Multivariate Statistical Methods

Assignment 2

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Question 1

```
data <- read.table("../data/T1-9.dat")
names(data) <- c("country", "100m", "200m", "400m", "800m", "1500m", "3000m", "marathon")
numeric_data <- data[, -1]

countries <- as.character(data$country)</pre>
```

a)

Consider again the data set containing National track records for women. In lab 1 we studied different distance measures between an observation and the mean vector. The most common multivariate residual is the Mahalanobis distance and we computed this distance for all 54 observations. a) The Mahalanobis distance has an approximate chi square distribution, if the data come from a multivariate normal distribution and the number of observations is fairly large. Use the approximate chi square distribution for testing each observation at significance level 0.1 %, and conclude which countries can be regarded as outliers.

```
X <- as.matrix(numeric_data)
means <- colMeans(X)
covariances <- cov(X)
X_central <- X - rep(1, nrow(X)) %*% t(means)

mdist_sq <- X_central %*% solve(covariances) %*% t(X_central)
country_mdist <- diag(mdist_sq)

significance_level <- 0.001
p <- ncol(X)
quantile <- qchisq(1 - significance_level, df=p)

outliers <- country_mdist > quantile
print("Outliers without correction")
#> [1] "Outliers without correction"
countries[outliers]
#> [1] "KORN" "PNG" "SAM"
```

Here are the countries that can be regarded as outliers since their Mahalanobis distance is greater than the critical set chi square distribution as 24.3218863. The countries are Cooks Island, North Korea, Papa New Guinea and Samoa Island.

No clue what the multiple-testing correction procedure refers to.

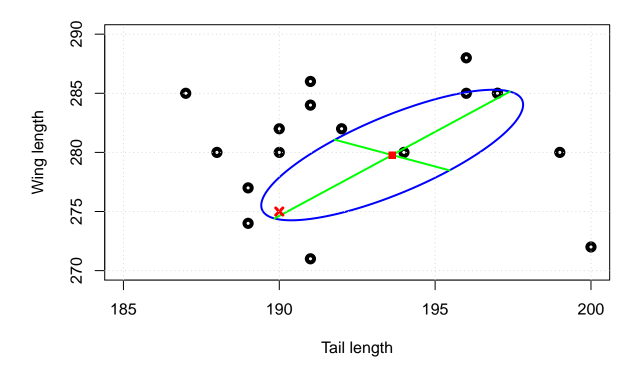
b)

The Mahalanobis takes the covariances into consideration so the distances lead to a elliptic decision boundary as opposed to the circular boundary by Euclidean distance. That indicates that North Korea is an outlier based on the covariances meaning their result do not follow the general trend.

Question 2

a)

```
bird <- read.table("../data/T5-12.DAT")</pre>
mu <- c(190, 275) #mus
x_bar <- colMeans(bird)</pre>
S <- cov(bird)
angles <- seq(0, 2 * pi, length.out=200) # make angles for circle
n <- nrow(bird)</pre>
p <- ncol(bird)</pre>
confidence level <- 0.05
\ensuremath{\mbox{\#\#}} eigenvalues and eigvactors from covariance matrix S
eigVal <- eigen(S)$values</pre>
eigVec <- eigen(S)$vectors</pre>
quantile <- qf(1 - confidence_level, df1=p, df2=n - p)
scale \leftarrow sqrt(eigVal * p * (n - 1) * quantile / (n * (n - p)))
scaled <- eigVec %*% diag(scale) # scale eigenvectors to length = square-root</pre>
        <- rbind(x_bar[1] + scaled[1, ], x_bar[1] - scaled[1, ])
        <- rbind(x_bar[2] + scaled[2, ], x_bar[2] - scaled[2, ])
ellBase <- cbind(scale[1]*cos(angles), scale[2]*sin(angles)) # making a circle base...
ellax <- eigVec %*% t(ellBase) # where the ellips axis goes through eigenvectors.
plot(bird, lwd="4", xlab="Tail length", ylab="Wing length", xlim=c(185, 200), ylim=c(270, 290))
lines((ellax + x_bar)[1, ], (ellax + x_bar)[2, ], asp=1, type="1", lwd=2, col="blue")
matlines(xMat, yMat, lty=1, lwd=2, col="green") #
points(mu[1], mu[2], pch=4, col="red", lwd=3)
grid()
points(mean(bird[,1]),mean(bird[,2]), type="p", col="red", pch=15)
```



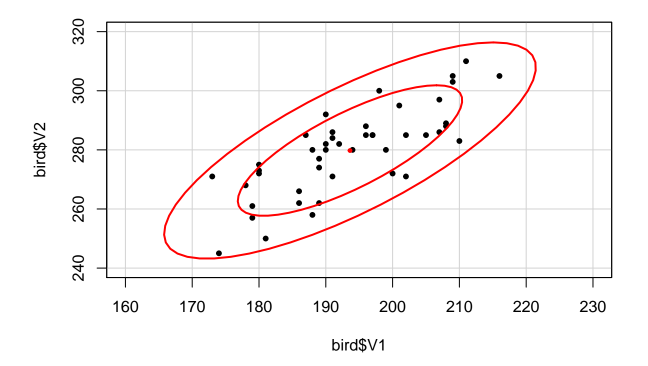
Since the male mean (red cross) is inside the confidence region we do not reject the hypothesis that males and females have the same mean.

b)

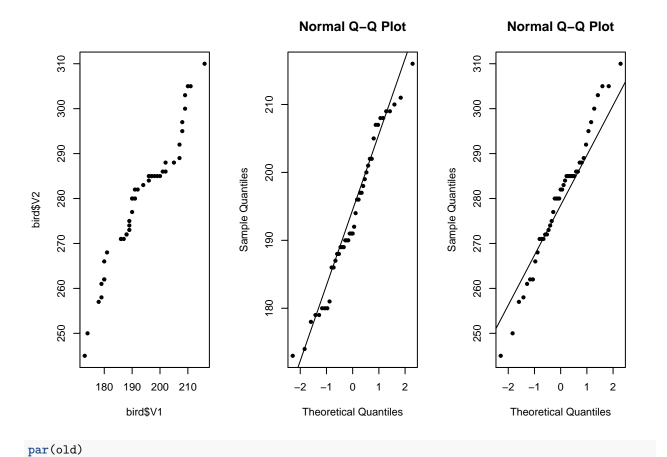
```
compute_tsq_intervals <- function(data, confidence=0.05) {</pre>
    n <- nrow(data)</pre>
    p <- ncol(data)</pre>
    x_bar <- colMeans(data)</pre>
    S <- cov(data)
    offset \leftarrow sqrt(p * (n - 1) * qf(1 - confidence, df1=p, df2=n - p) / (n - p) * diag(S) / n)
    rbind(x_bar - offset, x_bar + offset)
}
compute_bonferroni_intervals <- function(data, confidence=0.05) {</pre>
    n <- nrow(data)</pre>
    p <- ncol(data)</pre>
    x_bar <- colMeans(data)</pre>
    S <- cov(data)
    offset <- sqrt(diag(S) / n) * qt(1 - confidence / (2 * p), df=n - 1)
    rbind(x_bar - offset, x_bar + offset)
}
```

T-square test always gives wider confidence intervals since it takes the correlation between the measured variables into account. Bonferroni intervals are more precise if you are interested in the individual component means, but if you are interested in the overall data mean you should consider the T-square intervals.

c)



```
old <- par(mfrow=c(1, 3))
qqplot(bird$V1, bird$V2, pch=20)
qqnorm(bird$V1, pch=20)
qqline(bird$V1)
qqnorm(bird$V2, pch=20)
qqline(bird$V2)</pre>
```



A bivariate normal distribution would be a viable population model. The qqplots do not deviate to much from the straight lines and the scatter plot shows that the points could very well have been generated from a

bivariate normal distribution.

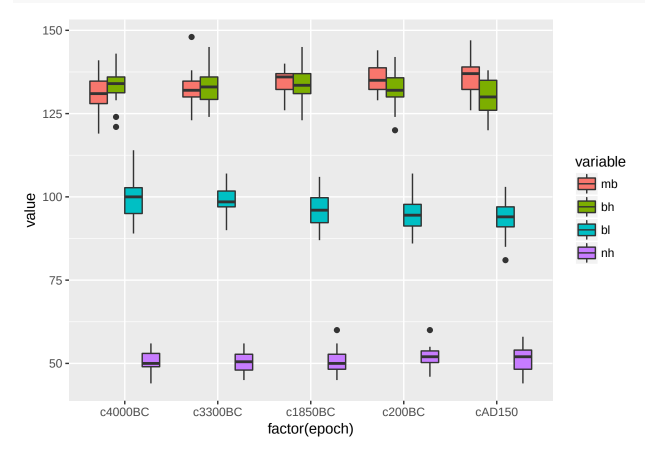
Question 3

```
library(heplots)
library(gplot2)
library(reshape2)

data <- Skulls
numeric_data <- data[, -1]
colors <- as.numeric(data$epoch)</pre>
```

a)

```
# pairs(numeric_data, col=colors)
mm <- melt(data, id="epoch")
ggplot(mm) +
    geom_boxplot(aes(x=factor(epoch), y=value, fill=variable))</pre>
```

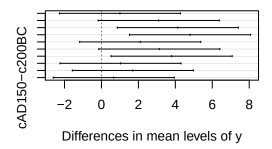


b)

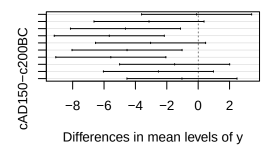
```
group_means <- data %>%
   group_by(epoch) %>%
   summarise_all(funs(mean(., na.rm=TRUE)))
print("Group means")
#> [1] "Group means"
group_means
#> # A tibble: 5 x 5
                       bh
#>
     epoch
                mb
                            bl
      <ord>
             <dbl>
                     <dbl>
                            <db1>
                                    <db1>
#> 1 c4000BC 131.3667 133.6000 99.16667 50.53333
#> 2 c3300BC 132.3667 132.7000 99.06667 50.23333
#> 3 c1850BC 134.4667 133.8000 96.03333 50.56667
#> 4 c200BC 135.5000 132.3000 94.53333 51.96667
#> 5 cAD150 136.1667 130.3333 93.50000 51.36667
fit <- manova(cbind(mb, bh, bl, nh) ~ data$epoch, data)
summary.aov(fit)
#> Response mb :
            Df Sum Sq Mean Sq F value
                                      Pr(>F)
#> Residuals 145 3061.07 21.111
#> ---
#> Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
#>
#> Response bh :
            Df Sum Sq Mean Sq F value Pr(>F)
#> Residuals 145 3405.3 23.485
#> ---
#> Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
#>
#> Response bl :
            Df Sum Sq Mean Sq F value
                                    Pr(>F)
#>
#> Residuals 145 3506.0 24.179
#> Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
#>
#> Response nh :
            Df Sum Sq Mean Sq F value Pr(>F)
            4 61.2 15.300 1.507 0.2032
#> data$epoch
#> Residuals 145 1472.1 10.153
print("T-square Confidence Intervals")
#> [1] "T-square Confidence Intervals"
compute_tsq_intervals(numeric_data)
        mb
              bh bl
#> [1,] 132.7147 131.2755 95.076 50.10776
#> [2,] 135.2320 133.8178 97.844 51.75890
```

```
X <- as.matrix(data[,2:5])</pre>
y <- as.factor(data[,1])
par(mfrow=c(2,2))
compareX1 = aov(X[,1] \sim y)
plot(TukeyHSD(compareX1))
TukeyHSD(compareX1)$y[2,]
          diff
                                               p adj
   3.10000000 -0.17713439 6.37713439 0.07323876
compareX2 = aov(X[,2] \sim y)
plot(TukeyHSD(compareX2))
compareX3 = aov(X[,3] \sim y)
plot(TukeyHSD(compareX3))
TukeyHSD(compareX3)$y[2,]
         diff
                      lwr
                                           p adj
#> -3.1333333 -6.6405432 0.3738765 0.1037609
compareX4 = aov(X[,4] \sim y)
plot(TukeyHSD(compareX4))
```

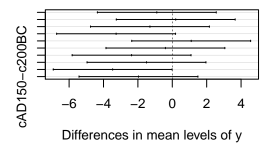
95% family-wise confidence level



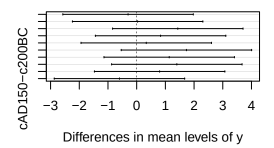
95% family-wise confidence level



95% family-wise confidence level



95% family-wise confidence level

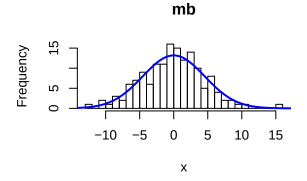


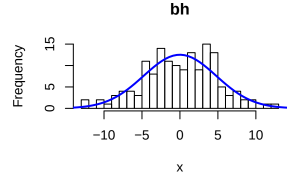
c)

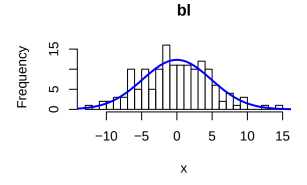
```
residuals <- fit$res
col_names <- c("mb", "bh", "bl", "nh")

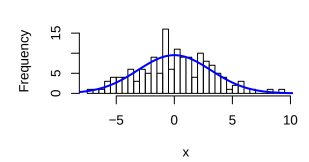
old <- par(mfrow=c(2, 2))

for (col in 1:ncol(residuals)) {
    x <- residuals[, col]
    main <- col_names[col]
    h <- hist(x, breaks=25, main=main)
    offset <- (max(x) - min(x)) / 2
    xfit <- seq(min(x) - offset, max(x) + offset, length = 100)
    yfit <- dnorm(xfit, mean = mean(x), sd = sd(x))
    yfit <- yfit * diff(h$mids[1:2]) * length(x)
    lines(xfit, yfit, col="blue", lwd=2)
}</pre>
```









nh

par(old)

Appendix

Code

```
# Question 1
data <- read.table("../data/T1-9.dat")</pre>
names(data) <- c("country", "100m", "200m", "400m", "800m", "1500m", "3000m", "marathon")</pre>
numeric_data <- data[, -1]</pre>
countries <- as.character(data$country)</pre>
X <- as.matrix(numeric_data)</pre>
means <- colMeans(X)</pre>
covariances <- cov(X)
X_central <- X - rep(1, nrow(X)) %*% t(means)</pre>
mdist_sq <- X_central %*% solve(covariances) %*% t(X_central)</pre>
country_mdist <- diag(mdist_sq)</pre>
significance_level <- 0.001
p <- ncol(X)
quantile <- qchisq(1 - significance_level, df=p)</pre>
outliers <- country_mdist > quantile
print("Outliers without correction")
countries[outliers]
# Question 2
bird <- read.table("../data/T5-12.DAT")</pre>
mu <- c(190, 275) #mus
x_bar <- colMeans(bird)</pre>
S <- cov(bird)
angles <- seq(0, 2 * pi, length.out=200) # make angles for circle
n <- nrow(bird)</pre>
p <- ncol(bird)
confidence_level <- 0.05</pre>
## eigenvalues and eigvactors from covariance matrix S
eigVal <- eigen(S)$values</pre>
eigVec <- eigen(S)$vectors</pre>
quantile <- qf(1 - confidence_level, df1=p, df2=n - p)
scale \leftarrow sqrt(eigVal * p * (n - 1) * quantile / (n * (n - p)))
scaled <- eigVec %*% diag(scale) # scale eigenvectors to length = square-root
        <- rbind(x_bar[1] + scaled[1, ], x_bar[1] - scaled[1, ])
xMat
        <- rbind(x_bar[2] + scaled[2, ], x_bar[2] - scaled[2, ])
ellBase <- cbind(scale[1]*cos(angles), scale[2]*sin(angles)) # making a circle base...
```

```
ellax <- eigVec %*% t(ellBase) # where the ellips axis goes through eigenvectors.
plot(bird, lwd="4", xlab="Tail length", ylab="Wing length", xlim=c(185, 200), ylim=c(270, 290))
lines((ellax + x_bar)[1, ], (ellax + x_bar)[2, ], asp=1, type="1", lwd=2, col="blue")
matlines(xMat, yMat, lty=1, lwd=2, col="green") #
points(mu[1], mu[2], pch=4, col="red", lwd=3)
points(mean(bird[,1]),mean(bird[,2]), type="p", col="red", pch=15)
compute tsq intervals <- function(data, confidence=0.05) {</pre>
    n <- nrow(data)</pre>
    p <- ncol(data)</pre>
    x_bar <- colMeans(data)</pre>
    S <- cov(data)
    offset \leftarrow sqrt(p * (n - 1) * qf(1 - confidence, df1=p, df2=n - p) / (n - p) * diag(S) / n)
    rbind(x_bar - offset, x_bar + offset)
}
compute_bonferroni_intervals <- function(data, confidence=0.05) {</pre>
   n <- nrow(data)</pre>
    p <- ncol(data)</pre>
   x_bar <- colMeans(data)</pre>
    S <- cov(data)
    offset <- sqrt(diag(S) / n) * qt(1 - confidence / (2 * p), df=n - 1)
    rbind(x_bar - offset, x_bar + offset)
}
print("T-square Confidence Intervals")
compute_tsq_intervals(bird, confidence_level)
print("Bonferroni Confidence Intervals")
compute_bonferroni_intervals(bird, confidence_level)
# Question 3
library(heplots)
library(dplyr)
library(ggplot2)
library(reshape2)
data <- Skulls
numeric_data <- data[, -1]</pre>
colors <- as.numeric(data$epoch)</pre>
# pairs(numeric_data, col=colors)
mm <- melt(data, id="epoch")</pre>
ggplot(mm) +
    geom_boxplot(aes(x=factor(epoch), y=value, fill=variable))
group_means <- data %>%
    group_by(epoch) %>%
    summarise_all(funs(mean(., na.rm=TRUE)))
print("Group means")
group_means
```

```
fit <- manova(cbind(mb, bh, bl, nh) ~ data$epoch, data)
summary.aov(fit)

print("T-square Confidence Intervals")
compute_tsq_intervals(numeric_data)</pre>
```