## R code Faraz

Faraz Younus 2023-05-28

## R Markdown

```
attach(df)
summary(df)
```

```
##
                         gender
                                         N/a_col
        cancer
                                                            st1
##
    Min.
           :1.000
                             :1.000
                                      Min.
                                             :38.00
                                                              :
                                                                 20.0
                     Min.
                                                       Min.
    1st Qu.:2.000
##
                     1st Qu.:1.000
                                      1st Qu.:56.50
                                                       1st Qu.: 119.5
##
    Median :3.000
                     Median :1.000
                                      Median :66.00
                                                       Median : 245.0
##
   Mean
           :2.968
                     Mean
                            :1.381
                                      Mean
                                                              : 475.6
                                             :64.08
                                                       Mean
                                                       3rd Qu.: 452.5
##
    3rd Qu.:4.000
                     3rd Qu.:2.000
                                      3rd Qu.:72.50
##
    Max.
           :6.000
                     Max.
                            :2.000
                                      Max.
                                             :93.00
                                                       Max.
                                                              :4288.0
##
         st2
                           st3
                                             st4
##
    Min.
           : 18.0
                      Min.
                                 8.0
                                        Min.
                                               :10.00
    1st Qu.: 109.5
                      1st Qu.:
                                50.0
                                        1st Qu.:18.50
##
                                        Median :30.00
##
    Median : 199.0
                      Median : 124.0
##
    Mean
           : 270.1
                             : 200.4
                                        Mean
                                               :33.97
                      Mean
##
    3rd Qu.: 392.0
                      3rd Qu.: 260.0
                                        3rd Qu.:41.00
##
    Max.
           :1056.0
                      Max.
                             :1267.0
                                        Max.
                                               :91.00
```

## **Including Plots**

You can also embed plots, for example: Below we can see that the distribution of many varaibles isn't normal.

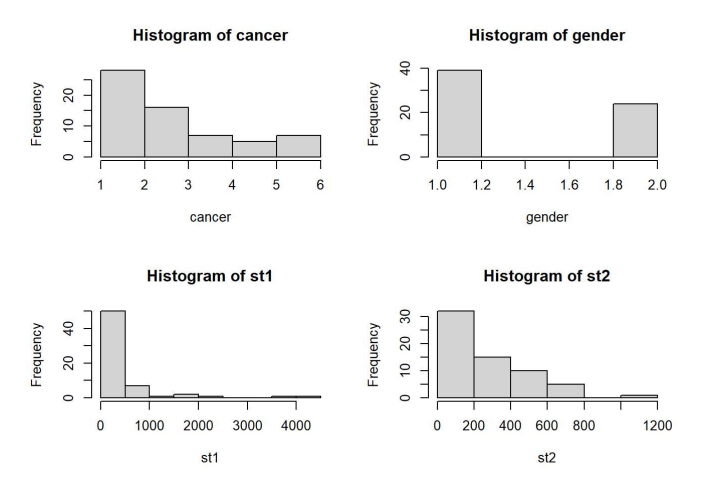
MANOVA in R can be used to determine whether there are significant differences between multiple groups on multiple dependent variables. However, it does not tell you which groups differ from the rest. This can be determined using a post-hoc test, such as Linear Discriminant Analysis (LDA).

MANOVA has several assumptions, including:

Multivariate normality: Each combination of independent and dependent variables should have a multivariate normal distribution. This can be verified using Shapiro-Wilk's test. Linearity: Dependent variables should have a linear relationship with each group (factor) of the independent variable. No multicollinearity: Dependent variables should not have very high correlations. No outliers: There should not be any outliers in the dependent variables.

This isn't important in our case.

```
par(mfrow = c(2, 2))
# This code allows use to get histograms of the distributions of different variables
hist(cancer)
hist(gender)
hist(st1)
hist(st2)
```



Note that the echo = FALSE parameter was added to the code chunk to prevent printing of the R code that generated the plot. To conduct a manova in R we can simply



```
# Create a factor variable for the independent variable
group <- factor(cancer)

#group<- levels(factor)

# Perform the MANOVA
result<-manova(cbind(st1, st2, st3, st4) ~ group, data = df)

# Print the MANOVA results
summary(result)</pre>
```

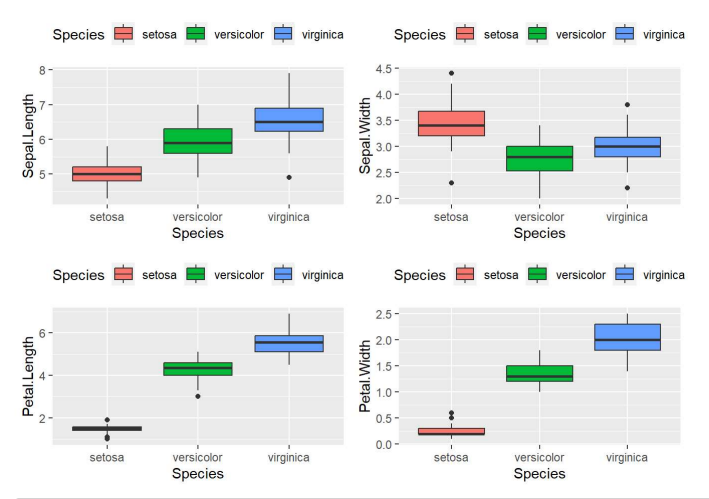
```
## Df Pillai approx F num Df den Df Pr(>F)
## group 5 0.85343 3.092 20 228 2.189e-05 ***
## Residuals 57
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Pillai: The Pillai's trace statistic is a multivariate test statistic used in MANOVA. It measures the overall effect of the grouping variable on the dependent variables. The reported Pillai's trace value is 0.85343.

Pr(>F): This column displays the p-value associated with the approximate F-test. It indicates the probability of observing the obtained F-statistic (or a more extreme value) under the null hypothesis of no group effect. In this case, the p-value is 2.189e-05, which is very small, suggesting strong evidence against the null hypothesis.

Overall, the MANOVA results suggest that there is a significant multivariate effect of the grouping variable on the dependent variables (st1, st2, st3, st4).

```
# Perform MANOVA
library(ggplot2)
library(gridExtra)
box s1 <- ggplot(iris, aes(x = Species, y = Sepal.Length, fill = Species)) +</pre>
  geom_boxplot() +
  theme(legend.position = "top")
box_sw \leftarrow ggplot(iris, aes(x = Species, y = Sepal.Width, fill = Species)) +
  geom boxplot() +
  theme(legend.position = "top")
box pl <- ggplot(iris, aes(x = Species, y = Petal.Length, fill = Species)) +</pre>
  geom_boxplot() +
  theme(legend.position = "top")
box_pw <- ggplot(iris, aes(x = Species, y = Petal.Width, fill = Species)) +</pre>
  geom boxplot() +
  theme(legend.position = "top")
grid.arrange(box sl, box sw, box pl, box pw, ncol = 2, nrow = 2)
```



dependent\_vars <- cbind(iris\$Sepal.Length, iris\$Sepal.Width, iris\$Petal.Length, iris\$Petal.Widt
h)
independent\_var <- iris\$Species

manova\_model <- manova(dependent\_vars ~ independent\_var, data = iris)
summary(manova\_model)</pre>

```
## Df Pillai approx F num Df den Df Pr(>F)
## independent_var 2 1.1919 53.466 8 290 < 2.2e-16 ***

## Residuals 147

## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

library(effectsize)

eta\_squared(manova\_model)

Parameter <chr></chr>	<b>Eta2_partial</b>	CI	CI_low	CI_high
	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>
independent_var	0.5959494	0.95	0.5363619	1

1 row

```
library(MASS)

iris_lda <- lda(independent_var ~ dependent_vars, CV = F)
iris_lda</pre>
```

```
## Call:
## lda(independent_var ~ dependent_vars, CV = F)
##
## Prior probabilities of groups:
##
       setosa versicolor virginica
   0.3333333 0.3333333 0.3333333
##
##
## Group means:
##
              dependent_vars1 dependent_vars2 dependent_vars3 dependent_vars4
## setosa
                        5.006
                                        3.428
                                                        1.462
                                                                        0.246
## versicolor
                        5.936
                                        2.770
                                                        4.260
                                                                        1.326
## virginica
                                        2.974
                                                                        2.026
                        6.588
                                                        5.552
##
## Coefficients of linear discriminants:
##
                          LD1
## dependent vars1 0.8293776 0.02410215
## dependent vars2 1.5344731 2.16452123
## dependent vars3 -2.2012117 -0.93192121
## dependent_vars4 -2.8104603 2.83918785
##
## Proportion of trace:
##
      LD1
             LD2
## 0.9912 0.0088
```

```
LDA_Dataframe <- data.frame(
   species = iris[, "Species"],
   lda = predict(iris_lda)$x
)
head(LDA_Dataframe)</pre>
```

	species	lda.LD1	Ida.LD2
	<fct></fct>	<dbl></dbl>	<dbl></dbl>
1	setosa	8.061800	0.3004206
2	setosa	7.128688	-0.7866604
3	setosa	7.489828	-0.2653845
4	setosa	6.813201	-0.6706311
5	setosa	8.132309	0.5144625

<dbl></dbl>
1.4617210
1.46172

```
ggplot(LDA_Dataframe) +
  geom_point(aes(x = lda.LD1, y = lda.LD2, color = species), size = 4) +
  theme_classic()
```

