

# DEB RNA HFX S100

## Load required packages

```
## corrplot 0.84 loaded
## -- Attaching packages -----
## v ggplot2 3.3.0    v purrr  0.3.4
## v tidyr  1.0.3    v dplyr  0.8.5
## v readr  1.3.1    v forcats 0.5.0
## -- Conflicts -----
## x dplyr::filter() masks stats::filter()
## x dplyr::lag()    masks stats::lag()
##
## Attaching package: 'kableExtra'
## The following object is masked from 'package:dplyr':
##
##   group_rows
##
## Attaching package: 'magrittr'
## The following object is masked from 'package:purrr':
##
##   set_names
## The following object is masked from 'package:tidyr':
##
##   extract
##
## Attaching package: 'gridExtra'
## The following object is masked from 'package:dplyr':
##
##   combine
##
## *****
## Note: As of version 1.0.0, cowplot does not change the
##   default ggplot2 theme anymore. To recover the previous
##   behavior, execute:
##   theme_set(theme_cowplot())
## *****
##
## Attaching package: 'cowplot'
```

```

## The following object is masked from 'package:ggpubr':
##
##   get_legend
## Loading required package: lattice
## Loading required package: survival
## Loading required package: Formula
##
## Attaching package: 'Hmisc'
## The following objects are masked from 'package:dplyr':
##
##   src, summarize
## The following objects are masked from 'package:base':
##
##   format.pval, units
##
## Attaching package: 'caret'
## The following object is masked from 'package:survival':
##
##   cluster
## The following object is masked from 'package:purrr':
##
##   lift
## Loading required package: usethis

```

## Loading RNP:XL data into the R environment

I included the manually evaluated datasets for my ASMS talk to see if machine learning can help me with identifying a possible score cutoff or other metric to help reduce the numbers of entries I have to go through after RNP:xl output. The datasets include a merged version of both the S100 and S100 fraction that were initially considered individually:

\*Ecoli DEB RNA

### Data wrangling

First, we need to add an extra column to each dataset, indicating which cross-linker was used in the experiment.

Then, we need to combine the dataframes.

### First filtering step

Remove any entry that does not contain an RNA adduct and fits to a target better than a decoy.

### Compute additional metrics (columns)

#### Peplength

#### classification of amino acids

Create new column with peptide length and classify identified amino acids in peptides as follows:

1. charged amino acids: R, K, D, E
2. polar amino acids: Q, N, H, S, T, Y, C, W
3. hydrophobic amino acids: A, I, L, M, V, P, G

### Peptide pI (isoelectric point)

Compute the isoelectric point for each peptide in each entry.

### aI (aliphatic index)

Compute the aliphatic Index for each peptide in each entry. See the following paper for more info:

*Thermostability and aliphatic index of globular proteins. J Biochem. 1980 Dec;88(6):1895-8.*

“A statistical analysis shows that the aliphatic index, which is defined as the relative volume of a protein occupied by aliphatic side chains (alanine, valine, isoleucine, and leucine), of proteins of thermophilic bacteria is significantly higher than that of ordinary proteins. The index may be regarded as a positive factor for the increase of thermostability of globular proteins.”

In general, an aI of greater 92.6 is particularly stable thermally as found in thermophiles. Now, since we are investigating tryptic peptides, I could only find one paper on that matter.

*GETTING INTIMATE WITH TRYPSIN, THE LEADING PROTEASE IN PROTEOMICS Published online 15 June 2013 in Wiley Online Library (wileyonlinelibrary.com). DOI 10.1002/mas.21376*

According to this paper, human proteins result in 61 tryptic peptides on average.

### proposed cross-linking amino acid

### Hydrophobicity index

### Molecular weight of peptide

*Note: returns the average masses, not the monoisotopic masses!*

## Machine learning

### Split dataset

Separate different species of nucleotides (RNA, DNA) and cross-linker (DEB, DEB).

### Setting up the dataset

Harmonize the number of trues and falses and dropping all categorical columns. 50% T and 50% F

```
set.seed(42)
drops_general<-c("precursor_purity", "RNPxl_a_ion_score")

ml_DEB_RNA<-data1 %>%
  filter(XLinker == "DEB", Curated == "0", Nucleotides == "RNA") %>%
  sample_n(459) %>%
  rbind(RNA_DEB)
rows_DEB_RNA <- sample(nrow(ml_DEB_RNA))
ml_DEB_RNA <- ml_DEB_RNA[rows_DEB_RNA, ] %>%
  dplyr::select_if(is.numeric)
ml_DEB_RNA<-ml_DEB_RNA[, !(colnames(ml_DEB_RNA) %in% drops_general)]
```

## Splitting into train and test data

I will use 80% of the data to train, and 20% to test the model

```
ml_DEB_RNA$Curated<-as.factor(ml_DEB_RNA$Curated)
ml_DEB_RNA$Curated<-factor(ml_DEB_RNA$Curated, levels = c("0", "1"), labels = c("False", "True"))
ml_DEB_RNA_train <- ml_DEB_RNA[1:734, ]
ml_DEB_RNA_test  <- ml_DEB_RNA[735:918, ]
```

## KNN

K nearest neighbor classification based on numeric variables, min-max normalized.

```
library(class)
library(gmodels)

knn_drop <- c("Curated", "isotope_error", "RNXPxl_a_ion_score", "RNXPxl_precursor_purity")

normalize <- function(x) {
  return((x-min(x))/(max(x)-min(x)))
}

ml_DEB_RNA_test_knn<-ml_DEB_RNA_test[,!(names(ml_DEB_RNA_test) %in% knn_drop)]
ml_DEB_RNA_test_knn<-as.data.frame(lapply(ml_DEB_RNA_test_knn[1:39], normalize))
ml_DEB_RNA_test_knn_labels<-ml_DEB_RNA_test[1:184, 5]

ml_DEB_RNA_train_knn<-ml_DEB_RNA_train[,!(names(ml_DEB_RNA_train) %in% knn_drop)]
ml_DEB_RNA_train_knn<-as.data.frame(lapply(ml_DEB_RNA_train_knn[1:39], normalize))
ml_DEB_RNA_train_knn_labels<-as.data.frame(ml_DEB_RNA_train[1:734, 5])
```

## Evaluate Model Performance DEB RNA

Here, I chose 5 neighbors:

```
DEB_RNA_knn_pred <- knn(train = ml_DEB_RNA_train_knn, test = ml_DEB_RNA_test_knn,
  cl = ml_DEB_RNA_train_knn_labels$Curated, k =5 )
CrossTable(x = ml_DEB_RNA_test_knn_labels$Curated, y = DEB_RNA_knn_pred, prop.chisq = F)
```

```
##
##
##   Cell Contents
## |-----|
## |                N |
## |      N / Row Total |
## |      N / Col Total |
## |      N / Table Total |
## |-----|
##
##
## Total Observations in Table:  184
##
##
##                | DEB_RNA_knn_pred
## ml_DEB_RNA_test_knn_labels$Curated |      False |      True | Row Total |
## -----|-----|-----|-----|
```

```
##           False |         79 |         17 |         96 |
##           |     0.823 |     0.177 |     0.522 |
##           |     0.859 |     0.185 |           |
##           |     0.429 |     0.092 |           |
## -----|-----|-----|-----|
##           True |         13 |         75 |         88 |
##           |     0.148 |     0.852 |     0.478 |
##           |     0.141 |     0.815 |           |
##           |     0.071 |     0.408 |           |
## -----|-----|-----|-----|
##           Column Total |         92 |         92 |         184 |
##           |     0.500 |     0.500 |           |
## -----|-----|-----|-----|
##
##
```

## Decision trees and classification rules

### C5.0

```
library(C50)

C5_drop <- c("isotope_error", "RNPx1_a_ion_score", "precursor_purity")

ml_DEB_RNA_train_C5<-ml_DEB_RNA_train[,!(names(ml_DEB_RNA_train) %in% C5_drop)]
ml_DEB_RNA_test_C5<-ml_DEB_RNA_test[,!(names(ml_DEB_RNA_test) %in% C5_drop)]
```

### DEB RNA

with boosting and penalty for false positives

penalize False Positives 4 times more than false negatives.

```
matrix_dimensions<- list(c("True", "False"), c("True", "False"))
names(matrix_dimensions) <- c("predicted", "actual")
error_cost <- matrix(c(0,1,4,0), nrow = 2, dimnames = matrix_dimensions)

C5_model_DEB_RNA_boost_pen<-C5.0(x=ml_DEB_RNA_train_C5[-5], y=ml_DEB_RNA_train_C5$Curated,
                                costs = error_cost, trials = 10)
summary(C5_model_DEB_RNA_boost_pen)
```

```
##
## Call:
## C5.0.default(x = ml_DEB_RNA_train_C5[-5], y =
## ml_DEB_RNA_train_C5$Curated, trials = 10, costs = error_cost)
##
##
## C5.0 [Release 2.07 GPL Edition]      Mon Jun 21 00:35:29 2021
## -----
##
## Class specified by attribute `outcome'
##
## Read 734 cases (40 attributes) from undefined.data
## Read misclassification costs from undefined.costs
```

```

##
## ----- Trial 0: -----
##
## Decision tree:
##
## RNPxl_total_MIC > 0.2365065:
## :...aa_class_polar > 4:
## :   :...RNPxl_pl_im_MIC <= 0.01192007: True (5)
## :   :   RNPxl_pl_im_MIC > 0.01192007: False (4/1)
## :   aa_class_polar <= 4:
## :   :...Hydro <= 0.6: True (205/3)
## :   :   Hydro > 0.6:
## :   :   :...pI <= 9.45: False (6/3)
## :   :   :   pI > 9.45: True (8)
## RNPxl_total_MIC <= 0.2365065:
## :...aa_class_hydrophobic > 3:
## :   :...RNPxl_pl_pc_MIC <= 0: False (322/26)
## :   :   RNPxl_pl_pc_MIC > 0:
## :   :   :...RNPxl_pl_pc_MIC > 0.04382946: True (8)
## :   :   :   RNPxl_pl_pc_MIC <= 0.04382946:
## :   :   :   :...RNPxl_modds <= 0.3741593: False (25/3)
## :   :   :   :   RNPxl_modds > 0.3741593:
## :   :   :   :   :...RNPxl_RNA_MASS_z0 <= 449.0948: False (5)
## :   :   :   :   :   RNPxl_RNA_MASS_z0 > 449.0948: True (14)
## aa_class_hydrophobic <= 3:
## :...aa_class_charged <= 2:
## :   :...RNPxl_pl_modds <= 0.4927427: False (14/9)
## :   :   RNPxl_pl_modds > 0.4927427:
## :   :   :...RNPxl_precursor_score <= 0.001193983: True (60)
## :   :   :   RNPxl_precursor_score > 0.001193983: False (1)
## aa_class_charged > 2:
## :...RT <= 818.5449: True (7)
## :   RT > 818.5449:
## :   :...RNPxl_pl_modds <= 0.6882493: False (19/2)
## :   :   RNPxl_pl_modds > 0.6882493:
## :   :   :...pI <= 4.43: True (10)
## :   :   :   pI > 4.43:
## :   :   :   :...RNPxl_pl_err <= 0.006021813: False (8/1)
## :   :   :   :   RNPxl_pl_err > 0.006021813:
## :   :   :   :   :...RNPxl_immonium_score <= 0.02881235: True (12)
## :   :   :   :   :   RNPxl_immonium_score > 0.02881235: False (1)
##
## ----- Trial 1: -----
##
## Decision tree:
##
## mw > 1856:
## :...RNPxl_pl_modds <= 0.8764216: False (128.9/3)
## :   RNPxl_pl_modds > 0.8764216:
## :   :...RNPxl_err <= 0.002779919: True (40.4/0.8)
## :   :   RNPxl_err > 0.002779919: False (45.6/22.1)
## mw <= 1856:
## :...RNPxl_pl_modds <= 0.04243729: False (18.2/2.3)
## :   RNPxl_pl_modds > 0.04243729:

```

```

##      :...RNPxl_err > 0.009156602: True (73.1/0.8)
##      RNPxl_err <= 0.009156602:
##      :...precursor > 973.7704: False (19/3)
##      precursor <= 973.7704:
##      :...RNPxl_total_MIC > 0.3725607: True (79.6)
##      RNPxl_total_MIC <= 0.3725607:
##      :...aa_class_hydrophobic > 7: False (31.1/14.4)
##      aa_class_hydrophobic <= 7:
##      :...pI > 11.13: False (16.7/5.3)
##      pI <= 11.13:
##      :...RNPxl_immonium_score > 0.008700988:
##      :...Peplength <= 7: True (51.7/3)
##      :   Peplength > 7:
##      :   :...RNPxl_marker_ions_score > 0.0006902145: True (22.8/1.5)
##      :   :   RNPxl_marker_ions_score <= 0.0006902145: [S1]
##      RNPxl_immonium_score <= 0.008700988:
##      :...RNPxl_partial_loss_score > 6.786638: True (100.5)
##      RNPxl_partial_loss_score <= 6.786638:
##      :...Peplength > 10: False (12.1/1.5)
##      Peplength <= 10:
##      :...aIndex > 128.12: False (5.3/0.8)
##      aIndex <= 128.12:
##      :...Peplength > 9: True (69.2)
##      Peplength <= 9:
##      :...pI <= 4.18: False (8.3/3.8)
##      pI > 4.18: [S2]
##
## SubTree [S1]
##
## RNPxl_partial_loss_score > 13.77283: True (9.9)
## RNPxl_partial_loss_score <= 13.77283:
## :...mw <= 861.97: True (9.9)
##   mw > 861.97:
##   :...RNPxl_pl_MIC <= 0.05950294: False (49.7/13.6)
##   RNPxl_pl_MIC > 0.05950294: True (15.2/1.5)
##
## SubTree [S2]
##
## RNPxl_Da_difference <= -0.0001694992: False (34.4/26.6)
## RNPxl_Da_difference > -0.0001694992: True (133.8/2.3)
##
## ----- Trial 2: -----
##
## Decision tree:
##
## mw > 2184.58: False (106.7/20.2)
## mw <= 2184.58:
## :...RNPxl_total_MIC > 0.2095862: True (384.2/11.5)
##   RNPxl_total_MIC <= 0.2095862:
##   :...score <= 0.0594036: True (70.1/0.6)
##   score > 0.0594036:
##   :...RNPxl_pl_modds <= 0.8817645:
##     :...RNPxl_precursor_score > 0.0002412407: True (30.2)
##     :   RNPxl_precursor_score <= 0.0002412407:

```

```

##           :   ...Peplength > 12: False (64.6/5)
##           :           Peplength <= 12:
##           :           ...RNPxl_Da_difference <= -0.0001225632: False (21.8/2.9)
##           :           RNPxl_Da_difference > -0.0001225632:
##           :           ...RNPxl_err <= 0.004044351: False (62.2/43.4)
##           :           RNPxl_err > 0.004044351: True (46/0.6)
## RNPxl_pl_modds > 0.8817645:
##           :...Hydro > 0.28: False (8.2/0.6)
##           Hydro <= 0.28:
##           :...RNPxl_immonium_score <= 0.008185891:
##           :           ...RNPxl_total_MIC <= 0.06030513: False (19.3/15.2)
##           :           RNPxl_total_MIC > 0.06030513: True (199/2.3)
##           RNPxl_immonium_score > 0.008185891:
##           :...RNPxl_pl_err <= 0.003974453: False (10.1)
##           RNPxl_pl_err > 0.003974453:
##           :...aa_class_polar <= 5: True (95.1/8.2)
##           aa_class_polar > 5: False (4.7)
##
## ----- Trial 3: -----
##
## Decision tree:
##
## aa_class_hydrophobic <= 3:
## :...RNPxl_RNA_MASS_z0 <= 391.0781: False (31.7/20)
## :   RNPxl_RNA_MASS_z0 > 391.0781:
## :   :...Hydro <= -2.13: False (32.3/24.7)
## :   Hydro > -2.13: True (395.6/9.2)
## aa_class_hydrophobic > 3:
## :...RNPxl_total_MIC > 0.3001701:
## :   ...RNPxl_mass_error_p <= 1.941169: True (126.3)
## :   RNPxl_mass_error_p > 1.941169: False (7.7/3.2)
## RNPxl_total_MIC <= 0.3001701:
## :...mw > 2405.7: False (50.9)
## mw <= 2405.7:
## :...RNPxl_pl_modds <= 0.07640828: False (21.3)
## RNPxl_pl_modds > 0.07640828:
## :...RNPxl_immonium_score > 0.008149372:
## :   ...RNPxl_mass_error_p <= 1.912772: False (89.1/21)
## :   RNPxl_mass_error_p > 1.912772: True (81.1/12.7)
## RNPxl_immonium_score <= 0.008149372:
## :...RNPxl_err <= 0.001154643: False (19.5/6.8)
## RNPxl_err > 0.001154643:
## :...score <= 0.813925: True (207.1/8.2)
## score > 0.813925:
## :...RNPxl_pl_Morph <= 8.029901: False (18)
## RNPxl_pl_Morph > 8.029901: True (49.3/5.9)
##
## ----- Trial 4: -----
##
## Decision tree:
##
## RNPxl_immonium_score > 0.02881235:
## :...RNPxl_pl_pc_MIC <= 0.01601188: False (45/1.1)
## :   RNPxl_pl_pc_MIC > 0.01601188: True (9.6)

```



```

## RNPxl_immonium_score <= 0.02881235:
## :...RNPxl_total_MIC > 0.2148431:
##   :...aa_class_polar <= 6: True (325/12.5)
##   :   aa_class_polar > 6: False (8/6.1)
## RNPxl_total_MIC <= 0.2148431:
##   :...aa_class_hydrophobic <= 3:
##   :   :...precursor <= 524.2557: False (8.2/3.7)
##   :   :   precursor > 524.2557:
##   :   :     :...RNPxl_immonium_score > 0.009534214: False (64.4/50)
##   :   :     :   RNPxl_immonium_score <= 0.009534214:
##   :   :     :     :...RNPxl_score <= 0.8732423: True (253.2/6.8)
##   :   :     :     :   RNPxl_score > 0.8732423: False (3.8)
##   aa_class_hydrophobic > 3:
##   :...aa_class_hydrophobic > 12: False (22.1)
##   :   aa_class_hydrophobic <= 12:
##   :     :...RNPxl_pl_pc_MIC > 0.04382946: True (35.6)
##   :     :   RNPxl_pl_pc_MIC <= 0.04382946:
##   :     :     :...RNPxl_pl_Morph > 12.05118: True (50.6/3.8)
##   :     :     :   RNPxl_pl_Morph <= 12.05118:
##   :     :     :     :...aa_class_charged > 5: False (24.5)
##   :     :     :     :   aa_class_charged <= 5:
##   :     :     :     :     :...aa_class_polar > 5: False (18.3)
##   :     :     :     :     :   aa_class_polar <= 5:
##   :     :     :     :     :     :...precursor <= 727.8291: False (28.6/9.3)
##   :     :     :     :     :     :   precursor > 727.8291:
##   :     :     :     :     :     :     :...aa_class_charged <= 1: True (25.1/0.7)
##   :     :     :     :     :     :     :   aa_class_charged > 1:
##   :     :     :     :     :     :     :     :...pI > 6.45: False (55.9/18.6)
##   :     :     :     :     :     :     :     :   pI <= 6.45:
##   :     :     :     :     :     :     :     :     :...RNPxl_pl_Morph <= 4.00529: False (8.3)
##   :     :     :     :     :     :     :     :     :   RNPxl_pl_Morph > 4.00529: [S1]
##   :     :     :     :     :     :     :     :     :     :
## SubTree [S1]
##
## RNPxl_pl_im_MIC <= 0.01466738: True (120.9/8.8)
## RNPxl_pl_im_MIC > 0.01466738: False (6.1/0.4)
##
## ----- Trial 5: -----
##
## Decision tree:
##
## aa_class_hydrophobic > 5:
## :...RNPxl_pl_pc_MIC > 0.01601188: True (40.4/1.3)
## :   RNPxl_pl_pc_MIC <= 0.01601188:
## :     :...score <= 0.145311: True (31.8/3.1)
## :     :   score > 0.145311:
## :     :     :...RNPxl_pl_MIC > 0.1462661: True (30.6/1.8)
## :     :     :   RNPxl_pl_MIC <= 0.1462661:
## :     :     :     :...RNPxl_pl_modds <= 1.37235: False (144/33.1)
## :     :     :     :   RNPxl_pl_modds > 1.37235:
## :     :     :     :     :...RT <= 1838.091: True (32.7)
## :     :     :     :     :   RT > 1838.091: False (65.9/30)
## aa_class_hydrophobic <= 5:
## :...RNPxl_total_MIC <= 0.04410198: False (54/33.5)

```

```

## RNPxl_total_MIC > 0.04410198:
## :...aIndex > 78.5:
## :...RNPxl_total_MIC > 0.3017294: True (34.4)
## : RNPxl_total_MIC <= 0.3017294:
## : :...RNPxl_marker_ions_score <= 0.001819908: False (55.7/31.2)
## : RNPxl_marker_ions_score > 0.001819908: True (10.1)
## aIndex <= 78.5:
## :...Peplength <= 6: True (92)
## Peplength > 6:
## :...aa_class_hydrophobic <= 1: False (26.6/18)
## aa_class_hydrophobic > 1:
## :...RNPxl_Morph <= 4.025711: True (206.3/1.1)
## RNPxl_Morph > 4.025711:
## :...precursor_intensity <= 1252038: False (70.7/51)
## precursor_intensity > 1252038:
## :...RNPxl_err <= 0.001018666: False (12.5/6.4)
## RNPxl_err > 0.001018666: True (204.2/3.4)
##
## ----- Trial 6: -----
##
## Decision tree:
##
## Peplength > 10:
## :...RNPxl_total_MIC > 0.2480816: True (68.5/2.2)
## : RNPxl_total_MIC <= 0.2480816:
## : :...aIndex <= 20: True (38.7/3.1)
## : aIndex > 20:
## : :...RNPxl_pl_modds <= 0.8905877: False (110.2/15.7)
## : RNPxl_pl_modds > 0.8905877:
## : :...precursor_error_ppm > 4.391985: True (29.9)
## : precursor_error_ppm <= 4.391985:
## : :...RNPxl_pl_pc_MIC > 0.01709038: True (17)
## : RNPxl_pl_pc_MIC <= 0.01709038:
## : :...RNPxl_immonium_score > 0.008237569: False (29.4/2.9)
## : RNPxl_immonium_score <= 0.008237569:
## : :...RNPxl_mass_error_p <= 1.497245: False (8.7)
## : RNPxl_mass_error_p > 1.497245: True (81.4/5.7)
## Peplength <= 10:
## :...score <= 0.0752907: True (87.8/0.2)
## score > 0.0752907:
## :...aIndex > 86.25:
## :...RNPxl_pl_Morph <= 2.227052: False (9.9)
## : RNPxl_pl_Morph > 2.227052:
## : :...RNPxl_Morph <= 4.530347: True (102.8/11)
## : RNPxl_Morph > 4.530347: False (9.7/2.1)
## aIndex <= 86.25:
## :...RNPxl_RNA_MASS_z0 > 758.1211: False (26.4/18.5)
## RNPxl_RNA_MASS_z0 <= 758.1211:
## :...aa_class_charged <= 2: True (335.1/9.3)
## aa_class_charged > 2:
## :...precursor_error_ppm <= 0.6376348: True (82.6)
## precursor_error_ppm > 0.6376348:
## :...RNPxl_precursor_score > 0.0007802289: False (3.9/0.4)
## RNPxl_precursor_score <= 0.0007802289:

```

```

##           :...RNPxl_pl_err > 0.01726561: False (3.1)
##           RNPxl_pl_err <= 0.01726561:
##           :...RNPxl_marker_ions_score > 0.001386256: False (2.6)
##           RNPxl_marker_ions_score <= 0.001386256:
##           :...RNPxl_pl_modds > 3.219504: False (3.4)
##           RNPxl_pl_modds <= 3.219504:
##           :...mw <= 1219.46: True (144.4/6.9)
##           mw > 1219.46: False (2.5)
##
## ----- Trial 7: -----
##
## Decision tree:
##
## aa_class_hydrophobic > 6:
## :...RNPxl_pl_pc_MIC > 0.01601188: True (30.1/1.7)
## :   RNPxl_pl_pc_MIC <= 0.01601188:
## :     :...OMS_precursor_mz_error_ppm <= 3.760296: False (188.2/80.2)
## :     :   OMS_precursor_mz_error_ppm > 3.760296: True (27.7/1.6)
## aa_class_hydrophobic <= 6:
## :...RNPxl_pl_MIC > 0.1492823: True (76.8)
## :   RNPxl_pl_MIC <= 0.1492823:
## :     :...RNPxl_RNA_MASS_z0 <= 335.123:
## :     :   :...score > 0.929751: True (22.6)
## :     :   :   score <= 0.929751:
## :     :     :...precursor_error_ppm <= 0.5690612: True (17.5/0.2)
## :     :     :   precursor_error_ppm > 0.5690612:
## :     :     :     :...RNPxl_Da_difference > 0.00196853: True (14.2/0.3)
## :     :     :     :   RNPxl_Da_difference <= 0.00196853:
## :     :     :     :     :...RNPxl_peptide_mass_z0 <= 827.5593: True (12.2/0.2)
## :     :     :     :     :   RNPxl_peptide_mass_z0 > 827.5593: False (37.5/0.3)
## :     :     :   RNPxl_RNA_MASS_z0 > 335.123:
## :     :     :     :...RNPxl_total_MIC <= 0.037681: False (41.6/21.6)
## :     :     :     :   RNPxl_total_MIC > 0.037681:
## :     :     :     :     :...aa_class_hydrophobic > 5: False (27.9/15.8)
## :     :     :     :     :   aa_class_hydrophobic <= 5:
## :     :     :     :     :     :...Peplength > 12: False (25.4/16.1)
## :     :     :     :     :     :   Peplength <= 12:
## :     :     :     :     :     :     :...OMS_precursor_mz_error_ppm > 2.227397: True (115/0.2)
## :     :     :     :     :     :     :   OMS_precursor_mz_error_ppm <= 2.227397:
## :     :     :     :     :     :     :     :...RNPxl_Da_difference <= -0.0007595894:
## :     :     :     :     :     :     :     :   :...precursor_error_ppm <= 3.304309: False (55.9/33.1)
## :     :     :     :     :     :     :     :   :   precursor_error_ppm > 3.304309: True (11.9)
## :     :     :     :     :     :     :     :   RNPxl_Da_difference > -0.0007595894:
## :     :     :     :     :     :     :     :     :...RNPxl_modds > 1.71208: False (17.2/12.3)
## :     :     :     :     :     :     :     :     :   RNPxl_modds <= 1.71208:
## :     :     :     :     :     :     :     :     :     :...RT <= 4286.273: True (391.1/13.1)
## :     :     :     :     :     :     :     :     :     :   RT > 4286.273: False (5.1/1.5)
##
## ----- Trial 8: -----
##
## Decision tree:
##
## RNPxl_MIC > 0.3118303: True (111.2)
## RNPxl_MIC <= 0.3118303:

```

```

## :...mw > 2184.58: False (73.9/29.7)
##   mw <= 2184.58:
##     :...RT <= 1262.902: True (274.2/14.8)
##       RT > 1262.902:
##         :...RNPxl_pl_modds <= 0.1330409: False (22)
##           RNPxl_pl_modds > 0.1330409:
##             :...RNPxl_RNA_MASS_z0 <= 351.1179:
##               :...RNPxl_pl_MIC > 0.08438432: True (39.5/1.2)
##                 :   RNPxl_pl_MIC <= 0.08438432:
##                   :     :...RNPxl_pl_modds <= 3.201042: False (68.3/24.2)
##                     :       RNPxl_pl_modds > 3.201042: True (10.7)
##                 RNPxl_RNA_MASS_z0 > 351.1179:
##                   :...pI > 9.3:
##                     :...precursor_intensity <= 953246.9: True (66.8/4)
##                       :   precursor_intensity > 953246.9:
##                         :     :...RNPxl_MIC <= 0.2795199: False (84.3/40.6)
##                           :       RNPxl_MIC > 0.2795199: True (18.6)
##                     pI <= 9.3:
##                       :...Hydro <= -1.89: False (7.5)
##                         Hydro > -1.89:
##                           :...aa_class_polar <= 1: False (29.3/21.2)
##                             aa_class_polar > 1:
##                               :...Peplength <= 9: True (205.9/0.1)
##                                 Peplength > 9:
##                                   :...precursor_error_ppm > 2.518739: False (5.9/0.4)
##                                     precursor_error_ppm <= 2.518739:
##                                       :...pI <= 3.88: False (3.1)
##                                         pI > 3.88: True (170.5/12)
##
## ----- Trial 9: -----
##
## Decision tree:
##
## RNPxl_total_MIC <= 0.2148431:
## :...aa_class_hydrophobic > 3:
## :   :...Peplength > 20: False (27.5)
## :   :   Peplength <= 20:
## :   :     :...RNPxl_score <= -0.640421: False (145.5/61.1)
## :   :     :   RNPxl_score > -0.640421:
## :   :     :     :...RNPxl_pl_pc_MIC > 0.0002567632: True (93.8/4.2)
## :   :     :     :   RNPxl_pl_pc_MIC <= 0.0002567632:
## :   :     :     :     :...RNPxl_immonium_score <= 0.008149372: True (112.5/9.9)
## :   :     :     :     :   RNPxl_immonium_score > 0.008149372: False (51.5/12.5)
## :   :   aa_class_hydrophobic <= 3:
## :   :     :...aa_class_charged <= 2: True (179.1/6)
## :   :     :   aa_class_charged > 2:
## :   :     :     :...RNPxl_pl_modds > 2.211792: True (27.6)
## :   :     :     :   RNPxl_pl_modds <= 2.211792:
## :   :     :     :     :...precursor_error_ppm <= 0.51688: True (48.3/3.4)
## :   :     :     :     :   precursor_error_ppm > 0.51688:
## :   :     :     :     :     :...RNPxl_peptide_mass_z0 <= 678.4429: True (7.7)
## :   :     :     :     :     :   RNPxl_peptide_mass_z0 > 678.4429:
## :   :     :     :     :     :     :...RNPxl_Morph <= 2.469005: True (11.3/0.6)
## :   :     :     :     :     :     :   RNPxl_Morph > 2.469005: False (37.1/8.2)

```

```

## RNPxl_total_MIC > 0.2148431:
## :...aa_class_polar > 6: False (13.2/9.6)
##   aa_class_polar <= 6:
##     :...RNPxl_total_MIC > 0.3725607: True (150.8)
##       RNPxl_total_MIC <= 0.3725607:
##         :...RNPxl_score <= -0.9358985: True (99.4)
##           RNPxl_score > -0.9358985:
##             :...Hydro <= -2.04: False (2.8)
##               Hydro > -2.04:
##                 :...RT > 2551.2: False (20/12.4)
##                   RT <= 2551.2:
##                     :...RNPxl_pl_pc_MIC <= 0.1841059: True (119.7/2.3)
##                       RNPxl_pl_pc_MIC > 0.1841059: False (3.1)
##
##
## Evaluation on training data (734 cases):
##
## Trial          Decision Tree
## -----
##   Size      Errors  Cost
##
##   0      19  48( 6.5%)  0.08
##   1      22  92(12.5%)  0.18
##   2      14  62( 8.4%)  0.21
##   3      13 107(14.6%)  0.39
##   4      19  74(10.1%)  0.26
##   5      16  87(11.9%)  0.20
##   6      21  63( 8.6%)  0.27
##   7      18  90(12.3%)  0.22
##   8      16  86(11.7%)  0.32
##   9      18  58( 7.9%)  0.21
## boost                2( 0.3%)  0.00  <<
##
##
##   (a)  (b)  <-classified as
##   ----  ----
##   363      (a): class False
##     2  369  (b): class True
##
##
## Attribute usage:
##
## 100.00% RNPxl_MIC
## 100.00% RNPxl_immonium_score
## 100.00% RNPxl_pl_modds
## 100.00% RNPxl_total_MIC
## 100.00% Peplength
## 100.00% aa_class_hydrophobic
## 100.00% mw
##  98.23% score
##  97.14% RNPxl_pl_MIC
##  96.73% aIndex
##  76.29% RNPxl_err
##  76.16% OMS_precursor_mz_error_ppm

```

```

## 74.52% RT
## 72.21% RNPxl_RNA_MASS_z0
## 70.98% precursor
## 70.30% aa_class_polar
## 65.94% aa_class_charged
## 64.31% RNPxl_pl_pc_MIC
## 61.72% Hydro
## 61.44% RNPxl_score
## 59.54% pI
## 43.32% RNPxl_Da_difference
## 41.14% RNPxl_precursor_score
## 40.87% RNPxl_Morph
## 40.19% precursor_error_ppm
## 38.69% RNPxl_pl_Morph
## 36.51% RNPxl_partial_loss_score
## 35.56% RNPxl_mass_error_p
## 28.47% RNPxl_modds
## 27.52% precursor_intensity
## 24.39% RNPxl_marker_ions_score
## 19.21% RNPxl_pl_err
## 8.31% RNPxl_peptide_mass_z0
## 7.08% RNPxl_pl_im_MIC
##
##
## Time: 0.2 secs

```

```

C5_DEB_RNA_predict_boost_pen<-predict(C5_model_DEB_RNA_boost_pen, ml_DEB_RNA_test_C5)
CrossTable(ml_DEB_RNA_test_C5$Curated, C5_DEB_RNA_predict_boost_pen, prop.chisq = FALSE,
            prop.c = FALSE, prop.r = FALSE, dnn = c("actual TRUE", "predicted TRUE"))

```

```

##
##
## Cell Contents
## |-----|
## |                N |
## |      N / Table Total |
## |-----|
##
##
## Total Observations in Table: 184
##
##
##      | predicted TRUE
## actual TRUE | False | True | Row Total |
## -----|-----|-----|-----|
##      False |      86 |     10 |          96 |
##           | 0.467 | 0.054 |           |
## -----|-----|-----|-----|
##      True |      5 |     83 |          88 |
##           | 0.027 | 0.451 |           |
## -----|-----|-----|-----|
## Column Total |      91 |     93 |          184 |
## -----|-----|-----|-----|
##
##

```

# Classifiers

## DEB RNA

### RIPPER

```
library(rJava)
library(RWeka)
```

```
RIPPER_model_DEB_RNA <- JRip(Curated ~ ., data = ml_DEB_RNA_train_C5)
RIPPER_model_DEB_RNA
```

```
## JRIP rules:
```

```
## =====
```

```
##
```

```
## (mw >= 1164.34) and (RNPxl_total_MIC <= 0.148872) and (RNPxl_pl_modds <= 0.848058) and (score >= 0.4
```

```
## (RNPxl_total_MIC <= 0.236507) and (Hydro >= -0.67) and (RNPxl_immonium_score >= 0.008186) => Curated
```

```
## (RNPxl_peptide_mass_z0 >= 1130.473637) and (RNPxl_total_MIC <= 0.193156) and (RNPxl_RNA_MASS_z0 <= 6
```

```
## (RNPxl_total_MIC <= 0.213968) and (precursor_intensity <= 1138125.875) and (RNPxl_pl_modds <= 0.6873
```

```
## (RNPxl_total_MIC <= 0.28572) and (aa_class_hydrophobic >= 4) and (OMS_precursor_mz_error_ppm >= -0.1
```

```
## (aIndex >= 70) and (RNPxl_total_MIC <= 0.213968) and (RNPxl_pl_MIC <= 0.01229) => Curated=False (12.
```

```
## => Curated=True (364.0/21.0)
```

```
##
```

```
## Number of Rules : 7
```

```
summary(RIPPER_model_DEB_RNA)
```

```
##
```

```
## === Summary ===
```

```
##
```

```
## Correctly Classified Instances          685           93.3243 %
```

```
## Incorrectly Classified Instances         49           6.6757 %
```

```
## Kappa statistic                        0.8665
```

```
## Mean absolute error                    0.1161
```

```
## Root mean squared error                0.2409
```

```
## Relative absolute error                23.2226 %
```

```
## Root relative squared error            48.1898 %
```

```
## Total Number of Instances              734
```

```
##
```

```
## === Confusion Matrix ===
```

```
##
```

```
##   a   b   <-- classified as
```

```
## 342  21 |   a = False
```

```
##  28 343 |   b = True
```

```
RIPPER_DEB_RNA_predict <- predict(RIPPER_model_DEB_RNA, ml_DEB_RNA_test_C5)
```

```
CrossTable(ml_DEB_RNA_test_C5$Curated, RIPPER_DEB_RNA_predict, prop.chisq = FALSE,
  prop.c = FALSE, prop.r = FALSE, dnn = c("actual TRUE", "predicted TRUE"))
```

```
##
```

```
##
```

```
##   Cell Contents
```

```
## |-----|
```

```
## |                                     N |
```

```
## |           N / Table Total |
```

```
## |-----|
```

```
##
##
## Total Observations in Table: 184
##
##
##      | predicted TRUE
## actual TRUE | False | True | Row Total |
## -----|-----|-----|-----|
##      False |      81 |     15 |         96 |
##           | 0.440 | 0.082 |           |
## -----|-----|-----|-----|
##      True |     15 |     73 |         88 |
##           | 0.082 | 0.397 |           |
## -----|-----|-----|-----|
## Column Total |     96 |     88 |        184 |
## -----|-----|-----|-----|
##
##
```

## DEB RNA HFX, charge state 2-7

Now, let us test KNN, C5, and RIPPER models on unknown data: Ecoli DEB HFX runs. Combine the triplicates into one giant dataframe first!

### Data setup DEB RNA HFX charge state 2-7

```
A_Wulf_020819_080819_Ecoli_DEB_S100_rep1_XL_table <- read_delim("A_Wulf_020819_080819_Ecoli_DEB_S100_rep1_XL_table.txt",
  delim = "\t")
```

```
## Parsed with column specification:
## cols(
##   .default = col_double(),
##   sequence = col_character(),
##   accessions = col_character(),
##   `RNPxl:NT` = col_character(),
##   `RNPxl:RNA` = col_character(),
##   `RNPxl:best_localization` = col_character(),
##   `RNPxl:localization_scores` = col_character(),
##   protein_references = col_character(),
##   target_decoy = col_character(),
##   `PeakAnnotations(mz|intensity|charge|annotation;)` = col_character()
## )
```

## See spec(...) for full column specifications.

```
A_Wulf_020819_080819_Ecoli_DEB_S100_rep2_XL_table <- read_delim("A_Wulf_020819_080819_Ecoli_DEB_S100_rep2_XL_table.txt",
  delim = "\t")
```

```
## Parsed with column specification:
## cols(
##   .default = col_double(),
##   sequence = col_character(),
##   accessions = col_character(),
##   `RNPxl:NT` = col_character(),
##   `RNPxl:RNA` = col_character(),
```



```

## `RNPxl:best_localization` = col_character(),
## `RNPxl:localization_scores` = col_character(),
## protein_references = col_character(),
## target_decoy = col_character(),
## `PeakAnnotations(mz|intensity|charge|annotation;)` = col_character()
## )
## See spec(...) for full column specifications.
A_Wulf_020819_080819_Ecoli_DEB_S100_rep3_XL_table <- read_delim("A_Wulf_020819_080819_Ecoli_DEB_S100_rep3_XL_table",
  delim = "\t")

## Parsed with column specification:
## cols(
##   .default = col_double(),
##   sequence = col_character(),
##   accessions = col_character(),
##   `RNPxl:NT` = col_character(),
##   `RNPxl:RNA` = col_character(),
##   `RNPxl:best_localization` = col_character(),
##   `RNPxl:localization_scores` = col_character(),
##   protein_references = col_character(),
##   target_decoy = col_character(),
##   `PeakAnnotations(mz|intensity|charge|annotation;)` = col_character()
## )
## See spec(...) for full column specifications.
DEB_RNA_S100_HFX_total_reps <- rbind(A_Wulf_020819_080819_Ecoli_DEB_S100_rep1_XL_table,
  A_Wulf_020819_080819_Ecoli_DEB_S100_rep2_XL_table, A_Wulf_020819_080819_Ecoli_DEB_S100_rep3_XL_table)

DEB_RNA_HFX_S100_KNN <- DEB_RNA_S100_HFX_total_reps
names(DEB_RNA_HFX_S100_KNN) <- gsub("\\.", "_", names(DEB_RNA_HFX_S100_KNN))
names(DEB_RNA_HFX_S100_KNN) <- gsub("\\:", "_", names(DEB_RNA_HFX_S100_KNN))
names(DEB_RNA_HFX_S100_KNN) <- gsub("\\ ", "_", names(DEB_RNA_HFX_S100_KNN))
names(DEB_RNA_HFX_S100_KNN) <- gsub("\\|", "_", names(DEB_RNA_HFX_S100_KNN))
DEB_RNA_HFX_S100_KNN$XLinker <- "DEB"
DEB_RNA_HFX_S100_KNN$Nucleotides <- "RNA"
DEB_RNA_HFX_S100_KNN$Sample <- "S100"
DEB_RNA_HFX_S100_KNN <- DEB_RNA_HFX_S100_KNN[DEB_RNA_HFX_S100_KNN$RNPxl_RNA != "none",
] %>% filter(target_decoy == "target")
DEB_RNA_HFX_S100_KNN$prepPep <- gsub("(Carbamyl)", "", as.character(DEB_RNA_HFX_S100_KNN$sequence))
DEB_RNA_HFX_S100_KNN$prepPep <- gsub("(Oxidation)", "", as.character(DEB_RNA_HFX_S100_KNN$prepPep))
DEB_RNA_HFX_S100_KNN$prepPep <- gsub("(Phospho)", "", as.character(DEB_RNA_HFX_S100_KNN$prepPep))
DEB_RNA_HFX_S100_KNN$prepPep <- gsub("\\(|\\|)", "", DEB_RNA_HFX_S100_KNN$prepPep)
DEB_RNA_HFX_S100_KNN$prepPep <- gsub("\\\\.\\.*", "", DEB_RNA_HFX_S100_KNN$prepPep)
DEB_RNA_HFX_S100_KNN$Pepseq <- DEB_RNA_HFX_S100_KNN$prepPep
DEB_RNA_HFX_S100_KNN$Peplength <- nchar(DEB_RNA_HFX_S100_KNN$Pepseq, type = "chars")
DEB_RNA_HFX_S100_KNN$aaclasscharged <- str_count(DEB_RNA_HFX_S100_KNN$Pepseq, paste("R|K|D|E",
collapse = "|"))
DEB_RNA_HFX_S100_KNN$aaclasspolar <- str_count(DEB_RNA_HFX_S100_KNN$Pepseq, paste("Q|N|H|S|T|Y|C|W",
collapse = "|"))
DEB_RNA_HFX_S100_KNN$aaclasshydrophobic <- str_count(DEB_RNA_HFX_S100_KNN$Pepseq,
paste("A|I|L|M|F|V|P|G", collapse = "|"))
DEB_RNA_HFX_S100_KNN$pI <- round(pI(DEB_RNA_HFX_S100_KNN$Pepseq), 2)
DEB_RNA_HFX_S100_KNN$aIndex <- round(aIndex(DEB_RNA_HFX_S100_KNN$Pepseq), 2)

```

```

## Warning: Sequence 31683 has unrecognized amino acid types. Output value might be
## wrong calculated

## Warning: Sequence 35472 has unrecognized amino acid types. Output value might be
## wrong calculated

## Warning: Sequence 36280 has unrecognized amino acid types. Output value might be
## wrong calculated

## Warning: Sequence 37383 has unrecognized amino acid types. Output value might be
## wrong calculated
DEB_RNA_HFX_S100_KNN$XL_aa <- gsub("[:A-Z:]", "", DEB_RNA_HFX_S100_KNN$RNPxl_best_localization)
DEB_RNA_HFX_S100_KNN$Hydro <- round(hydrophobicity(DEB_RNA_HFX_S100_KNN$Pepseq, scale = "KyteDoolittle",
2)

## Warning: Sequence 31683 has unrecognized amino acid types. Output value might be
## wrong calculated

## Warning: Sequence 35472 has unrecognized amino acid types. Output value might be
## wrong calculated

## Warning: Sequence 36280 has unrecognized amino acid types. Output value might be
## wrong calculated

## Warning: Sequence 37383 has unrecognized amino acid types. Output value might be
## wrong calculated
DEB_RNA_HFX_S100_KNN$mw <- round(mw(DEB_RNA_HFX_S100_KNN$Pepseq, monoisotopic = F),
2)

## Warning: Sequence 31683 has unrecognized amino acid types. Output value might be
## wrong calculated

## Warning: Sequence 35472 has unrecognized amino acid types. Output value might be
## wrong calculated

## Warning: Sequence 36280 has unrecognized amino acid types. Output value might be
## wrong calculated

## Warning: Sequence 37383 has unrecognized amino acid types. Output value might be
## wrong calculated
DEB_RNA_HFX_S100_KNN$cv <- c("35_45")

names(DEB_RNA_HFX_S100_KNN)[names(DEB_RNA_HFX_S100_KNN) == "precursor_m/z"] <- "precursor"
names(DEB_RNA_HFX_S100_KNN)[names(DEB_RNA_HFX_S100_KNN) == "precursor_error_(ppm)"] <- "precursor_err"
DEB_RNA_HFX_S100_KNN <- DEB_RNA_HFX_S100_KNN %>% filter(RNPxl_isPhospho == "0")

DEB_RNA_HFX_S100_KNN_model <- DEB_RNA_HFX_S100_KNN %>% select(colnames(ml_DEB_RNA_train_knn))

```

## Normalizing

```

DEB_RNA_HFX_S100_KNN_model_norm <- DEB_RNA_HFX_S100_KNN_model %>% select_if(is.numeric)
DEB_RNA_HFX_S100_KNN_model_norm <- as.data.frame(lapply(DEB_RNA_HFX_S100_KNN_model_norm,
normalize))
DEB_RNA_HFX_S100_KNN_model_norm[is.na(DEB_RNA_HFX_S100_KNN_model_norm)] <- 0

```

## Classifying DEB RNA HFX data

### KNN5

```
DEB_RNA_HFX_S100_KNN_pred <- knn(train = ml_DEB_RNA_train_knn, test = DEB_RNA_HFX_S100_KNN_model_norm,  
  cl = ml_DEB_RNA_train_knn_labels$Curated, k = 5)  
DEB_RNA_HFX_S100_KNN_model$Prediction <- DEB_RNA_HFX_S100_KNN_pred  
table(DEB_RNA_HFX_S100_KNN_model$Prediction)
```

```
##  
## False True  
## 17616 3365
```

```
DEB_RNA_HFX_S100_KNN$Prediction_KNN <- DEB_RNA_HFX_S100_KNN_pred
```

### C5.0 with boosting and penalty for false positives

penalize False Positives 4 times more than false negatives.

```
matrix_dimensions<- list(c("True", "False"), c("True", "False"))  
names(matrix_dimensions) <- c("predicted", "actual")  
error_cost <- matrix(c(0,1,4,0), nrow = 2, dimnames = matrix_dimensions)
```

```
ml_DEB_RNA_HFX_test_C5<-DEB_RNA_HFX_S100_KNN %>%  
  select(colnames(ml_DEB_RNA_train))
```

```
C5_DEB_RNA_HFX_predict_boost_pen<-predict(C5_model_DEB_RNA_boost_pen, ml_DEB_RNA_HFX_test_C5)
```

```
ml_DEB_RNA_HFX_test_C5$Prediction<-C5_DEB_RNA_HFX_predict_boost_pen  
table(ml_DEB_RNA_HFX_test_C5$Prediction)
```

```
##  
## False True  
## 16238 4743
```

```
DEB_RNA_HFX_S100_KNN$Prediction_C5<-C5_DEB_RNA_HFX_predict_boost_pen
```

### RIPPER

```
RIPPER_DEB_RNA_HFX_predict <- predict(RIPPER_model_DEB_RNA, ml_DEB_RNA_HFX_test_C5)  
ml_DEB_RNA_HFX_test_C5$Prediction_RIPPER <- RIPPER_DEB_RNA_HFX_predict  
table(ml_DEB_RNA_HFX_test_C5$Prediction_RIPPER)
```

```
##  
## False True  
## 15062 5919
```

```
DEB_RNA_HFX_S100_KNN$Prediction_RIPPER <- RIPPER_DEB_RNA_HFX_predict
```

## collapsed Proteins

### S100 HFX charge state 2-7 KNN

```
d<-DEB_RNA_HFX_S100_KNN %>%  
  filter(Prediction_KNN == "True") %>%  
  select(accessions) %>%
```

```
distinct()
nrow(d)
```

```
## [1] 1544
```

### S100 HFX charge state 2-7 C5.0

```
nrow(DEB_RNA_HFX_S100_KNN %>%
  filter(Prediction_C5 == "True") %>%
  select(accessions) %>%
  distinct())
```

```
## [1] 1909
```

### S100 HFX charge state 2-7 RIPPER

```
nrow(DEB_RNA_HFX_S100_KNN %>%
  filter(Prediction_RIPPER == "True") %>%
  select(accessions) %>%
  distinct())
```

```
## [1] 2179
```

### Export KNN predictions

```
HFX_DEB_RNA_S100_KNN_predictions <- DEB_RNA_HFX_S100_KNN %>%
  filter(Prediction_KNN == "True")

write_csv(HFX_DEB_RNA_S100_KNN_predictions, "HFX_DEB_RNA_S100_KNN_predictions.csv")

table(HFX_DEB_RNA_S100_KNN_predictions$accessions) %>%
  as.data.frame() %>%
  arrange(desc(Freq)) %>%
  head(10)
```

```
##           Var1 Freq
## 1 sp|P02359|RS7_ECOLI 38
## 2 sp|P37677|LYXK_ECOLI 23
## 3 sp|P43329|HRPA_ECOLI 23
## 4 sp|P09373|PFLB_ECOLI 18
## 5 sp|Q46888|LTND_ECOLI 16
## 6 sp|P0A7Z4|RPOA_ECOLI 15
## 7 sp|P16525|TUS_ECOLI 15
## 8 sp|P52143|YPJA_ECOLI 15
## 9 sp|P32690|YJBI_ECOLI 14
## 10 sp|P0A7J3|RL10_ECOLI 13
```