

class09

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```
##PDB Statistics
```

```
library(readr)
```

```
Warning: package 'readr' was built under R version 4.3.3
```

```
data <- 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.
```

```
data
```

```
# 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>
```

```
n.total <- sum(data$Total)
n.xray <- sum(data$`X-ray`)

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

```
[1] 81.43
```

```
n.em <- sum(data$EM)

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.

81.43% of structures in PDB are solved by X-Ray, and 12.27% are made through EM

```
n.protein <- c(176378, 20438, 12709, 342, 221, 83, 32)
n.protein2 <- sum(n.protein)
round(n.protein2/n.total)
```

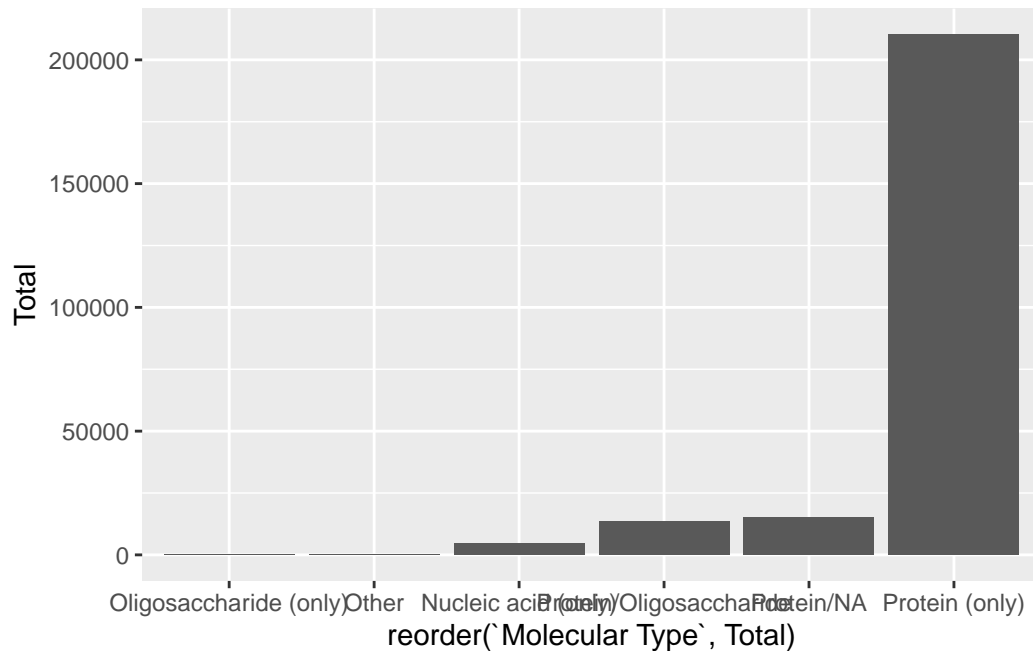
```
[1] 1
```

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

```
library(ggplot2)

# Can use the `` marks for columns with subtypes like "Molecular Weight"

ggplot(data, aes(x = reorder(`Molecular Type`, Total), y = Total)) + geom_col()
```



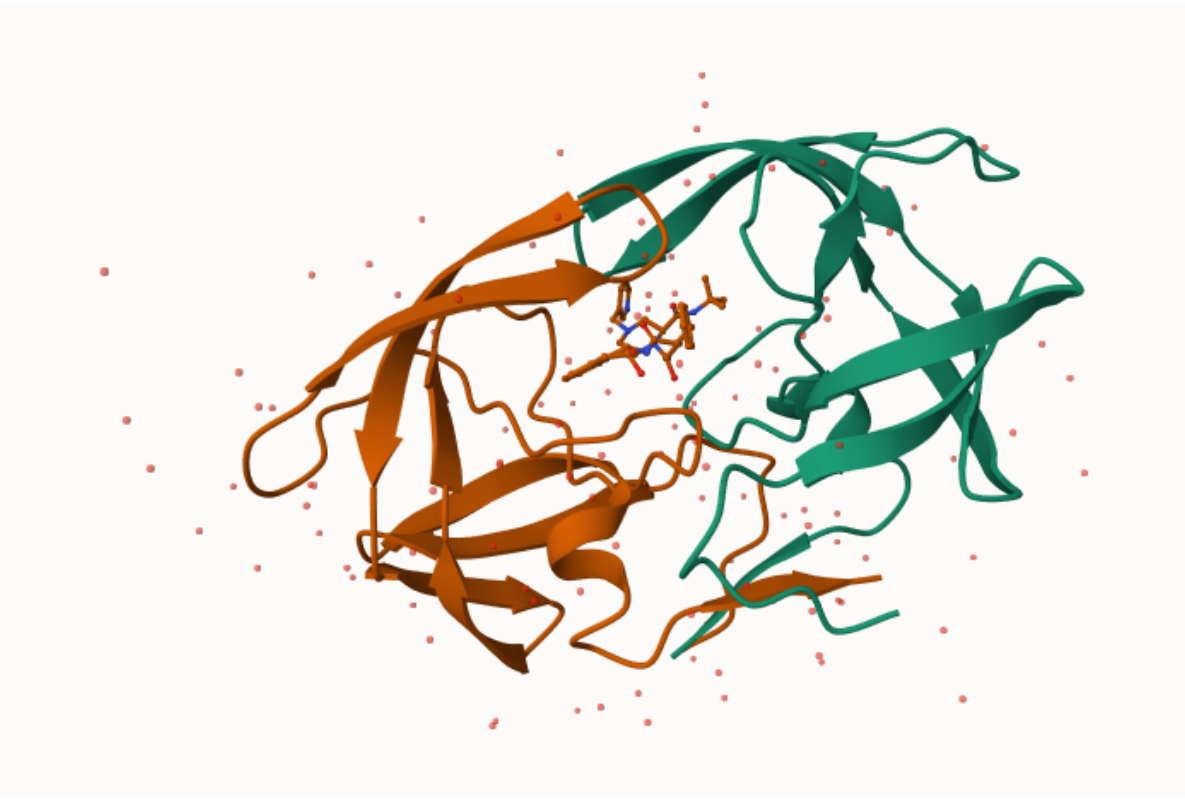
Q3: Type HIV in the PDB website search box on the home page and determine how many HIV-1 protease structures are in the current PDB?

There are 4,866 structures in the search.

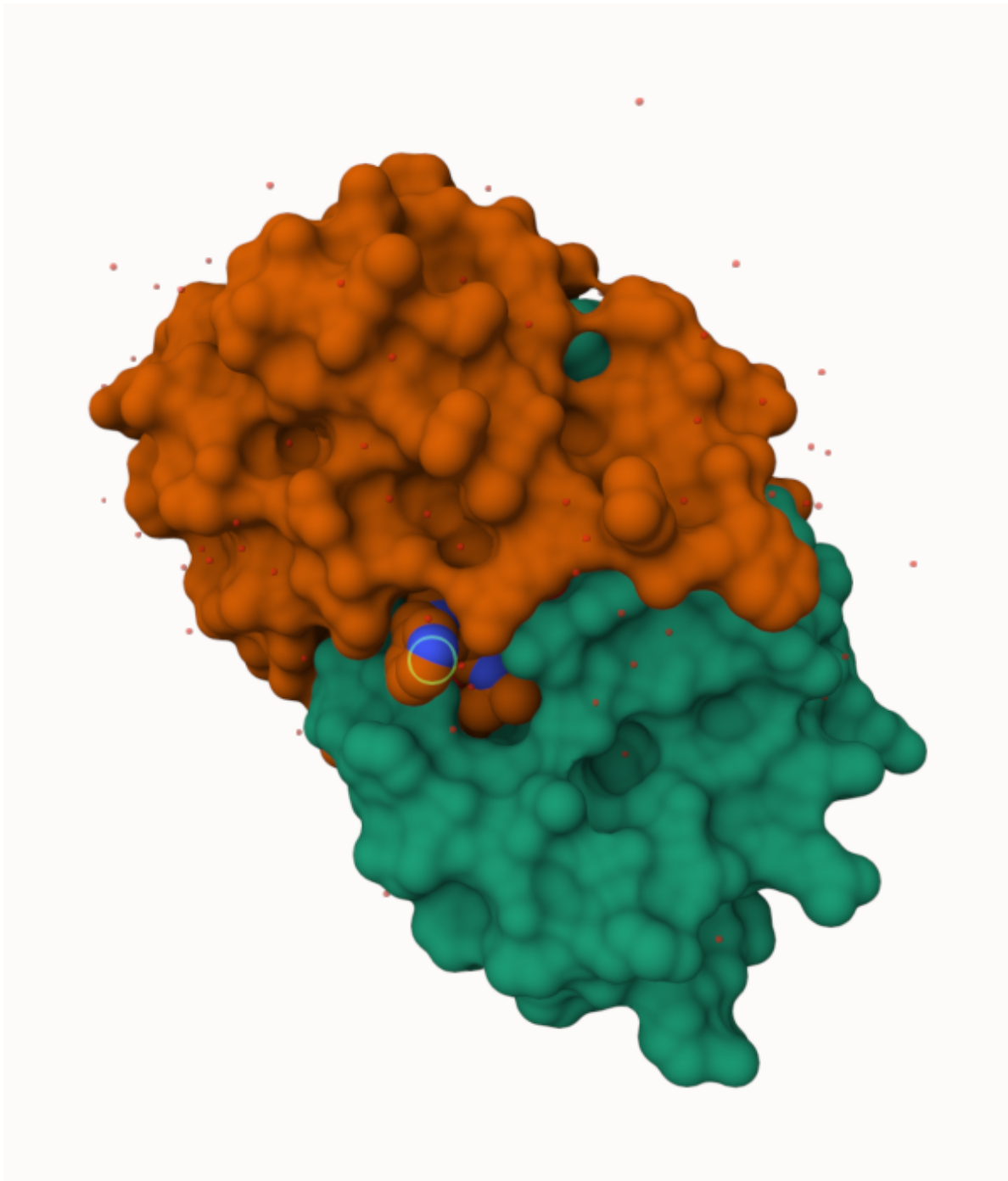
##Visualizing structure data

The Mol* viewer is embedded in many bioinformatics websites. The homepage is <https://molstar.org/>

I can insert any figure or image file using markdown format







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

We see one atom (oxygen) to represent water in these structures because X-ray crystallography

usually does not detect hydrogen because of low electron density.

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

This water molecule is number 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.

See 1HSG 2.png above.

Bio3D package for structural bioinformatics

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

```
library(bio3d)
```

```
Warning: package 'bio3d' was built under R version 4.3.3
```

```
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
```

Q7: How many amino acid residues are there in this pdb object?

There are 198 amino acid residues in this pdb object.

Q8: Name one of the two non-protein residues?

Water is one of the two non protein residues.

Q9: How many protein chains are in this structure?

There are 2 chains in the structure.

```
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" "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" "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 it's 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 = GB02ZHZF014
..
Reporting 249 hits

```

```

head(blast$hit.tbl)

```

	queryid	subjectids	identity	alignmentlength	mismatches	gapopens	q.start
1	Query_7763163	1W5V_A	100.00	99	0	0	1
2	Query_7763163	2FDE_A	100.00	99	0	0	1
3	Query_7763163	1AJV_A	100.00	99	0	0	1

```

4 Query_7763163      2R38_A      98.99          99          1          0          1
5 Query_7763163      2R3T_A      98.99          99          1          0          1
6 Query_7763163      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 1W5V_A
2     99       2     100 1.70e-67      198      100 153.7426 2FDE_A 2FDE_A
3     99       1      99 1.99e-67      198      100 153.5851 1AJV_A 1AJV_A
4     99       1      99 2.50e-67      198      100 153.3569 2R38_A 2R38_A
5     99       1      99 2.50e-67      198      100 153.3569 2R3T_A 2R3T_A
6     99       1      99 2.50e-67      198      100 153.3569 1HXB_A 1HXB_A

```

Plot a quick overview of blast results

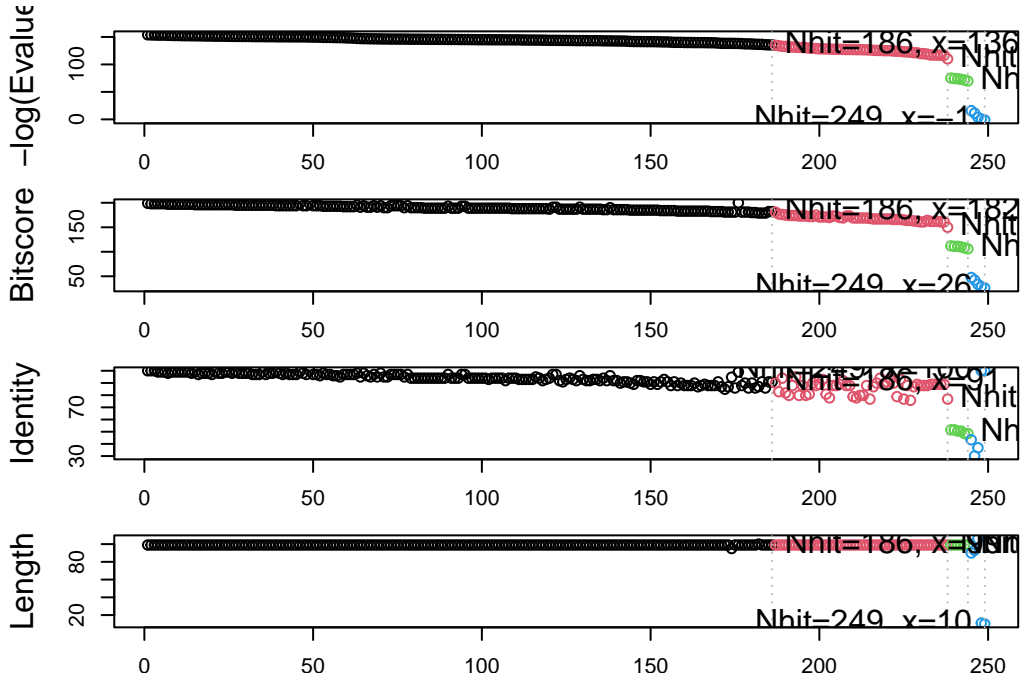
```
hits <- plot(blast)
```

```

* 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 an Normal Mode Analysis (NMA) to predict large scale motions/flexibility/dynamics of any biomolecule that we can read into R

Let's look at ADK and chain A only!

```
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:

```
MRIILLGAPGAGKGTQAQFIMEKYGIPQISTGMLRAAVKSGSELGKQAKDIMDAGKLVT  
DELVIALVKERIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVVDYVLEFDVPDELIVDRI  
VGRRVHAPSGRVYHVFKFNPPKVEGKDDVTGEELTRKDDQEETVRKRLVEYHQMTAPLIG  
YYSKAEAGNTKYAKVDGTPVAEVRADLEKILG
```

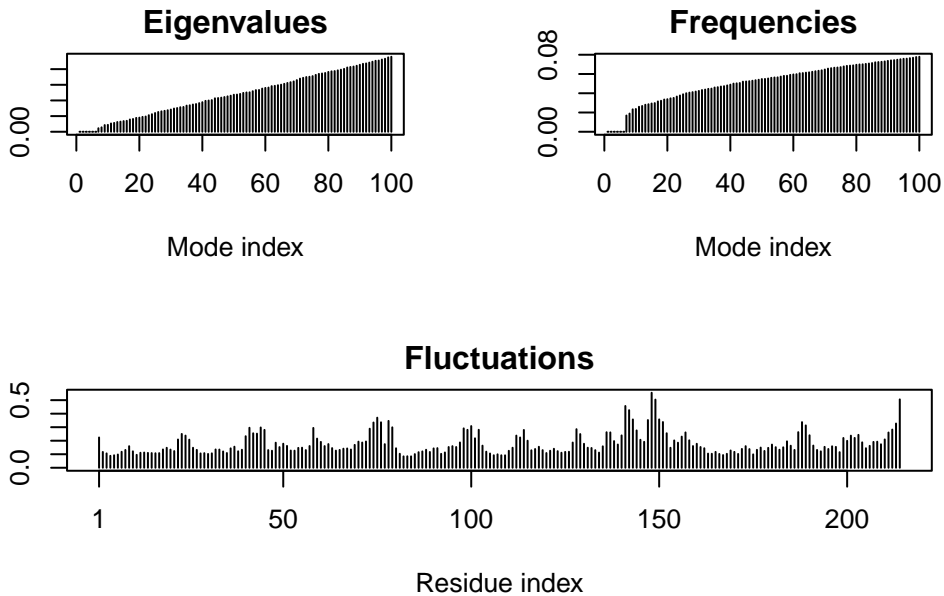
+ attr: atom, helix, sheet, seqres, xyz,
calpha, call

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

Building Hessian... Done in 0.02 seconds.

Diagonalizing Hessian... Done in 0.22 seconds.

```
plot(m)
```



Let's write out a “trajectory” of 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)
```

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

msa is found only on BioConductor and not CRAN

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

The package bio3d-view is not found on BioConductir or CRAN.

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
```

```
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	MRIILLGAPGAGKGTQAQFIMEKYGIPQISTGMLRAAVKSGSELGKQAKDIMDAGKLVT								
	1	60
	61	120
pdb 1AKE A	DELVIALVKERIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFDVPDELIVDRI								
	61	120
	121	180
pdb 1AKE A	VGRRVHAPSGRVYHVKFNPPKVEGKDDVTGEELTTRKDDQEETVRKRLVEYHQMTAPLIG								
	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?

There are 214 amino acids in this sequence.

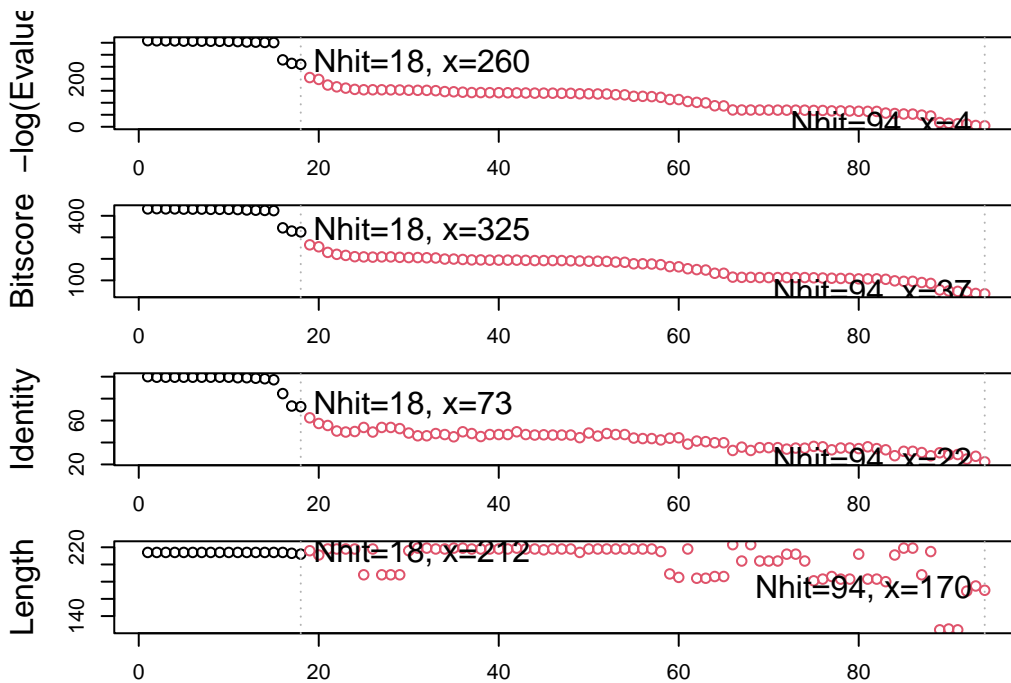
```
b <- blast.pdb(aa)
```

```
Searching ... please wait (updates every 5 seconds) RID = GB03J9GP014
.....
Reporting 94 hits
```

```
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"

hits <- NULL
hits$pdb.id <- c('1AKE_A', '6S36_A', '6RZE_A', '3HPR_A', '1E4V_A', '5EJE_A', '1E4Y_A', '3X2S_A', '6HA

files <- get.pdb(hits$pdb.id, path="pdbs", split=TRUE, gzip=TRUE)

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
```

A progress bar consisting of vertical bars and horizontal tick marks. The right side of each bar is labeled with a percentage value. The percentages increase sequentially from 0% to 100%, indicating the progress of a download or processing step.

Percentage
0%
8%
15%
23%
31%
38%
46%
54%
62%
69%
77%
85%
92%
100%

```
pdbs <- pdbaln(files, fit = TRUE, exefile="msa")
```

```
Reading PDB files:  
pdbs/split_chain/1AKE_A.pdb  
pdbs/split_chain/6S36_A.pdb
```

```
pdb/split_chain/6RZE_A.pdb
pdb/split_chain/3HPR_A.pdb
pdb/split_chain/1E4V_A.pdb
pdb/split_chain/5EJE_A.pdb
pdb/split_chain/1E4Y_A.pdb
pdb/split_chain/3X2S_A.pdb
pdb/split_chain/6HAP_A.pdb
pdb/split_chain/6HAM_A.pdb
pdb/split_chain/4K46_A.pdb
pdb/split_chain/3GMT_A.pdb
pdb/split_chain/4PZL_A.pdb

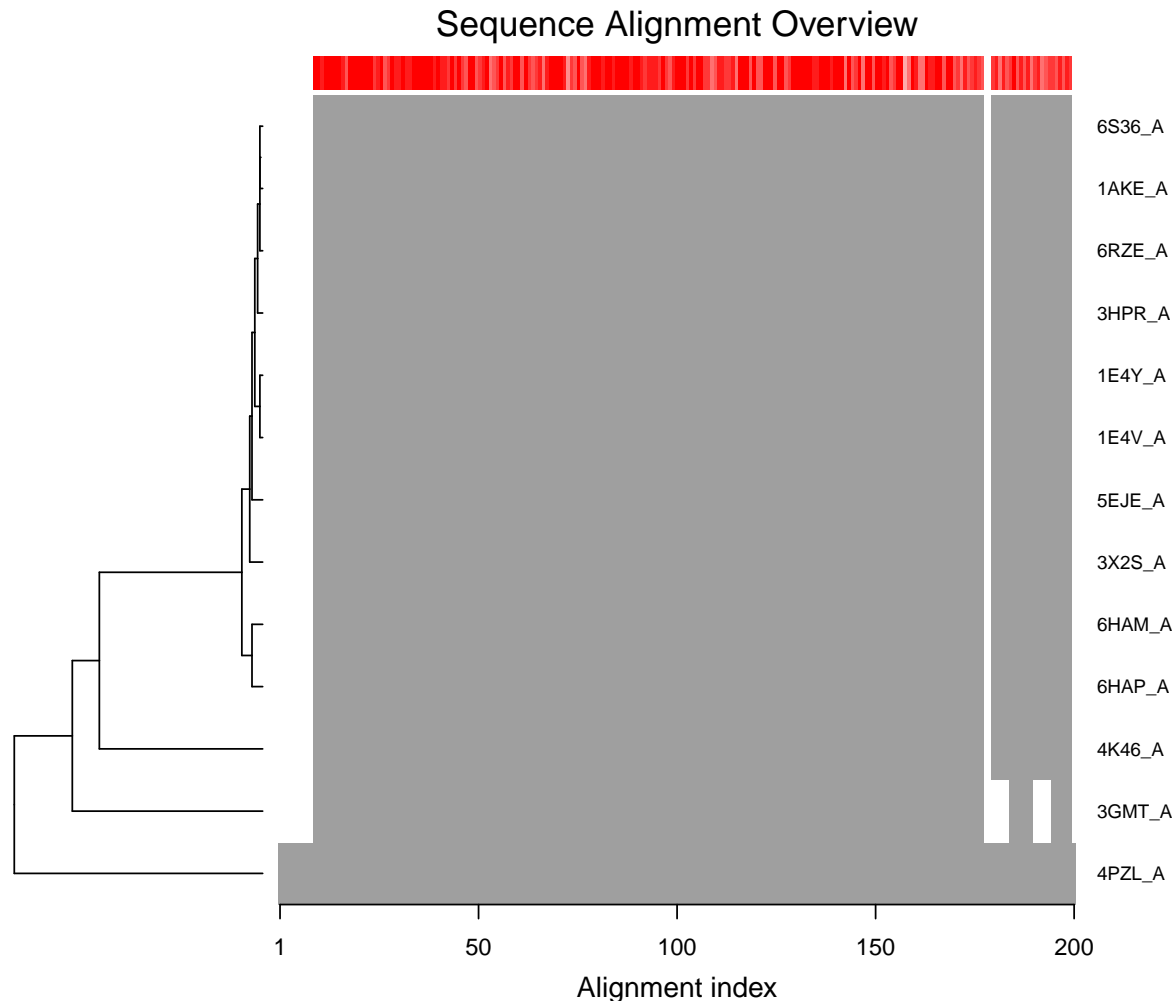
    PDB has ALT records, taking A only, rm.alt=TRUE
.    PDB has ALT records, taking A only, rm.alt=TRUE
.    PDB has ALT records, taking A only, rm.alt=TRUE
.    PDB has ALT records, taking A only, rm.alt=TRUE
..    PDB has ALT records, taking A only, rm.alt=TRUE
....    PDB has ALT records, taking A only,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

Extracting sequences

```
pdb/seq: 1    name: pdb/split_chain/1AKE_A.pdb
    PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 2    name: pdb/split_chain/6S36_A.pdb
    PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 3    name: pdb/split_chain/6RZE_A.pdb
    PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 4    name: pdb/split_chain/3HPR_A.pdb
    PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 5    name: pdb/split_chain/1E4V_A.pdb
pdb/seq: 6    name: pdb/split_chain/5EJE_A.pdb
    PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 7    name: pdb/split_chain/1E4Y_A.pdb
pdb/seq: 8    name: pdb/split_chain/3X2S_A.pdb
pdb/seq: 9    name: pdb/split_chain/6HAP_A.pdb
pdb/seq: 10   name: pdb/split_chain/6HAM_A.pdb
    PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 11   name: pdb/split_chain/4K46_A.pdb
    PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 12   name: pdb/split_chain/3GMT_A.pdb
pdb/seq: 13   name: pdb/split_chain/4PZL_A.pdb
```

```
ids <- basename.pdb(pdbs$id)

plot(pdbs, labels=ids)
```

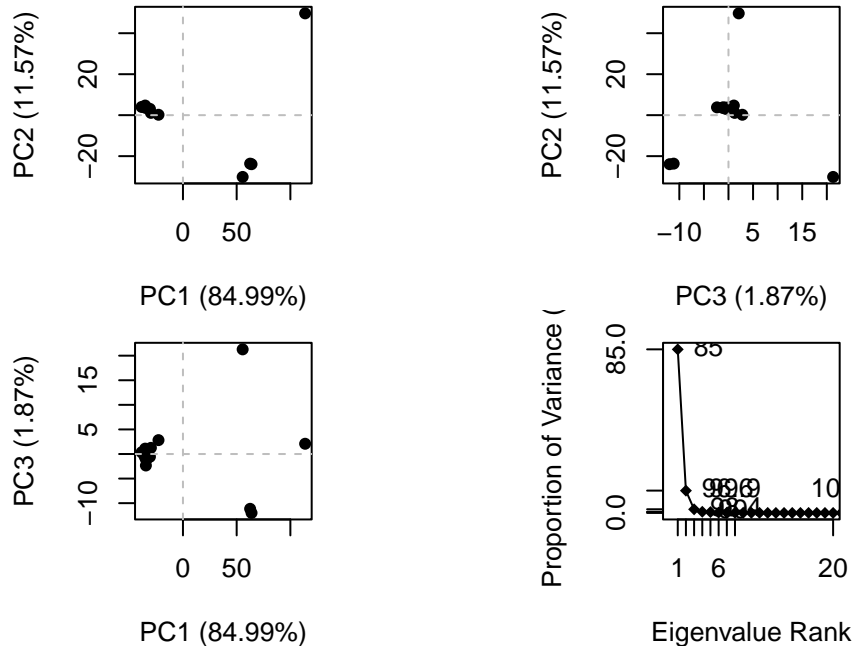


```
anno <- pdb.annotate(ids)
unique(anno$source)
```

```
[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] "Burkholderia pseudomallei 1710b"  
[7] "Francisella tularensis subsp. tularensis SCHU S4"
```

```
pc.xray <- pca(pdfs)  
plot(pc.xray)
```



```
rd <- rmsd(pdfs)
```

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

```
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)
```

