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Research questions

Which attributes best predict breast cancer malignancy?

Is it possible to create a machine learning model that can predict breast cancer malignancy with an accuracy of 80% or higher?

Load libraries and read data

```
library(tibble)
## Warning: package 'tibble' was built under R version 4.0.5
library(dplyr)
## Warning: package 'dplyr' was built under R version 4.0.5
##
## Attaching package: 'dplyr'
## The following objects are masked from 'package:stats':
##
##
       filter, lag
## The following objects are masked from 'package:base':
##
##
       intersect, setdiff, setequal, union
library(ggplot2)
## Warning: package 'ggplot2' was built under R version 4.0.5
library(ggrepel)
## Warning: package 'ggrepel' was built under R version 4.0.5
library(reshape2)
## Warning: package 'reshape2' was built under R version 4.0.5
library(scales)
```

This breast cancer database was obtained from the University of Wisconsin Hospitals, Madison from

Warning: package 'scales' was built under R version 4.0.5

Dr. William H. Wolberg.

The data provided does not heave clear column names, but a description of each attribute has been given in the *breast-cancer-wisconsin.names* file. This allows us to assign representative names to every column. The code to do this is shown below.

Basic look at the data

```
# Look at head of data
data
## # A tibble: 698 x 11
##
           id Clump_Thickness Cell_Size_Uniform~ Cell_Shape_Unifo~ Marginal_Adhesi~
##
        <int>
                        <int>
                                            <int>
                                                              <int>
                                                                                <int>
##
  1 1002945
                            5
                                                4
                                                                  4
                                                                                    5
## 2 1015425
                            3
                                                1
                                                                                    1
                                                                  1
## 3 1016277
                            6
                                                8
                                                                  8
                                                                                    1
## 4 1017023
                            4
                                                                                    3
                                                1
                                                                  1
## 5 1017122
                            8
                                               10
                                                                 10
                                                                                    8
## 6 1018099
                            1
                                                1
                                                                  1
                                                                                    1
## 7 1018561
                            2
                                                1
                                                                                    1
                            2
## 8 1033078
                                                1
                                                                  1
                                                                                    1
## 9 1033078
                                                                  1
                                                                                    1
## 10 1035283
                            1
                                                1
                                                                   1
                                                                                    1
## # ... with 688 more rows, and 6 more variables:
       Single_Epithelial_Cell_Size <int>, Bare_Nuclei <int>,
       Bland_Chromatin <int>, Normal_Nucleoli <int>, Mitoses <int>, Class <int>
```

```
# Summarize data
summary(data)
```

```
##
                     Clump_Thickness Cell_Size_Uniformity Cell_Shape_Uniformity
         id
                     Min. : 1.000
  Min.
         :
             61634
                                     Min. : 1.000
                                                         Min. : 1.000
  1st Qu.: 870258
                     1st Qu.: 2.000
                                     1st Qu.: 1.000
                                                         1st Qu.: 1.000
                     Median : 4.000
## Median : 1171710
                                     Median : 1.000
                                                         Median : 1.000
## Mean : 1071807
                     Mean : 4.417
                                     Mean : 3.138
                                                         Mean : 3.211
   3rd Qu.: 1238354
                     3rd Qu.: 6.000
                                     3rd Qu.: 5.000
                                                         3rd Qu.: 5.000
## Max. :13454352
                     Max. :10.000
                                     Max. :10.000
                                                         Max. :10.000
##
## Marginal_Adhesion Single_Epithelial_Cell_Size Bare_Nuclei
## Min. : 1.000
                    Min. : 1.000
                                              Min. : 1.000
## 1st Qu.: 1.000
                    1st Qu.: 2.000
                                              1st Qu.: 1.000
## Median : 1.000
                    Median : 2.000
                                              Median : 1.000
## Mean : 2.809
                    Mean : 3.218
                                              Mean : 3.548
## 3rd Qu.: 4.000
                    3rd Qu.: 4.000
                                              3rd Qu.: 6.000
## Max.
         :10.000
                    Max. :10.000
                                              Max.
                                                     :10.000
##
                                              NA's
                                                    :16
## Bland Chromatin Normal Nucleoli
                                     Mitoses
## Min. : 1.000
                   Min. : 1.00 Min. : 1.00 Min.
```

```
1st Qu.: 2.000
                     1st Qu.: 1.00
                                      1st Qu.: 1.00
                                                      1st Qu.:2.000
                                                      Median :2.000
   Median : 3.000
                     Median: 1.00
##
                                     Median : 1.00
  Mean
##
          : 3.438
                     Mean
                            : 2.87
                                     Mean
                                            : 1.59
                                                      Mean
                                                             :2.691
   3rd Qu.: 5.000
                     3rd Qu.: 4.00
                                      3rd Qu.: 1.00
                                                      3rd Qu.:4.000
##
##
   Max.
           :10.000
                     Max.
                            :10.00
                                     Max.
                                             :10.00
                                                      Max.
                                                             :4.000
##
```

All data has been normalized to fit a grading systems that grades the severity of each attribute. A detailed description of what each grade value means for each attribute can be found in the Breast Cancer Diagnosis Web User Interface.

Correlation

To get a sense of which attributes are best suited to predict the class, it is helpful to take a look at the correlation. Below is a piece of code that prints the column names of the columns with a correlation of 0.8 or higher to the Class attribute.

```
# Get correlation
data.cor <- cor(data[,-1], use = "complete.obs")
data.cor <- as.data.frame(data.cor)
# Keep all correlations that are higher than 0.8
cor.names <- names(data.cor)[which(data.cor$Class > 0.8)]
# Print
cat(cor.names)
```

Cell_Size_Uniformity Cell_Shape_Uniformity Bare_Nuclei Class

This gives us just three attributes of interest besides the Class attribute which obviously fully correlates to itself. This is good, because this means the model we will train later will likely only need these three attributes to accurately predict the Malignancy of a cancer clump. A simple model is always better, because less data needs to be acquired in order to use it. However, to make sure we do not miss any value information, it is useful to make a heat map of the correlation table. The heat map and corresponding code is shown below.

```
# Make correlation matrix
cormat <- cor(na.omit(data[, -1]))</pre>
# Melt to correct format
melted.cormat <- melt(cormat)</pre>
# head(melted.cormat) # <- uncomment this to look at the new matrix
# Make heatmap with ggplot
ggplot(data = melted.cormat, aes(x=Var1, y=Var2, fill=value)) +
  geom_tile() + # geom_tile makes the heatmap tiles
  scale_fill_gradient(low = rgb(0, 0, 0),
                      high = rgb(0, 1, 1),
                      guide = "colorbar") + # sets color the gradient for tiles
  ylab('') +
  xlab('') + # the x and y axes have no useful names, and are thus left blank
  labs(caption =
      "Figure 1 ~ Correlation Heatmap of breast cancer data set.
      The lighter the color the better, the correlation between the two columns.") + # adds caption
  theme_minimal() + # set theme to minimal
```

theme(axis.text.x = element_text(angle = 45, hjust = 1), # adjust x labels to be diagonal plot.caption = element_text(hjust = 0.7)) # adjust caption position

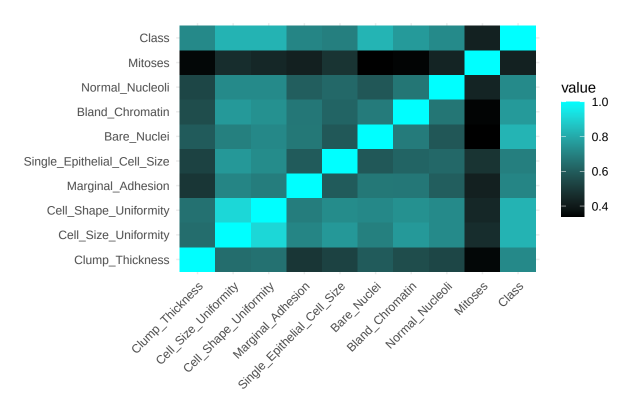


Figure 1 ~ Correlation Heatmap of breast cancer data set. The lighter the color the better, the correlation between the two columns.

Looking at 'Class' column of the heat map in figure 1, the Bland_Chromatin correlation is the lightest color visible besides the three attributes found earlier (Cell_Size_Uniformity, Cell_Shape_Uniformity Bare_Nuclei). The following code takes a look at the exact correlation of this attribute.

```
# Print correlation of Bland_Chromatin to Class
cormat["Bland_Chromatin", "Class"]
```

[1] 0.7583497

The correlation is roughly 0.76, which is close to the 0.8 threshold set earlier. Bland_Chromatin might be another attribute of interest.

Another thing that stands out in figure 1 is that Cell_Shape_Uniformity and Cell_Size_Uniformity show a high correlation as depicted by the light 4x4 square at the bottom left of the figure. This will be discussed further later on.

Class Distribution

Change Class to be factor

The code shown below substitutes the previously numeric data in the Class column of the data set to a factor with two labels: B for benign and M for malignant. This makes following visualizations easier and is also needed when eventually classifying the data with our model.

```
# Change class values to B for benign and M for Malignant instead of 2 and 4
data$Class[data$Class == 2] <- 'B'</pre>
data$Class[data$Class == 4] <- 'M'</pre>
data["Class"]
## # A tibble: 698 x 1
      Class
##
##
      <chr>>
##
   1 B
## 2 B
## 3 B
## 4 B
## 5 M
## 6 B
## 7 B
## 8 B
## 9 B
## 10 B
## # ... with 688 more rows
```

Pie Chart

To visualize the distribution of instances over the two classes a pie chart is made.

```
# Get class counts
m.count <- length(data$Class[data$Class == 'M'])</pre>
b.count <- length(data$Class[data$Class == 'B'])</pre>
# Combine in data frame
count.data <- data.frame(Class = c('Malignant', 'Benign'), Value = c(m.count, b.count))</pre>
count.data %>%
arrange(desc(Value)) %>%
mutate(prop = percent(Value / sum(Value), accuracy = 0.1)) -> count.data
# Make pie chart
ggplot(count.data, aes(x = "", y = Value, fill = Class)) +
  geom_bar(stat="identity", width=1, color="white") +
  coord_polar("y", start=0) +
  geom_label_repel(aes(label = prop), size=5, show.legend = F, nudge_x = -1, segment.colour= NA) +
 labs(caption="Figure 2 ~ A pie chart of the distrubtion of the classes in the dataset.") +
  theme_void() +
  theme(plot.caption = element_text(hjust = 1))
```

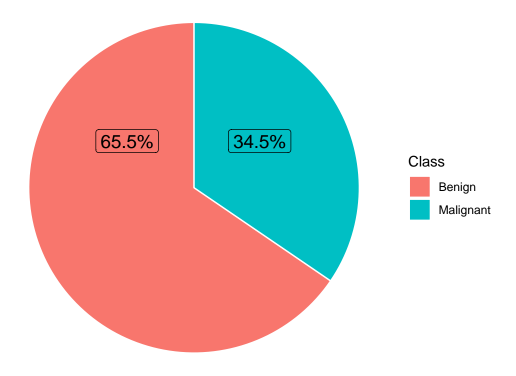


Figure 2 ~ A pie chart of the distrubtion of the classes in the dataset.

```
# Calculate exact percentage
m.count / (m.count + b.count) * 100
```

[1] 34.52722

Figure 2 shows that roughly two thirds of the data consists of Benign instances. When calculated, we find that only 34.5% of the instances are malignant. This should be kept in mind when testing the model. A ZeroR algorithm would for instance already classify 65.5% of the instances correctly by pure chance. Although this seems accurate, this would obviously not be a good model, since ZeroR ignores all predictors and simply picks the majority category.

Further Visualization

Correlation Cell Size, Cell Shape

The following graph shows the relation between Cell Size and Cell Shape as discussed earlier when looking at figure 1.

```
# Make plot
# coef(linear.model <- lm(Cell_Shape_Uniformity ~ Cell_Size_Uniformity, data = data))

ggplot(data = data, mapping = aes(x = Cell_Size_Uniformity, y = Cell_Shape_Uniformity)) +
    # Add points
geom_jitter(aes(col = Class), size = 0.7, alpha = 0.5, width = 0.36, height = 0.36) +</pre>
```

```
# Add linear regression line
geom_smooth(method = "lm", se = T, col='darkgrey') +
# Set axis labels
xlab("Cell Size Uniformity (grade 1-10)") +
ylab("Cell Shape Uniformity (grade 1-10)") +
# Scale x and y axis to represent the 1 - 10 grading
ylim(0, 10) +
scale_y_continuous(breaks = seq(0, 10, by = 2)) +
xlim(0, 10) +
scale_x_continuous(breaks = seq(0, 10, by = 2)) +
labs(caption="Figure 3 ~ Cell Shape as a function of Cell Size.
    The color of the points show their class, red being benign, blue being malignant.") +
# Make theme minimal
theme_minimal() +
theme(plot.caption = element_text(hjust = 0.5))
```

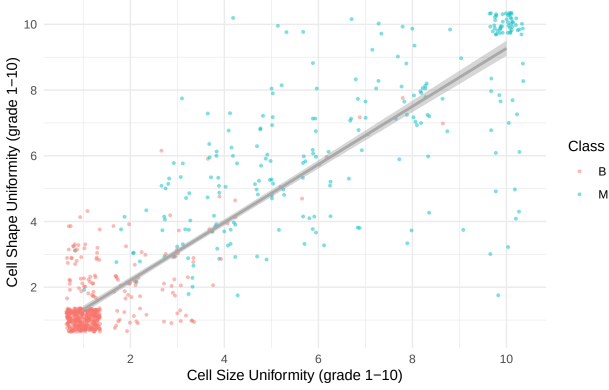


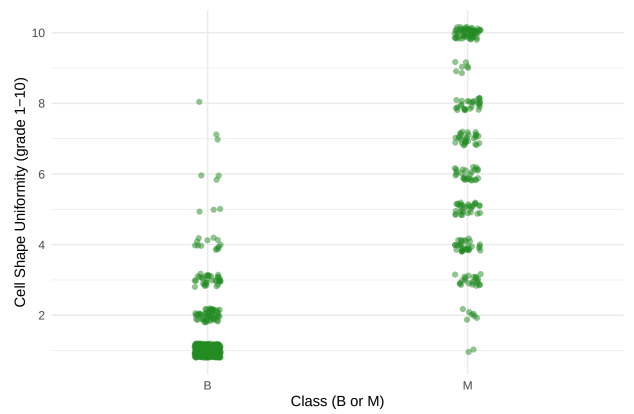
Figure 3 \sim Cell Shape as a function of Cell Size. The color of the points show their class, red being benign, blue being malignant.

Figure 3 shows the correlation between the attributes Cell_Shape_Uniformity and Cell_Size_Uniformity. When looking at the spread of the two colors in this graph, the benign instances seem to form a group in the bottom left, while the malignant cases are spread across the middle and the top right. This division suggests these attributes would be good predictors for breast cancer malignancy. A simple perceptron could divide the two classes with some accuracy based on these two attributes alone.

How does cell shape relate to malignancy

Below we see how cell shape relates to malignancy, by plotting points with jitter to show how many points represent each grade.

```
ggplot(data = data, mapping = aes(x = Class, y = Cell_Shape_Uniformity)) +
  geom_jitter(width = 0.05, height = 0.2, col = "forestgreen", alpha = 0.5) +
  ylim(0, 10) +
  xlab("Class (B or M)") +
  ylab("Cell Shape Uniformity (grade 1-10)") +
  scale_y_continuous(breaks = seq(0, 10, by = 2)) +
  labs(caption="Figure 4 ~ Cell Shape as a function of Class. Jitter and transparancy is added to show theme_minimal()
```



3 4 ~ Cell Shape as a function of Class. Jitter and transparancy is added to show the denisity of points at each grade.

Looking at figure 4, we can conclude that most of benign cases will get a cell shape grade of 2 or lower. The malignant cases, however, appear more spread out. This would cause inaccuracy when classifying with this attribute only. # Set data to csv

```
to.csv <- na.omit(data)
write.csv(to.csv, "breast_cancer_malignancy_dataset.csv", row.names = FALSE)</pre>
```

Trying machine learning algorithms

When testing different machine learning algorithms it is import a decision is made about sensitivity versus specificity. A very sensitive model may catch most if not all malignant cases, but might have a higher false positive rate. A very specific model will eliminate false positives, but may miss more malignant cases. In this case I will opt for sensitivity over specificity, because the consequences of a missed malignant case are higher than that of a falsely diagnosed benign case.

In order to achieve this, we can add a cost matrix to the uses machine learning algorithm that ways false negatives higher than Before we start optimizing for this however, let us first look which algorithms perform best on their own, before we add a cost matrix.

In most tested models shown below, only attributes with a correlation of 0.75 or higher to the Class column, as found in the exploratory data analysis above (see figure 1), where used as predictors unless stated otherwise. All models where trained and tested in Weka

Cost Insensitive Classifiers

J48

This model was trained using only the following attributes as predictors:

- Cell Size Uniformity
- Cell_Shape_Uniformity
- Bare_Nuclei
- Bland Chromatin

Accuracy: 9 correctly classified instances

Recall for M: 0.941 Precision for M: 0.930

Table 1: Confusion Matrix J48

| classified as -> | В | Μ |
|------------------|-----|-----|
| В | 426 | 17 |
| M | 14 | 225 |

Random Forest

This model was trained using only the following attributes as predictors: $\[$

- $\bullet \quad {\bf Cell_Size_Uniformity}$
- Cell_Shape_Uniformity
- $\bullet~$ Bare_Nuclei
- Bland_Chromatin

Accuracy: 96.04% correctly classified instances

Recall for M: 0.937 Precision for M: 0.949

Table 2: Confusion Matrix Random Forest

| classified as -> | В | Μ |
|------------------|-----|-----|
| B | 431 | 12 |
| M | 15 | 224 |

Logistic regression

This model was trained using only the following attributes as predictors:

- Cell_Size_Uniformity
- Cell_Shape_Uniformity
- \bullet Bland_Chromatin

Accuracy: 95.89% correctly classified instances

Recall for M: 0.933 Precision for M: 0.949

Table 3: Confusion Matrix Logistic

| classified as -> | В | Μ |
|------------------|-----|-----|
| В | 431 | 12 |
| M | 16 | 223 |

Naive Bayes

This model was trained using only the following attributes as predictors:

- $\bullet \quad Cell_Size_Uniformity$
- \bullet Cell_Shape_Uniformity
- Bare_Nuclei
- Bland_Chromatin

Accuracy: 95.31% correctly classified instances

Recall for M: 0.950 Precision for M: 0.919

Table 4: Confusion Matrix Naive Bayes

| classified as -> | В | Μ |
|------------------|-----|-----|
| В | 423 | 20 |
| M | 12 | 227 |

Logistic with more attributes

This model was trained using only the following attributes as predictors:

- Clump_Thickness
- Cell_Size_Uniformity
- Cell_Shape_Uniformity
- Marginal Adhesion
- Single_Epithelial_Cell_Size
- Bare Nuclei
- Bland_Chromatin
- $\bullet \quad Normal_Nucleoli$

Accuracy: 96.77% correctly classified instances

Recall for M: 0.954 Precision for M: 0.954

Table 5: Confusion Matrix Logistic with more attributes

| classified as -> | В | Μ |
|------------------|-----|-----|
| В | 432 | 11 |
| M | 11 | 228 |

Cost Sensitive Classifiers

Logistic Cost Sensitve Classifier

 dingen

ROC

```
roc_data <- read.table("ROC.arff",</pre>
                    sep = ",",
                    comment.char = "@")
names(roc_data) <- c("Instance_number",</pre>
                  "True Positives",
                  "False_Negatives",
                  "False_Positives",
                  "True_Negatives",
                  "False_Positive_Rate",
                  "True_Positive_Rate",
                  "Precision",
                  "Recall",
                  "Fallout",
                  "FMeasure",
                  "Sample_Size",
                  "Lift",
                  "Threshold")
```

head(roc_data)

```
Instance_number True_Positives False_Negatives False_Positives True_Negatives
## 1
                                239
                   0
                                                                 443
                                                                                  0
## 2
                                239
                                                   0
                   1
                                                                 431
                                                                                 12
## 3
                   2
                                239
                                                   0
                                                                 422
                                                                                 21
## 4
                   3
                                239
                                                   0
                                                                 413
                                                                                 30
## 5
                   4
                                239
                                                   0
                                                                 401
                                                                                 42
                   5
                                                   0
## 6
                                239
                                                                 388
                                                                                 55
##
    False_Positive_Rate True_Positive_Rate Precision Recall Fallout FMeasure
## 1
                                                            1 0.64956 0.519001
                1.000000
                                          1
                                              0.35044
## 2
                0.972912
                                          1 0.356716
                                                            1 0.643284 0.525853
## 3
                0.952596
                                          1 0.361573
                                                            1 0.638427 0.531111
## 4
                0.932280
                                          1 0.366564
                                                            1 0.633436 0.536476
## 5
                0.905192
                                          1 0.373437
                                                            1 0.626563
                                                                         0.5438
                0.875847
## 6
                                             0.38118
                                                            1 0.61882 0.551963
##
    Sample Size
                     Lift Threshold
## 1
        1.000000
                        1 0.005601
## 2
        0.982405 1.01791 0.006906
## 3
       0.969208 1.03177 0.007832
       0.956012 1.046012
                           0.008255
## 5
       0.938416 1.065625
                           0.008821
## 6
       0.919355 1.087719 0.008936
```

library(ggpubr)

Warning: package 'ggpubr' was built under R version 4.0.5

