$\operatorname{MSDS6372}$ Project 2 - Osteoporosis in Women

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Run Normal Logit Model with Identified Predictors
Add Interactions to Normal logit
Running Random Forest Fit
Running Conditional Random Forest Fit
LDA AND QDA Model fit
*** Appendix C: Test interaction - LDA ===================================
The main effects model
Add to main effects model
Final Interaction Model

Objective 1 - EDA and logistic regression model

Display the ability to perform EDA and build a logistic regression model

• Perform your logistic regression analysis and provide interpretation of the regression coefficients including hypothesis testing, and confidence intervals. For simplicity sake, you do not need to include interactions with this model. Comment on the practical vs statistical significance of the deemed important factors...

Logistical Considerations.

• Just like last time, this does not have to be extremely fancy in terms of the model building approach, let EDA, feature selection, and overall intuition guide you.

Introduction

Data Description

The data set provided is about predicting whether a woman with osteoporosis will have another bone fracture. Of course getting a bone fracture is somewhat circumstantial, but with this disease every day life could trigger a break if the progression of the disease is strong.

The dataset included a total of 14 variables: 3 ID variables which tell us the subject, doctor and physical location of each record, 4 continuous variables (BMI, Weight, Height, and Age), 6 categorical variables (PRIORFRAC, PREMENO, MOMFRAC, PREMEO, MOMFRAC, ARMASSIST, SMOKE, RATERISK), and the response (FRACTURE). We were unable to find a mapping the subjects with their location to understand the mix of countries represented.

We have 500 subjects in the dataset of which 33% of the subjects have/had fractures.

Missing values were not detected in dataset. Special characters were removed from column headings. What we know/don't know about the sample (500)

Exploratory Analysis

Assumptions

This is a prospective study which means it's a study over time of a group of similar individuals who differ with respect to certain factors under a study and how these factors affect rates of a certain outcome (Fracture vs No-Fracture) Linearity - Independence of errors - Based on SUB ID(Subject ID) we confirm each record is an independent sample. Multicollinearity - Weight and BMI are highly correlated but we will remove one from the analysis.

GLOW dataset:

Variable Name Type

```
integer 500 - Identification Code (1 - n)
    SUB_ID
  SITE ID
           integer
                      6 - Study Site (1 - 6)
   PHY ID
           integer 127 - Physician ID code (128 unique codes)
PRIORFRAC*
            factor
                      2 - History of Prior Fracture (1: No, 2: Yes)
                    36 - Age at Enrollment (Years)
      AGE integer
    WEIGHT
           numeric 128 - Weight at enrollment (Kilograms)
   HEIGHT
           integer
                    34 - Height at enrollment (Centimeters)
      BMI numeric 409 - Body Mass Index (Kg/m^2)
 PREMENO*
                      2 - Menopause before age 45 (1: No, 2: Yes)
```

#Unique

factor

```
MOMFRAC*
                       2 - Mother had hip fracture (1: No, 2: Yes)
             factor
                       2 - Arms are needed to stand from a chair (1: No, 2: Yes)
ARMASSIST*
             factor
                       2 - Former or current smoker (1: No, 2: Yes)
   SMOKE*
             factor
RATERISK*
                       3 - Self-reported risk of fracture (1: Less, 2: Same, 3: Greater)
             factor
FRACSCORE
           integer
                      12 - Fracture Risk Score (Composite Risk Score)
FRACTURE*
             factor
                       2 - Any fracture in first year (1: No, 2: Yes)
```

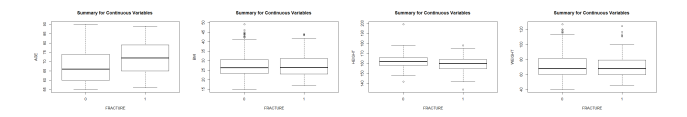


Figure 1 - Boxplots for Continuous Variables AGE, BMI, HEIGHT, WEIGHT

In Figure 1, we see the boxplots for the continuous variables AGE, WEIGHT, HEIGHT, BMI.

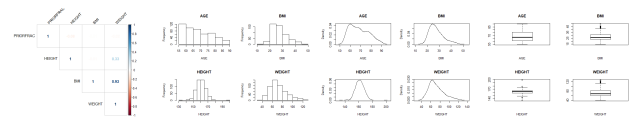


Figure 2 - Correlation, and Density plots for Continuous Variables AGE, BMI, HEIGHT, WEIGHT

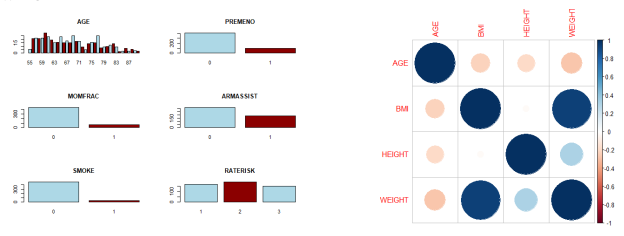


Figure 3 - Barplot (occurrances) and Multivariate Plots for Categorical and Continuous Variables

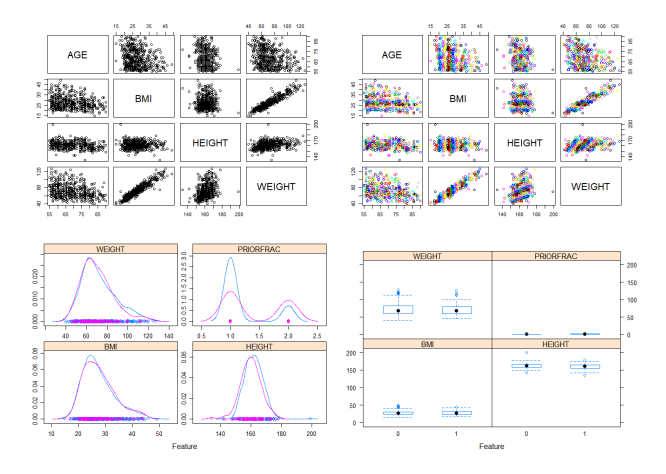


Figure 4 - Scatterplots

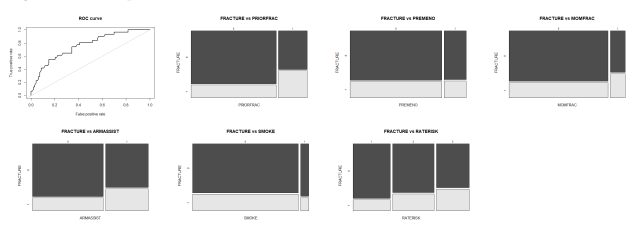


Figure 5 - ROC and 2-way Tables

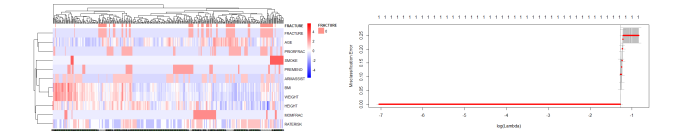


Figure 6 - Clustering

Restatement of Problem and the overall approach to solve it.

Model Selection

Checking Assumptions

Parameter Interpretation

Final conclusions from the analysis of Objective 1

to do:

take writeup from EDA and place here with reference to figures, etc

Objective 2 - additional competing models

- Record the predictive performance metrics from your simple, highly interpretable model from Objective 1.
- You must include one additional model which is also a more complicated logistic regression model than in Objective 1. By complicated, I do not mean that you include more predictors (that will be somewhat sorted out in Objective 1), but rather model complexity through interaction terms, new variables created by the group, or transformations.
- Create another competing model using just the continuous predictors and use LDA or QDA.
- Use a nonparameteric model approach as a competing model. Random forest for predictors that are both categorical and continuous or a k-nearest neighbors approach if just working with continuous predictors.
- Provide a summary table of the performance across the competing methods. Summarize the overall findings. A really great report will also give insight as to why the "best" model won out. This is where a thorough EDA will always help.

Record predictive performance from objective 1

Another competing model (interaction terms)

Another competing model (just continuous) LDA

nonparametric model (random forest)

another nonparametric model (conditional random forest)

Summary table of performance

Model	Predictors	Accurac	$\mathrm{cy}95\%$ CI	Sensitivi	ty Specificity	AUC
Logistic Regression (logit)	7	74.5%	(66.7%, 81.3%)	94.6%	13.5%	67.8%
Logistic Regression w/Interactions (logit)	7+3 interactions	75.2%	(67.4%, 81.9%)	92.9%	21.6%	72.2%
Random Forest*	10	74.5%	(66.7%, 81.3%)	99.1%	0.0%	74.2%
*RF Lower Cutoff (decreasing the probability from 50% to 30%)	10	73.8%	(66%, 80.7%)	91.1%	21.6%	
Conditional Random Forest	10	75.2%	(67.4%, 81.9%)	94.6%	16.2%	69.3%
LDA	4	73.2%	(65.3%, 80.1%)	94.6%	8.1%	60.2%

Conclusion/Discussion

to do:

take writeup from EDA and place here with reference to figures, etc

Data Set 1: Osteoporosis in Women

From Hosmer, Lemeshow, and Sturdivant (2013), Applied Logistic Regression, 3rd Edition. The Global Longitudinal Study of Osteoporosis in Women (GLOW) is an international study of osteoporosis in women aged 55 years and over. The major goals of the study are to examine prevention and treatment of fractures and distribution of risk factors among older women. Complete details on the study as well as a list of GLOW publications may be found at the Center for Outcomes Research web site, http://www.outcomes-umassmed.org/glow. There are over 60K observations in the original data set. This data set contains a sample of 500 of them. The link below is to a website with the data set and description of the variables. The data set in question is called "glow500".

https://www.umass.edu/statdata/statdata/glow/index.html Note: If you choose this data set, you MAY NOT use the Hosmer, Lemeshow, and Sturdivant text to help you in your analysis. You may only use Chapter 1 in order to obtain a description of the data.

Of course if you dont have the book

https://www.umass.edu/statdata/statdata/data/glow/glow.pdf provides definitions to the variables.

The Global Longitudinal Study of Osteoporosis in Women (GLOW) (2005-2014) was a prospective cohort study of physician practices in the provision of prophylaxis and treatment against osteoporotic fractures. The goal of this research was to improve understanding of the risk and prevention of osteoporosis-related fractures among female residents of 10 countries who were 55 years of age and older. GLOW enrolled over 60,000 women through over 700 physicians in 10 countries, and conducted annual follow-up for up to 5 years through annual patient questionnaires.

Setup:

Data Import and Cleaning

Missing values were not detected in dataset. Special characters were removed from column headings. What we know/don't know about the sample (500): 1. We do not know if the subjects are distributed equally around the world. We will assume that the same percentage from each region was selected for the sample in this dataset. 2. Based on the Sub_ID(Subject ID), we can assume that the datat is independent sample of participants.

```
glow_data_file <- here::here("data", "glow500.csv")
dataset_loc <-
dataset <- read.csv(glow_data_file, sep=",", stringsAsFactors = TRUE, header=TRUE,na.strings=c(""))
# List rows of data that have missing values
Missing_values <- dataset[!complete.cases(dataset),]
# Create new dataset without missing data
dataset <- na.omit(dataset)
#remove FRACSCORE feature per professor Turner
drops <- c("FRACSCORE")
dataset <- dataset[, !(names(dataset) %in% drops)]</pre>
```

```
#Cleanup column names
colnames(dataset)[colnames(dataset)=="i..SUB_ID"] <- "SUB_ID"</pre>
#set categorical variables as factors
dataset$PRIORFRAC <- factor(dataset$PRIORFRAC,labels=c("0","1"))</pre>
dataset$PREMENO <- factor(dataset$PREMENO,labels=c("0","1"))</pre>
dataset$MOMFRAC <- factor(dataset$MOMFRAC,labels=c("0","1"))</pre>
dataset$ARMASSIST <- factor(dataset$ARMASSIST,labels=c("0","1"))</pre>
dataset$SMOKE <- factor(dataset$SMOKE,labels=c("0","1"))</pre>
dataset$RATERISK <- factor(dataset$RATERISK,labels=c("1","2","3"))</pre>
dataset$FRACTURE <- factor(dataset$FRACTURE,labels=c("0","1"))</pre>
#rearrange columns
dataset <- dataset[c("SUB_ID", "SITE_ID", "PHY_ID", "AGE", "BMI", "HEIGHT", "WEIGHT", "PRIORFRAC", "PREMENO", "M</pre>
str(dataset)
## 'data.frame':
                   500 obs. of 14 variables:
## $ SUB_ID : int 1 2 3 4 5 6 7 8 9 10 ...
## $ SITE_ID : int 1 4 6 6 1 5 5 1 1 4 ...
## $ PHY_ID : int 14 284 305 309 37 299 302 36 8 282 ...
## $ AGE
            : int 62 65 88 82 61 67 84 82 86 58 ...
## $ BMI : num 28.2 34 20.6 24.3 29.4 ...
## $ HEIGHT : int 158 160 157 160 152 161 150 153 156 166 ...
## $ WEIGHT : num 70.3 87.1 50.8 62.1 68 68 50.8 40.8 62.6 63.5 ...
## $ PRIORFRAC: Factor w/ 2 levels "0","1": 1 1 2 1 1 2 1 2 2 1 ...
## $ PREMENO : Factor w/ 2 levels "0","1": 1 1 1 1 1 1 1 1 1 1 ...
## $ MOMFRAC : Factor w/ 2 levels "0","1": 1 1 2 1 1 1 1 1 1 1 ...
## $ ARMASSIST: Factor w/ 2 levels "0","1": 1 1 2 1 1 1 1 1 1 1 ...
## $ SMOKE : Factor w/ 2 levels "0", "1": 1 1 1 1 1 2 1 1 1 1 ...
## $ FRACTURE : Factor w/ 2 levels "0","1": 1 1 1 1 1 1 1 1 1 1 1 ...
```

Exploratory Data Analysis

Grouping Variables as Continuous, Categorical, and ID

```
numericVar <- dataset[,4:7]
ID_var <- dataset[,c(1:3)]
set_noID <- dataset[4:14]
categoricalVar <- dataset[8:14]</pre>
```

Create Train and Validation Datasets

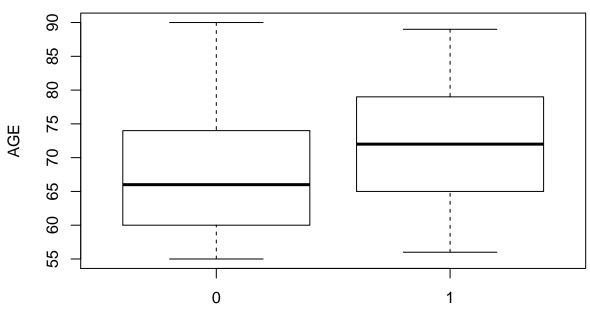
```
validation_index = createDataPartition(dataset$FRACTURE, p=0.70, list=FALSE)
validationData = dataset[-validation_index,c(4:14)]
trainingData = dataset[validation_index,c(4:14)]
```

Summary Statistics

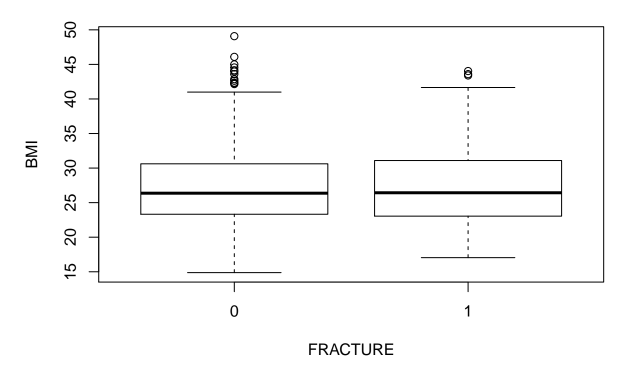
Assumptions This is a prospective study which means its a study over time of a group of similar individuals who differ with respect to certain factors under a study and how these factors affect rates of a certain outcome (Fracture vs No-Fracture) Linearity - Independence of errors - Based on SUB_ID(Subject ID) we confirm each record is an independent sample. Multicollinearity - Weight and BMI are highly correlated but we will remove one from the

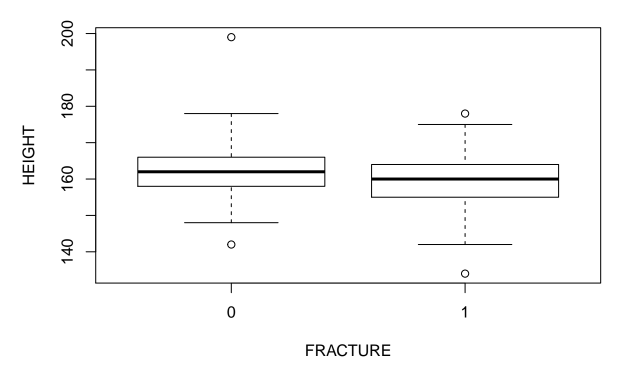
```
#Summary stats by groups for continous predictors
t(aggregate(AGE~FRACTURE, data=dataset, summary))
##
                           [,2]
               [,1]
               "0"
                           "1"
## FRACTURE
               "55.00000" "56.00000"
## AGE.Min.
## AGE.1st Qu. "60.00000" "65.00000"
               "66.00000" "72.00000"
## AGE.Median
## AGE.Mean
               "67.48533" "71.79200"
## AGE.3rd Qu. "74.00000" "79.00000"
               "90.00000" "89.00000"
## AGE.Max.
t(aggregate(BMI~FRACTURE,data=dataset,summary))
##
               [,1]
                           [,2]
## FRACTURE
               "0"
                           "1"
## BMI.Min.
               "14.87637" "17.04223"
## BMI.1st Qu. "23.32087" "23.04688"
## BMI.Median "26.36709" "26.43080"
               "27.50140" "27.70793"
## BMI.Mean
## BMI.3rd Qu. "30.61756" "31.09282"
## BMI.Max.
               "49.08241" "44.03628"
t(aggregate(WEIGHT~FRACTURE,data=dataset,summary))
##
                  [,1]
                               [,2]
                               "1"
## FRACTURE
                  " 39.90000" " 45.80000"
## WEIGHT.Min.
## WEIGHT.1st Qu. " 60.30000" " 59.90000"
                  " 68.00000" " 68.00000"
## WEIGHT.Median
                  " 72.16693" " 70.79200"
## WEIGHT.Mean
## WEIGHT.3rd Qu. " 81.60000" " 79.40000"
                  "127.00000" "124.70000"
## WEIGHT.Max.
t(aggregate(HEIGHT~FRACTURE,data=dataset,summary))
##
                   [,1]
                             [,2]
                  "0"
                             "1"
## FRACTURE
## HEIGHT.Min.
                  "142.000" "134.000"
## HEIGHT.1st Qu. "158.000" "155.000"
## HEIGHT.Median
                  "162.000" "160.000"
## HEIGHT.Mean
                  "161.864" "159.864"
## HEIGHT.3rd Qu. "166.000" "164.000"
                  "199.000" "178.000"
## HEIGHT.Max.
```

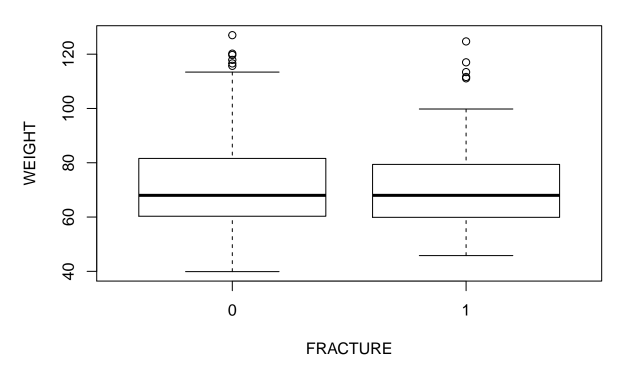
```
#par(mfrow=c(2,2)) # put four figures in a row (2*4)
for (i in 4:7) {
  boxplot(dataset[,i] ~ dataset$FRACTURE,ylab=names(dataset)[i],xlab="FRACTURE", main="Summary for Cont}
}
```



FRACTURE







```
#create an nicer summary table
index<-which(sapply(dataset,is.numeric))
tab.cont<-c()
for (i in index){
   tab.cont<-rbind(tab.cont,summary(dataset[,i]))
}
rownames(tab.cont)<-names(dataset)[index]
View(tab.cont)
tab.cont</pre>
### Min. 1st Qu. Median Mean 3rd Qu. Max.
```

```
## SUB_ID
            1.00000 125.75000 250.50000 250.50000 375.25000 500.00000
## SITE_ID
            1.00000
                      2.00000
                                3.00000
                                          3.43600
                                                   5.00000
## PHY_ID
            1.00000 57.75000 182.50000 178.55000 298.00000 325.00000
## AGE
           55.00000 61.00000 67.00000 68.56200
                                                  76.00000 90.00000
## BMI
           14.87637 23.26889
                               26.41898
                                        27.55303
                                                  30.79205 49.08241
## HEIGHT
          134.00000 157.00000 161.50000 161.36400 165.00000 199.00000
           39.90000 59.90000 68.00000 71.82320 81.30000 127.00000
## WEIGHT
```

```
# display the first 20 rows
print(head(dataset, n=20))
```

```
## SUB_ID SITE_ID PHY_ID AGE BMI HEIGHT WEIGHT PRIORFRAC PREMENO ## 1 1 14 62 28.16055 158 70.3 0 0 ## 2 2 4 284 65 34.02344 160 87.1 0 0
```

```
## 3
                          305 88 20.60936
                                                157
                                                       50.8
                     6
                                                                      1
                                                                              0
## 4
            4
                     6
                          309
                                82 24.25781
                                                160
                                                       62.1
                                                                      0
                                                                              0
## 5
            5
                                61 29.43213
                                                       68.0
                           37
                                                152
                                                                      0
                                                                              0
## 6
            6
                                67 26.23356
                                                       68.0
                                                                              0
                     5
                          299
                                                161
                                                                      1
            7
## 7
                     5
                          302
                                84 22.57778
                                                150
                                                       50.8
                                                                      0
                                                                              0
## 8
            8
                     1
                           36
                                82 17.42919
                                                153
                                                       40.8
                                                                      1
                                                                              0
## 9
            9
                     1
                            8
                                86 25.72321
                                                156
                                                       62.6
                                                                      1
                                                                              0
                                58 23.04398
## 10
           10
                                                166
                                                       63.5
                                                                      0
                     4
                          282
                                                                              0
## 11
           11
                     6
                          315
                                67 28.87778
                                                153
                                                       67.6
                                                                      0
                                                                              0
## 12
           12
                           34
                                56 42.27473
                                                167
                                                     117.9
                                                                      0
                                                                              0
                     1
## 13
           13
                     6
                          315
                               59 25.56775
                                                162
                                                       67.1
                                                                      0
                                                                              0
                               72 21.15702
                                                                              0
## 14
           14
                           33
                                                165
                                                       57.6
                                                                      0
                     1
## 15
                           23
                                64 23.90625
           15
                     1
                                                160
                                                       61.2
                                                                      0
                                                                              1
## 16
           16
                     3
                          179
                                68 30.09143
                                                161
                                                       78.0
                                                                      0
                                                                              0
## 17
           17
                     4
                          284
                                67 38.82461
                                                165
                                                      105.7
                                                                      0
                                                                              0
## 18
           18
                     4
                          283
                                69 25.07240
                                                162
                                                       65.8
                                                                      0
                                                                              0
## 19
           19
                     3
                          179
                                78 31.09282
                                                162
                                                       81.6
                                                                              0
                                                                      1
                                60 23.00296
                                                                      0
                                                                              0
## 20
           20
                          313
                                                157
                                                       56.7
##
      MOMFRAC ARMASSIST SMOKE RATERISK FRACTURE
                                         2
## 1
             0
                        0
                               0
                                                   0
## 2
             0
                        0
                               0
                                         2
                                                   0
## 3
             1
                        1
                               0
                                         1
                                                   0
             0
                        0
                               0
## 4
                                         1
                                                   0
## 5
             0
                        0
                               0
                                         2
                                                   0
             0
                        0
                                         2
## 6
                               1
                                                   0
## 7
             0
                        0
                               0
                                         1
                                                   0
## 8
             0
                        0
                               0
                                         2
                                                   0
## 9
             0
                        0
                               0
                                         2
                                                   0
## 10
             0
                        0
                               0
                                         1
                                                   0
## 11
                        0
                                         1
             1
                               1
                                                   0
                                         2
## 12
             0
                        1
                               1
                                                   0
## 13
             0
                        0
                               1
                                         1
                                                   0
## 14
             0
                               0
                                         1
                                                   0
                        1
                                         2
## 15
             0
                        0
                               0
                                                   0
## 16
             0
                        1
                               0
                                         1
                                                   0
## 17
             0
                        0
                               0
                                         1
                                                   0
## 18
             0
                        0
                               0
                                         2
                                                   0
## 19
             0
                        1
                               0
                                         3
                                                   0
                                         2
## 20
             0
                        0
                               0
                                                   0
```

```
# display the dimensions of the dataset
print(dim(dataset))
```

```
## [1] 500 14
```

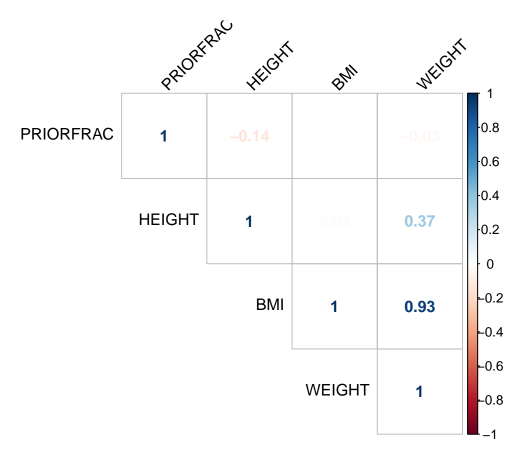
```
# list types for each attribute
print(sapply(dataset, class))
```

```
SUB_ID
              SITE_ID
                         PHY_ID
                                       AGE
                                                 BMI
                                                        HEIGHT
                                                                  WEIGHT
## "integer" "integer" "integer" "integer" "numeric" "integer" "numeric"
## PRIORFRAC
              PREMENO
                        MOMFRAC ARMASSIST
                                               SMOKE
                                                      RATERISK
                                                                FRACTURE
  "factor"
             "factor"
                       "factor" "factor"
                                           "factor"
                                                      "factor"
                                                                "factor"
```

```
# Standard Deviations for the non-categorical columns
std=sapply(set_noID,sd)
## Warning in var(if (is.vector(x) || is.factor(x)) x else as.double(x), na.rm = na.rm): Calling var(x)
    Use something like 'all(duplicated(x)[-1L])' to test for a constant vector.
## Warning in var(if (is.vector(x) || is.factor(x)) x else as.double(x), na.rm = na.rm): Calling var(x)
    Use something like 'all(duplicated(x)[-1L])' to test for a constant vector.
## Warning in var(if (is.vector(x) || is.factor(x)) x else as.double(x), na.rm = na.rm): Calling var(x)
    Use something like 'all(duplicated(x)[-1L])' to test for a constant vector.
## Warning in var(if (is.vector(x) || is.factor(x)) x else as.double(x), na.rm = na.rm): Calling var(x)
    Use something like 'all(duplicated(x)[-1L])' to test for a constant vector.
## Warning in var(if (is.vector(x) || is.factor(x)) x else as.double(x), na.rm = na.rm): Calling var(x)
    Use something like 'all(duplicated(x)[-1L])' to test for a constant vector.
## Warning in var(if (is.vector(x) || is.factor(x)) x else as.double(x), na.rm = na.rm): Calling var(x)
    Use something like 'all(duplicated(x)[-1L])' to test for a constant vector.
## Warning in var(if (is.vector(x) || is.factor(x)) x else as.double(x), na.rm = na.rm): Calling var(x)
    Use something like 'all(duplicated(x)[-1L])' to test for a constant vector.
print('The standard deviations are:')
## [1] "The standard deviations are:"
print(std)
##
         AGF.
                    BMT
                            HEIGHT
                                       WEIGHT PRIORFRAC
                                                           PREMENO
##
   8.9895372 5.9739583 6.3554928 16.4359918 0.4345961 0.3958249
##
     MOMFRAC ARMASSIST
                             SMOKE
                                    RATERISK
                                               FRACTURE
```

Correlations

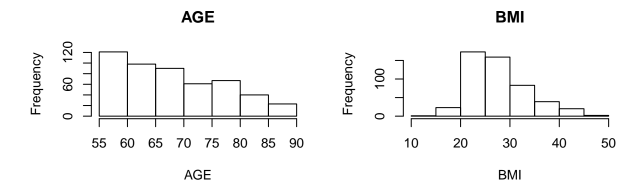
BMI and Weight show to be highly correlation which makes sense since weight is a factor in calculation of BMI. We will remove Weight from models in order to meet assumptions.

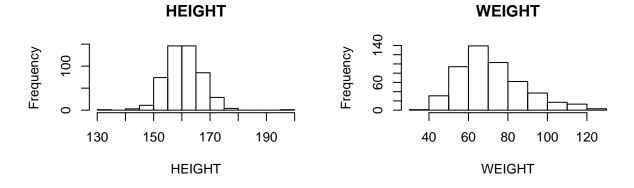


Visualization of Continuous Variables For the categorical variables, we show an unbalanced dataset of subjects with majority false PRIORFRAC, PREMENO, MOMFRAC, ARMASSIST, and SMOKE. There was a good balance of subjects in the 3 levels of RATERISK. An unblanced dataset will cause a model to favor the skewed numbers.

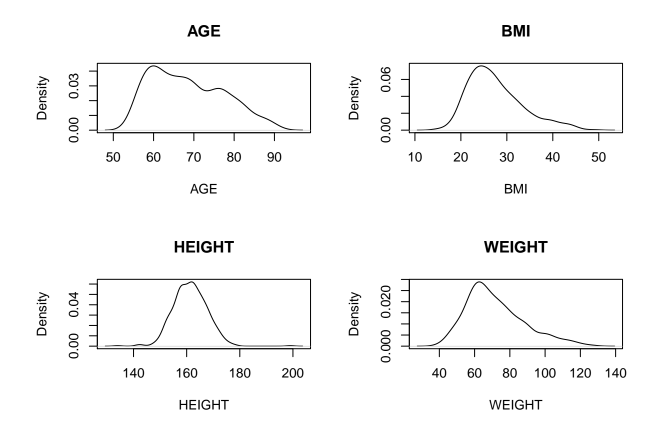
For the continous variables, we can see that BMI and Weight are highly correlated and weight and height are also correlated. When building the model, we will remove Weight as to meet the assumptions of logistic regression.

```
# Data visualizations
dataset_numeric = numericVar
#Histograms
par(mfrow=c(2,2)) # put four figures in a row (2*4)
for (i in 1:4) {
   hist(dataset_numeric[,i],xlab=names(dataset_numeric)[i],main=names(dataset_numeric)[i])
}
```

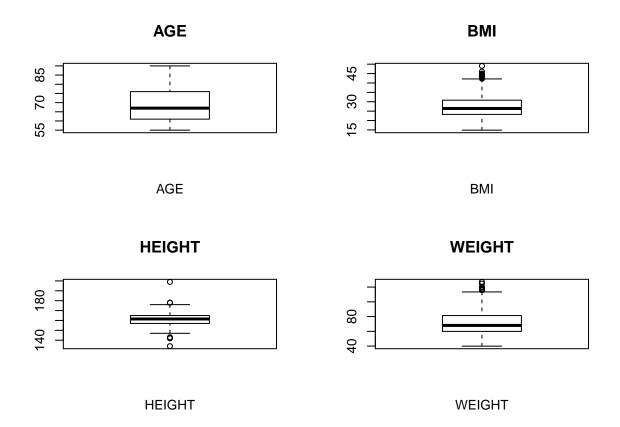




```
#Density Plots
par(mfrow=c(2,2))
for(i in 1:4) {
    plot(density(dataset_numeric[,i]), xlab=names(dataset_numeric)[i], main=names(dataset_numeric)[i])
}
```

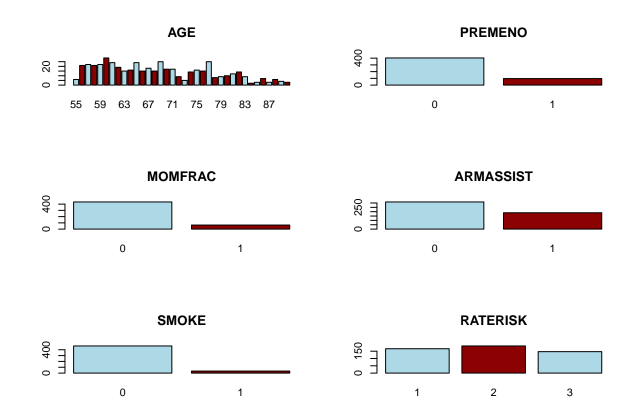


```
#Box And Whisker Plots
par(mfrow=c(2,2))
for(i in 1:4) {
   boxplot(dataset_numeric[,i], xlab=names(dataset_numeric)[i], main=names(dataset_numeric)[i])
}
```

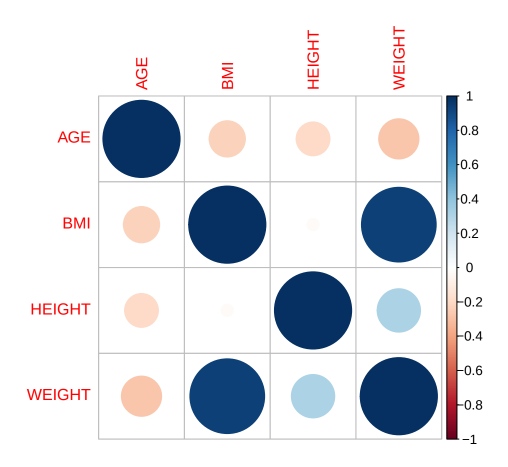


```
#Barplots, which is used to count the accurances for categorical attributes
dataset_categorical = set_noID[,-c(2:5,11)]

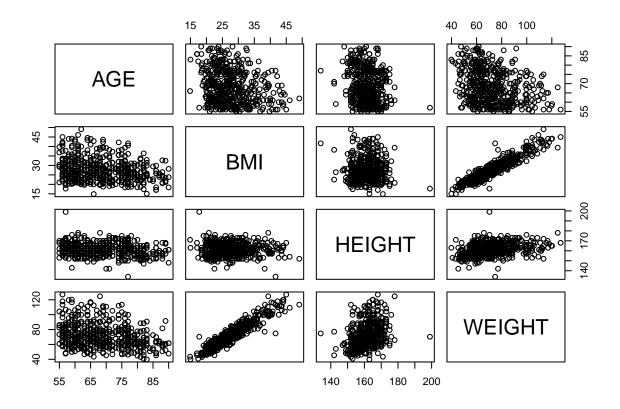
par(mfrow=c(3,2))
for(i in 1:6) {
   counts <- table(dataset_categorical[,i]) # get the count for each categorical value
   name <- names(dataset_categorical)[i]
   barplot(counts, main=name,col=c("lightblue","darkred"))
}</pre>
```



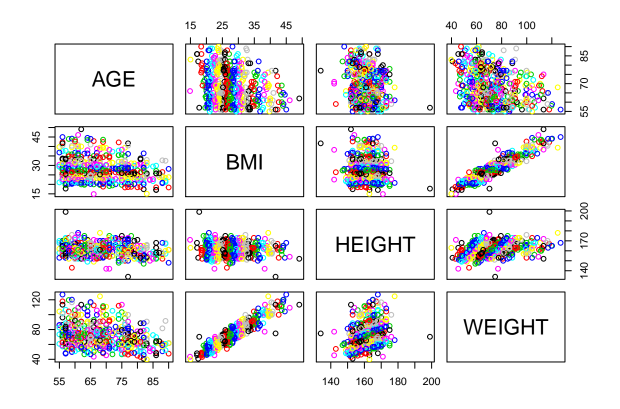
```
#Multivariate Visualization
correlations1=cor(dataset_numeric)
print(correlations1)
##
                 AGE
                            BMI
                                      HEIGHT
                                                 WEIGHT
          1.0000000 -0.22125651 -0.19264861 -0.2715964
## AGE
         -0.2212565 1.00000000 -0.02437689
                                              0.9373360
## HEIGHT -0.1926486 -0.02437689 1.00000000
                                              0.3159691
## WEIGHT -0.2715964 0.93733603 0.31596915
                                             1.0000000
par(mfrow=c(1,1))
corrplot(correlations1, methods="circle")
## Warning in text.default(pos.xlabel[, 1], pos.xlabel[, 2], newcolnames, srt
## = tl.srt, : "methods" is not a graphical parameter
## Warning in text.default(pos.ylabel[, 1], pos.ylabel[, 2], newrownames, col
## = tl.col, : "methods" is not a graphical parameter
## Warning in title(title, ...): "methods" is not a graphical parameter
```



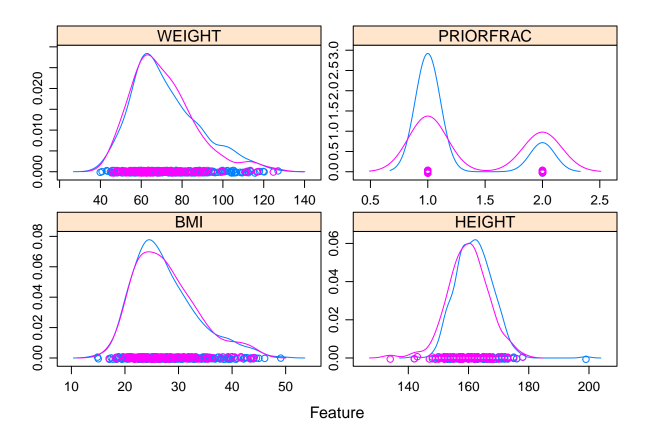
pair-wise scatterplots of the numeric attributes
par(mfrow=c(1,1))
pairs(dataset_numeric)



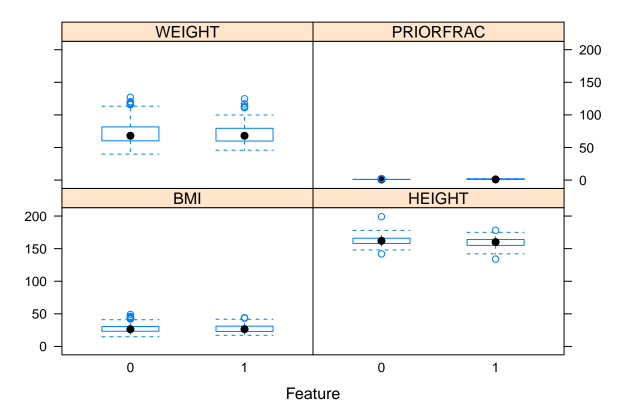
#Scatterplot Matrix By Class (use different color to distinguish different class)
par(mfrow=c(1,1))
pairs(dataset_numeric, col=dataset[,5])



```
# density plots for each attribute by class value
X <- set_noID[2:5]
Y <- set_noID$FRACTURE
X$PRIORFRAC <- as.numeric(X$PRIORFRAC)
scales <- list(x=list(relation="free"), y=list(relation="free"))
par(mfrow=c(1,1))
featurePlot(x=X, y=set_noID$FRACTURE, plot="density", scales=scales)</pre>
```



```
#Box And Whisker Plots By Class
par(mfrow=c(1,1))
featurePlot(x=X, y=set_noID$FRACTURE, plot="box")
```



Checking the Balance of the Full dataset

The current sample dataset containes a larger proportion of subjects that did not develop fracture. Building a model against this dataset could produce bias towards the majority class. Below you will see how many subjects with(1)/without(0) Fractures as well as the proportion percentage for each. After splitting the dataset into training and validation(test) sets, we noticed the proportion of the training and test was not any better.

We fit a logistic model on the unbalanced training dataset with a threshold of .05. It shows a Precision of 1 which says there are no false positives. Recall equals 0.20 is low and indicates that we have higher number of false negatives. The F equals 0.20 is also low and suggests weak accuracy of this model.

We also plotted a ROC curve to visualize the model. The AUC equals 0.764 which is low and shows the data is not balanced.

We will attempt to balance the dataset in order to create a more balanced distribution of and a better prediction.

table(dataset\$FRACTURE)

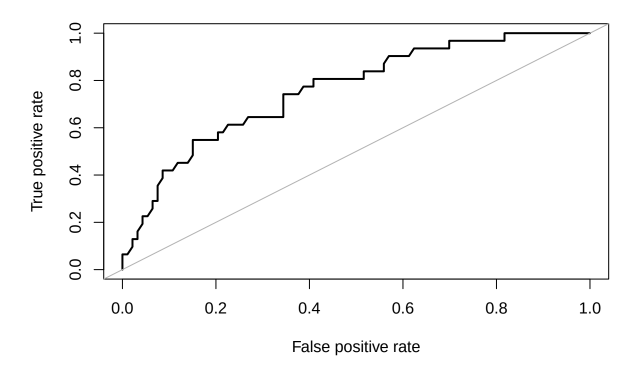
```
## 0 1
## 375 125
```

prop.table(table(dataset\$FRACTURE))

```
## 0 1
## 0.75 0.25
```

```
# split the data into training and validation sets
set.seed(84)
validation index = createDataPartition(dataset$FRACTURE, p=0.75, list=FALSE)
validationData = dataset[-validation_index,c(4:14)]
trainingData = dataset[validation_index,c(4:14)]
prop.table(table(validationData$FRACTURE))
##
##
      0
## 0.75 0.25
prop.table(table(trainingData$FRACTURE))
##
##
      0
          1
## 0.75 0.25
#fit a logistic regressio to unblanced training set
fit.dataset <- glm(formula=FRACTURE~ ., data = trainingData, family="binomial")</pre>
pred.fit.dataset <- predict(fit.dataset, newdata = validationData, type="response")</pre>
#Check Accuracy of fitted model.
accuracy.meas(validationData$FRACTURE,pred.fit.dataset, threshold=.05)
##
## accuracy.meas(response = validationData$FRACTURE, predicted = pred.fit.dataset,
       threshold = 0.05)
##
##
## Examples are labelled as positive when predicted is greater than 0.05
## precision: 0.250
## recall: 1.000
## F: 0.200
#Check Accuracy of Test dataset using ROC curve
roc.curve(validationData$FRACTURE, pred.fit.dataset, plotit = TRUE)
```

ROC curve



Area under the curve (AUC): 0.760

##Create a vector of all categorical variables and run frequency 2X2s with Mosaic plots.

Chi-Square Test For the 2-way tables the chisq test independence will show if 2 categorical variables are related in some population. Null Hypothesis: The two categorical variables are independent. Alternative Hypothesis: The two categorical variables are dependent

Variable: PRIORFRAC 41% of subjects with Prior Franctures also had current Fractures but only make up 25% of the overall subjects in the sample that had prior fractures. The Chi-squared p-value favors overwhemingly the alternative hypothesis that the PRIORFRAC variable is dependent on Fracture variable.

Variable: PREMENO 80% of the sample subjects are not in Pre-Menopausehad of which 24% had fractures. The same frequency of 25% Premenopausal women had fractures. The Chi-squared p-value favors the null hypothesis that the PREMENO variable is independent on Fracture variable.

Variable: MOMFRAC 13% of subjects have Mothers with a history of fractures. Out of those 13%, 36% of subjects also had fractures. The Chi-squared p-value favors the alternative hypothesis that the MOMFRAC variable is probably dependent on Fracture variable.

Variable: ARMASSIST 62% (312/500) subjects do not have Armassist of which 20% had fractures. Of those with Armassist, 33% had fractures. The Chi-squared p-value favors the alternative hypothesis that the ARMASSIST variable is most likely dependent on Fracture variable.

Variable: SMOKE In the dataset, 93% of subjects are non-smokers of which 26% had fractures. 7% of the subjects who were smokers of which 26% had no fractures. Although the subjects are not balance in smoker vs non-smoker category, the p-value for Chi-squared test shows .47 we favor the alternative hypothesis that the Smoke variable is dependent on the Fracture.

Variable: RATERISK Raterisk shows the frequency of subjects in each Raterisk level is between 29%-33%. This is pretty even in terms of how many subjects are within each Raterisk. For those that did have Fractures, their probability of a fracture increased with the level of Raterisk. This makes sense.

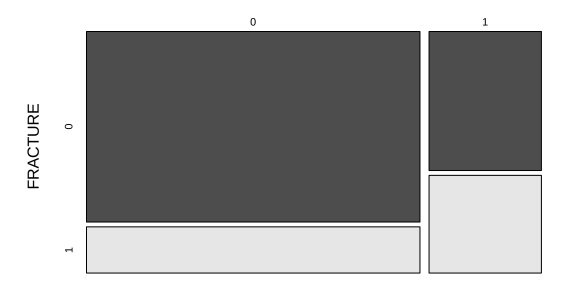
categoricalVarVec <- c("PRIORFRAC","PREMENO","MOMFRAC","ARMASSIST","SMOKE","RATERISK")</pre>

```
for(categoricalVar in categoricalVarVec){
 CrossTable(dataset[,categoricalVar], dataset$FRACTURE, chisq = TRUE , expected = TRUE, dnn=c(categori
 mosaicplot(CrossTable(dataset[ ,categoricalVar], dataset$FRACTURE)$t, main=paste("FRACTURE vs",categoricalVar]
}
##
##
##
     Cell Contents
##
      Expected N |
## |
## | Chi-square contribution |
    N / Row Total |
           N / Col Total |
        N / Table Total |
## |-----|
##
##
## Total Observations in Table: 500
##
       | FRACTURE
##
     PRIORFRAC | 0 |
                        1 | Row Total |
##
##
           0 | 301 | 73 |
          | 280.500 | 93.500 |
##
           | 1.498 | 4.495 | |
| 0.805 | 0.195 | 0.748 |
| 0.803 | 0.584 | |
| 0.602 | 0.146 |
##
##
##
           ---|------|------|
          1 | 74 | 52 | 126 |
##
          | 94.500 | 31.500 |
                4.447 | 13.341 |
            ##
           | 0.587 | 0.413 | 0.252 |
| 0.197 | 0.416 | |
| 0.148 | 0.104 | |
##
## -----|----|
## Column Total | 375 | 125 |
                0.750 | 0.250 |
   ----|-----|-----|
##
##
## Statistics for All Table Factors
##
##
## Pearson's Chi-squared test
## -----
## Chi^2 = 23.78123 d.f. = 1 p = 1.079299e-06
```

```
##
## Pearson's Chi-squared test with Yates' continuity correction
## -----
## Chi^2 = 22.63532 d.f. = 1 p = 1.958512e-06
##
##
##
  Cell Contents
## |-----|
## | Chi-square contribution |
## | N / Row Total |
         N / Col Total |
## |
## |
     N / Table Total |
## |-----|
##
##
## Total Observations in Table: 500
##
                  | dataset$FRACTURE
## dataset[, categoricalVar] | 0 | 1 | Row Total |
## -----|----|-----|
                  0 | 301 | 73 |
##
                      1.498 | 4.495 |
                  - 1
                             0.195 |
0.584 |
##
                   0.805 |
                                      0.748 |
                       0.803 |
                      0.602 | 0.146 |
                       74 | 52 | 126 |
4.447 | 13.341 | |
                  1 |
##
##
##
                      0.587 | 0.413 |
##
                      0.197 |
                              0.416 |
                             0.104 |
                   0.148 |
## -----|----|-----|
                      375 | 125 |
         Column Total |
                       0.750 | 0.250 |
##
                  ## -----|-----|-----|
##
##
```

30

FRACTURE vs PRIORFRAC



PRIORFRAC

```
##
##
##
     Cell Contents
## |-----|
           N I
## |
      Expected N |
## | Chi-square contribution |
## | N / Row Total | ## | N / Col Total | ## | N / Table Total |
## |-----|
##
##
## Total Observations in Table: 500
##
##
        | FRACTURE
##
      PREMENO | 0 | 1 | Row Total |
##
           0 | 303 | 100 | 403 |
| 302.250 | 100.750 | |
| 0.002 | 0.006 |
##
             | 0.002 | 0.006 | | |
| 0.752 | 0.248 | 0.806 |
| 0.808 | 0.800 | |
##
##
##
              | 0.606 | 0.200 |
```

```
1 | 72 | 25 | 97 |
| 72.750 | 24.250 |
##
##
           0.008 | 0.023 |
##
              0.742 |
                       0.258 |
                                0.194 |
##
           0.192 |
                       0.200 |
                       0.050 l
          - 1
              0.144 |
             375 |
                       125 |
## Column Total |
               0.750 | 0.250 |
   1
    -----|-----|
## Statistics for All Table Factors
##
##
## Pearson's Chi-squared test
## Chi^2 = 0.038372 d.f. = 1 p = 0.844698
## Pearson's Chi-squared test with Yates' continuity correction
## -----
## Chi^2 = 0.004263556 d.f. = 1 p = 0.9479384
##
##
##
##
   Cell Contents
## | Chi-square contribution |
## | N / Row Total | ## | N / Col Total |
        N / Table Total |
##
##
## Total Observations in Table: 500
##
##
                     | dataset$FRACTURE
##
## dataset[, categoricalVar] | 0 | 1 | Row Total |
  -----|
                        303 | 100 | 403 |
0.002 | 0.006 | |
0.752 | 0.248 | 0.806 |
                   0 |
##
                    0.800 |
0.200 |
                        0.808 |
##
                        0.606 |
  -----|-----|-----|
                         72 | 25 | 97 |
                   1 |
                    | 0.008 | 0.023 | |
| 0.742 | 0.258 | 0.194 |
##
##
                        0.192 | 0.200 |
##
                    | 0.144 | 0.050 |
## -----|-----|
```

##	Column Total	375	125	500
##	1	0.750	0.250	1
##	I			
##				
##				

FRACTURE vs PREMENO



PREMENO

```
##
##
##
  Cell Contents
## |-----|
## |
     Expected N |
## | Chi-square contribution |
## | N / Row Total | ## | N / Col Total |
   N / Table Total |
##
## Total Observations in Table: 500
##
##
##
       | FRACTURE
   MOMFRAC | 0 | 1 | Row Total |
## -----|
    0 | 334 | 101 | 435 |
##
```

```
| 326.250 | 108.750 |
##
           0.184 | 0.552 |
##
          -
            0.768 |
                    0.232 |
##
          -
                            0.870 |
            0.891 |
                    0.808 |
##
          0.668 | 0.202 |
##
          ##
 -----|-----|
       1 | 41 | 24 | 65 |
           48.750 | 16.250 |
##
         3.696
            1.232 |
##
          1
##
          1
             0.631 | 0.369 |
                             0.130 |
          1
            0.109 |
                     0.192 |
            0.082 |
                     0.048 |
          ## Column Total | 375 | 125 |
      | 0.750 | 0.250 |
   -----|----|
##
##
## Statistics for All Table Factors
##
## Pearson's Chi-squared test
## -----
## Chi^2 = 5.664604 d.f. = 1 p = 0.01731063
##
## Pearson's Chi-squared test with Yates' continuity correction
## -----
## Chi^2 = 4.957265 d.f. = 1 p = 0.02598127
##
##
##
##
##
   Cell Contents
## |
## | Chi-square contribution |
## | N / Row Total |
        N / Col Total |
    N / Table Total |
## |-----|
##
## Total Observations in Table: 500
##
##
##
                  | dataset$FRACTURE
## dataset[, categoricalVar] | 0 | 1 | Row Total |
 -----|-----|-----|
                      334 | 101 |
##
                 0 |
                            ##
                      0.184 |
                  ##
                  0.768 |
                     0.891 | 0.808 |
##
                      0.668 | 0.202 |
## -----|----|-----|
```

##	1	41	l 24	l 65 l
##		1.232	3.696	l I
##		0.631	0.369	0.130
##		0.109	0.192	
##		0.082	0.048	
##				
##	Column Total	375	l 125	J 500 J
##		0.750	0.250	
##				
##				
##				

FRACTURE vs MOMFRAC



MOMFRAC

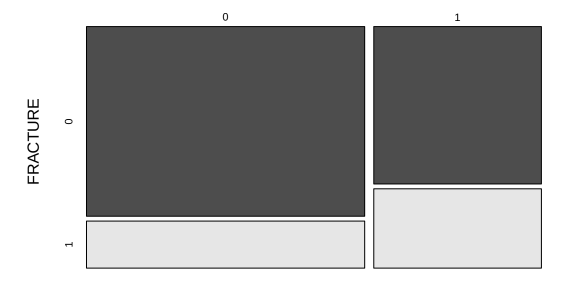
```
##
##
##
      Cell Contents
##
                         N I
## |
## |
                Expected N |
## | Chi-square contribution |
## |
            N / Row Total |
              N / Col Total |
## |
## |
           N / Table Total |
##
## Total Observations in Table: 500
```

```
##
      | FRACTURE
##
   ARMASSIST | 0 |
                      1 | Row Total |
##
## -----|-----|
        0 | 250 | 62 | 312 |
##
         | 234.000 | 78.000 |
            1.094 | 3.282 | |
0.801 | 0.199 | 0.624 |
##
          ##
         - 1
##
         - 1
            0.667 | 0.496 |
             0.500 l
                     0.124 |
  -----|-----|
##
       1 | 125 | 63 | 188 |
         | 141.000 | 47.000 |
##
             ##
          - 1
             0.665 |
##
          ##
          - 1
            0.333 |
             0.250
                     0.126
  -----|-----|
            375 | 125 |
0.750 | 0.250 |
## Column Total |
    1
## -----|-----|
##
## Statistics for All Table Factors
##
## Pearson's Chi-squared test
## Chi^2 = 11.63848 d.f. = 1 p = 0.0006460138
## Pearson's Chi-squared test with Yates' continuity correction
 -----
## Chi^2 = 10.92244 d.f. = 1 p = 0.0009500637
##
##
##
##
  Cell Contents
## |-----|
## | Chi-square contribution |
## | N / Row Total |
## |
        N / Col Total |
     N / Table Total |
## |-----|
##
##
## Total Observations in Table: 500
##
##
                  | dataset$FRACTURE
## dataset[, categoricalVar] | 0 | 1 | Row Total |
## -----|-----|
```

##

##	0	250	l 62	312
##	I	1.094	3.282	1
##	I	0.801	0.199	0.624
##	I	0.667	0.496	1
##	I	0.500	0.124	1
##				
##	1	125	l 63	188
##	I	1.816	5.447	1
##	I	0.665	0.335	0.376
##	I	0.333	0.504	1
##	I	0.250	0.126	1
##				
##	Column Total	375	125	J 500 J
##	I	0.750	0.250	1
##				
##				
##				

FRACTURE vs ARMASSIST



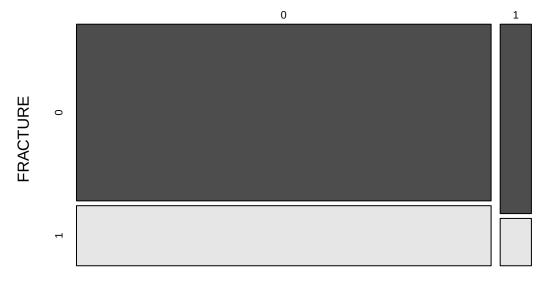
ARMASSIST

```
##
##
## Cell Contents
## |------|
## | N |
## | Expected N |
## | Chi-square contribution |
## | N / Row Total |
```

```
## | N / Col Total | ## | N / Table Total |
##
## Total Observations in Table: 500
##
##
           | FRACTURE
       SMOKE | 0 |
                        1 | Row Total |
     -----|----|
          0 | 347 | 118 |
                                   465 l
##
             348.750 | 116.250 |
##
          - 1
              0.009 | 0.026 |
0.746 | 0.254 |
0.925 | 0.944 |
0.694 | 0.236 |
##
           - 1
##
           ##
            ##
           1
                28 | 7 |
##
         1 |
              26.250 | 8.750 | |
0.117 | 0.350 | |
0.800 | 0.200 | 0.070 |
##
           ##
           - 1
           0.075 |
                        0.056 |
##
           0.014 |
           - 1
                0.056 l
## -----|-----|
## Column Total | 375 |
                         125 |
  | 0.750 | 0.250 |
       -----|------|------|
##
## Statistics for All Table Factors
##
##
## Pearson's Chi-squared test
## -----
## Chi^2 = 0.5017921 d.f. = 1 p = 0.4787137
## Pearson's Chi-squared test with Yates' continuity correction
## -----
## Chi^2 = 0.2560164 d.f. = 1 p = 0.6128703
##
##
##
    Cell Contents
## |-----|
## | Chi-square contribution |
## | N / Row Total | ## | N / Col Total |
     N / Table Total |
## |-----|
##
##
```

## ## ##						
##	dataset\$FRACTURE					
## ##	dataset[, categoricalVar]	0	1 	Row Total		
##	0	347	118	465		
##	!	0.009	0.026			
##	!	0.746	0.254	0.930		
##	!	0.925	0.944			
##		0.694	0.236			
##						
##	1	28	7	35		
##	l	0.117	0.350			
##	I	0.800	0.200	0.070		
##	I	0.075	0.056	l I		
##	I	0.056	0.014	l l		
##						
##	Column Total	375	125	500		
##	I	0.750	0.250			
##						
##						
##						

FRACTURE vs SMOKE



SMOKE

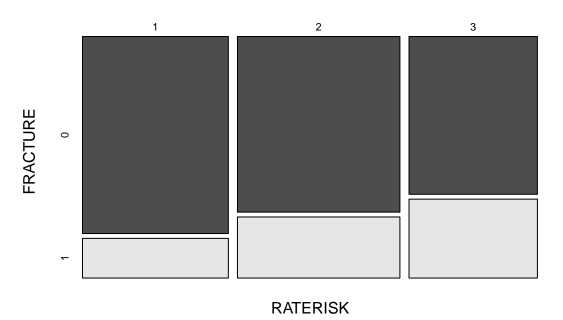
##

```
## Cell Contents
## |
    Expected N |
## |
## | Chi-square contribution |
## | N / Row Total |
         N / Col Total |
      N / Table Total |
## |
## |-----|
##
##
## Total Observations in Table: 500
##
##
      | FRACTURE
                     1 | Row Total |
##
    RATERISK | 0 |
##
  -----|----|
         1 | 139 | 28 |
##
##
         | 125.250 | 41.750 |
             1.509 | 4.528 | |
0.832 | 0.168 | 0.334 |
##
          ##
          - 1
##
          - 1
             0.371 | 0.224 |
          | 0.278 | 0.056 |
##
##
         ---|------|------|
         2 | 138 | 48 | 186 |
##
                                | 139.500 | 46.500 |
             0.016 | 0.048 |
0.742 | 0.258 |
0.368 | 0.384 |
##
          1
                             0.372 |
          0.276 | 0.096 |
## -----|-----|
         3 | 98 | 49 | 147 |
##
         | 110.250 | 36.750 |
##
          | 1.361 | 4.083 |
| 0.667 | 0.333 |
| 0.261 | 0.392 |
##
          0.294 |
##
##
          0.196 |
                      0.098 l
## -----|----|
             375 |
                      125 |
                                500 l
## Column Total |
              0.750 |
                      0.250 |
       1
  -----|-----|
##
## Statistics for All Table Factors
##
## Pearson's Chi-squared test
## -----
## Chi^2 = 11.54688 d.f. = 2 p = 0.003109037
##
##
##
##
##
```

```
## Cell Contents
## |-----|
## | Chi-square contribution |
## | N / Row Total | ## | N / Col Total |
    N / Table Total |
## |-----|
##
##
## Total Observations in Table: 500
##
##
                     | dataset$FRACTURE
##
## dataset[, categoricalVar] | 0 | 1 | Row Total |
  -----|-
                                  28 | 167 |
4.528 | |
                         139 |
##
                     1 |
                          1.509 |
##
                                             0.334 l
##
                          0.832 |
                                   0.168 |
                         0.371 |
                                   0.224
##
##
                           0.278 |
                                    0.056 l
                     --|-----|---|----|--
##
                     2 |
                           138 |
                                   48 |
                                              ĺ
                     0.016 l
                                   0.048 l
##
                                   0.258 |
                      0.742 |
                                              0.372 |
                           0.368 |
                                    0.384 |
##
                           0.276 |
                                    0.096 |
                                  49 |
                           98 |
                     3 |
                                              - 1
##
                           1.361 |
                                   4.083 |
                                   0.333 |
##
                           0.667 |
                                  0.392 |
##
                           0.261 |
                           0.196 |
                                    0.098 |
                           375 | 125 |
0.750 | 0.250 |
            Column Total |
                                                500 I
##
                     - 1
## -----|----|-----|
##
##
```

41

FRACTURE vs RATERISK



Logistic Regression

Training set will be 70% of dataset and Test set will be remaining 30%

Build Model using Training Data

Question of Interest? What are the odds of getting a fracture, given certain conditions?

```
set.seed(84)
model <- glm(FRACTURE ~ AGE + WEIGHT + HEIGHT + BMI + PRIORFRAC + PREMENO + MOMFRAC + ARMASSIST + SMOKE
##
  Call: glm(formula = FRACTURE ~ AGE + WEIGHT + HEIGHT + BMI + PRIORFRAC +
##
##
       PREMENO + MOMFRAC + ARMASSIST + SMOKE + RATERISK, family = "binomial",
       data = trainingData)
##
##
## Coefficients:
##
  (Intercept)
                        AGE
                                  WEIGHT
                                                HEIGHT
                                                                BMI
     -12.04673
                    0.03168
                                 -0.10711
                                               0.04735
                                                            0.29193
##
    PRIORFRAC1
                   PREMENO1
                                MOMFRAC1
##
                                            ARMASSIST1
                                                             SMOKE1
                                 0.35482
                                                           -0.08005
##
       0.73265
                    0.04114
                                               0.30067
##
     RATERISK2
                  RATERISK3
##
       0.38692
                    0.57786
##
## Degrees of Freedom: 375 Total (i.e. Null); 364 Residual
```

```
## Null Deviance:
                        422.9
## Residual Deviance: 385.4
                                AIC: 409.4
summary(model)
##
## Call:
  glm(formula = FRACTURE ~ AGE + WEIGHT + HEIGHT + BMI + PRIORFRAC +
##
       PREMENO + MOMFRAC + ARMASSIST + SMOKE + RATERISK, family = "binomial",
##
       data = trainingData)
##
## Deviance Residuals:
##
       Min
                 1Q
                      Median
                                   3Q
                                           Max
## -1.4739
           -0.7388 -0.5757
                             -0.1189
                                         2.1597
##
## Coefficients:
##
                Estimate Std. Error z value Pr(>|z|)
## (Intercept) -12.04673
                           13.81668
                                    -0.872
                                            0.38326
                 0.03168
                            0.01715
                                      1.847
                                             0.06472
## AGE
## WEIGHT
                -0.10711
                            0.09271
                                     -1.155
                                            0.24793
## HEIGHT
                 0.04735
                            0.08516
                                      0.556 0.57823
## BMI
                 0.29193
                            0.23882
                                      1.222
                                             0.22157
                                      2.582
## PRIORFRAC1
                 0.73265
                            0.28371
                                             0.00981 **
## PREMENO1
                 0.04114
                            0.32545
                                      0.126
                                             0.89940
## MOMFRAC1
                 0.35482
                            0.36197
                                      0.980 0.32697
## ARMASSIST1
                 0.30067
                            0.29666
                                      1.014 0.31080
## SMOKE1
                -0.08005
                            0.50041
                                     -0.160 0.87290
                 0.38692
## RATERISK2
                            0.32506
                                      1.190 0.23393
## RATERISK3
                 0.57786
                            0.34936
                                      1.654 0.09812 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
  (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 422.88 on 375 degrees of freedom
##
## Residual deviance: 385.45 on 364 degrees of freedom
## AIC: 409.45
##
## Number of Fisher Scoring iterations: 4
h1 <- hoslem.test(model$y, fitted(model), g = 10) #number of groups to divide dataset into is 10
h1
##
##
   Hosmer and Lemeshow goodness of fit (GOF) test
```

Interpretation of logistic regression model: Weight, height, BMI, Premeno, Armassist, and Smoke are not statistically significant variables. Priorfrac and Age are statistically significant variables and have the lowest p-value indicating a strong association with having a Fracture.

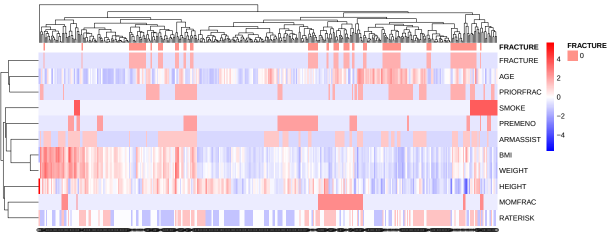
##

data: model\$y, fitted(model)

X-squared = 7.8006, df = 8, p-value = 0.4532

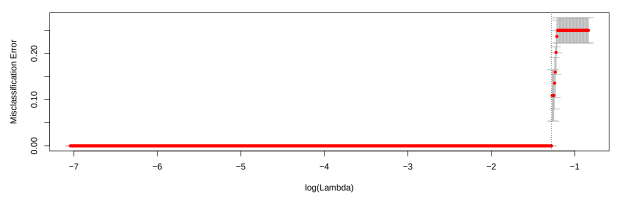
Clustering

```
#Lets look at a heatmap using hierarchical clustering to see if the
#response naturually clusters out using the predictors
#Transposting the predictor matrix and giving the response categories its
#row names.
#Get Training Set
# convert factors to numeric for pheatmap
temp <- trainingData</pre>
indx <- sapply(temp, is.factor)</pre>
temp[indx] <- lapply(temp[indx], function(x) as.numeric(as.character(x)))</pre>
dat.train <- temp
dat.train.x <- dat.train[,1:ncol(dat.train)]</pre>
dat.train.y <- dat.train$FRACTURE</pre>
dat.train.y <- as.factor(as.character(dat.train.y))</pre>
#Heatmap
x<-t(dat.train.x)
colnames(x)<-dat.train.y</pre>
pheatmap(x,annotation_col=data.frame(FRACTURE=dat.train.y),scale="row",legend=T,color=colorRampPalette(
```



```
##logistic regression
dat.train.x <- as.matrix(dat.train.x)

cvfit <- cv.glmnet(dat.train.x, dat.train.y, family = "binomial", type.measure = "class", nlambda = 100
plot(cvfit)</pre>
```



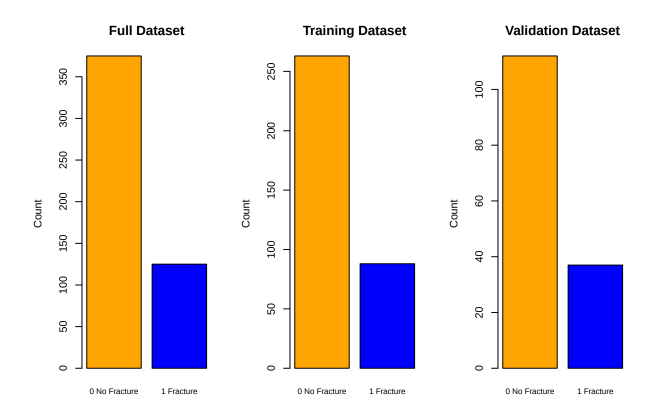
```
coef(cvfit, s = "lambda.min")
```

*** Appendix B: Model Comparison - Analysis ========

```
## 'data.frame':
                   500 obs. of 14 variables:
## $ SUB_ID
             : int 1 2 3 4 5 6 7 8 9 10 ...
## $ SITE_ID : int 1 4 6 6 1 5 5 1 1 4 ...
## $ PHY_ID
              : int 14 284 305 309 37 299 302 36 8 282 ...
  $ PRIORFRAC: Factor w/ 2 levels "0", "1": 1 1 2 1 1 2 1 2 2 1 ...
              : int 62 65 88 82 61 67 84 82 86 58 ...
##
  $ AGE
   $ WEIGHT
              : num 70.3 87.1 50.8 62.1 68 68 50.8 40.8 62.6 63.5 ...
              : int 158 160 157 160 152 161 150 153 156 166 ...
##
  $ HEIGHT
              : num 28.2 34 20.6 24.3 29.4 ...
  $ PREMENO : Factor w/ 2 levels "0","1": 1 1 1 1 1 1 1 1 1 1 ...
##
   $ MOMFRAC : Factor w/ 2 levels "0","1": 1 1 2 1 1 1 1 1 1 1 ...
  $ ARMASSIST: Factor w/ 2 levels "0","1": 1 1 2 1 1 1 1 1 1 1 ...
             : Factor w/ 2 levels "0", "1": 1 1 1 1 1 2 1 1 1 1 ...
  $ SMOKE
   $ RATERISK : Factor w/ 3 levels "1","2","3": 2 2 1 1 2 2 1 2 2 1 ...
   $ FRACTURE : Factor w/ 2 levels "0","1": 1 1 1 1 1 1 1 1 1 1 ...
```

Create Train and Validation Datasets

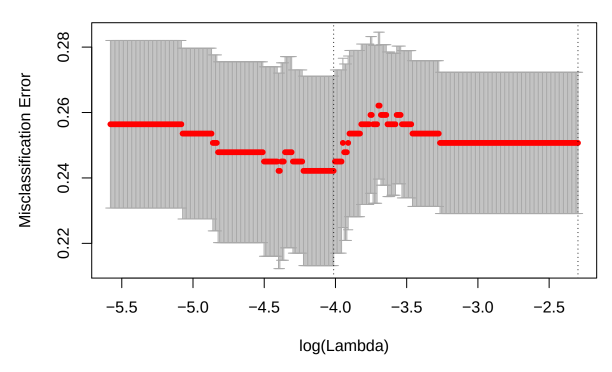
```
set.seed(999)
validation_index = createDataPartition(dataset$FRACTURE, p=0.70, list=FALSE)
validationData = dataset[-validation_index,c(4:14)]
trainingData = dataset[validation_index,c(4:14)]
table(dataset$FRACTURE)
##
##
    0 1
## 375 125
table(trainingData$FRACTURE)
##
##
    0
        1
## 263 88
table(validationData$FRACTURE)
##
##
   0
## 112 37
#BarPlots of Fracture counts between full, training and validation datasets.
par(mfrow=c(1,3))
\#par(mar=c(5,8,4,2)) \# increase y-axis margin.
count_full <- table(dataset$FRACTURE)</pre>
count_trn <- table(trainingData$FRACTURE)</pre>
count_test <- table(validationData$FRACTURE)</pre>
barplot(count_full,main="Full Dataset", ylab="Count", col=c("orange","blue"),names.arg=c("O No Fracture
barplot(count_trn,main="Training Dataset", ylab="Count", col=c("orange","blue"),names.arg=c("O No Fract
barplot(count_test,main="Validation Dataset", ylab="Count", col=c("orange", "blue"),names.arg=c("O No Fr
```



```
## Formatting Test Data Set
# Recode Rate Risk Variable since its ordinal and we donot want to loose its info if it gets
# coded as nominal variable before running the Model
validationData$RATERISK <- factor(validationData$RATERISK, levels = c(1,2,3), ordered = T)

xfactors_test <- model.matrix(validationData$FRACTURE ~ validationData$PRIORFRAC + validationData$PREME
x_test <- as.matrix(data.frame(validationData$AGE, validationData$WEIGHT, validationData$HEIGHT, validat
## Formatting Training Data Set
trainingData$RATERISK <- factor(trainingData$RATERISK, levels = c(1,2,3), ordered = T)
xfactors_train <- model.matrix(trainingData$FRACTURE ~ trainingData$PRIORFRAC + trainingData$PREMENO +
x_train <- as.matrix(data.frame(trainingData$AGE, trainingData$WEIGHT, trainingData$HEIGHT, trainingData
## Doing Cross validation to find the best fitting model based upon Lasso
cvfit <- cv.glmnet(x_train, y=trainingData$FRACTURE, family = "binomial", type.measure = "class", nlamb
plot(cvfit)</pre>
```

9 9 9 8 8 8 8 8 8 6 6 5 5 5 5 5 5 4 3 2 1



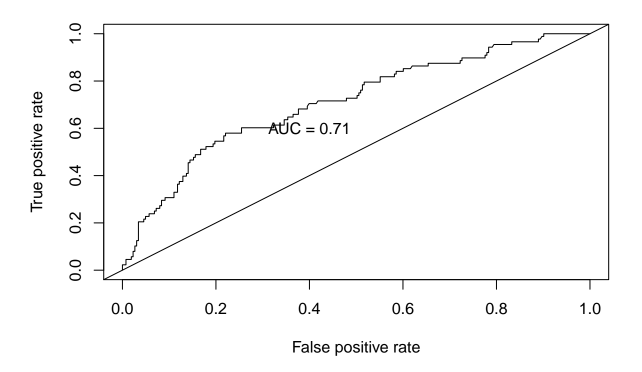
```
# Model with Lowest Lambda is shrinking all the coefficients, hence selecting lambda based upon # Test Set AUC and EDA Results #cvfitplanet.fit coef(cvfit, s="lambda.min")
```

```
## 12 x 1 sparse Matrix of class "dgCMatrix"
##
## (Intercept)
                            1.52902669
## trainingData.AGE
                            0.03598544
## trainingData.WEIGHT
## trainingData.HEIGHT
                           -0.03340871
## trainingData.BMI
## trainingData.PRIORFRAC1
                            0.15180349
## trainingData.PREMENO1
## trainingData.MOMFRAC1
                            0.04035537
## trainingData.ARMASSIST1
                            0.52512963
## trainingData.SMOKE1
                           0.33991586
## trainingData.RATERISK.L
## trainingData.RATERISK.Q
```

```
# Fitting the best model based upon selected lambda
fit <- glmnet(x_train, y=trainingData$FRACTURE, family="binomial", alpha = 1, lambda = cvfit$lambda.min
# First Predicting the responses on training data set itself
fit.pred <- predict(fit, newx = x_train, type = "response")</pre>
```

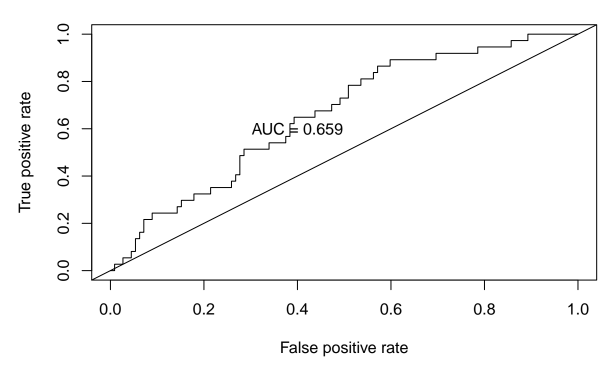
```
#Create ROC curves for training Data Set
pred <- prediction(fit.pred[,1], trainingData$FRACTURE)
roc.perf = performance(pred, measure = "tpr", x.measure = "fpr")
auc.train <- performance(pred, measure = "auc")
auc.train <- auc.train@y.values

##Plot ROC for training Set
plot(roc.perf)
abline(a=0, b= 1) #Ref line indicating poor performance
text(x = .40, y = .6,paste("AUC = ", round(auc.train[[1]],3), sep = ""))</pre>
```



```
#Run model from training set on validation Set
fit.pred1 <- predict(fit, newx = x_test, type = "response")

#ROC curves
pred1 <- prediction(fit.pred1[,1], validationData$FRACTURE)
roc.perf1 = performance(pred1, measure = "tpr", x.measure = "fpr")
auc.val1 <- performance(pred1, measure = "auc")
auc.val1 <- auc.val1@y.values
plot(roc.perf1)
abline(a=0, b= 1)
text(x = .40, y = .6,paste("AUC = ", round(auc.val1[[1]],3), sep = ""))</pre>
```



```
#confusion matrix
pdata <- predict(fit, newx = x_test, type = "response")</pre>
pdata_logical <- pdata[, 1] > 0.5
confusionMatrix(data = as.factor(as.numeric(pdata_logical)), reference = as.factor(as.numeric(validation))
## Confusion Matrix and Statistics
##
##
             Reference
                0
                    1
## Prediction
##
            0 108
                   35
                    2
##
                4
##
##
                  Accuracy: 0.7383
##
                    95% CI: (0.66, 0.8068)
       No Information Rate: 0.7517
##
       P-Value [Acc > NIR] : 0.6866
##
##
##
                      Kappa : 0.0255
##
    Mcnemar's Test P-Value : 1.556e-06
##
##
               Sensitivity: 0.96429
##
               Specificity: 0.05405
##
            Pos Pred Value: 0.75524
            Neg Pred Value: 0.33333
##
```

Prevalence: 0.75168
Detection Rate: 0.72483

##

##

```
## Detection Prevalence : 0.95973
## Balanced Accuracy : 0.50917
##
## 'Positive' Class : 0
##

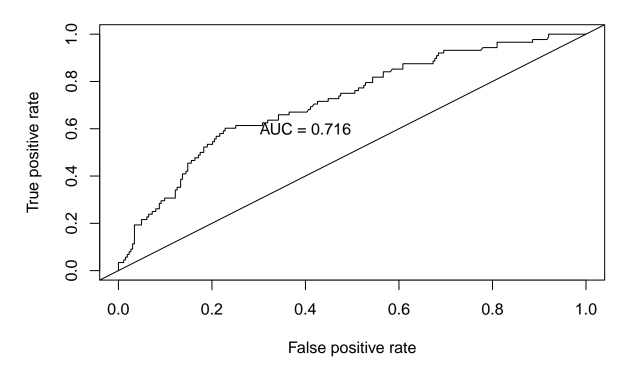
#mydata <- dataset[, c(4:14)] %>% dplyr::select_if(is.numeric)
#predictors <- colnames(mydata)
#mydata <- mydata %>%
# mutate(logit = log(probabilities/(1-probabilities))) %>%
# gather(key = "predictors", value = "predictor.value", -logit)
```

Run Normal Logit Model with Identified Predictors

```
set.seed(999)
logit.fit <- glm(FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC + ARMASSIST + RATERISK , data = training
summary(logit.fit)
##
## glm(formula = FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC +
      ARMASSIST + RATERISK, family = binomial(link = "logit"),
##
##
      data = trainingData)
##
## Deviance Residuals:
      Min
                10
                    Median
                                 3Q
                                         Max
## -1.5491 -0.7377 -0.5763
                            0.2298
                                      2.2214
##
## Coefficients:
              Estimate Std. Error z value Pr(>|z|)
## (Intercept) 3.33365 3.80104 0.877 0.38047
                                  2.755 0.00587 **
## AGE
              0.04347
                         0.01578
## HEIGHT
              -0.04881
                         0.02165 -2.254 0.02418 *
                         0.30097
## PRIORFRAC1 0.22281
                                  0.740 0.45912
## MOMFRAC1
              0.33522
                         0.38263
                                  0.876 0.38097
## ARMASSIST1 0.68418
                         0.27861
                                   2.456 0.01406 *
## RATERISK.L 0.50762
                         0.24656
                                   2.059 0.03951 *
## RATERISK.Q -0.06727
                         0.22219 -0.303 0.76209
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 395.31 on 350 degrees of freedom
## Residual deviance: 355.55 on 343 degrees of freedom
## AIC: 371.55
##
## Number of Fisher Scoring iterations: 4
```

```
# To exponentiate the log ODDS to make it ODDS Ratio and also get corresponding 95% CIs
exp(cbind(ODDs_Ratio = coef(logit.fit), confint(logit.fit)))
## Waiting for profiling to be done...
##
               ODDs Ratio
                              2.5 %
## (Intercept) 28.0403767 0.0172620 5.378401e+04
                1.0444253 1.0128322 1.077662e+00
## HEIGHT
                0.9523628 0.9118527 9.929059e-01
## PRIORFRAC1 1.2495774 0.6857875 2.237992e+00
               1.3982464 0.6449924 2.917750e+00
## MOMFRAC1
## ARMASSIST1 1.9821443 1.1469582 3.428352e+00
## RATERISK.L 1.6613370 1.0283916 2.713240e+00
## RATERISK.Q 0.9349462 0.6060378 1.451499e+00
# First Predicting the responses on training data set itself
logistic.fit.pred.train <- predict(logit.fit, newdata=trainingData, type = "response")</pre>
#Create ROC curves for training Data Set
pred.train <- prediction(logistic.fit.pred.train, trainingData$FRACTURE)</pre>
roc.perf = performance(pred.train, measure = "tpr", x.measure = "fpr")
auc.train <- performance(pred.train, measure = "auc")</pre>
auc.train <- auc.train@y.values</pre>
##Plot ROC for training Set
plot(roc.perf, main="Logistic Reg Training Data Set")
abline(a=0, b= 1) #Ref line indicating poor performance
text(x = .40, y = .6, paste("AUC = ", round(auc.train[[1]],3), sep = ""))
```

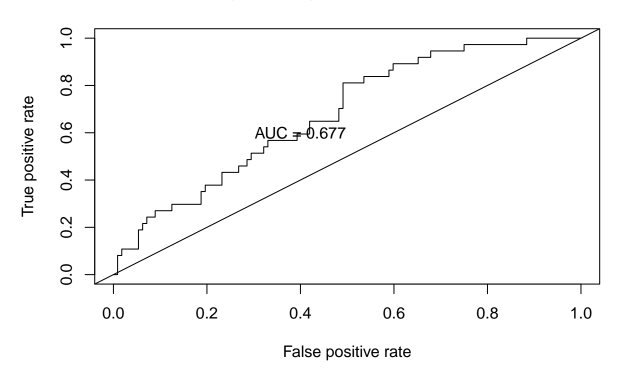
Logistic Reg Training Data Set



```
#Run model from training set on validation Set
logistic.fit.pred.test <- predict(logit.fit, newdata=validationData, type = "response")

#ROC curves
pred.test <- prediction(logistic.fit.pred.test, validationData$FRACTURE)
roc.perf1 = performance(pred.test, measure = "tpr", x.measure = "fpr")
auc.val1 <- performance(pred.test, measure = "auc")
auc.val1 <- auc.val1@y.values
plot(roc.perf1, main="Logistic Reg Validation Data Set")
abline(a=0, b= 1)
text(x = .40, y = .6,paste("AUC = ", round(auc.val1[[1]],3), sep = ""))</pre>
```

Logistic Reg Validation Data Set



```
#confusion matrix
pdata_logical <- logistic.fit.pred.test > 0.5
confusionMatrix(data = as.factor(as.numeric(pdata_logical)), reference = as.factor(as.numeric(validation))
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction
                0
##
            0 106
                   32
##
                6
                    5
##
##
                  Accuracy: 0.745
                    95% CI: (0.6672, 0.8128)
##
##
       No Information Rate: 0.7517
       P-Value [Acc > NIR] : 0.6175
##
##
##
                     Kappa : 0.1067
    Mcnemar's Test P-Value : 5.002e-05
##
##
##
               Sensitivity: 0.9464
##
               Specificity: 0.1351
##
            Pos Pred Value : 0.7681
##
            Neg Pred Value: 0.4545
##
                Prevalence: 0.7517
##
            Detection Rate: 0.7114
      Detection Prevalence: 0.9262
##
```

```
## Balanced Accuracy : 0.5408
##
## 'Positive' Class : 0
##
```

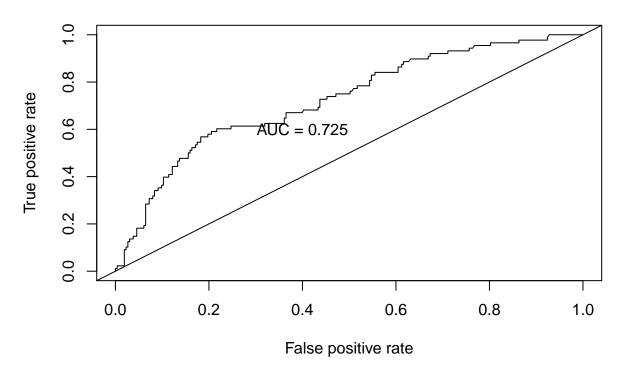
Add Interactions to Normal logit

```
set.seed(999)
# Since top 3 predictors are Age, PriorFrac and RISK, adding model complexity
# via interactions
logit.fit.interactions <- glm(FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC + ARMASSIST + RATERISK + AG
summary(logit.fit.interactions)
##
## Call:
## glm(formula = FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC +
      ARMASSIST + RATERISK + AGE:PRIORFRAC + RATERISK:AGE + MOMFRAC:ARMASSIST,
##
##
      family = binomial(link = "logit"), data = trainingData)
##
## Deviance Residuals:
##
      Min
                10
                     Median
                                  30
                                          Max
## -1.5521 -0.7592 -0.5543 0.2845
                                       2.3802
##
## Coefficients:
##
                      Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                       1.65488 3.98301 0.415 0.67779
                                  0.02058 3.273 0.00107 **
## AGE
                       0.06735
## HEIGHT
                      -0.04913
                                  0.02191 -2.242 0.02495 *
## PRIORFRAC1
                       3.83397
                                  2.26015 1.696 0.08982 .
## MOMFRAC1
                       0.74167
                                0.50795
                                          1.460 0.14426
## ARMASSIST1
                                  0.29990 2.687 0.00721 **
                       0.80585
                      1.64141
## RATERISK.L
                                  2.00121 0.820 0.41210
## RATERISK.Q
                      -1.25322 1.78216 -0.703 0.48193
                      -0.05038
## AGE:PRIORFRAC1
                                  0.03124 -1.612 0.10688
## AGE:RATERISK.L
                      -0.01548
                                  0.02788 -0.555
                                                  0.57869
## AGE:RATERISK.Q
                       0.01632
                                  0.02498
                                          0.653 0.51355
## MOMFRAC1:ARMASSIST1 -0.74229
                                  0.76220 -0.974 0.33011
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 395.31 on 350 degrees of freedom
## Residual deviance: 351.13 on 339 degrees of freedom
## AIC: 375.13
## Number of Fisher Scoring iterations: 5
# First Predicting the responses on training data set itself
logistic.fit.pred.train.interaction <- predict(logit.fit.interactions, newdata=trainingData, type = "re</pre>
```

```
#Create ROC curves for training Data Set
pred.train.interaction <- prediction(logistic.fit.pred.train.interaction, trainingData$FRACTURE)
roc.perf = performance(pred.train.interaction, measure = "tpr", x.measure = "fpr")
auc.train <- performance(pred.train.interaction, measure = "auc")
auc.train <- auc.train@y.values

##Plot ROC for training Set
plot(roc.perf, main="Logistic Reg With Interactions Training Data Set")
abline(a=0, b= 1) #Ref line indicating poor performance
text(x = .40, y = .6,paste("AUC = ", round(auc.train[[1]],3), sep = ""))</pre>
```

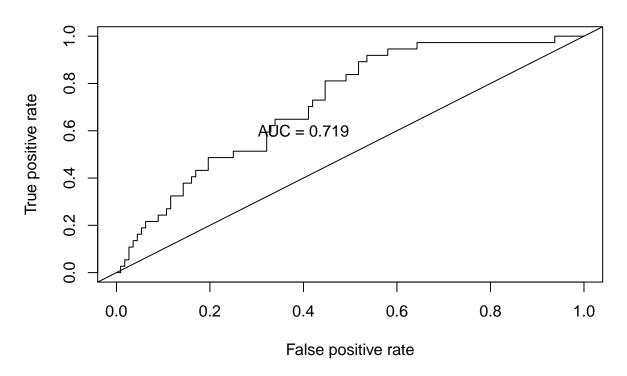
Logistic Reg With Interactions Training Data Set



```
#Run model from training set on validation Set
logistic.fit.pred.test.interaction <- predict(logit.fit.interactions, newdata=validationData, type =

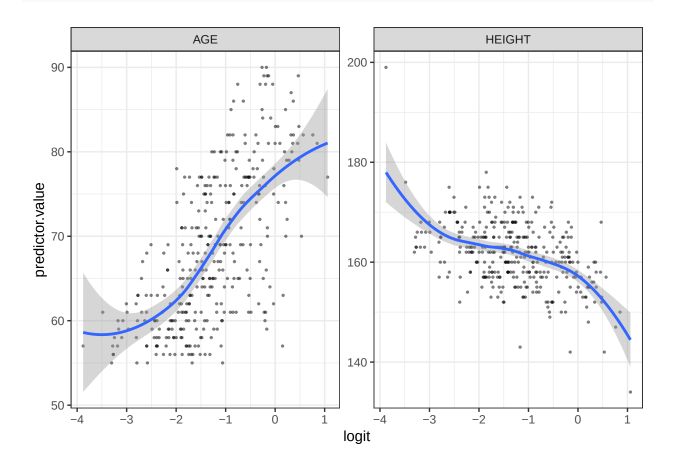
#ROC curves
pred.test.interaction <- prediction(logistic.fit.pred.test.interaction, validationData$FRACTURE)
roc.perf1 = performance(pred.test.interaction, measure = "tpr", x.measure = "fpr")
auc.val1 <- performance(pred.test.interaction, measure = "auc")
auc.val1 <- auc.val1@y.values
plot(roc.perf1, main="Logistic Reg With Interactions Validations Data Set")
abline(a=0, b= 1)
text(x = .40, y = .6,paste("AUC = ", round(auc.val1[[1]],3), sep = ""))</pre>
```

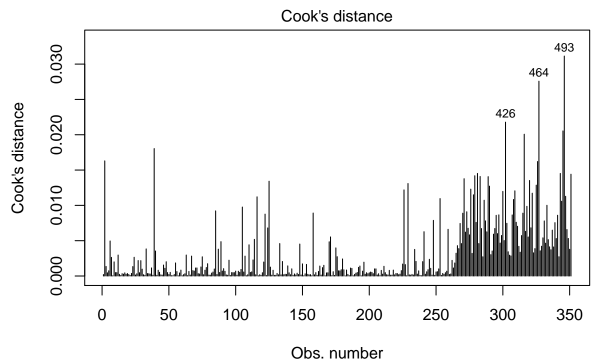
Logistic Reg With Interactions Validations Data Set



```
#confusion matrix
pdata_logical <- logistic.fit.pred.test.interaction > 0.5
confusionMatrix(data = as.factor(as.numeric(pdata_logical)), reference = as.factor(as.numeric(validation))
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction
                0
            0 105
                   30
##
##
##
                  Accuracy : 0.7517
##
                    95% CI: (0.6743, 0.8187)
##
##
       No Information Rate: 0.7517
       P-Value [Acc > NIR] : 0.5440183
##
##
##
                     Kappa : 0.16
    Mcnemar's Test P-Value : 0.0002983
##
##
##
               Sensitivity: 0.9375
##
               Specificity: 0.1892
##
            Pos Pred Value : 0.7778
##
            Neg Pred Value: 0.5000
##
                Prevalence: 0.7517
##
            Detection Rate: 0.7047
##
      Detection Prevalence: 0.9060
```

```
Balanced Accuracy: 0.5633
##
##
##
          'Positive' Class : 0
##
# Checking the assumptions
probabilities <- predict(logit.fit.interactions, type = "response")</pre>
predicted.classes <- ifelse(probabilities > 0.5, "pos", "neg")
head(predicted.classes)
       1
## "neg" "neg" "neg" "neg" "neg" "neg"
# Linearity assumption
subNumericPred <- trainingData %>% dplyr::select(AGE, HEIGHT)
predictors <- colnames(subNumericPred)</pre>
subNumericPred <- subNumericPred %>%
                  mutate(logit = log(probabilities/(1-probabilities))) %>%
                  gather(key = "predictors", value = "predictor.value", -logit)
ggplot(subNumericPred, aes(logit, predictor.value)) +
                geom_point(size = 0.5, alpha = 0.5) +
                geom_smooth(method = "loess") +
                theme_bw() +
                facet_wrap(~predictors, scales = "free_y")
```





1(FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC + ARMASSIST + RATERI

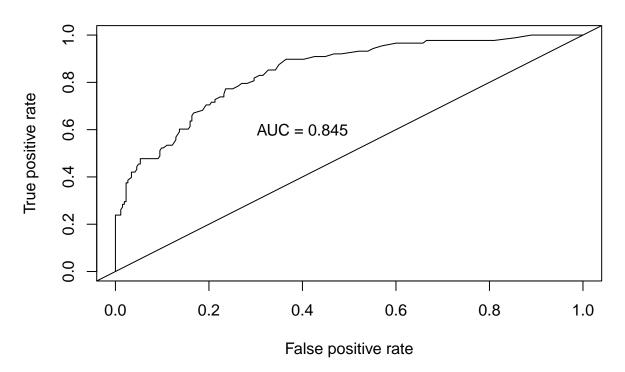
Running Random Forest Fit

```
set.seed(999)
str(trainingData)
```

```
'data.frame':
                    351 obs. of 11 variables:
##
   \ PRIORFRAC: Factor w/ 2 levels "0","1": 1 2 1 1 2 1 2 1 1 1 ...
               : int 62 88 82 61 67 84 86 58 67 56 ...
   $ WEIGHT
                      70.3 50.8 62.1 68 68 ...
               : num
##
   $ HEIGHT
                     158 157 160 152 161 150 156 166 153 167 ...
                     28.2 20.6 24.3 29.4 26.2 ...
##
   $ BMI
               : num
   $ PREMENO
              : Factor w/ 2 levels "0", "1": 1 1 1 1 1 1 1 1 1 ...
   $ MOMFRAC : Factor w/ 2 levels "0","1": 1 2 1 1 1 1 1 2 1 ...
##
   $ ARMASSIST: Factor w/ 2 levels "0","1": 1 2 1 1 1 1 1 1 1 2 ...
##
              : Factor w/ 2 levels "0", "1": 1 1 1 1 2 1 1 1 2 2 ...
   $ SMOKE
   $ RATERISK : Ord.factor w/ 3 levels "1"<"2"<"3": 2 1 1 2 2 1 2 1 1 2 ...</pre>
   $ FRACTURE : Factor w/ 2 levels "0", "1": 1 1 1 1 1 1 1 1 1 1 ...
```

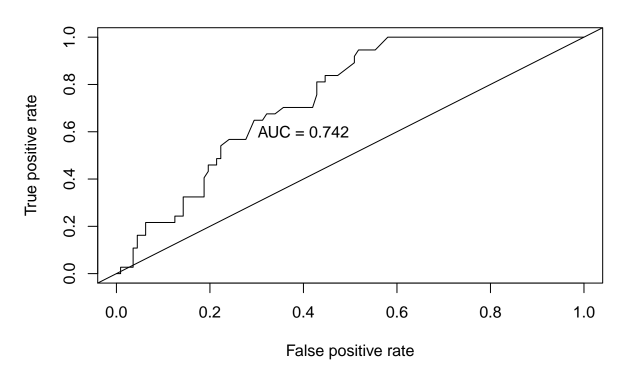
```
rf.fit <- randomForest(FRACTURE ~ ., data=trainingData, mtry=4, ntree=500, maxnodes = 12, importance=T)
rf.fit
##
## Call:
   Type of random forest: classification
                      Number of trees: 500
##
## No. of variables tried at each split: 4
##
          OOB estimate of error rate: 24.79%
##
## Confusion matrix:
      0 1 class.error
## 0 260 3 0.01140684
## 1 84 4 0.95454545
rf.fit.pred.train <- predict(rf.fit, newdata=trainingData, type="prob")</pre>
pred.train.rf <- prediction(rf.fit.pred.train[,2], trainingData$FRACTURE)</pre>
roc.perf = performance(pred.train.rf, measure = "tpr", x.measure = "fpr")
auc.train <- performance(pred.train.rf, measure = "auc")</pre>
auc.train <- auc.train@y.values</pre>
plot(roc.perf, main="Random Forest Training Data Set")
abline(a=0, b= 1)
text(x = .40, y = .6,paste("AUC = ", round(auc.train[[1]],3), sep = ""))
```

Random Forest Training Data Set



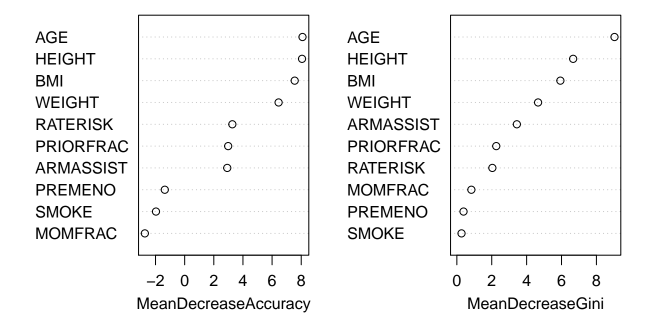
```
#confusion matrix Training
pdata_logical_train <- (rf.fit.pred.train[,2] >= 0.5)
confusionMatrix(data = as.factor(as.numeric(pdata logical train)), reference = as.factor(as.numeric(tra
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction 0 1
            0 263 76
##
##
               0 12
##
##
                  Accuracy : 0.7835
                    95% CI: (0.7367, 0.8254)
##
##
       No Information Rate: 0.7493
       P-Value [Acc > NIR] : 0.07672
##
##
##
                     Kappa : 0.1913
##
   Mcnemar's Test P-Value : < 2e-16
##
##
               Sensitivity: 1.0000
##
               Specificity: 0.1364
            Pos Pred Value: 0.7758
##
##
            Neg Pred Value: 1.0000
##
                Prevalence: 0.7493
##
            Detection Rate: 0.7493
##
      Detection Prevalence: 0.9658
##
         Balanced Accuracy: 0.5682
##
          'Positive' Class: 0
##
##
rf.fit.pred.test <- predict(rf.fit, newdata=validationData, type="prob")</pre>
pred.test.rf <- prediction(rf.fit.pred.test[,2], validationData$FRACTURE)</pre>
roc.perf = performance(pred.test.rf, measure = "tpr", x.measure = "fpr")
auc.train <- performance(pred.test.rf, measure = "auc")</pre>
auc.train <- auc.train@y.values</pre>
plot(roc.perf, main="Random Forest Validation Data Set")
abline(a=0, b= 1)
text(x = .40, y = .6,paste("AUC = ", round(auc.train[[1]],3), sep = ""))
```

Random Forest Validation Data Set



```
#confusion matrix Test
pdata_logical <- rf.fit.pred.test[,2] > 0.5
confusionMatrix(data = as.factor(as.numeric(pdata_logical)), reference = as.factor(as.numeric(validation))
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction
                0
            0 111 37
##
##
##
                  Accuracy: 0.745
##
                    95% CI: (0.6672, 0.8128)
##
##
       No Information Rate: 0.7517
       P-Value [Acc > NIR] : 0.6175
##
##
                     Kappa: -0.0132
##
    Mcnemar's Test P-Value : 1.365e-08
##
##
##
               Sensitivity: 0.9911
##
               Specificity: 0.0000
            Pos Pred Value : 0.7500
##
##
            Neg Pred Value: 0.0000
##
                Prevalence: 0.7517
##
            Detection Rate: 0.7450
##
      Detection Prevalence: 0.9933
```

```
##
         Balanced Accuracy: 0.4955
##
##
          'Positive' Class : 0
##
#confusion matrix Test Lower Cutoff
pdata_logical_lowercf <- rf.fit.pred.test[,2] >= 0.3
confusionMatrix(data = as.factor(as.numeric(pdata_logical_lowercf)), reference = as.factor(as.numeric(v))
## Confusion Matrix and Statistics
##
##
            Reference
## Prediction 0 1
           0 102 29
##
##
           1 10
##
##
                  Accuracy : 0.7383
##
                    95% CI: (0.66, 0.8068)
       No Information Rate: 0.7517
##
       P-Value [Acc > NIR] : 0.686582
##
##
##
                     Kappa : 0.1533
   Mcnemar's Test P-Value : 0.003948
##
##
##
               Sensitivity: 0.9107
##
               Specificity: 0.2162
            Pos Pred Value : 0.7786
##
##
            Neg Pred Value: 0.4444
##
                Prevalence: 0.7517
##
            Detection Rate: 0.6846
      Detection Prevalence: 0.8792
##
##
         Balanced Accuracy: 0.5635
##
##
          'Positive' Class : 0
##
varImpPlot(rf.fit)
```

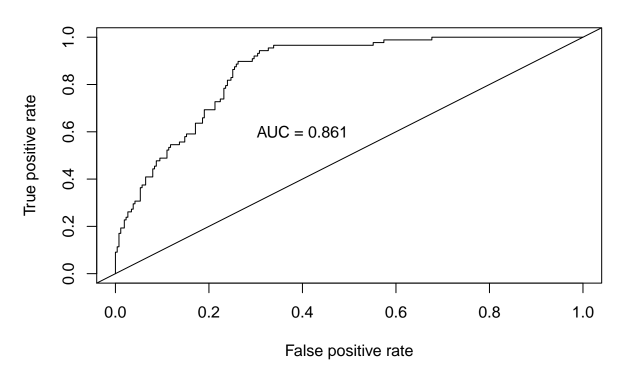


Running Conditional Random Forest Fit

```
set.seed(999)
crf.fit <- cforest(FRACTURE ~ ., data=trainingData, control=cforest_unbiased(ntree=500))</pre>
crf.fit
##
     Random Forest using Conditional Inference Trees
##
##
## Number of trees: 500
##
## Response: FRACTURE
## Inputs: PRIORFRAC, AGE, WEIGHT, HEIGHT, BMI, PREMENO, MOMFRAC, ARMASSIST, SMOKE, RATERISK
## Number of observations: 351
crf.fit.pred.train <- predict(crf.fit, newdata=trainingData, 00B = TRUE, type="prob")</pre>
unlist.Pred.train <- matrix(unlist(crf.fit.pred.train), ncol=2, byrow = TRUE)
pred.train.crf <- prediction(unlist.Pred.train[,2], trainingData$FRACTURE)</pre>
roc.perf = performance(pred.train.crf, measure = "tpr", x.measure = "fpr")
auc.train <- performance(pred.train.crf, measure = "auc")</pre>
auc.train <- auc.train@y.values</pre>
plot(roc.perf, main="Conditional Random Forest Training Data Set")
```

```
abline(a=0, b= 1)
text(x = .40, y = .6,paste("AUC = ", round(auc.train[[1]],3), sep = ""))
```

Conditional Random Forest Training Data Set

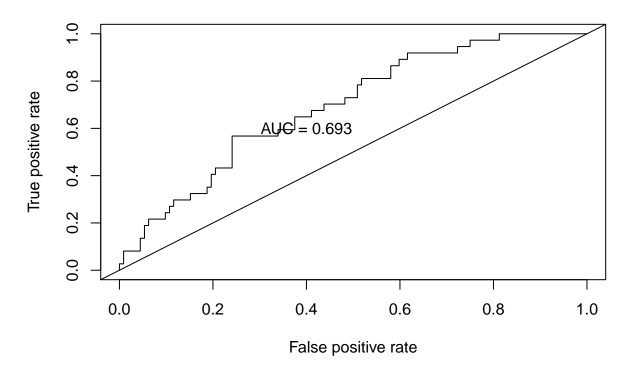


```
#confusion matrix Training
pdata_logical_train <- (unlist.Pred.train[,2] >= 0.5)
confusionMatrix(data = as.factor(as.numeric(pdata_logical_train)), reference = as.factor(as.numeric(tra
```

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction
##
            0 258 70
##
                5
                  18
##
##
                  Accuracy : 0.7863
                    95% CI : (0.7397, 0.8281)
##
       No Information Rate: 0.7493
##
       P-Value [Acc > NIR] : 0.06
##
##
##
                     Kappa: 0.246
##
   Mcnemar's Test P-Value: 1.467e-13
##
##
               Sensitivity: 0.9810
##
               Specificity: 0.2045
            Pos Pred Value: 0.7866
##
```

```
##
            Neg Pred Value: 0.7826
##
                Prevalence: 0.7493
            Detection Rate: 0.7350
##
      Detection Prevalence: 0.9345
##
##
         Balanced Accuracy: 0.5928
##
##
          'Positive' Class: 0
##
crf.fit.pred.test <- predict(crf.fit, newdata=validationData, 00B = T, type="prob")</pre>
unlist.Pred.test <- matrix(unlist(crf.fit.pred.test), ncol=2, byrow = TRUE)
pred.test.crf <- prediction(unlist.Pred.test[,2], validationData$FRACTURE)</pre>
roc.perf = performance(pred.test.crf, measure = "tpr", x.measure = "fpr")
auc.train <- performance(pred.test.crf, measure = "auc")</pre>
auc.train <- auc.train@y.values</pre>
plot(roc.perf, main="Conditional Random Forest Validation Data Set")
abline(a=0, b= 1)
text(x = .40, y = .6, paste("AUC = ", round(auc.train[[1]],3), sep = ""))
```

Conditional Random Forest Validation Data Set



```
#confusion matrix
pdata_logical <- unlist.Pred.test[,2] > 0.5
confusionMatrix(data = as.factor(as.numeric(pdata_logical)), reference = as.factor(as.numeric(validation))
```

Confusion Matrix and Statistics
##

```
##
             Reference
               0
                   1
## Prediction
##
            0 106 31
                6
                    6
##
            1
##
                  Accuracy : 0.7517
##
##
                    95% CI: (0.6743, 0.8187)
##
       No Information Rate: 0.7517
##
       P-Value [Acc > NIR] : 0.544
##
##
                     Kappa : 0.1403
    Mcnemar's Test P-Value: 7.961e-05
##
##
               Sensitivity: 0.9464
##
##
               Specificity: 0.1622
##
            Pos Pred Value: 0.7737
##
            Neg Pred Value: 0.5000
##
                Prevalence: 0.7517
##
            Detection Rate: 0.7114
##
      Detection Prevalence: 0.9195
##
         Balanced Accuracy: 0.5543
##
          'Positive' Class : 0
##
##
relativeImp <- varimp(crf.fit)</pre>
sort(relativeImp, decreasing = T)
##
             AGE
                         HEIGHT
                                    ARMASSIST
                                                         BMI
                                                                 PRIORFRAC
                  7.581395e-03 6.124031e-03 1.674419e-03
##
  8.372093e-03
                                                             4.496124e-04
##
          WEIGHT
                      RATERISK
                                        SMOKE
                                                     PREMENO
                                                                   MOMFRAC
   2.790698e-04 - 7.751938e-05 - 7.751938e-05 - 3.255814e-04 - 7.751938e-04
```

LDA AND QDA Model fit

```
library(MASS)
library(gridExtra)

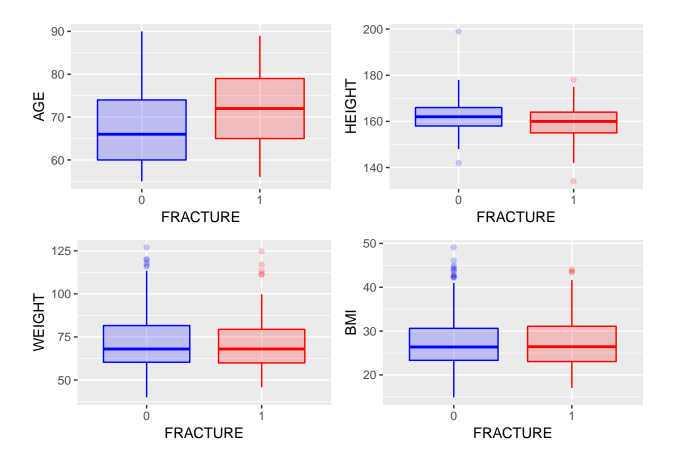
## Assumption of Eq Variance / CoVariance
box.AGE <- ggplot(dataset, aes(x = FRACTURE, y = AGE, col = FRACTURE, fill = FRACTURE)) +
    geom_boxplot(alpha = 0.2) +
    theme(legend.position = "none") +
    scale_color_manual(values = c("blue", "red")) +
    scale_fill_manual(values = c("blue", "red"))

box.HEIGHT <- ggplot(dataset, aes(x = FRACTURE, y = HEIGHT, col = FRACTURE, fill = FRACTURE)) +
    geom_boxplot(alpha = 0.2) +
    theme(legend.position = "none") +
    scale_color_manual(values = c("blue", "red")) +
    scale_fill_manual(values = c("blue", "red"))</pre>
```

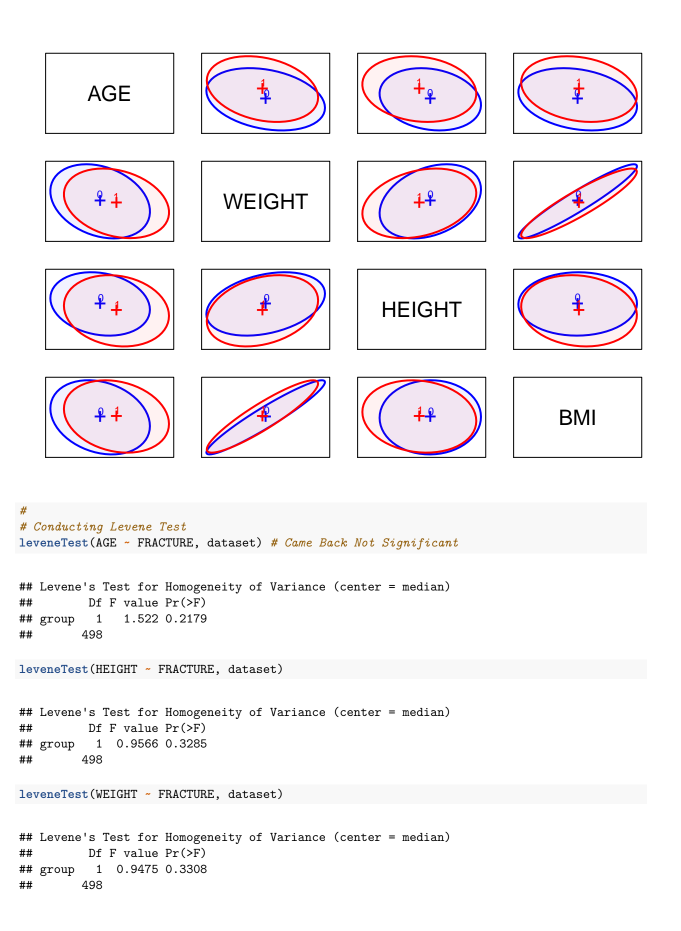
```
box.WEIGHT <- ggplot(dataset, aes(x = FRACTURE, y = WEIGHT, col = FRACTURE, fill = FRACTURE)) +
    geom_boxplot(alpha = 0.2) +
    theme(legend.position = "none") +
    scale_color_manual(values = c("blue", "red")) +
    scale_fill_manual(values = c("blue", "red"))

box.BMI <- ggplot(dataset, aes(x = FRACTURE, y = BMI, col = FRACTURE, fill = FRACTURE)) +
    geom_boxplot(alpha = 0.2) +
    theme(legend.position = "none") +
    scale_color_manual(values = c("blue", "red")) +
    scale_fill_manual(values = c("blue", "red"))

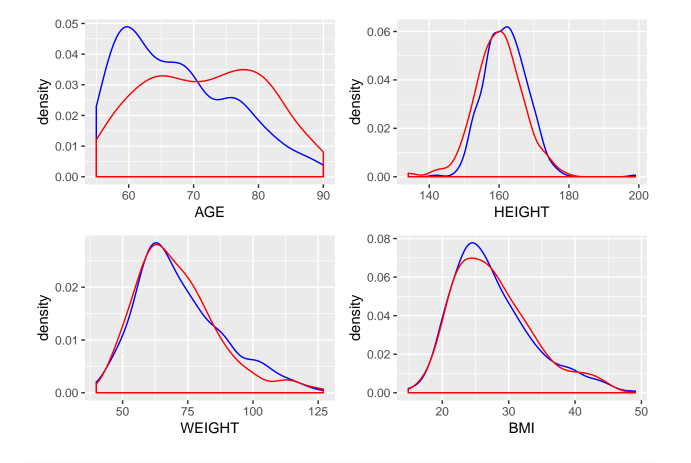
grid.arrange(box.AGE, box.HEIGHT, box.WEIGHT, box.BMI, nrow = 2, ncol = 2)</pre>
```



covEllipses(dataset[,c(5:8)], dataset\$FRACTURE, fill = TRUE, pooled = FALSE, col = c("blue", "red"), v.

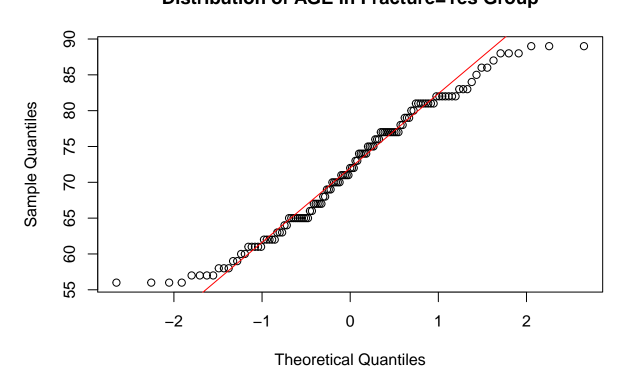


```
leveneTest(BMI ~ FRACTURE, dataset)
## Levene's Test for Homogeneity of Variance (center = median)
          Df F value Pr(>F)
## group 1 0.0188 0.8911
         498
##
# Came Back Not Significant, Confirms findings from previous plots
density.AGE <- ggplot(dataset, aes(x = AGE, y = ..density.., col = FRACTURE)) +</pre>
  geom_density(aes(y = ..density..)) +
  scale_color_manual(values = c("blue", "red")) +
  theme(legend.position = "none")
density.HEIGHT <- ggplot(dataset, aes(x = HEIGHT, y = ..density.., col = FRACTURE)) +</pre>
  geom density(aes(y = ..density..)) +
  scale_color_manual(values = c("blue", "red")) +
  theme(legend.position = "none")
density.WEIGHT <- ggplot(dataset, aes(x = WEIGHT, y = ..density.., col = FRACTURE)) +</pre>
  geom_density(aes(y = ..density..)) +
  scale_color_manual(values = c("blue", "red")) +
  theme(legend.position = "none")
density.BMI <- ggplot(dataset, aes(x = BMI, y = ..density.., col = FRACTURE)) +</pre>
  geom_density(aes(y = ..density..)) +
  scale_color_manual(values = c("blue", "red")) +
  theme(legend.position = "none")
grid.arrange(density.AGE, density.HEIGHT, density.WEIGHT, density.BMI, nrow = 2, ncol = 2)
```



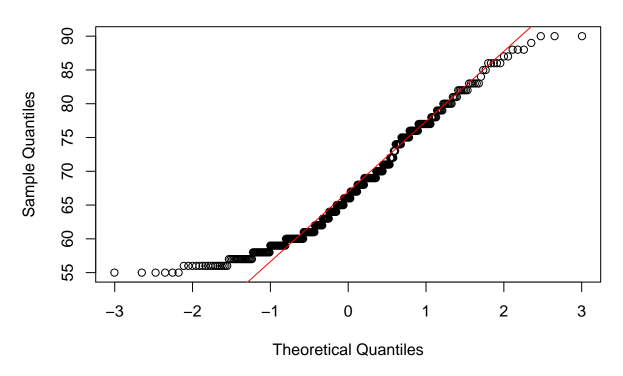
```
\textit{\# Check QQ Plot for AGE to ascertain Normality in BOTH Groups}
frac.yes <- subset(dataset, FRACTURE == 1)</pre>
frac.no <- subset(dataset, FRACTURE == 0)</pre>
# Plot
qqnorm(frac.yes$AGE, main = "Distribution of AGE in Fracture=Yes Group"); qqline(frac.yes$AGE, col = 2)
```

Distribution of AGE in Fracture=Yes Group



qqnorm(frac.no\$AGE, main = "Distribution of AGE in Fracture=No Group"); qqline(frac.no\$AGE, col = 2)

Distribution of AGE in Fracture=No Group

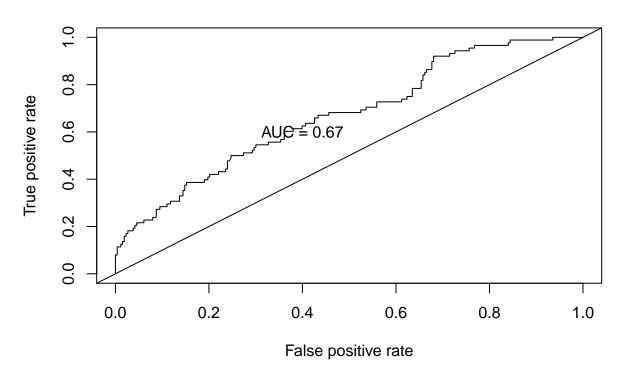


```
## Assumptions for Normality and of Equal Variance-Coavariance matrices Are Successfully Met.
## Run the LDA Now
set.seed(999)
lda.fit <- lda(FRACTURE ~ AGE + HEIGHT + WEIGHT + BMI, data = trainingData)</pre>
lda.fit
## Call:
## lda(FRACTURE ~ AGE + HEIGHT + WEIGHT + BMI, data = trainingData)
## Prior probabilities of groups:
## 0.7492877 0.2507123
##
## Group means:
          AGE
                HEIGHT
                         WEIGHT
## 0 67.16730 162.2129 72.01559 27.31879
## 1 71.95455 159.7614 70.53409 27.69421
##
## Coefficients of linear discriminants:
##
                  LD1
## AGE
           0.08790497
## HEIGHT 0.20784982
## WEIGHT -0.31576637
           0.86125425
## BMI
```

```
#ROC on training data set
ldaprd <- predict(lda.fit, newdata = trainingData)$posterior
ldaprd <- ldaprd[,2]
pred.train <- prediction(ldaprd, trainingData$FRACTURE)
roc.perf = performance(pred.train, measure = "tpr", x.measure = "fpr")
auc.train <- performance(pred.train, measure = "auc")
auc.train <- auc.train@y.values

#Plot ROC on Training Data
plot(roc.perf,main="LDA Training Data Set")
abline(a=0, b= 1) #Ref line indicating poor performance
text(x = .40, y = .6,paste("AUC = ", round(auc.train[[1]],3), sep = ""))</pre>
```

LDA Training Data Set



```
prd <- predict(lda.fit, newdata = trainingData)$class
confusionMatrix(data = prd, reference = trainingData$FRACTURE)</pre>
```

```
## Confusion Matrix and Statistics

##

## Reference

## Prediction 0 1

## 0 258 76

## 1 5 12

##

##

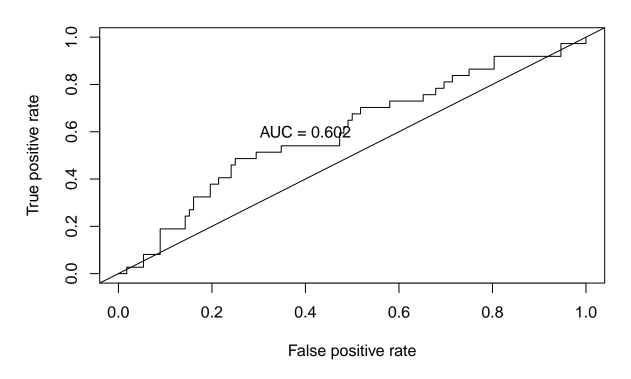
Accuracy: 0.7692

##

95% CI: (0.7216, 0.8123)
```

```
No Information Rate: 0.7493
##
       P-Value [Acc > NIR] : 0.2128
##
##
##
                     Kappa : 0.1604
   Mcnemar's Test P-Value: 7.381e-15
##
##
##
               Sensitivity: 0.9810
               Specificity: 0.1364
##
##
            Pos Pred Value: 0.7725
##
            Neg Pred Value: 0.7059
##
                Prevalence: 0.7493
##
            Detection Rate: 0.7350
##
      Detection Prevalence: 0.9516
##
         Balanced Accuracy: 0.5587
##
          'Positive' Class : 0
##
##
#ROC on test data set
ldaprd.test <- predict(lda.fit, newdata = validationData)$posterior</pre>
ldaprd.test <- ldaprd.test[,2]</pre>
pred.test <- prediction(ldaprd.test, validationData$FRACTURE)</pre>
roc.perf = performance(pred.test, measure = "tpr", x.measure = "fpr")
auc.train <- performance(pred.test, measure = "auc")</pre>
auc.train <- auc.train@y.values
#Plot ROC on Training Data
plot(roc.perf,main="LDA Validation Data Set")
abline(a=0, b= 1) #Ref line indicating poor performance
text(x = .40, y = .6,paste("AUC = ", round(auc.train[[1]],3), sep = ""))
```

LDA Validation Data Set

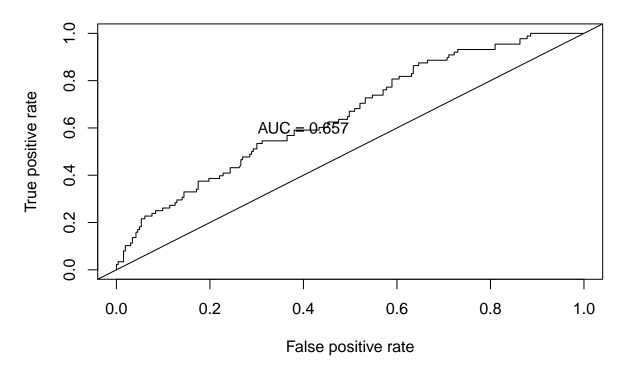


```
prd.test <- predict(lda.fit, newdata = validationData)$class
confusionMatrix(data = prd.test, reference = validationData$FRACTURE)</pre>
```

```
Confusion Matrix and Statistics
##
##
             Reference
  Prediction
##
                0
                    1
##
            0 106
                   34
##
            1
                6
                    3
##
                  Accuracy: 0.7315
##
##
                    95% CI: (0.6529, 0.8008)
##
       No Information Rate: 0.7517
       P-Value [Acc > NIR] : 0.7493
##
##
##
                     Kappa: 0.0368
    Mcnemar's Test P-Value : 1.963e-05
##
##
               Sensitivity: 0.94643
##
               Specificity: 0.08108
##
            Pos Pred Value: 0.75714
##
            Neg Pred Value: 0.33333
##
                Prevalence: 0.75168
##
##
            Detection Rate: 0.71141
##
      Detection Prevalence: 0.93960
```

```
##
         Balanced Accuracy: 0.51375
##
          'Positive' Class : 0
##
##
## Running QDA to see if it improves AUC
qda.fit <- qda(FRACTURE ~ AGE + HEIGHT + WEIGHT + BMI, data = trainingData)</pre>
#ROC on training data set
qdaprd <- predict(qda.fit, newdata = trainingData)$posterior</pre>
qdaprd <- qdaprd[,2]
pred.train <- prediction(qdaprd, trainingData$FRACTURE)</pre>
roc.perf = performance(pred.train, measure = "tpr", x.measure = "fpr")
auc.train <- performance(pred.train, measure = "auc")</pre>
auc.train <- auc.train@y.values</pre>
#Plot ROC on Training Data
plot(roc.perf,main="QDA Training Data Set")
abline(a=0, b= 1) #Ref line indicating poor performance
text(x = .40, y = .6, paste("AUC = ", round(auc.train[[1]],3), sep = ""))
```

QDA Training Data Set

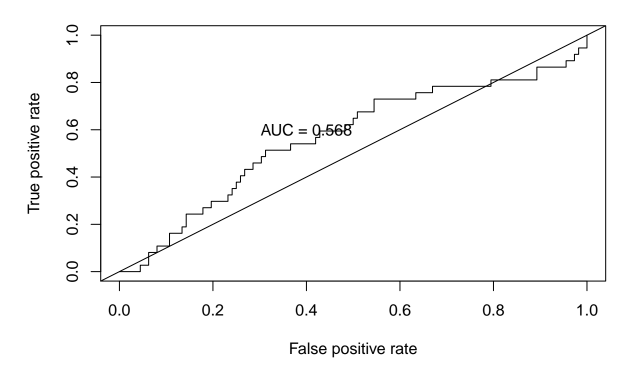


```
prd <- predict(qda.fit, newdata = trainingData)$class
confusionMatrix(data = prd, reference = trainingData$FRACTURE)</pre>
```

Confusion Matrix and Statistics

```
##
             Reference
##
               0 1
## Prediction
##
            0 249 71
            1 14 17
##
##
##
                  Accuracy : 0.7578
                    95% CI : (0.7095, 0.8017)
##
##
       No Information Rate: 0.7493
##
       P-Value [Acc > NIR] : 0.3827
##
##
                     Kappa: 0.1784
##
   Mcnemar's Test P-Value : 1.247e-09
##
##
               Sensitivity: 0.9468
               Specificity: 0.1932
##
##
            Pos Pred Value: 0.7781
            Neg Pred Value: 0.5484
##
##
                Prevalence: 0.7493
            Detection Rate: 0.7094
##
      Detection Prevalence : 0.9117
##
##
         Balanced Accuracy: 0.5700
##
          'Positive' Class : 0
##
##
#ROC on test data set
qdaprd.test <- predict(qda.fit, newdata = validationData)$posterior</pre>
qdaprd.test <- qdaprd.test[,2]</pre>
pred.test <- prediction(qdaprd.test, validationData$FRACTURE)</pre>
roc.perf = performance(pred.test, measure = "tpr", x.measure = "fpr")
auc.train <- performance(pred.test, measure = "auc")</pre>
auc.train <- auc.train@y.values</pre>
#Plot ROC on Training Data
plot(roc.perf,main="QDA Validation Data Set")
abline(a=0, b= 1) #Ref line indicating poor performance
text(x = .40, y = .6, paste("AUC = ", round(auc.train[[1]],3), sep = ""))
```

QDA Validation Data Set



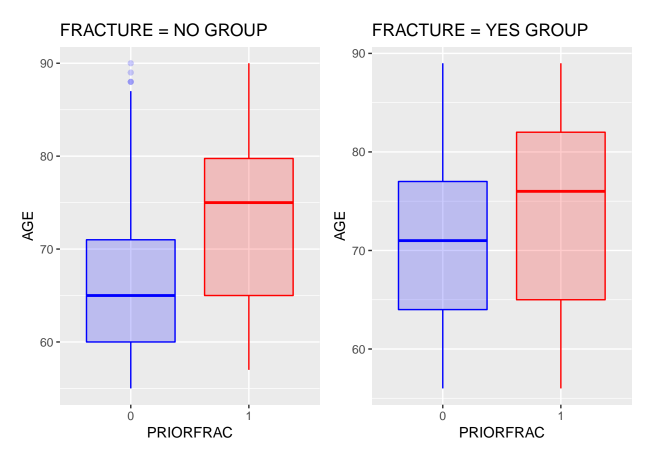
```
prd.test <- predict(qda.fit, newdata = validationData)$class
confusionMatrix(data = prd.test, reference = validationData$FRACTURE)</pre>
```

```
Confusion Matrix and Statistics
##
##
             Reference
  Prediction
##
                0
                    1
##
            0 103
                   33
##
            1
                9
                    4
##
                  Accuracy: 0.7181
##
##
                    95% CI : (0.6387, 0.7887)
##
       No Information Rate: 0.7517
       P-Value [Acc > NIR] : 0.8512971
##
##
##
                     Kappa: 0.0355
    Mcnemar's Test P-Value: 0.0003867
##
##
               Sensitivity: 0.9196
##
               Specificity: 0.1081
##
            Pos Pred Value: 0.7574
##
            Neg Pred Value: 0.3077
##
                Prevalence: 0.7517
##
##
            Detection Rate: 0.6913
##
      Detection Prevalence: 0.9128
```

```
##
          'Positive' Class : 0
##
##
frac.yes <- subset(dataset, FRACTURE == 1)</pre>
frac.no <- subset(dataset, FRACTURE == 0)</pre>
box.Prior.Age.Frac.Yes <- ggplot(frac.yes, aes(x = PRIORFRAC, y = AGE, col = PRIORFRAC, fill = PRIORFRA
  geom_boxplot(alpha = 0.2) +
  theme(legend.position = "none") +
  scale_color_manual(values = c("blue", "red")) +
  scale_fill_manual(values = c("blue", "red")) +
  ggtitle("FRACTURE = YES GROUP")
box.Prior.Age.Frac.No <- ggplot(frac.no, aes(x = PRIORFRAC, y = AGE, col = PRIORFRAC, fill = PRIORFRAC
  geom_boxplot(alpha = 0.2) +
  theme(legend.position = "none") +
  scale_color_manual(values = c("blue", "red")) +
  scale_fill_manual(values = c("blue", "red"))+
   ggtitle("FRACTURE = NO GROUP")
grid.arrange(box.Prior.Age.Frac.No, box.Prior.Age.Frac.Yes, nrow = 1, ncol = 2)
```

##

Balanced Accuracy: 0.5139



```
#MOMFRAC: ARMASSIST
par(mfrow = c(1, 2))
mosplot.Frac.No <- mosaicplot(CrossTable(frac.no$MOMFRAC, frac.no$ARMASSIST)$t, main = "FRACTURE = NO G
##
##
   Cell Contents
## |-----|
## | Chi-square contribution |
## | N / Row Total |
          N / Col Total |
      N / Table Total |
## |-----|
##
##
## Total Observations in Table: 375
##
         | frac.no$ARMASSIST
## frac.no$MOMFRAC | 0 | 1 | Row Total |
## -----|-----|
             0 | 225 | 109 |
                          0.326 | 0.891 |
0.872 |
             0.024
             | 0.674 | 0.326 |
| 0.900 | 0.872 |
| 0.600 | 0.291 |
##
##
## -----|-----|
              1 | 25 | 16 | 41 |
| 0.199 | 0.398 | |
| 0.610 | 0.390 | 0.109 |
           1 l
##
              | 0.100 | 0.128 |
| 0.067 | 0.043 |
##
## -----|-----|
    Column Total | 250 | 125 |
       0.667 | 0.333 |
     -----|-----|
##
##
mosplot.Frac.Yes <- mosaicplot(CrossTable(frac.yes$MOMFRAC, frac.yes$ARMASSIST)$t, main = "FRACTURE = Y.
##
##
    Cell Contents
## | Chi-square contribution |
## | N / Row Total | ## | N / Col Total |
```

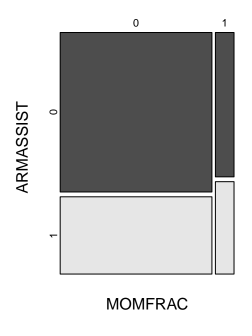
| N / Table Total |

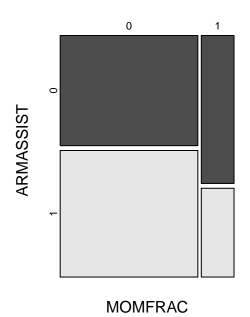
## ## ##	Total Observations		125	
##		111 14010.	120	
##				
##	frac.yes\$ARMASSIST			
##	<pre>frac.yes\$MOMFRAC </pre>	0	1	Row Total
##				
##	0	47	J 54	101
##		0.191	0.188	
##		0.465	0.535	0.808
##		0.758	0.857	
##		0.376	0.432	
##				
##	1	15	9	24
##		0.805	0.792	
##		0.625	0.375	0.192
##		0.242	0.143	! !
##		0.120	0.072	!
##				
##	Column Total	62	63	125
##	ļ	0.496	0.504	I I
##				
##				

FRACTURE = NO GROUP

##

FRACTURE = YES GROUP





*** Appendix C: Test interaction - LDA ==============

Get Glow dataset

```
glow <- read_glow_dataset()</pre>
```

model interactions - main effects

```
model_z1 <- glm(FRACTURE ~ AGE, family = binomial, data = glow)</pre>
model_z2 <- glm(FRACTURE ~ WEIGHT, family = binomial, data = glow)</pre>
model_z3 <- glm(FRACTURE ~ HEIGHT, family = binomial, data = glow)</pre>
model_z4 <- glm(FRACTURE ~ BMI, family = binomial, data = glow)</pre>
model_z5 <- glm(FRACTURE ~ PRIORFRAC, family = binomial, data = glow)</pre>
model_z6 <- glm(FRACTURE ~ PREMENO, family = binomial, data = glow)
model_z7 <- glm(FRACTURE ~ MOMFRAC, family = binomial, data = glow)</pre>
model_z8 <- glm(FRACTURE ~ ARMASSIST, family = binomial, data = glow)</pre>
model_z9 <- glm(FRACTURE ~ SMOKE, family = binomial, data = glow)</pre>
model_z10 <- glm(FRACTURE ~ RATERISK, family = binomial, data = glow)</pre>
## AGE
                0.05289
                            0.01163
                                      4.548 5.42e-06 ***
## WEIGHT
               -0.005197
                            0.006415 -0.810
                                                0.418
## HEIGHT
               -0.05167
                            0.01709 -3.022 0.00251 **
## BMI
                0.005758
                            0.017185 0.335 0.73760
## PRIORFRACYes 1.0638
                            0.2231
                                      4.769 1.85e-06 ***
               0.05077
                                     0.196
## PREMENOYes
                            0.25921
                                                0.845
## MOMFRACYes
              0.6605
                            0.2810
                                      2.351
                                              0.0187 *
## ARMASSISTYes 0.7091
                            0.2098
                                      3.381 0.000723 ***
                            0.4358
## SMOKEYes
              -0.3077
                                     -0.706
## RATERISKSame
                                 0.2664
                                           2.050
                                                   0.0404 *
                      0.5462
## RATERISKGreater
                     0.9091
                                 0.2711
                                           3.353
                                                   0.0008 ***
```

> code below:

This leads us to consider the covariates above that are significant in the univariate results above at the 25% level

AGE, HEIGHT, PRIORFRAC, MOMFRAC, ARMASSIST, RATERISK {SAME, GREATER}

```
# fit a univariate logistic regression model for each covariate
# continuous - AGE WEIGHT HEIGHT BMI
# categorical - PRIORFRAC PREMENO MOMFRAC ARMASSIST SMOKE RATERISK

# model0
#model_z1 <- glm(FRACTURE ~ AGE, family = binomial, data = glow)
#model_z2 <- glm(FRACTURE ~ WEIGHT, family = binomial, data = glow)
#model_z3 <- glm(FRACTURE ~ HEIGHT, family = binomial, data = glow)
#model_z4 <- glm(FRACTURE ~ BMI, family = binomial, data = glow)
#model_z5 <- glm(FRACTURE ~ PRIORFRAC, family = binomial, data = glow)
#model_z6 <- glm(FRACTURE ~ MOMFRAC, family = binomial, data = glow)
#model_z7 <- glm(FRACTURE ~ MOMFRAC, family = binomial, data = glow)
#model_z8 <- glm(FRACTURE ~ ARMASSIST, family = binomial, data = glow)
#model_z9 <- glm(FRACTURE ~ SMOKE, family = binomial, data = glow)</pre>
```

```
#model_z10 <- glm(FRACTURE ~ RATERISK, family = binomial, data = glow)</pre>
#summary(model_z1)
#summary(model_z2)
#summary(model_z3)
#summary(model_z4)
#summary(model_z5)
#summary(model z6)
#summary(model z7)
#summary(model z8)
#summary(model_z9)
#summary(model_z10)
# not interesting due to all variables (i.e. SUB_ID, SITE_ID, PHY_ID)
# model00 <- qlm(FRACTURE ~ ., family = binomial, data = qlow)
# summary(model00)
# full model, order by continuous, then factor
model0 <- glm(FRACTURE ~ AGE + WEIGHT + HEIGHT + BMI + PRIORFRAC + PREMENO + MOMFRAC + ARMASSIST + SMOK
summary(model0)
##
## Call:
  glm(formula = FRACTURE ~ AGE + WEIGHT + HEIGHT + BMI + PRIORFRAC +
##
       PREMENO + MOMFRAC + ARMASSIST + SMOKE + RATERISK, family = binomial,
##
       data = glow)
##
## Deviance Residuals:
##
      Min
              1Q
                     Median
                                  3Q
                                          Max
## -1.6811 -0.7228 -0.5639 -0.1008
                                       2.2182
##
## Coefficients:
##
                   Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                  -15.74709 12.67053 -1.243 0.21394
## AGE
                    0.03895
                               0.01476
                                        2.640 0.00829 **
## WEIGHT
                               0.08664 -1.407 0.15949
                   -0.12189
## HEIGHT
                    0.06620
                               0.07825
                                         0.846 0.39755
                             0.22339
## BMI
                    0.33181
                                         1.485 0.13745
## PRIORFRACYes
                    0.67577 0.25012
                                         2.702 0.00690 **
## PREMENOYes
                    0.10080
                             0.28540
                                         0.353 0.72395
## MOMFRACYes
                    0.63438
                               0.30784
                                         2.061 0.03933 *
## ARMASSISTYes
                    0.36102 0.25647
                                         1.408 0.15924
## SMOKEYes
                   -0.31228
                               0.46216 -0.676 0.49923
                                         1.501 0.13324
                    0.42256
                               0.28144
## RATERISKSame
## RATERISKGreater
                    0.75645
                               0.29944
                                         2.526 0.01153 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 562.34 on 499 degrees of freedom
## Residual deviance: 503.84 on 488 degrees of freedom
## AIC: 527.84
```

```
##
## Number of Fisher Scoring iterations: 4
# fit model # note - should remove below model1
model0_fitted <- update(model0, . ~ . - WEIGHT - BMI - PREMENO - SMOKE)</pre>
summary(model0 fitted)
##
## Call:
## glm(formula = FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC +
      ARMASSIST + RATERISK, family = binomial, data = glow)
##
## Deviance Residuals:
       Min
                  1Q
                        Median
                                      3Q
                                               Max
## -1.66692 -0.72502 -0.56338 -0.03841
## Coefficients:
##
                  Estimate Std. Error z value Pr(>|z|)
                              3.22992 0.839 0.40157
## (Intercept)
                   2.70935
## AGE
                   0.03434
                              0.01305
                                       2.632 0.00848 **
                              0.01827 -2.400 0.01640 *
## HEIGHT
                  -0.04383
## PRIORFRACYes
                   0.64526
                              0.24606
                                       2.622 0.00873 **
                   0.62122
                              0.30698
                                      2.024 0.04300 *
## MOMFRACYes
## ARMASSISTYes
                   0.44579
                              0.23281
                                       1.915 0.05551 .
## RATERISKSame
                   0.42202
                              0.27925
                                       1.511 0.13071
## RATERISKGreater 0.70692
                              0.29342
                                       2.409 0.01599 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 562.34 on 499 degrees of freedom
## Residual deviance: 507.50 on 492 degrees of freedom
## AIC: 523.5
##
## Number of Fisher Scoring iterations: 4
# build model with following covariates (drop WEIGHT, BMI, PREMENO, SMOKE)
# AGE, HEIGHT, PRIORFRAC, MOMFRAC, ARMASSIST, RATERISK {SAME, GREATER}
model1 <- glm(FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC + ARMASSIST + RATERISK, family = binomial,
summary(model1)
##
## Call:
## glm(formula = FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC +
      ARMASSIST + RATERISK, family = binomial, data = glow)
##
## Deviance Residuals:
       Min
                  1Q
                        Median
                                      3Q
                                               Max
## -1.66692 -0.72502 -0.56338 -0.03841
```

Coefficients:

```
Estimate Std. Error z value Pr(>|z|)
                          3.22992 0.839 0.40157
## (Intercept)
                 2.70935
## AGE
                 ## HEIGHT
## PRIORFRACYes
                 0.64526
                          0.24606 2.622 0.00873 **
                ## MOMFRACYes
                0.44579 0.23281 1.915 0.05551 .
## ARMASSISTYes
                 0.42202 0.27925 1.511 0.13071
## RATERISKSame
## RATERISKGreater 0.70692
                           0.29342 2.409 0.01599 *
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 562.34 on 499 degrees of freedom
## Residual deviance: 507.50 on 492 degrees of freedom
## AIC: 523.5
##
## Number of Fisher Scoring iterations: 4
# from above result, adding back the removed covariates we see they are not needed to keep the remainin
# this becomes the model, adding back removed covariates WEIGHT, BMI, PREMENO, SMOKE the coefficients d
# this becomes the main effects model
\textit{\# need to check scale of logit for remaining continous variables} \textit{ AGE HEIGHT}
# assume HEIGHT is linearin logit
```

The main effects model

```
model1 <- glm(FRACTURE \sim AGE + HEIGHT + PRIORFRAC + MOMFRAC + ARMASSIST + RATERISK, family = binomial, data = glow)
```

use lrtest from package lmtest

test interactions for the following:

- 5. AGE: [HEIGHT, PRIORFRAC, MOMFRAC, ARMASSIST, RATERISK]
- 6. HEIGHT: [PRIORFRAC, MOMFRAC, ARMASSIST, RATERISK]
- 7. PRIORFRAC: [MOMFRAC, ARMASSIST, RATERISK]
- 8. MOMFRAC: [ARMASSIST, RATERISK]
- 9. ARMASSIST: RATERISK

total 15 interactions

```
library(lmtest)
# model AGE* , HEIGHT* , PRIORFRAC*

model_effects <- glm(FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC + ARMASSIST + RATERISK, family = bin
lrtest(model effects)</pre>
```

```
## Likelihood ratio test
##
## Model 1: FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC + ARMASSIST + RATERISK
## Model 2: FRACTURE ~ 1
   #Df LogLik Df Chisq Pr(>Chisq)
## 1 8 -253.75
## 2 1 -281.17 -7 54.835 1.608e-09 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
# (5) AGE: [HEIGHT, PRIORFRAC, MOMFRAC, ARMASSIST, RATERISK]
test <- model effects
test <- update(test, . ~ . + AGE:HEIGHT)</pre>
summary(test)
##
## Call:
## glm(formula = FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC +
      ARMASSIST + RATERISK + AGE: HEIGHT, family = binomial, data = glow)
##
## Deviance Residuals:
                       Median
       Min
                 1Q
## -1.66848 -0.73323 -0.56252 0.02069
                                          2.23640
##
## Coefficients:
                   Estimate Std. Error z value Pr(>|z|)
                 14.749125 23.931667 0.616 0.5377
## (Intercept)
                  -0.135869 0.335087 -0.405 0.6851
## AGE
## HEIGHT
                 -0.119095 0.149402 -0.797 0.4254
## PRIORFRACYes
                  0.634947 0.246751 2.573 0.0101 *
                 ## MOMFRACYes
## ARMASSISTYes 0.447271 0.232895 1.920 0.0548 .
## RATERISKSame 0.435127 0.280319 1.552 0.1206
## RATERISKGreater 0.707865 0.293394 2.413 0.0158 *
## AGE:HEIGHT
                0.001065 0.002095 0.508 0.6113
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 562.34 on 499 degrees of freedom
## Residual deviance: 507.24 on 491 degrees of freedom
## AIC: 525.24
##
## Number of Fisher Scoring iterations: 4
lrtest(test)
## Likelihood ratio test
## Model 1: FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC + ARMASSIST + RATERISK +
      AGE: HEIGHT
##
```

```
## Model 2: FRACTURE ~ 1
## #Df LogLik Df Chisq Pr(>Chisq)
## 1 9 -253.62
    1 -281.17 -8 55.096 4.23e-09 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
test <- model effects
test <- update(test, . ~ . + AGE:PRIORFRAC)</pre>
summary(test)
##
## Call:
## glm(formula = FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC +
##
      ARMASSIST + RATERISK + AGE: PRIORFRAC, family = binomial,
##
      data = glow)
##
## Deviance Residuals:
      \mathtt{Min}
                1Q
                     Median
                                  3Q
                                          Max
## -1.48423 -0.74080 -0.53895 -0.00078
                                      2.26588
##
## Coefficients:
##
                 Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                0.63708 3.35881 0.190 0.849565
                 ## AGE
## HEIGHT
                 ## PRIORFRACYes
                ## MOMFRACYes
                 0.41887 0.23395 1.790 0.073391 .
## ARMASSISTYes
                  0.43496 0.28053 1.551 0.121014
## RATERISKSame
## RATERISKGreater 0.72044 0.29561 2.437 0.014804 *
## AGE:PRIORFRACYes -0.05864 0.02583 -2.270 0.023188 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
      Null deviance: 562.34 on 499 degrees of freedom
## Residual deviance: 502.34 on 491 degrees of freedom
## AIC: 520.34
## Number of Fisher Scoring iterations: 4
lrtest(test)
## Likelihood ratio test
## Model 1: FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC + ARMASSIST + RATERISK +
      AGE: PRIORFRAC
## Model 2: FRACTURE ~ 1
   #Df LogLik Df Chisq Pr(>Chisq)
## 1 9 -251.17
## 2 1 -281.17 -8 59.991 4.679e-10 ***
```

```
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
test <- model_effects</pre>
test <- update(test, . ~ . + AGE:MOMFRAC)</pre>
summary(test)
##
## Call:
## glm(formula = FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC +
      ARMASSIST + RATERISK + AGE: MOMFRAC, family = binomial, data = glow)
##
## Deviance Residuals:
      Min
                1Q
                      Median
                                  3Q
                                          Max
## -1.58376 -0.72859 -0.56182 -0.02562
##
## Coefficients:
                Estimate Std. Error z value Pr(>|z|)
##
                2.59055 3.24368 0.799 0.42450
## (Intercept)
                 ## AGE
## HEIGHT
                ## PRIORFRACYes
                ## MOMFRACYes
                1.57119 2.31121 0.680 0.49662
                 0.45447 0.23374 1.944 0.05185 .
## ARMASSISTYes
                 0.42505 0.27940 1.521 0.12819
## RATERISKSame
## RATERISKGreater 0.71044 0.29363 2.420 0.01554 *
## AGE:MOMFRACYes -0.01353 0.03264 -0.414 0.67854
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 562.34 on 499 degrees of freedom
## Residual deviance: 507.33 on 491 degrees of freedom
## AIC: 525.33
## Number of Fisher Scoring iterations: 4
lrtest(test)
## Likelihood ratio test
## Model 1: FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC + ARMASSIST + RATERISK +
      AGE: MOMFRAC
## Model 2: FRACTURE ~ 1
## #Df LogLik Df Chisq Pr(>Chisq)
## 1 9 -253.66
## 2 1 -281.17 -8 55.005 4.406e-09 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

```
test <- model_effects</pre>
test <- update(test, . ~ . + AGE:ARMASSIST)</pre>
summary(test)
##
## glm(formula = FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC +
      ARMASSIST + RATERISK + AGE: ARMASSIST, family = binomial,
##
      data = glow)
##
## Deviance Residuals:
     Min
          10
                  Median
                            3Q
## -1.6352 -0.7272 -0.5646 -0.0295
                                   2.2329
## Coefficients:
##
                 Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                 2.33972 3.33003 0.703 0.48230
                 0.03990 0.01785
                                    2.235 0.02542 *
## AGE
## HEIGHT
                 ## PRIORFRACYes
                ## MOMFRACYes
                 0.63376 0.30795
                                   2.058 0.03959 *
                 1.24419 1.76410 0.705 0.48063
## ARMASSISTYes
                  0.42815 0.27964
## RATERISKSame
                                    1.531 0.12575
## RATERISKGreater 0.71996 0.29494 2.441 0.01464 *
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 562.34 on 499 degrees of freedom
## Residual deviance: 507.29 on 491 degrees of freedom
## AIC: 525.29
## Number of Fisher Scoring iterations: 4
lrtest(test)
## Likelihood ratio test
## Model 1: FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC + ARMASSIST + RATERISK +
     AGE: ARMASSIST
## Model 2: FRACTURE ~ 1
## #Df LogLik Df Chisq Pr(>Chisq)
## 1 9 -253.65
     1 -281.17 -8 55.043 4.331e-09 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
test <- model_effects</pre>
test <- update(test, . ~ . + AGE:RATERISK)</pre>
summary(test)
```

```
##
## Call:
## glm(formula = FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC +
      ARMASSIST + RATERISK + AGE: RATERISK, family = binomial, data = glow)
## Deviance Residuals:
           10
                      Median
                                   30
                                           Max
## -1.68668 -0.74463 -0.56590 -0.02638
                                       2.34976
##
## Coefficients:
                    Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                     0.53632 3.53444 0.152 0.87939
## AGE
                     0.06673
                               0.02473 2.698 0.00697 **
## HEIGHT
                    ## PRIORFRACYes
                     ## MOMFRACYes
                     0.65241
                               0.30765
                                         2.121 0.03395 *
                               0.23443 2.072 0.03828 *
## ARMASSISTYes
                   0.48569
## RATERISKSame
                    3.28427 2.27575 1.443 0.14898
## RATERISKGreater
                    4.25804
                             2.28873 1.860 0.06282 .
## AGE:RATERISKSame -0.03999
                              0.03151 -1.269 0.20438
## AGE:RATERISKGreater -0.05021
                             0.03202 -1.568 0.11690
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 562.34 on 499 degrees of freedom
## Residual deviance: 504.79 on 490 degrees of freedom
## AIC: 524.79
##
## Number of Fisher Scoring iterations: 5
lrtest(test)
## Likelihood ratio test
## Model 1: FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC + ARMASSIST + RATERISK +
      AGE: RATERISK
## Model 2: FRACTURE ~ 1
## #Df LogLik Df Chisq Pr(>Chisq)
## 1 10 -252.40
## 2 1 -281.17 -9 57.54 3.982e-09 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
# (4) HEIGHT: [PRIORFRAC, MOMFRAC, ARMASSIST, RATERISK]
test <- model_effects</pre>
test <- update(test, . ~ . + HEIGHT:PRIORFRAC)</pre>
summary(test)
##
## Call:
```

```
## glm(formula = FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC +
##
      ARMASSIST + RATERISK + HEIGHT: PRIORFRAC, family = binomial,
       data = glow)
##
##
## Deviance Residuals:
                    Median
      Min
                1Q
                                  3Q
                                          Max
## -1.6670 -0.7274 -0.5615 -0.0037
##
## Coefficients:
##
                      Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                       3.79297
                                 3.89138 0.975 0.32970
                                           2.597 0.00941 **
                       0.03395
                                 0.01307
## AGE
## HEIGHT
                      -0.05041
                                0.02253 -2.238 0.02524 *
                               6.03699 -0.401 0.68869
## PRIORFRACYes
                      -2.41864
## MOMFRACYes
                       0.63692
                                0.30850
                                          2.065 0.03896 *
## ARMASSISTYes
                       0.43526
                                 0.23394
                                           1.861
                                                  0.06281 .
                               0.27946
## RATERISKSame
                       0.42634
                                           1.526 0.12711
## RATERISKGreater
                       0.70410
                               0.29356 2.399 0.01646 *
## HEIGHT:PRIORFRACYes 0.01915
                               0.03770 0.508 0.61146
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
       Null deviance: 562.34 on 499 degrees of freedom
## Residual deviance: 507.24 on 491 degrees of freedom
## AIC: 525.24
## Number of Fisher Scoring iterations: 4
lrtest(test)
## Likelihood ratio test
## Model 1: FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC + ARMASSIST + RATERISK +
      HEIGHT: PRIORFRAC
## Model 2: FRACTURE ~ 1
## #Df LogLik Df Chisq Pr(>Chisq)
## 1 9 -253.62
      1 -281.17 -8 55.092 4.236e-09 ***
## 2
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
test <- model_effects</pre>
test <- update(test, . ~ . + HEIGHT:MOMFRAC)</pre>
summary(test)
##
## Call:
## glm(formula = FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC +
##
       ARMASSIST + RATERISK + HEIGHT: MOMFRAC, family = binomial,
##
       data = glow)
##
```

```
## Deviance Residuals:
      Min
           10
                     Median
                                  30
                                          Max
## -1.62068 -0.74163 -0.55649 0.06604
## Coefficients:
                   Estimate Std. Error z value Pr(>|z|)
##
## (Intercept)
                   4.73834 3.51132 1.349 0.17719
                   ## AGE
                   -0.05646 0.02021 -2.794 0.00521 **
## HEIGHT
## PRIORFRACYes
                   ## MOMFRACYes
                  -11.35526 7.64959 -1.484 0.13770
                   ## ARMASSISTYes
                   0.42455 0.28002 1.516 0.12949
## RATERISKSame
                    ## RATERISKGreater
## HEIGHT:MOMFRACYes 0.07401 0.04718 1.569 0.11675
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 562.34 on 499 degrees of freedom
## Residual deviance: 505.08 on 491 degrees of freedom
## AIC: 523.08
## Number of Fisher Scoring iterations: 4
lrtest(test)
## Likelihood ratio test
## Model 1: FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC + ARMASSIST + RATERISK +
      HEIGHT: MOMFRAC
## Model 2: FRACTURE ~ 1
   #Df LogLik Df Chisq Pr(>Chisq)
## 1 9 -252.54
## 2 1 -281.17 -8 57.258 1.603e-09 ***
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
test <- model_effects</pre>
test <- update(test, . ~ . + HEIGHT:ARMASSIST)</pre>
summary(test)
##
## Call:
## glm(formula = FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC +
##
      ARMASSIST + RATERISK + HEIGHT: ARMASSIST, family = binomial,
##
      data = glow)
##
## Deviance Residuals:
      Min
               1Q
                  Median
                              3Q
                                     Max
## -1.6742 -0.7177 -0.5638 -0.1472
                                   2.1734
##
```

```
## Coefficients:
##
                     Estimate Std. Error z value Pr(>|z|)
                    -0.57428 4.12234 -0.139 0.88920
## (Intercept)
## AGE
                      0.03401
                                 0.01308 2.601 0.00931 **
## HEIGHT
                     -0.02318
                                 0.02432 -0.953 0.34051
## PRIORFRACYes
                     0.67913 0.24841
                                          2.734 0.00626 **
                      0.58729 0.30807 1.906 0.05660 .
## MOMFRACYes
                               5.77628 1.305 0.19179
## ARMASSISTYes
                      7.53985
## RATERISKSame
                      0.41583
                                 0.27981
                                           1.486 0.13725
## RATERISKGreater
                      0.70729
                                 0.29369 2.408 0.01603 *
## HEIGHT: ARMASSISTYes -0.04419
                                 0.03594 -1.229 0.21890
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 562.34 on 499 degrees of freedom
## Residual deviance: 505.98 on 491 degrees of freedom
## AIC: 523.98
##
## Number of Fisher Scoring iterations: 4
lrtest(test)
## Likelihood ratio test
##
## Model 1: FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC + ARMASSIST + RATERISK +
      HEIGHT: ARMASSIST
## Model 2: FRACTURE ~ 1
   #Df LogLik Df Chisq Pr(>Chisq)
## 1 9 -252.99
## 2 1 -281.17 -8 56.352 2.409e-09 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
test <- model_effects</pre>
test <- update(test, . ~ . + HEIGHT:RATERISK)</pre>
summary(test)
##
## Call:
## glm(formula = FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC +
##
      ARMASSIST + RATERISK + HEIGHT: RATERISK, family = binomial,
##
      data = glow)
##
## Deviance Residuals:
       Min
                  1Q
                        Median
                                     3Q
                                              Max
## -1.64936 -0.72375 -0.57251 -0.05841
##
## Coefficients:
##
                         Estimate Std. Error z value Pr(>|z|)
                         3.25641 5.81516 0.560 0.57549
## (Intercept)
                                    0.01310 2.536 0.01122 *
## AGE
                          0.03321
```

```
## HEIGHT
                     ## PRIORFRACYes
## MOMFRACYes
                     ## ARMASSISTYes
                     0.44610 0.23290 1.915 0.05544 .
## RATERISKSame
                      2.93823
                               7.29965 0.403 0.68730
## RATERISKGreater
                    -3.15056 7.29448 -0.432 0.66581
## HEIGHT:RATERISKSame -0.01577 0.04550 -0.347 0.72890
## HEIGHT:RATERISKGreater 0.02394
                              0.04528 0.529 0.59695
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
     Null deviance: 562.34 on 499 degrees of freedom
##
## Residual deviance: 506.55 on 490 degrees of freedom
## AIC: 526.55
## Number of Fisher Scoring iterations: 4
lrtest(test)
## Likelihood ratio test
## Model 1: FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC + ARMASSIST + RATERISK +
     HEIGHT: RATERISK
##
## Model 2: FRACTURE ~ 1
  #Df LogLik Df Chisq Pr(>Chisq)
## 1 10 -253.28
## 2
    1 -281.17 -9 55.786 8.624e-09 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
# (3) PRIORFRAC: [MOMFRAC, ARMASSIST, RATERISK]
test <- model_effects</pre>
test <- update(test, . ~ . + PRIORFRAC:MOMFRAC)</pre>
summary(test)
##
## Call:
## glm(formula = FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC +
     ARMASSIST + RATERISK + PRIORFRAC: MOMFRAC, family = binomial,
##
     data = glow)
##
##
## Deviance Residuals:
            1Q
                     Median
                                3Q
      Min
                                        Max
## -1.52616 -0.73215 -0.54992 0.02399
                                     2.25279
## Coefficients:
                     Estimate Std. Error z value Pr(>|z|)
##
## (Intercept)
                      2.97592 3.23781 0.919 0.35804
## AGE
                      ## HEIGHT
```

```
## PRIORFRACYes
                         0.80102 0.26285
                                             3.047 0.00231 **
                         ## MOMFRACYes
## ARMASSISTYes
                         0.43294 0.23384 1.851 0.06411 .
                         0.41959
## RATERISKSame
                                             1.497 0.13437
                                   0.28027
## RATERISKGreater
                         0.71282
                                    0.29401
                                             2.425 0.01533 *
## PRIORFRACYes: MOMFRACYes -1.07823 0.65021 -1.658 0.09726 .
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
      Null deviance: 562.34 on 499 degrees of freedom
##
## Residual deviance: 504.75 on 491 degrees of freedom
## AIC: 522.75
##
## Number of Fisher Scoring iterations: 4
lrtest(test)
## Likelihood ratio test
##
## Model 1: FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC + ARMASSIST + RATERISK +
      PRIORFRAC: MOMFRAC
## Model 2: FRACTURE ~ 1
   #Df LogLik Df Chisq Pr(>Chisq)
## 1 9 -252.37
## 2 1 -281.17 -8 57.59 1.382e-09 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
test <- model_effects</pre>
test <- update(test, . ~ . + PRIORFRAC:ARMASSIST)</pre>
summary(test)
##
## Call:
## glm(formula = FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC +
      ARMASSIST + RATERISK + PRIORFRAC: ARMASSIST, family = binomial,
##
##
      data = glow)
##
## Deviance Residuals:
##
       Min
                 1Q
                       Median
                                    3Q
                                            Max
## -1.69860 -0.71874 -0.56691 -0.04199
                                        2.21033
##
## Coefficients:
                          Estimate Std. Error z value Pr(>|z|)
                                   3.25923 0.892 0.37241
## (Intercept)
                           2.90711
## AGE
                           0.03434
                                     0.01306 2.630 0.00854 **
                                    0.01842 -2.436 0.01487 *
## HEIGHT
                          -0.04486
## PRIORFRACYes
                           0.52412
                                     0.34418
                                               1.523 0.12780
## MOMFRACYes
                           0.63247 0.30798
                                               2.054 0.04001 *
## ARMASSISTYes
                           0.36456
                                   0.28322 1.287 0.19803
                           ## RATERISKSame
```

```
## RATERISKGreater
                            0.68837
                                    0.29591
                                               2.326 0.02000 *
## PRIORFRACYes: ARMASSISTYes 0.24587
                                   0.48467 0.507 0.61194
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 562.34 on 499 degrees of freedom
## Residual deviance: 507.24 on 491 degrees of freedom
## AIC: 525.24
##
## Number of Fisher Scoring iterations: 4
lrtest(test)
## Likelihood ratio test
##
## Model 1: FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC + ARMASSIST + RATERISK +
      PRIORFRAC: ARMASSIST
## Model 2: FRACTURE ~ 1
   #Df LogLik Df Chisq Pr(>Chisq)
## 1 9 -253.62
## 2 1 -281.17 -8 55.093 4.235e-09 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
test <- model_effects</pre>
test <- update(test, . ~ . + PRIORFRAC:RATERISK)</pre>
summary(test)
##
## Call:
## glm(formula = FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC +
      ARMASSIST + RATERISK + PRIORFRAC: RATERISK, family = binomial,
##
      data = glow)
##
## Deviance Residuals:
                1Q
       Min
                       Median
                                    3Q
                                            Max
## -1.69776 -0.71989 -0.56384 -0.03822
## Coefficients:
                              Estimate Std. Error z value Pr(>|z|)
##
## (Intercept)
                              2.733523 3.235459 0.845 0.39819
                                                  2.642 0.00823 **
## AGE
                              0.034508
                                       0.013060
## HEIGHT
                             -0.043896
                                       0.018313 -2.397 0.01653 *
## PRIORFRACYes
                              0.564292
                                        0.497212
                                                   1.135 0.25641
                              0.623104 0.307302
                                                 2.028 0.04260 *
## MOMFRACYes
## ARMASSISTYes
                              0.429891 0.236033 1.821 0.06856 .
## RATERISKSame
                              ## RATERISKGreater
                              0.632806
                                       0.355571
                                                   1.780 0.07513 .
## PRIORFRACYes:RATERISKSame
                              ## PRIORFRACYes:RATERISKGreater 0.208811 0.624586
                                                  0.334 0.73814
## ---
```

```
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 562.34 on 499 degrees of freedom
## Residual deviance: 507.32 on 490 degrees of freedom
## AIC: 527.32
## Number of Fisher Scoring iterations: 4
lrtest(test)
## Likelihood ratio test
## Model 1: FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC + ARMASSIST + RATERISK +
      PRIORFRAC: RATERISK
## Model 2: FRACTURE ~ 1
## #Df LogLik Df Chisq Pr(>Chisq)
## 1 10 -253.66
## 2 1 -281.17 -9 55.015 1.21e-08 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
# (2) MOMFRAC: [ARMASSIST, RATERISK]
test <- model_effects</pre>
test <- update(test, . ~ . + MOMFRAC:ARMASSIST)
summary(test)
##
## Call:
## glm(formula = FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC +
      ARMASSIST + RATERISK + MOMFRAC: ARMASSIST, family = binomial,
##
      data = glow)
## Deviance Residuals:
       Min
                10
                      Median
                                   30
                                            Max
## -1.65273 -0.72683 -0.55140 0.03367
                                        2.27218
##
## Coefficients:
                        Estimate Std. Error z value Pr(>|z|)
                                   3.25148 0.912 0.36160
## (Intercept)
                         2.96640
## AGE
                         0.03760
                                   0.01323
                                           2.842 0.00448 **
## HEIGHT
                        ## PRIORFRACYes
                         0.61633
                                   0.24770
                                           2.488 0.01284 *
## MOMFRACYes
                         1.17111
                                   0.38940
                                           3.007 0.00263 **
## ARMASSISTYes
                         ## RATERISKSame
                         0.41386 0.28032 1.476 0.13985
                         0.71051
                                   0.29445 2.413 0.01582 *
## RATERISKGreater
                                 0.62405 -2.144 0.03201 *
## MOMFRACYes: ARMASSISTYes -1.33817
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
```

```
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 562.34 on 499 degrees of freedom
## Residual deviance: 502.83 on 491 degrees of freedom
## AIC: 520.83
##
## Number of Fisher Scoring iterations: 4
lrtest(test)
## Likelihood ratio test
##
## Model 1: FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC + ARMASSIST + RATERISK +
      MOMFRAC: ARMASSIST
## Model 2: FRACTURE ~ 1
## #Df LogLik Df Chisq Pr(>Chisq)
## 1 9 -251.41
      1 -281.17 -8 59.509 5.818e-10 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
test <- model effects
test <- update(test, . ~ . + MOMFRAC:RATERISK)</pre>
summary(test)
##
## Call:
## glm(formula = FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC +
      ARMASSIST + RATERISK + MOMFRAC: RATERISK, family = binomial,
##
      data = glow)
##
## Deviance Residuals:
                       Median
                 1Q
                                             Max
## -1.73530 -0.73156 -0.56262 -0.02886
                                         2.20217
##
## Coefficients:
                           Estimate Std. Error z value Pr(>|z|)
                            2.76974 3.23715
                                               0.856 0.39221
## (Intercept)
## AGE
                            0.03436 0.01308
                                               2.627 0.00861 **
## HEIGHT
                            ## PRIORFRACYes
                            0.83795
                                               0.032 0.97479
## MOMFRACYes
                            0.02648
                            0.44890
## ARMASSISTYes
                                      0.23340
                                                1.923 0.05444 .
## RATERISKSame
                            0.29742
                                       0.29700
                                                1.001 0.31663
                                                2.253 0.02428 *
## RATERISKGreater
                            0.70206
                                       0.31167
## MOMFRACYes:RATERISKSame
                             1.04615
                                       0.95957
                                                1.090 0.27561
## MOMFRACYes:RATERISKGreater 0.36775
                                       0.96207
                                                0.382 0.70227
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 562.34 on 499 degrees of freedom
```

```
## Residual deviance: 505.79 on 490 degrees of freedom
## AIC: 525.79
## Number of Fisher Scoring iterations: 4
lrtest(test)
## Likelihood ratio test
## Model 1: FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC + ARMASSIST + RATERISK +
      MOMFRAC: RATERISK
## Model 2: FRACTURE ~ 1
   #Df LogLik Df Chisq Pr(>Chisq)
## 1 10 -252.90
    1 -281.17 -9 56.542 6.183e-09 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
# (1) ARMASSIST: RATERISK
test <- model_effects</pre>
test <- update(test, . ~ . + ARMASSIST:RATERISK)</pre>
summary(test)
##
## Call:
## glm(formula = FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC +
      ARMASSIST + RATERISK + ARMASSIST: RATERISK, family = binomial,
##
##
      data = glow)
##
## Deviance Residuals:
          1Q Median
                               3Q
      Min
                                      Max
## -1.6586 -0.7419 -0.5544 -0.0470
                                    2.2531
##
## Coefficients:
                            Estimate Std. Error z value Pr(>|z|)
##
## (Intercept)
                            2.38363 3.22529 0.739 0.45988
                             0.03534
                                       0.01314 2.691 0.00713 **
## AGE
## HEIGHT
                            -0.04274
                                     0.01819 -2.349 0.01883 *
## PRIORFRACYes
                             ## MOMFRACYes
                             0.44193 1.375 0.16906
## ARMASSISTYes
                             0.60776
## RATERISKSame
                             ## RATERISKGreater
                             0.98400 0.38373
                                                2.564 0.01034 *
## ARMASSISTYes:RATERISKSame
                                     0.56723 0.190 0.84956
                             0.10760
## ARMASSISTYes:RATERISKGreater -0.60953
                                       0.58200 -1.047 0.29496
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 562.34 on 499 degrees of freedom
## Residual deviance: 505.42 on 490 degrees of freedom
```

```
## AIC: 525.42
##
## Number of Fisher Scoring iterations: 4
lrtest(test)
## Likelihood ratio test
##
## Model 1: FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC + ARMASSIST + RATERISK +
      ARMASSIST: RATERISK
## Model 2: FRACTURE ~ 1
   #Df LogLik Df Chisq Pr(>Chisq)
## 1 10 -252.71
## 2 1 -281.17 -9 56.912 5.253e-09 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Results from interactions
             0.001065 0.002095 0.508 0.6113
## AGE:HEIGHT
  AGE:PRIORFRACYes -0.05864 0.02583 -2.270 0.023188 *
                   -0.01353 0.03264 -0.414 0.67854
## AGE:MOMFRACYes
## AGE:ARMASSISTYes -0.01132 0.02479 -0.457 0.64802
## AGE:RATERISKSame -0.03999 0.03151 -1.269 0.20438
## AGE:RATERISKGreater -0.05021 0.03202 -1.568 0.11690
## HEIGHT:PRIORFRACYes 0.01915 0.03770 0.508 0.61146
## HEIGHT:MOMFRACYes 0.07401 0.04718 1.569 0.11675
## HEIGHT: ARMASSISTYes -0.04419 0.03594 -1.229 0.21890
## HEIGHT:RATERISKSame -0.01577
                                  0.04550 -0.347 0.72890
## HEIGHT:RATERISKGreater 0.02394 0.04528 0.529 0.59695
  PRIORFRACYes: MOMFRACYes -1.07823 0.65021 -1.658 0.09726 .
## PRIORFRACYes: ARMASSISTYes 0.24587 0.48467 0.507 0.61194
## PRIORFRACYes:RATERISKSame 0.001597
                                        0.625563
                                                0.003 0.99796
## PRIORFRACYes:RATERISKGreater 0.208811
                                        0.624586
                                                0.334 0.73814
  MOMFRACYes: ARMASSISTYes -1.33817 0.62405 -2.144 0.03201 *
## MOMFRACYes:RATERISKSame 1.04615 0.95957
                                              1.090 0.27561
## MOMFRACYes:RATERISKGreater 0.36775 0.96207 0.382 0.70227
## ARMASSISTYes:RATERISKSame 0.10760 0.56723 0.190 0.84956
## ARMASSISTYes:RATERISKGreater -0.60953 0.58200 -1.047 0.29496
```

Add to main effects model

we find three interactions, AGE:PRIORFRACYes, PRIORFRACYes:MOMFRACYes, MOMFRACYes:ARMASSISTYes

##

```
## Call:
## glm(formula = FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC +
      ARMASSIST + RATERISK + AGE:PRIORFRAC + PRIORFRAC:MOMFRAC +
      MOMFRAC: ARMASSIST, family = binomial, data = glow)
##
## Deviance Residuals:
      Min 10
                     Median
                                  30
                                          Max
## -1.47664 -0.74929 -0.51571 0.07753
                                      2.33224
##
## Coefficients:
                       Estimate Std. Error z value Pr(>|z|)
                        1.20626 3.38765 0.356 0.721785
## (Intercept)
## AGE
                        0.05949 0.01677
                                          3.547 0.000389 ***
## HEIGHT
                       -0.04610 0.01854 -2.487 0.012886 *
## PRIORFRACYes
                        4.63031 1.88158 2.461 0.013860 *
                        ## MOMFRACYes
                        ## ARMASSISTYes
## RATERISKSame
                        0.42125 0.28217 1.493 0.135462
                        ## RATERISKGreater
                       ## AGE:PRIORFRACYes
## PRIORFRACYes:MOMFRACYes -0.83184 0.64852 -1.283 0.199606
## MOMFRACYes:ARMASSISTYes -1.15254 0.61838 -1.864 0.062350 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 562.34 on 499 degrees of freedom
## Residual deviance: 496.53 on 489 degrees of freedom
## AIC: 518.53
## Number of Fisher Scoring iterations: 4
\# create final model with interactions terms AGE:PRIORFRAC + MOMFRAC:ARMASSIST
model_effects_final <- glm(FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC + ARMASSIST + RATERISK +
                    AGE:PRIORFRAC + MOMFRAC:ARMASSIST, family = binomial, data = glow)
summary(model effects final)
##
## Call:
## glm(formula = FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC +
      ARMASSIST + RATERISK + AGE:PRIORFRAC + MOMFRAC:ARMASSIST,
##
      family = binomial, data = glow)
##
## Deviance Residuals:
      Min
             1Q
                  Median
                              3Q
                                     Max
## -1.6995 -0.7459 -0.5238 0.0620
##
## Coefficients:
                       Estimate Std. Error z value Pr(>|z|)
##
                        0.96955 3.38252 0.287 0.774392
## (Intercept)
## AGE
                        0.05890
                                 0.01666 3.535 0.000408 ***
## HEIGHT
                       4.65073 1.88342 2.469 0.013538 *
## PRIORFRACYes
```

```
## MOMFRACYes
                          1.19902
                                     0.39487
                                              3.036 0.002393 **
## ARMASSISTYes
                          0.61423
                                    0.25358
                                              2.422 0.015426 *
## RATERISKSame
                          0.42626 0.28154
                                              1.514 0.130012
                          0.72116
## RATERISKGreater
                                    0.29660
                                              2.431 0.015040 *
                         -0.05610
## AGE:PRIORFRACYes
                                     0.02600 -2.158 0.030950 *
## MOMFRACYes:ARMASSISTYes -1.26534
                                     0.62377 -2.029 0.042507 *
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 562.34 on 499 degrees of freedom
## Residual deviance: 498.17 on 490 degrees of freedom
## AIC: 518.17
##
## Number of Fisher Scoring iterations: 4
```

Final Interaction Model

```
(Intercept)
AGE
                     0.05890
                               0.01666 3.535 0.000408 ***
HEIGHT
                    -0.04413
                               0.01848 -2.388 0.016949 *
PRIORFRACYes
                     4.65073
                              1.88342 2.469 0.013538 *
MOMFRACYes
                     1.19902
                              0.39487
                                       3.036 0.002393 **
ARMASSISTYes
                     RATERISKGreater
                     0.72116
                              0.29660
                                       2.431 0.015040 *
AGE:PRIORFRACYes
                    -0.05610
                               0.02600 -2.158 0.030950 *
MOMFRACYes: ARMASSISTYes -1.26534
                               0.62377 -2.029 0.042507 *
```

FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC + ARMASSIST + RATERISK + AGE:PRIORFRAC + MOMFRAC:ARMASSIS

library(pROC)

```
## Type 'citation("pROC")' for a citation.

##
## Attaching package: 'pROC'

## The following object is masked from 'package:gmodels':
##
## ci

## The following object is masked from 'package:glmnet':
##
## auc

## The following objects are masked from 'package:stats':
##
## cov, smooth, var
```

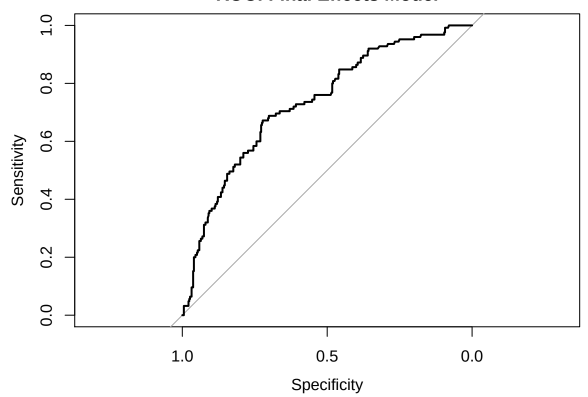
```
library(vcdExtra)
## Loading required package: gnm
##
## Attaching package: 'gnm'
## The following object is masked from 'package:modeltools':
##
##
       parameters
## The following object is masked from 'package:lattice':
##
##
       barley
##
## Attaching package: 'vcdExtra'
## The following object is masked from 'package:carData':
##
##
       Burt
## The following object is masked from 'package:plyr':
##
##
       summarise
## The following object is masked from 'package:dplyr':
##
##
       summarise
# vcov(model_effects_final)
HLtest(model_effects_final)
## Hosmer and Lemeshow Goodness-of-Fit Test
##
## Call:
## glm(formula = FRACTURE ~ AGE + HEIGHT + PRIORFRAC + MOMFRAC +
##
       ARMASSIST + RATERISK + AGE:PRIORFRAC + MOMFRAC:ARMASSIST,
##
       family = binomial, data = glow)
## ChiSquare df
                  P_value
    7.268011 8 0.5080118
##
glow$predict_mfinal <- predict(model_effects_final, type = "response")</pre>
with(glow, addmargins(table(glow$predict_mfinal > 0.5, glow$FRACTURE)))
##
##
            No Yes Sum
##
     FALSE 354 97 451
##
     TRUE
           21 28 49
           375 125 500
##
     Sum
```

```
(roc_final_model <- roc(glow$FRACTURE ~ glow$predict_mfinal, data = glow))

##
## Call:
## roc.formula(formula = glow$FRACTURE ~ glow$predict_mfinal, data = glow)
##
## Data: glow$predict_mfinal in 375 controls (glow$FRACTURE No) < 125 cases (glow$FRACTURE Yes).
## Area under the curve: 0.7331

plot(roc_final_model, main = "ROC: Final Effects Model")</pre>
```

ROC: Final Effects Model



 misc

##

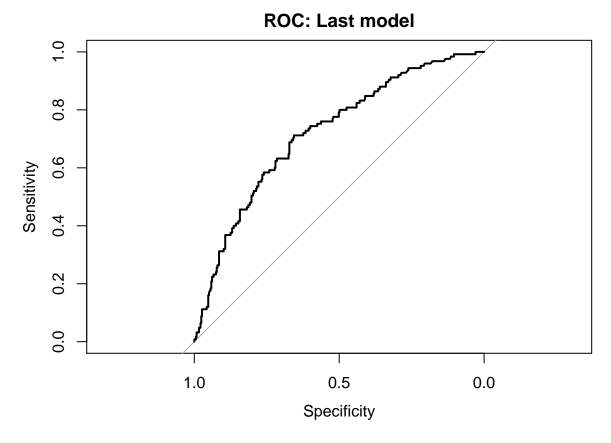
3.10152 8 0.9278259

```
## restart with clean data
glow <- read_glow_dataset()

model_last <- glm(FRACTURE ~ AGE:PRIORFRAC + HEIGHT + MOMFRAC:ARMASSIST + I(as.integer(RATERISK) == 3),
HLtest(model_last)

## Hosmer and Lemeshow Goodness-of-Fit Test
##
## Call:
## glm(formula = FRACTURE ~ AGE:PRIORFRAC + HEIGHT + MOMFRAC:ARMASSIST +
## I(as.integer(RATERISK) == 3), family = binomial, data = glow)
## ChiSquare df P_value</pre>
```

```
summary(HLtest(model_last))
## Partition for Hosmer and Lemeshow Goodness-of-Fit Test
##
##
                  cut total obs
                                                chi
                                     exp
## 1 [0.0243,0.0967]
                        50 47 45.97075 0.1518032
## 2
       (0.0967, 0.123]
                         50 46 44.35914 0.2463653
## 3
       (0.123,0.152] 50 42 43.19969 -0.1825284
## 4
        (0.152, 0.18]
                         50 41 41.81265 -0.1256745
                         50 42 40.34124 0.2611609
## 5
        (0.18, 0.213]
        (0.213,0.251]
## 6
                         50 36 38.55936 -0.4121599
## 7
        (0.251, 0.292]
                         50 38 36.55362 0.2392312
                         50 32 33.67421 -0.2885110
## 8
        (0.292, 0.372]
## 9
         (0.372, 0.47]
                         50 28 29.25398 -0.2318447
## 10
         (0.47, 0.724]
                         50 23 21.27536 0.3739034
## Hosmer and Lemeshow Goodness-of-Fit Test
## Call:
## glm(formula = FRACTURE ~ AGE:PRIORFRAC + HEIGHT + MOMFRAC:ARMASSIST +
##
       I(as.integer(RATERISK) == 3), family = binomial, data = glow)
##
   ChiSquare df
                 P_{value}
##
      3.10152 8 0.9278259
# classification table
glow$predict_last <- predict(model_last, type = "response")</pre>
with(glow, addmargins(table(predict_last > 0.5, FRACTURE)))
          FRACTURE
##
##
            No Yes Sum
##
     FALSE 355 103 458
##
     TRUE
          20 22 42
##
     Sum
           375 125 500
# Sensitivy, specificity, ROC (using pROC)
roc_model_last <- roc(glow$FRACTURE ~ glow$predict_last, data = glow)</pre>
plot(roc_model_last, main = "ROC: Last model")
```



```
# create table
vars <- c("thresholds", "sensitivities", "specificities")
model_table <- data.frame(roc_model_last[vars])

findIndex <- function(x, y) which.min( (x-y)^2 )
cutPoints <- seq(0.05, 0.75, by = 0.05)

tableIndex <- mapply(findIndex, y = cutPoints, MoreArgs = list(x = roc_model_last$thresholds))
model_table[tableIndex, ]</pre>
```

```
##
       thresholds sensitivities specificities
## 3
       0.05165803
                           1.000
                                   0.005333333
## 43 0.09905744
                           0.976
                                   0.128000000
## 120 0.15054070
                           0.880
                                   0.349333333
## 202 0.20014367
                           0.760
                                   0.549333333
## 259 0.25035362
                           0.640
                                   0.674666667
## 316 0.29918726
                           0.520
                                   0.789333333
## 349 0.34952747
                           0.416
                                   0.842666667
## 379 0.40012793
                           0.320
                                   0.893333333
## 401 0.44487716
                           0.240
                                   0.925333333
## 416 0.49400138
                           0.176
                                   0.944000000
## 426 0.55045683
                           0.120
                                   0.954666667
                           0.056
                                   0.978666667
## 443 0.59996390
## 452 0.65801860
                           0.024
                                   0.992000000
## 455 0.69417409
                           0.008
                                   0.994666667
```

```
# plot
plot(specificities ~ thresholds, xlim = c(0, 1), type = "l",
xlab = "probability cutoff", ylab = "sensitivity / specificity",
ylim = c(0, 1), data = model_table, main = "probability sensitivity")
with(model_table, lines(thresholds, sensitivities, col = "red"))
legend(x = 0.75, y = 0.55, legend = c("Sensitivity", "Specificity"),
lty = 1, col = c("red", "black"))
abline(h = c(0, 1), col = "grey80", lty = "dotted")
```

probability sensitivity

