Series 2

1. In a study on the contribution of air pollution to mortality, General Motors collected data from 60 US Standard Metropolitan Statistical Areas (SMSAs). The dependent variable is the age adjusted mortality. The data includes variables measuring demographic characteristics of the cities, variables measuring climate characteristics and variables recording the pollution potential.

Mortality Age Adjusted mortality

JanTemp Mean January temperature (degrees Fahrenheit)

JulyTemp Mean July temperature (degrees Fahrenheit)

RelHum Relative Humidity
Rain Annual rainfall (inches)
Educ Median education
Dens Population density
NonWhite Percentage of non whites

WhiteCollar Percentage of white collar workers

Pop Population Income Median income

HC Hydrocarbon pollution potential
 NOx Nitrous Oxide pollution potential
 SO2 Sulfur Dioxide pollution potential

Try to find a good linear model that represents the data. Therefore, take the following steps:

- 1. Produce diagnostic plots such as scatterplots of the response **Mortality** versus the predictors and residual plots. Are the model assumptions valid? Which variables need to be log-transformed?
- 2. Perform a stepwise variable selection. Do forward selection and backward elimination lead to the same model? Which variables have a significant influence on the **Mortality**-level and in which direction? Compare these results to an all-subsets regression using Mallows- C_p .

R-hints:

```
> ## Reading the dataset
> url <- "http://stat.ethz.ch/Teaching/Datasets/mortality.csv"
> mortality <- read.csv(url,header = TRUE)</pre>
> mortality <- mortality[,-1]</pre>
> ##Create pairs plot using the splom() function of the ``lattice" package
> library(lattice)
> splom(~mortality,pscales=0,cex=0.5)
> ## Fit the full model
> mortal.full <- lm(Mortality ~. , data=mortality)</pre>
> ## Fit the empty model. This is not very useful in itself, but is required
> ## as a starting model for stepwise forward variable selection
> mortal.empty <- lm(Mortality ~ 1, data = mortality)</pre>
> ## Backward elimination, starting from the full model
               <- step(mortal.full, direction = "backward")
> mortal.bw
> ## Forward selection, starting from the empty model
               <- step(mortal.empty, direction = "forward",
> mortal.fw
                     scope = list(upper=mortal.full,lower=mortal.empty))
> ## Loading the package for all-subsets regression
> library(leaps)
```

```
## All subsets model choice, compare to the stepwise methods
mortal.alls <- regsubsets(...)

## Load function to produce a nice figure of C_p versus p
source("ftp://stat.ethz.ch/Teaching/maechler/CompStat/cp-plot.R")
p.regsubsets(mortal.alls)</pre>
```

2. a) The artificial dataset shown in Figure 1 is generated from a mixture of two normal distributions (see part b)). Imagine you wouldn't know anything about the distribution and you'd want to estimate the density by use of a kernel density estimator with Gaussian kernel. Guess a good bandwidth.

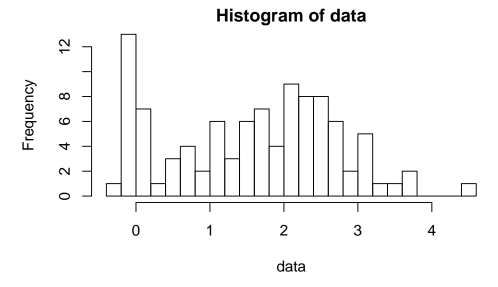


Figure 1: Artificial data from mixture of normal distributions

- b) Carry out a simulation with R to evaluate the quality of bandwidths for kernel density estimation for a mixture of normal distributions where a point is generated by a $\mathcal{N}(0,0.01)$ -distribution with probability 0.2 and by a $\mathcal{N}(2,1)$ -distribution with probability 0.8. Repeat the following code 200 times for multiple bandwidth values: h = 0.02, 0.1, 0.3, 0.6, 1, 1.5, as well as h equal to your own guess from part a) and to a plug-in estimate of the optimal bandwidth (see below):
 - 1. Generate a dataset of 100 points from the mixture distribution from above:

```
> set.seed(1)
> data <- numeric(100)
> for(i in 1:100){
   p <- runif(1, min = 0, max = 1)
   if (p < 0.2)
     data[i] <- rnorm(1, mean = 0, sd = sqrt(0.01))
   else
     data[i] <- rnorm(1, mean = 2, sd = 1)
}</pre>
```

2. Compute the kernel density estimator with the function density. Consult help(density) to understand the syntax.

```
> ke < -density(data, bw = 0.1, n = 61, from = -1, to = 5)
An estimated plug-in bandwidth is obtained by using bw = "sj".
```

3. In the lecture, we saw that the goodness of a kernel estimate can be measured using the integrated mean squared error (IMSE). In practice, the IMSE can be approximated by summing the squared difference between the kernel estimator and the true density values (up to a constant factor).

Note that the commands given above are suitable for a single execution, but not necessarily for the full simulation. For example, you will need a vector of the qualities of all simulation runs. You may define a matrix for the qualities to store the results for all different bandwidths. Consider help(matrix), help(apply).

Compute and compare the averaged quality of the kernel estimators for the different bandwidths.

c) (Optional) Repeat the whole project with the Epanechnikov kernel, which can be obtained by specifying kernel="epanechnikov" in density.

Preliminary discussion: Friday, March 06.

Deadline: Friday, March 13.