#### **Assignment 2**

### BISC577A, Unit 3

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ChIP-seq, which is in vivo experiment, is method that analyzes protein-DNA interactions, measuring specific biological modifications along genome such as detecting DNA-binding proteins, e.g. TF (transcription factor), binding sites. The experimental procedure is almost the same as ChIP-chip except the last step where sequencing is used to replace microarray. Unlike in vitro experiments like SELEX-seq and PBM, Chlp-seq isolate chromatin, use antibody to immunoprecipitate target factor in chromatin, and DNA bound to this factor gets then sequenced, to produce target DNA site library bound to target protein in vivo, eventually revealing which genomic regions were bound by the factor at chromatin isolation. Also, unlike ChIP-chip, the enriched DNA segments is rather sequenced directly than dependent on hybridization in tiling arrays. It doesn't require much starting material and has quite high spatial resolution and good signal to noise ratio as well as dynamic range. But, it's quite costly and needs a lot of tissue, which might be problematic for some types of samples. SELEX-seq is in vitro highthroughput technique that produces oligonucleotides of ssDNA or RNA that target ligand specifically bind to so as to determine TF-DNA sequence specificity by using classical protein-DNA SELEX assays with massive parallel sequencing. Its latent specificity could be precedent for distinguishment of similar TF specificities between in vivo and in vitro. Also, it could determine relative affinities to DNA sequence for any TF. But, the product should be purified after PCR, such as from dsDNA or other left-overs. Also, its range over readout (to be analyzed) is known to be limited, and prior sequence-based alignment of different oligomers are needed. PBM (protein binding microarray) also can characterize TF sequence specificity in a high-throughput fashion. It can provide whole comprehensive table with preference of particular TF for any sequence variant though it required prior background knowledge of the protein of interest and affinity reagents.

Two different sequence models "1-mer" and "1-mer+shape" are built as prediction models for three different in vitro data of Mad, Max, and Myc.

```
7534 samples
144 predictor
No pre-processing
Resampling: Cross-Validated (10 fold)
Summary of sample sizes: 6782, 6780, 6780, 6779, 6780, 6781, ...
Resampling results across tuning parameters:
 lambda
              RMSE
                         Rsquared
 3.051758e-05 0.3809422 0.7748270
 6.103516e-05 0.3809422 0.7748270
 1.220703e-04 0.3809422 0.7748270
  2.441406e-04 0.3809422 0.7748270
 4.882812e-04 0.3809422 0.7748270
  9.765625e-04 0.3809422 0.7748270
  1.953125e-03 0.3809422 0.7748270
  3.906250e-03 0.3809422 0.7748270
  7.812500e-03 0.3809422 0.7748270
 1.562500e-02 0.3809422 0.7748270
 3.125000e-02 0.3811609 0.7747944
 6.250000e-02 0.3842917 0.7743338
 1.250000e-01 0.3946565 0.7727535
 2.500000e-01 0.4222592 0.7679309
 5.000000e-01 0.4771609 0.7555835
 1.000000e+00 0.5544751 0.7312023
 2.000000e+00 0.6344850 0.6950211
 4.000000e+00 0.6991414 0.6551491
 8.000000e+00 0.7432415 0.6208768
 1.600000e+01 0.7698562 0.5968907
 3.200000e+01 0.7846970 0.5823424
 6.400000e+01 0.7925893 0.5742313
 1.280000e+02 0.7966512 0.5699719
 2.560000e+02 0.7988001 0.5677692
 5.120000e+02 0.8008122 NaN
                              NaN
 1.024000e+03 0.8008122
                             NaN
 2.048000e+03 0.8008122
 4.096000e+03 0.8008122
                              NaN
  8.192000e+03 0.8008122
                              NaN
 1.638400e+04 0.8008122
                              NaN
 3.276800e+04 0.8008122
                               NaN
Tuning parameter 'alpha' was held constant at a value of 0
RMSE was used to select the optimal model using the smallest value.
The final values used for the model were alpha = 0 and lambda = 0.015625.
> average_Rsquared5 <- mean(na.omit(model5$results$Rsquared))</pre>
> head(average_Rsquared5)
Γ17 0.71613
```

Figure 1. L2-regularized MLR model for "1-mer" on Mad

```
7534 samples
274 predictor
No pre-processing
Resampling: Cross-Validated (10 fold)
Summary of sample sizes: 6780, 6781, 6781, 6779, 6782, 6782, ...
Resampling results across tuning parameters:
             RMSE
 lambda
                          Rsquared
 3.051758e-05 0.3014281 0.8634341
  6.103516e-05 0.3014281 0.8634341
 1.220703e-04 0.3014281 0.8634341
 2.441406e-04 0.3014281 0.8634341
 4.882812e-04 0.3014281 0.8634341
 9.765625e-04 0.3014281 0.8634341
 1.953125e-03 0.3014281 0.8634341
 3.906250e-03 0.3014281 0.8634341
 7.812500e-03 0.3014281 0.8634341
 1.562500e-02 0.3014281 0.8634341
 3.125000e-02 0.3014281 0.8634341
 6.250000e-02 0.3161308 0.8499889
 1.250000e-01 0.3507648 0.8171492
  2.500000e-01 0.3837904 0.7871111
  5.000000e-01 0.4209726 0.7595674
  1.000000e+00 0.4715008 0.7293100
  2.000000e+00 0.5382431 0.6882220
  4.000000e+00 0.6111136 0.6317770
  8.000000e+00 0.6760116 0.5670342
  1.600000e+01 0.7249594 0.5099457
  3.200000e+01 0.7578188 0.4692471
  6.400000e+01 0.7776414 0.4448170
 1.280000e+02 0.7888304 0.4311913
 2.560000e+02 0.7947790 0.4240596
 5.120000e+02 0.8010289
 1.024000e+03 0.8010289
 2.048000e+03 0.8010289
 4.096000e+03 0.8010289
                             NaN
 8.192000e+03 0.8010289
                             NaN
 1.638400e+04 0.8010289
                             NaN
 3.276800e+04 0.8010289
                             NaN
Tuning parameter 'alpha' was held constant at a value of 0
RMSE was used to select the optimal model using the smallest value.
The final values used for the model were alpha = 0 and lambda = 0.03125.
> average_Rsquared6 <- mean(na.omit(model6$results$Rsquared))</pre>
> head(average_Rsquared6)
[1] 0.7336331
```

Figure 2. L2-regularized MLR model for "1-mer+shape" on Mad

```
8568 samples
144 predictor
No pre-processing
Resampling: Cross-Validated (10 fold)
Summary of sample sizes: 7712, 7711, 7711, 7711, 7712, 7712, ...
Resampling results across tuning parameters:
 lambda
              RMSE
                          Rsquared
 3.051758e-05 0.09942387 0.7858999
 6.103516e-05 0.09942387 0.7858999
 1.220703e-04 0.09942387 0.7858999
 2.441406e-04 0.09942387 0.7858999
 4.882812e-04 0.09942387 0.7858999
 9.765625e-04 0.09942387 0.7858999
 1.953125e-03 0.09942387 0.7858999
 3.906250e-03 0.09942387 0.7858999
 7.812500e-03 0.09943377 0.7858939
 1.562500e-02 0.10015205 0.7854702
 3.125000e-02 0.10255425 0.7840143
 6.250000e-02 0.10915894 0.7794362
 1.250000e-01 0.12290319 0.7669988
 2.500000e-01 0.14323516 0.7399677
 5.000000e-01 0.16517483 0.6947949
 1.000000e+00 0.18351620 0.6392346
 2.000000e+00 0.19645176 0.5877872
 4.000000e+00 0.20450912 0.5505702
 8.000000e+00 0.20913087 0.5276691
 1.600000e+01 0.21163308 0.5148803
 3.200000e+01 0.21293695 0.5081473
 6.400000e+01 0.21360578 0.5046590
 1.280000e+02 0.21428316
 2.560000e+02 0.21428316
                               NaN
 5.120000e+02 0.21428316
                               NaN
 1.024000e+03 0.21428316
                               NaN
 2.048000e+03 0.21428316
                               NaN
 4.096000e+03 0.21428316
                               NaN
 8.192000e+03 0.21428316
                               NaN
 1.638400e+04 0.21428316
                              NaN
 3.276800e+04 0.21428316
                               NaN
Tuning parameter 'alpha' was held constant at a value of 0
RMSE was used to select the optimal model using the smallest value.
The final values used for the model were alpha = 0 and lambda = 0.00390625.
> average_Rsquared3 <- mean(na.omit(model3$results$Rsquared))</pre>
> head(average_Rsquared3)
[1] 0.7025783
```

Figure 3. L2-regularized MLR model for "1-mer" on Max

```
8568 samples
274 predictor
No pre-processing
Resampling: Cross-Validated (10 fold)
Summary of sample sizes: 7712, 7710, 7710, 7712, 7712, 7711, ...
Resampling results across tuning parameters:
             RMSE
 lambda
                          Rsquared
 3.051758e-05 0.07983256 0.8641997
  6.103516e-05 0.07983256 0.8641997
 1.220703e-04 0.07983256 0.8641997
  2.441406e-04 0.07983256 0.8641997
 4.882812e-04 0.07983256 0.8641997
 9.765625e-04 0.07983256 0.8641997
 1.953125e-03 0.07983256 0.8641997
 3.906250e-03 0.07983256 0.8641997
 7.812500e-03 0.07983256 0.8641997
 1.562500e-02 0.08293992 0.8539958
 3.125000e-02 0.08989418 0.8312003
 6.250000e-02 0.09770819 0.8074273
 1.250000e-01 0.10777506 0.7813824
  2.500000e-01 0.12195142 0.7483146
 5.000000e-01 0.14044348 0.6999057
  1.000000e+00 0.16037194 0.6323146
  2.000000e+00 0.17804236 0.5566054
  4.000000e+00 0.19170558 0.4908092
  8.000000e+00 0.20111950 0.4454490
  1.600000e+01 0.20705466 0.4181150
  3.200000e+01 0.21046769 0.4032364
  6.400000e+01 0.21232627 0.3953695
 1.280000e+02 0.21429740 NaN
 2.560000e+02 0.21429740
                              NaN
 5.120000e+02 0.21429740
                              NaN
 1.024000e+03 0.21429740
                              NaN
 2.048000e+03 0.21429740
                             NaN
 4.096000e+03 0.21429740
                             NaN
 8.192000e+03 0.21429740
                              NaN
 1.638400e+04 0.21429740
                             NaN
 3.276800e+04 0.21429740
                              NaN
Tuning parameter 'alpha' was held constant at a value of 0
RMSE was used to select the optimal model using the smallest value.
The final values used for the model were alpha = 0 and lambda = 0.0078125.
> average_Rsquared4 <- mean(na.omit(model4$results$Rsquared))</pre>
> head(average_Rsquared4)
[1] 0.7200874
```

Figure 4. L2-regularized MLR model for "1-mer+shape" on Max

```
6926 samples
144 predictor
No pre-processing
Resampling: Cross-Validated (10 fold)
Summary of sample sizes: 6234, 6234, 6234, 6234, 6233, 6232, ...
Resampling results across tuning parameters:
  lambda
               RMSE
                          Rsquared
  3.051758e-05 0.3741327 0.7782157
  6.103516e-05 0.3741327 0.7782157
  1.220703e-04 0.3741327 0.7782157
  2.441406e-04 0.3741327 0.7782157
  4.882812e-04 0.3741327 0.7782157
  9.765625e-04 0.3741327 0.7782157
  1.953125e-03 0.3741327 0.7782157
  3.906250e-03 0.3741327 0.7782157
  7.812500e-03 0.3741327 0.7782157
  1.562500e-02 0.3741327 0.7782157
  3.125000e-02 0.3744965 0.7781523
  6.250000e-02 0.3775201 0.7776340
 1.250000e-01 0.3875459 0.7758580
  2.500000e-01 0.4143209 0.7704810
  5.000000e-01 0.4676630 0.7570520
  1.000000e+00 0.5434125 0.7315101
  2.000000e+00 0.6228073 0.6956054
  4.000000e+00 0.6879572 0.6582451
  8.000000e+00 0.7330583 0.6273191
 1.600000e+01 0.7604372 0.6061893
  3.200000e+01 0.7758067 0.5934025
  6.400000e+01 0.7839728 0.5863221
 1.280000e+02 0.7881927 0.5825823
  2.560000e+02 0.7924668 0.5340876
  5.120000e+02 0.7925094
                               NaN
 1.024000e+03 0.7925094
                               NaN
  2.048000e+03 0.7925094
                               NaN
  4.096000e+03 0.7925094
                               NaN
  8.192000e+03 0.7925094
                               NaN
  1.638400e+04 0.7925094
                               NaN
  3.276800e+04 0.7925094
                                NaN
Tuning parameter 'alpha' was held constant at a value of 0
RMSE was used to select the optimal model using the smallest value.
The final values used for the model were alpha = 0 and lambda = 0.015625.
> average_Rsquared <- mean(na.omit(model2$results$Rsquared))</pre>
> head(average_Rsquared)
Γ17 0.7190249
```

Figure 5. L2-regularized MLR model for "1-mer" on Myc

```
No pre-processing
Resampling: Cross-Validated (10 fold)
Summary of sample sizes: 6233, 6234, 6234, 6234, 6234, 6232, ...
Resampling results across tuning parameters:
             RMSE
 lambda
                         Rsquared
 3.051758e-05 0.3044091 0.8551475
  6.103516e-05 0.3044091 0.8551475
 1.220703e-04 0.3044091 0.8551475
  2.441406e-04 0.3044091 0.8551475
 4.882812e-04 0.3044091 0.8551475
 9.765625e-04 0.3044091 0.8551475
 1.953125e-03 0.3044091 0.8551475
 3.906250e-03 0.3044091 0.8551475
 7.812500e-03 0.3044091 0.8551475
 1.562500e-02 0.3044091 0.8551475
 3.125000e-02 0.3044091 0.8551475
 6.250000e-02 0.3237244 0.8372757
 1.250000e-01 0.3475603 0.8157906
  2.500000e-01 0.3750757 0.7942302
 5.000000e-01 0.4129851 0.7705247
  1.000000e+00 0.4686106 0.7390194
  2.000000e+00 0.5398607 0.6931013
  4.000000e+00 0.6138105 0.6317843
  8.000000e+00 0.6768448 0.5657804
  1.600000e+01 0.7231997 0.5094154
  3.200000e+01 0.7535558 0.4698781
  6.400000e+01 0.7716196 0.4459729
 1.280000e+02 0.7816803 0.4326430
 2.560000e+02 0.7869946 0.4256434
 5.120000e+02 0.7925509
 1.024000e+03 0.7925509
                              NaN
 2.048000e+03 0.7925509
                             NaN
 4.096000e+03 0.7925509
                             NaN
 8.192000e+03 0.7925509
 1.638400e+04 0.7925509
                             NaN
 3.276800e+04 0.7925509
                              NaN
Tuning parameter 'alpha' was held constant at a value of 0
RMSE was used to select the optimal model using the smallest value.
The final values used for the model were alpha = 0 and lambda = 0.03125.
> average_Rsquared <- mean(na.omit(model2$results$Rsquared))</pre>
> head(average_Rsquared)
[1] 0.7307367
```

6926 samples 274 predictor

Figure 6. L2-regularized MLR model for "1-mer+shape" on Myc

# Comparison of Two Models on Mad

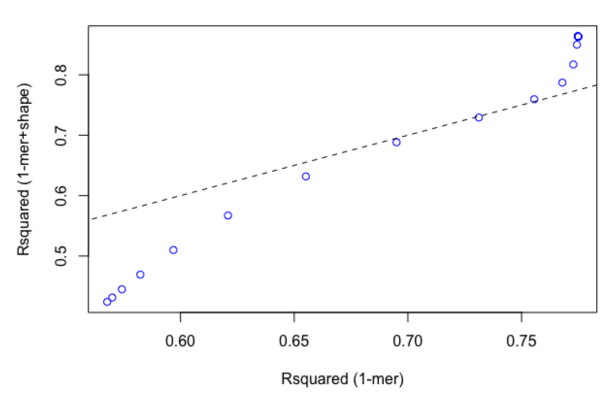


Figure 7. Comparison "1-mer" vs. "1-mer+shape" on Mad

## Comparison of Two Models on Max

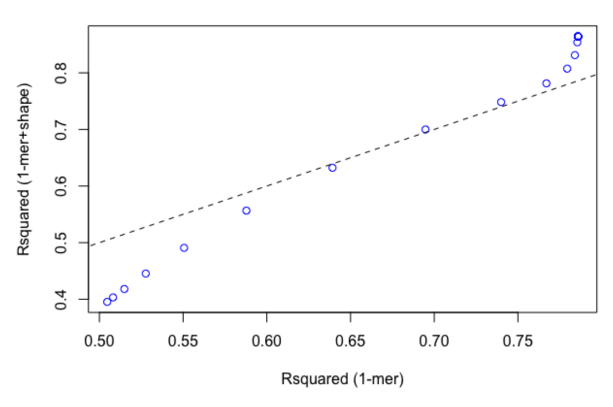


Figure 8. Comparison "1-mer" vs. "1-mer+shape" on Max

# Comparison of Two Models on Myc

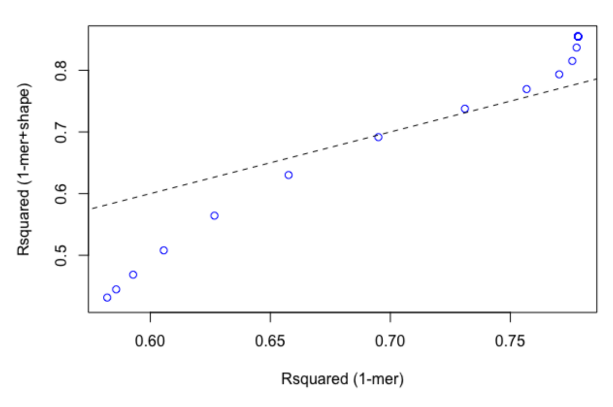


Figure 9. Comparison "1-mer" vs. "1-mer+shape" on Myc

### Comparison of Two Models on Mad, Max, Myc

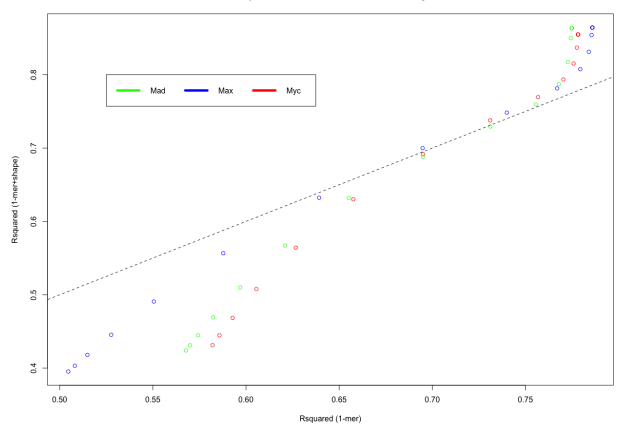


Figure 10. Comparison "1-mer" vs. "1-mer+shape" on Mad, Max, Myc

To see if the shape-augmented model outperforms significantly enough, we compute p-value with null hypothesis that two models perform the same (no outperformance). Note that R-squared is measure of how close the data are to fitted regression, i.e. how well the variation in response is explained by the fitted model. The improvement looks quite modest here, e.g. 0.716 vs. 0.733 of average R-squared between 1-mer and 1-mer+shape on Mad, but we want to know if this is statistically significant. One idea could be utilizing one-tail or two-tail t-test. For example, on data set Mad, we can set up the null hypothesis that there is no difference between two R-squared means of 1-mer and 1-mer+shape, and run t-test on list of generated R-squared of 1-mer and 1-mer+shape since if there's no difference between two average R-squared, then neither model outperforms the other. Then, we can obtain, in this case, p-value = 0.6475, which is way greater than 0.05. So, we can accept the null, i.e., there's not much statistically significant difference between two. This can be also confirmed by running one-tail test on 1-mer+shape with fixed mean = 0.716, which will yield p-value=0.6122. Hence, again, we can't reject the null hypothesis that average R-squared of 1-mer+shape is statistically significantly greater than average of R-squared of 1-mer. The results are similar on the remaining data sets Max and Myc. Even running t-test on combined data sets, i.e., three R-squared means from each data for 1-mer and the same for 1-mer+shape, yielded the similar result.

But, another approach by using Fisher's exact test yielded different result. In general, if we have contingency table, say 2 X 2 matrix with rows = student A, B and columns = Pass MATH577, fail MATH577, the null hypothesis states that the proportions/ratios of passing MATH577 are not different for two students. In our case, if we construct contingency table with rows = 1-mer, 1-mer+shape, and columns = averaged R-squared of Mad, Max, and Myc, the null hypothesis states that the proportions of mean R-squared on Mad (or Max, Myc) are not different between 1-mer and 1-mer+shape. Since the table should be integer-valued for computation, Fisher's test can be run by updating values multiplied by power of 10. The resulting p-value was 0.0005, so null hypothesis can be rejected, i.e. there's significant difference in R-squared on Mad/Max/Myc between two models. (But, this approach might not be entirely correct because unlike t-test, proportion of R-squared on one data over the all doesn't necessarily represent the performance)

Or, though not actually tested here, but like Zhou et al's PNAS paper, we can assess Spearman's rank correlation, take the differences in this values between models to get null distribution of rank correlation differences, then calculate empirical p-values.

Recently, it'd be important to note that the first approach would probably not be entirely accurate, either because we're not sure if the data is normally distributed (and it doesn't look like normal anyway). Hence, the alternative way might be Wilcox's rank sum test. With similar null and alternative hypothesis, we can run the test and get p-value = 0.0505 on, for instance, on data Mad. We can also run the same test on all data set including Mad, Max, and Myc and obtain p-value = 0.05.

```
> t.test(na.omit(model5$results$Rsquared),na.omit(model6$results$Rsquared), var.equal=TRUE, paired=FALSE)
        Two Sample t-test
data: na.omit(model5$results$Rsquared) and na.omit(model6$results$Rsquared)
t = -0.46033, df = 46, p-value = 0.6475
alternative hypothesis: true difference in means is not equal to {\bf 0}
95 percent confidence interval:
-0.09403996 0.05903378
sample estimates:
mean of x mean of y
0.7161300 0.7336331
> t.test(na.omit(model6$results$Rsquared),mu=0.7161300)
       One Sample t-test
data: na.omit(model6$results$Rsquared)
t = 0.51394, df = 23, p-value = 0.6122
alternative hypothesis: true mean is not equal to 0.71613
95 percent confidence interval:
0.6631813 0.8040850
sample estimates:
mean of x
0.7336331
> t.test(rsq[1,],rsq[2,], var.equal=TRUE, paired=FALSE)
       Two Sample t-test
data: rsq[1, ] and rsq[2, ]
t = -2.3846, df = 4, p-value = 0.07561
alternative hypothesis: true difference in means is not equal to 0
95 percent confidence interval:
-0.033708500 0.002559166
sample estimates:
mean of x mean of y
0.7125777 0.7281523
> t.test (rsq[2,], mu=0.7125777)
       One Sample t-test
data: rsq[2, ]
t = 3.7817, df = 2, p-value = 0.06335
alternative hypothesis: true mean is not equal to 0.7125777
95 percent confidence interval:
0.7104322 0.7458724
sample estimates:
mean of x
0.7281523
> r_contingency
                  [,2]
         [,1]
[1,] 0.716130 0.702578 0.719025
[2,] 0.733633 0.720087 0.730737
> r_fisher <- fisher.test(1000000*r_contingency, simulate.p.value=TRUE)</pre>
> r_fisher$p.value
```

[1] 0.0004997501

> wilcox.exact(na.omit(model6\$results\$Rsquared),na.omit(model5\$results\$Rsquared), p
aired=FALSE, alternative="greater",exact = TRUE)

Exact Wilcoxon rank sum test

data: na.omit(model6\$results\$Rsquared) and na.omit(model5\$results\$Rsquared) W = 367, p-value = 0.05056

alternative hypothesis: true mu is greater than 0

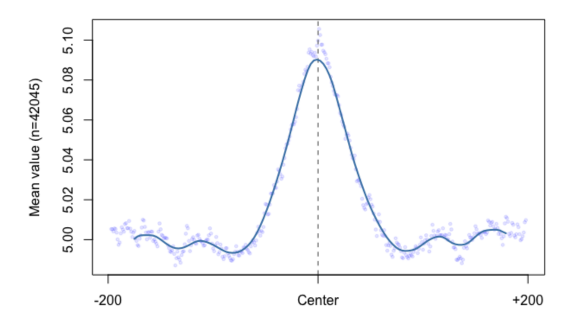
> wilcox.exact(rsq[2,],rsq[1,], paired=FALSE, alternative="greater",exact = TRUE)

Exact Wilcoxon rank sum test

data: rsq[2, ] and rsq[1, ] W = 9, p-value = 0.05

alternative hypothesis: true mu is greater than 0

CTCF transcription factor is actually transcriptional repressor, which is known as 11-zinc finger protein. It's also often referred as CCCTC-binding factor because it binds as transcription factor to repeats of the core sequence CCCTC. This CFCT binds to the consensus sequence that is defined by its zinc finger motifs. This zinc motif in them typically consists of alpha helix pair linked by zinc coordination. One helix binds in major groove at each half-site, while the second helix helps maintaining the overall structure. And, even though some local distortions to roll or propeller twist could be observed, DNA itself retains B-DNA form with mean helical twist of ~34°, usually quite low propeller twist in range between -16° and -20° in average, as well as deep, narrow minor groove. Indeed, average 5Å MGW is quite close to 5.2 Å of 1bna, recalling the previous assignment. Even if the mean propeller twist is not really within the typical bound, if we only consider dinucleotide values for helical parameter, for example,  $\approx$ -8° for CC, it seems to make sense (Or, some twist might have happened when binding). Helical twist is very close to that of typical B-DNA.



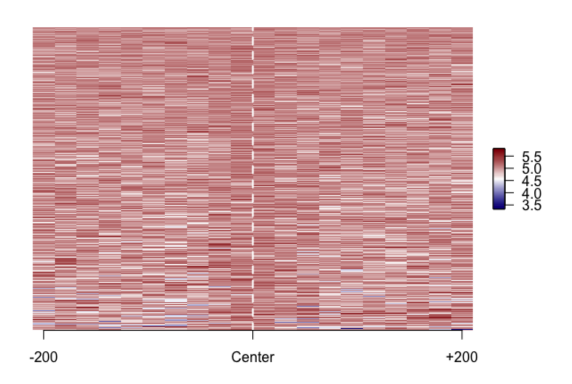
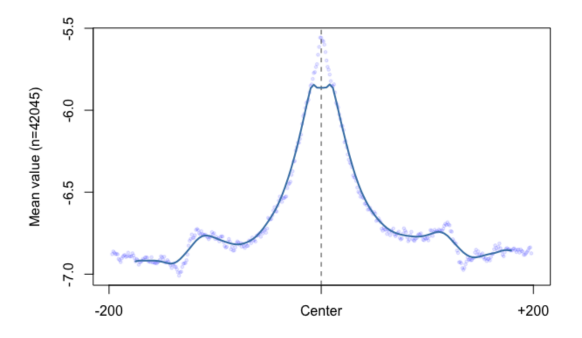


Figure 11. Ensemble plot for minor groove width (MGW)



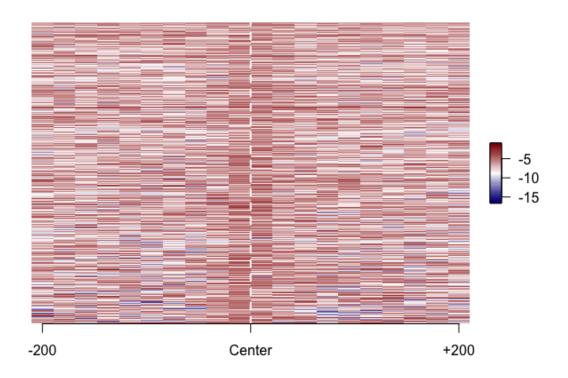


Figure 12. Ensemble plot for propeller twist (ProT)

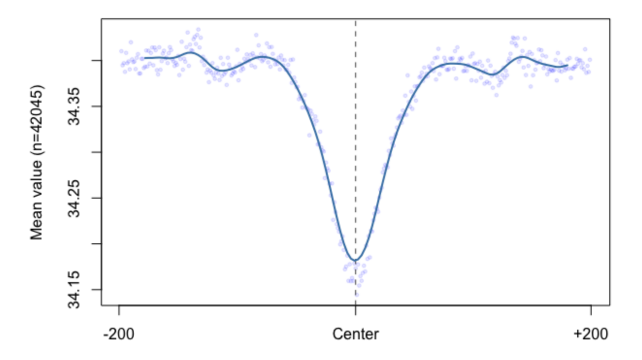


Figure 13. Ensemble plot for helical twist (HeIT)

Two different logistic models for "1-mer" and "1-mer+shape" are built as prediction models for three different in vivo ChIP-seq data of CTCF TF of Mus musculus. Comparison can be done via receiver operating characteristic curve (ROC), which is a plot of true positive rate against the false positive rate with different possible cutpoints. The closer the ROC curve follows left borderline and then top border, the more accuracy, i.e., how well the test separates the tested groups into two, we attain. On the other hand, diagonal-like curve will imply less accuracy. This accuracy can be measured by area under ROC curve (AUC) where area 1 represents indeed a perfect test whereas area 0.5 represents meaningless. AS we have 0.827 for "1-mer" and 0.813 for "1-mer+shape", the models are considered to be "good"/"accurate."

Deviance Residuals:						
Min	1Q	Median	3	Q Mo	ХX	
-2.9476	-0.6794	0.0243	0.736	8 3.48	15	
Coefficients: (29 not defined because of singularities)						
	Esti	mate St	d. Error	z value	Pr(> z )	
(Interce	ot) -15.56	6068 27	0.260537	-0.058	0.954070	
X1	8.15	9449 27	0.261038		0.975915	
X2	8.32	4427 27	0.261047	0.031	0.975428	
X3	8.21	5030 27	0.261059	0.030	0.975751	
X4	7.97	8090 27	0.261021	0.030	0.976450	
X5	0.07	7150	0.171087	0.451	0.652033	
X6	0.29	9716	0.176777	1.695	0.089991	
X7	0.32	4966	0.178338	1.822	0.068426	
X8		NA	NA		NA	
X9	0.02		0.174384		0.897901	
X10	0.61		0.177934		0.000532	
X11	0.40		0.175001		0.021642	*
X12		NA	NA		NA	
X13	-0.27		0.172028		0.115671	
X14	0.51		0.175926		0.003550	
X15	0.33		0.176926		0.056661	
X16		NA	NA		NA	
X17	0.00		0.176162		0.994843	
X18	0.48		0.177939		0.006324	**
X19	0.80		0.181178		9.95e-06	***
X20		NA	NA		NA	
X21	0.17				0.326428	
X22	0.41		0.172061		0.016911	
X23	0.51		0.179185		0.003963	**
X24		NA	NA		NA	
X25	0.00		0.172591		0.991593	
X26	0.60		0.179119		0.000664	***
X27	0.12		0.180620		0.471896	
X28	0.03	NA	NA 0 477020		NA O OFFI	
X29	0.03		0.177020		0.853724	**
X30	0.51		0.176469		0.003422	***
X31	0.77		0.177214		1.23e-05	***
X32	0.43	NA	NA 0 173010		NA 0.012000	
X33	-0.42		0.172019		0.013099	*
X34	0.22		0.177160		0.196232	*
X35	0.42		0.174006		0.015630	~
X36	0.17	NA =110	NA 0 177751		NA 0 222169	
X37	0.17		0.177251		0.323168	***
X38	0.61		0.183990		0.000840	***
X39	0.69		0.176304		8.77e-05	***
X40		NA	NA	NA.	NA	

Doviance Posiduals:

```
X41
        0.143905   0.177280   0.812   0.416941
X42
        0.763439
              0.179883
                     4.244 2.19e-05 ***
        0.510925
              0.173215
                     2.950 0.003181 **
X43
        NA
               NA
                      NA NA
X44
X45
       -0.063388
              0.176826 -0.358 0.719989
X46
        0.427036
               0.172069
                     2.482 0.013073 *
X47
        0.603167
               0.173456
                     3.477 0.000506 ***
        NA
              NA
X48
                      NA NA
X49
        0.107923 0.172627
                     0.625 0.531852
              0.177900
                     3.263 0.001100 **
X50
        0.580576
X51
        0.524668
              0.177986
                     2.948 0.003200 **
X52
        NΔ
              NA
                     NA NA
X53
       X54
       4.123 3.75e-05 ***
X55
       0.736248 0.178589
                     NA NA
        NA
              NA
       -0.455561 0.179616 -2.536 0.011203 *
X58
       X59
       0.871557   0.182687   4.771   1.84e-06 ***
        NA
              NA
                     NA NA
X60
X61
       X62
X63
        X64
        NA
              NA NA NA
X65
       0.194903 0.176912 1.102 0.270596
                     3.788 0.000152 ***
X66
        0.688738 0.181809
X67
        0.741696
              0.182024
                     4.075 4.61e-05 ***
              NA
                     NA NA
X68
        NA
X69
        0.089602
              0.173799
                     0.516 0.606170
X70
        0.511750
              0.173679
                     2.947 0.003214 **
X71
        0.643368
               0.175808
                     3.659 0.000253 ***
X72
        NA
              NA
                     NA NA
        0.033207 0.170084 0.195 0.845206
X73
       0.448757 0.176714
                     2.539 0.011103 *
X74
        0.238920 0.181575
                     1.316 0.188235
X75
              NA
        NA
X76
                     NA NA
       0.318275   0.173792   1.831   0.067048   .
X77
X78
       0.304644 0.178536 1.706 0.087945 .
X79
        X80
        NA NA NA NA
       NA
              NA
                     NA NA
X85
       -0.190459 0.169043 -1.127 0.259873
       X86
        0.399426   0.181048   2.206   0.027371 *
X87
```

```
NA
                                   NA
         0.500252   0.176687   2.831   0.004636 **
X90
         0.877815   0.177609   4.942   7.72e-07 ***
X91
X92
          NA NA
                           NA
                                   NA
         0.110890 0.173606 0.639 0.522989
X93
X94
         X95
         0.560911 0.180685 3.104 0.001907 **
                         NA
X96
          NA NA
                                 NA
X97
         0.059087 0.169690 0.348 0.727685
X98
         0.362830 0.176932 2.051 0.040299 *
X99
         0.429126 0.176359 2.433 0.014964 *
X100
            NA
                   NA
                          NA
                                   NA
X101
         -0.268958
                 0.172029 -1.563 0.117949
X102
          0.003304
                 0.181251
                         0.018 0.985457
X103
         0.421132
                 0.181142
                         2.325 0.020079 *
X104
            NA
                   NA
                           NA
                                  NA
X105
         -0.051988 0.170083 -0.306 0.759862
         X106
         X107
X108
           NA
                  NA
                          NA NA
         0.116783 0.173790 0.672 0.501598
X109
        0.290683 0.179422 1.620 0.105209
X110
X111
        0.406056 0.181603 2.236 0.025355 *
X112
           NA
                   NA
                          NA
                                   NA
        0.071766 0.171743 0.418 0.676043
X113
        0.547531 0.183024 2.992 0.002775 **
X114
        0.475032 0.179792 2.642 0.008239 **
X115
             NA
                     NA
X117
         0.418484 0.168056 2.490 0.012769 *
          0.555931   0.178077   3.122   0.001797 **
X118
          X119
X120
              NA
                      NA
                           NA
                                   NA
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' '1
(Dispersion parameter for binomial family taken to be 1)
   Null deviance: 2772.6 on 1999 degrees of freedom
Residual deviance: 1767.1 on 1908 degrees of freedom
AIC: 1951.1
Number of Fisher Scoring iterations: 14
```

Figure 14. Logistic regression model on bound and unbound data (1-mer)

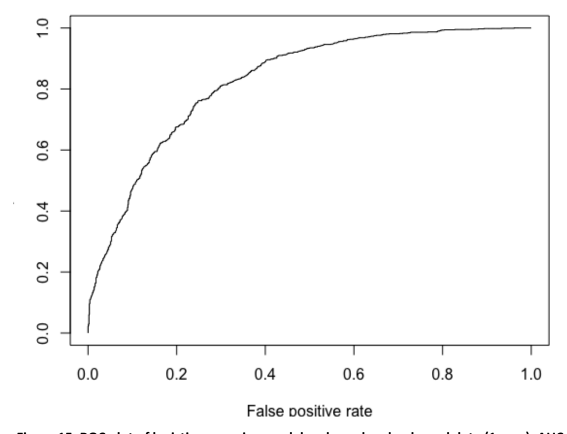


Figure 15. ROC plot of logistic regression model on bound and unbound data (1-mer). AUC = 0.827536

```
Deviance Residuals:
  Min 1Q Median
                       30
                                  Max
-2.9728 -0.5996 0.0183 0.6266 3.8727
Coefficients: (29 not defined because of singularities)
           Estimate Std. Error z value Pr(>|z|)
(Intercept) 1.266e+02 4.464e+02 0.284 0.776729
X1
          -1.726e+02 4.467e+02 -0.386 0.699167
X2
          -1.727e+02 4.467e+02 -0.387 0.699118
Х3
          -1.727e+02 4.467e+02 -0.387 0.699115
X4
          -1.730e+02 4.467e+02 -0.387 0.698531
                              1.152 0.249190
X5
           4.446e-01 3.859e-01
                              0.814 0.415360
           2.499e-01 3.068e-01
Х6
           3.023e-01 2.644e-01
                              1.143 0.252858
X7
X8
               NA
                     NA
                               NA NA
                              1.672 0.094510 .
χ9
           8.235e-01 4.925e-01
X10
           1.466e+00 5.210e-01
                              2.813 0.004904 **
X11
          2.010e-01 5.272e-01 0.381 0.702980
X12
               NA
                         NA
                              NA
          -5.002e-01 4.986e-01 -1.003 0.315731
X14
          -5.318e-01 5.183e-01 -1.026 0.304845
X15
          -2.686e-01 5.110e-01 -0.526 0.599099
X16
               NA
                         NA
                              NA
X17
          -5.371e-01 5.176e-01 -1.038 0.299397
          -7.005e-02 5.378e-01 -0.130 0.896374
X18
          -1.600e-01 5.161e-01 -0.310 0.756514
X19
X20
                NA
                         NA
                               NA
                              0.583 0.560102
X21
           3.064e-01 5.258e-01
X22
          -1.094e-01 5.275e-01 -0.207 0.835743
X23
           8.676e-01 5.384e-01
                              1.611 0.107081
X24
                         NA
                               NA
                NA
X25
           4.545e-01 5.426e-01
                              0.838 0.402254
X26
           1.177e+00 5.611e-01
                               2.098 0.035928 *
           5.683e-01 5.308e-01
X27
                              1.071 0.284368
               NA
                     NA
X28
                               NA NA
X29
          -5.251e-01 4.911e-01 -1.069 0.284963
           3.492e-02 5.116e-01
X30
                              0.068 0.945582
           3.979e-01 4.938e-01 0.806 0.420357
X31
X32
               NA
                     NA
                              NA
X33
          -6.963e-01 5.193e-01 -1.341 0.179934
           2.051e-01 5.368e-01 0.382 0.702419
           1.713e-01 5.173e-01
                              0.331 0.740629
                NA
                        NA
                              NA
X37
           7.768e-01 5.451e-01
                              1.425 0.154108
X38
           4.929e-01 5.476e-01 0.900 0.368052
           7.552e-01 5.278e-01 1.431 0.152515
X39
X40
                 NA
                         NA
                              NA NA
```

```
X41
        -3.824e-01 5.214e-01 -0.733 0.463356
X42
         6.242e-01 5.616e-01 1.111 0.266371
         -5.193e-01 5.313e-01 -0.977 0.328332
X43
           NA NA
X44
                            NA NA
                            2.812 0.004930 **
          1.449e+00 5.155e-01
X45
                            0.186 0.852317
X46
          9.992e-02 5.367e-01
X47
          6.200e-01 5.223e-01
                            1.187 0.235245
X48
           NA NA
                            NA NA
         -3.656e-01 5.330e-01 -0.686 0.492776
X49
X50
         -1.981e-01 5.499e-01 -0.360 0.718704
X51
          4.166e-01 5.415e-01
                            0.769 0.441658
X52
           NA
                   NA
                            NA NA
X53
         -3.844e-01 5.175e-01 -0.743 0.457594
                            1.468 0.142094
X54
          7.991e-01 5.443e-01
X55
         -1.113e-01 5.254e-01 -0.212 0.832161
          NA
X56
                   NA
                            NA NA
          2.383e-01 5.134e-01 0.464 0.642503
X57
X58
          1.409e-01 5.414e-01 0.260 0.794642
X59
          6.142e-01 5.093e-01
                            1.206 0.227829
X60
          NA NA
                            NA NA
X61
         -4.188e-03 5.221e-01 -0.008 0.993600
          1.500e-01 5.310e-01 0.282 0.777595
X63
          5.790e-03 5.228e-01 0.011 0.991162
X64
          NA NA NA NA
X65
         -5.231e-01 5.248e-01 -0.997 0.318904
X66
         -4.069e-02 5.550e-01 -0.073 0.941552
X67
          1.729e-01 5.275e-01 0.328 0.743028
X68
          NA NA NA NA
X69
         -1.075e+00 5.155e-01 -2.086 0.036954 *
X70
         -6.830e-01 5.467e-01 -1.249 0.211549
X71
          1.435e-02 5.138e-01 0.028 0.977724
                            NA NA
X72
          NA NA
X73
          3.362e-01 5.115e-01
                            0.657 0.510980
X74
          5.766e-01 5.099e-01
                            1.131 0.258169
X75
          7.637e-01 5.228e-01
                            1.461 0.144095
X76
          NA NA
                             NA NA
          1.350e+00 5.163e-01 2.615 0.008926 **
X77
                            2.203 0.027583 *
X78
          1.214e+00 5.512e-01
X79
          1.260e+00 5.406e-01
                             2.330 0.019792 *
          NA NA
X80
                            NA NA
X81
         -9.743e-01 5.222e-01 -1.866 0.062084 .
X82
          1.613e-01 5.382e-01 0.300 0.764460
XX3
         -3.621e-01 5.211e-01 -0.695 0.487085
          NA NA NA NA
X84
X85
          9.043e-01 5.062e-01 1.787 0.074003 .
X86
          7.261e-01 5.293e-01 1.372 0.170157
          1.059e+00 5.097e-01 2.078 0.037721 *
```

```
X88
                  NA
                                            NA
                          NA
                                  NA
X89
           2.883e-01 5.072e-01
                                0.568 0.569796
X90
            9.804e-01 5.246e-01
                                1.869 0.061651
X91
            5.441e-01 5.132e-01
                                1.060 0.289084
X92
                 NA
                        NA
                                  NA NA
           -1.432e-01 5.057e-01
                                -0.283 0.777065
X93
X94
            5.291e-02
                      5.146e-01
                                 0.103 0.918109
X95
            2.173e-01
                     5.090e-01
                                 0.427 0.669469
X96
                 NA
                          NA
                                   NA
X97
            4.378e-01 5.131e-01
                                 0.853 0.393486
X98
            8.407e-02
                      5.406e-01
                                 0.156 0.876409
X99
            2.933e-01 5.287e-01
                                 0.555 0.579057
X100
                NA
                       NA
                                  NA NA
X101
           -7.678e-01 5.254e-01 -1.461 0.143913
X102
                                0.256 0.797930
            1.422e-01 5.554e-01
X103
            6.235e-01 5.231e-01
                                1.192 0.233288
X104
                                  NA NA
                NΑ
                        NΑ
X105
            3.502e-01 5.011e-01
                                 0.699 0.484647
X106
            1.645e+00 5.299e-01
                                3.105 0.001905 **
            7.813e-01 4.921e-01
                                1.588 0.112390
                 NA
                           NA
                                  NA
X109
            4.334e-01 4.839e-01
                                0.896 0.370491
X110
           -4.023e-01 5.251e-01
                                -0.766 0.443578
X111
           2.668e-01 4.816e-01
                                0.554 0.579645
X112
                NA
                           NA
                                 NA
                                           NA
X113
           -9.224e-01 3.887e-01 -2.373 0.017643 *
                                0.249 0.803743
X114
           8.250e-02 3.320e-01
X115
           -1.809e-01 3.023e-01 -0.598 0.549545
X116
                  NΑ
                            NA
                                 NA
                                            NΑ
                                1.886 0.059306 .
X117
            4.771e-01 2.530e-01
X118
            7.559e-01 2.522e-01
                                 2.997 0.002729 **
X119
            1.068e+00 2.625e-01
                                 4.068 4.75e-05 ***
X120
                  NA
                            NA
                                  NA
                                            NA
X121
            1.392e+00 1.531e+00
                                 0.909 0.363397
X122
            4.643e+00 1.662e+00
                                 2.794 0.005203 **
X123
           -2.261e-02 1.645e+00
                                -0.014 0.989034
X124
            2.915e+00 1.682e+00
                                1.733 0.083059 .
X125
            1.399e+00 1.709e+00
                                 0.818 0.413093
X126
                                 0.132 0.894666
            2.255e-01 1.703e+00
X127
           7.565e-01 1.686e+00
                                 0.449 0.653590
X128
           7.542e-02 1.769e+00
                                 0.043 0.965992
X129
           -6.275e-01 1.760e+00
                                -0.357 0.721411
X130
           -7.346e-01 1.697e+00
                                -0.433 0.665108
X131
           3.897e+00 1.697e+00
                                2.297 0.021637 *
X132
           -1.363e+00 1.741e+00
                               -0.783 0.433531
X133
            8.729e-01 1.702e+00 0.513 0.608058
X134
            6.421e-01 1.747e+00 0.368 0.713231
```

```
1.487 0.137078
X135
            2.625e+00 1.766e+00
X136
            7.926e-02 1.660e+00 0.048 0.961912
X137
           -2.494e+00 1.708e+00 -1.461 0.144102
X138
           1.331e+00 1.762e+00 0.755 0.450100
X139
           -2.142e-01 1.751e+00 -0.122 0.902618
X140
           1.227e+00 1.722e+00 0.713 0.475896
           2.249e+00 1.665e+00 1.351 0.176761
X141
X142
           -4.386e-01 1.740e+00 -0.252 0.800977
X143
           -5.009e-01 1.779e+00 -0.282 0.778220
X144
            2.451e+00 1.653e+00
                                 1.482 0.138259
X145
           1.026e+00 1.652e+00
                                 0.621 0.534556
X146
           -1.804e-01 1.606e+00 -0.112 0.910555
X147
            3.322e-01 1.086e+00
                                  0.306 0.759785
X148
            2.386e+00 1.095e+00
                                  2.178 0.029375 *
X149
            9.826e-01 1.093e+00
                                  0.899 0.368849
X150
            1.002e+00 1.123e+00
                                  0.892 0.372146
X151
           -9.601e-01 1.107e+00 -0.867 0.385906
X152
                                 0.697 0.485927
            7.380e-01 1.059e+00
                                 0.567 0.570817
X153
            6.106e-01 1.077e+00
X154
           1.224e+00 1.071e+00 1.144 0.252830
X155
           -5.522e-02 1.086e+00 -0.051 0.959452
X156
           3.907e+00 1.126e+00 3.470 0.000520 ***
X157
            2.552e-01 1.118e+00 0.228 0.819467
X158
            8.416e-01 1.146e+00 0.734 0.462671
X159
            1.986e+00 1.097e+00 1.811 0.070174 .
X160
            1.521e+00 1.080e+00 1.408 0.159084
X161
           1.098e+00 1.113e+00 0.986 0.323929
X162
           1.547e+00 1.066e+00 1.452 0.146421
X163
           -6.634e-01 1.083e+00 -0.612 0.540325
X164
           -1.089e+00 1.118e+00 -0.975 0.329736
           1.626e-01 1.120e+00 0.145 0.884614
X165
X166
           9.596e-01 1.090e+00
                                 0.881 0.378486
X167
           1.543e-01 1.092e+00
                                 0.141 0.887630
X168
           7.783e-01 1.053e+00
                                  0.739 0.459844
X169
           8.224e-01 1.093e+00
                                 0.753 0.451709
X170
           -6.880e-01 1.158e+00 -0.594 0.552325
X171
           -8.968e-01 1.073e+00 -0.835 0.403501
X172
           1.725e+00 1.091e+00
                                 1.581 0.113971
X173
           -1.418e-01 1.224e+00 -0.116 0.907781
X174
           -2.184e+00 1.435e+00 -1.522 0.128123
           -8.784e-01 1.448e+00 -0.607 0.544105
X175
X176
           -3.279e+00 1.586e+00 -2.068 0.038671 *
X177
           -1.043e+00 1.553e+00 -0.671 0.502045
X178
           1.346e-01 1.529e+00 0.088 0.929824
X179
           -1.149e-02 1.551e+00 -0.007 0.994092
X180
           -6.474e-01 1.596e+00 -0.406 0.685050
           2.554e+00 1.627e+00 1.570 0.116387
X181
```

```
-2.062e-01 1.578e+00 -0.131 0.895999
X182
           1.394e+00 1.565e+00 0.891 0.372993
X183
           1.255e+00 1.577e+00 0.796 0.426132
X184
          -9.422e-01 1.579e+00 -0.597 0.550676
X185
          1.032e+00 1.604e+00 0.643 0.519987
X186
          6.324e-01 1.567e+00 0.404 0.686575
X187
X188
          -1.387e-01 1.560e+00 -0.089 0.929120
X189
          -6.548e-01 1.563e+00 -0.419 0.675224
X190
          6.980e-01 1.574e+00 0.443 0.657519
          1.260e+00 1.586e+00 0.795 0.426800
X192
          -2.413e+00 1.566e+00 -1.541 0.123395
X193
          7.788e-01 1.540e+00 0.506 0.612967
X194
          9.956e-02 1.491e+00 0.067 0.946774
          8.369e-01 1.571e+00 0.533 0.594244
X195
X196
          5.572e-01 1.572e+00 0.354 0.723042
          -1.248e+00 1.559e+00 -0.800 0.423428
X197
X198
          -2.590e-01 1.519e+00 -0.171 0.864615
           2.830e+00 1.320e+00 2.144 0.032071 *
X199
X200
          -3.267e-02 8.381e-01 -0.039 0.968902
X201
           3.283e+00 9.959e-01 3.297 0.000978 ***
X202
           2.051e+00 1.003e+00
                                2.046 0.040800 *
X203
           2.016e+00 1.000e+00
                                2.015 0.043889 *
X204
           4.282e-01 9.924e-01
                                0.431 0.666152
X205
           1.335e+00
                     1.005e+00
                                1.328 0.184085
X206
           8.355e-01 9.958e-01
                                0.839 0.401442
           9.288e-01 9.752e-01 0.952 0.340900
X207
           1.367e+00 1.026e+00 1.332 0.182931
X208
           2.078e+00 9.859e-01 2.108 0.035047 *
X209
           3.356e+00 9.981e-01 3.363 0.000772 ***
X210
           4.990e-01 9.761e-01 0.511 0.609231
X211
           1.823e+00 9.982e-01 1.826 0.067864
X212
X213
          3.085e+00 1.022e+00 3.017 0.002549 **
X214
          3.000e+00 9.950e-01 3.016 0.002565 **
X215
          1.920e+00 9.903e-01 1.939 0.052549
X216
          -7.898e-01 9.597e-01 -0.823 0.410530
          -1.170e+00 9.755e-01 -1.200 0.230263
          7.398e-01 1.002e+00 0.739 0.460201
X219
          9.754e-02 9.842e-01 0.099 0.921051
X220
          6.804e-01 9.941e-01 0.684 0.493678
X221
          1.195e+00 9.850e-01 1.213 0.225030
X222
          9.760e-01 1.001e+00 0.975 0.329624
X223
          7.381e-01 9.754e-01 0.757 0.449221
          -3.562e-01 9.625e-01 -0.370 0.711340
X224
X225
          1.502e+00 9.605e-01 1.564 0.117734
           1.461e+00 8.403e-01 1.739 0.082104
X226
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' '1
(Dispersion parameter for binomial family taken to be 1)
   Null deviance: 2772.6 on 1999 degrees of freedom
Residual deviance: 1587.4 on 1802 degrees of freedom
AIC: 1983.4
```

Figure 16. Logistic regression model on bound and unbound data (1-mer+shape)

Number of Fisher Scoring iterations: 15

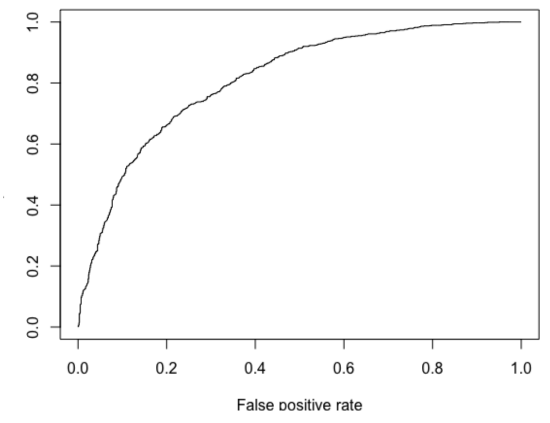


Figure 17. ROC plot of logistic regression model on bound and unbound data (1-mer+shape). AUC = 0.813343