CancerDataR

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Tasks:

Summary of data, Introduction - Truman

Organizing presentation - everyone

Presenting - Truman, Minhaz, Vincent

Anaylizing data/Coding - Truman, Minhaz, Vincent

Putting everything together/Conclusions/Report - Navin, Bobak

Introduction

Breast cancer is a malignant cell growth in the breast. If it is left untreated the cancer can spread to other parts of the human body and it can be very deadly. There are generally two type of tumors non-cancerous and cancerous and the difference between the two is important, Benign tumor is non-cancerous and not dangerous on its own, but a malignant tumor, means the mass is cancerous.

Our goal for this project is to predict whether the cancer is benign or malignant and to determine what actually contribute to the classification of the two types.

We are given the following: Attribute Information: 1) ID number 2) Diagnosis (M = malignant, B = benign) 3-32)

- a) radius (mean of distances from center to points on the perimeter)
- b) texture (standard deviation of gray-scale values)
- c) perimeter
- d) area
- e) smoothness (local variation in radius lengths) f)compactness (perimeter^2 / area 1.0)
- f) concavity (severity of concave portions of the contour)
- g) concave points (number of concave portions of the contour)
- h) symmetry
- i) fractal dimension("coastline approximation" 1)

The mean, standard error and "worst" or largest (mean of the three largest values) of these features were computed for each image, resulting in 30 features Class distribution: 357 benign, 212 malignant

Summary of the data

We first take a small look at the data set and loading library/files we will need.

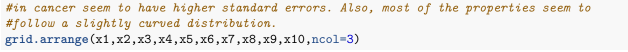
```
# all the library and files we'll be using
library(tidyverse)
library(gridExtra)
library(ICSNP)
library(MASS)
```

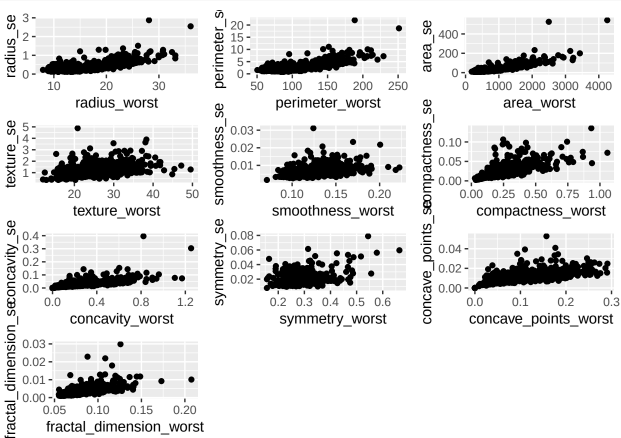
```
library(klaR)
source("Box_M.R")
# preview of the data
cancer = read.csv("Project3-Data.csv")
head(cancer[1:5])
##
           id diagnosis radius_mean texture_mean perimeter_mean
## 1
       842302
                      М
                               17.99
                                            10.38
                                                           122.80
## 2
                                            17.77
       842517
                               20.57
                                                           132.90
                      Μ
## 3 84300903
                      М
                               19.69
                                            21.25
                                                           130.00
## 4 84348301
                      М
                               11.42
                                            20.38
                                                           77.58
## 5 84358402
                      Μ
                               20.29
                                            14.34
                                                           135.10
## 6
      843786
                      М
                               12.45
                                            15.70
                                                            82.57
# number of variables we have
num_var = ncol(cancer) - 1
num var
## [1] 31
# number of observation we have
num obs = nrow(cancer)
num obs
## [1] 569
# the number of each type of tumor
table(cancer$diagnosis)
##
##
     В
## 357 212
```

Some visuals of the data

Standard errors vs worst cases

```
# filter out the first 2 columns
filcancer1=cancer[-1:-2]
x1<-ggplot(filcancer1, aes(x=radius worst, y = radius se))+geom point()</pre>
x2<-ggplot(filcancer1, aes(x=perimeter_worst, y = perimeter_se))+geom_point()</pre>
x3<-ggplot(filcancer1, aes(x=area_worst, y = area_se))+geom_point()</pre>
#If we look at the first plot of texture's worst against standard errors, we can see
#non constant variance due to the cone shape of the data.
x4<-ggplot(filcancer1, aes(x=texture worst, y = texture se))+geom point()
#Again we have a cone shape in the plot of smoothness worst versus standard error.
x5<-ggplot(filcancer1, aes(x=smoothness_worst, y = smoothness_se))+geom_point()
x6<-ggplot(filcancer1, aes(x=compactness_worst, y = compactness_se))+geom_point()
x7<-ggplot(filcancer1, aes(x=concavity_worst, y = concavity_se))+geom_point()</pre>
x8<-ggplot(filcancer1, aes(x=symmetry_worst, y = symmetry_se))+geom_point()
x9<-ggplot(filcancer1, aes(x=concave_points_worst, y = concave_points_se))+geom_point()</pre>
x10<-ggplot(filcancer1, aes(x=fractal_dimension_worst, y = fractal_dimension_se))+geom_point()
#Looking at the data, most points are near the the origin. However, the further stages
```

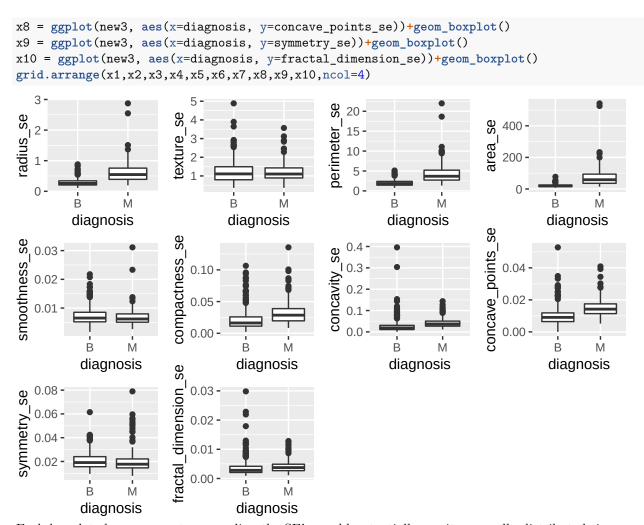




If we look at all the graphs for each category for measuring breast cancer, we can see that most data points cluster quite near to the origin, but not exactly on it. There are no data points at the origin since this would mean that the person has no reason to suspect that they have breast cancer. If we now look at the data points that are close to the center but not exactly, these points might represent either benign tumours or early stage malignant tumours, since these breasts don't really deviate that much from the average. If we look at the points further out, these points are likely middle stage malignant tumours, since they deviate a sizeable amount. Now, looking at the outliers, we have reason to suspect that these are late stage malignant tumours too, given the deformity of the breasts.

Boxplot of the SEs

```
new = cancer[-1]
new2 = new[,-c(2:11)]
new3 = new2[,-c(12:21)]
x1 = ggplot(new3, aes(x=diagnosis, y=radius_se))+geom_boxplot()
x2 = ggplot(new3, aes(x=diagnosis, y=texture_se))+geom_boxplot()
x3 = ggplot(new3, aes(x=diagnosis, y=perimeter_se))+geom_boxplot()
x4 = ggplot(new3, aes(x=diagnosis, y=area_se))+geom_boxplot()
x5 = ggplot(new3, aes(x=diagnosis, y=smoothness_se))+geom_boxplot()
x6 = ggplot(new3, aes(x=diagnosis, y=compactness_se))+geom_boxplot()
x7 = ggplot(new3, aes(x=diagnosis, y=concavity_se))+geom_boxplot()
```



Each boxplot shows symmetry regarding the SE's, could potentially say its normally distributed since we have large sample size to back up the outliers.

Methods

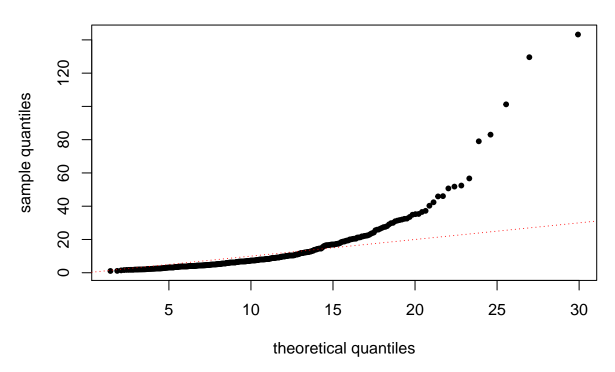
Chi Square plot

before we start any analysis we want to verify the normallity of the data.

```
source("ChisqPlot.R")
can = cancer[,2:12]

# setting the independent variables into a matrix
can.matrix = as.matrix(can[,2:11])
chisqplot(can.matrix)
```





Because our data have a very large sample, I would say the normality assumption here is fine.

Fitting a generalized linear model

we first starts of by fitting a generalized linear model to assess the significance of each of the variables. here we are only dealing with the "mean" variables as we believe that the other two category "standard error" and "worst" will not give us much information regarding the type of cancer.

```
# taking the mean values
can = cancer[2:12]
# changing diagnosis from chr to factor so is easy for model fitting
can$diagnosis = as.factor(can$diagnosis)
# fitting the generalized linear model
glm.fit = glm(diagnosis ~ ., data=can, family=binomial)
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred
summary(glm.fit)
##
  glm(formula = diagnosis ~ ., family = binomial, data = can)
##
## Deviance Residuals:
                         Median
       Min
                   1Q
                                       30
                                                Max
## -1.95590 -0.14839 -0.03943
                                  0.00429
                                             2.91690
##
## Coefficients:
```

```
##
                            Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                                                 -0.573
                            -7.35952
                                       12.85259
                                                           0.5669
## radius mean
                            -2.04930
                                        3.71588
                                                  -0.551
                                                           0.5813
                                                   5.961
                                                          2.5e-09 ***
## texture_mean
                             0.38473
                                        0.06454
## perimeter_mean
                            -0.07151
                                        0.50516
                                                  -0.142
                                                           0.8874
## area mean
                             0.03980
                                        0.01674
                                                   2.377
                                                           0.0174 *
## smoothness mean
                            76.43227
                                       31.95492
                                                   2.392
                                                           0.0168 *
## compactness_mean
                            -1.46242
                                       20.34249
                                                  -0.072
                                                           0.9427
## concavity_mean
                             8.46870
                                        8.12003
                                                   1.043
                                                           0.2970
## concave_points_mean
                            66.82176
                                       28.52910
                                                   2.342
                                                           0.0192 *
## symmetry_mean
                            16.27824
                                       10.63059
                                                   1.531
                                                           0.1257
## fractal_dimension_mean -68.33703
                                       85.55666
                                                 -0.799
                                                           0.4244
##
## Signif. codes:
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 751.44
                               on 568
                                       degrees of freedom
## Residual deviance: 146.13
                              on 558
                                       degrees of freedom
  AIC: 168.13
##
## Number of Fisher Scoring iterations: 9
```

Theres various information in the summary but look at the coefficients, we have estimate, SE, z-score, and p-value, the p-value that is less than 0.05 indicates significance, that is those variable has an impact on either cancer being M or B. Example: for a unit increase in texture mean the odd of cancer being M (vs B) increases by exp(0.38473).

Now that we know which of the variables are actually significant, we will fit the model again with only those significant variables.

```
# new model after removing insignificant variables
glm.fit2 = glm(diagnosis ~ ., data=can[,c(1,3,5,6,9)], family=binomial)
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred
summary(glm.fit2)
##
## Call:
  glm(formula = diagnosis ~ ., family = binomial, data = can[,
##
       c(1, 3, 5, 6, 9)])
##
##
## Deviance Residuals:
                   1Q
##
        Min
                         Median
                                        3Q
                                                 Max
## -2.31798 -0.15623 -0.04212
                                   0.01662
                                             2.84201
##
## Coefficients:
                         Estimate Std. Error z value Pr(>|z|)
##
## (Intercept)
                        -23.677816
                                     3.882774
                                               -6.098 1.07e-09 ***
## texture_mean
                         0.362687
                                     0.060544
                                                5.990 2.09e-09 ***
## area_mean
                         0.010342
                                     0.002002
                                                5.165 2.40e-07 ***
## smoothness_mean
                         59.471304
                                    25.965153
                                                2.290
                                                          0.022 *
## concave_points_mean 76.571210
                                    16.427864
                                                4.661 3.15e-06 ***
##
```

Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1

```
##
## (Dispersion parameter for binomial family taken to be 1)
##
## Null deviance: 751.44 on 568 degrees of freedom
## Residual deviance: 156.44 on 564 degrees of freedom
## AIC: 166.44
##
## Number of Fisher Scoring iterations: 8
```

Taking a look at the different AIC values of the two model, we see that the AIC values for the reduced model are smaller than the full model, this tells us that not only are some of the variable are insignificant but it also will effect the accuracy of our result.

With the following we can get a rough probability of type of cancer with given values.

```
# a function to get the probability of cancer being type M
prob = function(x1,x2,x3,x4){
  x = exp(-23.677816 + 0.362687*x1 + 0.010342*x2 + 59.471304*x3 + 76.571210*x4)
  pix = x/(1+x)
  return(pix)
}
```

Discriminant Analysis

data: cancer1 and cancer2

Now we shall take a look at another method, here we use discriminant analysis, Discriminant analysis is a technique that is used to analyze the research data when the criterion or the dependent variable is categorical and the predictor or the independent variable is interval in nature (which is what we have here).

First we have to compute a two-sample Hotelling T-Squared test and compute Bartlett's test for homogeneous covariance matrices. with this we can determine whether or not to use Linear DA or quadratic DA as one requires equal covariance and the other one does not (LDA require equal covariance).

```
# again we are only working with the means
can = cancer[,2:12]
# setting the independent variables into a matrix
can.matrix = as.matrix(can[,2:11])
fit=manova(can.matrix ~ can$diagnosis)
summary(fit, test="Hotelling-Lawley")
                  Df Hotelling-Lawley approx F num Df den Df
                                                                 Pr(>F)
## can$diagnosis
                               2.1522
                                        120.09
                                                    10
                                                          558 < 2.2e-16 ***
## Residuals
                 567
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
# create separate data sets for Benign and Malignant tumors.
cancer1 <- can[can[,1]=="M",2:11]</pre>
cancer2 <- can[can[,1] == "B",2:11]
HotellingsT2(cancer1, cancer2)
##
##
   Hotelling's two sample T2-test
##
```

```
## T.2 = 120.09, df1 = 10, df2 = 558, p-value < 2.2e-16
## alternative hypothesis: true location difference is not equal to c(0,0,0,0,0,0,0,0,0)
n1 = dim(cancer1)[1]
n2 = dim(cancer2)[1]

Box_M(can.matrix, n=c(n1, n2))

## Test result:
## [,1]
## Box.M-C 221.7243
## p.value 0.0000</pre>
```

Here we see that we do not have equal covariance and so we'll be using QDA instead of LDA for better performance/accuracy.

Discriminant analysis with all 10 variables

We first start with all 10 variables just so we can have a comparison later with the reduced model.

```
# spliting the data into 2 set, training and testing
training_sample <- sample(c(TRUE, FALSE), nrow(can), replace = T, prob = c(0.6,0.4))
cantrain <- can[training_sample, ]</pre>
cantest <- can[!training_sample, ]</pre>
# the model
cancer.qda <- qda(diagnosis ~ ., data=cantrain)</pre>
cancer.qda
## Call:
## qda(diagnosis ~ ., data = cantrain)
## Prior probabilities of groups:
##
## 0.6454294 0.3545706
##
## Group means:
     radius_mean texture_mean perimeter_mean area_mean smoothness_mean
## B
        12.14709
                     17.88893
                                     78.08618 462.6901
                                                              0.09209579
        17.60867
                     21.44117
                                    116.44977 995.2336
                                                              0.10305633
##
     compactness_mean concavity_mean concave_points_mean symmetry_mean
## B
           0.07951146
                           0.04716042
                                                0.02608572
                                                                0.1722215
           0.14691539
                           0.16412391
                                                0.08904148
                                                               0.1926086
## M
     fractal_dimension_mean
## B
                 0.06275897
## M
                 0.06276938
#Confusion test
set.seed(1)
confusionTest <- table(cantest$diagnosis, predict(cancer.qda, newdata=cantest)$class)</pre>
confusionTest
##
##
         В
             Μ
##
     B 119
             5
##
    M 15
```

```
n <- sum(confusionTest)
aer <- (n - sum(diag(confusionTest))) / n
aer
## [1] 0.09615385</pre>
```

Discriminant analysis with the significant variables

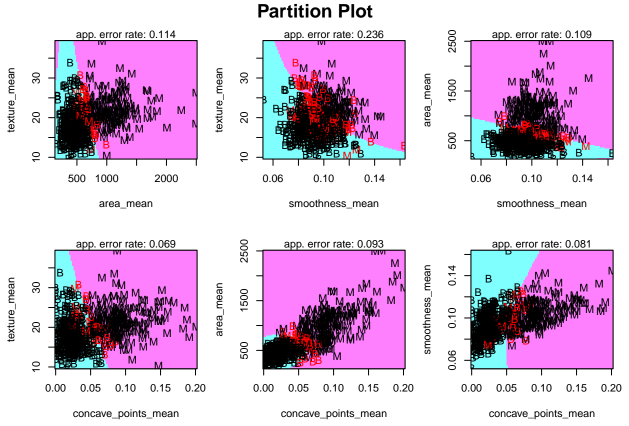
We now do the same thing but with the reduced model.

```
# splitting data into 2sets, training and testing
can2 = can[,c(1,3,5,6,9)]
training_sample2 <- sample(c(TRUE, FALSE), nrow(can2), replace = T, prob = c(0.6,0.4))
cantrain2 <- can2[training_sample2, ]</pre>
cantest2 <- can2[!training_sample2, ]</pre>
# the model
cancer.qda2 <- qda(diagnosis ~ ., data=cantrain2, CV=FALSE)</pre>
cancer.qda2
## Call:
## qda(diagnosis ~ ., data = cantrain2, CV = FALSE)
##
## Prior probabilities of groups:
           В
## 0.6340058 0.3659942
##
## Group means:
     texture_mean area_mean smoothness_mean concave_points_mean
## B
         17.98995 457.4986
                                   0.09226236
                                                        0.02516845
         21.73795 983.6764
                                   0.10292252
                                                        0.08891394
# testing the accuracy of our model
set.seed(1)
qda.test <- predict(cancer.qda2,cantest2)</pre>
cantest2$qda <- qda.test$class</pre>
confusionTest <-table(cantest2$qda,cantest2$diagnosis)</pre>
confusionTest
##
##
         В
##
     B 130 10
        7 75
n <- sum(confusionTest)</pre>
aer <- (n - sum(diag(confusionTest))) / n</pre>
aer
```

[1] 0.07657658

Taking a look at the errors of the two model, full vs reduced we see that the difference between the 2 is negligible (the AER for the two are very close). This also however tells us that with only 4 variables, texture mean, area mean, smoothness mean and concave points mean we can accurately predict about 93% of the class of observation which is very good.

```
# here are just some more visuals of the data
partimat(diagnosis ~ ., data=can2, method="qda")
```



From this we can also confirm that the accuracy of our model is indeed good.

Conclusion

From the two analysis we done, from discriminant analysis to simply fitting a generalized linear model we can clearly conclude that the dependent variable cancer type or diagnosis hugely depend on simply four variables, that is, it's mainly depend on texture mean, area mean, smoothness mean and concave points mean and from these four variables we can determine the odds patient's cancer type and so from that can determine whether treatment are necessary.

Reference

Fayed, L., & Paul, D. (n.d.). Differences Between a Malignant and Benign Tumor. Retrieved from https://www.verywellhealth.com/what-does-malignant-and-benign-mean-514240

Sign In. (n.d.). Retrieved from https://rpubs.com/Nolan/298913