Comparative Structure Analysis and Structure_pca Lab Class 11 Alphafold

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Here we run through a complete "pipeline" of structure analysis that begins with a single sequence identifier and ends in a PCA analysis.

library(bio3d)

Step 1. Get a sequence

Retreive a sequence for the protein we are interested in. We will take ADK "lake_A"

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

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

Fetching... Please wait. Done.

aa

	1	•	•				60		
pdb 1AKE A	MRIILLGAPGAGKGTQAQFIMEKYGIPQISTGDMLRAAVKSGSELGKQAKDIMDAGKLVT								
	1	•		•			60		
	61						120		
pdb 1AKE A	DELVIALVKERIAQEDCRNGFLLDGFPRTIPQADAMKEAGINVDYVLEFDVPDELIVDRI								
	61	•		•		•	120		

```
121
                                                                         180
pdb|1AKE|A
             VGRRVHAPSGRVYHVKFNPPKVEGKDDVTGEELTTRKDDQEETVRKRLVEYHQMTAPLIG
           121
                                                                         180
           181
                                               214
pdb|1AKE|A
             YYSKEAEAGNTKYAKVDGTKPVAEVRADLEKILG
           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
```

Step 2. Blast search

Run a BLAST search of the PDB for all related sequences to our input aa

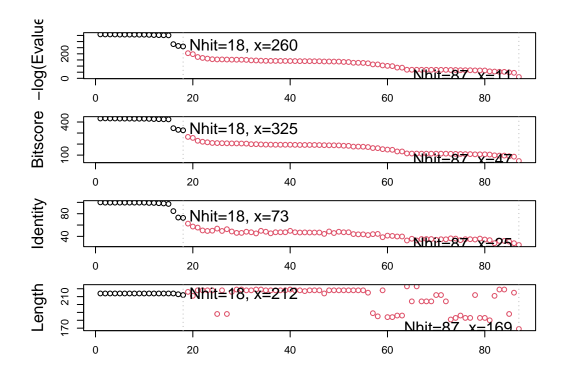
```
blast <- blast.pdb(aa)

Searching ... please wait (updates every 5 seconds) RID = V5F0C46Y013
......
Reporting 87 hits</pre>
```

hits <- plot(blast)</pre>

* Possible cutoff values: 260 11 Yielding Nhits: 18 87

* Chosen cutoff value of: 260 Yielding Nhits: 18



Step 3. Download all structure

These are our "top hits" i.e. all the structures in the PDB database related to our input sequence.

hits\$pdb.id

```
[1] "1AKE_A" "8BQF_A" "4X8M_A" "6S36_A" "8Q2B_A" "8RJ9_A" "6RZE_A" "4X8H_A" [9] "3HPR_A" "1E4V_A" "5EJE_A" "1E4Y_A" "3X2S_A" "6HAP_A" "6HAM_A" "8PVW_A" [17] "4K46_A" "4NP6_A"
```

```
# 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/8BQF.pdb.gz exists. Skipping download

Warning in get.pdb(hits\$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE): pdbs/4X8M.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/8Q2B.pdb.gz exists. Skipping download

Warning in get.pdb(hits\$pdb.id, path = "pdbs", split = TRUE, gzip = TRUE): pdbs/8RJ9.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/4X8H.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/8PVW.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/4NP6.pdb.gz exists. Skipping download

	I	0%
 ====	I	6%
 ======	1	11%
 ========	1	17%
 ===========	ı	22%
 ===================================	I	28%
 ===================================	ı	33%
 	1	39%
 ==================================	i	44%
 -==================================	· 1	50%
' 	' 1	56%
	'	61%
	1	
=====================================		67%
=====================================	1	72%
 		78%
		83%

	 ===================================		89%
	 ===================================		94%
1	 	1	100%

Step 4. Align and superpose

pdbs/split_chain/1AKE_A.pdb

Reading PDB files:

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

```
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

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/8BQF_A.pdb PDB has ALT records, taking A only, rm.alt=TRUE pdb/seq: 3 name: pdbs/split_chain/4X8M_A.pdb name: pdbs/split_chain/6S36_A.pdb pdb/seq: 4 PDB has ALT records, taking A only, rm.alt=TRUE name: pdbs/split_chain/8Q2B_A.pdb pdb/seq: 5 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 name: pdbs/split_chain/6HAM_A.pdb pdb/seq: 15 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 name: pdbs/split_chain/4K46_A.pdb pdb/seq: 17 PDB has ALT records, taking A only, rm.alt=TRUE name: pdbs/split_chain/4NP6_A.pdb pdb/seq: 18

pdbs

[Truncated_Name:5]8Q2B_A.pdb	MRIILLGAP	GAGKGTQAQFIMEK	YGIPQISTGDMLRAA
[Truncated_Name:6]8RJ9_A.pdb	MRIILLGAP	GAGKGTQAQFIMEK	YGIPQISTGDMLRAA
[Truncated_Name:7]6RZE_A.pdb	MRIILLGAP	GAGKGTQAQFIMEK	YGIPQISTGDMLRAA
[Truncated_Name:8]4X8H_A.pdb	MRIILLGAF	GAGKGTQAQFIMEK	YGIPQISTGDMLRAA
[Truncated_Name:9]3HPR_A.pdb	MRIILLGAF	GAGKGTQAQFIMEK	YGIPQISTGDMLRAA
[Truncated_Name:10]1E4V_A.pdb	MRIILLGAF	VAGKGTQAQFIMEK	YGIPQISTGDMLRAA
[Truncated_Name:11]5EJE_A.pdb	MRIILLGAP	GAGKGTQAQFIMEK	YGIPQISTGDMLRAA
[Truncated_Name:12]1E4Y_A.pdb	MRIILLGAL	VAGKGTQAQFIMEK	YGIPQISTGDMLRAA
[Truncated_Name:13]3X2S_A.pdb	MRIILLGAP	GAGKGTQAQFIMEK	YGIPQISTGDMLRAA
[Truncated_Name:14]6HAP_A.pdb	MRIILLGAP	GAGKGTQAQFIMEK	YGIPQISTGDMLRAA
[Truncated_Name:15]6HAM_A.pdb	MRIILLGAF	GAGKGTQAQFIMEK	YGIPQISTGDMLRAA
[Truncated_Name:16]8PVW_A.pdb	MRIILLGAF	GAGKGTQAQFIMEK	YGIPQISTGDMLRAA
[Truncated_Name:17]4K46_A.pdb	MRIILLGAF	GAGKGTQAQFIMAK	FGIPQISTGDMLRAA
[Truncated_Name:18]4NP6_A.pdb	NAMRIILLGAF	GAGKGTQAQFIMEK	FGIPQISTGDMLRAA
	*****	*******	^*****
	1 .		. 40
	41 .		. 80
[Truncated_Name:1]1AKE_A.pdb	VKSGSELGKQA	KDIMDAGKLVTDEL	VIALVKERIAQEDCR
[Truncated_Name:2]8BQF_A.pdb	VKSGSELGKQA	KDIMDAGKLVTDEL	VIALVKERIAQE
[Truncated_Name:3]4X8M_A.pdb	VKSGSELGKQA	KDIMDAGKLVTDEL	VIALVKERIAQEDCR
[Truncated_Name:4]6S36_A.pdb	VKSGSELGKQA	KDIMDAGKLVTDEL	VIALVKERIAQEDCR
[Truncated_Name:5]8Q2B_A.pdb	VKSGSELGKQA	KDIMDAGKLVTDEL	VIALVKERIAQEDCR
[Truncated_Name:6]8RJ9_A.pdb	VKSGSELGKQA	KDIMDAGKLVTDEL	VIALVKERIAQEDCR
[Truncated_Name:7]6RZE_A.pdb	VKSGSELGKQA	KDIMDAGKLVTDEL	VIALVKERIAQEDCR
[Truncated_Name:8]4X8H_A.pdb	VKSGSELGKQA	KDIMDAGKLVTDEL	VIALVKERIAQEDCR
[Truncated_Name:9]3HPR_A.pdb	VKSGSELGKQA	KDIMDAGKLVTDEL	VIALVKERIAQEDCR
[Truncated_Name:10]1E4V_A.pdb	VKSGSELGKQA	KDIMDAGKLVTDEL	VIALVKERIAQEDCR
[Truncated_Name:11]5EJE_A.pdb	VKSGSELGKQA	KDIMDACKLVTDEL	VIALVKERIAQEDCR
[Truncated_Name:12]1E4Y_A.pdb	VKSGSELGKQA	KDIMDAGKLVTDEL	VIALVKERIAQEDCR
[Truncated_Name:13]3X2S_A.pdb	VKSGSELGKQA	KDIMDCGKLVTDEL	VIALVKERIAQEDSR
[Truncated_Name:14]6HAP_A.pdb	VKSGSELGKQA	KDIMDAGKLVTDEL	VIALVRERICQEDSR
[Truncated_Name:15]6HAM_A.pdb	IKSGSELGKQA	KDIMDAGKLVTDEI	IIALVKERICQEDSR
[Truncated_Name:16]8PVW_A.pdb	VKSGSELGKQA	KDIMDAGKLVTDEL	VIALVKERIAQEDCR
[Truncated_Name:17]4K46_A.pdb	IKAGTELGKQA	KSVIDAGQLVSDDI	ILGLVKERIAQDDCA
[Truncated_Name:18]4NP6_A.pdb			ILGLIKERIAQADCE
	^* *^****	* ^^* **^*	^^*^***
	41 .	•	. 80
	0.4		40/

[Truncated_Name:1]1AKE_A.pdb [Truncated_Name:2]8BQF_A.pdb [Truncated_Name:3]4X8M_A.pdb NGFLLDGFPRTIPQADAMKEAGINVDYVLEFDVPDELIVD
-GFLLDGFPRTIPQADAMKEAGINVDYVIEFDVPDELIVD
NGFLLDGFPRTIPQADAMKEAGINVDYVLEFDVPDELIVD

[Truncated_Name:4]6S36_A.pdb	NGFLLDO	GFPRTIPQADAM	KEAGINVDY	/VLEFDVPDE!	LIVD
[Truncated_Name:5]8Q2B_A.pdb	NGFLLD(GFPRTIPQADAM	KEAGINVDY	/VLEFDVPDE	LIVD
[Truncated_Name:6]8RJ9_A.pdb	NGFLLA	GFPRTIPQADAM	KEAGINVDY	/VLEFDVPDE	LIVD
[Truncated_Name:7]6RZE_A.pdb	NGFLLD	GFPRTIPQADAM	KEAGINVDY	/VLEFDVPDE!	LIVD
[Truncated_Name:8]4X8H_A.pdb	NGFLLD	GFPRTIPQADAM	KEAGINVDY	/VLEFDVPDE!	LIVD
[Truncated_Name:9]3HPR_A.pdb	NGFLLD	GFPRTIPQADAM	KEAGINVDY	/VLEFDVPDE!	LIVD
[Truncated_Name:10]1E4V_A.pdb	NGFLLD	GFPRTIPQADAM	KEAGINVDY	/VLEFDVPDE!	LIVD
[Truncated_Name:11]5EJE_A.pdb	NGFLLD	GFPRTIPQADAM	KEAGINVDY	/VLEFDVPDE!	LIVD
[Truncated_Name:12]1E4Y_A.pdb		GFPRTIPQADAM			
[Truncated_Name:13]3X2S_A.pdb		GFPRTIPQADAM			
[Truncated_Name:14]6HAP_A.pdb		GFPRTIPQADAM			
[Truncated_Name:15]6HAM_A.pdb		GFPRTIPQADAM			
[Truncated_Name:16]8PVW_A.pdb		GFPRTIPQADAM			
[Truncated_Name:17]4K46_A.pdb		GFPRTIPQADGL			
[Truncated_Name:18]4NP6_A.pdb		GFPRTIPQADGL			

	81			•	120
	121			•	160
[Truncated_Name:1]1AKE_A.pdb	RIVGRRV	/HAPSGRVYHVK	FNPPKVEG	KDDVTGEELT"	TRKD
[Truncated_Name:2]8BQF_A.pdb	RIVGRRV	/HAPSGRVYHVK	FNPPKVEG	KDDVTGEELT"	TRKD
[Truncated_Name:3]4X8M_A.pdb	RIVGRRV	/HAPSGRVYHVK	FNPPKVEG	KDDVTGEELT"	TRKD
[Truncated_Name:4]6S36_A.pdb	KIVGRRV	/HAPSGRVYHVK	FNPPKVEG	KDDVTGEELT"	TRKD
[Truncated_Name:5]8Q2B_A.pdb	RIVGRRV	/HAPSGRVYHVK	FNPPKVEG	KDDVTGEELT"	TRKA
[Truncated_Name:6]8RJ9_A.pdb	RIVGRRV	/HAPSGRVYHVK	FNPPKVEG	KDDVTGEELT"	TRKD
[Truncated_Name:7]6RZE_A.pdb	AIVGRRV	/HAPSGRVYHVK	FNPPKVEG	KDDVTGEELT"	TRKD
[Truncated_Name:8]4X8H_A.pdb	RIVGRRV	/HAPSGRVYHVK	FNPPKVEG	KDDVTGEELT"	TRKD
[Truncated_Name:9]3HPR_A.pdb	RIVGRRV	/HAPSGRVYHVK	FNPPKVEG	KDDGTGEELT"	TRKD
[Truncated_Name:10]1E4V_A.pdb	RIVGRRV	/HAPSGRVYHVK	FNPPKVEG	KDDVTGEELT"	TRKD
[Truncated_Name:11]5EJE_A.pdb	RIVGRRV	/HAPSGRVYHVK	FNPPKVEG	KDDVTGEELT"	TRKD
[Truncated_Name:12]1E4Y_A.pdb	RIVGRRV	/HAPSGRVYHVK	FNPPKVEG	KDDVTGEELT"	TRKD
[Truncated_Name:13]3X2S_A.pdb	RIVGRRV	/HAPSGRVYHVK	FNPPKVEG	KDDVTGEELT"	TRKD
[Truncated_Name:14]6HAP_A.pdb	RIVGRRV	/HAPSGRVYHVK	FNPPKVEG	KDDVTGEELT"	TRKD
[Truncated_Name:15]6HAM_A.pdb	RIVGRRV	/HAPSGRVYHVK	FNPPKVEG	KDDVTGEELT'	TRKD
[Truncated_Name:16]8PVW_A.pdb		ETSGRV			
[Truncated_Name:17]4K46_A.pdb		AHLASGRTYHNV			
[Truncated_Name:18]4NP6_A.pdb		AHLPSGRTYHVV			
	^ *	***		,	
	121				160

[Truncated_Name:1]1AKE_A.pdb [Truncated_Name:2]8BQF_A.pdb 161 200
DQEETVRKRLVEYHQMTAPLIGYYSKEAEAGNTKYAKVDG
DQEETVRKRLVEYHQMTAPLIGYYSKEAEAGNTKYAKVDG

```
[Truncated_Name:3]4X8M_A.pdb
                                DQEETVRKRLVEWHQMTAPLIGYYSKEAEAGNTKYAKVDG
[Truncated_Name:4]6S36_A.pdb
                                DQEETVRKRLVEYHQMTAPLIGYYSKEAEAGNTKYAKVDG
[Truncated_Name:5]8Q2B_A.pdb
                                DQEETVRKRLVEYHQMTAPLIGYYSKEAEAGNTKYAKVDG
[Truncated_Name:6]8RJ9_A.pdb
                                DQEETVRKRLVEYHQMTAPLIGYYSKEAEAGNTKYAKVDG
[Truncated Name:7]6RZE A.pdb
                                DQEETVRKRLVEYHQMTAPLIGYYSKEAEAGNTKYAKVDG
[Truncated Name:8]4X8H A.pdb
                                DQEETVRKRLVEYHQMTAALIGYYSKEAEAGNTKYAKVDG
[Truncated Name:9]3HPR A.pdb
                                DQEETVRKRLVEYHQMTAPLIGYYSKEAEAGNTKYAKVDG
[Truncated_Name:10]1E4V_A.pdb
                                DQEETVRKRLVEYHQMTAPLIGYYSKEAEAGNTKYAKVDG
[Truncated_Name:11]5EJE_A.pdb
                                DQEECVRKRLVEYHQMTAPLIGYYSKEAEAGNTKYAKVDG
[Truncated_Name: 12] 1E4Y_A.pdb
                                DQEETVRKRLVEYHQMTAPLIGYYSKEAEAGNTKYAKVDG
[Truncated_Name:13]3X2S_A.pdb
                                DQEETVRKRLCEYHQMTAPLIGYYSKEAEAGNTKYAKVDG
[Truncated_Name:14]6HAP_A.pdb
                                DQEETVRKRLVEYHQMTAPLIGYYSKEAEAGNTKYAKVDG
[Truncated_Name: 15] 6HAM_A.pdb
                                DQEETVRKRLVEYHQMTAPLIGYYSKEAEAGNTKYAKVDG
[Truncated_Name:16]8PVW_A.pdb
                                DNEETVRKRLVEYHQMTAPLIGYYSKEAEAGNTKYAKVDG
[Truncated_Name:17]4K46_A.pdb
                                DKEETVLARLGVYHNQTAPLIAYYGKEAEAGNTQYLKFDG
[Truncated_Name:18]4NP6_A.pdb
                                DKEETVRARLNVYHTQTAPLIEYYGKEAAAGKTQYLKFDG
                                            ^* ** ** ** ** ** * * * *
                              161
                                                                        200
                              201
                                               216
[Truncated_Name:1]1AKE_A.pdb
                                TKPVAEVRADLEKILG
[Truncated Name:2]8BQF A.pdb
                                TKPVAEVRADLEKIL-
[Truncated_Name:3]4X8M_A.pdb
                                TKPVAEVRADLEKILG
[Truncated Name: 4] 6S36 A.pdb
                                TKPVAEVRADLEKILG
[Truncated_Name:5]8Q2B_A.pdb
                                TKPVAEVRADLEKILG
[Truncated_Name:6]8RJ9_A.pdb
                                TKPVAEVRADLEKILG
[Truncated_Name:7]6RZE_A.pdb
                                TKPVAEVRADLEKILG
[Truncated_Name:8]4X8H_A.pdb
                                TKPVAEVRADLEKILG
[Truncated_Name:9]3HPR_A.pdb
                                TKPVAEVRADLEKILG
[Truncated_Name:10]1E4V_A.pdb
                                TKPVAEVRADLEKILG
[Truncated_Name:11]5EJE_A.pdb
                                TKPVAEVRADLEKILG
[Truncated_Name:12]1E4Y_A.pdb
                                TKPVAEVRADLEKILG
[Truncated_Name:13]3X2S_A.pdb
                                TKPVAEVRADLEKILG
[Truncated_Name:14]6HAP_A.pdb
                                TKPVCEVRADLEKILG
[Truncated Name: 15] 6HAM A.pdb
                                TKPVCEVRADLEKILG
[Truncated_Name:16]8PVW_A.pdb
                                TKPVAEVRADLEKILG
[Truncated Name:17]4K46 A.pdb
                                TKAVAEVSAELEKALA
[Truncated_Name:18]4NP6_A.pdb
                                TKQVSEVSADIAKALA
                                ** * ** *^^ * *
                              201
                                                216
```

Call:

pdbaln(files = files, fit = TRUE, exefile = "msa")

```
Class:
  pdbs, fasta

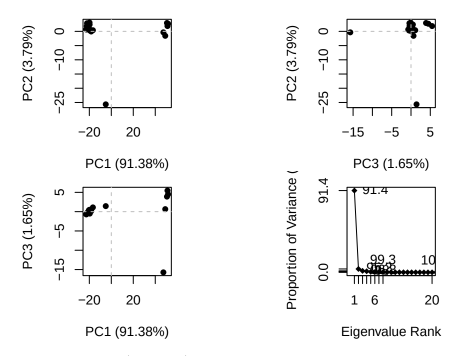
Alignment dimensions:
  18 sequence rows; 216 position columns (182 non-gap, 34 gap)
```

+ attr: xyz, resno, b, chain, id, ali, resid, sse, call

Step 5. PCA

Let's use our old friend PCA to make sense of these confusing, complicated structure relationships

```
pc <- pca(pdbs)
plot(pc)</pre>
```



Let's make a trajectory (or movie) of the main conformational changes captured by PC1. We will use the mktrj() function for this...

```
mktrj(pc, file="pca_result.pdb")
```

Back of the envelope comparison of the PDB size to UniProt

```
uniprot <- 253206171
pdb <- 231029
pdb/uniprot * 100
```

[1] 0.09124146