Class 9: Structural Bioinformatics 1

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1. The RCSB Protein Data Bank (PDB)

Protein structures by X-ray crystalography dominate this database. We are skipping Q1-Q3 because the website was too slow for us.

2. Visualizing the HIV-1 protease structure

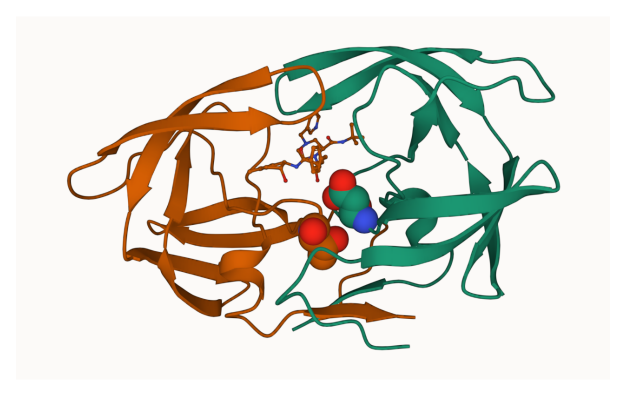


Figure 1: HIV-Pr structure from 1hsg

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

Because the hydrogen atoms are so small that with this high resolution, we cannot visualize them in the image.

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?

The water is located between the ligand and binding site. 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 (we recommend "Ball & Stick" for these side-chains). Add this figure to your Quarto document.

Discussion Topic: Can you think of a way in which indinavir, or even larger ligands and substrates, could enter the binding site?

One way that indinavir or even larger ligands and substrates could enter the binding site is to make the polymers more flexible so that it could allow them to enter. We can also break bonds to make the protein smaller that could allows them to enter.

3. Introduction to Bio3D in R

Bio3D is an R package for structural bioinformatics. To use it we need to call it up with the library() function

```
library(bio3d)
To read a PDB file we can use read.pdb()
pdb <- read.pdb("1hsg")
Note: Accessing on-line PDB file</pre>
```

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
The ATOM records of a PDB file are stored in pdb$atom
  head(pdb$atom)
 type eleno elety alt resid chain resno insert
                                                     X
                                                                  z o
                                                            У
1 ATOM
          1
                N < NA >
                         PRO
                                 Α
                                       1 <NA> 29.361 39.686 5.862 1 38.10
2 ATOM
          2
               CA <NA>
                         PRO
                                       1 <NA> 30.307 38.663 5.319 1 40.62
                                 Α
3 ATOM
          3
               C <NA>
                         PRO
                                 Α
                                       1 <NA> 29.760 38.071 4.022 1 42.64
4 ATOM
          4
               O <NA>
                         PRO
                                      1 <NA> 28.600 38.302 3.676 1 43.40
                                 Α
5 ATOM
          5
               CB <NA>
                         PRO
                                 Α
                                       1 <NA> 30.508 37.541 6.342 1 37.87
                                       1 <NA> 29.296 37.591 7.162 1 38.40
6 ATOM
          6
               CG <NA>
                         PRO
                                 Α
 segid elesy charge
1 <NA>
           N
               <NA>
2 <NA>
           С
               <NA>
3 <NA>
           C <NA>
```

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

4 <NA>

5 <NA> 6 <NA>

O <NA>
C <NA>

<NA>

С

198

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

MK1

Q9: How many protein chains are in this structure?

2

4. Comparative structure analysis of Adenylate Kinase

```
# Install packages in the R console NOT your Rmd/Quarto file
#install.packages("ggrepel")
#install.packages("devtools")
#install.packages("BiocManager")

#BiocManager::install("msa")
#devtools::install_bitbucket("Grantlab/bio3d-view")
```

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?:

bio3d-view

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

TRUE

Compare analysis of Adenylate kinase (ADK)

We will start our analysis with a single PDB id (code from the PDB database): 1AKE First we get its primary sequence:

```
aa <- get.seq("1ake_a")</pre>
```

```
Fetching... Please wait. Done.
  aa
                                                                         60
pdb|1AKE|A
             \tt MRIILLGAPGAGKGTQAQFIMEKYGIPQISTGDMLRAAVKSGSELGKQAKDIMDAGKLVT
                                                                         60
                                                                          120
            DELVIALVKERIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFDVPDELIVDRI
                                                                         120
           121
                                                                         180
            VGRRVHAPSGRVYHVKFNPPKVEGKDDVTGEELTTRKDDQEETVRKRLVEYHQMTAPLIG
pdb|1AKE|A
           121
                                                                         180
           181
                                               214
pdb|1AKE|A
             YYSKEAEAGNTKYAKVDGTKPVAEVRADLEKILG
               . . . . . . 214
           181
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>
```

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

Searching ... please wait (updates every 5 seconds) RID = NGEXRXRH013 Reporting 98 hits

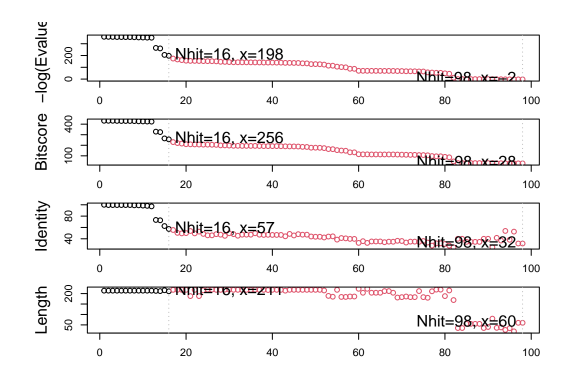
Plot a summary of search results hits <- plot(b)

* Possible cutoff values: 197 -3

Yielding Nhits: 16 98

* Chosen cutoff value of: 197

Yielding Nhits: 16



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

[1] "1AKE_A" "4X8M_A" "6S36_A" "6RZE_A" "4X8H_A" "3HPR_A"

Use these ADK structures for analysis:

```
hits <- NULL
  hits$pdb.id <- c('1AKE_A','6S36_A','6RZE_A','3HPR_A','1E4V_A','5EJE_A','1E4Y_A','3X2S_A','
Download all these PDB files from the online database...
  # Download related PDB files
  files <- get.pdb(hits$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% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15% | 15%
```

```
# Align related PDBs
pdbs <- pdbaln(files, fit = TRUE)#, exefile="msa")</pre>
```

```
Reading PDB files:
pdbs/split_chain/1AKE_A.pdb
pdbs/split_chain/6S36_A.pdb
pdbs/split_chain/6RZE_A.pdb
pdbs/split chain/3HPR A.pdb
pdbs/split_chain/1E4V_A.pdb
pdbs/split_chain/5EJE_A.pdb
pdbs/split_chain/1E4Y_A.pdb
pdbs/split_chain/3X2S_A.pdb
pdbs/split_chain/6HAP_A.pdb
pdbs/split_chain/6HAM_A.pdb
pdbs/split_chain/4K46_A.pdb
pdbs/split_chain/3GMT_A.pdb
pdbs/split_chain/4PZL_A.pdb
   PDB has ALT records, taking A only, rm.alt=TRUE
     PDB has ALT records, taking A only, rm.alt=TRUE
       PDB has ALT records, taking A only, rm.alt=TRUE
    PDB has ALT records, taking A only, rm.alt=TRUE
```

Extracting sequences

```
name: pdbs/split_chain/1AKE_A.pdb
pdb/seq: 1
   PDB has ALT records, taking A only, rm.alt=TRUE
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
             name: pdbs/split chain/5EJE A.pdb
pdb/seq: 6
   PDB has ALT records, taking A only, rm.alt=TRUE
pdb/seq: 7
             name: pdbs/split_chain/1E4Y_A.pdb
pdb/seq: 8
             name: pdbs/split_chain/3X2S_A.pdb
pdb/seq: 9
             name: pdbs/split_chain/6HAP_A.pdb
pdb/seq: 10
              name: pdbs/split_chain/6HAM_A.pdb
   PDB has ALT records, taking A only, rm.alt=TRUE
              name: pdbs/split_chain/4K46_A.pdb
pdb/seq: 11
   PDB has ALT records, taking A only, rm.alt=TRUE
```

pdbs [Truncated_Name:1]1AKE_A.pdb ----MRIILLGAPGAGKGTQAQFIMEKYGIPQIS [Truncated_Name:2]6S36_A.pdb -----MRIILLGAPGAGKGTQAQFIMEKYGIPQIS [Truncated_Name:3]6RZE_A.pdb -----MRIILLGAPGAGKGTQAQFIMEKYGIPQIS [Truncated_Name: 4] 3HPR_A.pdb -----MRIILLGAPGAGKGTQAQFIMEKYGIPQIS [Truncated_Name:5]1E4V_A.pdb -----MRIILLGAPVAGKGTQAQFIMEKYGIPQIS [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 [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

name: pdbs/split_chain/3GMT_A.pdb

name: pdbs/split_chain/4PZL_A.pdb

pdb/seq: 12

pdb/seq: 13

RIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFD

120

81

[Truncated_Name:1]1AKE_A.pdb

[Truncated_Name:2]6S36_A.pdb	RI	AQEDCR	NGFLLDO	GFPRT:	IPQAD <i>I</i>	AMKEAGI	INVDYV	LEFD
[Truncated_Name:3]6RZE_A.pdb	RI	AQEDCR	NGFLLDO	GFPRT:	IPQAD <i>I</i>	AMKEAG	INVDYV	LEFD
[Truncated_Name:4]3HPR_A.pdb	RI	AQEDCR	NGFLLDO	GFPRT:	IPQAD <i>I</i>	AMKEAG	INVDYV	LEFD
[Truncated_Name:5]1E4V_A.pdb	RI	AQEDCR	NGFLLDO	GFPRT:	IPQAD <i>A</i>	AMKEAGI	INVDYV	LEFD
[Truncated_Name:6]5EJE_A.pdb		•	NGFLLDO					
[Truncated_Name:7]1E4Y_A.pdb		•	NGFLLDO					
[Truncated_Name:8]3X2S_A.pdb		•	NGFLLDO					
[Truncated_Name:9]6HAP_A.pdb		-	NGFLLDO		-			
[Truncated_Name:10]6HAM_A.pdb		•	NGFLLDO		•			
[Truncated_Name:11]4K46_A.pdb			KGFLLDO					
[Truncated_Name:12]3GMT_A.pdb		-	NGYLFDO		-			
[Truncated_Name:13]4PZL_A.pdb			NGFLLDO					
	*^	*	*^* **				` ^**^	
	81							120
	01		•		•	•		120
	121							160
[Truncated_Name:1]1AKE_A.pdb		DELIVD	RIVGRRV	/HAPS	GRVYHV	/KFNPP	(VEGKD	DVTG
[Truncated_Name:2]6S36_A.pdb			KIVGRRV					
[Truncated_Name:3]6RZE_A.pdb			AIVGRRV					
[Truncated_Name:4]3HPR_A.pdb			RIVGRRV					
[Truncated_Name:5]1E4V_A.pdb			RIVGRRV					
[Truncated_Name:6]5EJE_A.pdb			RIVGRRV					
[Truncated_Name:7]1E4Y_A.pdb			RIVGRRV					
[Truncated_Name:8]3X2S_A.pdb			RIVGRRV					
[Truncated_Name:9]6HAP_A.pdb			RIVGRRV					
[Truncated_Name:10]6HAM_A.pdb			RIVGRRV					
[Truncated_Name:11]4K46_A.pdb			RMAGRR <i>A</i>					
[Truncated_Name: 12] 3GMT_A.pdb			RMSGRR7					
[Truncated_Name: 13] 4PZL_A.pdb			RITGRRI					
[II ansassa_name. 10] II EE_III pas	*		^ ***			^***		* **
	121							160
			·		•	·		
	161							200
[Truncated_Name:1]1AKE_A.pdb	EE	LTTRKD	DQEETVE	RKRLV	EYHQM1	CAPLIGY	YSKEA	EAGN
[Truncated_Name:2]6S36_A.pdb			DQEETVF		-			
[Truncated_Name:3]6RZE_A.pdb	EE	LTTRKD	DQEETVE	RKRLV	EYHQMT	[APLIGY	YSKEA	EAGN
[Truncated_Name:4]3HPR_A.pdb	EE	LTTRKD	DQEETVE	RKRLV	EYHQM1	CAPLIGY	YSKEA	EAGN
[Truncated_Name:5]1E4V_A.pdb	EE	LTTRKD	DQEETVE	RKRLV	EYHQM1	CAPLIGY	YSKEA	EAGN
[Truncated_Name:6]5EJE_A.pdb	EE	LTTRKD	DQEECVE	RKRLV	EYHQM1	CAPLIGY	YSKEA	EAGN
[Truncated_Name:7]1E4Y_A.pdb			DQEETVF					
[Truncated_Name:8]3X2S_A.pdb			DQEETVF					
[Truncated_Name:9]6HAP_A.pdb			DQEETVF					
			• -		٠,			

[Truncated_Name:10]6HAM_A.pdb

EELTTRKDDQEETVRKRLVEYHQMTAPLIGYYSKEAEAGN

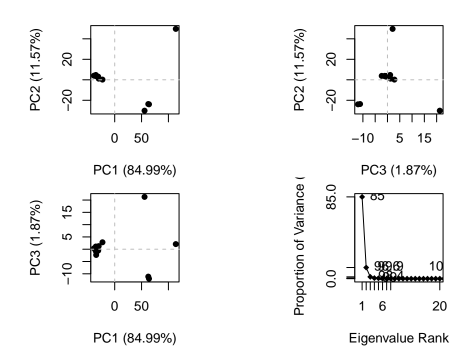
```
[Truncated_Name:11]4K46_A.pdb
                                EDLVIREDDKEETVLARLGVYHNQTAPLIAYYGKEAEAGN
[Truncated_Name:12]3GMT_A.pdb
                                EPLVQRDDDKEETVKKRLDVYEAQTKPLITYYGDWARRGA
[Truncated_Name: 13] 4PZL_A.pdb
                                EPLITRTDDNEDTVKQRLSVYHAQTAKLIDFYRNFSSTNT
                                     * ** *^ *
                                               **
                              161
                                                                        200
                              201
                                                           227
[Truncated_Name:1]1AKE_A.pdb
                                T--KYAKVDGTKPVAEVRADLEKILG-
[Truncated_Name:2]6S36_A.pdb
                                T--KYAKVDGTKPVAEVRADLEKILG-
[Truncated_Name:3]6RZE_A.pdb
                                T--KYAKVDGTKPVAEVRADLEKILG-
[Truncated_Name:4]3HPR_A.pdb
                                T--KYAKVDGTKPVAEVRADLEKILG-
[Truncated_Name:5]1E4V_A.pdb
                                T--KYAKVDGTKPVAEVRADLEKILG-
[Truncated_Name:6]5EJE_A.pdb
                                T--KYAKVDGTKPVAEVRADLEKILG-
[Truncated_Name:7]1E4Y_A.pdb
                                T--KYAKVDGTKPVAEVRADLEKILG-
[Truncated_Name:8]3X2S_A.pdb
                                T--KYAKVDGTKPVAEVRADLEKILG-
                                T--KYAKVDGTKPVCEVRADLEKILG-
[Truncated_Name:9]6HAP_A.pdb
[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)
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
  \#par(mar=c(1,1,1,1))
  #plot(pdbs, labels=ids)
```

Annotate collected PDB structures

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

Jump to PCA

```
# 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