

# **SLLD - Module 1**

## **Resampling**

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```
library(boot) #Bootstrap Functions
library(coin) #Conditional Inference Procedures in a
# Permutation Test Framework
library(ggplot2) #Create Elegant Data Visualisations Using
# the Grammar of Graphics
```

## Data

We generate a sample from a binomial distribution with parameters  $(25, 0.6)$ .

```
options(width = 60)
set.seed(123)
x <- rbinom(n=50,           # sample size
             size=25,         # num. of trials
             prob=0.6)       # prob. of success per trial
x
```

```
## [1] 16 13 16 12 11 19 15 12 15 15 11 15 14 15 18 12 17 19
## [19] 16 11 12 14 14 9 14 14 15 14 16 18 11 12 14 13 20 15
## [37] 13 17 16 17 18 16 16 16 18 18 17 15 17 12
```

```
n <- length(x)
summary(x)
```

```
##      Min. 1st Qu. Median      Mean 3rd Qu.      Max.
## 9.00   13.00  15.00  14.86  16.75  20.00
```

We can automatically perform non-parametric bootstrapping using the **boot** package. The main bootstrapping function is **boot()** and has the following syntax:

- **data**: The data as a vector, matrix or data frame.
- **statistic**: A function which when applied to data returns a vector containing the statistic(s) of interest.
- **R**: The number of bootstrap replicates.

**Remark:** it is mandatory to pass a "user-defined function" in the field statistic. In the case of the 95th percentile, our estimation function is:

```
# x: vector
# d: set of indexes
# prob: quantile
sampleperc <- function(x, d, prob=0.95) {
  return(quantile(x[d], probs=prob))
}
```

Once we have written a function like this, here is how we obtain bootstrap estimates of the standard error for the 95th percentile of the distribution:

```
set.seed(123)
b = boot(x, sampleperc, R=1000)
print(b)

##
## ORDINARY NONPARAMETRIC BOOTSTRAP
##
##
## Call:
## boot(data = x, statistic = sampleperc, R = 1000)
##
##
## Bootstrap Statistics :
##      original    bias   std. error
## t1*    18.55 -0.03435   0.5642082
```

# Parametric Bootstrapping

We have a sample of size  $n = 50$  from a binomial distribution with parameters ( $N = 25$ ,  $p = 0.6$ ).

```
set.seed(123)
n = 50
N <- 25
x <- rbinom(n = n, size = N, prob = 0.6)
```

Let's assume that we know the underlying distribution, where the actual parameter  $p$  is unknown, and we want to estimate the 95th percentile as before.

```
p_hat <- mean(x)/N
p_hat
```

```
## [1] 0.5944
```

We use parametric bootstrap and compute B samples of size n from the known distribution.

```
B <- 1000 # number of bootstrap samples  
tempdata <- rbinom(B*n,  
size = N,  
prob = p_hat)  
bootstrapsample <- matrix(tempdata, nrow = n, ncol = B)  
dim(bootstrapsample)
```

```
## [1] 50 1000
```

```
bootstrapsample[1:3, 1:5]
```

```
##      [,1] [,2] [,3] [,4] [,5]  
## [1,]    19   14   12   17   16  
## [2,]    15   16   15   10   17  
## [3,]    13   15   16   14   14
```

Now we can compute the statistic of interest on each of these 1000 samples – producing B bootstrap values. In our case, we are interested in the 95th percentile.

```
B_values <- apply(bootstrapsample, 2, quantile, prob=0.95)  
summary(B_values)
```

```
##      Min. 1st Qu. Median      Mean 3rd Qu.      Max.  
## 17.00 18.00 18.55 18.60 19.00 20.55
```

So we have the following estimate and standard error:

```
resultsParam <- data.frame("mean" = mean(B_values),  
"SD" = sd(B_values))  
rownames(resultsParam) <- "Manual_Parametric"  
resultsParam
```

```
##                      mean             SD  
## Manual_Parametric 18.60495 0.6487479
```

Let's compare it with the boot function:

```
set.seed(123)  
B = 1000  
p.rg <- function(data, mle, N=25) {  
  out <- rbinom(length(data),  
  size = N,  
  prob = mle)  
  out  
}  
bBoot <- boot(x, sampleperc, R = B, sim = "parametric",  
ran.gen = p.rg, mle = (mean(x)/N))
```

bBoot

```
##  
## PARAMETRIC BOOTSTRAP  
##  
##  
## Call:  
## boot(data = x, statistic = sampleperc, R = B, sim = "parametric",  
##       ran.gen = p.rg, mle = (mean(x)/N))  
##  
##  
## Bootstrap Statistics :  
##      original   bias   std. error  
## t1*    18.55  0.0555  0.6484676
```

```
# notice how close this is to our previous computation  
resultsParam <- rbind.data.frame(resultsParam,  
data.frame("mean" = bBoot$t0,  
"SD" = sd(bBoot$t)))  
rownames(resultsParam)[2] <- "boot_Parametric"  
resultsParam
```

```
##               mean        SD  
## Manual_Parametric 18.60495 0.6487479  
## boot_Parametric   18.55000 0.6484676
```

# Permutation Test

Permutation tests are particularly relevant in experimental studies, where we are often interested in the sharp null hypothesis of no difference between treatment groups.

Let's generate a dataset divided into treatment (1) and control group (0).

```
# they have a difference in mean equal to 1
set.seed(1)
n <- 80
tr <- rbinom(n, 1, 0.3)
y <- 1 + tr + rnorm(n, 0, 2)
```

Let us compute the difference in mean between the two groups. The difference in means is, as we would expect (since we made it up), about 1:

```
means <- by(y, tr, mean)
diff0 <- diff(means)
diff0
```

```
## [1] 1.198222
```

To obtain a single permutation of the data, we simply resample without replacement and calculate the difference again:

```
s <- sample(tr, length(tr), FALSE) # shuffle the labels  
by(y, s, mean) # compute mean in vector y according to class s
```

```
## s: 0  
## [1] 1.732339  
## -----  
## s: 1  
## [1] 1.147594
```

```
diff(by(y, s, mean)) # difference between 2 means
```

```
## [1] -0.584745
```

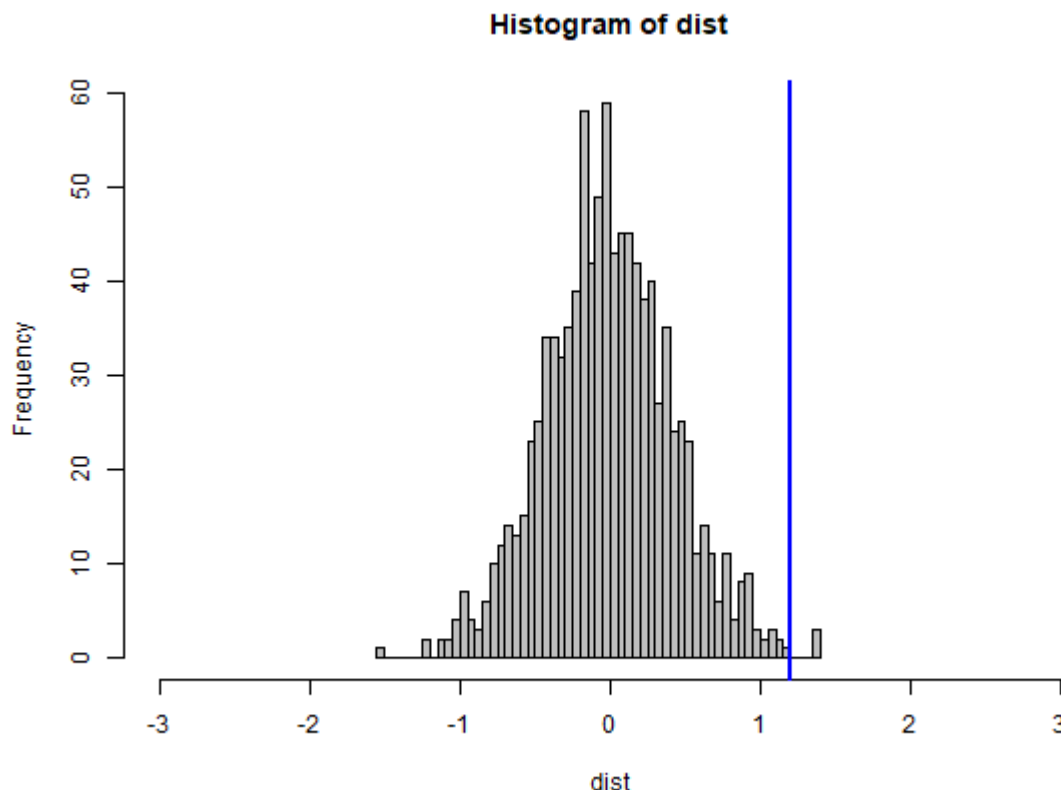
If we repeat this process a large number of times, we can build our approximate permutation distribution (i.e., the sampling distribution for the mean-difference). We use the function **replicate** to repeat our permutation process.

```
set.seed(123)
nperm = 1000
# nperm = 200 # try this too :)
# nperm = 5000
dist <- replicate(nperm, diff(by(y, sample(tr, length(tr),
                                         FALSE), mean)))
head(dist)
```

```
##           1           1           1           1
## 0.158136857 -0.128001168 -0.455300572 -1.004146337
##           1           1
## 0.007316022  0.094566325
```

We can look at our distribution using an histogram indicating with a vertical line the observed difference:

```
hist(dist, xlim = c(-3, 3), col = "grey", breaks = 100)
abline(v = diff(by(y, tr, mean)), col = "blue", lwd = 2)
```



Now, we can use the distribution to obtain a p-value for our mean-difference by counting how many permuted mean-differences are larger than the one we observed in our actual data. We can then divide this by the number of items in our permutation distribution (i.e., nperm=1000 from our call to replicate, above):

```
sum(dist > diff0)/nperm # one-tailed test
```

```
## [1] 0.003
```

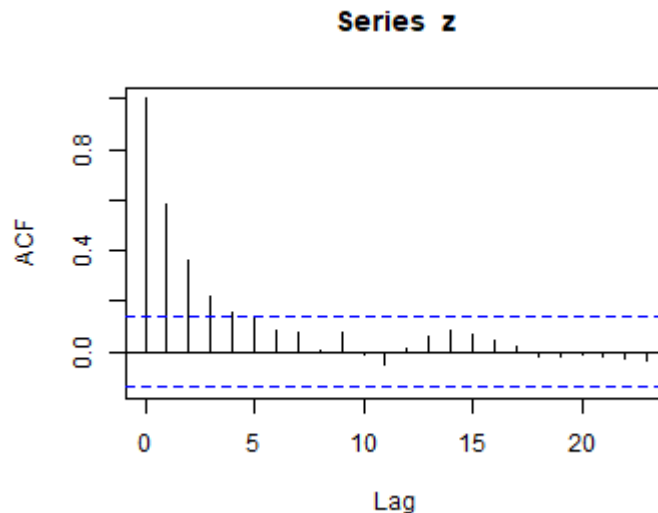
```
sum(abs(dist) > abs(diff0))/nperm # two-tailed test
```

```
## [1] 0.006
```

# Time Series Bootstrap

The function **tsboot** (in the **boot** package) generates R bootstrap replicates of a statistic applied to a time series. The replicate time series can be generated using fixed or random block lengths or can be model based replicates.

```
# they have a difference in mean equal to 1
set.seed(1)
z <- arima.sim(n = 200, list(ar = 0.6))
acf(z)
```



```
ar.fun <- function(fun) {  
    ar.fit <- ar(fun, order.max = 1)  
    c(ar.fit$ar[1])  
}  
prova <- tsboot(z, ar.fun, R = 1000, l = 6, sim = "fixed")  
prova$t0
```

```
## [1] 0.5804858
```

```
ar.fun(z)
```

```
## [1] 0.5804858
```

```
prova$t[1:5]
```

```
## [1] 0.4245009 0.4276346 0.4033080 0.4816728 0.4838219
```

```
sd(prova$t)
```

```
## [1] 0.05240388
```

# **Now it's your turn!!!**