simTool

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1 Note

This Sweave File is part of μ Toss package. But because some parts of this document are computational intensiv the extension of the Sweave file is ".actual-lyRnw".

2 Introduction

This document will give an introduction to the use of simTool. We will start with a very simple application then raise the degree of complexity in a few steps and in the end reproduce some of the results from Benjamini, Krieger, Yekutieli (2006). Basically for a simulation we need the following things

- 1. a function that generates the data (for example p-values)
- some procedures that evaluates the generated data (for example the bonferroni correction)
- statistics we want to calculate. For example the the false discovery proportion (FDP).

If we are speak for example of 1000 replications, we mean that these 3 steps were repeated 1000 times. That means, after 1000 replications there have been 1000 data sets generated. Every procedure was applied to these 1000 data sets. If we have specified for example 3 procedures then every of these procedures will give us an output, this means 3000 "output objects". And every specified statistic is applied to these 3000 "output objects".

2.1 Motivation

2.1.1 Example 1

Suppose we want to compare some characteristics of the two procedures BH and holm. Denote by $V_n(BH)$ and $V_n(holm)$ the number of true hypotheses rejected. For example we are interested in the distribution of $V_n(BH)$ and $V_n(holm)$ and also in $E[V_n(BH)]$ and $E[V_n(holm)]$. Lets have a look on the arguments of theses procedures:

> args(BH)

```
function (pValues, alpha, silent = FALSE)
NULL
```

> args(holm)

```
function (pValues, alpha, silent = FALSE)
NULL
```

Both have the argument alpha, which stands for the niveau of the corresponding error measure. BH controls the false discovery rate (FDR) and holm controls the family-wise error rate (FWER). One question may be how must alpha be set in holm such that for fixed alpha = 0.1 in BH we have $E[V_n(BH)] \approx E[V_n(lndm)]$? And perhaps we are also interested in how a dependence structure affects the distribution of $V_n(BH)$ and $V_n(lndm)$.

2.1.2 Example 2

Suppose you have developed a new procedure controlling some error measure and you want to compare this new procedure with some other already established procedures. Then a simulation study will consist of creating data and gathering statistics for different sample sizes, dependence structures and parameter constellations.

Basically it is always the same story. Data is generated by a specific function, perhaps this depends on some parameters, and we want to apply different procedures, again depending on some parameters, to this generated data. Nearly everyone will say, "don't bother me with the details of programming, just do the simulation with the above information and give me a nice data.frame which I can analyze." And this is exactly the purpose of the simTool.

Data generating function

MIII I

For now we only want to use the procedures BH and holm. Again, lets have a look on the arguments of these procedures:

```
> args(BH)
function (pValues, alpha, silent = FALSE)
NULT.
> args(holm)
function (pValues, alpha, silent = FALSE)
```

Not much has to be specified. The parameter alpha is the level at which the corresponding error rate should be controlled and pValues are the p-values of the hypotheses that should be tested. In general, if we say that we reject a p-value this means that we reject the corresponding hypotheses. The following function generates data and will be used throughout the whole document. The data generating function must have an entry \$procInput. All in \$procInput will be used as input for the specified procedures. In our situation we will generate a list with 2 entries. \$procInput will only consist of the generated p-values . And \$groundTruth indicates which p-value corresponds to a true or false hypothesis.

```
> pValFromEquiCorrData <- function(sampleSize, rho, mu, pi0) {
     nmbOfFalseHyp <- round(sampleSize * (1 - pi0))
     nmbOfTrueHvp <- sampleSize - nmbOfFalseHvp
     muVec <- c(rep(mu, nmbOfFalseHyp), rep(0, nmbOfTrueHyp))
     Y <- sqrt(rho) * rnorm(1) + sqrt(1 - rho) * rnorm(sampleSize) +
          muVec
     return(list(procInput = list(pValues = 1 - pnorm(Y)), groundTruth = muVec ==
         0))
+ }
```

2.2.1 Illustration of generated Data

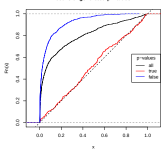
We will now generate 1000 p-Values independently (rho = 0). sampleSize * (1-pi0) = 700 of them correspond to false hypotheses and 300 to true hypothe-202

```
> set.seed(123)
> data <- pValFromEquiCorrData(sampleSize = 1000, rho = 0, mu = 2,
      pi0 = 0.3)
```

Lets visualize the different p-values .

```
> local({
     pValues = data$procInput$pValues
     groundTruth = data$groundTruth
     plot(ecdf(pValues), do.points = FALSE, verticals = TRUE,
          main = "ecdf's of generated p-values")
```

ecdf's of generated p-values



2.3 A very simple example

Lets directly start with the function call and then analysing what happend. Just for now we implement a simple version of the data generating function in section 2.2 and a simple version of the BH.

The following happend:

- 1. Call myGen
- 2. Append generated data set to sim\$data
- 3. Call BH.05 with the generated p-values
- Add to the results of BH.05 the parameter constellation used by myGen and BH.05 and also the position of the used data in sim\$data
- 5. Append this extended results of BH.05 to sim\$results
- 6. repeat this 9 more times

Since replications = 10 in simulation the object sim consists of:

- > names(sim)
- [1] "data" "results"
- > length(sim\$data)
- [1] 10
- > length(sim\$results)
- [1] 10

The structure of one object in sim\$data coincides with the structure of the return of myGen:

- > names(myGen())
 - [1] "procInput" "groundTruth"
 - > names(sim\$data[[1]])
 - [1] "procInput" "groundTruth"

Lets have a look at the additional information added by simulation to the object returned by BH.05. First the entries of the pure return value of BH.05 and then of sim\$results

- > names(BH.05(runif(100)))
- [1] "adjPValues" "criticalValues" "rejected" "errorControl"
- > names(sim\$results[[1]])
- [1] "data.set.number" "parameters" "adjPValues" "criticalValues"
- [5] "rejected" "errorControl"

data.set.number is the number of the used data set in sim\$data. Since every generated data set is used only once we have

- > sapply(sim\$results, function(x) x\$data.set.number)
- [1] 1 2 3 4 5 6 7 8 9 10

Thus it is possible to reproduce any result directly. Let us reproduce the results of the 6th replication

```
> idx <- sim%results[[6]]$data.set.number

> pValues <- sim%data[[idx]]$procInput$pValues

> all(BH.05(pValues)$adjPValues == sim%results[[6]]$adjPValues)

fil TRUE
```

Since our data generating function and procedure did not really have any parameter, there is not much information in:

> sim\$results[[1]]\$parameters

\$funName

[1] "myGen"

\$method

[1] "BH.simple"

Note, BH.05 technically has the parameter pValues but since this parameter is the "input channel" for the output data of the data generating function it is not really regarded as parameter that affect the behaviour of the procedure like for example the level α for the error measure.

2.4 A simple example

Indeed, for the simple example above the simTool is not very helpful. Let us increase the complexity of our simulation. As a first step, we want to increase the parameter rho of the data generating function gradually and investigate $EV_n(BH)$ and $EV_n^2(BH)$.

```
> set.seed(I23)

> sim <- simulation(replications = 10, list(funName = "pVGen",

+ fun = pValFromEquiCorrData, sampleSize = 100, rho = seq(0,

+ 1, by = 0.2), mu = 2, pio = 0.5), list(list(funName = "BH",

+ fun = BH, alpha = (0.5, 0.25), silent = TREU,
```

The following happend:

- 1. Call myGen with rho = 0
- Append generated data set to sim\$data
- Call BH with alpha = 0.5 and the generated p-values
- Add to the results the parameter constellation used by myGen and BH and also the position of the used data in sim\$data
- 5. Append this extended results to sim\$results
- Call BH with alpha = 0.25 and the same p-values as in 3.
- Add to the results the parameter constellation used by myGen and BH and also the position of the used data in sim\$data

- 8. Append this extended results of to sim\$results
- 9, repeat this 9 more times
- Call myGen with rho = 0.2
- 11. and so on

So we have now a bunch of data sets and results.

> length(sim\$data)

[1] 60

> length(sim\$results)

Γ17 120

60 data sets have been generated because we have rho = 0, 0.2, 0.4, 0.6, 0.8, 1 and for each rho we generated 10 data sets. We have applied BH with alpha = 0.5 to all 60 data sets and BH with alpha = 0.25, yielding 120 objects in results. We see from the data.set.number that after generating a data set it is used two times in series

> sapply(sim\$results, function(x) x\$data.set.number)

```
[1] 1 1 2 2 3 3 4 4 5 5 6 6 7 7 8 8 8 9 9 10 10 11 11 12 12 13 [26] 13 14 14 15 15 16 16 17 17 18 18 19 19 20 20 21 21 22 22 23 23 24 24 25 25 [51] 26 26 27 27 28 28 29 29 30 30 31 31 32 32 33 33 34 34 35 35 36 36 37 37 38 67 38 16 38 39 40 40 41 41 42 42 43 43 44 44 45 45 46 46 47 47 48 48 49 49 50 50
```

[101] 51 51 52 52 53 53 54 54 55 55 56 56 57 57 58 58 59 59 60 60

In order to gather how many true hypotheses have been rejected corresponding to the different parameter constellations we need a function that is able to calculate it.

- > NumberOfType1Error <- function(data, result) sum(data\$groundTruth *
 + result\$rejected)</pre>
- > V2 <- function(data, result) NumberOfType1Error(data, result)^2

Lets us calculate for the first result object the number of rejected true hypotheses explicitly.

```
> idx = sim$results[[1]]$data.set.number
> data = sim$data[[idx]]
> NumberOfType1Error(data, sim$results[[1]])
```

wumberorryperError(data, Simpresurts[[1]]

[1] 15

This means for the following parameter constellation

> sim\$results[[1]]\$parameters

\$funName

[1] "pVGen"

\$sampleSize

[1] 100

\$rho

[1] 0

\$mu

[1] 2

\$pi0

[1] 0.5

\$method

[1] "BH"

\$alpha

[1] 0.5

\$silent

[1] TRUE

we one time observed

[1] 15

rejected true null hypotheses. In order to estimate $EV_n(BH)$ and $EV_n^2(BH)$ we have to search in sim\$results for the other 9 results using the same parameter constellation. In order to facilitate this task we provide a function.

```
> result <- gatherStatistics(sim, list(V = NumberOfType1Error,
+ V2 = V2), mean)
> result
```

\$statisticDF

	funName	sampleSize	rho	mu	pi0	method	alpha	silent	V.mean	V2.mean
1	pVGen	100	0	2	0.5	BH	0.5	TRUE	15.1	245.3
2	pVGen	100	0	2	0.5	BH	0.25	TRUE	5.9	40.9
3	pVGen	100	0.2	2	0.5	BH	0.5	TRUE	21.5	591.9
4	pVGen	100	0.2	2	0.5	BH	0.25	TRUE	9.2	163.2
5	pVGen	100	0.4	2	0.5	BH	0.5	TRUE	14.6	401.6
6	pVGen	100	0.4	2	0.5	BH	0.25	TRUE	6.2	126.6
7	pVGen	100	0.6	2	0.5	BH	0.5	TRUE	22.3	860.7
8	pVGen	100	0.6	2	0.5	BH	0.25	TRUE	9.3	265.7
9	pVGen	100	0.8	2	0.5	BH	0.5	TRUE	18.1	711.1
10	pVGen	100	0.8	2	0.5	BH	0.25	TRUE	8.0	328.6
11	pVGen	100	1	2	0.5	BH	0.5	TRUE	20.0	1000.0
12	pVGen	100	1	2	0.5	BH	0.25	TRUE	15.0	750.0

\$name.statistics

[1] "V.mean" "V2.mean"

\$name.avgFun

[1] "mean"

As you can see every, parameter constellation has its own row in **\$statisticDF**. By the way, again, we see that p-values is a parameter of BH but since it is contained in **\$procInput** it is not considered as "real parameter".

If we are interested in more than one statistics, then we simply provide a list with "average functions"

```
> result <- gatherStatistics(sim, list(V = NumberOfType1Error,
+ V2 = V2), list(mean = mean, sd = function(vec) round(sd(vec),
+ 1)))</pre>
```

> result\$statisticDF

```
funName sampleSize rho mu piO method alpha silent V.mean V2.mean V.sd V2.sd
1
    pVGen
                100 0 2 0.5
                                BH 0.5 TRUE
                                                15.1
                                                      245.3 4.4 131.7
2
    pVGen
               100
                    0 2 0.5
                                BH 0.25
                                         TRUE
                                                 5.9
                                                       40.9 2.6
3
               100 0.2 2 0.5
                                                      591.9 12.0 597.3
    pVGen
                                BH 0.5
                                         TRUE
                                                21.5
4
    pVGen
               100 0.2 2 0.5
                                BH 0.25 TRUE
                                                 9.2
                                                     163.2 9.3 329.0
5
    pVGen
               100 0.4 2 0.5
                                BH 0.5 TRUE
                                                14.6
                                                     401.6 14.5 684.8
               100 0.4 2 0.5
                                BH 0.25
                                                     126.6 9.9 318.5
6
    pVGen
                                          TRUE
                                                 6.2
                                                     860.7 20.1 981.2
    pVGen
               100 0.6 2 0.5
                                BH 0.5
                                          TRUE
                                                22.3
               100 0.6 2 0.5
8
    pVGen
                                BH 0.25
                                          TRUE
                                                 9.3
                                                      265.7 14.1 508.8
                                                      711.1 20.6 1051.5
9
    pVGen
               100 0.8 2 0.5
                                BH
                                   0.5
                                          TRUE
                                                18.1
               100 0.8 2 0.5
                                BH 0.25
                                         TRUE
                                                 8.0
                                                      328.6 17.1 801.7
10
    pVGen
    pVGen
               100 1 2 0.5
                                BH 0.5 TRUE
                                                20.0 1000.0 25.8 1291.0
12
    pVGen
               100 1 2 0.5
                                BH 0.25 TRUE
                                                15.0 750.0 24.2 1207.6
```

Another possibility is to call gatherStatistics without and "average function".

Then every result get his own row in \$statisticDF

```
> result <- gatherStatistics(sim, list(V = NumberOfType1Error,
+ V2 = V2))</pre>
```

> head(result\$statisticDF)

```
funName sampleSize rho mu pi0 method alpha silent V V2
   pVGen
               100 0 2 0.5
                                BH 0.5 TRUE 15 225
2
   pVGen
               100
                   0 2 0.5
                                BH
                                     0.5
                                          TRUE 10 100
3
   pVGen
               100
                   0 2 0.5
                                BH
                                     0.5
                                         TRUE 20 400
                    0 2 0.5
                                    0.5 TRUE 11 121
Δ
   pVGen
               100
                                RH
                    0 2 0.5
                                     0.5 TRUE 19 361
   pVGen
               100
                                BH
               100
                    0 2 0.5
                                BH
                                    0.5 TRUE 16 256
   pVGen
```

> tail(result\$statisticDF)

```
funName sampleSize rho mu pi0 method alpha silent V
                                          TRUE 0
115
    pVGen
               100 1 2 0.5
                                BH 0.25
                                                    ٥
116
    pVGen
                100 1 2 0.5
                                BH 0.25
                                          TRUE 50 2500
117
     pVGen
                100
                     1 2 0.5
                                BH 0.25
                                          TRUE 50 2500
                                BH 0.25
118
    pVGen
                100 1 2 0.5
                                          TRUE 50 2500
119
                100 1 2 0.5
                                BH 0.25
                                         TRUE 0
    pVGen
120
    pVGen
                100 1 2 0.5
                               BH 0.25
                                         TRUE 0
```

2.5 Plotting a bit

```
> if (!is.element("sim.plot", ls())) {
    isim.plot <- sim.plot (sim.plot cash) {
        im.plot <- sim.plot (sim.plot cash) {
            tun = pValFromEqui(orrData, sampleSize = 100, rho = seq(0, + 1, by = 0.2), mu = 0, pi0 = 1), list(list(fumName = "BH", + fun = function(pValues, alpha) BH(pValues, alpha, silent = TRUE), + alpha = c(0.5, 0.75))))
    }
} length(sim.plot$data)</pre>
```

[1] 6000

> length(sim.plot\$results)

100 0.2 0 1 100 0.4 0 1

[1] 12000

3

5

The are already different kind of R functions that take data.frames and generate histograms, boxplots and so on from them. For some plot example we will need the lattice package.

> require(lattice)

First we calculate V for every MutossSim object and then plot a histogram and a boxplot of V, see Figure 1 (p. 11), Figure 2 (p. 12).

```
> result.all <- gatherStatistics(sim.plot, list(V = NumberOfType1Error))
```

Also after the "average process" we again get an data.frame which can be used to generate plots.

BH 0.5 8.833764 11.544236 10.189 21.428169

BH 0.5 12.131991 15.860009 13.996 29.472574

> print(histogram(~V | rho, data = subset(result.all\$statisticDF,
+ alpha == "0.5")))

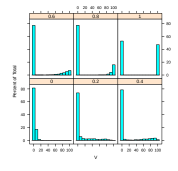


Figure 1: Histogram of the number of rejected true hypotheses for alpha equals $0.5\,$

> print(bwplot(V ~ rho | alpha, data = subset(result.all\$statisticDF)))

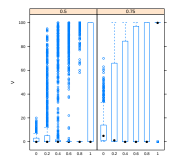


Figure 2: Boxplot of the number of rejected true hypotheses

```
> print(xyplot(V.M1 + V.M + V.Mu ~ rho | alpha, data = result$statisticDF,
+ type = "a", auto.key = list(space = "right", points = FALSE,
+ lines = TRUE())
```

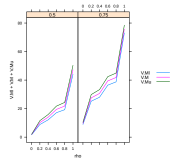


Figure 3: XYPlot of the mean number of rejected true hypotheses with asymptotic 95% confidence intervalls (pointwise)

2.6 Memory considerations

Up to now we have kept any information in memory. Somtimes, it is useful not to discard any generated Information. For example, if some objects in results appear odd, they can directly be reproduced and of course with all information available we can investigate anything like distribution or any statistic we did not consider before the simulation. The price for this liberty is in general an extensive memory usage which of course restrict the size of our simulation. On the other hand if we exactly know that we are only interested in EV_n the mean number of true rejected hypotheses, why should we keep the generated p-values or the index of the rejected p-values? Thus it would be enough that an object in \$results contains V_n the number of rejected true hypotheses. We will know show an example were only the information needed is kept in memory. The following is basically the 'simple example' from section 2.4 The main issue is to save memory thus we will calculate the V_n right after applying the procedure to the generated data set. If done so, we can also discard the generated data!

```
> V.BH <- function(pValues, groundTruth, alpha) {
   out <- BH(pValues, alpha, silent = TRUE)
   + V <- sum(outfrejected * groundTruth)
   + return(list(V = V))
   + }
```

As already mentioned anything in \$procInput will be used as an input for the procedures. Since our procedure now has a parameter groundTruth our data generated function has to be adepted. We just move groundTruth in \$procInput.

Obviously, it is unnecessary to store this generated data, because the number of Type I Errors we are interested in will be directly calculated. Setting discardProcInput = TRUE in simulation removes \$procInput from the object generated by the data generating function after \$procInput has been used by all specified procedures. In our case theses will be the 2 procedures V.BH with albha = 0.5 and albha = 0.25.

```
alpha = 0.5 and alpha = 0.25.
> set.seed(123)
> sim <- simulation(replications = 10, list(funName = "pVGen2",
+ fun = pValFromEquiforrData2, sampleSize = 100, rho = seq(0,
+ 1, by = 0.2), sun = 2, pio = 0.50, list(sit(funName = "V.BH",
+ fun = V.BH, alpha = c(0.5, 0.25))), discardProcInput = TRUE)
> result <- gatherStatistics(sim, list(V = function(data, results) results$V),
+ mean)</pre>
```

\$statisticDF

```
funName sampleSize rho mu piO method alpha V.mean
  pVGen2
              100 0 2 0.5 V.BH 0.5 15.1
               100 0 2 0.5 V.BH 0.25
   pVGen2
   pVGen2
               100 0.2 2 0.5
                             V.BH 0.5 21.5
4
  pVGen2
               100 0.2 2 0.5 V.BH 0.25
               100 0.4 2 0.5 V.BH 0.5 14.6
5
  pVGen2
  pVGen2
               100 0.4 2 0.5 V.BH 0.25 6.2
  pVGen2
              100 0.6 2 0.5 V.BH 0.5 22.3
  pVGen2
               100 0.6 2 0.5 V.BH 0.25
                                         9.3
               100 0.8 2 0.5 V.BH 0.5 18.1
100 0.8 2 0.5 V.BH 0.25 8.0
  pVGen2
10 pVGen2
11 pVGen2
               100 1 2 0.5 V.BH 0.5 20.0
              100 1 2 0.5 V.BH 0.25 15.0
12 pVGen2
```

\$name.parameters

[1] "funName" "sampleSize" "rho" "m11" "pi0" [6] "method" "alpha"

\$name.statistics [1] "V.mean"

\$name.avgFun

[1] "mean"

Here the corresponding table from section 2.4.

```
> set.seed(123)
```

```
> sim.simple <- simulation(replications = 10, list(funName = "pVGen",
     fun = pValFromEquiCorrData, sampleSize = 100, rho = seq(0,
          1, by = 0.2), mu = 2, pi0 = 0.5), list(list(funName = "BH",
```

fun = BH, alpha = c(0.5, 0.25), silent = TRUE)))

> result.simple <- gatherStatistics(sim.simple, list(V = NumberOfType1Error),

mean) > print(result.simple)

\$statisticDF

	funName	sampleSize	rho	mu	pi0	method	alpha	silent	V.mean
1	pVGen	100	0	2	0.5	BH	0.5	TRUE	15.1
2	pVGen	100	0	2	0.5	BH	0.25	TRUE	5.9
3	pVGen	100	0.2	2	0.5	BH	0.5	TRUE	21.5
4	pVGen	100	0.2	2	0.5	BH	0.25	TRUE	9.2
5	pVGen	100	0.4	2	0.5	BH	0.5	TRUE	14.6
6	pVGen	100	0.4	2	0.5	BH	0.25	TRUE	6.2
7	pVGen	100	0.6	2	0.5	BH	0.5	TRUE	22.3
8	pVGen	100	0.6	2	0.5	BH	0.25	TRUE	9.3
9	pVGen	100	0.8	2	0.5	BH	0.5	TRUE	18.1
10	pVGen	100	0.8	2	0.5	BH	0.25	TRUE	8.0
11	pVGen	100	1	2	0.5	BH	0.5	TRUE	20.0
12	pVGen	100	1	2	0.5	BH	0.25	TRUE	15.0

```
$name.parameters
                                             "mu"
[1] "funName"
                 "sampleSize" "rho"
                                                          "pi0"
[6] "method"
                 "alpha"
                              "silent"
$name.statistics
[1] "V.mean"
$name.avgFun
[1] "mean"
The same results but the memory usage differ much:
> print(s1 <- object.size(sim))
127192 bytes
> print(s2 <- object.size(sim.simple))
565912 bytes
> unclass(s2/s1)
[1] 4.449274
```

Reproducing some results from BKY (2006)

Now, we will reproduce figure 1 from the publication. For this we need to estimate the FDR. This will be done be calculating the FDP for every object in \$results and then calculating the empirical mean. In this simulation where we exactly know what we want and thus retain only the necessary information. To be able to calculate the FDP right after applying the procedure we need to know which pValue belongs to a true or false hypothesis. Thus, like in the foregoing section, \$groundTruth will be an element of \$procInput.

```
> pValFromEquiCorrData2 <- function(sampleSize, rho, mu, pi0) {
     nmbOfFalseHyp <- round(sampleSize * (1 - pi0))
     nmbOfTrueHyp <- sampleSize - nmbOfFalseHyp
     muVec <- c(rep(mu, nmbOfFalseHyp), rep(0, nmbOfTrueHyp))
     Y <- sqrt(rho) * rnorm(1) + sqrt(1 - rho) * rnorm(sampleSize) +
          muVec
     return(list(procInput = list(pValues = 1 - pnorm(Y), groundTruth = muVec ==
         0)))
+ }
```

We also need a function that calculates the FDP.

```
> FDP <- function(pValues, groundTruth, proc = c("BH", "M-S-HLF")) {
     if (proc == "BH")
         out <- BH(pValues, alpha = 0.05, silent = TRUE)
     else out <- adaptiveSTS(pValues, alpha = 0.05, silent = TRUE)
     R <- sum(out$rejected)
     if (R == 0)
```

```
return(list(FDP = 0))
     V <- sum(out$rejected * groundTruth)</pre>
     return(list(FDP = V/R))
+ }
Now, the (partial) reproduction of the results can start!
> set.seed(123)
> date()
[1] "Thu Dec 01 18:05:29 2011"
> sim <- simulation(replications = 10000, list(funName = "pVGen2",
     fun = pValFromEquiCorrData2, sampleSize = c(16, 32, 64, 128,
         256, 512), rho = c(0, 0.1, 0.5), mu = 5, pi0 = c(0.25,
         0.75)), list(list(funName = "FDP", fun = FDP, proc = c("BH",
      "M-S-HLF"))), discardProcInput = TRUE)
> date()
[1] "Thu Dec 01 18:39:30 2011"
> result <- gatherStatistics(sim, list(FDP = function(data, results) results$FDP),
     mean)> result
$statisticDF
  funName sampleSize rho mu pi0 method
                                        proc FDP.mean
                 16 0 5 0.25
                                 FDP
                                        BH 0.01243507
   pVGen2
                 16 0 5 0.25
2
   pVGen2
                                  FDP M-S-HLF 0.04680482
   nVGen2
                 32 0 5 0.25
                                 FDP
                                          BH 0.01254447
                 32 0 5 0.25
                                 FDP M-S-HLF 0.04933171
   pVGen2
   pVGen2
5
                64 0 5 0.25
                                 FDP
                                          BH 0.01245073
                64 0 5 0 25
                                 FDP M-S-HLF 0.05027779
   pVGen2
   pVGen2
               128 0 5 0.25
                                 FDP BH 0.01250933
   pVGen2
               128 0 5 0.25
                                 FDP M-S-HLF 0.05000333
   pVGen2
                256 0 5 0.25
                                  FDP BH 0.01259937
                256 0 5 0.25
10 pVGen2
                                  FDP M-S-HLF 0.05021692
11 pVGen2
                512 0 5 0.25
                                  FDP BH 0.01246457
12 pVGen2
                512 0 5 0.25
                                  FDP M-S-HLF 0.04993815
13 pVGen2
                16 0.1 5 0.25 FDP BH 0.01243852
                16 0.1 5 0.25
14 pVGen2
                                 FDP M-S-HLF 0.04929676
15 pVGen2
                32 0.1 5 0.25 FDP BH 0.01290907
16 pVGen2
                32 0.1 5 0.25
                                 FDP M-S-HLF 0.05355816
17 pVGen2
                64 0.1 5 0.25
                                FDP
                                        BH 0.01271804
                64 0.1 5 0.25
18 pVGen2
                                  FDP M-S-HLF 0.05626510
19 pVGen2
                128 0.1 5 0.25
                                  FDP
                                          BH 0.01233749
20 pVGen2
               128 0.1 5 0.25
                                  FDP M-S-HLF 0.05637963
21 pVGen2
                256 0.1 5 0.25
                                 FDP
                                          BH 0.01231818
22 pVGen2
                256 0.1 5 0.25
                                  FDP M-S-HLF 0.05619898
```

16 0.5 5 0.25 FDP M-S-HLF 0.06255181

FDP BH 0.01248149

FDP M-S-HLF 0.05719747

FDP BH 0.01269308

512 0.1 5 0.25

512 0.1 5 0.25

16 0.5 5 0.25

23 pVGen2

24 pVGen2

25 pVGen2

26 pVGen2

```
27 pVGen2
                 32 0.5 5 0.25
                                   FDP
                                           BH 0.01227550
28 pVGen2
                 32 0.5 5 0.25
                                  FDP M-S-HLF 0.06710637
29 pVGen2
                 64 0.5 5 0.25
                                   FDP
                                           BH 0.01183768
   pVGen2
                 64 0.5 5 0.25
                                   FDP M-S-HLF 0.07187446
31 pVGen2
                 128 0.5 5 0.25
                                   FDP
                                           BH 0.01200182
32 pVGen2
                 128 0.5 5 0.25
                                   FDP M-S-HLF 0.07523846
33
  pVGen2
                 256 0.5 5 0.25
                                  FDP
                                           BH 0.01245500
  pVGen2
                 256 0.5 5 0.25
                                   FDP M-S-HLF 0.07724869
34
35
   pVGen2
                 512 0.5 5 0.25
                                  FDP
                                         BH 0.01193485
36
   pVGen2
                 512 0.5 5 0.25
                                  FDP M-S-HLF 0.07817799
                 16 0 5 0.75
37
   pVGen2
                                   FDP
                                           BH 0.03687048
                 16 0 5 0.75
38
   pVGen2
                                  FDP M-S-HLF 0.05066989
39
  pVGen2
                 32 0 5 0.75
                                  FDP
                                           BH 0.03771246
                 32 0 5 0.75
                                  FDP M-S-HLF 0.04999444
40
  pVGen2
41
   pVGen2
                 64 0 5 0.75
                                  FDP
                                           BH 0.03736944
   pVGen2
                 64 0 5 0.75
                                  FDP M-S-HLF 0.04979089
  pVGen2
                 128 0 5 0.75
                                  FDP
                                           BH 0.03716345
44
   pVGen2
                 128 0 5 0.75
                                  FDP M-S-HLF 0.04963708
   pVGen2
                 256
                     0
                         5 0.75
                                   FDP
                                           BH 0.03812883
   pVGen2
                 256
                     0
                         5 0.75
                                   FDP M-S-HLF 0.05079616
   pVGen2
47
                 512 0 5 0.75
                                  FDP
                                           BH 0.03731167
48 pVGen2
                 512 0 5 0.75
                                  FDP M-S-HLF 0.05002008
  pVGen2
                 16 0.1 5 0.75
                                  FDP
                                           BH 0.03624063
50
   pVGen2
                 16 0.1 5 0.75
                                  FDP M-S-HLF 0.06107526
   pVGen2
                 32 0.1 5 0.75
                                  FDP
51
                                           BH 0.03702100
                 32 0.1 5 0.75
52
   pVGen2
                                  FDP M-S-HLF 0.06206310
   pVGen2
                 64 0.1 5 0.75
                                   FDP
                                           BH 0.03771594
53
  pVGen2
                 64 0.1 5 0.75
                                   FDP M-S-HLF 0.06404561
55 pVGen2
                 128 0.1 5 0.75
                                  FDP
                                           BH 0.03677325
                                  FDP M-S-HLF 0.06185353
   pVGen2
                 128 0.1 5 0.75
   pVGen2
                 256 0.1 5 0.75
                                  FDP
                                           BH 0.03694803
  pVGen2
                 256 0.1 5 0.75
                                  FDP M-S-HLF 0.06317781
                 512 0.1 5 0.75
                                   FDP
                                           BH 0.03694764
59
  pVGen2
   pVGen2
                 512 0.1 5 0.75
                                   FDP M-S-HLF 0.06267116
61
  pVGen2
                 16 0.5 5 0.75
                                   FDP
                                           BH 0.03533002
62 pVGen2
                 16 0.5 5 0.75
                                   FDP M-S-HLF 0.11075864
63 pVGen2
                 32 0.5 5 0.75
                                  FDP
                                           BH 0.03501351
64 pVGen2
                 32 0.5 5 0.75
                                   FDP M-S-HLF 0.11841870
65 pVGen2
                 64 0.5 5 0.75
                                  FDP
                                           BH 0.03331830
66
   pVGen2
                 64 0.5 5 0.75
                                  FDP M-S-HLF 0.12028660
67
   pVGen2
                 128 0.5 5 0.75
                                  FDP
                                           BH 0.03413304
   pVGen2
                 128 0.5 5 0.75
                                   FDP M-S-HLF 0.12464657
   pVGen2
                 256 0.5 5 0.75
                                  FDP
                                           BH 0.03561711
70
                 256 0.5 5 0.75
                                   FDP M-S-HLF 0.12532959
  pVGen2
71 pVGen2
                 512 0.5 5 0.75
                                  FDP
                                           BH 0.03463151
72 pVGen2
                 512 0.5 5 0.75
                                  FDP M-S-HLF 0.12510925
$name.parameters
                "sampleSize" "rho"
[1] "funName"
                                         "mu"
                                                     "pi0"
```

[6] "method"

"proc"

```
$name.statistics
[1] "FDP.mean"
```

\$name.avgFun

[1] "mean"

> print(xyplot(FDP.mean ~ sampleSize | pi0 * rho, data = result\$statisticDF,
+ group = proc, type = "a", auto.key = list(space = "right",

+ points = FALSE, lines = TRUE)))

