### Project

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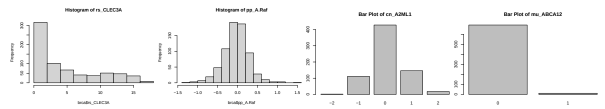
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#### Summary Statistics and Data Preprocessing

#### **Data Overview**

The dataset has 705 observations and 1941 features (1936 predictors and 5 outcomes). There are four different kinds of predictors: rs (gene expression), cn (copy number variations), mu (mutations), and pp (protein levels). Among them, rs and pp are continuous variables, and cn and mu are categorical variables.



#### Remove Missing Values

According to the instruction, we dropped vital.status, and we only considered each response variable as a binary variable. Therefore, we treated the observations that had other outcomes as missing values and removed them from our dataset.

Then the dataset sub had 507 observations and 1940 features.

```
dim(sub)
```

## [1] 507 1940

#### Deal with Multicollinearity

One of the noticeable characteristics of the data is its high dimensionality. There are 1936 predictors, almost four times as many as there are observations. Therefore, it is essential to check correlation.

Since there are four kinds of predictors, it is unlikely that two variables coming from different kinds would be highly correlated. Also, to reduce the computational cost, we split the data into four subsets: rs, cn, mu, and pp, each of which contained only ond kind of predictors.

Then, we created the correlation matrix for each subset, and extracted variables that are highly-correlated with at least one other variable. Take rs as an example. The dataframe idx stores all matrix indices of highly-correlated variables and the corresponding correlation coefficients. If the i-th variable is highly-correlated with the j-th variable, then we only need one of them. Thus, we removed all variables with indices i. For rs, 94 predictors were removed. We applied the same process to the other three subsets. In total, 882 predictors were removed. There are 1059 predictors remained.

```
names(idx) = c("i", "j", "corr")
idx[1:3,]

## i j corr
## 2 3 4 0.94
## 3 5 56 0.84
## 4 9 10 0.95

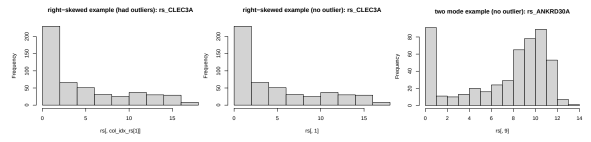
# remove highly-correlated variables
rmv = unique(idx[,1])
length(rmv)
```

## [1] 94

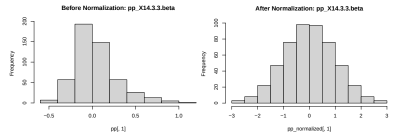
#### Continuous Variables

As mentioned before, **rs** and **pp** are continuous variables, so we should examine if there are any outliers. We first normalized the variable, and stored row and column indices if the data point was three standard deviations away from the mean. For the two subsets, **rs** had 100 outliers, and **pp** had no outlier.

We further looked into rs predictors that included outliers, and we found the vast majority of them had a long tail, mostly right and some left. In addition, a number of rs predictors that did not contain outliers also had a non-standard distribution. As a result, a log transformation of rs predictors would be beneficial.



Unlike rs, pp variables were distributed quite normally. However, many of the variables would contain outliers without normalization. Therefore, we normalized pp variables.

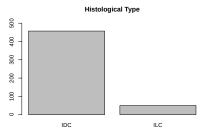


#### Categorical Variables

All four outcomes were more or less imbalanced, among which histological.type was the most imbalanced. Only 10% of the responses were ILC. In some cases, such imbalance would be problematic since models would learn nothing from the minority class. If our data also suffer from problems like this, we should resolve the imbalance by techniques like undersampling. However, if our models perform well enough on current data, no further action needs to be taken.

Fortunately, we trained our models first, and found the models obtained high enough accuracy and AUC scores. Therefore, we decided to not address imbalanced outcomes.

```
##
## Negative Positive
## 0.34714 0.65286
##
## Negative Positive
## 0.2445759 0.7554241
##
## infiltrating ductal carcinoma infiltrating lobular carcinoma
## 0.90138067 0.09861933
```



#### Modeling PR Status

Before modeling, we split the train and test datasets. We used 25% of the samples (126) for testing and 75% of the samples (381) for training.

#### Support Vector Machine (SVM)

The goal of the project was to make classifications. Plus, we needed to alleviate "the curse of dimensionality". Therefore, we should choose classification models that perform well on high-dimensional data. Support vector machines are famous for its capability in high-dimensional spaces, so we first fitted a basic linear SVM, with the default cost = 1 to see how it worked.

As the confusion matrix showed, the in-sample accuracy was 1.0, which implied that we might prefer the linear kernel to the radial kernel.

```
## actual
## predicted Negative Positive
## Negative 133 0
## Positive 0 248
```

Then we constructed two grids of tuning parameters for both linear and radial kernels, and we used 5-fold cross-validation to tune the parameters and to determine which kernel was better.

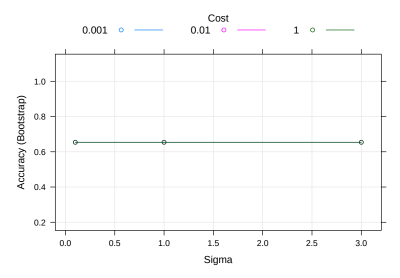
For the linear kernel, the best C was 0.001 with an in-sample accuracy of 0.8324.

```
## C
## 1 0.001

## C Accuracy Kappa AccuracySD KappaSD
## 1 0.001 0.8336029 0.6058770 0.02730027 0.06541415
## 2 0.010 0.7999388 0.5464954 0.02526173 0.06257166
## 3 0.050 0.8014261 0.5503720 0.02442031 0.06150767
## 4 0.100 0.8014261 0.5503720 0.02442031 0.06150767
## 5 0.500 0.8014261 0.5503720 0.02442031 0.06150767
## 6 1.000 0.8014261 0.5503720 0.02442031 0.06150767
```

For the radial kernel, the best C was 0.001 and the best sigma was 3. However, the figure showed that the radial SVM fitted poorly, since the accuracies remained the same and were merely 0.6677. The fact verified our hypothesis that a linear kernel would work better. Thus, we picked the linear SVM with C equals 0.001 to make classifications.

```
## sigma C
## 3 3 0.001
```



We made predictions for the test data, and printed the confusion table below. The accuracy was 0.9048. Thus, the linear SVM performed quite well.

```
## actual
## predicted Negative Positive
## Negative 33 2
## Positive 10 81
## [1] 0.9047619
```

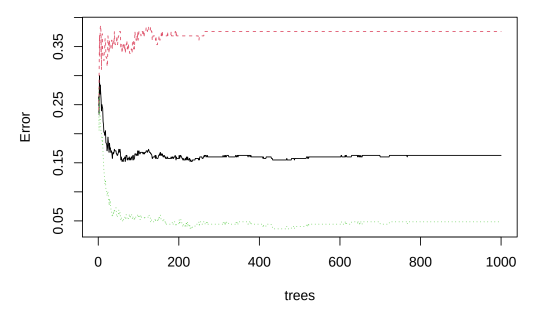
#### **Random Forest**

Random forests are another classification model that is less vulnerable to "the curse of dimensionality". Since there were many parameters that needed to be tuned, we again utilized caret package to cross validate the model.

Before applying cross-validation, we should first determine num.trees, because it could not be included in the grid of parameters. Therefore, we first fitted a random forest using the randomForest() method with ntree = 1000, and plotted the error against trees.

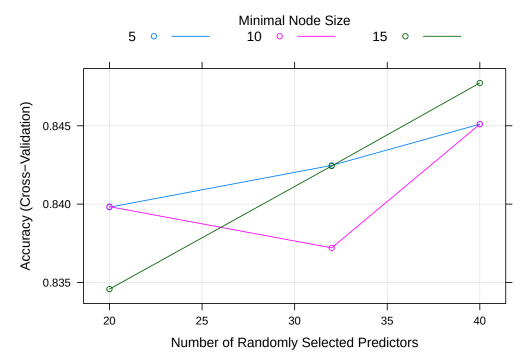
As the plot demonstrated, the error stopped decreasing after the number of trees reached around 500. Thus, ntree = 500 should be sufficient for training.

#### **Random Forest ntree Selection**



In a 5-fold cross-validation, we tuned mtry and min.node.size. The output showed that the best parameters were mtry = 40 and min.node.size = 15 when num.trees = 500 according to the test accuracy.

##		$\mathtt{mtry}$	splitrule	min.node.size				
##	9	40	gini	15				
##		mtry	splitrule	min.node.size	Accuracy	Kappa	${\tt AccuracySD}$	KappaSD
##	1	20	gini	5	0.8398027	0.6199996	0.01816714	0.04744967
##	2	20	gini	10	0.8398378	0.6191721	0.01181501	0.02631224
##	3	20	gini	15	0.8345746	0.6097325	0.01575637	0.03413845
##	4	32	gini	5	0.8424694	0.6272972	0.01375275	0.03444365
##	5	32	gini	10	0.8372062	0.6150392	0.01262152	0.02909492
##	6	32	gini	15	0.8424343	0.6268844	0.01948896	0.05058240
##	7	40	gini	5	0.8451009	0.6341821	0.01487847	0.03698468
##	8	40	gini	10	0.8451009	0.6356210	0.02177987	0.05139481
##	9	40	gini	15	0.8477325	0.6408985	0.01796807	0.04555969



The confusion matrix and the highest test accuracy, 0.9048, were shown here.

```
## actual
## predicted Negative Positive
## Negative 33 2
## Positive 10 81
## [1] 0.9047619
```

# Histological Type (hcluster and knn regression) (est 2-3 pages, pt.20)

To establish the Modeling for histological, We first use logistical regression, However, when we build the model, we found that the logistic model's algorithm is not coverge. The reason that this error occur, because the variable x can divide the reponse variable y into 0 and 1 perfectly. The accurate will be 100%. To solve the problem we decided to use penalized regression. Thus, we choose the modle of logistic regession with ridge penalty

#### Logistic Regression

To performance

```
y = as.factor(sub$histological.type)

y = as.factor(ifelse(y == "infiltrating lobular carcinoma", 1, 0))
sub3 = cbind(rs, cn, mu, pp, y) # cleaned dataset with PR.status as response

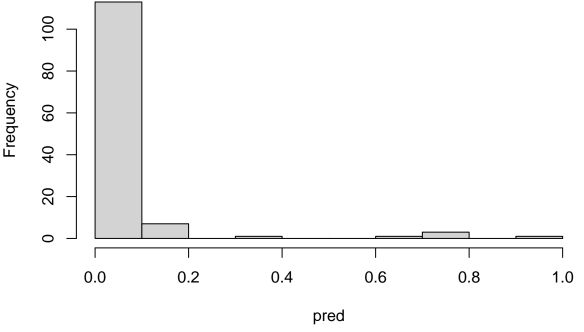
set.seed(651978735)
n = dim(sub)[1]
test_size = as.integer(0.25 * n)
test_idx = sample(1:n, test_size) # 25% of the sample size
```

```
Xtest = sub3[test_idx, -ncol(sub2)]
Xtrain = sub3[-test_idx, -ncol(sub2)]

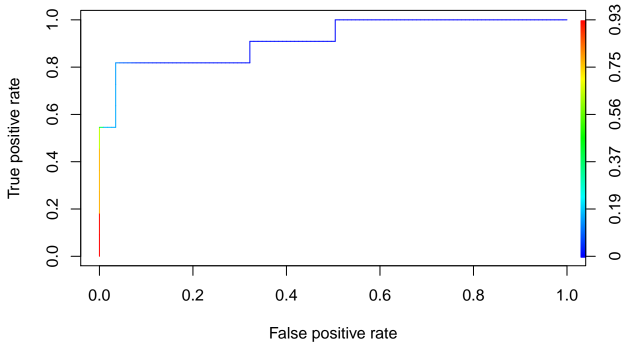
ytest = sub3[test_idx, ncol(sub2)]
ytrain = sub3[-test_idx, ncol(sub2)]
```

logistic regression with ridge penalty and 10 fold corss validation.

#### Histogram of pred



```
library(ROCR)
roc2 <- prediction(pred, ytest)
# calculates the ROC curve
perf2 <- performance(roc2,"tpr","fpr")
plot(perf2,colorize=TRUE)</pre>
```

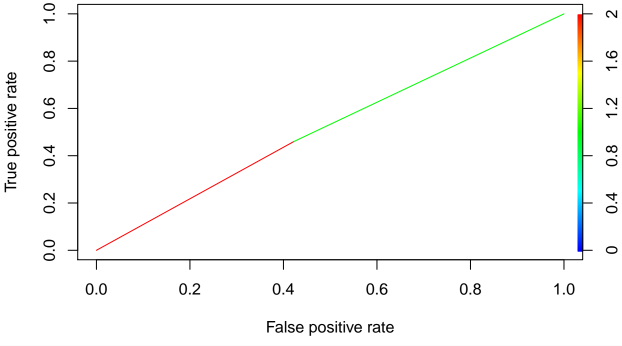


```
performance(roc2, measure = "auc")@y.values[[1]]
```

## [1] 0.915415

#### hcluster clustering

```
\# sub4 = sub3[,-1099]
# kmeanfit <- kmeans(sub4, 2)</pre>
# table((kmeanfit$cluster - 1), sub3$y)
data1 = dist(sub3)
complete_hc <- hclust(data1,method = "complete")</pre>
# plot(complete_hc)
single_hc <- hclust(data1,method = "single")</pre>
# plot(single_hc)
average_hc <- hclust(data1,method = "average")</pre>
# plot(average_hc)
clusters = cutree(complete_hc, k = 2)
table((clusters - 1), sub3$y)
##
##
         0
              1
##
     0 264
            27
##
     1 193
            23
roc2 <- prediction((clusters - 1), sub3$y)</pre>
# calculates the ROC curve
perf2 <- performance(roc2, "tpr", "fpr")</pre>
plot(perf2,colorize=TRUE)
```



```
performance(roc2, measure = "auc")@y.values[[1]]
```

## [1] 0.5188403

## Variable Selection for All Outcomes (random forest?) (est. 2-3 pages. pt.20)

Using Random Forest, select the most important 50 variables, and make predictions based on these variables.

```
impt = rf.fit$importance[order(rf.fit$importance[,3], decreasing=TRUE),][1:50,]
vars = rownames(impt)
# sub4 is the cleaned dataset with all four response variables
sub4 = subset(sub, select = vars)
sub4 = cbind(sub4, sub[1937:1940])
sub4$PR.Status = as.factor(sub4$PR.Status)
sub4$histological.type = as.factor(sub4$histological.type)
sub4$ER.Status = as.factor(sub4$ER.Status)
sub4$HER2.Final.Status = as.factor(sub4$HER2.Final.Status)
Xtest = sub4[test_idx, 1:50]
Xtrain = sub4[-test_idx, 1:50]
ytest = sub4$ER.Status[test_idx]
ytrain = sub4$ER.Status[-test_idx]
svm.fit = svm(ytrain ~., data=Xtrain,
              type="C-classification", kernel="linear", scale=F, cost=1)
table("predicted" = svm.fit$fitted, "actual" = ytrain)
             actual
## predicted Negative Positive
    Negative
                    75
```

```
##
     Positive
                     15
                             282
pred = predict(svm.fit, newdata = Xtest)
table("predicted" = pred, "actual" = ytest)
##
             actual
## predicted Negative Positive
##
     Negative
                     27
     Positive
                               83
##
                      7
library(ROCR)
roc = prediction(as.numeric(pred), ytest)
performance(roc, measure = "auc")@y.values[[1]]
## [1] 0.8481458
perf = performance(roc, "tpr", "fpr")
plot(perf, colorize = T)
      0.8
                                                                                            9
True positive rate
      9.0
      0.4
      0.2
      0.0
             0.0
                           0.2
                                         0.4
                                                        0.6
                                                                      8.0
                                                                                    1.0
                                        False positive rate
rf.fit = randomForest(Xtrain, ytrain,
                       ntree=500,
                       mtry=7,
                       nodesize=10,
                       samplesize=400,
                       importance=TRUE)
pred = predict(rf.fit, Xtest)
table("predicted" = pred, "actual" = ytest)
##
             actual
## predicted Negative Positive
     Negative
                     26
                                3
##
     Positive
                      8
                              89
```

```
roc = prediction(as.numeric(pred), ytest)
performance(roc, measure = "auc")@y.values[[1]]
## [1] 0.8660486
perf = performance(roc, "tpr", "fpr")
plot(perf, colorize = T)
       0.8
                                                                                                2.6
True positive rate
       9.0
       0.4
       0.2
       0.0
                            0.2
             0.0
                                           0.4
                                                          0.6
                                                                         8.0
                                                                                        1.0
                                          False positive rate
```