Non-Parametric Statistics - Problem Sets

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Introduction

In the following document, the solutions to three problem sets from Prof. Eduardo García-Portugués book on nonparametric Statistics (https://bookdown.org/egarpor/NP-UC3M/) as final assignment for the Course in Nonparametric Statistics at Universidad Carlos III de Madrid. Contributions made to this document are made are equal among each of the three authors

```
# general libraries
library(ggplot2)
library(np)
library(dplyr)
```

Excercise 4.21

Consider the data(sunspots_births, package = "rotasym") dataset. This dataset contains recorded sunspots births during 1872–2018. The sunspots appear in groups.

The variables avaiable in the dataset are:

- date: UTC date, of the first observation of a group of sunspots.
- cycle: solar cycle in which the group of sunspots was observed.
- total_area: total whole spot area of the group, measured in millionths of the solar hemisphere.
- dist_sun_disc: distance from the center of Sun's disc, measured in units of the solar radius.
- theta: mean longitude angle $\theta \in [0, 2\pi)$ of the group position.
- **phi:** mean latitude angle $\phi \in [-\pi/2, \pi/2)$ of the group position.

a)

\$ theta

\$ phi

Filter the dataset to account only for the 23rd solar cycle.

: num 0.1012 -0.2164 0.1379 -0.0646 0.1134 ...

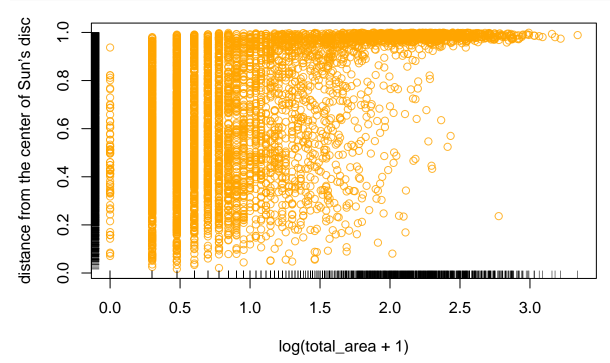
: num 1.98 5.17 5.79 4.27 2.99 ...

```
# Filter the dataset to account only for the 23rd solar cycle.
data_23 <- sunspots_births %>% filter(cycle==23)
```

b)

Inspect the graphical relation between dist $_$ sun $_$ disc (response) and log10(total $_$ area + 1) (predictor).

In the solar cycle 23, we see that for small group areas, there is not a clear relation with the distance from the center of the Sun's disc. But as the area is being wider, the distance achieve the length of the solar radius.



c)

Compute the CV bandwidth for the above regression, for the local linear estimator.

```
bw1 <- np::npregbw(formula = dist_sun_disc ~ log10(total_area + 1), data = data_23, regtype = "ll")
## Multistart 1 of 1 | Multistart 1 of 1
```

CV bandwidth for the LLE: 0.1392041

d)

Compute and plot the local linear estimator using CV bandwidth. Comment on the fit.

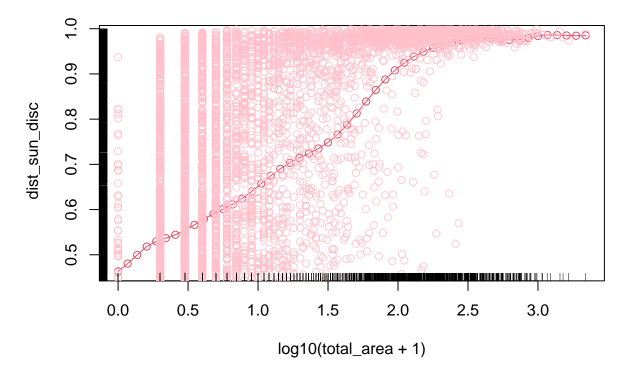
The local linear fit shows a higher slope for middle values of the area, and have a behavior near to a constant function with large values of the area. That means, that specially for large values of the area is doing a good job.

Also, as commented before, for small values of the area there is not an evident relation, so it would be difficult to find a fit that takes that into account.

```
kre1 <- np::npreg(bws = bw1)

# Plot
plot(kre1, col = 2, type = "o")
points(log10(data_23$total_area + 1), data_23$dist_sun_disc, col = adjustcolor("pink", alpha.f = 0.8))
rug(log10(data_23$total_area + 1), side = 1)
rug(data_23$dist_sun_disc, side = 2)</pre>
```

Warning in rug(data_23\$dist_sun_disc, side = 2): some values will be clipped



e)

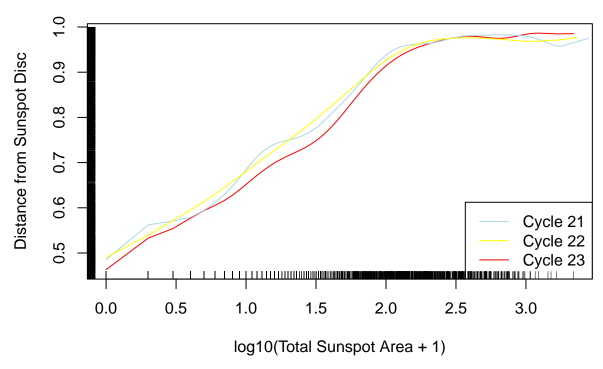
Plot the local linear fits for the 23rd, 22rd, and 21st solar cycles. Do they have a similar pattern?

```
data_21 <- sunspots_births %>% filter(cycle == 21)
data_22 <- sunspots_births %>% filter(cycle == 22)
```

```
bw1_21 <- np::npregbw(formula = dist_sun_disc ~ log10(total_area + 1),</pre>
                       data = data_21, regtype = "11")
bw1_22 <- np::npregbw(formula = dist_sun_disc ~ log10(total_area + 1),</pre>
                       data = data_22, regtype = "11")
kre1_21 <- np::npreg(bws = bw1_21)</pre>
kre1_22 <- np::npreg(bws = bw1_22)</pre>
kre1 21 %>% saveRDS("kre1 21.RDS")
kre1_22 %>% saveRDS("kre1_22.RDS")
kre1 21 <- readRDS("kre1 21.RDS")</pre>
kre1_22 <- readRDS("kre1_22.RDS")</pre>
# Extract x-values and fitted values for manual plotting
x 21 <- log10(data 21$total area + 1)
yhat_21 <- fitted(kre1_21)</pre>
x_22 <- log10(data_22$total_area + 1)</pre>
yhat_22 <- fitted(kre1_22)</pre>
x_23 <- log10(data_23$total_area + 1)</pre>
yhat_23 <- fitted(kre1)</pre>
ord_21 <- order(x_21)
ord_22 <- order(x_22)
ord_23 <- order(x_23)
# Base plot with Cycle 23
plot(x_23[ord_23], yhat_23[ord_23], type = "1", col = "red",
     xlab = "log10(Total Sunspot Area + 1)",
     ylab = "Distance from Sunspot Disc",
     main = "Local Linear Fits for Solar Cycles 21, 22, and 23")
# Add lines manually for Cycle 21 and 22
lines(x_21[ord_21], yhat_21[ord_21], col = "lightblue")
lines(x_22[ord_22], yhat_22[ord_22], col = "yellow")
# Add rug plots
rug(log10(data_23$total_area + 1), side = 1)
rug(data_23$dist_sun_disc, side = 2)
```

Warning in rug(data_23\$dist_sun_disc, side = 2): some values will be clipped

Local Linear Fits for Solar Cycles 21, 22, and 23



Yes, all of the three fits starts at 0 for those who does not have area, then have a steep slope and ends in a almost constant function.

f)

Randomly split the dataset for the 23rd cycle into a training sample (80%) and testing sample (20%). Repeat 10 times these random splits. Then, compare the average mean square errors of the predictions by the local constant and local linear estimators with CV bandwidths. Comment on the results.

```
library(tidymodels)
bw0_c23 <- numeric(10)
bw1_c23 <- numeric(10)

kre0_c23 <- list()
kre1_c23 <- list()

MSE0_c23 <- numeric(10)

MSE1_c23 <- numeric(10)

data_23
for (i in 1:10) {

# Train-Test Split
set.seed(934 + i)
sunspot_split <- initial_split(data_23, prop = 0.80, strata = dist_sun_disc)
training_sunspot <- training(sunspot_split)
testing_sunspot <- testing(sunspot_split)</pre>
```

```
# Obtain LCE and LLE CV bandwith
  bw0 <- np::npregbw(formula = dist_sun_disc ~ log10(total_area + 1),</pre>
                       data = training sunspot, regtype = "lc")
  bw1 <- np::npregbw(formula = dist_sun_disc ~ log10(total_area + 1),</pre>
                       data = training_sunspot, regtype = "11")
  # Store the bandwidths
  bw0 c23[i] \leftarrow bw0
  bw1_c23[i] \leftarrow bw1
  # Fit the np regression with the selected bandwidth
  kre0 <- np::npreg(bws = bw0)</pre>
  kre1 <- np::npreg(bws = bw1)</pre>
  # Store the kernel regression
  kre0_c23[[i]] <- kre0</pre>
  kre1_c23[[i]] <- kre1</pre>
  # Make predictions
  pred LC <- predict(kre0,</pre>
                      newdata = data.frame(total area = log10(testing sunspot$total area + 1)))
  pred_LL <- predict(kre1,</pre>
                      newdata = data.frame(total_area= log10(testing_sunspot$total_area + 1)))
  # Store the MSE
  MSEO_c23[i] <- mean((testing_sunspot$dist_sun_disc - pred_LC)^2)
  MSE1_c23[i] <- mean((testing_sunspot$dist_sun_disc - pred_LL)^2)</pre>
}
results <- data.frame(LC_MSE = MSEO_c23, LL_MSE = MSE1_c23)
results %>% saveRDS("MSE_LL_LC.RDS")
```

```
results <- readRDS("MSE LL LC.RDS")</pre>
colMeans(results)
```

```
LC_MSE
##
                  LL_MSE
## 0.08652581 0.08764039
```

The average of the mean square error of the local linear fit is slightly higher than the local constant fit. It can be related with the behavior for small values of the area, where no trend is is seen. # Excercises 5.10 Investigate the accuracy of the naive bootstrap confidence intervals implemented in np::npplot. To do so:

1. Simulate M = 500 samples of size n = 100 from the regression model $Y = m(X) + \varepsilon$, where m(x) = 100 $0.25x^2 - 0.75x + 3$, $X \sim N(0, 1.5^2)$, and $\varepsilon \sim N(0, 0.75^2)$.

```
library(np)
set.seed(1234)
M = 500
n = 100
x m = matrix(0, ncol = n, nrow = M)
y_muestras = matrix(0,ncol = n,nrow = M)
```

```
m = function(a){
    0.25*a^2-0.75*a+3
}
for (i in 1:500){
    x = rnorm(100,0,1.5)
    epsilon = rnorm(100,0,0.75)
    y_muestras[i,] = m(x)+epsilon
    x_m[i,] = x
}
```

2. Compute the 95% confidence intervals for m(x) along $x \leftarrow seq(-5, 5, by = 0.1)$, for each of the M samples. Do it for the normal approximation and quantile-based confidence intervals. **Solution:** Using the function to estimate the interval from the book we are applying this for each sample and calculate the CI for each of those.

```
x grid = seq(-5,5,by=0.1)
np_pred_CI <- function(npfit, exdat, B = 200, conf = 0.95,
                         type_CI = c("standard", "quantiles")[1]) {
  # Extract predictors
  xdat <- npfit$eval
  # Extract response, using a trick from np::npplot.rbandwidth
  tt <- terms(npfit$bws)
  tmf <- npfit$bws$call[c(1, match(c("formula", "data"),</pre>
                                      names(npfit$bws$call)))]
  tmf[[1]] <- as.name("model.frame")</pre>
  tmf[["formula"]] <- tt</pre>
  tmf <- eval(tmf, envir = environment(tt))</pre>
  ydat <- model.response(tmf)</pre>
  # Predictions
  m_hat <- np::npreg(txdat = xdat, tydat = ydat, exdat = exdat,</pre>
                      bws = npfit$bws)$mean
  # Function for performing Step 3
  boot_function <- function(data, indices) {</pre>
    np::npreg(txdat = xdat[indices,], tydat = ydat[indices],
               exdat = exdat, bws = npfit$bws)$mean
  # Carry out Step 3
  m_hat_star <- boot::boot(data = data.frame(xdat), statistic = boot_function,</pre>
                             R = B)$t
  # Confidence intervals
  alpha <- 1 - conf
  if (type_CI == "standard") {
    z \leftarrow qnorm(p = 1 - alpha / 2)
    se <- apply(m_hat_star, 2, sd)</pre>
    lwr <- m_hat - z * se</pre>
    upr \leftarrow m hat + z * se
  } else if (type_CI == "quantiles") {
```

```
q <- apply(m_hat_star, 2, quantile, probs = c(alpha / 2, 1 - alpha / 2))
lwr <- q[1, ]
upr <- q[2, ]
} else {
    stop("Incorrect type_CI")
}
# Return evaluation points, estimates, and confidence intervals
    return(data.frame("exdat" = exdat, "m_hat" = m_hat, "lwr" = lwr, "upr" = upr))
}</pre>
```

Now with the function for calculate CI normal and quantile we create a matrix to save the values for the interval of each sample.

```
set.seed(42234)
# Storage
normal_ci_up = matrix(NA, nrow= M,ncol = length(x_grid))
normal_ci_low = matrix(NA, nrow= M,ncol = length(x_grid))
m_hat_normal = matrix(NA, nrow = M, ncol = length(x_grid))

for (i in 1:M) {
    # Fit nonparametric regression to give to the function
    np_fit <- npreg(bws = npregbw(y_muestras[i,] ~ x_m[i,]), gradients = FALSE)

# Compute CI using function first for normal CI
    ci_normal_result = np_pred_CI(np_fit, exdat = x_grid, B = 500, type_CI = "standard")

m_hat_normal[i,] = ci_normal_result$m_hat
    normal_ci_up[i,] = ci_normal_result$upr
    normal_ci_low[i,] = ci_normal_result$lwr
}</pre>
```

Create CI with quantile formula.

```
set.seed(42234)
# Storage
quantile_ci_up = matrix(NA, ncol = length(x_grid),nrow = M)
quantile_ci_low = matrix(NA, ncol = length(x_grid),nrow = M)
m_hat_quantile <- matrix(NA, ncol = length(x_grid), nrow = M)

for (i in 1:M) {
    # Fit nonparametric regression to give to the function
    np_fit <- npreg(bws = npregbw(y_muestras[i,] ~ x_m[i,]), gradients = FALSE)

# Compute CI using function first for normal CI
    ci_normal_result = np_pred_CI(np_fit, exdat = x_grid, B = 500, type_CI = "quantiles")

m_hat_quantile[i,] = ci_normal_result$m_hat
    quantile_ci_up[i,] = ci_normal_result$upr
    quantile_ci_low[i,] = ci_normal_result$lwr
}</pre>
```

```
normal_ci_low = readRDS("normal_ci_low.RDS")
normal_ci_up = readRDS("normal_ci_up.RDS")
quantile_ci_low = readRDS("quantile_ci_low.RDS")
quantile_ci_up = readRDS("quantile_ci_low.RDS")
```

3. Check if m(x) belongs to each of the confidence intervals, for each x. Solution:

To check if m(x) belongs to each confidence intervals we need to see if m(x) belongs to each sample. Then the save value is going to be true if belong and false if it doesn't.

```
belong_normal = matrix(NA,ncol = length(x_grid), nrow = M)
belong_quantile = matrix(NA,ncol = length(x_grid), nrow = M)
m.x = sapply(x_grid,FUN = m)
for (i in 1:M){
  for (j in 1:length(x_grid)){
    if (m.x[j]>=normal_ci_low[i,j] & m.x[j] <= normal_ci_up[i,j]){</pre>
      belong normal[i,j] = TRUE
    }
    else{
      belong_normal[i,j] = FALSE
    if (m.x[j]>=quantile_ci_low[i,j] & m.x[j] <= quantile_ci_up[i,j]){</pre>
      belong_quantile[i,j] = TRUE
    }
    else{
      belong_quantile[i,j] = FALSE
    }
 }
}
```

4. Approximate the actual coverage of the confidence intervals. **Solution:** To estimate the coverage we are calculating the % of belonging of m(x) for each sample and the do the mean of this value. We can do this apply mean by row because the data is boolean.

```
coverage.normal = apply(belong_normal, 1, mean)
coverage.quantile = apply(belong_quantile, 1, mean)
cat("the mean coverage of the normal CI is :",mean(coverage.normal),"and ","the mean coverage of the qu
```

the mean coverage of the normal CI is : 0.765604 and the mean coverage of the quantile CI is : 0

```
#n = 200
set.seed(1234)
n2 = 200
x_m2 = matrix(0,ncol = n2,nrow = M)
y_muestras2 = matrix(0,ncol = n2,nrow = M)
for (i in 1:500){
    x = rnorm(n2,0,1.5)
    epsilon = rnorm(n2,0,0.75)
    y_muestras2[i,] = m(x)+epsilon
    x_m2[i,] = x
}
```

```
# Storage
normal_ci_up2 = matrix(NA, nrow= M,ncol = length(x_grid))
normal_ci_low2 = matrix(NA, nrow= M,ncol = length(x_grid))
m_hat_normal2 = matrix(NA, nrow = M, ncol = length(x_grid))
for (i in 1:M) {
  # Fit nonparametric regression to give to the funciton
  np_fit <- npreg(bws = npregbw(y_muestras2[i,] ~ x_m2[i,]), gradients = FALSE)</pre>
  # Compute CI using function first for normal CI
  ci_normal_result2 = np_pred_CI(np_fit, exdat = x_grid, B = 500, type_CI = "standard")
  m_hat_normal2[i,] = ci_normal_result2$m_hat
  normal_ci_up2[i,] = ci_normal_result2$upr
  normal_ci_low2[i,] = ci_normal_result2$lwr
}
quantile_ci_up2 = matrix(NA, ncol = length(x_grid),nrow = M)
quantile_ci_low2 = matrix(NA, ncol = length(x_grid),nrow = M)
m_hat_quantile2 <- matrix(NA, ncol = length(x_grid), nrow = M)</pre>
for (i in 1:M) {
  # Fit nonparametric regression to give to the funciton
  np_fit <- npreg(bws = npregbw(y_muestras2[i,] ~ x_m2[i,]), gradients = FALSE)</pre>
  # Compute CI using function first for normal CI
  ci_normal_result2 = np_pred_CI(np_fit, exdat = x_grid, B = 500, type_CI = "quantiles")
  m_hat_quantile2[i,] = ci_normal_result2$m_hat
  quantile_ci_up2[i,] = ci_normal_result2$upr
  quantile_ci_low2[i,] = ci_normal_result2$lwr
}
normal_ci_low2 = readRDS("normal_ci_low2.RDS")
normal_ci_up2 = readRDS("normal_ci_up2.RDS")
quantile ci low2 = readRDS("quantile ci low2.RDS")
quantile_ci_up2 = readRDS("quantile_ci_low2.RDS")
belong_normal2 = matrix(NA,ncol = length(x_grid), nrow = M)
belong_quantile2 = matrix(NA,ncol = length(x_grid), nrow = M)
m.x2 = sapply(x_grid,FUN = m)
for (i in 1:M){
  for (j in 1:length(x_grid)){
    if (m.x2[j]>=normal_ci_low2[i,j] & m.x2[j] <= normal_ci_up2[i,j]){</pre>
      belong_normal2[i,j] = TRUE
    }
    else{
      belong_normal2[i,j] = FALSE
    if (m.x2[j]>=quantile ci low2[i,j] & m.x2[j] <= quantile ci up2[i,j]){
      belong_quantile2[i,j] = TRUE
```

```
else{
      belong_quantile2[i,j] = FALSE
    }
 }
coverage.normal2 = apply(belong_normal2, 1, mean)
coverage.quantile2 = apply(belong_quantile2, 1, mean)
cat("the mean coverage of the normal CI is : ", mean(coverage.normal2), "and ", "the mean coverage of the q
## the mean coverage of the normal CI is : 0.8064158 and the mean coverage of the quantile CI is : 0
#n = 500
set.seed(1234)
n3 = 500
x_m3 = matrix(0, ncol = n3, nrow = M)
y_muestras3 = matrix(0,ncol = n3,nrow = M)
for (i in 1:500){
 x = rnorm(n3,0,1.5)
  epsilon = rnorm(n3,0,0.75)
  y_{muestras3[i,]} = m(x) + epsilon
  x_m3[i,] = x
}
# Storage
normal_ci_up3 = matrix(NA, nrow= M,ncol = length(x_grid))
normal_ci_low3 = matrix(NA, nrow= M,ncol = length(x_grid))
m_hat_normal3 = matrix(NA, nrow = M, ncol = length(x_grid))
for (i in 1:M) {
  # Fit nonparametric regression to give to the funciton
  np_fit <- npreg(bws = npregbw(y_muestras3[i,] ~ x_m3[i,]), gradients = FALSE)</pre>
  # Compute CI using function first for normal CI
  ci_normal_result3 = np_pred_CI(np_fit, exdat = x_grid, B = 500, type_CI = "standard")
  m_hat_normal3[i,] = ci_normal_result3$m_hat
 normal_ci_up3[i,] = ci_normal_result3$upr
  normal_ci_low3[i,] = ci_normal_result3$1wr
quantile_ci_up3 = matrix(NA, ncol = length(x_grid),nrow = M)
quantile_ci_low3 = matrix(NA, ncol = length(x_grid), nrow = M)
m_hat_quantile3 <- matrix(NA, ncol = length(x_grid), nrow = M)</pre>
for (i in 1:M) {
  # Fit nonparametric regression to give to the funciton
  np_fit <- npreg(bws = npregbw(y_muestras3[i,] ~ x_m3[i,]), gradients = FALSE)</pre>
  # Compute CI using function first for normal CI
  ci_normal_result3 = np_pred_CI(np_fit, exdat = x_grid, B = 500, type_CI = "quantiles")
  m_hat_quantile3[i,] = ci_normal_result3$m_hat
  quantile_ci_up3[i,] = ci_normal_result3$upr
```

```
quantile_ci_low3[i,] = ci_normal_result3$lwr
}
normal_ci_low3 = readRDS("normal_ci_low3.RDS")
normal_ci_up3 = readRDS("normal_ci_up3.RDS")
quantile_ci_low3 = readRDS("quantile_ci_low3.RDS")
quantile_ci_up3 = readRDS("quantile_ci_low3.RDS")
belong_normal3 = matrix(NA, ncol = length(x_grid), nrow = M)
belong_quantile3 = matrix(NA,ncol = length(x_grid), nrow = M)
m.x3 = sapply(x_grid, FUN = m)
for (i in 1:M){
  for (j in 1:length(x_grid)){
    if (m.x3[j]>=normal_ci_low3[i,j] & m.x3[j] <= normal_ci_up3[i,j]){</pre>
      belong_normal3[i,j] = TRUE
    }
    else{
      belong_normal3[i,j] = FALSE
    }
    if (m.x3[j]>=quantile_ci_low3[i,j] & m.x3[j] <= quantile_ci_up3[i,j]){</pre>
      belong_quantile3[i,j] = TRUE
    else{
      belong_quantile3[i,j] = FALSE
    }
  }
}
coverage.normal3 = apply(belong_normal3, 1, mean)
coverage.quantile3 = apply(belong_quantile3, 1, mean)
cat("the mean coverage of the normal CI is : ", mean(coverage.normal3), "and ", "the mean coverage of the q
```

the mean coverage of the normal CI is : 0.8580792 and the mean coverage of the quantile CI is : 0

When you increase the number in the n the covarege increase almost 5% in each time. This is because having a large sample increase the knowlege in the population given a better CI. Also the Normal CI have a larger coverage for the central limit theorem and this would increase just by increasing the sample size.

```
ci_normal_result %>% saveRDS("ci_normal_result.RDS")
ci_normal_result2 %>% saveRDS("ci_normal_result2.RDS")
ci_normal_result3 %>% saveRDS("ci_normal_result3.RDS")
m_hat_quantile %>% saveRDS("m_hat_quantile.RDS")
m_hat_quantile2 %>% saveRDS("m_hat_quantile2.RDS")
m_hat_quantile3 %>% saveRDS("m_hat_quantile3.RDS")
normal_ci_low %>% saveRDS("normal_ci_low.RDS")
normal_ci_low2 %>% saveRDS("normal_ci_low2.RDS")
normal_ci_low3 %>% saveRDS("normal_ci_low3.RDS")
normal_ci_up %>% saveRDS("normal_ci_up.RDS")
normal_ci_up2 %>% saveRDS("normal_ci_up3.RDS")
normal_ci_up3 %>% saveRDS("normal_ci_up3.RDS")
quantile_ci_low %>% saveRDS("quantile_ci_low.RDS")
```

```
quantile_ci_low2 %>% saveRDS("quantile_ci_low2.RDS")
quantile_ci_low3 %>% saveRDS("quantile_ci_low3.RDS")
quantile_ci_up %>% saveRDS("quantile_ci_up.RDS")
quantile_ci_up2 %>% saveRDS("quantile_ci_up2.RDS")
quantile_ci_up3 %>% saveRDS("quantile_ci_up3.RDS")
```

Excercise 6.8

The following exercise revolves around the comparison of power between the three nonparametric normality tests:

- 1. Kolmogorov-Smirnov
- 2. Anderson-Darling
- 3. Cramer von Mises tests.

with the test hypothesis:

```
H_0: X \sim N(0,1) \text{ vs } H_1: X \not\sim N(0,1)
```

The comparison is performed by performing these tests M times for samples of size n from a chosen normal or t-student's and comparing the distribution of the resulting M p-values the corresponding rejection rate for significance level α .

```
# Load Libraries Required for Excercises
library(goftest)
library(nortest)
library(latex2exp)
```

We begin by setting up a versatile function that performs three steps:

- 1. Draw M samples of size n from either a normal or t-student's distribution, perform the three nonparametric tests and store the resulting M p-values.
- 2. For the three normality tests with , calculate the estimated rejection rate (Power of the test) for significance level α .

$$\mbox{Rejection Rate} = \frac{\mbox{Number of rejections of}\ H_0}{\mbox{Total number of tests}}$$

3. Plot the distribution of the M p-values for each test.

```
# Function for Simulation of p-values
simulate_pvalues <- function(mu_samp=0, sd_samp=1 , M, n, alpha, dist = "normal") {

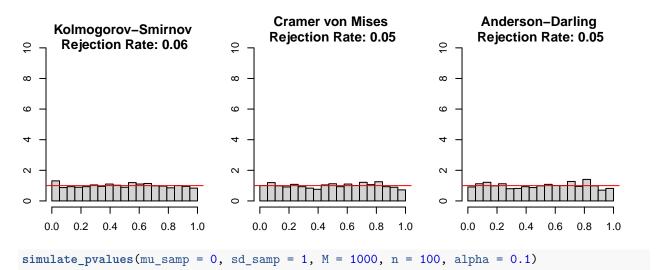
# generate p-Values for samples from Normal
pvalues <- sapply(1:M, function(i) {

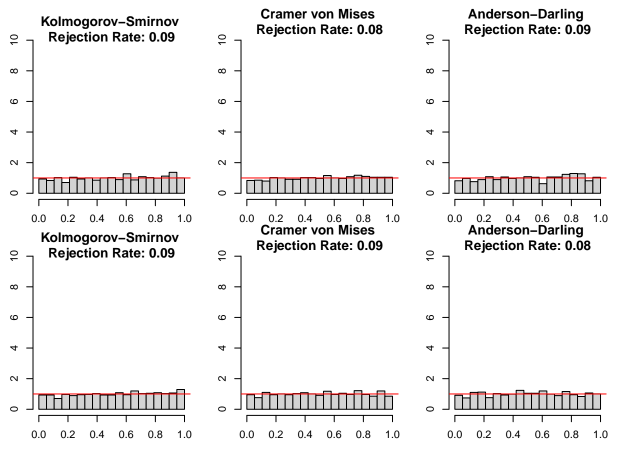
if(dist == "normal"){
    x = rnorm(n, mean = mu_samp, sd = sd_samp)
}</pre>
```

```
if(dist == "t-students"){
      x = rt(n, df = 10, ncp = mu_samp)
    pval_ks = ks.test(x, "pnorm", mean = 0, sd = 1)$p.value
    pval_cvm = goftest::cvm.test(x = x, null = "pnorm")$p.value
    pval_ad = goftest::ad.test(x = x, null = "pnorm")$p.value
    return(c(pval ks, pval cvm, pval ad))
  })
  # get rejection rates
  ks_rejection_rate = sum(pvalues[1,] < alpha) / M</pre>
  cvm_rejection_rate = sum(pvalues[2,] < alpha) / M</pre>
  ad_rejection_rate = sum(pvalues[3,] < alpha) / M</pre>
  # plot
  par(mfrow = c(2, 3), mar = c(2, 2, 2, 2)) # Increase margins
  hist(pvalues[1,],
       breaks = seq(0, 1, 1 = 20),
       probability = TRUE,
       main = paste("\nKolmogorov-Smirnov", "\nRejection Rate:", round(ks_rejection_rate, 2)),
       ylim = c(0, 10),
       ylab = "p-value")
  abline(h = 1, col = "red")
  hist(pvalues[2,],
       breaks = seq(0, 1, 1 = 20),
       probability = TRUE,
       main = paste("Cramer von Mises", "\nRejection Rate:", round(cvm_rejection_rate, 2)),
       ylim = c(0, 10),
       ylab = "p-value")
  abline(h = 1, col = "red")
  hist(pvalues[3,],
       breaks = seq(0, 1, 1 = 20),
       probability = TRUE,
       main = paste("Anderson-Darling", "\nRejection Rate:", round(ad_rejection_rate, 2)),
       ylim = c(0, 10),
       ylab = "p-value")
  abline(h = 1, col = "red")
}
```

With the function in place, we can now simulate the p-values for the three tests for samples drawn from a normal distribution N(0,1) for different sample sizes n and significance levels α . With the sample drawn from a normal distribution, we expect the three test to overwhelmingly not reject the null Hypothesis $H \in F_N$ and the distribution of the p-Values to be a Uniform U(0,1). In fact, for both n=25 and n=100 and as well as $\alpha=0.05$ and $\alpha=0.1$, theses expectations are met, with the rejection rates corresponding to the chosen significance levels.

```
# HO True
set.seed(123)
simulate_pvalues(mu_samp = 0, sd_samp = 1, M = 1000, n = 25, alpha = 0.05)
```

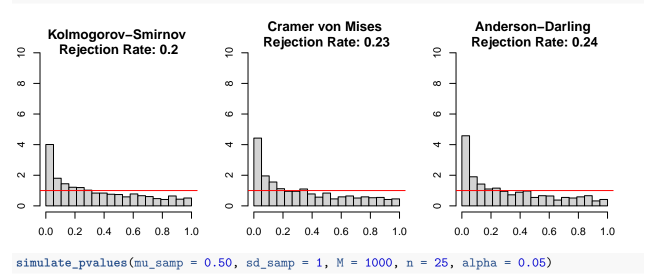


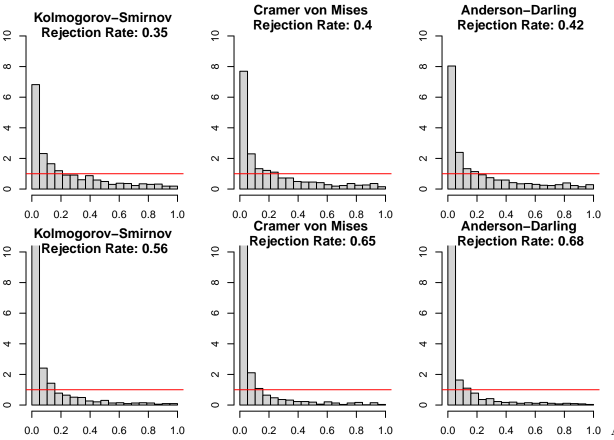


Performing the test for a sample drawn from a normal distribution (μ, σ) where $\mu \neq 0$, $\sigma \neq 1$, again for different sample sizes and a constant significance level $\alpha = 0.05$, results are more mixed. With the sample not drawn from a N(0,1) we expect rejection to happen at a greater rate compared to the previous results. In addition, we expect both Cramer von Mises and Anderson Darling to deliver greater rejections rates as they are in general terms more powerful than the Kolomogorov-Smirnov. This is in line with the results acquired from the three parameter combinations as can be seen in the shown rejection rates. Furthermore,

increasing the difference of μ from 0, appears to increase rejection rates more than an increase in sample size.

```
# HO False
simulate_pvalues(mu_samp = 0.25, sd_samp = 1, M = 1000, n = 25, alpha = 0.05)
simulate_pvalues(mu_samp = 0.25, sd_samp = 1, M = 1000, n = 50, alpha = 0.05)
```

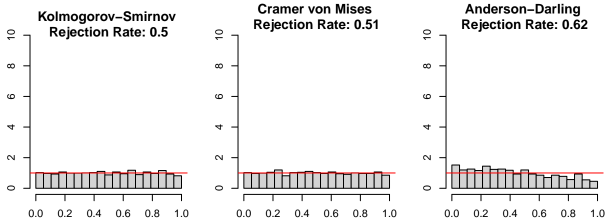




last experiment, we now draw the samples from a t-students distribution with $\mu = 0$ and df = 10. Like before, we expect rejection rates to be high as the null hypothesis $H_0: F = F_N$ is false. This is inline with the results, with an additional observation that the rejection rate of the Anderson-Darling test is higher

than those of the other two test. This can be explained with the emphasises of the Anderson-Darling test on the tails of the distribution, picking up better the different behaviors Normal and t-students distribution have at the tails.

```
simulate_pvalues(mu_samp = 0,M = 1000, n = 50, alpha = 0.5, dist = "t-students")
```



This difference in the tails can be seen by comparing both distributions as done in the plot below.

