

omicsExample1.R

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```
# omicsExample1.R
#
# Code for Example 1 for R intro in course Omics Data Analysis: a simple
# qPCR data set (not very omic, but we have to start somewhere)
#
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# We assume that the file is in this directory and set it as working directory
# Modify as required
setwd("~/OmicsDataAnalysis/Data/")
dat <- read.table("qPCR.txt", header = TRUE, stringsAsFactors = TRUE)

# Look at the top of the data: seems ok
head(dat)

##      Sample Con DeltaCt
## 1 Control   10  3.3687
## 2 Control   10  3.4063
## 3 Control   10  3.5066
## 4 Control    2  4.5963
## 5 Control    2  3.7704
## 6 Control    2  3.4906

# Structure of the data: one grouping variable, four numeric variables
str(dat)

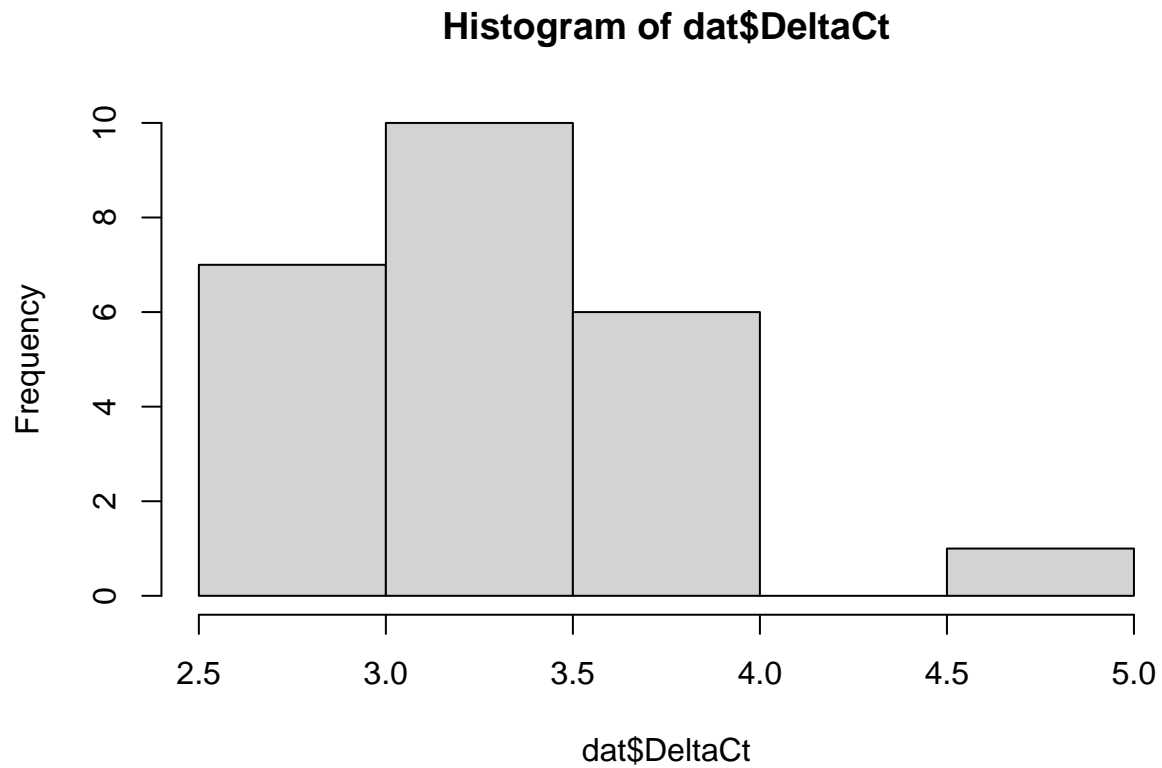
## 'data.frame':   24 obs. of  3 variables:
##  $ Sample : Factor w/ 2 levels "Control","Treatment": 1 1 1 1 1 1 1 1 1 1 ...
##  $ Con     : num  10 10 10 2 2 2 0.4 0.4 0.4 0.08 ...
##  $ DeltaCt: num  3.37 3.41 3.51 4.6 3.77 ...

# Simple numerical summaries of the data
summary(dat)

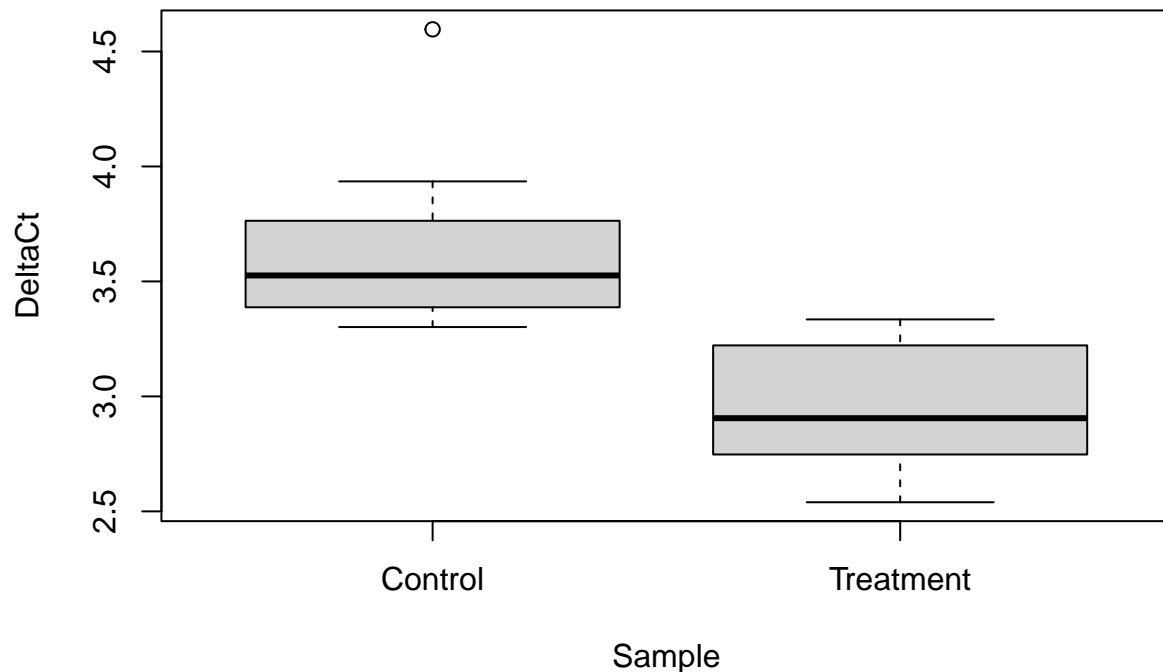
##           Sample           Con           DeltaCt
## Control :12   Min.      : 0.08   Min.      :2.540
## Treatment:12  1st Qu.: 0.32   1st Qu.:2.919
##              Median : 1.20   Median :3.335
##              Mean   : 3.12   Mean    :3.298
##              3rd Qu.: 4.00   3rd Qu.:3.516
##              Max.   :10.00   Max.    :4.596

# We want to study how DeltaCt (difference in time until detection threshold
# between sequence of interest and reference sequence) differs between
# treated / controls
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# Let's start by looking at the distribution of DeltaCt
# Hm, seems like we have a fairly large value
hist(dat$DeltaCt)
```



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# Direct comparison: treatment vs controls
# Looks like treated samples have lower DeltaCt,
# ergo higher level of SOI
boxplot(DeltaCt ~ Sample, data = dat)
```



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# Statistical test: is DeltaCt on average the same?
# t-test without equal variance assumption (i.e. Welch t-test)
# Null hypothesis: same average DeltaCt in both groups
# Based on p-value and confidence interval, we can reject this
t.test(DeltaCt ~ Sample, data = dat)
```

```
##
## Welch Two Sample t-test
##
## data: DeltaCt by Sample
## t = 5.2577, df = 20.684, p-value = 3.429e-05
## alternative hypothesis: true difference in means between group Control and group Treatment is not eq
## 95 percent confidence interval:
## 0.413698 0.955952
## sample estimates:
## mean in group Control mean in group Treatment
## 3.640408 2.955583
```

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# Sensitivity analysis: does our result depend on the one
# large values among the controls? We re-run without that
# observation
dat2 <- subset(dat, DeltaCt < 4)
str(dat2) # We removed exactly one observation
```

```
## 'data.frame': 23 obs. of 3 variables:
## $ Sample : Factor w/ 2 levels "Control","Treatment": 1 1 1 1 1 1 1 1 1 1 ...
## $ Con : num 10 10 10 2 2 0.4 0.4 0.4 0.08 0.08 ...
```

```
## $ DeltaCt: num 3.37 3.41 3.51 3.77 3.49 ...
# Yes, this holds up well; the original result (to be reported)
# is not unduly affected by just one observation
t.test(DeltaCt ~ Sample, data = dat2)

##
## Welch Two Sample t-test
##
## data: DeltaCt by Sample
## t = 5.9708, df = 20.05, p-value = 7.644e-06
## alternative hypothesis: true difference in means between group Control and group Treatment is not eq
## 95 percent confidence interval:
## 0.3890666 0.8067849
## sample estimates:
## mean in group Control mean in group Treatment
## 3.553509 2.955583
```