 82 83 84 85 86 87 88 89 90 91 92 93 94 95 96 97 98 99 1
"P" "V" "N" "I" "I" "G" "R" "N" "L" "L" "T" "Q" "I" "G" "C" "T" "L" "N" "F" "P"
2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21
"Q" "I" "T" "L" "W" "Q" "R" "P" "L" "V" "T" "I" "K" "I" "G" "G" "Q" "L" "K" "E"
22 23 24 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41
"A" "L" "L" "D" "T" "G" "A" "D" "D" "T" "V" "L" "E" "E" "M" "S" "L" "P" "G" "R"
42 43 44 45 46 47 48 49 50 51 52 53 54 55 56 57 58 59 60 61
"W" "K" "P" "K" "M" "I" "G" "G" "I" "G" "F" "I" "K" "V" "R" "Q" "Y" "D" "Q"
62 63 64 65 66 67 68 69 70 71 72 73 74 75 76 77 78 79 80 81
"I" "L" "I" "E" "I" "C" "G" "H" "K" "A" "I" "G" "T" "V" "L" "V" "G" "P" "T" "P"
82 83 84 85 86 87 88 89 90 91 92 93 94 95 96 97 98 99
"V" "N" "I" "I" "G" "R" "N" "L" "L" "T" "Q" "I" "G" "C" "T" "L" "N" "F"

```

Let's trim to chain A and get just its sequence:

```

chainA <- trim.pdb( hiv, chain="A" )
chainA.seq <- pdbseq(chainA)

```

Let's blast

```

blast <- blast.pdb(chainA.seq)

```

```

Searching ... please wait (updates every 5 seconds) RID = G54Y2PN4016
.....
Reporting 249 hits

```

```

head(blast$hit.tbl)

```

	queryid	subjectids	identity	alignmentlength	mismatches	gapopens	q.start
1	Query_1980777	1W5V_A	100.00	99	0	0	1
2	Query_1980777	2FDE_A	100.00	99	0	0	1
3	Query_1980777	1AJV_A	100.00	99	0	0	1
4	Query_1980777	2R38_A	98.99	99	1	0	1
5	Query_1980777	2R3T_A	98.99	99	1	0	1
6	Query_1980777	1HXB_A	98.99	99	1	0	1

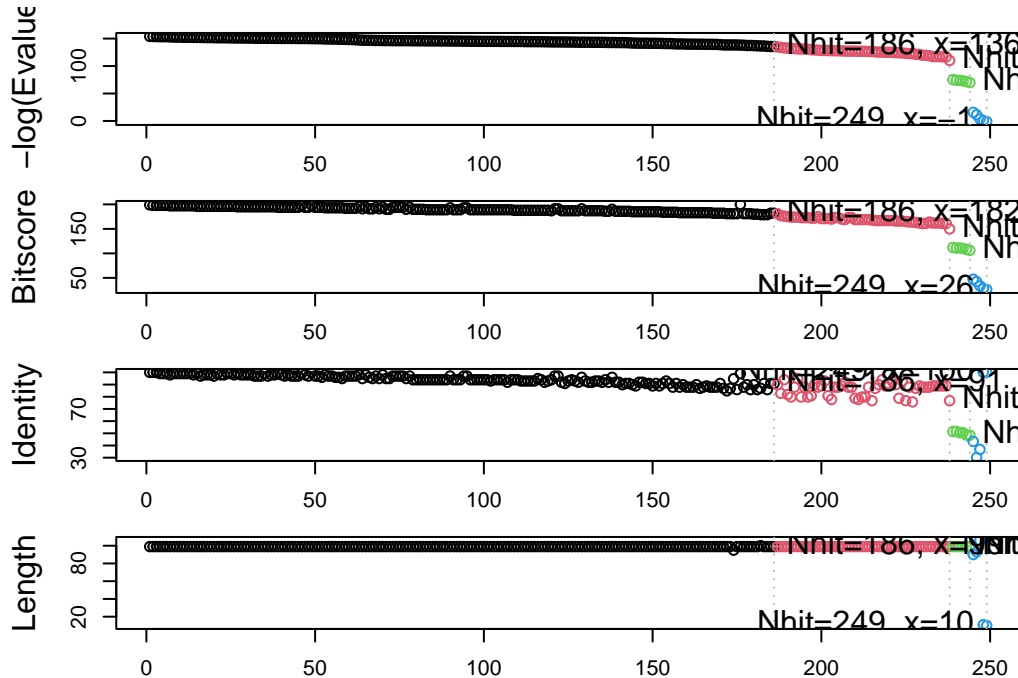
	q.end	s.start	s.end	evalue	bitscore	positives	mlog.evalue	pdb.id	acc
1	99		12	110	1.38e-67	199	100	153.9511	1W5V_A
2	99		2	100	1.70e-67	198	100	153.7426	2FDE_A
3	99		1	99	1.99e-67	198	100	153.5851	1AJV_A
4	99		1	99	2.50e-67	198	100	153.3569	2R38_A
5	99		1	99	2.50e-67	198	100	153.3569	2R3T_A
6	99		1	99	2.50e-67	198	100	153.3569	1HXB_A

Plot a quick overview of blast results

```
hits <- plot(blast)
```

```
* Possible cutoff values: 135 110 69 -2
Yielding Nhits: 186 238 244 249

* Chosen cutoff value of: 69
Yielding Nhits: 244
```



```
hits$pdb.id
```

```

[1] "1W5V_A" "2FDE_A" "1AJV_A" "2R38_A" "2R3T_A" "1HXB_A" "1BV9_A" "1AAQ_A"
[9] "1AXA_A" "1HVS_A" "1ZP8_A" "2QHC_A" "1A8G_A" "204L_A" "5COK_A" "1TCX_A"
[17] "2Z54_A" "1D4S_A" "1BV7_A" "1BWA_A" "1A9M_A" "2FLE_A" "1ODY_A" "1GNN_A"
[25] "1GNM_A" "5YRS_B" "1HEF_E" "10DX_A" "4QGI_A" "1BVE_A" "2AZ8_A" "1A30_A"
[33] "6DH6_A" "6DHO_A" "2I4D_A" "600S_A" "1RL8_A" "5YRS_A" "1ZSF_A" "2Q64_A"
[41] "6DH3_A" "2NPH_A" "2Q63_A" "1LZQ_A" "1FB7_A" "1G6L_A" "1HIV_A" "600U_A"
[49] "1HVC_A" "2I4V_A" "2AZ9_A" "600T_A" "2P3B_B" "5KAO_A" "2WLO_A" "6OPT_A"
[57] "1IZI_A" "1MRX_A" "2PYM_A" "2PYN_A" "1DMP_A" "4K4P_A" "1LV1_A" "1AID_A"
[65] "1LV1_A" "1ZBG_A" "3TKG_A" "1HVC_A" "5YOK_A" "1G6L_A" "1FGC_C" "3K4V_A"
[73] "3KT5_A" "3KT5_A" "4QLH_A" "4QLH_A" "2F3K_A" "4Q5M_A" "2AOC_A" "3B80_A"
[81] "3VF5_A" "2AVQ_A" "1DW6_C" "1KZK_A" "2HS1_A" "1K6C_A" "1MTB_A" "4Q1X_A"
[89] "4Q1W_A" "4Q5M_A" "3D1X_A" "2AVM_A" "3PWM_A" "3KT2_A" "3KT2_A" "1SDV_A"
[97] "3JVW_A" "3OY4_A" "1A94_A" "2HS2_A" "4EJ8_A" "2FGU_A" "2AVV_A" "3JW2_A"
[105] "3BVA_A" "1FFF_C" "3S43_B" "2NXD_A" "1FG6_C" "1EBK_C" "4Q1Y_A" "3EL4_A"
[113] "1F7A_A" "1K2B_A" "2FGV_A" "1Z8C_A" "2G69_A" "3EL9_A" "30XV_A" "1BDR_A"
[121] "3N3I_A" "3N3I_A" "30XW_A" "3S43_A" "3EM3_A" "3CYW_A" "5KQX_A" "2B60_A"
[129] "7DOZ_A" "1K2C_A" "1MT7_A" "3EM4_A" "4QJ9_A" "1BDL_A" "3LZS_A" "5T84_A"
[137] "4DQB_A" "7DOZ_A" "4QJ2_A" "3LZV_A" "1SGU_A" "2FXE_A" "1BDQ_A" "3U71_A"
[145] "2R5P_A" "40BD_A" "7MAS_A" "3IXO_A" "3D3T_A" "5YOJ_A" "3LZU_A" "4NJS_A"
[153] "3EKP_A" "1B6J_A" "3EKQ_A" "2RKF_A" "1C6X_A" "7MAR_A" "4DQF_A" "1RPI_A"
[161] "3OU1_B" "3PJ6_A" "2P3A_A" "60GQ_A" "30Q7_A" "5KR1_A" "30QD_A" "4RVI_A"
[169] "3OQA_A" "1B6K_A" "3OUD_B" "6MK9_A" "3S09_A" "1Q9P_A" "6I45_A" "7SEP_A"
[177] "4NJT_A" "3BXR_A" "4YOA_A" "4DQC_A" "2FDD_A" "2RKG_A" "4DQH_A" "2P3C_A"
[185] "4EP2_A" "4EP2_A" "4EQO_A" "4NPT_A" "60PU_A" "4NPU_A" "3U7S_A" "3HAW_A"
[193] "2AZB_A" "3TPP_A" "3HBO_A" "3GGU_A" "7N6T_A" "60PV_A" "4EQO_A" "60PX_A"
[201] "204N_A" "5T2E_A" "3UCB_A" "3KA2_A" "3FSM_A" "60PW_A" "2AZC_A" "3FSM_A"
[209] "3HLO_A" "2P3D_A" "3T3C_A" "7MYP_A" "6054_X" "60PY_A" "4Z4X_A" "60PZ_A"
[217] "2JE4_A" "1DAZ_C" "7MAP_A" "7MAQ_A" "1K1U_A" "2B7Z_A" "3MWS_A" "1K1T_A"
[225] "8DCH_A" "3I2L_A" "6P9A_A" "2FXD_A" "2J9J_A" "3DCK_A" "2J9J_B" "3NXE_A"
[233] "2040_A" "2040_A" "3NXE_A" "3KA2_A" "3HLO_A" "5B18_A" "1SIP_A" "2SAM_A"
[241] "1AZ5_A" "1SIV_A" "1HII_A" "1IVP_A"

```

## Prediction of functional motions

We can run a Normal Mode Analysis (NMA) to predict large scale motions/flexibility/dynamics of any biomolecule that we can read into R.

```
adk <- read.pdb("1ake")
```

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

```
adk_A <- trim.pdb(adk, chain="A")
adk_A
```

```
Call: trim.pdb(pdb = adk, chain = "A")

Total Models#: 1
Total Atoms#: 1954, XYZs#: 5862 Chains#: 1 (values: A)

Protein Atoms#: 1656 (residues/Calpha atoms#: 214)
Nucleic acid Atoms#: 0 (residues/phosphate atoms#: 0)

Non-protein/nucleic Atoms#: 298 (residues: 242)
Non-protein/nucleic resid values: [ AP5 (1), HOH (241) ]

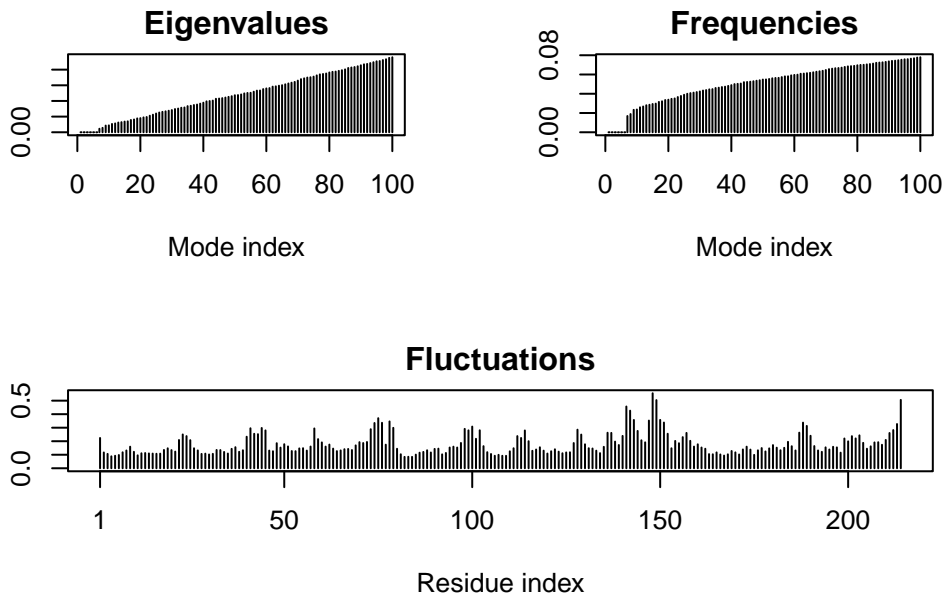
Protein sequence:
MRIILLGAPGAGKGTQAQFIMEKYGIPQISTGDMRLRAAVKSGSELGKQAKDIDMAGKLVT
DELVIALVKERIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVVDYVLEFDVPDELIVDRI
VGRRVHAPSGRVYHVKFNPPKVEGKDDVTGEELTTRKDQEETVRKRLVEYHQMTAPLIG
YYSKEAEAGNTKYAKVDGTPVAEVRADLEKILG

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

```
m <- nma(adk_A)
```

```
Building Hessian...      Done in 0.037 seconds.
Diagonalizing Hessian... Done in 0.424 seconds.
```

```
plot(m)
```



Let's write out a “trajectory” or predicted motion

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

## Play with 3D viewing in R

We can use the new **bio3dview** package, which is not yet on CRAN, to render interactive 3D views in R and HTML quarto output reports.

To install from GitHub we can use the **pak** package.

```
# library(bio3dview)
#
# view.pdb(adk)
```

## 4. Comparative structure analysis of Adenylate Kinase

The goal of this section is to perform **principal component analysis** (PCA) on the complete collection of Adenylate kinase structures in the protein data-bank (PDB).

**Adenylate kinase** (often called simply Adk) is a ubiquitous enzyme that functions to maintain the equilibrium between cytoplasmic nucleotides essential for many cellular processes. Adk operates by catalyzing the reversible transfer of a phosphoryl group from ATP to AMP. This reaction requires a rate limiting conformational transition (i.e. change in shape). Here we analyze all currently available Adk structures in the PDB to reveal detailed features and mechanistic principles of these essential shape changing transitions.

The **bio3d** package `pca()` function provides a convenient interface for performing PCA of biomolecular structure data. As we have discussed in previous classes, PCA is a statistical approach used to transform large data-sets down to a few important components that usefully describe the directions where there is most variance. In terms of protein structures PCA can be used to capture major structural variations within a set of structures (a.k.a. structure ensemble). This can make interpreting major conformational states (such as ‘active’ and ‘inactive’ or ‘ligand bound’ and ‘un-bound’ states) and structural mechanisms for activation or regulation more clear.

### Overview

Starting from only one Adk PDB identifier (**PDB ID: 1AKE**) we will search the entire PDB for related structures using BLAST, fetch, align and superpose the identified structures, perform PCA and finally calculate the normal modes of each individual structure in order to probe for potential differences in structural flexibility.

### Setup

We will begin by first installing the packages we need for today’s session.

*Install packages in the R console NOT your Rmd/Quarto file*

**Q10:** Which of the packages above is found only on BioConductor and not CRAN?

**Answer:** msa

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

**Answer:** 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

Below we perform a blast search of the PDB database to identify related structures to our query Adenylate kinase (ADK) sequence. In this particular example we use function `get.seq()` to fetch the query sequence for chain A of the PDB ID 1AKE and use this as input to `blast.pdb()`. Note that `get.seq()` would also allow the corresponding UniProt identifier.

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

```
Warning in get.seq("1ake_A"): Removing existing file: seqs.fasta
```

```
Fetching... Please wait. Done.
```

```
aa
```

	1	.	.	.	.	60
pdb 1AKE A	MRIILLGAPGAGKGTQAQFIMEKYGIPQISTGMLRAAVKGSELGKQAKDIDMAGKLVT					
	1	.	.	.	.	60
	61	.	.	.	.	120
pdb 1AKE A	DELVIALVKERIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFDVPDELVDR					
	61	.	.	.	.	120
	121	.	.	.	.	180
pdb 1AKE A	VGRRVHAPSGRVYHVKFNPPKVEGKDDVTGEELTRKDDQEETVRKRLVEYHQMTAPLIG					
	121	.	.	.	.	180
	181	.	.	.	214	
pdb 1AKE A	YYSKAEAGNTKYAKVDGTPVAEVRADLEKILG					
	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?

**Answer:** 214

Now we can use this sequence as a query to BLAST search the PDB to find similar sequences and structures.

```
# Blast or hmmer search  
b <- blast.pdb(aa)
```

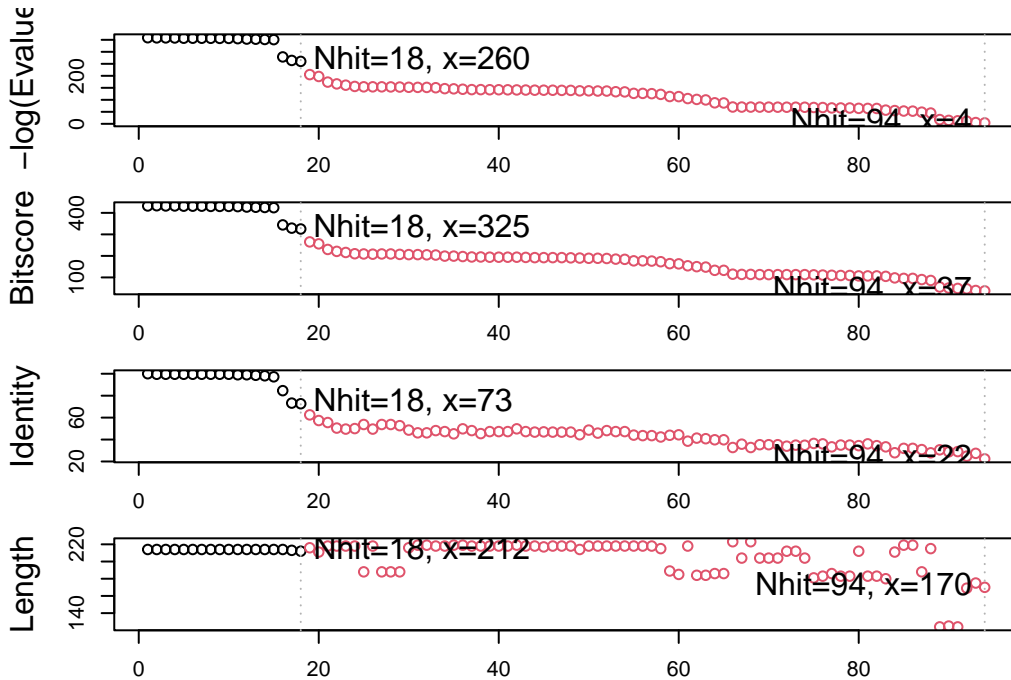
```
Searching ... please wait (updates every 5 seconds) RID = GG07S70C014  
.....  
Reporting 94 hits
```

The function `plot.blast()` facilitates the visualization and filtering of the Blast results. It will attempt to set a seed position to the point of largest drop-off in normalized scores (i.e. the biggest jump in E-values). In this particular case we specify a cutoff (after initial plotting) of to include only the relevant E.coli structures:

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

```
* Possible cutoff values:    260 3  
Yielding Nhits:          18 94  
  
* Chosen cutoff value of:  260  
Yielding Nhits:          18
```



```
# List out some 'top hits'
head(hits$pdb.id)
```

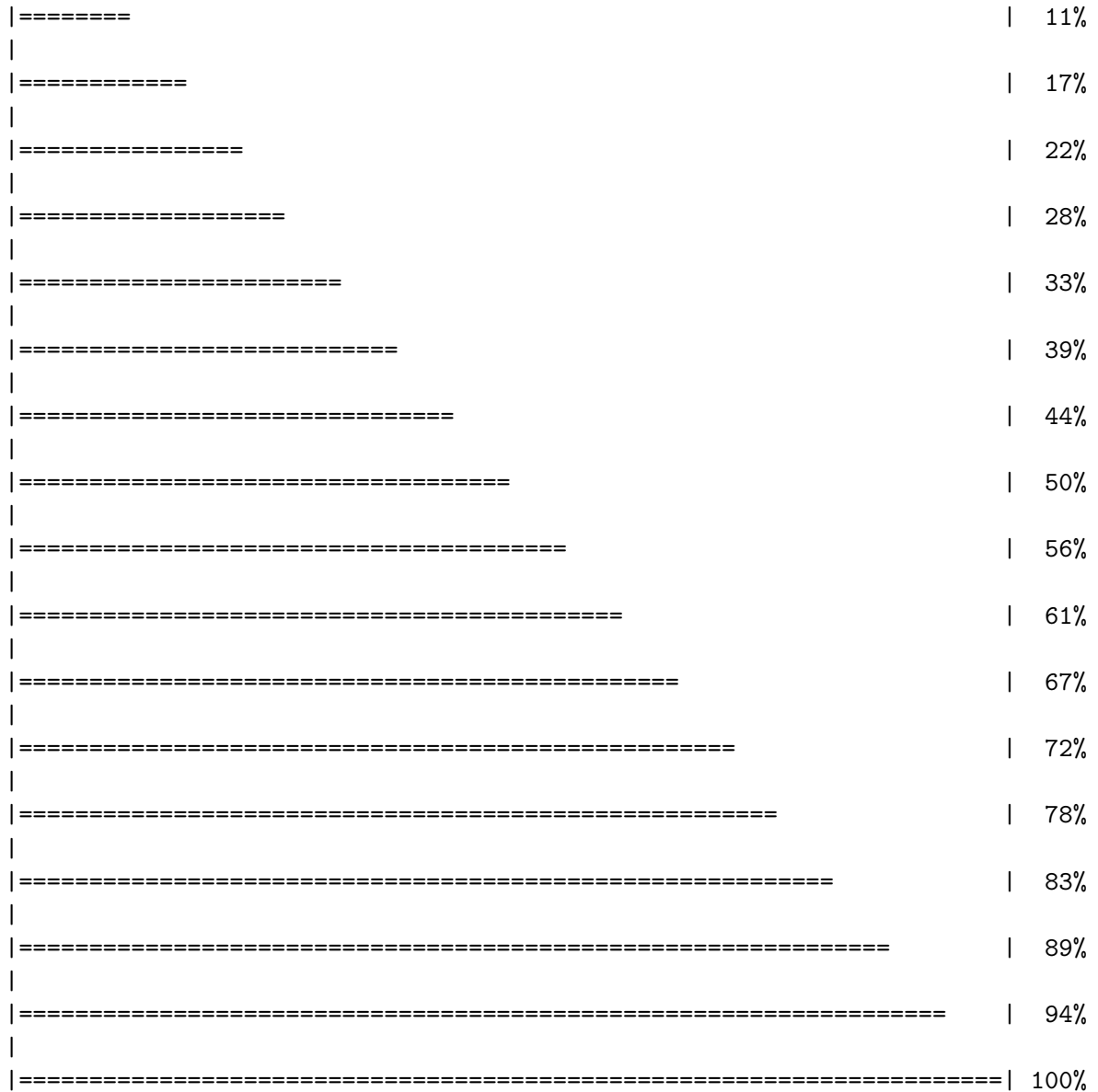
```
[1] "1AKE_A" "8BQF_A" "4X8M_A" "6S36_A" "8Q2B_A" "8RJ9_A"
```

The Blast search and subsequent filtering identified a total of 13 related PDB structures to our query sequence. The PDB identifiers of this collection are accessible through the `$pdb.id` attribute to the `hits` object (i.e. `hits$pdb.id`). Note that adjusting the cutoff argument (to `plot.blast()`) will result in a decrease or increase of hits.

We can now use function `get.pdb()` and `pdbslit()` to fetch and parse the identified structures.

```
# Download releated PDB files
files <- get.pdb(hits$pdb.id, path="pdbs", split=TRUE, gzip=TRUE)
```

```
|
|=====
|=====
```



## Align and superpose structures

Next we will use the `pdbaln()` function to align and also optionally fit (i.e. superpose) the identified PDB structures.

```
# Align releted PDBs
pdbs <- pdbaln(files, fit = TRUE, exefile="msa")
```

Reading PDB files:  
pdbs/split\_chain/1AKE\_A.pdb  
pdbs/split\_chain/8BQF\_A.pdb  
pdbs/split\_chain/4X8M\_A.pdb  
pdbs/split\_chain/6S36\_A.pdb  
pdbs/split\_chain/8Q2B\_A.pdb  
pdbs/split\_chain/8RJ9\_A.pdb  
pdbs/split\_chain/6RZE\_A.pdb  
pdbs/split\_chain/4X8H\_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/8PVW\_A.pdb  
pdbs/split\_chain/4K46\_A.pdb  
pdbs/split\_chain/4NP6\_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
- .. 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

```
pdb/seq: 1    name: pdbs/split_chain/1AKE_A.pdb
PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 2    name: pdbs/split_chain/8BQF_A.pdb
PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 3    name: pdbs/split_chain/4X8M_A.pdb
pdb/seq: 4    name: pdbs/split_chain/6S36_A.pdb
PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 5    name: pdbs/split_chain/8Q2B_A.pdb
PDB has ALT records, taking A only, rm.alt=TRUE
```

```

pdb/seq: 6    name: pdbs/split_chain/8RJ9_A.pdb
    PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 7    name: pdbs/split_chain/6RZE_A.pdb
    PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 8    name: pdbs/split_chain/4X8H_A.pdb
pdb/seq: 9    name: pdbs/split_chain/3HPR_A.pdb
    PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 10   name: pdbs/split_chain/1E4V_A.pdb
pdb/seq: 11   name: pdbs/split_chain/5EJE_A.pdb
    PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 12   name: pdbs/split_chain/1E4Y_A.pdb
pdb/seq: 13   name: pdbs/split_chain/3X2S_A.pdb
pdb/seq: 14   name: pdbs/split_chain/6HAP_A.pdb
pdb/seq: 15   name: pdbs/split_chain/6HAM_A.pdb
    PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 16   name: pdbs/split_chain/8PVW_A.pdb
    PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 17   name: pdbs/split_chain/4K46_A.pdb
    PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 18   name: pdbs/split_chain/4NP6_A.pdb

```

```

# Vector containing PDB codes for figure axis
ids <- basename.pdb(pdbs$id)

# Draw schematic alignment
#plot(pdbs, labels=ids)

# Plot is too large to render so I took a screenshot and inserted it manually:

```

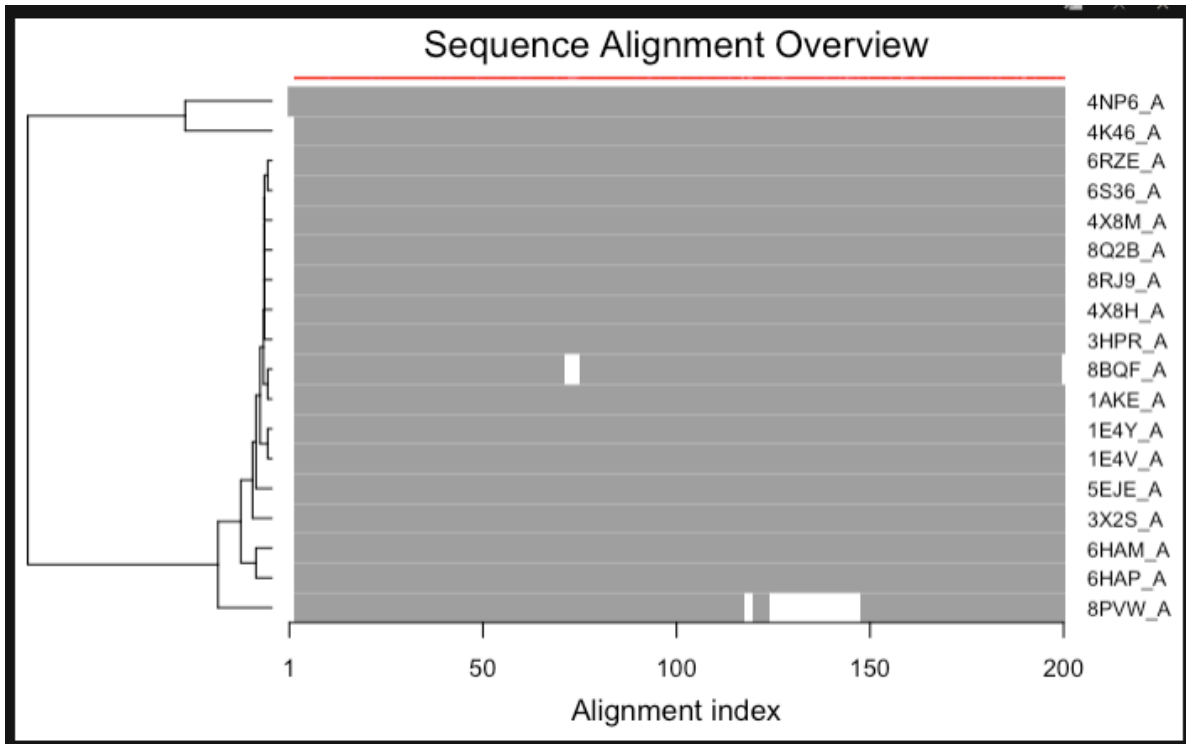


Figure 4: sequence alignment plot

### **Optional: Viewing our superposed structures**

We can view our superposed results with the new `bio3d.view view()` function:

```
library(bio3d.view)
library(rgl)

view.pdbs(pdbs)
```



```
## Annotate collected PDB structures
```

The function `pdb.annotate()` provides a convenient way of annotating the PDB files we have collected. Below we use the function to annotate each structure to its source species. This will come in handy when annotating plots later on:

```
anno <- pdb.annotate(ids)
unique(anno$source)
```

```
[1] "Escherichia coli"
[2] "Escherichia coli K-12"
[3] "Escherichia coli O139:H28 str. E24377A"
[4] "Escherichia coli str. K-12 substr. MDS42"
[5] "Photobacterium profundum"
[6] "Vibrio cholerae O1 biovar El Tor str. N16961"
```

We can view all available annotation data:

## anno

	structureId	chainId	macromoleculeType	chainLength	experimentalTechnique
1AKE_A	1AKE	A	Protein	214	X-ray
8BQF_A	8BQF	A	Protein	234	X-ray
4X8M_A	4X8M	A	Protein	214	X-ray
6S36_A	6S36	A	Protein	214	X-ray
8Q2B_A	8Q2B	A	Protein	214	X-ray
8RJ9_A	8RJ9	A	Protein	214	X-ray
6RZE_A	6RZE	A	Protein	214	X-ray
4X8H_A	4X8H	A	Protein	214	X-ray
3HPR_A	3HPR	A	Protein	214	X-ray
1E4V_A	1E4V	A	Protein	214	X-ray
5EJE_A	5EJE	A	Protein	214	X-ray
1E4Y_A	1E4Y	A	Protein	214	X-ray
3X2S_A	3X2S	A	Protein	214	X-ray
6HAP_A	6HAP	A	Protein	214	X-ray
6HAM_A	6HAM	A	Protein	214	X-ray
8PVW_A	8PVW	A	Protein	187	X-ray
4K46_A	4K46	A	Protein	214	X-ray
4NP6_A	4NP6	A	Protein	217	X-ray
	resolution	scopDomain			pfam
1AKE_A	2.000	Adenylate kinase			Adenylate kinase (ADK)
8BQF_A	2.050	<NA>			Adenylate kinase (ADK)
4X8M_A	2.600	<NA>			Adenylate kinase (ADK)
6S36_A	1.600	<NA>	Adenylate kinase, active site lid (ADK_lid)		
8Q2B_A	1.760	<NA>	Adenylate kinase, active site lid (ADK_lid)		
8RJ9_A	1.590	<NA>	Adenylate kinase, active site lid (ADK_lid)		
6RZE_A	1.690	<NA>			Adenylate kinase (ADK)
4X8H_A	2.500	<NA>			Adenylate kinase (ADK)
3HPR_A	2.000	<NA>			Adenylate kinase (ADK)
1E4V_A	1.850	Adenylate kinase			Adenylate kinase (ADK)
5EJE_A	1.900	<NA>			Adenylate kinase (ADK)
1E4Y_A	1.850	Adenylate kinase			Adenylate kinase (ADK)
3X2S_A	2.800	<NA>			Adenylate kinase (ADK)
6HAP_A	2.700	<NA>	Adenylate kinase, active site lid (ADK_lid)		
6HAM_A	2.550	<NA>			Adenylate kinase (ADK)
8PVW_A	2.000	<NA>	Adenylate kinase, active site lid (ADK_lid)		
4K46_A	2.010	<NA>			Adenylate kinase (ADK)
4NP6_A	2.004	<NA>			Adenylate kinase (ADK)
	ligandId				
1AKE_A		AP5			

8BQF_A	AP5	
4X8M_A	<NA>	
6S36_A	CL (3), NA, MG (2)	
8Q2B_A	AP5, SO4, MPO	
8RJ9_A	ADP (2)	
6RZE_A	CL (2), NA (3)	
4X8H_A	<NA>	
3HPR_A	AP5	
1E4V_A	AP5	
5EJE_A	AP5, CO	
1E4Y_A	AP5	
3X2S_A	JPY (2), AP5, MG	
6HAP_A	AP5	
6HAM_A	AP5	
8PVW_A	AP5	
4K46_A	ADP, AMP, PO4	
4NP6_A	<NA>	
		ligandName
1AKE_A	BIS(ADENOSINE)-5'-PENTAPHOSPHATE	
8BQF_A	BIS(ADENOSINE)-5'-PENTAPHOSPHATE	
4X8M_A	<NA>	
6S36_A	CHLORIDE ION (3), SODIUM ION, MAGNESIUM ION (2)	
8Q2B_A	BIS(ADENOSINE)-5'-PENTAPHOSPHATE, SULFATE ION, 3[N-MORPHOLINO]PROPANE SULFONIC ACID	
8RJ9_A	ADENOSINE-5'-DIPHOSPHATE (2)	
6RZE_A	CHLORIDE ION (2), SODIUM ION (3)	
4X8H_A	<NA>	
3HPR_A	BIS(ADENOSINE)-5'-PENTAPHOSPHATE	
1E4V_A	BIS(ADENOSINE)-5'-PENTAPHOSPHATE	
5EJE_A	BIS(ADENOSINE)-5'-PENTAPHOSPHATE, COBALT (II) ION	
1E4Y_A	BIS(ADENOSINE)-5'-PENTAPHOSPHATE	
3X2S_A	N-(pyren-1-ylmethyl)acetamide (2), BIS(ADENOSINE)-5'-PENTAPHOSPHATE, MAGNESIUM ION	
6HAP_A	BIS(ADENOSINE)-5'-PENTAPHOSPHATE	
6HAM_A	BIS(ADENOSINE)-5'-PENTAPHOSPHATE	
8PVW_A	BIS(ADENOSINE)-5'-PENTAPHOSPHATE	
4K46_A	ADENOSINE-5'-DIPHOSPHATE, ADENOSINE MONOPHOSPHATE, PHOSPHATE ION	
4NP6_A	<NA>	
		source
1AKE_A	Escherichia coli	
8BQF_A	Escherichia coli	
4X8M_A	Escherichia coli	
6S36_A	Escherichia coli	
8Q2B_A	Escherichia coli	
8RJ9_A	Escherichia coli	

6RZE_A	Escherichia coli
4X8H_A	Escherichia coli
3HPR_A	Escherichia coli K-12
1E4V_A	Escherichia coli
5EJE_A	Escherichia coli 0139:H28 str. E24377A
1E4Y_A	Escherichia coli
3X2S_A	Escherichia coli str. K-12 substr. MDS42
6HAP_A	Escherichia coli 0139:H28 str. E24377A
6HAM_A	Escherichia coli K-12
8PVW_A	Escherichia coli K-12
4K46_A	Photobacterium profundum
4NP6_A	Vibrio cholerae O1 biovar El Tor str. N16961

1AKE\_A STRUCTURE OF THE COMPLEX BETWEEN ADENYLATE KINASE FROM ESCHERICHIA COLI AND THE INHIBITOR

8BQF_A	E. coli Adenylate Kinase variant D158A (ΔD158A)
4X8M_A	E. coli adenylate kinase Asp84Δ
6S36_A	
8Q2B_A	
8RJ9_A	
6RZE_A	Crys
4X8H_A	
3HPR_A	
1E4V_A	
5EJE_A	
1E4Y_A	
3X2S_A	
6HAP_A	
6HAM_A	
8PVW_A	
4K46_A	
4NP6_A	

		citation	rObserved	rFree
1AKE_A	Muller, C.W., et al. J Mol Biology (1992)	0.19600	NA	
8BQF_A	Scheerer, D., et al. Proc Natl Acad Sci U S A (2023)	0.22073	0.25789	
4X8M_A	Kovermann, M., et al. Nat Commun (2015)	0.24910	0.30890	
6S36_A	Rogne, P., et al. Biochemistry (2019)	0.16320	0.23560	
8Q2B_A	Nam, K., et al. J Chem Inf Model (2024)	0.18320	0.22440	
8RJ9_A	Nam, K., et al. Sci Adv (2024)	0.15190	0.20290	
6RZE_A	Rogne, P., et al. Biochemistry (2019)	0.18650	0.23500	
4X8H_A	Kovermann, M., et al. Nat Commun (2015)	0.19610	0.28950	
3HPR_A	Schrank, T.P., et al. Proc Natl Acad Sci U S A (2009)	0.21000	0.24320	
1E4V_A	Muller, C.W., et al. Proteins (1993)	0.19600	NA	
5EJE_A	Kovermann, M., et al. Proc Natl Acad Sci U S A (2017)	0.18890	0.23580	

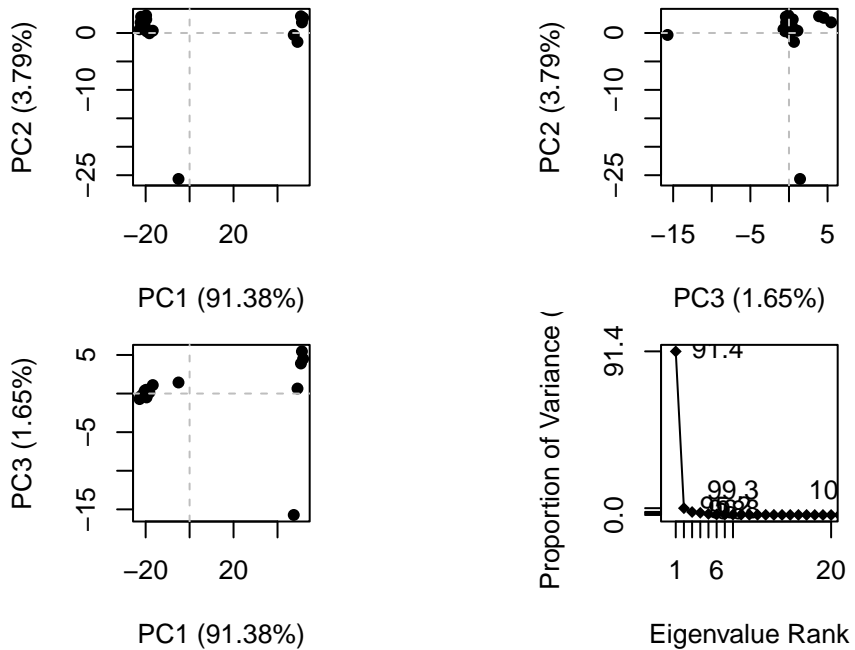
1E4Y_A	Muller, C.W., et al. Proteins (1993)	0.17800	NA
3X2S_A	Fujii, A., et al. Bioconjug Chem (2015)	0.20700	0.25600
6HAP_A	Kantaev, R., et al. J Phys Chem B (2018)	0.22630	0.27760
6HAM_A	Kantaev, R., et al. J Phys Chem B (2018)	0.20511	0.24325
8PVW_A	Rodriguez, J.A., et al. To be published	0.18590	0.23440
4K46_A	Cho, Y.-J., et al. To be published	0.17000	0.22290
4NP6_A	Kim, Y., et al. To be published	0.18800	0.22200
rWork spaceGroup			
1AKE_A	0.19600 P 21 2 21		
8BQF_A	0.21882 P 2 21 21		
4X8M_A	0.24630 C 1 2 1		
6S36_A	0.15940 C 1 2 1		
8Q2B_A	0.18100 P 1 21 1		
8RJ9_A	0.15010 P 21 21 2		
6RZE_A	0.18190 C 1 2 1		
4X8H_A	0.19140 C 1 2 1		
3HPR_A	0.20620 P 21 21 2		
1E4V_A	0.19600 P 21 2 21		
5EJE_A	0.18630 P 21 2 21		
1E4Y_A	0.17800 P 1 21 1		
3X2S_A	0.20700 P 21 21 21		
6HAP_A	0.22370 I 2 2 2		
6HAM_A	0.20311 P 43		
8PVW_A	0.18340 P 2 21 21		
4K46_A	0.16730 P 21 21 21		
4NP6_A	0.18600 P 43		

## Principal component analysis

Function `pca()` provides principal component analysis (PCA) of the structure data. PCA is a statistical approach used to transform a data set down to a few important components that describe the directions where there is most variance. In terms of protein structures PCA is used to capture major structural variations within an ensemble of structures.

PCA can be performed on the structural ensemble (stored in the `pdbs` object) with the function `pca.xyz()`, or more simply `pca()`.

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



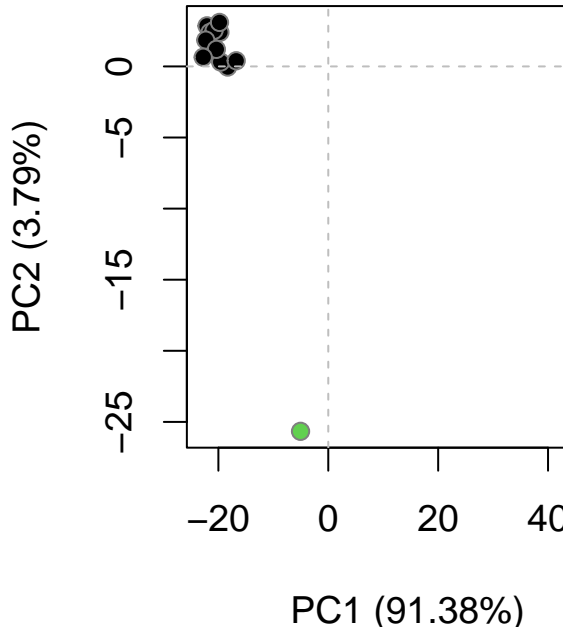
Function `rmsd()` will calculate all pairwise RMSD values of the structural ensemble. This facilitates clustering analysis based on the pairwise structural deviation:

```
# Calculate RMSD
rd <- rmsd(pdbs)
```

Warning in `rmsd(pdbs)`: No indices provided, using the 182 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)
```



The plot shows a conformer plot – a low-dimensional representation of the conformational variability within the ensemble of PDB structures. The plot is obtained by projecting the individual structures onto two selected PCs (e.g. PC-1 and PC-2). These projections display the inter-conformer relationship in terms of the conformational differences described by the selected PCs.

## 5. Optional further visualization

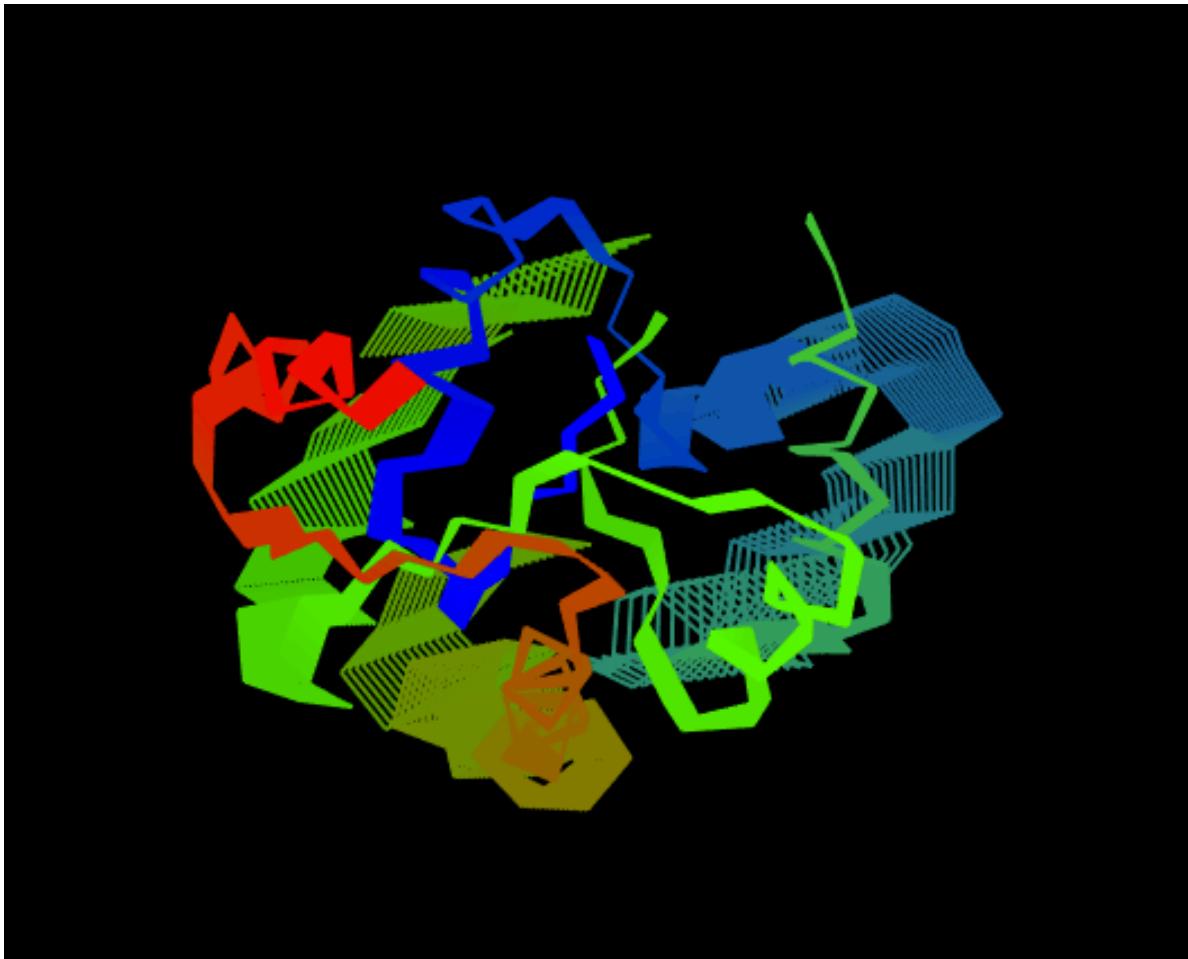
To visualize the major structural variations in the ensemble the function `mktrj()` can be used to generate a trajectory PDB file by interpolating along a give PC (eigenvector):

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

We can also view our results with the new `bio3d.view view()` function:

```
view.xyz(pc1)
```

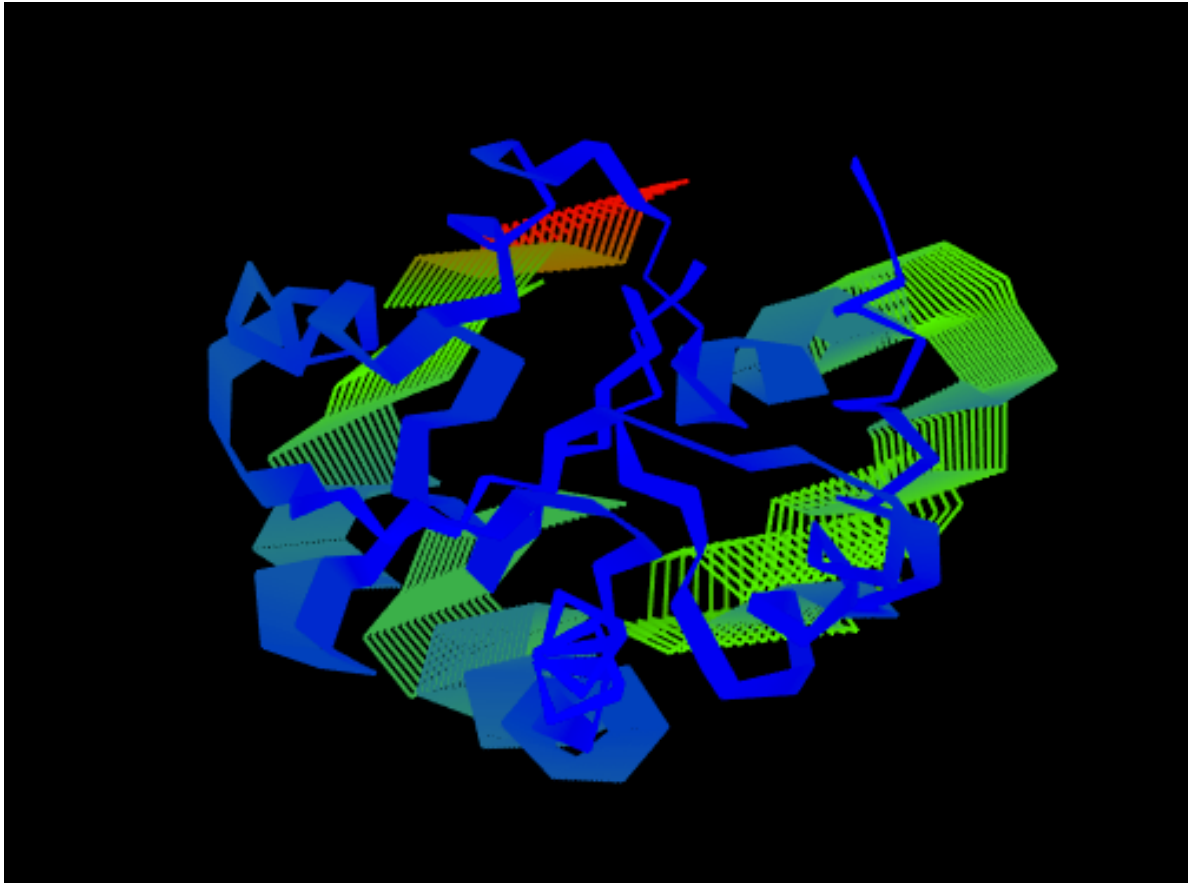
```
Potential all C-alpha atom structure(s) detected: Using calpha.connectivity()
```



We could set the color to highlight the most variable regions like so:

```
view.xyz(pc1, col=vec2color( rmsf(pc1) ))
```

```
Potential all C-alpha atom structure(s) detected: Using calpha.connectivity()
```



We can also plot our main PCA results with ggplot:

```
#Plotting results with ggplot2
library(ggplot2)
library(ggrepel)

df <- data.frame(PC1=pc.xray$z[,1],
                  PC2=pc.xray$z[,2],
                  col=as.factor(grps.rd),
                  ids=ids)

p <- ggplot(df) +
  aes(PC1, PC2, col=col, label=ids) +
  geom_point(size=2) +
  geom_text_repel(max.overlaps = 20) +
  theme(legend.position = "none")
p
```

