Urinary Biomarkers for Pancreatic Cancer

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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:

- ggplot2
- tidyr
- dplyr
- readr

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	Name Full Name		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
TFF1	$\mathrm{TFF}1$	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

Name	Description
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
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
${ m TFF1}$	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.

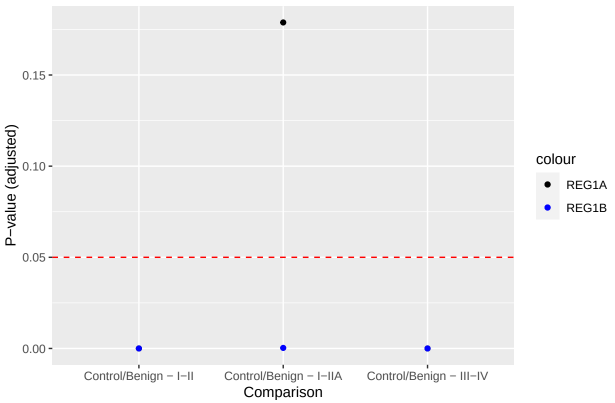
```
# Change the empty strings to NA
dataset[dataset == ""] <- NA</pre>
# Remove unnecessary columns
drop <- c("sample_id", "patient_cohort", "sample_origin", "benign_sample_diagnosis")</pre>
dataset <- dataset[,!(names(dataset) %in% drop)]</pre>
# Group the samples
dataset <- dataset %>%
 mutate(
    ## Factor for order of age
    diagnosis_group = factor(
      dplyr::case_when(
        diagnosis == 1 ~ "Control/Benign",
        diagnosis == 2 ~ "Control/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")
 )
```

3.1 REG1A vs. REG1B

```
REG1A <- dunnTest(dataset$REG1A ~ dataset$diagnosis_group)
REG1B <- dunnTest(dataset$REG1B ~ dataset$diagnosis_group)

ggplot(REG1A$res[c(1,2,4),], aes(x = Comparison, y = P.adj)) +
    geom_point(aes(color = "REG1A")) +
    geom_point(data = REG1B$res[c(1,2,4),], aes(color = "REG1B")) +
    geom_hline(yintercept = 0.05, color = "red", linetype = "dashed") +
    labs(y = "P-value (adjusted)", title = "Comparison REG1A and REG1B") +
    scale_color_manual(values = c("black", "blue"), limits = c("REG1A", "REG1B"))</pre>
```

Comparison REG1A and REG1B



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

Using a Kruskal-Wallis test with Dunn's multiple comparisons, REG1B shows clearly a higher performance than REG1A when the control/benign samples were compared with the stage I-IIA PDAC samples. Although very similar in performance, REG1B was used further on and therefor, column 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 3: 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	${\rm diagnosis_group}$
Min. :0.05655	Min.: 0.000129	Min.: 0.0011	Min.: 0.005	Control/Benign:391
1st Qu.:0.37323	1st Qu.: 0.167179	1st Qu.: 10.7572	1st Qu.: 43.961	I-IIA: 27
Median: 0.72384	Median: 1.649862	Median: 34.3034	Median: 259.874	I-II: 75
Mean $:0.85538$	Mean: 3.063530	Mean: 111.7741	Mean: 597.869	III-IV:97
3rd Qu.:1.13948	3rd Qu.: 5.205037	3rd Qu.: 122.7410	3rd Qu.: 742.736	NA
Max. :4.11684	Max. :23.890323	Max. :1403.8976	Max. :13344.300	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 quicker access of the different samples. Table 5 shows the different amounts of samples per diagnosis and the amount of which are also blood samples. After the blood samples are seperated the column can be dropped.

```
# Different diagnosis and blood groups
control <- subset(dataset, diagnosis == 1)</pre>
benign <- subset(dataset, diagnosis == 2)</pre>
pdac <- subset(dataset, diagnosis == 3)</pre>
blood <- subset(dataset, plasma_CA19_9 >= 0)
# Drop the "plasma" column
dataset <- dataset[,-5]</pre>
# Demographics
demograph <- data.frame(c(sum(control$sex == "F"), sum(control$sex == "M")),</pre>
                         c(sum(benign$sex == "F"), sum(benign$sex == "M")),
                         c(sum(pdac$sex == "F"), sum(pdac$sex == "M")))
blood.demo <- data.frame(c(sum(blood$sex == "F" & blood$diagnosis == 1),
                            sum(blood$sex == "M" & blood$diagnosis == 1)),
                          c(sum(blood$sex == "F" & blood$diagnosis == 2),
                            sum(blood$sex == "M" & blood$diagnosis == 2)),
                          c(sum(blood$sex == "F" & blood$diagnosis == 3),
                            sum(blood$sex == "M" & blood$diagnosis == 3)))
colnames(blood.demo) <- c("Control", "Benign", "PDAC")</pre>
colnames(demograph) <- c("Control", "Benign", "PDAC")</pre>
demograph <- rbind(demograph, blood.demo)</pre>
rownames(demograph) <- c("Female total", "Male total", "Female blood", "Male blood")</pre>
pander(demograph, booktabs = T, caption = "Demographic of the samples",
       justify = c("left", "center", "center", "center"))
```

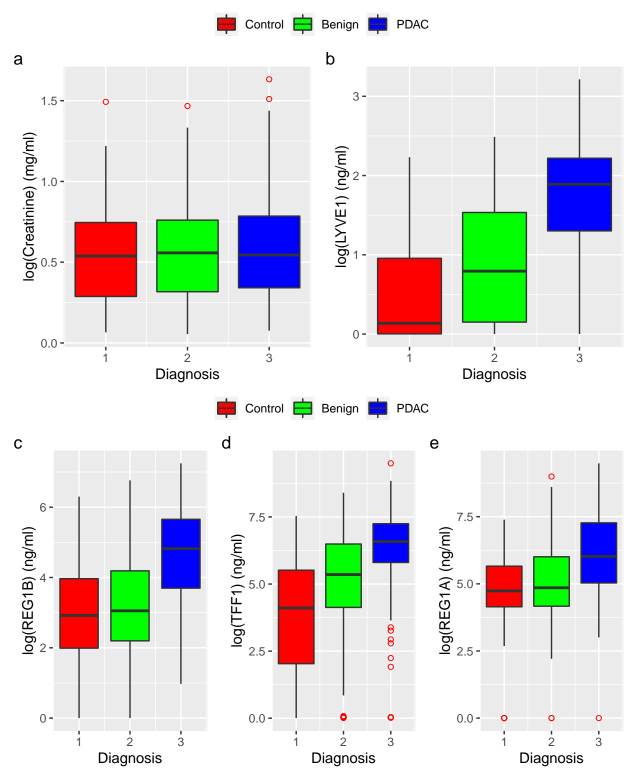
Table 5: Demographic of the samples

	Control	Benign	PDAC
Female total	115	101	83
Male total	68	107	116
Female blood	58	57	64
Male blood	34	51	86

4 Analyse the data

4.1 Boxplots

```
# Boxplot function
create.plots <- function(y.values, y.label, plt.tag) {</pre>
  list(ggplot(data = control, aes(x = diagnosis, y = !!sym(y.values))) +
    geom_boxplot(outlier.color = "red", outlier.shape = 1, aes(fill = "Control")) +
    geom_boxplot(data = benign, outlier.color = "red", outlier.shape = 1,
                 aes(fill = "Benign")) +
    geom_boxplot(data = pdac, outlier.color = "red", outlier.shape = 1,
                 aes(fill = "PDAC")) +
    labs(x = "Diagnosis", y = y.label, tag = plt.tag) +
    scale_fill_manual(values = c("red", "green", "blue"),
                      limits = c("Control", "Benign", "PDAC"),
                      name = ""))
}
# Create the boxplots for the different columns
y.values <- names(dataset[5:9])</pre>
y.labs <- c("log(Creatinine) (mg/ml)", "log(LYVE1) (ng/ml)", "log(REG1B) (ng/ml)",
            "log(TFF1) (ng/ml)", "log(REG1A) (ng/ml)")
plt.tag <- c("a", "b", "c", "d", "e")
plts <- mapply(create.plots, y.values, y.labs, plt.tag)</pre>
# Grid and print the plots
p1 <- ggarrange(plotlist = plts[1:2], ncol = 2,
                common.legend = TRUE, legend = "top")
p2 <- ggarrange(plotlist = plts[3:5], ncol = 3,</pre>
                common.legend = TRUE, legend = "top")
my.grid <- ggarrange(p1, p2, nrow = 2)</pre>
print(annotate_figure(my.grid))
```

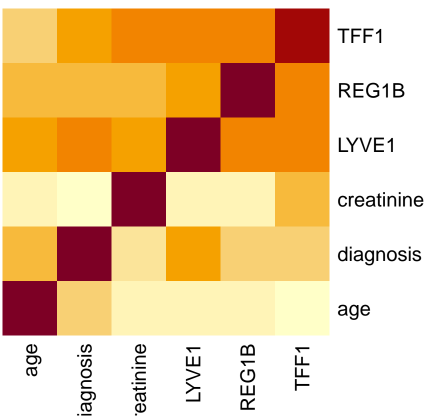


The outliers are not localized in a specific diagnosis group, but rather spread over the groups.

4.2 Correlation matrix

```
cor_matrix <- cor(dataset[,c(1, 3, 5:8)])
heatmap(cor_matrix, scale = "column", Colv = NA, Rowv = NA, main = "Correlation matrix")</pre>
```

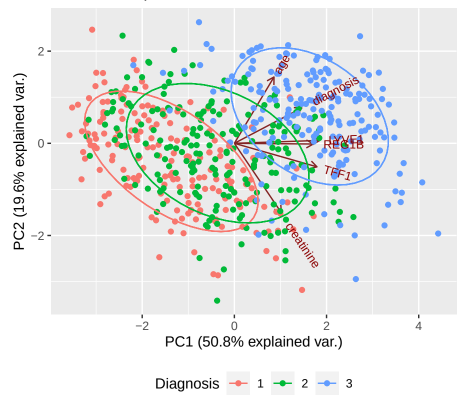
Correlation matrix



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.

4.3 PCA

PCA of complete dataset



The PCA shows that there is a clustering on the right upper side of the PDAC diagnosis, while the control and benign groups can be considered the same cluster. It is also very clear that creatinine has no correlation - as shown in the previous heatmap - but TFF1 does not seem as close to LYVE1 and REG1B as predicited. In fact, the latter two have a higher correlation with each other. Every point close to the origin have values close to the mean for all variables.