Urinary Biomarkers for Pancreatic Cancer

Log Theme
09 - Introduction Machine Learning

Lisa Hu

414264

Bio-Informatics

Hanzehogeschool Groningen, ILST

Dave Langers (LADR) & Bart Barnard (BABA)

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1 Data description

The data can be found on kaggle.com: Urinary biomarkers for pancreatic cancer The files are saved as Data.csv and Documentation.csv for easier access.

The following packages were used:

- \bullet dplyr
- readr
- pander
- FSA
- \bullet ggplot2
- ggpubr
- ggbiplot

2 Reading the data

We first want to create an insight of our data:

```
dataset <- read.csv("../data/Data.csv")
codebook <- read_delim("../data/codebook.txt", delim = "|")
pander(codebook[1:4], booktabs = T, caption = "Data values", split.tables = 100)</pre>
```

Table 1: Data values

| Name | Full Name | Type | Unit |
|---------------------------------|---------------------------|----------------------|-------|
| sample_id | Sample ID | chr | - |
| $patient_cohort$ | Patient's Cohort | chr | - |
| $\operatorname{sample_origin}$ | Sample Origin | chr | - |
| age | Age of subject | dbl | - |
| sex | Sex of subject | chr | - |
| diagnosis | Diagnosis | dbl | - |
| stage | Stage | chr | - |
| benign_sample_diagnosis | Benign Sample's Diagnosis | chr | - |
| plasma_CA19_9 | Blood plasma CA19-9 | dbl | U/ml |
| creatinine | Creatinine | dbl | mg/ml |
| LYVE1 | LYVE1 | dbl | ng/ml |
| REG1B | REG1B | dbl | ng/ml |
| ${ m TFF1}$ | ${ m TFF1}$ | dbl | ng/ml |
| REG1A | REG1A | dbl | ng/ml |

Table 2: Description

| Name | Description |
|-------------------------|---|
| sample_id | Unique string identifying each subject |
| patient_cohort | Cohort $1 = $ previously used samples; Cohort 2 |
| | = newly added samples |
| $sample_origin$ | BPTB: Barts Pancreas Tissue Bank, London, |
| | UK; ESP: Spanish National Cancer Research |
| | Centre, Madrid, Spain; LIV: Liverpool |
| | University, UK; UCL: University College |
| | London, UK |
| age | Age in years |
| sex | M = male; F = female |
| diagnosis | 1 = control (no cancer); 2 = benign |
| | hepatobiliary disease; $3 = PDA$ (pancreatic |
| | cancer) |
| stage | The stage of the disease (IA, IB, IIA, IIB, III, |
| | IV) |
| benign_sample_diagnosis | The diagnosis for those with a benign diagnosis |

| Name | Description |
|---------------|--|
| plasma_CA19_9 | Blood plasma levels of CA19-9 monoclonal antibody, usually elevated when pancreatic cancer |
| creatinine | Urinary biomarker of kidney function |
| LYVE1 | Urinary levels of Lymphatic Vessel Endothelial |
| | Hyaluronan receptor 1 |
| REG1B | Urinary levels of Regenerating Family Member |
| | 1 Beta |
| TFF1 | Urinary levels of Trefoil Factor 1 |
| REG1A | Urinary levels of Regenerating Family Member |
| | 1 Alpha |

The information given in the codebook originates from the <code>Documentation.csv</code>. This file was given with the data file and can be found on the website.

3 Manipulate the data

A lot of the rows contain empty strings instead of NA, which has to be fixed first. Besides that, the columns sample_id, patient_cohort, sample_origin, and benign_sample_diagnosis in the dataset significant value for the analysis and are therefor dropped. A column diagnosis_group was added for a comparison test.

```
# Change the empty strings to NA
dataset[dataset == ""] <- NA</pre>
# Remove unnecessary columns
drop <- c("sample_id", "patient_cohort", "sample_origin", "benign_sample_diagnosis")</pre>
dataset <- dataset[,!(names(dataset) %in% drop)]</pre>
# Group the samples
dataset <- dataset %>%
  mutate(
    ## Factor for order of age
    diagnosis_group = factor(
      dplyr::case_when(
        diagnosis == 1 ~ "Control",
        diagnosis == 2 ~ "Benign",
        stage == "I" ~ "I-IIA",
        stage == "IA" ~ "I-IIA",
        stage == "IB" ~ "I-IIA",
        stage == "II" ~ "I-II",
        stage == "IIA" ~ "I-IIA",
        stage == "IIB" ~ "I-II",
        stage == "III" ~ "III-IV",
        stage == "IV" ~ "III-IV" ),
      level = c("Control", "Benign", "I-IIA", "I-II", "III-IV")
    ),
    ## Factor if there's a blood sample or not
    blood = factor(
      dplyr::case_when(
        plasma_CA19_9 >= 0 \sim "yes",
        TRUE ~ "no"),
      level = c("yes", "no")
```

3.1 REG1A vs. REG1B

Table 3: Adjusted p-values of Kruskal-Wallis test, Dunn's multiple comparisons; ns - not significant. The header shows the groups that were compared.

| | Control - I-II | Control - I-IIA | Control - III-IV |
|-------|----------------|-----------------|------------------|
| REG1A | 1.928479e-05 | ns | 4.837915e-07 |
| REG1B | 3.864924e-15 | 0.0002123534 | 5.789369e-17 |

```
pander(comparison[,c(1,3,5)], split.tables = 100, booktabs = T)
```

| | Benign - I-II | Benign - I-IIA | Benign - III-IV |
|-------|---------------|----------------|-----------------|
| REG1A | 0.000768779 | ns | 4.494778e-05 |
| REG1B | 1.200471e-12 | 0.001777207 | 3.927231e-14 |

```
dataset <- dataset[,!(names(dataset) %in% "REG1A")]</pre>
```

Although performance between the two is similar, a Kruskal-Wallis test with Dunn's multiple comparisons shows that REG1B outperforms REG1A when the control and benign samples are compared to the I-IIA PDAC samples. Therefor, REG1B was used further on in the experiments and REG1A is dropped.

3.2 Log transformation

A summary of the data shows very high maximum values, but rather low medians. A log-transformation is applied to correct this.

```
pander(summary(dataset), split.table = 100)
```

Table 5: Table continues below

| age | sex | diagnosis | stage | plasma_CA19_9 |
|-----------------|-----------------|----------------|-----------------|-----------------|
| Min. :26.00 | Length:590 | Min. :1.000 | Length:590 | Min.: 0.0 |
| 1st Qu.:50.00 | Class:character | 1st Qu.:1.000 | Class:character | 1st Qu.: 8.0 |
| Median $:60.00$ | Mode :character | Median : 2.000 | Mode :character | Median: 26.5 |
| Mean $:59.08$ | NA | Mean $:2.027$ | NA | Mean: 654.0 |
| 3rd Qu.:69.00 | NA | 3rd Qu.: 3.000 | NA | 3rd Qu.: 294.0 |
| Max. $:89.00$ | NA | Max. $:3.000$ | NA | Max. $:31000.0$ |
| NA | NA | NA | NA | NA's :240 |

| creatinine | LYVE1 | REG1B | TFF1 | diagnosis_group | blood |
|-----------------|-------------------|-------------------|------------------|-----------------|---------|
| Min. :0.05655 | Min.: 0.000129 | Min.: 0.0011 | Min.: 0.005 | Control:183 | yes:350 |
| 1st Qu.:0.37323 | 1st Qu.: 0.167179 | 1st Qu.: 10.7572 | 1st Qu.: 43.961 | Benign: 208 | no :240 |
| Median | Median: | Median: 34.3034 | Median: | I-IIA:27 | NA |
| :0.72384 | 1.649862 | | 259.874 | | |
| Mean $:0.85538$ | Mean: 3.063530 | Mean: 111.7741 | Mean: 597.869 | I-II: 75 | NA |
| 3rd Qu.:1.13948 | 3rd Qu.: | 3rd Qu.: | 3rd Qu.: 742.736 | III-IV : 97 | NA |
| | 5.205037 | 122.7410 | | | |
| Max. :4.11684 | Max. $:23.890323$ | Max. $:1403.8976$ | Max. :13344.300 | NA | NA |
| NA | NA | NA | NA | NA | NA |

```
log.data <- log(dataset[5:9] +1)
dataset[5:9] <- log.data</pre>
```

The samples are then grouped by diagnosis for easier access of the different samples. Table 5 shows the different amounts of samples per diagnosis and the amount of which are also blood samples.

```
# Different diagnosis and blood groups
control <- subset(dataset, diagnosis == 1)
benign <- subset(dataset, diagnosis == 2)
pdac <- subset(dataset, diagnosis == 3)

# Demographics
demograph <- dataset %>%
    group_by(sex, diagnosis, stage) %>% tally()
demograph.blood <- dataset %>%
    group_by(sex, blood) %>% tally()
pander(demograph, booktabs = T, caption = "Demographic of the samples")
```

Table 7: Demographic of the samples

| sex | diagnosis | stage | n |
|--------------|-----------|-------|-----|
| F | 1 | NA | 115 |
| \mathbf{F} | 2 | NA | 101 |
| \mathbf{F} | 3 | I | 1 |
| \mathbf{F} | 3 | IB | 6 |
| \mathbf{F} | 3 | II | 3 |
| \mathbf{F} | 3 | IIA | 6 |
| F | 3 | IIB | 33 |
| F | 3 | III | 27 |
| F | 3 | IV | 7 |
| M | 1 | NA | 68 |
| ${ m M}$ | 2 | NA | 107 |
| ${ m M}$ | 3 | IA | 3 |
| ${ m M}$ | 3 | IB | 6 |
| ${ m M}$ | 3 | II | 4 |
| M | 3 | IIA | 5 |
| M | 3 | IIB | 35 |
| ${ m M}$ | 3 | III | 49 |
| M | 3 | IV | 14 |

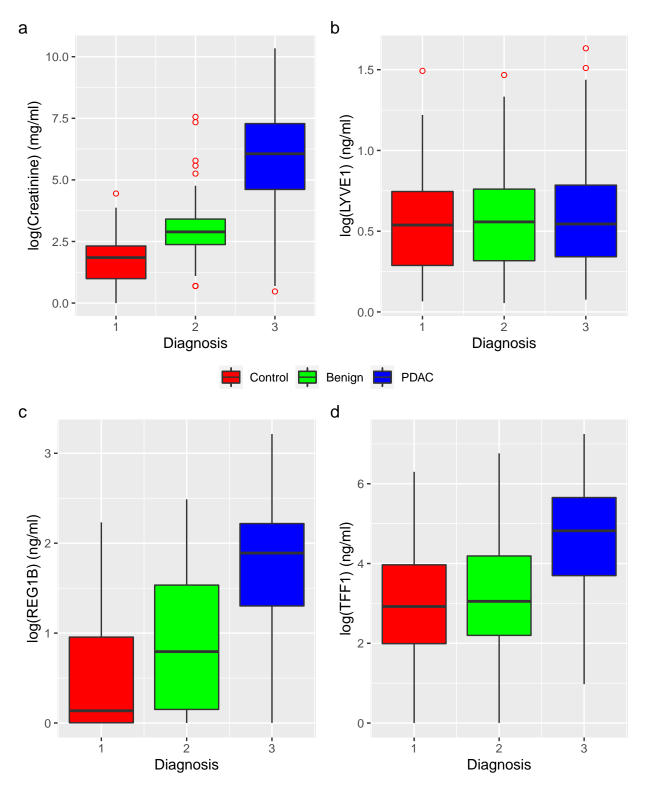
```
pander(demograph.blood, booktabs = T, caption = "Demographic of the blood samples")
```

Table 8: Demographic of the blood samples

| sex | blood | n |
|--------------|-------|-----|
| F | yes | 179 |
| \mathbf{F} | no | 120 |
| ${ m M}$ | yes | 171 |
| ${ m M}$ | no | 120 |

4 Analyse the data

4.1 Boxplots

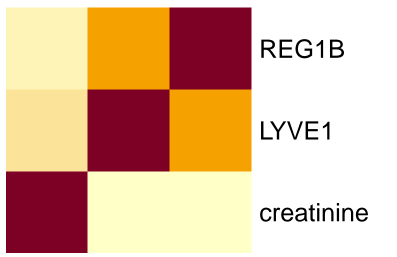


The outliers are not localized in a specific diagnosis group, but rather spread over the groups.

4.2 Correlation matrix

```
cor_matrix <- cor(dataset[,5:8])
heatmap(cor_matrix, scale = "column", Colv = NA, Rowv = NA, main = "Correlation matrix")</pre>
```

Correlation matrix



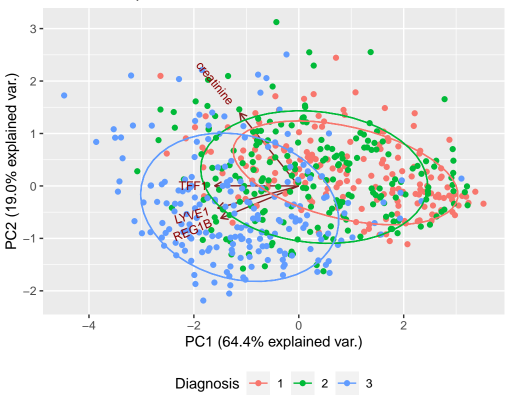
plasma_CA19_{

A19_9 atinine LYVE1

The heatmap shows that there is not much correlation between creatinine and the other variables. The other outstanding one has to be the TFF1 biomarker, being the most correlated variable to others.

4.3 PCA

PCA of complete dataset



While the control and benign group show relative distance from the PDAC group, there is a still a lot of overlapping samples with the benign and PDAC groups. As earlier concluded from the heatmap, the creatinine biomarker does not show much relativeness with the other biomarkers. LYVE1 is nicely in between the TFF1 and REG1B biomarkers. Every point close tho the origin have values close to the mean for all variables.

5 Machine Learning

5.1 Quality Metrics

Accuracy is the most important quality metric to measure the performance of an algorithm. Though it is easy to choose an algorithm this way, there are always multiple algorithms one can take as final option. Hence, other quality metrics have to be taken in account to make the optimal choice.

For this project, the model is a finished product, trained and tested with the already collected data. New samples for this model are manually inserted by the acting physician, thus speed is not a relevant metric. Naturally, it is of more importance a patient with a malignant cancer should not be classified as benign, rather than a patient with a benign case being classified as malignant. These errors can be visualized in a confusion matrix, which almost all algorithms output.

5.1.1 Confusion Matrix

A standard confusion matrix is a 2x2 matrix which shows all the correct hits and rejections, and errors. In Weka, a confusion matrix looks a bit like this:

Table 9: Example of a confusion matrix

$$\begin{array}{c|ccc} a & b & <\text{-classified as} \\ \hline TP & FP & a = Malignant \\ FN & TN & b = Benign \\ \end{array}$$

In this example, the correctly classified malignant instances are true positive (TP) and correctly classified benign instances are true negative (TN). The benign instances that were classified as malignant are false positive (FP) and malignant instances that were classified as benign are false negative (FN).

The example shows a situation where the algorithm can choose between two classes. In this project, there are 4 different classes, so what does a confusion matrix like that look like? Since there is no one TP or one FN, the different values are given per class. To get the confusion matrix of one class, the chosen class is the hit instance (positive), whereas all the other classes are the rejections (negatives).

The confusion matrices for this project will look a bit like this:

Table 10: Example of a project confusion matrix

| \mathbf{a} | b | \mathbf{c} | d | <- classified as |
|--------------|-------------|--------------|-------------|------------------|
| hit | error | error | error | a = Control |
| error | $_{ m hit}$ | error | error | b = Benign |
| error | error | $_{ m hit}$ | error | c = I-II |
| error | error | error | $_{ m hit}$ | d = III-IV |

5.1.2 Sensitivity and specificity

Generally important for machine learning algorithms, but also for this project: Sensitivity and specificity. Also known as the true positive rate (TPR), sensitivity is calculated as $\frac{TP}{TP+FN}$ whereas specificity - or the true negative rate (TNR) - is calculated as $\frac{TN}{TN+FP}$.

5.1.3 Area under ROC

A receiver operating characteristic curve, or ROC curve, is a curve when the TPR is plotted against the false positive rate (FPR). The curve describes how well the algorithm classified and can be manipulated with various cutoffs.

why important metric?

5.2 Weka: Model exploration

For the exploration of the model, the data is cleaned it contains only the biomarkers and the classification labels (Control, Benign, I-II, and III-IV).

```
# Set working directory to this folder
setwd("../data")
# Read dataset
dataset <- read.csv("../data/Data.csv")</pre>
# Change the empty strings to NA
dataset[dataset == ""] <- NA</pre>
# Group the samples
dataset$diagnosis <- factor(dataset$diagnosis, levels = unique(dataset$diagnosis),</pre>
                              labels = c("Control", "Benign", "PDAC"))
dataset$sex <- factor(dataset$sex)</pre>
# Drop unnecessary columns
drop <- c("sample_id", "patient_cohort", "sample_origin", "benign_sample_diagnosis",</pre>
          "REG1A", "stage")
dataset <- dataset[,!(names(dataset) %in% drop)]</pre>
# Move diagnosis to last column
dataset <- dataset %>% select(-3, everything())
# Log transform and meann centering
log.data <- log(dataset[3:7] +1)</pre>
dataset[3:7] <- log.data</pre>
```

To set a baseline, the data is run through different types of algorithms in Weka with 10-fold cross validation:

Table 11: Algorithm comparison: '*' = significantly worse; 'v' = significantly better

| Algorithm | Accuracy | Sensitivity | Specificity | AUROC |
|------------|----------|-------------|-------------|--------|
| ZeroR | 35.25 | 0 | 1 | 0.5 |
| OneR | 49.51 | 0.4612 | 0.7953 | 0.6283 |
| NaiveBayes | 60.07 | 0.5645 | 0.8017 | 0.8165 |
| Logistic | 65.2 | 0.528 | 0.8449 | 0.8173 |

| Algorithm | Accuracy | Sensitivity | Specificity | AUROC |
|----------------------|----------|-------------|-------------|--------|
| SimpleLogistic | 64.31 | 0.5296 | 0.8371 | 0.814 |
| SMO | 62.81 | 0.4258 | 0.8835 | 0.7721 |
| IBk | 52.37 | 0.3905 | 0.8915 | 0.641 |
| J48 | 59.08 | 0.5536 | 0.8065 | 0.7674 |
| RandomForest | 65.61 | 0.6581 | 0.8399 | 0.8502 |

With the data/base.exp file, all the options used for this run can be imported into Weka. Add the desired data and fill in the Results Destination for the run.

These results show a relative low accuracy and sensitivity. Some algorithms also have a low ROC value, putting the cutoff at 0.8: OneR, SMO, IBk and J48 will not be used. As for the remaining three: NaiveBayes has by far the lowest accuracy of them and is therefor also dropped. Leaving the options Logistic and RandomForest. Since earlier shown there is a linear correlation between the different variables, the Logistic algorithm would be more fitting for this type of data.

5.2.1 Data imbalance

To prepare the data for the optimization, the data needs to be split into groups: one file containing Control and PDAC samples, another file containing Benign and PDAC samples. These are each split up into a train and test set.

```
# Random split for training and test sets (50/50)
set.seed(391)
train.rows <- sort(sample(seq_len(nrow(dataset))), size = floor(0.7*nrow(dataset))))
training <- dataset[train.rows,]</pre>
test <- dataset[-train.rows,]</pre>
control.train <- subset(training, training$diagnosis == "Control" | training$diagnosis == "PDAC")</pre>
benign.train <- subset(training, training$diagnosis == "Benign" | training$diagnosis == "PDAC")
control.test <- subset(test, test$diagnosis == "Control" | training$diagnosis == "PDAC")</pre>
benign.test <- subset(test, test$diagnosis == "Benign" | training$diagnosis == "PDAC")
# Export dataset
write.csv(dataset, "../data/cleaned_data.csv", row.names = F, quote = F, na="")
write.csv(control.train, "../data/control_train.csv", row.names = F, quote = F, na="")
write.csv(control.test, "../data/control_test.csv", row.names = F, quote = F, na="")
write.csv(benign.train, "../data/benign_train.csv", row.names = F, quote = F, na="")
write.csv(benign.test, "../data/benign_test.csv", row.names = F, quote = F, na="")
# Set working directory back to log folder
setwd("../log")
```

5.2.2 Optimize algorithm

For the optimization of the algorithm, Weka's ThresholdSelector classifier will be used. The algorithm supplies a threshold on the probability output of the given classifier. The options for the ThresholdSelector looks like the following: