

# A Bootstrap Based Multiple Hypothesis Testing Procedure

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We propose a new methodology for multiple hypothesis testing based on bootstrap distribution of p values. We know that under null hypothesis p value is an uniform(0, 1) random variable. If the hypothesis are independent, then so are the p values. We use this intuition to construct a statistic that can be used to test multiple hypothesis.

We consider the case of coordinate wise testing of a data vector. If the data matrix  $X$  is of dimension  $I \times J$  then without loss of generality, we consider the following set of hypothesis for  $j = 1, 2, \dots, J$ .

$$\begin{aligned} H_0^{(j)} &: x_j = 0 \\ H_1^{(j)} &: x_j \neq 0 \end{aligned}$$

Where  $x_j$  denotes the  $j$ th coordinate of  $X$ . This set of tests can be performed in a variety of ways, for the purpose of illustration, we choose to use one sample t-tests.

The general procedure for testing multiple hypothesis is testing independently then use a correction method to correct for family wise error rate. Here we are not using any such correction, as we are using a bootstrap based approach.

We first generate bootstrap samples of our data, let us denote the bootstrap samples by  $X^{\{b\}}$  for  $b \in \{1, 2, \dots, B\}$ , where  $B$  is the bootstrap sample size. For each of these samples we can perform  $J$  tests in parallel and collect the p values. We denote the p values from  $j$ th coordinate of  $b$ th bootstrap sample by  $p_j^{\{b\}}$ . We then use a monotone transformation of the p values for better visualizing. The transformed order statistics are collected as shown.

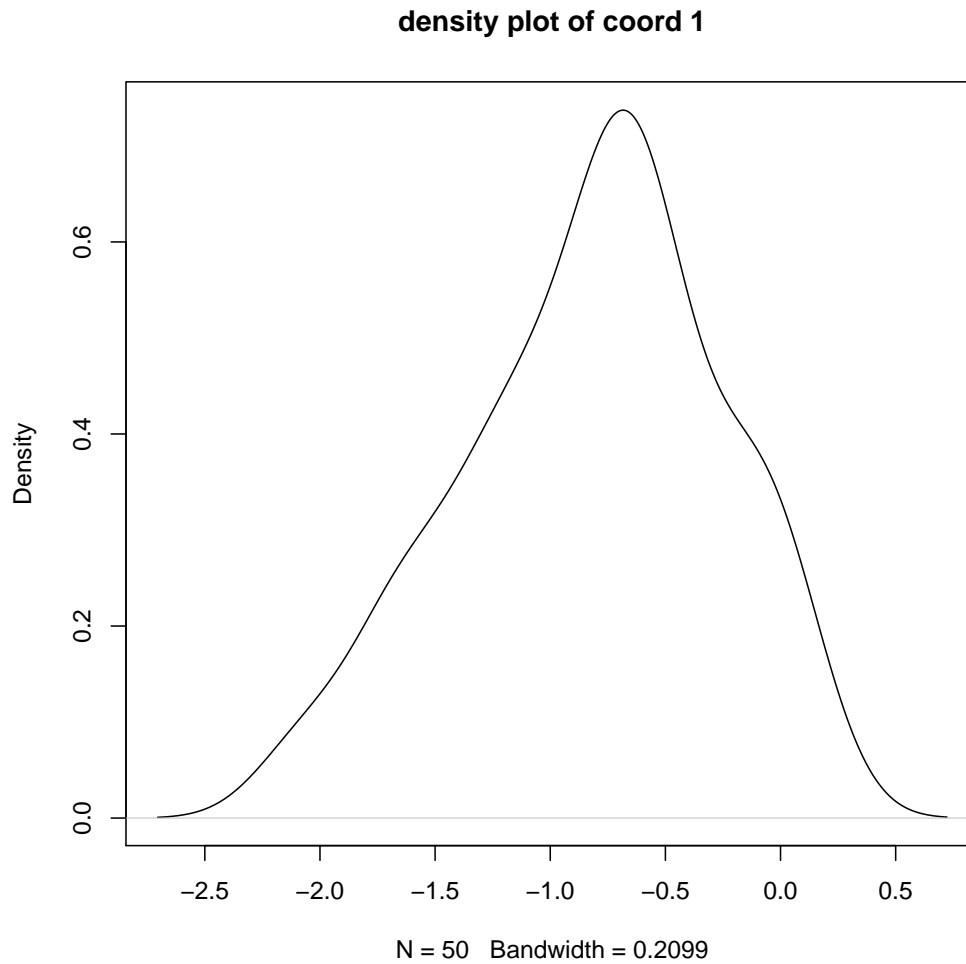
$$\begin{aligned}
Z_{(0)}^{\{b\}} &\stackrel{\text{def}}{=} 0 \\
Z_{(j)}^{\{b\}} &= -\log(1 - p_j^{\{b\}}) \\
W_j^{\{b\}} &\stackrel{\text{def}}{=} Z_{(j)}^{\{b\}} - Z_{(j-1)}^{\{b\}} - (n + 1 - j)^{-1}
\end{aligned}$$

We demonstrate the procedure below with a simulated example. The function `datgen` is used to simulate data from a multivariate normal distribution.

```

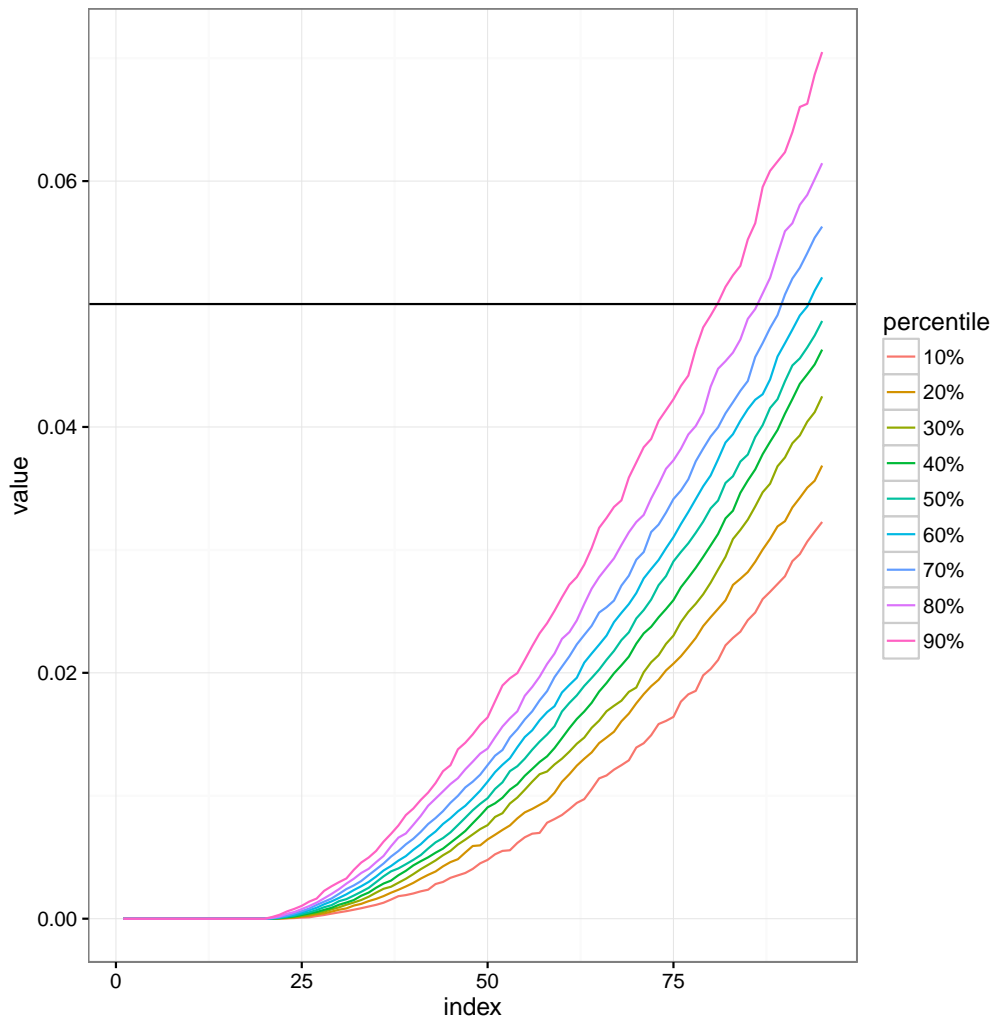
set.seed(12345)
suppressMessages(library(mhtboot))
n = 50;m = 500;m0 = 20;
sigeff = 1;
Sigma <- 0.25*diag(m)
X <- datgen(n,m,m0,sigeff,Sigma = Sigma)
plot(density(X[,1]),main="density plot of coord 1")

```



We then generate the distribution of the p values using bootstrap. This is implemented in the function series `pboot`. Here we are using the one sample version of the function. The `plotpboot` function is used to generate the quantile plot of the distribution of the p values.

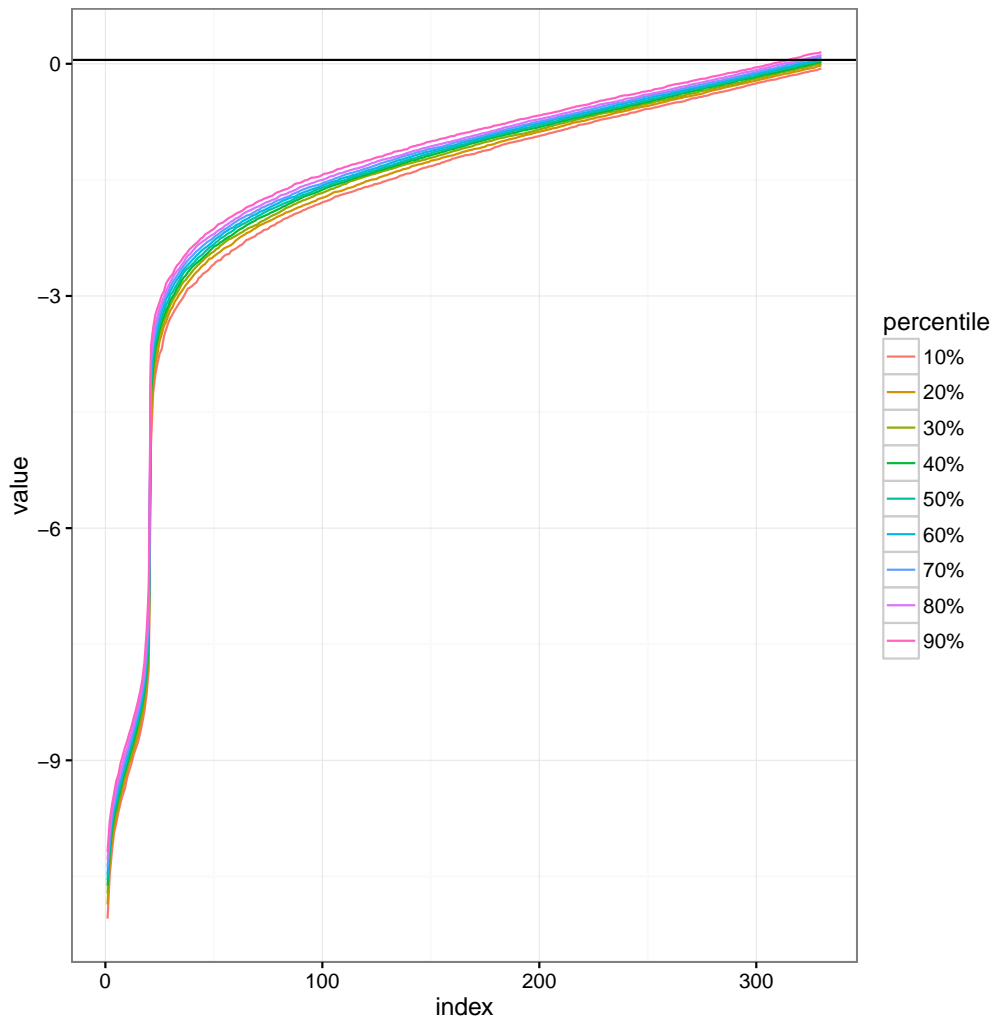
```
porder <- pboot.1sample(X=X,B=250,ncpus = 1)
plotpboot(porder = porder)
```



This approach can be extended to any set of hypothesis, in the package we also provide a function for two sample tests. Both of these functions can be used for user given tests as they accept test statistic as a parameter.

We can transform the order statistics of the p values using a monotone transformation. We show here the transformation using inverse normal cdf function.

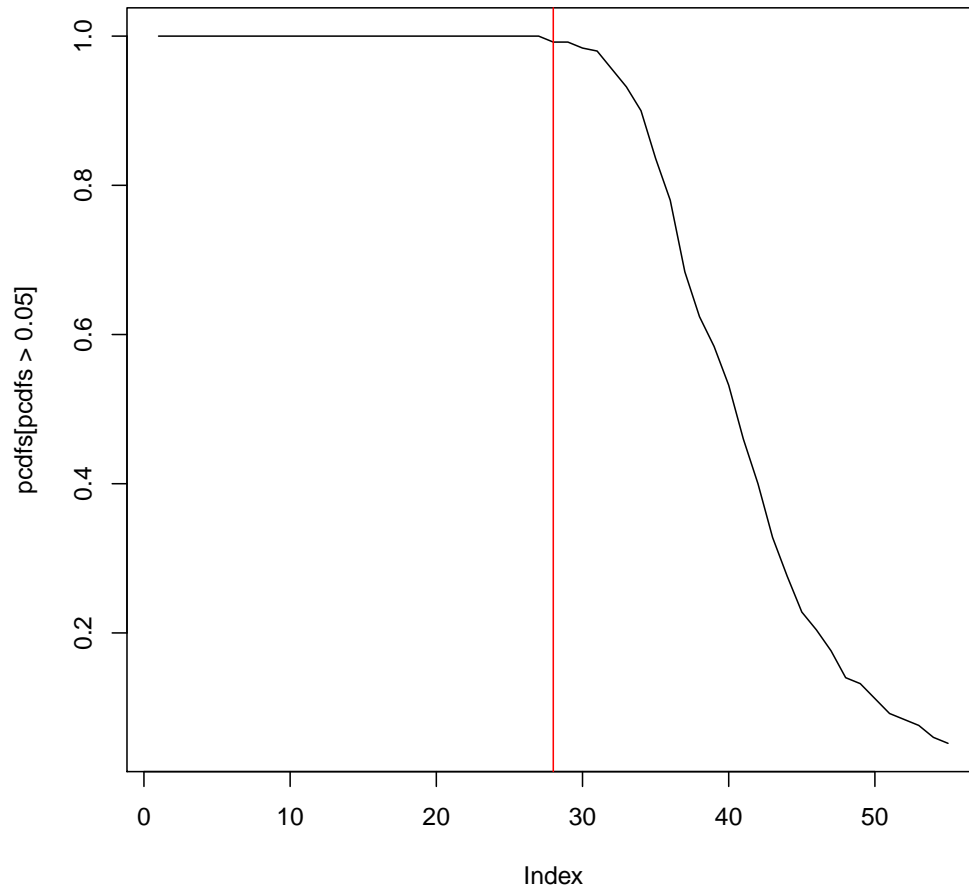
```
porder.tr <- ptrans(porder, trans="normal")
plotpboot(porder.tr)
```



We can also look at the points where each coordinates hit a certain probability.

```
porder <- ptrans(porder = porder)
hitplots(porder = porder, alpha = 0.005)

## [1] 28
```



Once we have the distribution of the p values, we can use them to detect the change point in their distribution. This is done through qelbow function.

```

out <- qelbow(porder = porder)
out

## dav dlm
## 34 21

```

All the above process is implemented in one single function for all one sample tests. The final cut off point is chosen by taking a minimum of two

detection methods.

```
out1 <- mht.1sample(X,ncpus = 1)
out1$cutoff

## [1] 26

out1$signal

## [1] 13 20 23 26 33 47 54 61 70 108 111 116 131 162 205 219 276 387 43
## [22] 462 467 471 477 494
```