Log Journal Project Theme 9

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EDA of yeast data: Protein Localization

The data has been retrived from uci. The dataset has been gathered of 1484 yeast sequences from SWISS-PROT using the annotations from YPD.

Attributes information:

- Sequence Name: Accession number for the SWISS-PROT database.
- mcg: McGeoch's method for signal sequence recognition.
- gvh: von Heijne's method for signal sequence recognition.
- alm: Score of the ALOM membrane spanning region prediction program.
- mit: Score of discriminant analysis of the amino acid content of the N-terminal region (20 residues long) of mitochondrial and non-mitochondrial proteins.
- erl: Presence of "HDEL" substring (thought to act as a signal for retention in the endoplasmic reticulum lumen). Binary attribute.
- pox: Peroxisomal targeting signal in the C-terminus.
- vac: Score of discriminant analysis of the amino acid content of vacuolar and extracellular proteins.
- nuc: Score of discriminant analysis of nuclear localization signals of nuclear and non-nuclear proteins.
- Class Distribution: The class is the localization site. Consisting of (abbreviation (full name) the amount):

CYT	(cytosolic or cytoskeletal)	463
NUC	(nuclear)	429
MIT	(mitochondrial)	244
ME3	(membrane protein, no N-terminal signal)	163
ME2	(membrane protein, uncleaved signal)	51
ME1	(membrane protein, cleaved signal)	44
EXC	(extracellular)	37
VAC	(vacuolar)	30
POX	(peroxisomal)	20
ERL	(endoplasmic reticulum lumen)	5

Codebook

Since there is not a codebook. I created my own codebook.

Name	Fullname	Datatypes
seq.name	Accession number for the SWISS-PROT	str
	database	
mcg	McGeoch's method for signal sequence	float
	recognition	
gvh	von Heijne's method for signal sequence	float
	recognition	
$_{ m alm}$	Score of the ALOM membrane spanning region	float
	prediction program	
mit	Score of discriminant analysis of the amino	float
	acid content of the N-terminal region	
erl	Presence of 'HDEL' substring	float
pox	Peroxisomal targeting signal in the C-terminus	bool
vac	Score of discriminant analysis of the amino acid	float
	content of vacuolar and extracellular proteins	
nuc	Score of discriminant analysis of nuclear	float
	localization signals of nuclear and non-nuclear	
	proteins	
loc.site	The class is the localization site	factor

Here is the codebook. With the attributes abbreviation, explanation and data types. As you can see there are many float datatypes.

Load the data

Now we need to load in the data. And change the column names to the attributes abbreviation, since these are non-existent. Let's give all the columns a name. And let's take a quick look at the data.

```
<chr>>
             <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
                                             <dbl> <dbl> <dbl> <chr>
1 ADT1_YEAST
              0.58
                                 0.13
                                         0.5
                                               0
                                                    0.48
                                                          0.22 MIT
                    0.61
                           0.47
2 ADT2 YEAST
              0.43
                     0.67
                                 0.27
                                         0.5
                                                    0.53
                                                          0.22 MIT
                           0.48
                                               0
3 ADT3_YEAST
              0.64
                                                          0.22 MIT
                     0.62
                           0.49
                                 0.15
                                         0.5
                                               0
                                                    0.53
4 AAR2 YEAST
                                                    0.54
                                                          0.22 NUC
              0.58
                    0.44
                           0.57
                                 0.13
                                         0.5
                                               0
5 AATM YEAST
             0.42
                    0.44
                           0.48
                                 0.54
                                         0.5
                                               0
                                                    0.48 0.22 MIT
6 AATC YEAST 0.51
                    0.4
                           0.56 0.17
                                         0.5
                                               0.5
                                                   0.49 0.22 CYT
```

The yeast data set contains scores per cellular localization sites.

Clean the data

We can drop the first columns since it is not necessary. Since the sequence names contribute nothing to create a prediction model.

```
data <- data[, -1]</pre>
  head(data)
# A tibble: 6 x 9
                                                                  gvh
                                                                                                         alm
                                                                                                                                                 mit
                                                                                                                                                                                        erl
                                                                                                                                                                                                                              pox
                                                                                                                                                                                                                                                                       vac
                                                                                                                                                                                                                                                                                                             nuc loc.site
              <dbl> 
                 0.58
                                                          0.61
                                                                                                 0.47
                                                                                                                                         0.13
                                                                                                                                                                                        0.5
                                                                                                                                                                                                                              0
                                                                                                                                                                                                                                                               0.48
                                                                                                                                                                                                                                                                                                     0.22 MIT
                                                                                                                                                                                                                                                                                                     0.22 MIT
                 0.43
                                                          0.67
                                                                                                  0.48
                                                                                                                                         0.27
                                                                                                                                                                                        0.5
                                                                                                                                                                                                                              0
                                                                                                                                                                                                                                                               0.53
3
                 0.64
                                                          0.62
                                                                                                  0.49
                                                                                                                                         0.15
                                                                                                                                                                                        0.5
                                                                                                                                                                                                                              0
                                                                                                                                                                                                                                                               0.53
                                                                                                                                                                                                                                                                                                     0.22 MIT
4
               0.58
                                                        0.44
                                                                                                                                                                                                                                                                                                     0.22 NUC
                                                                                                 0.57
                                                                                                                                         0.13
                                                                                                                                                                                        0.5
                                                                                                                                                                                                                              0
                                                                                                                                                                                                                                                               0.54
               0.42
                                                        0.44
                                                                                                 0.48
                                                                                                                                         0.54
                                                                                                                                                                                        0.5
                                                                                                                                                                                                                              0
                                                                                                                                                                                                                                                               0.48
                                                                                                                                                                                                                                                                                                     0.22 MIT
             0.51
                                                     0.4
                                                                                                  0.56
                                                                                                                                         0.17
                                                                                                                                                                                       0.5
                                                                                                                                                                                                                              0.5 0.49 0.22 CYT
```

The first column has been successfully dropped from the data.

Are there any missing values? Let's take a quick look.

```
# Count all the NA values
sum(is.na(data))
```

[1] 0

There are not any missing values. So nothing needs to be changed for this.

Now we need to take a clearer look at the data set using str().

```
str(data)
```

As you can see, the last column, loc.site, consists of characters, but this one needs to be converted to factors for clarity and complexity.

```
# Transform the last column to a factor
data$loc.site <- as.factor(data$loc.site)
str(data$loc.site)</pre>
```

```
Factor w/ 10 levels "CYT", "ERL", "EXC", ...: 7 7 7 8 7 1 7 8 7 1 ...
```

As you can see the data type of the column loc.site has been successfully changed to factors.

There are alot of rows. Let's visualize the amount of each classification.

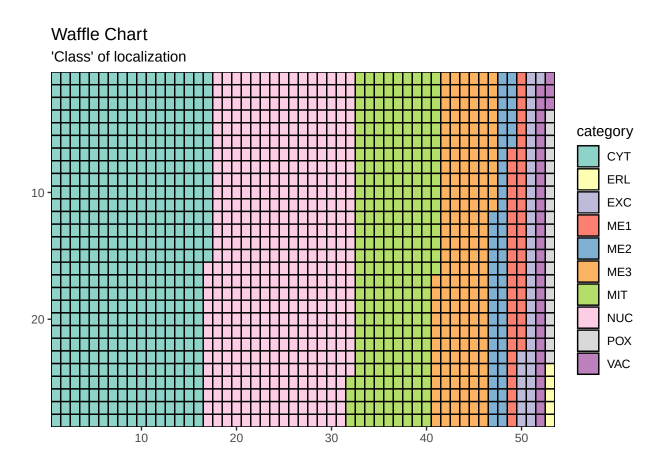


Figure 1: A waffle chart of the categorical composition

There are alot of CYT and NUC localization. This probably means that CYT has a greater variation in localisation than ERL.

The ERL variable also should be changed. As you can see, it is now a num data type but I needs to be a bool/binary datatype. Let's take a closer look at th ERL column.

summary(data\$erl)

```
Min. 1st Qu. Median Mean 3rd Qu. Max. 0.5000 0.5000 0.5000 0.5047 0.5000 1.0000
```

This looks weird because the value should be 0 or 1.

There are alot of 0.5 values. But should not exist in this column. The variable need to be a bool, or in other words it must be a 0 or a 1. So all the 0.5 values needs to be changed to a 0.

```
table(data[, "erl"])
```

0.5 1 1470 14

Before the data transformation there are 1470 counts of the value 0.5 and 14 of 1.

```
# Change every 0.5 value to a 0
data$erl[data$erl == 0.5] <- 0
table(data[,"erl"])</pre>
```

0 1 1470 14

The data has been successfully transformed. Now the datatype has to be changed to a bool.

```
# Change the datatype to a logical
data$erl <- as.logical(data$erl)
str(data$erl)</pre>
```

```
logi [1:1484] FALSE FALSE FALSE FALSE FALSE ...
```

Now every column has the right datatype.

Univariate Analysis

Let's take a quick look at the data using summary(). We can drop column 5 and 9 for this. Column 5, ERL, is a logical datatype and column 9, loc.site, is a string datatype.

summary(data[, -c(5,9)])

```
gvh
                                          alm
                                                          mit
     mcg
       :0.1100
                  Min.
                          :0.1300
                                    Min.
                                            :0.21
                                                     Min.
                                                            :0.0000
                  1st Qu.:0.4200
1st Qu.:0.4100
                                    1st Qu.:0.46
                                                     1st Qu.:0.1700
Median : 0.4900
                  Median :0.4900
                                    Median:0.51
                                                     Median :0.2200
                                            :0.50
Mean
       :0.5001
                  Mean
                          :0.4999
                                    Mean
                                                     Mean
                                                            :0.2612
3rd Qu.:0.5800
                  3rd Qu.:0.5700
                                     3rd Qu.:0.55
                                                     3rd Qu.:0.3200
Max.
        :1.0000
                  Max.
                          :1.0000
                                    Max.
                                            :1.00
                                                     Max.
                                                            :1.0000
     pox
                        vac
                                          nuc
Min.
       :0.0000
                          :0.0000
                                            :0.0000
                  Min.
                                    Min.
                  1st Qu.:0.4800
1st Qu.:0.0000
                                    1st Qu.:0.2200
Median :0.0000
                  Median :0.5100
                                    Median :0.2200
Mean
       :0.0075
                  Mean
                          :0.4999
                                    Mean
                                            :0.2762
```

```
3rd Qu.:0.0000 3rd Qu.:0.5300 3rd Qu.:0.3000 Max. :0.8300 Max. :0.7300 Max. :1.0000
```

As you can see all the datapoints are between 0 and 1. Se the data already has been transformed with a min-max normalization.

Let's visualise this with ggplot. Using jitterpoints and a violing plot.

```
p1 <- ggplot(data, mapping = aes(x = "", y = mcg)) + geom_violin(alpha=0.2) +
  geom_jitter(width = 0.2, alpha = 0.25, height = 0, color = "red") + xlab(NULL)
p2 <- ggplot(data, mapping = aes(x = "", y = gvh)) + geom_violin(alpha=0.2) +
 geom_jitter(width = 0.2, alpha = 0.25, height = 0, color = "blue") + xlab(NULL)
p3 <- ggplot(data, mapping = aes(x = "", y = alm)) + geom_violin(alpha=0.2) +
  geom_jitter(width = 0.2,alpha = 0.25, height = 0, color = "purple") + xlab(NULL)
p4 <- ggplot(data, mapping = aes(x = "", y = mit)) + geom_violin(alpha=0.2) +
  geom_jitter(width = 0.2, alpha = 0.25, height = 0, color = "brown") + xlab(NULL)
p5 <- ggplot(data, mapping = aes(x = "", y = pox)) + geom_violin(alpha=0.2) +
  geom_jitter(width = 0.2,alpha = 0.25, height = 0, color = "orange") + xlab(NULL)
p6 <- ggplot(data, mapping = aes(x = "", y = vac)) + geom_violin(alpha=0.2) +
 geom_jitter(width = 0.2, alpha = 0.25, height = 0, color = "green") + xlab(NULL)
p7 <- ggplot(data, mapping = aes(x = "", y = nuc)) + geom_violin(alpha=0.2) +
  geom_jitter(width = 0.2,alpha = 0.25, height = 0, color = "orange") + xlab(NULL)
plot <- ggarrange(p1, p2, p3, p4, p5, p6, p7, nrow = 4, ncol = 2)
annotate_figure(plot, top = text_grob("Boxplots", face = "bold", size = 14))
```

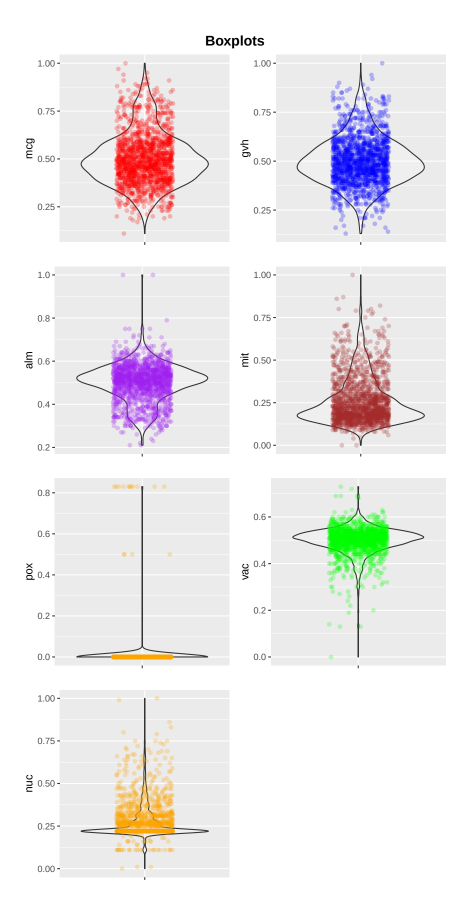


Figure 2: Boxplot comparing basic statistic for all columns

Mcg and gvh are normally distributed. Alm and mit are skewed but nothing crazy to worry about. Vac and nuc are really skewed however, not yet bad enough to worry about it. Pox is looks weird let's take a detailed look at pox.

```
table(data[, "pox"])
```

```
0 0.5 0.83
1469 4 11
```

As you can see almost all the numbers are 0. This does not mean that we just discord this column. Maybe it is very significant if the number is not 0.

```
subset(data, pox > 0, select=loc.site)
# A tibble: 15 x 1
   loc.site
   <fct>
 1 CYT
 2 MIT
3 POX
 4 POX
5 POX
 6 POX
7 POX
8 POX
9 POX
10 MIT
11 POX
12 MIT
13 POX
14 POX
15 POX
```

If the number is non-zero then the probability is very high that the protein is localised in peroxisomal region.

Bivariate Analysis

With a heatmap, you can easily see where there are correlations between variables. First let's create a correlation matrix

```
# Create a cor matrix
cor_matrix <- cor(data[, -c(5,9)])
cor_matrix <- as_tibble(cor_matrix)
# Add a column with the varibale names
cor_matrix <- cor_matrix %>% mutate(varnames = all_of(attr_names)[-c(1,6,10)])
kbl(cor_matrix, booktabs = T, align = "c") %>%
    kable_styling(full_width = T)
```

$\overline{\mathrm{mcg}}$	gvh	alm	mit	pox	vac	nuc	varnames
1.0000000	0.5816314	-0.1639513	0.1581755	0.0055970	0.0750427	-0.1245404	mcg
0.5816314	1.0000000	-0.2718000	0.1403139	0.0003918	0.0887594	-0.1029840	gvh
-0.1639513	-0.2718000	1.0000000	0.0596683	0.0093779	-0.1858054	-0.0220428	$_{ m alm}$
0.1581755	0.1403139	0.0596683	1.0000000	-0.0090398	-0.1035914	-0.0547965	mit
0.0055970	0.0003918	0.0093779	-0.0090398	1.0000000	0.0208997	-0.0356586	pox
0.0750427	0.0887594	-0.1858054	-0.1035914	0.0208997	1.0000000	0.0896904	vac
-0.1245404	-0.1029840	-0.0220428	-0.0547965	-0.0356586	0.0896904	1.0000000	nuc

The calculated correlation matrix. This needs to be tranformed to a long matrix.

varnames	variable	cor
mcg	mcg	1
mcg	gvh	0.5816
mcg	alm	-0.164
mcg	mit	0.1582
mcg	pox	0.005597
mcg	vac	0.07504

The long calculated correlation matrix.

```
ggplot(data = cor_matrix_long, aes(x=varnames, y=variable, fill=cor)) +
    geom_tile() +
    labs(x=NULL, y=NULL, title="Heatmap Correlation") +
    scale_fill_gradient(high = "purple", low = "white")
```

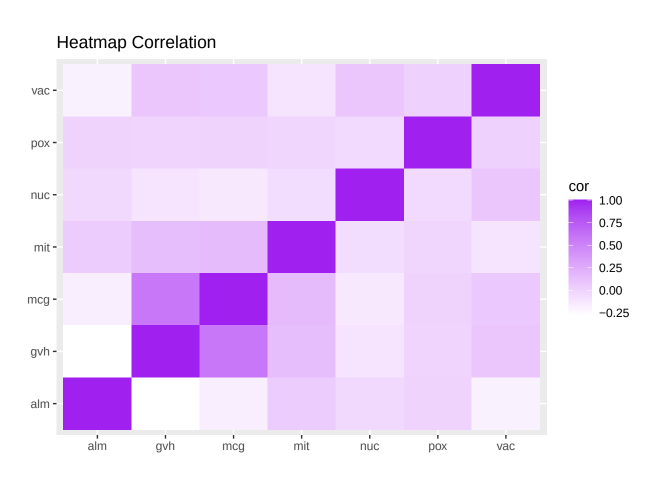


Figure 3: A heatmap pairwise correlation of selected numeric variables

As seen from the heatmap there is some correlation between mcg and gvh. Let's visualise this.

Scatterplot and trendline

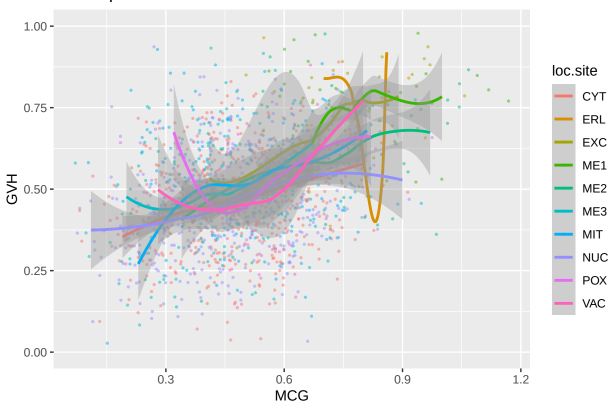


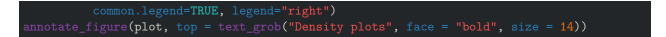
Figure 4: Scatterplot with trendline with the dependent variables

Every variable goes with a slow upward trend except erl this one has a weird curve.

Class labels

Now we need to look at how the data correlates with the classes. The height of the peak doesn't matter much, shifted peaks do.

```
p1 <- ggplot(data, aes(x=mcg)) + geom_density(aes(color=loc.site))
p2 <- ggplot(data, aes(x=gvh)) + geom_density(aes(color=loc.site))
p3 <- ggplot(data, aes(x=alm)) + geom_density(aes(color=loc.site))
p4 <- ggplot(data, aes(x=mit)) + geom_density(aes(color=loc.site))
p5 <- ggplot(data, aes(x=pox)) + geom_density(aes(color=loc.site))
p6 <- ggplot(data, aes(x=vac)) + geom_density(aes(color=loc.site))
p7 <- ggplot(data, aes(x=nuc)) + geom_density(aes(color=loc.site))
plot <- ggarrange(p1, p2, p3, p4, p5, p6, p7, ncol=4, nrow=2,</pre>
```



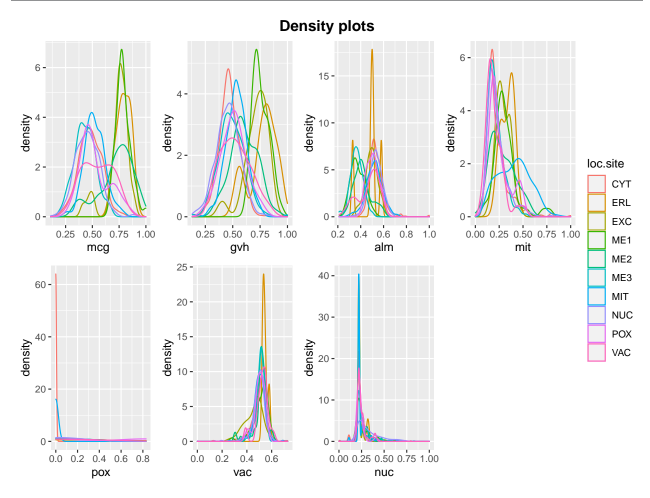


Figure 5: Density plots shows class distinction

At pox, vac and nuc, you don't see shifted peaks. At mcg and gvh you really see shifted peaks this shows a distribution of the different classes. There are small shifted peaks at alm en mit, this shows that there is difference but not so much. Let's look at mcg and gvh using ridge lines plot.

```
p1 <- ggplot(data, aes(x=mcg, y=loc.site, fill = loc.site)) +
    geom_density_ridges2(rel_min_height = 0.005) + theme_minimal() +
    coord_cartesian(clip = "off")
p2 <- ggplot(data, aes(x=gvh, y=loc.site, fill = loc.site)) +
    geom_density_ridges2(rel_min_height = 0.005) + theme_minimal() +
    coord_cartesian(clip = "off")
plot <- ggarrange(p1, p2, common.legend = TRUE)
annotate_figure(plot, top = text_grob("Ridge line plot", face = "bold", size = 14))</pre>
```

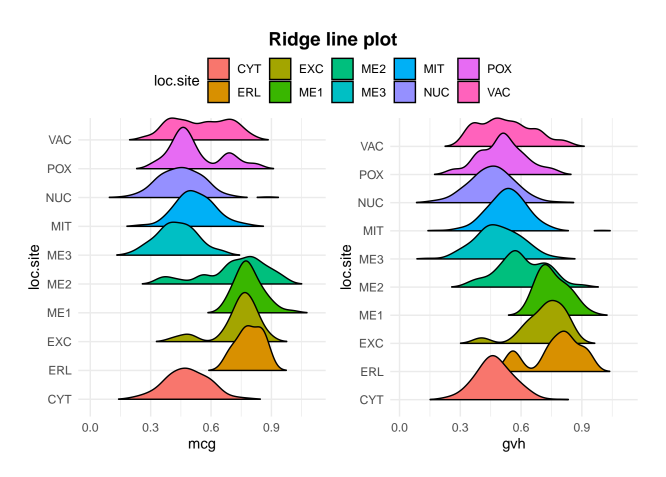


Figure 6: Ridge line plot between mcg and gvh

Now you can really see that the peaks are shifted.

We need to test the data to see if there is significant difference between it. Using a 1-way ANOVA test.

```
# Perform a one way anova test
res.aov <- summary(aov(mcg ~ loc.site, data = data))
res.aov[[1]]$`Pr(>F)`[1]
```

[1] 1.203676e-146

The P-value is < 0.05. So there is a significant difference.

Multivariate analysis

Here is the distance matrix.

One way to see if groups are clustered is to MDS plot the groups and calculated the distance matrix.

```
# Create a matrix
matrix <- with(data, rbind(mcg, gvh, alm, mit, pox, vac, nuc))</pre>
(distmat <- dist(matrix))</pre>
          mcg
                    gvh
                               alm
                                         mit
                                                   pox
                                                              vac
    4.623700
gvh
alm 6.699455
               6.524822
mit 11.476977 11.321051 11.025910
pox 19.910020 19.776585 19.479594 11.495734
vac 5.580672 5.083611 4.342165 10.946100 19.312255
nuc 11.161711 10.858508 10.144550 6.884693 11.545921 9.715282
```

autoplot(cmdscale(distmat, eig = TRUE), shape = FALSE, label = TRUE, label.size = 4)

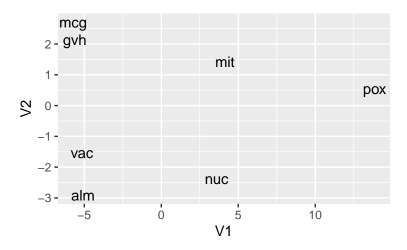


Figure 7: Classical (Metric) Multidimensional Scaling

As seen from above mcg and gvh are the clustered group.

A other plot to show is using PCA.

```
df <- data[-c(5,9)]
pca_res <- prcomp(df, scale. = TRUE)
autoplot(pca_res, data = data, colour = 'loc.site', loadings = TRUE, loadings.label = TRUE)</pre>
```

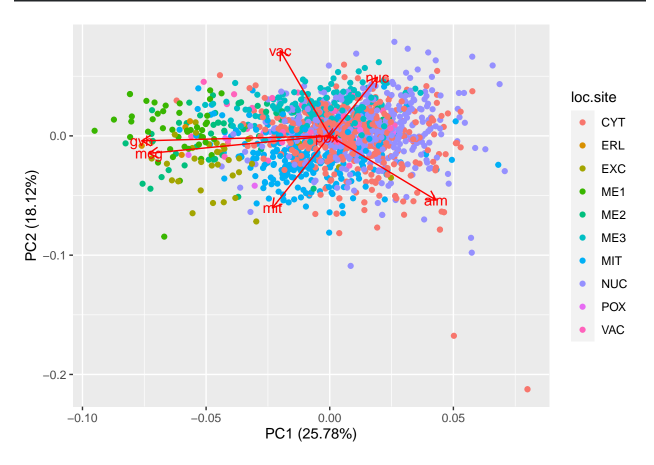


Figure 8: PCA plot showing the groups

As earlier shown mcg and gvh are grouped together.

MIL algorithms

First we need to investigate which standard ML algorithms is the best to use for this dataset using Weka.

Weka accepts .arff files. You can convert a .csv to a .arff file in Weka. So we need to export the clean dataset to a .csv file.

```
write.table(data, "data.csv", quote = FALSE, sep = ",", row.names=FALSE)
```

Open the .csv in Weka under tools -> ArffViewer. And saved the file as .arff. The first 15 lines of the .arff file needs to look like this.

```
cat(readLines("data.arff", n = 15), sep = "\n")

@relation data

@attribute mcg numeric
@attribute gvh numeric
@attribute alm numeric
@attribute mit numeric
@attribute erl {FALSE,TRUE}
@attribute pox numeric
@attribute vac numeric
@attribute nuc numeric
@attribute nuc numeric
@attribute loc.site {MIT,NUC,CYT,ME1,EXC,ME2,ME3,VAC,POX,ERL}

@data
0.58,0.61,0.47,0.13,FALSE,0,0.48,0.22,MIT
0.43,0.67,0.48,0.27,FALSE,0,0.53,0.22,MIT
```

Criteria for MIL algorithms

An accurate algorithm is needed that accurately predicts the dataset. It needs a high percentage of correct classifications on the dataset. An algorithm that is fast. What if there is a dataset of 1 million observations. And it should also not contain too many false negatives and false positives.

Research has been done on which algorithm is the best. From every representatives of all classifier categories. And ZeroR and OneR for a baseline performance. Eg: Decision Trees (C4.5, J48, RandomForest), Nearest Neighbor (IBk), SVM (SMO), NaiveBayes (Naïve Bayes) and Linear Logistic (SimpleLogistic). And carried out classifications 10-fold cross validation.

The experimenter can save the results in a csv file.

```
weka.result <- read.table("Result.csv", header = TRUE, sep = ",")
# Remove weka.classifiers.* from classifier name
for (i in 1:length(weka.result$Key_Scheme)) {
  name <- weka.result$Key_Scheme[i]
  tweaked.name <- strsplit(name, split = ".", fixed = TRUE)[[1]][4]
  weka.result$Key_Scheme[i] <- tweaked.name
}</pre>
```

Now the results of the csv can been plotted with ease of use. Like the ROC and precision.

```
# Create a dataframe of the mean of the ROC of every classifier
res <- aggregate(Area_under_ROC ~ Key_Scheme, data = weka.result, mean)

ggplot(data=res, aes(x=Key_Scheme, y=Area_under_ROC, fill=Key_Scheme)) +
    geom_bar(stat="identity") +
    theme(axis.text.x = element_text(angle = 45, vjust = 1, hjust=1)) +</pre>
```

Area under the curve for different classifiers

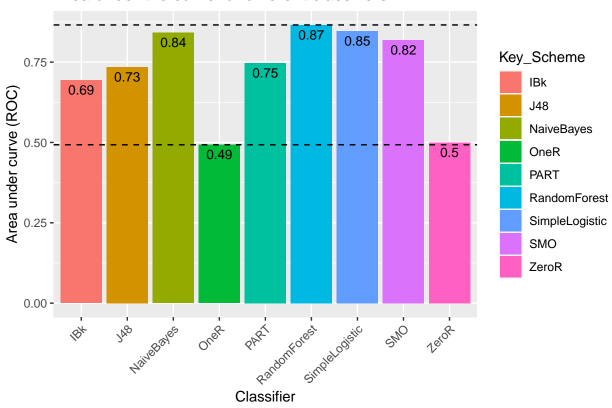


Figure 9: Area under the curve for different classifiers

RandomForest has the highest area under the curve, while OneR has the lowest with 0.49 and ZeroR with a close second last with 0.5.

```
res <- aggregate(IR_precision ~ Key_Scheme, data = weka.result, mean)

ggplot(data=res, aes(x=Key_Scheme, y=IR_precision, fill=Key_Scheme)) +
    geom_bar(stat="identity") +
    theme(axis.text.x = element_text(angle = 45, vjust = 1, hjust=1)) +
    labs(x="Classifier", y="Precision",
        title="Average precision for different classifiers") +
    geom_hline(aes(yintercept=min(IR_precision)), linetype = 'dashed') +
    geom_hline(aes(yintercept=max(IR_precision)), linetype = 'dashed') +
    geom_text(aes(label=round(IR_precision, digits=2)), vjust=1.5, size=3.5)</pre>
```

Average precision for different classifiers

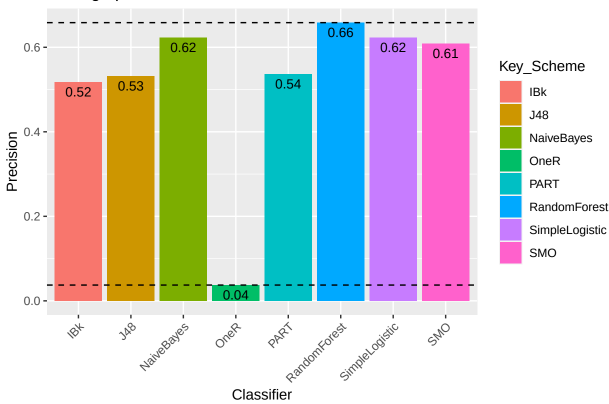


Figure 10: Average precision for different classifiers

RandomForest also has the highest average precision with 0.66. And OneR has the lowest with 0.04. ZeroR is not in this plots because it only consist of NaN.

Several algorithms cannot make a prediction for the classification VAC (0.000) and for some they get an error (?). 1 algorithm IBk has a very low prediction for VAC. All the algorithm above in the figure except for ZeroR, OneR, SMO and SimpleLogistic were used in the Weka Experimenter to compare.

```
>-< > < Resultset
4 4 0 trees.RandomForest '-P 100 -I 100 -num-slots 1 -K 0 -M 1.0 -V 0.001 -S 1' 1116839470751428698
1 2 1 bayes.NaiveBayes '' 5995231201785697655
0 1 1 trees.J48 '-C 0.25 -M 2' -217733168393644444
-2 0 2 rules.PART '-C 0.25 -M 2' 8121455039782598361
-3 0 3 lazy.IBk '-K 1 -W 0 -A \"weka.core.neighboursearch.LinearNNSearch -A \\\"weka.core.EuclideanDistance -R first-last\\\\"\"' -3080186098777067172
```

Figure 11: Ranking of chosen algorithms

This shows the ranking of the algorithms by the number of times a given algorithm beat the other algorithms. And RandomForest beat all 4 algorithms.

Dataset	(1)	trees.Ra	I	(2) rules	(3) tree	s (4)	lazy.	(5) bay	yes
data	(100)	61.25	I	54.70 *	56.38	* 5	2.61 *	57.88	8 *
		(v/ /*)	ī	(0/0/1)	(0/0/1	.) (0/0/1)	(0/0,	/1)

Figure 12: Ranking of chosen algorithms by SD

RandomForest has a 'v' next to their result. This means that the difference in the accuracy for these algorithms compared to all the other algorithms is significant.

After testing multiple algorithms in Weka, the algorithm RandomForest was chosen. It has the highest average precision of 0.611. And this was one of the few algorithms where the precision for VAC was not a '?'.

But for the ratings, no algorithm could make a prediction except 1 for VAC, however, it was also very low. So we looked at whether there is significant difference if we remove this classifier.

Dataset	(1)	trees.Ra	nd
data	(100)	61.25	1
data-weka.filters.unsuper	(100)	62.66	-1
(∀/ /*)			1

Figure 13: Removed class difference

And by the looks of it, there is no significant difference. So VAC is not removed from the dataset.

RandomForest has seven attributes. Below is a table with each attribute and its meaning.

Attribute	Description
P	Size of each bag, as a percentage of the training
	set size.
I	Number of iterations.
num-slots	Number of execution slots.
K	Number of attributes to randomly investigate.
${ m M}$	Number of attributes to randomly investigate.
V	Set minimum numeric class variance proportion.
S	Seed for random number generator.

Weka will find the optimal parameters for a classifier. With CVParameterSelection in Weka were the best parameters found.

Attribute	Old.Values	New.Values
P	100	50
I	100	1000
num-slots	1	0
K	0	1
${ m M}$	1	1

Attribute	Old.Values	New.Values
V	0.001	0.001
S	1	1

This table shows the parameters with their default settings and their new. P, I, num-slots and K were different than the default settings. The build will take more time because it does 10 times more I(terations).

Dataset	(2)	trees.Ra	I	(1) trees
data	(100)	62.51	ı	63.56

Figure 14: Iterations ranked

Number 1 in the figure above is with 1000 I(terations) and number 2 is 100 I(terations). And there is not a significant difference. So for performance, the value for I will be 100.

Weka Meta learners were used to get a better idea of an optimal classifier.

Classifier	TP Rate	FP Rate	Precision	ROC Area	
RandomForest.Standard	0.611	0.132	0.603	0.844	Weighted.Avg
RandomForest.tweaked	0.62	0.132	0.610	0.846	Weighted.Avg
Boosting	0.586	0.142	0.586	0.8	Weighted.Avg
Stacking	0.544	0.157	0.538	0.758	Weighted.Avg
Bagging	0.623	0.13	?	0.853	Weighted.Avg
Vote	0.625	0.128	?	0.851	Weighted.Avg

As it can be seen in this table, RandomForest with tweaked parameters is the best. This one has no '?' at VAC and precision.