# Lab 2 732A97 Multivariate Statistical Methods

Raymond Sseguya 2019-11-29

#### Inference about mean vectors

## Question 1: Test of outliers

Consider again the data set from the T1-9.dat file, National track records for women. In the first assignment we studied different distance measures between an observation and the sample average vector. The most common multivariate residual is the Mahalanobis distance and we computed this distance for all observations.

a) The Mahalanobis distance is approximately chi-square distributed, if the data comes from a multivariate normal distribution and the number of observations is large. Use this chi-square approximation for testing each observation at the 0.1% significance level and conclude which countries can be regarded as outliers. Should you use a multiple-testing correction procedure? Compare the results with and without one. Why is (or maybe is not) 0.1% a sensible significance level for this task?

```
trackrcs <- read.table("T1-9.dat",</pre>
  col.names = c("countries", "x100m", "x200m",
        "x400m", "x800m", "x1500m", "x3000m", "marathon"))
trackrcs2 <- (trackrcs)[,-1]
rownames(trackrcs2) <- trackrcs[,1]</pre>
C <- cov((trackrcs)[,-1])</pre>
x_bar = apply(trackrcs2,1,mean)
d0 = as.matrix(trackrcs2-x_bar)
deviation = sqrt(d0%*%t(d0))
d_{sq_m} \leftarrow d0\%*\%solve(C)\%*\%t(d0)
diagonal_vector3 <- diag(d_sq_m)</pre>
deviation countries3 <-
  cbind.data.frame(countries = as.vector(trackrcs[,1]),diagonal_vector3)
deviation countries ordered3 <-
   deviation_countries3[order(-deviation_countries3$diagonal_vector3), ]
named_Mahalanobis <- as.vector(deviation_countries_ordered3[,2])</pre>
names(named_Mahalanobis) <- rownames(deviation_countries_ordered3)</pre>
ch s <- combn(x=named Mahalanobis, m=2,
    FUN = function(c){
```

```
sg <<- 0.1/100
      pv <- chisq.test(x=c, p=rep(sg,2), rescale.p = TRUE)$p.value</pre>
      pvname <- paste0(names(c)[1]," ",names(c)[2])</pre>
      assign(pvname, pv)
      return(list(pv, pvname, xx=names(c)[1]))
outliers_unsorted <- unlist(ch_s[3, which(ch_s[1,] == 0)])
summary(as.data.frame.character(outliers_unsorted), maxsum = 10)
    outliers unsorted
##
##
    COK
          : 52
##
  PNG
           : 52
## SAM
           : 51
## GUA
           : 49
##
  BER
          : 48
##
  MR.I
           : 44
##
  CRC
           : 43
```

### print(deviation\_countries\_ordered3[1:5,])

```
countries diagonal_vector3
##
## COK
             COK
                           2094451
## PNG
             PNG
                           2083382
## SAM
             SAM
                           1793652
## GUA
             GUA
                           1485602
## BER
             BER
                           1472580
```

: 43

: 43

(Other):632

## DOM

##

## MAS

We can see from the summary that "COK", "PNG", "SAM", "GUA" and "BER" remain the top oultiers even with using the chi-square test.

#### possibly using a multiple-testing correction procedure

We can see that a multiple-testing correction procedure is clearly a bad idea as it does not actually pinpoint the actual outliers but it tells us only that there are indeed some outliers.

#### correct way

```
### the critical value
which(deviation_countries3[,2] > qchisq(p=0.1/100, df=54))

## [1] 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23
## [24] 24 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41 42 43 44 45 46
## [47] 47 48 49 50 51 52 53 54
```

b) One outlier is North Korea. This country is not an outlier with the Euclidean distance. Try to explain these seemingly contradictory results.

It seems in our results, North Korea is an outlier throughout.

## Question 2: Test, confidence region and confidence intervals for a mean vector

Look at the bird data in file T5-12.dat and solve Exercise 5:20 of Johnson, Wichern. Do not use any extra R package or built-in test but code all required matrix calculations. You MAY NOT use loops!

```
birds <- read.table("T5-12.DAT", col.names = c("taillength", "winglength") )</pre>
```

a) Find and sketch the 95% confidence ellipse for the population means  $\mu_1$  and  $\mu_2$ . Suppose it is known that  $\mu_1 = 190$  mm and  $\mu_2 = 275$  mm for male hook-billed kites. Are these plausible values for the mean tail length and mean wing length for the female birds? Explain.

```
mu0 < -c(190, 275)
S <- cov(birds); eigen(S)
## eigen() decomposition
## $values
## [1] 294.60898 34.62637
##
## $vectors
##
              [,1]
                         [,2]
## [1,] 0.5753739 -0.8178905
## [2,] 0.8178905 0.5753739
p <- ncol(birds); n <- nrow(birds)</pre>
alpha \leftarrow 0.05
c = sqrt(p*(n-1)/(n-p)*qf(p=(1-alpha), df1=p, df2=n-p))
(eigen(S)$values)*c/sqrt(n)
## [1] 112.64249 13.23924
```

b) Construct the simultaneous 95%  $T^2$ -intervals for  $\mu_1$  and  $\mu_2$  and the 95% Bonferroni intervals for  $\mu_1$  and  $\mu_2$ . Compare the two sets of intervals. What advantage, if any, do the  $T^2$ -intervals have over the Bonferroni intervals?

95%  $T^2$ -intervals

Yes. These are plausible values.

```
n <- nrow(birds)
# mu <- apply(birds, 2, mean)
# T_sq <- as.vector(n*t(mu-mu0)%*%solve(S)%*%(mu-mu0)); T_sq < c
c(mu0[1] - (sqrt(c)*sqrt(diag(S)[1]/n) ), mu0[1] + (sqrt(c)*sqrt(diag(S)[1]/n) ))
## taillength taillength
## 187.3772 192.6228</pre>
```

```
c(mu0[2] - (sqrt(c)*sqrt(diag(S)[2]/n) ), mu0[2] + (sqrt(c)*sqrt(diag(S)[2]/n) ))

## winglength winglength
## 271.5524 278.4476

95% Bonferroni intervals

t <- qt(p=( 1-(alpha/(2*p)) ), df=n-1)

c(mu0[1] - (t*sqrt(diag(S)[1]/n) ), mu0[1] + (t*sqrt(diag(S)[1]/n) ))

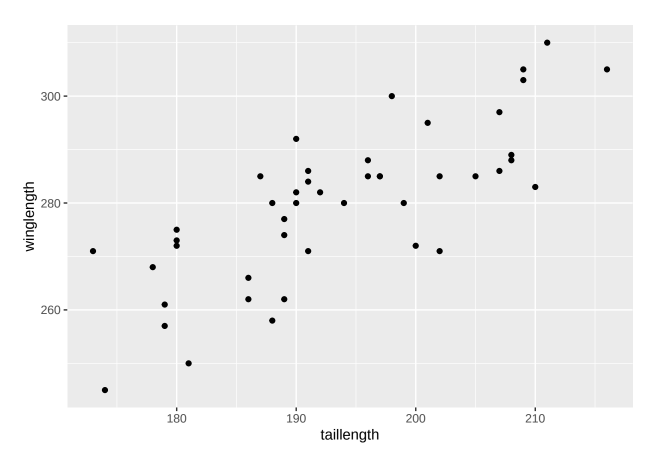
## taillength taillength
## 186.1993 193.8007

c(mu0[2] - (t*sqrt(diag(S)[2]/n) ), mu0[2] + (t*sqrt(diag(S)[2]/n) ))

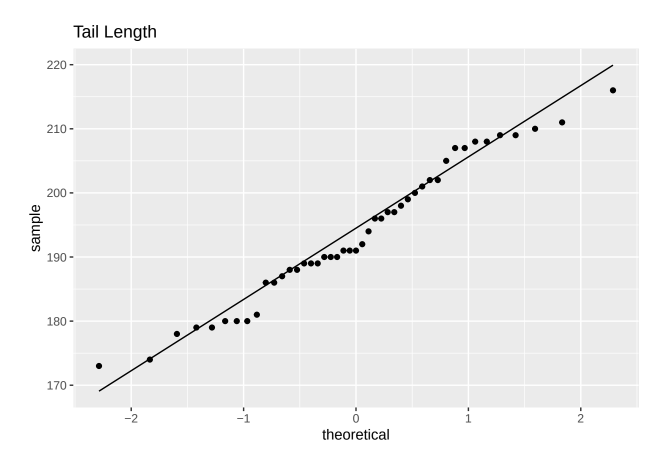
## winglength winglength
## 270.0041 279.9959</pre>
```

(c) Is the bivariate normal distribution a viable population model? Explain with reference to Q-Q plots and a scatter diagram.

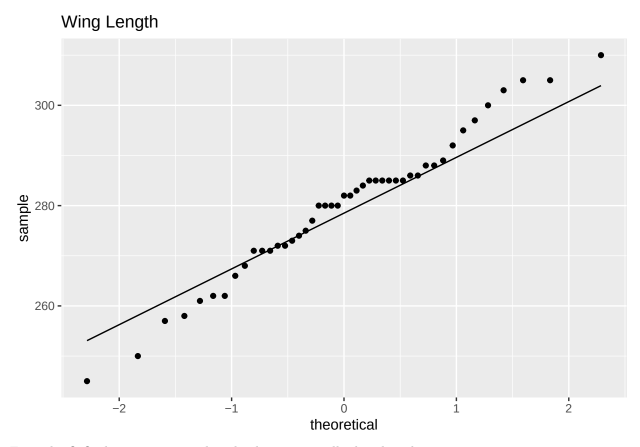
```
ggplot(data = birds)+aes(x=taillength, y=winglength)+
geom_point(stat = "identity")
```



```
ggplot(data = birds)+aes(sample=taillength)+stat_qq()+stat_qq_line()+
labs(title = "Tail Length")
```



```
ggplot(data = birds)+aes(sample=winglength)+stat_qq()+stat_qq_line()+
labs(title = "Wing Length")
```



From the Q-Q plots, we can say that the data is normally distributed.

# Question 3: Comparison of mean vectors (one-way MANOVA)

We will look at a data set on Egyptian skull measurements (published in 1905 and now in heplots R package as the object Skulls). Here observations are made from five epochs and on each object the maximum breadth (mb), basibregmatic height (bh), basialiveolar length (bl) and nasal height (nh) were measured.