

Class09_Structural1

Zixuan Zeng (A16142927)

Table of contents

PDB statistics	1
Introduction to Bio3D in R	6
Prediction of functional motions (Normal mode analysis)	10

PDB statistics

```
PDBstat <- read.csv("Data_Export_Summary.csv", row.names = 1)
PDBstat
```

	X.ray	EM	NMR	Integrative	Multiple.methods
Protein (only)	176,204	20,299	12,708	342	218
Protein/Oligosaccharide	10,279	3,385	34	8	11
Protein/NA	9,007	5,897	287	24	7
Nucleic acid (only)	3,066	200	1,553	2	15
Other	173	13	33	3	0
Oligosaccharide (only)	11	0	6	0	1
	Neutron	Other	Total		
Protein (only)	83	32	209,886		
Protein/Oligosaccharide	1	0	13,718		
Protein/NA	0	0	15,222		
Nucleic acid (only)	3	1	4,840		
Other	0	0	222		
Oligosaccharide (only)	0	4	22		

```

# Convert numeric-looking columns from character to numeric
PDBstat[] <- lapply(PDBstat, function(x) as.numeric(gsub(", ", "", x)))

# Check structure
str(PDBstat)

'data.frame':   6 obs. of  8 variables:
 $ X.ray          : num  176204 10279 9007 3066 173 ...
 $ EM             : num  20299 3385 5897 200 13 ...
 $ NMR            : num  12708 34 287 1553 33 ...
 $ Integrative    : num  342 8 24 2 3 0
 $ Multiple.methods: num  218 11 7 15 0 1
 $ Neutron         : num  83 1 0 3 0 0
 $ Other           : num  32 0 0 1 0 4
 $ Total           : num  209886 13718 15222 4840 222 ...

X.ray <- as.numeric(sub(",","",PDBstat$X.ray))
EM <- as.numeric(sub(",","",PDBstat$EM))
Total <- as.numeric(sub(",","",PDBstat$Total))

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)    176204 20299 12708        342            218      83
2 Protein/Oligosacch~ 10279 3385   34          8             11       1

```

```

3 Protein/NA          9007   5897   287      24          7    0
4 Nucleic acid (only) 3066     200   1553      2         15    3
5 Other                173     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(stats$Total)
n.xray <- sum(stats$'X-ray')
n.em <- sum(stats$EM)
round(n.xray/n.total * 100, 2) + round(n.em/n.total * 100, 2)

```

[1] 93.7

Q1. 93.7% of the structures in the PDB were determined by X-ray and EM.

```
stats$Total[1]/sum(stats$Total) * 100
```

[1] 86.05059

Q2. 86.05% of the structures in the PDB contain protein only.

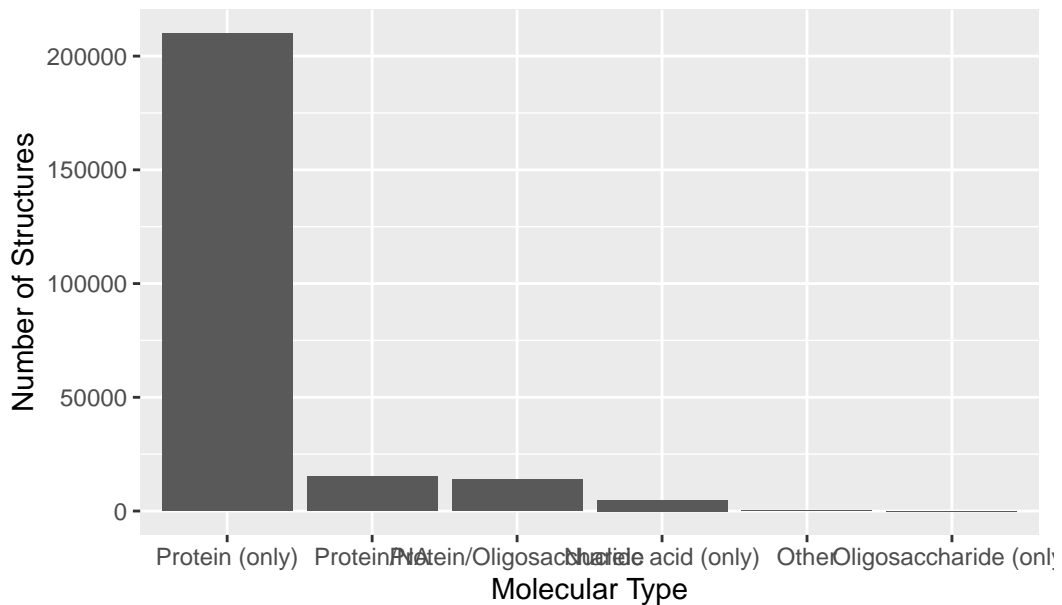
Q2b. Figure shown below

```

library(ggplot2)
ggplot(stats, aes(x=reorder(`Molecular Type`, -Total), Total)) + geom_bar(stat="identity") +

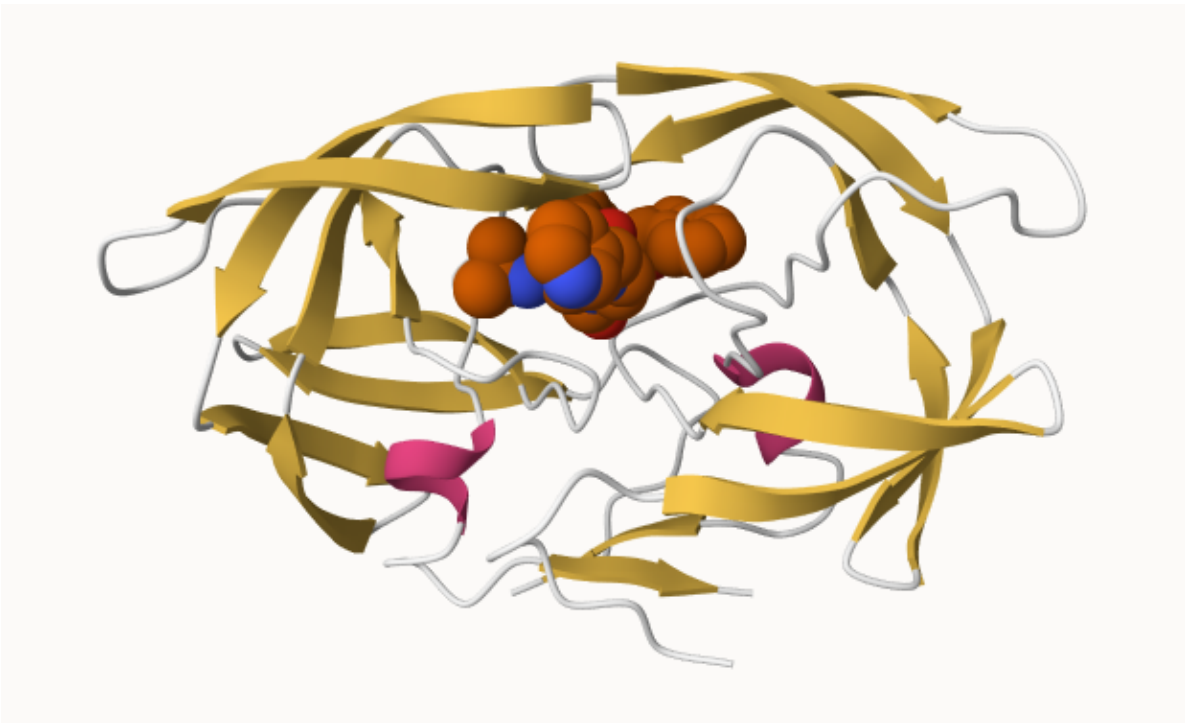
```

PDB Molecular Type Overview

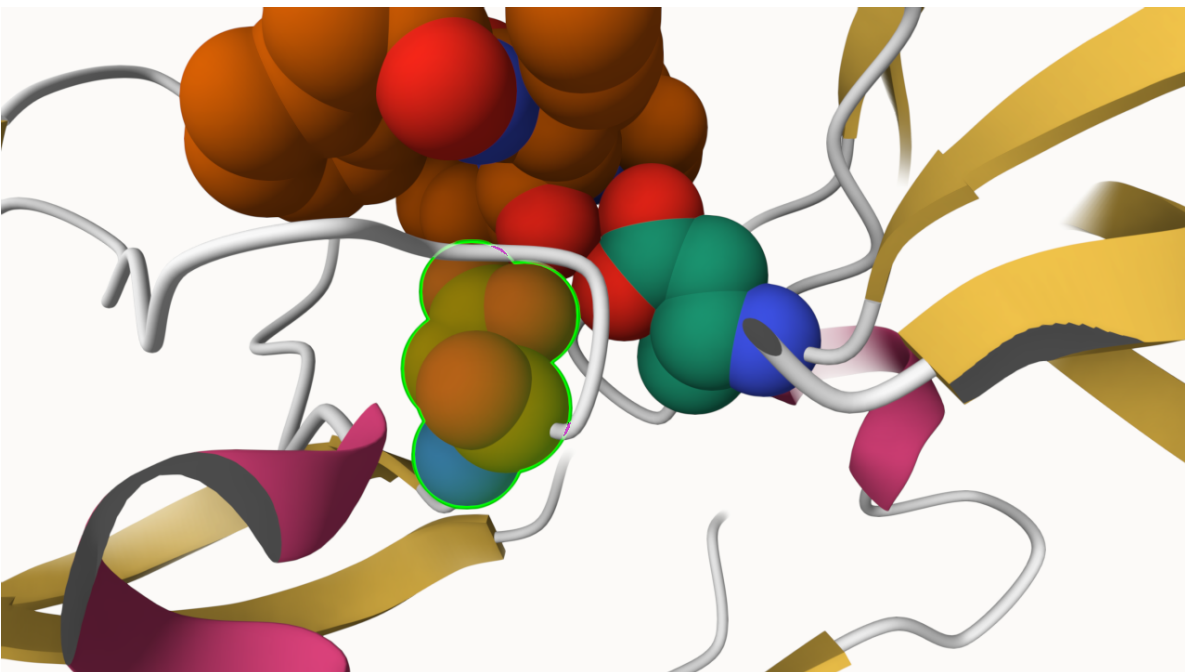


Q3. 4866 structures found for HIV-1 protease in the current pdb.

```
knitr::include_graphics("1HSG.png")
```

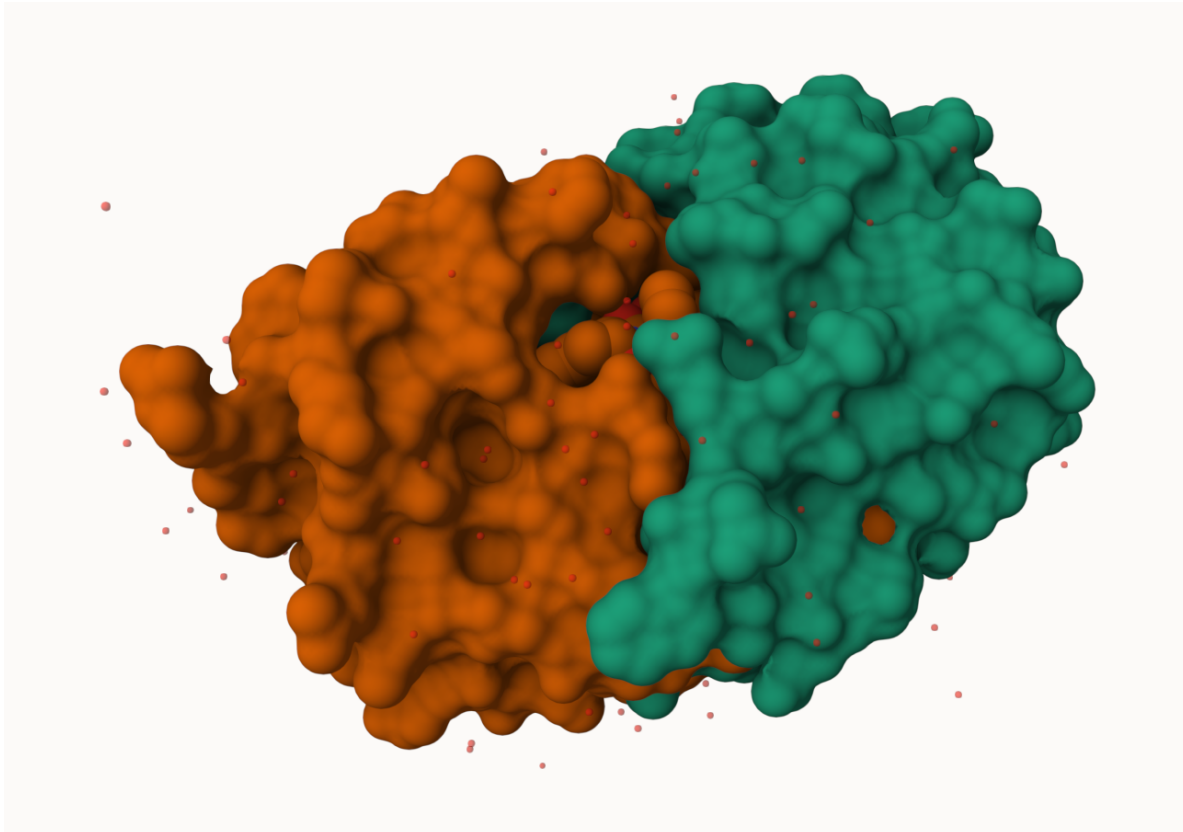


```
knitr::include_graphics("1HSG_(1).png")
```



Q4. We can only see the oxygen atom from a water molecule because of the resolution of the structure. Hydrogen atom is not observed at this resolution.

```
knitr::include_graphics("1HSG (2).png")
```



Introduction to Bio3D 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:
PQITLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWPQMIGGIGGFIKVRQYD
QILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNFPQITLWQRPLVTIKIGGQLKE
ALLDTGADDTVLEEMSLPGRWPQMIGGIGGFIKVRQYDQILIEICGHKAIGTVLVGPTP
VNIIGRNLLTQIGCTLNF

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

```

attributes(hiv)

```

$names
[1] "atom"    "xyz"     "seqres"  "helix"   "sheet"   "calpha"  "remark" "call"

$class
[1] "pdb" "sse"

```

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>
```

```
  pdbseq(hiv) #for both chain A and B
```

```
 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" "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"
```

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

```
#Blast chain A
blast <- blast.pdb(chainA.seq)
```

```
Searching ... please wait (updates every 5 seconds) RID = G7EPRK26016
.
.
.
Reporting 249 hits
```

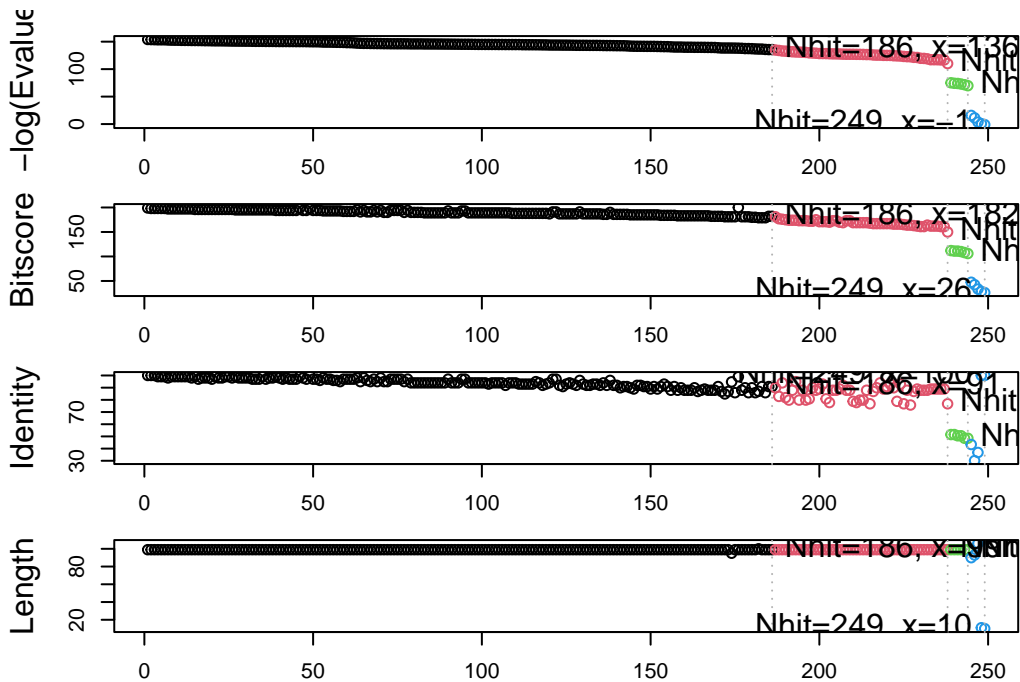
```
head(blast$hit.tbl)
```

	queryid	subjectids	identity	alignmentlength	mismatches	gapopens	q.start	q.end	s.start	s.end	evalue	bitscore	positives	mlog.evalue	pdb.id	acc	
1	Query_4119375	1W5V_A	100.00		99	0	0		99	12	110	1.38e-67	199	100	153.9511	1W5V_A	1W5V_A
2	Query_4119375	2FDE_A	100.00		99	0	0		99	2	100	1.70e-67	198	100	153.7426	2FDE_A	2FDE_A
3	Query_4119375	1AJV_A	100.00		99	0	0		99	1	99	1.99e-67	198	100	153.5851	1AJV_A	1AJV_A
4	Query_4119375	2R38_A	98.99		99	1	0		99	1	99	2.50e-67	198	100	153.3569	2R38_A	2R38_A
5	Query_4119375	2R3T_A	98.99		99	1	0		99	1	99	2.50e-67	198	100	153.3569	2R3T_A	2R3T_A
6	Query_4119375	1HXB_A	98.99		99	1	0		99	1	99	2.50e-67	198	100	153.3569	1HXB_A	1HXB_A

```
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] "3JWV_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" "4EQ0_A" "4NPT_A" "60PU_A" "4NPU_A" "3U7S_A" "3HAW_A"  
[193] "2AZB_A" "3TT_P_A" "3HBO_A" "3GGU_A" "7N6T_A" "60PV_A" "4EQ0_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" "3NZE_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 (Normal mode analysis)

```
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 0.012 seconds.
Diagonalizing Hessian... Done in 0.266 seconds.

```
plot(m)
```

