2024 SISBID High-Dimensional Hypothesis Testing Lab

Genevera I. Allen and Yufeng Liu

Load Packages

```
library(sda)
library(ggplot2)
```

H_0: feature is not associated with the response.## Data set 1 - Simulated Data

Small simulated data set to demonstrate multiple testing when all null hypthesis hold.

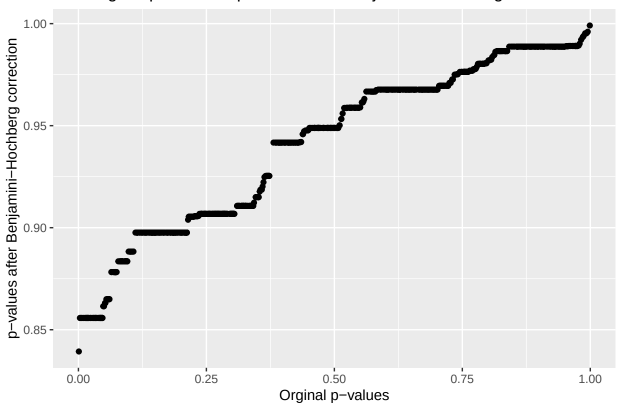
```
#simulate data
x <- matrix(rnorm(1000*50),ncol=50)
y <- sample(c(0,1),50,rep=TRUE)
ps <- NULL
for(i in 1:1000){
   ps <- c(ps,t.test(x[i,y==0],x[i,y==1])$p.value)
}
cat("Around 5% of p-values are below 0.05:",mean(ps<.05),fill=TRUE)</pre>
```

Around 5% of p-values are below 0.05: 0.058

Benjamini-Hochberg Algorithm for FDR Control

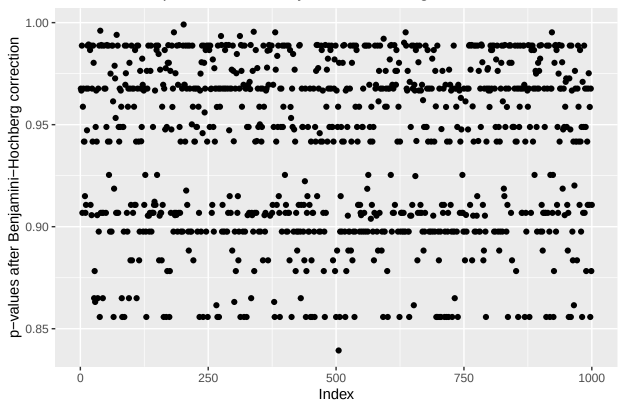
```
fdrs.bh <- p.adjust(ps, method="BH")
BHData = data.frame(cbind(ps,fdrs.bh))
colnames(BHData) = c("OriginalP","BH.P")
ggplot(BHData) +
   geom_point(mapping = aes(x = OriginalP, y = BH.P)) +
   ggtitle("Original p-value vs p-value after Benjamini-Hochberg correction") +
   theme(plot.title = element_text(hjust = 0.5)) +
   xlab("Orginal p-values") + ylab("p-values after Benjamini-Hochberg correction")</pre>
```

Original p-value vs p-value after Benjamini-Hochberg correction



```
BHData$index = 1:nrow(BHData)
ggplot(BHData) +
  geom_point(mapping = aes(x = index, y = BH.P)) +
  ggtitle("p-value after Benjamini-Hochberg correction") +
  theme(plot.title = element_text(hjust = 0.5)) +
  xlab("Index") + ylab("p-values after Benjamini-Hochberg correction")
```





Data set 2 - Simulated Data

Small simulated data set to demonstrate multiple testing when **not all null hypthesis hold**.

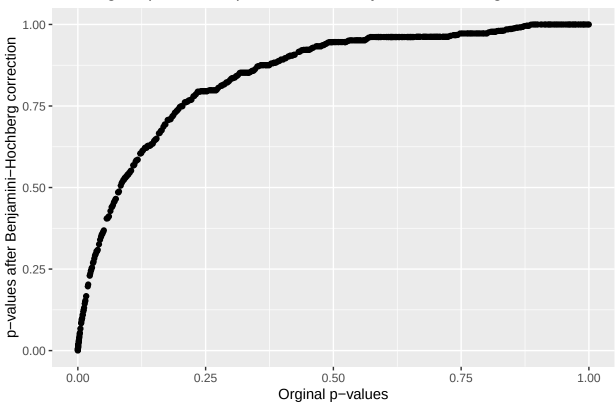
```
#simulate data
x <- matrix(rnorm(1000*50),ncol=50)
y <- sample(c(0,1),50,rep=TRUE)
x[1:100,y==0] <- x[1:100,y==0] + 1
ps <- NULL
for(i in 1:1000) {
   ps <- c(ps,t.test(x[i,y==0],x[i,y==1])$p.value)
}
cat("Way more than 5% of p-values are below 0.05:",mean(ps<.05),fill=TRUE)</pre>
```

Way more than 5% of p-values are below 0.05: 0.136

```
fdrs.bh <- p.adjust(ps, method="BH")
# plot

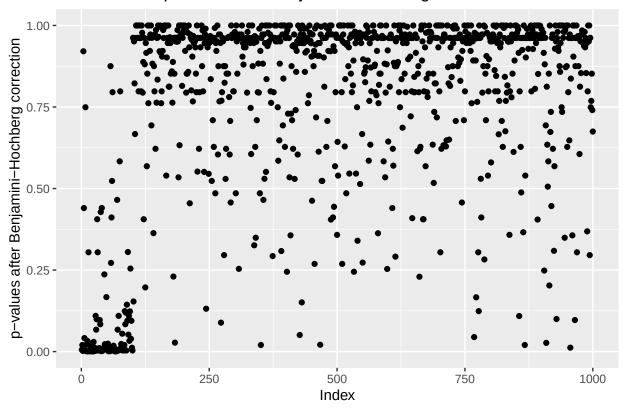
BHData = data.frame(cbind(ps,fdrs.bh))
colnames(BHData) = c("OriginalP","BH.P")
ggplot(BHData) +
  geom_point(mapping = aes(x = OriginalP, y = BH.P)) +
  ggtitle("Original p-value vs p-value after Benjamini-Hochberg correction") +
  theme(plot.title = element_text(hjust = 0.5)) +
  xlab("Orginal p-values") + ylab("p-values after Benjamini-Hochberg correction")</pre>
```

Original p-value vs p-value after Benjamini-Hochberg correction



```
BHData$index = 1:nrow(BHData)
ggplot(BHData) +
  geom_point(mapping = aes(x = index, y = BH.P)) +
  ggtitle("p-value after Benjamini-Hochberg correction") +
  theme(plot.title = element_text(hjust = 0.5)) +
  xlab("Index") + ylab("p-values after Benjamini-Hochberg correction")
```

p-value after Benjamini-Hochberg correction



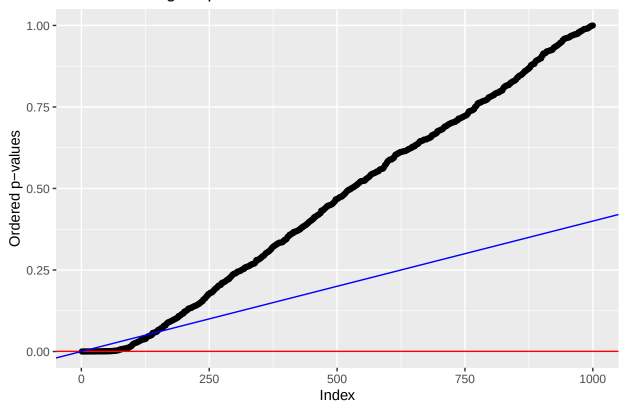
```
cat("Number of Tests with FDR below 0.4:",sum(fdrs.bh<0.4), fill=TRUE)</pre>
```

Number of Tests with FDR below 0.4: 138

Compute the BH FDR Directly: 138

```
BHData = BHData[order(ps,decreasing = FALSE),]
BHData$index = 1:nrow(BHData)
# plot
ggplot(BHData) +
    geom_point(mapping = aes(x = index, y = OriginalP)) +
    ggtitle("Original p-values with different correction methods") +
    geom_abline(intercept = 0.4/1000,slope = 0,col= "red") + #Bonferroni
    geom_abline(intercept = 0 ,slope = 0.4/1000,col= "blue") + #BH procedure
    theme(plot.title = element_text(hjust = 0.5)) +
    xlab("Index") + ylab("Ordered p-values")
```

Original p-values with different correction methods



Data set 3, Real Data: Prostate Data (Singh et al. 2002). This data set consists of gene expression levels for 6033 genes among 102 men.

The dataset is available from the R package "sda"

- * Problem 1 We wish to identify important genes to differentiate cancer or healthy patients. What kind of tests are reasonable?
- * Problem 2 In order to adjust for multiple comparisons, which procedures should one use?
- * Problem 3 Examine the list of genes identified.

```
## import data
data(singh2002)
x = singh2002$x
y = singh2002$y

n1 = sum(y == "healthy")
n2 = length(y) - n1
```

```
ps<-NULL
for(i in 1:ncol(x)) {
   ps <- c(ps, t.test(x[1:n1,i], x[(n1+1):(n1+n2),i])$p.value)
}
## ordered p-values names(ps)<-seq(1,ncol(x),1)
p1 =sort (ps)

## plot ordered p-values
plot(p1[1:100], pch=rep('*',100),ylim=c(0,0.003), ylab="ordered p-values")</pre>
```

```
## rejection boundry of Benjamini-Hochberg's procedure
abline(a=0, b=0.1/ncol(x), col="red")
## rejection boundary of Bonferroni at 0.1
abline(a=0.1/ncol(x), b=0, col="blue",lty=5)

cat("Compute the no. rejection by Bonferroni:",
    max(which(sort(ps,decreasing=FALSE) < .1/ncol(x))), fill=TRUE)</pre>
```

Compute the no. rejection by Bonferroni: 6

Compute the BH FDR Directly: 57

