Logistic Regression For Binary Classification

Introduction

For students who have studied cell biology and biochemistry, logistic regression may already be familiar. It may also be known as a *sigmoidal curve*, *dose-response curve*, *4-parameter fit*, a *median lethal dose curve* (LD-50) or even an exponential growth curve is given limited resources. However, it can also be thought of a binary classifier that can toggle between off and on, zero or one.

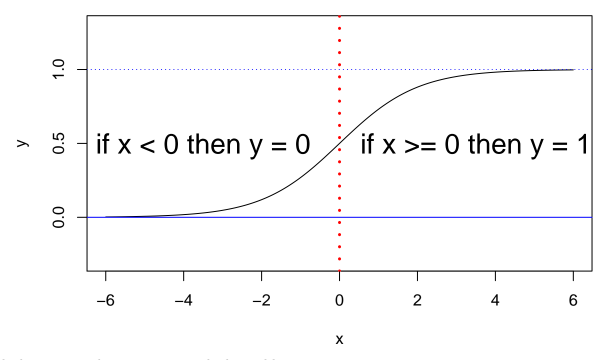
Logistic regression derives its name from the similarity to linear regression, for reasons you will see below. In this circumstance it does not calculate a dependent variable (y) as regression is normally thought to do but instead, Logistic Regression is used for classification between two states as follows.

$$f(x) = \begin{cases} 0 & for \ x < 0 \\ 1 & for \ x \ge 0 \end{cases}$$

In this simple form we can use the generation of a zero or one as a binary or logical value, hence logistic, zero indicating the absence of some quality or item and one indicating its presence.

The domain (x) of our logistic equation may be $(-\infty \text{ to } \infty)$, but at the **decision boundary** of x=0 (for this example) our system changes from *zero*, absence, to *one*, the presence of a quality or item. In the Logistic/Sigmoidal Curve figure below, the **decision boundary** is denoted by the *red dotted line*.

Logistic / Sigmoidal Curve



The logistic growth curve is commonly denoted by:

$$f(x) = \frac{M}{1 + Ae^{-r(x-x_0)}}$$

where M is the curve's maximum value, r is the maximum growth rate (also called the Malthusian parameter¹),

¹https://en.wikipedia.org/wiki/Malthusian_growth_model

 x_0 is the midpoint of the curve, A is the number of times that the initial population must double to reach M^2

In the specific case of Logistic Regression for Binary Classification where we have a probability between 0 and 1, M and A take on the value one.

Since the logistic equation is exponential it is easier to consider working with the formula in terms of the odds or log odds. Odds are the probabilities of success over failure denoted as $\frac{p}{1-p}$ or log-odds as $\ln\left(\frac{p}{1-p}\right)$.

In logistic regression, we find that the log-odds may be expressed as a set of linear equations in x.³ This is simply a transformation of the exponential curve to make it linear. The set of linear equations is similar to the idea inherent in Linear Regression.

$$ln\left(\frac{Pr(y_i = 1|x_i)}{Pr(y_i = 0|x_i)}\right) = \beta_0 + \beta_1 x_1 + \dots + \beta_n x_n$$

Substituting p for $Pr(y_i = 1|x_i)$ and 1-p for $Pr(y_i = 0|x_i)$ we have:

$$ln\left(\frac{p}{1-p}\right) = \sum_{i=1}^{k} \beta_i x_i$$

Eliminate the natural log by taking the exponent on both sides:

$$\frac{p}{1-p} = exp\left(\sum_{i=1}^{k} \beta_{i} x_{i}\right)$$

$$\frac{p}{1-p} = e^u : where u = \sum_{i}^{k} \beta_i x_i$$

Rearrange to solve for p we find:

$$p(u) = \frac{e^u}{1 + e^u}$$

Take the derivative of both sides using quotient rule:

$$p'(u) = \frac{(e^u)(1+e^u) - (e^u)(e^u)}{(1+e^u)^2}$$

Simplify:

$$p'(u) = \frac{e^u}{(1 + e^u)^2}$$

Separate out to produce two fractions:

$$p'(u) = \left(\frac{e^u}{1+e^u}\right) \cdot \left(\frac{1}{1+e^u}\right)$$

²https://en.wikipedia.org/wiki/Logistic_function

³http://juangabrielgomila.com/en/logistic-regression-derivation/

Substitute success and failure variables back into place:

```
p'(u) = p(u) \cdot (1 - p(u))
```

Note: Exploratory Data Analysis (EDA) has been carried out on the file c_m_TRANSFORMED.csv. It can be found in the EDA chapter.******(link to EDA)

```
# Partition data into training and testing sets
set.seed(1000)
index <- createDataPartition(c_m_TRANSFORMED$Class, p = 0.8, list = FALSE)
training_set.1 <- c_m_TRANSFORMED[index, ]</pre>
```

The test.set.1 and Class.test data sets are not produced since that the Logit run with 20 features was not deemed useful. The reason for its dismissal was that is contained extraneous features.

Logit Training #1 using 20 Features

```
set.seed(1000)
registerDoMC(cores = 3) # Start multi-processor mode
start_time <- Sys.time() # Start timer</pre>
# Create model, 10X fold CV repeated 5X
tcontrol <- trainControl(method = "repeatedcv",</pre>
                          number = 10,
                          repeats = 5)
model_obj.1 <- train(Class ~ .,</pre>
                      data = training_set.1,
                      trControl = tcontrol,
                      method = "glm",
                      family = "binomial")
end_time <- Sys.time() # End timer</pre>
end_time - start_time # Display time
## Time difference of 1.695754 secs
registerDoSEQ() # Stop multi-processor mode
```

Logit Results #1

```
summary(model_obj.1)
##
## Call:
## NULL
##
## Deviance Residuals:
##
       Min
                 1Q
                      Median
                                    3Q
                                            Max
##
  -5.9372
           -0.2835
                     -0.0194
                                0.0516
                                         3.6884
##
## Coefficients:
##
               Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                 8.0525
                             9.2156
                                      0.874 0.382234
## A
                 5.0438
                             9.6899
                                      0.521 0.602699
## C
               -14.2228
                             2.6949
                                     -5.278 1.31e-07 ***
## D
               -36.2676
                             8.0845
                                     -4.486 7.25e-06 ***
## E
                27.6016
                            11.1292
                                      2.480 0.013135 *
## F
                             5.2654
                                      1.067 0.286034
                 5.6174
## G
               -22.1970
                            10.3043
                                     -2.154 0.031229 *
## H
                90.1101
                            12.1105
                                      7.441 1.00e-13 ***
## I
                -5.9795
                             4.3945
                                     -1.361 0.173610
                -2.8961
                                     -0.294 0.768669
## K
                             9.8468
## L
                -3.7417
                             9.2217
                                     -0.406 0.684926
                            12.0747
                                     -0.012 0.990570
## M
                -0.1427
                 3.3478
                             9.6749
                                      0.346 0.729319
## N
## P
               -39.7466
                            11.1010
                                     -3.580 0.000343 ***
## Q
                -5.6804
                            11.2516
                                     -0.505 0.613664
               -83.6045
                            11.8104
                                     -7.079 1.45e-12 ***
## R.
## S
                -9.9745
                            10.0872
                                     -0.989 0.322750
## T
               -36.5980
                             9.2791
                                     -3.944 8.01e-05 ***
## V
                16.3411
                             9.7859
                                      1.670 0.094946
                            13.8870
                                      0.649 0.516141
## W
                 9.0169
## Y
               -31.9282
                            11.1167
                                     -2.872 0.004078 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
##
  (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 2593.68
                                on 1872
                                         degrees of freedom
## Residual deviance: 657.72
                                on 1852 degrees of freedom
## AIC: 699.72
##
## Number of Fisher Scoring iterations: 8
```

The Akaike information criterion (AIC) for model #1 is 699.72. This will be used later to compare the models generated to rate their ability to best utilize the features. - The list of probabilities for the estimates leaves us with only 9 important features to try re-modeling, R, H, P, C, E, Y, T, D, G.

Logit Training #2 using 9 Features

• Using **ONLY** features: (R, H, P, C, E, Y, T, D, G)

```
# Data import & handling
c_m_9aa <- read_csv("../00-data/02-aac_dpc_values/c_m_TRANSFORMED.csv",</pre>
                      col_types = cols(Class = col_factor(levels = c("0", "1")),
                                        A = col_skip(),
                                        F = col_skip(),
                                        I = col_skip(),
                                        K = col_skip(),
                                        L = col_skip(),
                                        M = col skip(),
                                        N = col_skip(),
                                        PID = col_skip(),
                                        Q = col_skip(),
                                        V = col_skip(),
                                        S = col_skip(),
                                        TotalAA = col_skip(),
                                        W = col_skip()))
# Partition data into training and testing sets
set.seed(1000)
index <- createDataPartition(c_m_9aa$Class, p = 0.8, list = FALSE)</pre>
training_set.2 <- c_m_9aa[ index, ]</pre>
test_set.2
            <- c_m_9aa[-index, ]</pre>
Class_test.2 <- as.factor(test_set.2$Class)</pre>
```

Logit Training #2 with 9 Features

```
set.seed(1000)
registerDoMC(cores = 3) # Start multi-core
start_time <- Sys.time() # Start timer</pre>
# Create model, 10X fold CV repeated 5X
fitControl <- trainControl(method = "repeatedcv",</pre>
                            number = 10,
                            repeats = 5,
                            savePredictions = "final") # IMPORTANT: Saves predictions
model_obj.2 <- train(Class ~ .,</pre>
                      data = training_set.2,
                      trControl = fitControl,
                      method = "glm",
                      family = "binomial")
end_time <- Sys.time()</pre>
                         # End timer
end_time - start_time
                          # Display time
```

```
## Time difference of 1.39758 secs
registerDoSEQ()  # Stop multi-core
```

Logit Summary #2

```
summary(model_obj.2)
```

```
##
## Call:
## NULL
##
## Deviance Residuals:
      Min
                1Q
                     Median
                                  3Q
                                          Max
## -6.2083 -0.2984 -0.0204
                              0.0601
                                       3.5666
## Coefficients:
##
              Estimate Std. Error z value Pr(>|z|)
                 8.306
                            1.007
## (Intercept)
                                   8.245 < 2e-16 ***
               -14.755
                            1.908 -7.733 1.05e-14 ***
## D
               -31.411
                            4.949 -6.347 2.20e-10 ***
## E
                            5.092
                                   4.307 1.66e-05 ***
                21.932
## G
               -23.259
                            5.071 -4.587 4.49e-06 ***
## H
                94.580
                            8.431 11.218 < 2e-16 ***
## P
               -29.394
                            6.264 -4.692 2.70e-06 ***
## R
               -82.809
                            6.363 -13.015 < 2e-16 ***
## T
               -40.915
                            5.624 -7.275 3.45e-13 ***
## Y
               -37.860
                            6.291 -6.018 1.77e-09 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 2593.68 on 1872 degrees of freedom
## Residual deviance: 688.96 on 1863 degrees of freedom
## AIC: 708.96
## Number of Fisher Scoring iterations: 8
```

Logit Confusion Matrix #2

```
Predicted_test_vals <- predict(model_obj.2, test_set.2[, -1])</pre>
confusionMatrix(Predicted_test_vals, Class_test.2, positive = "1")
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction
            0 235 18
##
                8 206
##
##
##
                  Accuracy: 0.9443
##
                    95% CI: (0.9195, 0.9633)
##
       No Information Rate: 0.5203
##
       P-Value [Acc > NIR] : < 2e-16
##
##
                     Kappa: 0.8883
##
##
    Mcnemar's Test P-Value: 0.07756
##
##
               Sensitivity: 0.9196
##
               Specificity: 0.9671
##
            Pos Pred Value: 0.9626
            Neg Pred Value: 0.9289
##
##
                Prevalence: 0.4797
            Detection Rate: 0.4411
##
##
      Detection Prevalence: 0.4582
##
         Balanced Accuracy: 0.9434
##
##
          'Positive' Class : 1
```

- The Akaike information criterion (AIC) for model #2 is 708.96. This will be used later to compare the models generated to rate their ability to best utilize the features.
- The number of unique false-positives and false-negatives is 26.

Obtain List of False Positives & False Negatives

536 proteins listed although they are NOT UNIQUE NOR SORTED.

Conclusion

Logit is easy to implement and understand and can be used for parameter importance measurements.

Considering the Table Logit Models, below, it is clear that model #2 with 9 features best describes the better of the two models.

Akaike Information Criterion:

$$AIC = 2K - ln(\widehat{L})$$

Where $ln(\widehat{L})$ is the log-likelihood, K is the number of parameters.

Model #	Features	AIC
1	20	699.72
2	9	708.96

Logit is a common machine learning method. It is easy to understand and explain. This supervised binary classification method is very useful for determining the importance of the features which can be applied. As we saw in Model#1 there were 11 features that had probabilities of the estimates used above the 5% threshold cut-off. In Model#2 only 9 features were used to describe the model and the AIC increased by 9.24.

The nine features which best described the logistic regression model were R, H, P, C, E, Y, T, D, G. If we compare this to the Boruta test carried out in the EDA we find the overlap interesting.

Test Model / Order of Importance (From Left To Right)

Test Model																				
Order	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20
Boruta	\mathbf{R}	Η	Ρ	K	\mathbf{C}	\mathbf{E}	Y	\mathbf{T}	\mathbf{S}	A	V	U	Ι	F	D	G	N	\mathbf{L}	M	Q
Logit	\mathbf{R}	Η	Ρ		\mathbf{C}	\mathbf{E}	Y	\mathbf{T}							D	G				

The first 7 out of 8 amino acid features are seen in the proper order as described by the Boruta Random Forest model. This is confirmation that Logit can pick up the importance of features similar to Boruta.

Logit produced 536 proteins which are false-negatives or false-positives. It should be noted that the 536 are NOT UNIQUE NOR SORTED. The number of FN/FP from the confusion matrix is 26. These proteins will be investigated further in the Outliers section which compares these FN/FP proteins to the PCA outliers.