

Comparison between different local tests: Simes, Simes with Storey and Wilcoxon-Mann-Whitney

m=20

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The aim is to compare the performance of three closed testing procedures, which respectively use Simes local test with and without Storey estimator for the proportion of true null hypotheses and Wilcoxon-Mann-Whitney local test.

We consider a null distribution F from which inliers come and we consider outliers which come from the alternative distribution

$$F^k$$

with $k > 0$, especially if $k \in \mathbb{N}_{>0}$ we know that F^k is the distribution of the random variable defined as the maximum of k observations drawn from F . So, we consider a sample drawn from the mixture distribution

$$G = (1 - \theta)F + \theta F^k$$

where $\theta \in [0, 1]$ is the proportion of inliers.

Since we deal with conformal p -values, we are interested not really in the sample distribution, but rather in the scores sample distribution. In our simulation study we draw n observations for the train set, l for the calibration set and mk for the test set. All observations are drawn from a d -multivariate standard normal distribution with $d = 3$ and using the algorithm of *isolation forest* trained on the training samples we compute the scores for the calibration and test samples. In order to generate scores associated to outlier observations, we consider the m blocks of k observations which create the test set. For each block i with $i = 1, \dots, m$, we draw from a Bernoulli random variable with success probability equal to θ . If the value is 1 then the i -th observation of the test set is inlier and we randomly sample its score value from the k scores of the i -th block, otherwise it is an outlier and its score value will be the maximum of the scores of the i -th block.

```
library(Cairo)
library(nout)

scores_from_mixture = function(k, raw_scores, m, m1){

  if(m1==0) scores = raw_scores

  if(m1==m){
    scores = sapply(0:(m1-1), function(i) max(raw_scores[(i*k+1):(i*k+k)]))
  }

  if(0<m1 & m1<m){
    scores.out = sapply(0:(m1-1), function(i) max(raw_scores[(i*k+1):(i*k+k)]))
    scores.in = raw_scores[(k*m1+1):length(raw_scores)]
    scores = c(scores.out, scores.in)
  }
}
```

```

    return("scores"=scores)
}

simuLMPI = function(B=10^4, n, l, m, d = 3, k = 2, theta, alpha = m/(l+1)){

  train = mvtnorm::rmvnorm(n=n, mean=rep(0,d))
  iso.fo = isotree::isolation.forest(train, ndim=d, ntrees=10, nthreads=1,
                                     scoring_metric = "depth", output_score = TRUE)

  crit=critWMW(m=m, n=n, alpha=alpha)
  m1 = round(theta*m)
  m0 = m-m1

  d_WMW = rep(0,B)
  d_Simes = rep(0,B)
  d_StoSimes = rep(0,B)
  d_BH = rep(0,B)
  d_StoBH = rep(0,B)

  for(b in 1:B){
    cal = mvtnorm::rmvnorm(n=1, mean=rep(0,d))
    te = mvtnorm::rmvnorm(n=k*m1+m0, mean=rep(0,d))

    S_cal = isotree::predict.isolation_forest(iso.fo$model, cal, type = "score")
    rawS_te = isotree::predict.isolation_forest(iso.fo$model, te, type = "score")
    S_te = scores_from_mixture(k=k, raw_scores=rawS_te, m=m, m1=m1)

    d_WMW[b] = d_mannwhitney(S_X=S_cal, S_Y=S_te, crit=crit)
    d_Simes[b] = d_Simes(S_X=S_cal, S_Y=S_te, alpha=alpha)
    d_StoSimes[b] = d_StoreySimes(S_X=S_cal, S_Y=S_te, alpha=alpha)
    d_BH[b] = d_benjhoch(S_X=S_cal, S_Y=S_te, alpha=alpha)
    d_StoBH[b] = d_StoreyBH(S_X=S_cal, S_Y=S_te, alpha=alpha)
  }

  discov = as.data.frame(cbind("d_BH"=d_BH, "d_StoBH"=d_StoBH, "d_Simes"=d_Simes,
                              "d_StoSimes"=d_StoSimes, "d_WMW"=d_WMW))
  colnames(discov) = c("BH", "BHSto", "CTSim", "CTSimSto", "CTWMW")
  mean.discov = apply(discov, MARGIN = 2, FUN = mean)

  powerGlobalNull = as.data.frame(cbind("d_BH"=d_BH>0, "d_StoBH"=d_StoBH>0, "d_Simes"=d_Simes>0,
                                         "d_StoSimes"=d_StoSimes>0, "d_WMW"=d_WMW>0))
  colnames(powerGlobalNull) = c("BH", "BHSto", "CTSim", "CTSimSto", "CTWMW")
  mean.powerGlobalNull = apply(powerGlobalNull, MARGIN = 2, FUN = mean)

  return(list("discoveries"=discov, "mean.discoveries" = mean.discov,
             "powerGlobalNull"=powerGlobalNull, "mean.powerGlobalNull"=mean.powerGlobalNull,
             "theta"=theta, "alpha"=alpha))
}

```

K=2

```
set.seed(321)

# Initializing parameters
B=10^5
n = 199
l = 199
m = 20
d = 3
k = 2
alpha = m/(l+1)
m1s = seq(from=0, to=m, by=1)
thetas = m1s/m

# Results
res = lapply(thetas, function(theta) simulMPI(B=B, n=n, l=l, m=m, d = d,
                                              k = k, theta, alpha = m/(l+1)))

# Storing results
store_res = list("mean.discov" = matrix(nrow=length(thetas), ncol = 5),
                 "mean.powerGlobalNull" = matrix(nrow=length(thetas), ncol = 5))
row.names = rep(NA, times=length(thetas))
for(i in 1:length(thetas)){
  row.names[i] = paste("theta =", thetas[i])
}
rownames(store_res$mean.discov) = row.names
colnames(store_res$mean.discov) = c("BH", "StoBH", "Simes", "StoSimes", "WMW")
rownames(store_res$mean.powerGlobalNull) = row.names
colnames(store_res$mean.powerGlobalNull) = c("BH", "StoBH", "Simes", "StoSimes", "WMW")

for(i in 1:length(res)){
  store_res$mean.discov[i,] = res[[i]]$mean.discov
  store_res$mean.powerGlobalNull[i,] = res[[i]]$mean.powerGlobalNull
}

store_res$mean.discov
```

##		BH	StoBH	Simes	StoSimes	WMW
##	theta = 0	0.13356	0.09157	0.11082	0.06563	0.21213
##	theta = 0.05	0.14100	0.10234	0.11524	0.07262	0.26859
##	theta = 0.1	0.14899	0.11757	0.12114	0.08232	0.32724
##	theta = 0.15	0.15389	0.13257	0.12440	0.09150	0.40271
##	theta = 0.2	0.16579	0.15410	0.13340	0.10454	0.49208
##	theta = 0.25	0.17264	0.17324	0.13706	0.11488	0.59937
##	theta = 0.3	0.18608	0.20098	0.14583	0.13120	0.72182
##	theta = 0.35	0.19511	0.23312	0.15234	0.14765	0.85528
##	theta = 0.4	0.20225	0.25870	0.15663	0.16016	1.03694
##	theta = 0.45	0.21792	0.30215	0.16842	0.18271	1.22042
##	theta = 0.5	0.22351	0.34121	0.17100	0.19780	1.42025
##	theta = 0.55	0.23429	0.38396	0.17711	0.21653	1.64145
##	theta = 0.6	0.24546	0.45028	0.18376	0.24067	1.92308

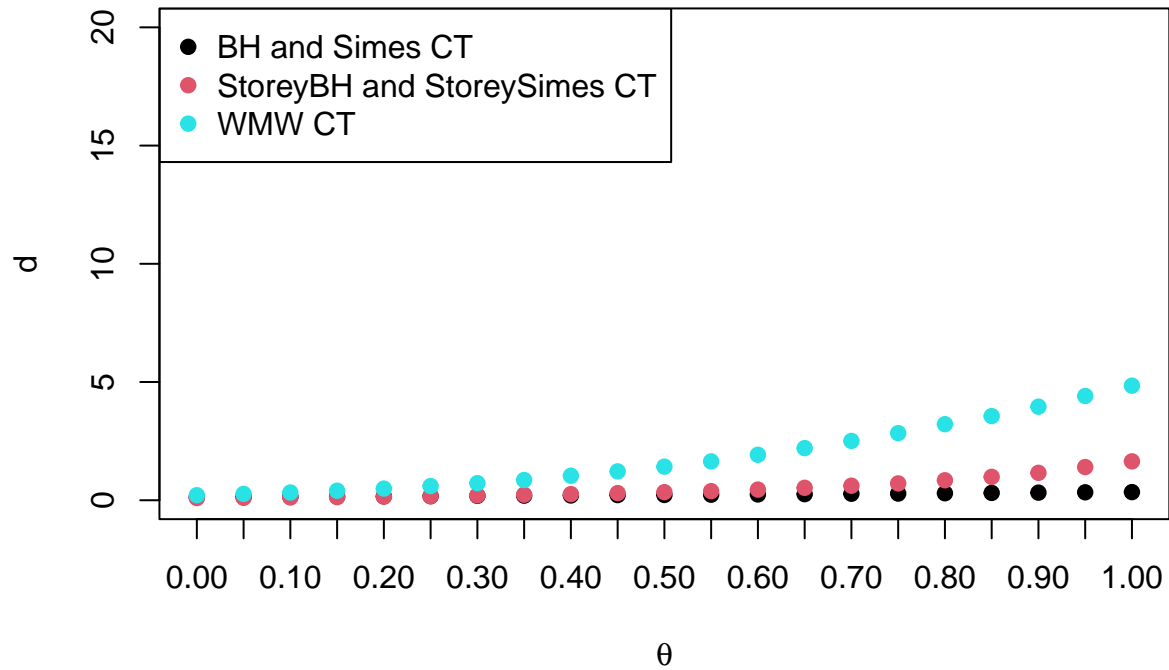
```
## theta = 0.65 0.25760 0.52609 0.18959 0.26993 2.20497
## theta = 0.7 0.27530 0.61463 0.20113 0.30469 2.50856
## theta = 0.75 0.28149 0.71146 0.20527 0.33734 2.83785
## theta = 0.8 0.29599 0.84480 0.21384 0.38104 3.21570
## theta = 0.85 0.31002 0.98900 0.21952 0.42882 3.55927
## theta = 0.9 0.32099 1.15951 0.22669 0.48246 3.95647
## theta = 0.95 0.33594 1.40212 0.23521 0.55729 4.40942
## theta = 1 0.34395 1.64439 0.24006 0.63501 4.84623
```

```
store_res$mean.powerGlobalNull
```

```
##          BH    StoBH    Simes StoSimes    WMW
## theta = 0    0.09980 0.05741 0.09980 0.05741 0.10159
## theta = 0.05 0.10383 0.06301 0.10383 0.06301 0.12445
## theta = 0.1 0.10862 0.07142 0.10862 0.07142 0.14879
## theta = 0.15 0.11100 0.07839 0.11100 0.07839 0.17604
## theta = 0.2 0.11858 0.08931 0.11858 0.08931 0.20884
## theta = 0.25 0.12181 0.09746 0.12181 0.09746 0.24668
## theta = 0.3 0.12854 0.11009 0.12854 0.11009 0.28695
## theta = 0.35 0.13295 0.12125 0.13295 0.12125 0.32800
## theta = 0.4 0.13659 0.13090 0.13659 0.13090 0.38177
## theta = 0.45 0.14550 0.14627 0.14550 0.14627 0.43038
## theta = 0.5 0.14773 0.15645 0.14773 0.15645 0.47972
## theta = 0.55 0.15237 0.16841 0.15237 0.16841 0.53285
## theta = 0.6 0.15602 0.18200 0.15602 0.18200 0.58794
## theta = 0.65 0.16042 0.19815 0.16042 0.19815 0.64008
## theta = 0.7 0.16979 0.21676 0.16979 0.21676 0.69153
## theta = 0.75 0.17205 0.23239 0.17205 0.23239 0.74023
## theta = 0.8 0.17830 0.25447 0.17830 0.25447 0.78702
## theta = 0.85 0.18177 0.27217 0.18177 0.27217 0.82263
## theta = 0.9 0.18711 0.29360 0.18711 0.29360 0.86126
## theta = 0.95 0.19359 0.32127 0.19359 0.32127 0.89091
## theta = 1 0.19628 0.34326 0.19628 0.34326 0.91813
```

```
plot(x = thetas, y = store_res$mean.discov[,1], col = 1, ylab = "d",
     xlab = expression(theta), ylim=c(0,m), pch=19, xaxt = "n",
     main = "Mean of the number of discoveries on B replications")
points(x = thetas, y = store_res$mean.discov[,2], col = 2, pch=19)
points(x = thetas, y = store_res$mean.discov[,5], col = 5, pch=19)
axis(side = 1, at = thetas)
legend("topleft", pch = 19, col = c(1,2,5),
      legend = c("BH and Simes CT", "StoreyBH and StoreySimes CT", "WMW CT"))
```

Mean of the number of discoveries on B replications



```
plot(x = thetas, y = store_res$mean.powerGlobalNull[,1], col = 1, ylab = "power",
     xlab = expression(theta), ylim=c(0,m), pch = 19, xaxt = "n",
     main = "Mean of the power on B replications")
points(x = thetas, y = store_res$mean.powerGlobalNull[,2], col = 2, pch=19)
points(x = thetas, y = store_res$mean.powerGlobalNull[,5], col = 5, pch=19)
abline(h=alpha, col = "darkgreen")
text(x=0.05, y=alpha+0.05, labels = paste("\u03B1", "=", alpha), col = "darkgreen")
axis(side = 1, at = thetas)
legend("topleft", pch = 19, col = c(1,2,5),
      legend = c("BH and Simes CT", "StoreyBH and StoreySimes CT", "WMW CT"))
```

Mean of the power on B replications

