Sample Size and Power Analysis for Microarray Studies

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1 Introduction

This document shows the functionality of the R-package SSPA. The package performs power and sample size analysis using a method described by [1, 2]. Our implementation allows for fast and realistic estimates of power and sample size for microarray experiments, given pilot data. By means of two simple commands (pilotData(), sampleSize()), a researcher can read their data in and compute the desired estimates. Other functions are provided to facilitate interpretation of results.

Given a set of test statistics from the pilot data, the knowledge of their distribution under the null hypothesis and the sample size used to compute them, the method estimates the power for a given false discovery rate. The multiple

testing problem is controlled through the adaptive version of the Benjamini and Hochberg method [3]. For more details about the implementation and method we refer to [4, 1, 2]. In [5] we describe two biological case studies using the packageSSPA.

For comparison we implement the power and sample size estimation method proposed by Ruppert [6] which uses a different estimation approach. Two additional packages need to be installed for using this method namely quadprog and splines.

2 Basic Example

We demonstrate the functionality of this package by using the preprocessed gene expression data from the leukemia ALL/AML study of [7] from the multtest package. To load the leukemia dataset, use data(golub), and to view a description of the experiment and data, type ?golub. The number of samples per group are shown by table(golub.cl) where ALL is class 0 and AML class 1.

```
> library(multtest)
> data(golub)
> table(golub.cl)

golub.cl
0 1
27 11
```

The required input for the sample size and power analysis is a vector of test-statistics and the sample sizes used to compute them. The test-statistics are obtained by a differential gene expression analysis using one of the available packages like limma, maanova, multtest (it is also possible to import the vector of test-statistics in R if they are calculated using another software package). Here we will use the function mt.teststat from the multtest package to obtain a vector of test-statistics from the leukemia data.

```
> tst <- mt.teststat(golub, golub.cl)
```

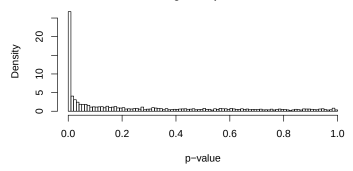
The first step in doing the sample size and power analysis is creating a object of class PilotData which will contain all the necessary information needed for the following power and sample size analysis; a vector of test-statistics and the sample sizes used to compute them. A user-friendly interface for creating an object of PilotData is available as pilotData().

```
> library(SSPA)
> pd <- pilotData(name="ALL/AML",testStatistics=tst,sampleSizeA=11,sampleSizeB=27)</pre>
```

Several ways of viewing the content of the PilotData object are possible either graphically or using a show-method by just typing the name of the created object of PilotData:

```
> par(mar=c(4.1, 4.1, 2.1, 2.1))
> layout(matrix(c(1,2),nrow=2))
> hist(pd, cex.main=1)
> plot(pd, cex.main=1)
```

Histogram of p-values



empirical cumulative distribution of p-values

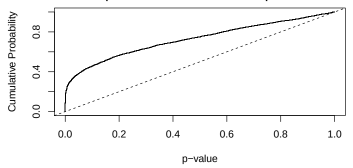


Figure 1: Histogram and ECDF plot of p-values.

> pd

An object of class "PilotData"

Experiment name: ALL/AML
Number of test-statistics: 3051
Effective sample size: 7.82
Null distribution: normal

A histogram of p-values is obtained by just calling hist(pd) and an empirical cumulative distribution of p-values by plot(pd). The accumulation of p-values near zero indicates that some genes are differentially expressed, as the deviation from the uniform distribution in the lower panel.

Now we can create an object of class SampleSize which will perform the estimation of the proportion of non-differentially expressed genes and the density of effect sizes. Several options are available see ?sampleSize. The default

method for estimation of the proportion of non-differentially expressed genes is the method proposed by [8] as implemented by convest() function from the package limma. Additionally the method by [9] and [2] are available, also a userdefined proportion is allowed. Again a generic show-method is implemented.

```
> ss <- sampleSize(pd)
> ss
```

An object of class "SampleSize"
Distribution of effect sizes is estimated from -6 to 6 using 1024 points.
Method for estimation of the proportion of non-differentially expressed: "Langaas Fraction of non-differentially expressed genes: 0.4702 (adjusted=0.4719).
Kernel used in the deconvolution is "fan" with bandwidth 0.353.

The density of effect size can be shown by a call to plotEffectSize(). When there are both up- and down-regulated genes a bimodal density is observed.

> plotEffectSize(ss, type='l')

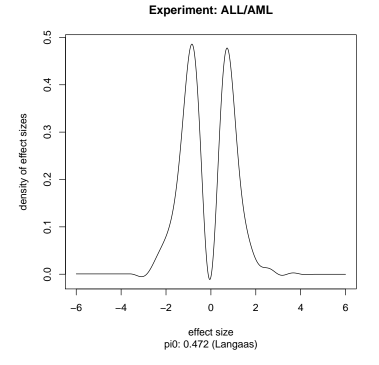


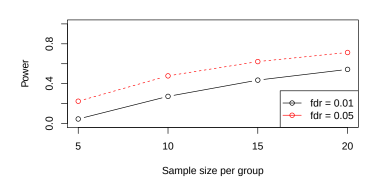
Figure 2: Density of effect-sizes.

Estimating the average power for other sample sizes then that of the pilotdata can be performed with the Power()-function. The user can also give the desired false discovery rate level or possible multiple false discovery rate levels as a vector.

```
> layout(matrix(c(1:2), nrow=2))
> par(mar=c(4.1, 4.1, 2.1, 2.1))
> pwr <- Power(ss, plot = FALSE, samplesizes = c(5, 10, 15, 20), fdr=0.01)
> plot(c(5, 10, 15, 20), pwr, ylim = c(0, 1), type = "b", ylab = "Power", xlab = "Sample siz"
> legend("bottomright", colnames(pwr), col=c(1:ncol(pwr)), pch=1, lty=1)
> pwr <- Power(ss, plot = FALSE, samplesizes = c(5, 10, 15, 20), fdr=c(0.01, 0.05))
> matplot(c(5, 10, 15, 20), pwr, ylim = c(0, 1), type = "b", pch=1, ylab = "Power", xlab = "Power
```

- fdr = 0.01

20



Sample size per group

10

Figure 3: Estimated average power for different sample sizes and FDR-levels.

15

3 SSPA adapted for the moderated t-statistics of limma

3.1 Approximated t-distribution

Sofar the estimations obtained are based on assumption that the test-statistics follow a normal distribution. Here we describe how we implemented an approximated t-distribution. Assume the t-statistics for each gene is calculated by:

$$T = \frac{\bar{X} - \bar{Y}}{\hat{\sigma}/\sqrt{N}} = \frac{\bar{X} - \bar{Y} - \Delta}{\hat{\sigma}/\sqrt{N}} + \frac{\Delta}{\hat{\sigma}/\sqrt{N}},\tag{1}$$

where $N = (1/n_X + 1/n_Y)^{-1}$, Δ is the difference between the means of the two groups and $\hat{\sigma}^2$ the pooled sample variance,

$$\hat{\sigma}^2 = \frac{n_X - 1}{n_X + n_Y - 2} S_X^2 + \frac{n_Y - 1}{n_X + n_Y - 2} S_Y^2. \tag{2}$$

If $\Delta=0$, then T has exactly a Student t-distribution with $\nu=n_X+n_Y-2$ degrees of freedom. If $\Delta\neq 0$, then T is the sum of Student random variable with n_X+n_Y-2 degrees of freedom plus

$$\frac{\Delta}{\hat{\sigma}/\sqrt{N}} = \frac{\Delta}{\sigma} \sqrt{N} \frac{\sigma}{\hat{\sigma}} \tag{3}$$

In other words, T is a Student random variable plus the square root of the effective sample size N times an effect size given by

$$\theta = \frac{\Delta}{\sigma} \frac{\sigma}{\hat{\sigma}} \tag{4}$$

This effect size is random, but since $\sigma/\hat{\sigma}$ converges a.s. to 1 it seems allright to view θ as a slightly perturbed effect size. The random variable T becomes $T + \sqrt{N}\theta$, when $\Delta \neq 0$. We can still use the model of Ferreira and Zwinderman (2006) but where the standard normal is replaced by the Student t-distribution.

3.2 moderated t-statistics

The moderated t-statistics calculated by limma follows a t-distribution with moderated degrees of freedom $\tilde{\nu}$. So we only have to plug-in the moderated degrees of freedom and we can use the moderated t-statistics in our power and sample size calculations. Because we use moderated degrees of freedom we also have a moderated sample size for each group.

The moderated degrees of freedom $\tilde{\nu}$ constists of two parts: ν_0 is the *prior* degrees of freedom estimated by limma using a empirical Bayesian approach and the residual degrees of freedom; for a two-group comparison this is given by $\nu = n_X + n_Y - 2$. Now we can define a moderated sample size for each group:

$$\tilde{\nu} = n_X + \nu_0/2 + n_Y + \nu_0/2 - 2 = \tilde{n_X} + \tilde{n_Y} - 2,\tag{5}$$

where the $\nu_0/2$ is added to the original sample sizes of each group.

Like before, first we created an object of class pilotData but now with test-Statistics the moderated t-statistics of the contrast of interest, sampleSizeA the moderated sample size of group A plus half the prior degrees of freedom (from a limma eBayes-fit object using fit\$df.prior). Internally the degrees of freedom are defined as fit\$df.prior + fit\$df.residual. By default the standard normal distribution is assumed so we need specifically the argument nullDist = "student".

3.3 Example using moderated t-statistics from limma

For illustration we will use the Swirl example from the *limma*-userguide but the Swirl data if obtained from the *marray*-package and using *convert* converted to a *limma* RGList. In the following bit of code the data is loaded, normalized and an empirical Bayesian linear model is fitted.

```
> library(marray)
> library(convert)
> library(limma)
> data(swirl)
> swirl <- as(swirl, "RGList")
> MA <- normalizeWithinArrays(swirl) #print tip loess
> design <- c(-1,1,-1,1)
> fit <- lmFit(MA, design)
> ordinary.t <- fit$coef/fit$stdev.unscaled/fit$sigma
> fitModerated <- eBayes(fit)</pre>
```

As described by the limma userguide; we can inspect the difference between moderated and original t test-statistics using qq-plots:

Now we load the SSPA-package and create two pilotData-objects, one for the ordinary t test-statistics and one for the moderated t test-statistics with the appropriate degrees of freedom and sample sizes.

```
> par(mfcol=c(1,2))
> qqt(fitModerated$t, df=fitModerated$df.prior + fitModerated$df.residual, pch=16, cex=0.5,
> abline(0,1)
> qqt(ordinary.t, df=fit$df.residual, pch=16, cex=0.5, main="Ordinary t")
> abline(0,1)
```

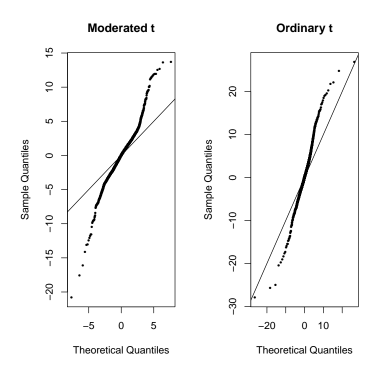


Figure 4: qq-plots: showing the difference between moderated and orignal t test-statistics.

An object of class "PilotData"

Experiment name: Swirl

Number of test-statistics: 8448

Effective sample size: 2.01

Null distribution: student

Degree of Freedom: 7.024394

The data comes from a dye-swap experiment with four microarrays, thus for each RNA source we have two biological and two technical replicates. Because this is an dye-swap experiment we can estimate the difference between the biological sources, leaving three degrees of freedom for estimation of the error.

The pdMod-object has a moderated sample sizes and moderated degrees of freedom, the moderated Student t-distribution will be used for the analysis. The effective sample size was defined by: $N = (1/n_X + 1/n_Y)^{-1}$, thus with both four samples gives two. The moderated sample size is 2.01.

The next figure show the difference between the ordinary t test-statistics and moderated t test-statistics (left-panel). The right-panel show the difference in p-values where one is calculated using the ordinary degrees of freedom and the other using the moderated degrees of freedom.

```
> ss <- sampleSize(pd)
> ss

An object of class "SampleSize"
Distribution of effect sizes is estimated from -6 to 6 using 1024 points.
Method for estimation of the proportion of non-differentially expressed: "Langaas Fraction of non-differentially expressed genes: 0.5957 (adjusted=0.6151).
Kernel used in the deconvolution is "fan" with bandwidth 0.3326.

> ssMod <- sampleSize(pdMod)
> ssMod

An object of class "SampleSize"
Distribution of effect sizes is estimated from -6 to 6 using 1024 points.
Method for estimation of the proportion of non-differentially expressed: "Langaas Fraction of non-differentially expressed genes: 0.6117 (adjusted=0.6256).
Kernel used in the deconvolution is "fan" with bandwidth 0.3326.
```

Once the proportion of non-differentially expressed genes is estimated the density of effect sizes can be estimated.

The dip below zero for the density of effect sizes based on the moderated test-statistics is unwanted. We implemented the suggestion by Efron *et al.* [10] for better upper bound estimates of π_0 . The default value 0.5 this value could be tuned to get a valid density.

Again estimation of the average power for other sample sizes then that of the pilot-data can be performed:

```
> par(mfcol=c(1,2))
> plot(abs(pd@testStatistics), abs(pdMod@testStatistics),
    xlab="|ordinary t test statistics|",
    ylab="|moderated t test statistics|", pch=16, cex=0.5)
> abline(a=0, b=1, col="red", lwd=2, lty=2)
> plot(-log10(pd@pValues), -log10(pdMod@pValues), xaxt="n", yaxt='n', xlab=expression(paste(ylab=expression(paste(-log[10],"(moderated p-values)")), pch=16, cex=0.5)
> at <- axTicks(2)
> axis(2, at=at, tcl=-1, labels=parse(text=c("10^0", paste("10^-", at[-1], sep="")));
> at <- axTicks(1)
> axis(1, at=at, tcl=-1, labels=parse(text=c("10^0", paste("10^-", at[-1], sep="")));
> abline(a=0, b=1, col="red", lwd=2, lty=2)
```

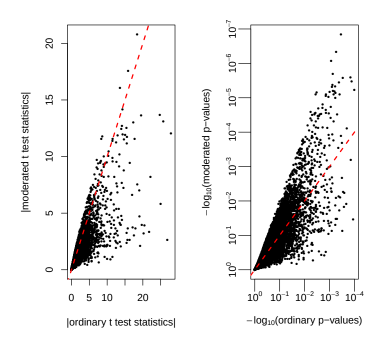


Figure 5: scatter plots: showing the difference between moderated and orignal t test-statistics.

```
> plotEffectSize(ssMod, type='1' , 1wd=2, sub=NULL)
> lines(ss@theta, ss@lambda, col="red", 1wd=2)
> piOMod <- paste("piO: ", signif(ssMod@piO[[2]], 3))
> piO <- paste("piO: ", signif(ss@piO[[2]], 3))
> labels <- c(paste("moderated ", piOMod), paste("ordinary ", piO))
> legend("topleft", labels, col=c("black", "red"), pch=1)
```

Experiment: Swirl

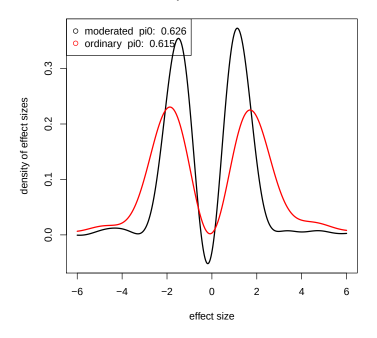


Figure 6: Density of effect size for moderated and orignal t test-statistics.

```
> samplesizes <- c(2, 4, 6, 8, 10)
> pwr <- Power(ss, plot = FALSE, samplesizes = samplesizes, fdr=c(0.05, 0.1))
> pwrMod <- Power(ssMod, plot = FALSE, samplesizes = samplesizes + nu0/2, fdr=c(0.05, 0.1))
> matplot(samplesizes, pwr, ylim = c(0, 1), type = "b", col=1, pch=1, ylab = "Power", xlab = "matlines(samplesizes, pwrMod, col=2, type="b", pch=1)
> legend("topleft", colnames(pwr), col=1, lty=c(1, 2))
> legend("bottomright", c("ordinary", "moderated"), col=1:2, lty=1)
```

Power Curves

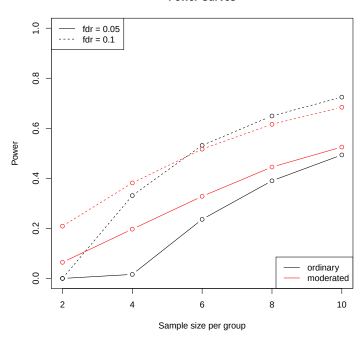


Figure 7: Estimated average power for other sample sizes then that of the pilot-data for both moderated and original t test-statistics and for different false discovery rate thresholds.

For small sample sizes the moderated test-statistics has higher power for larger sample sizes both statistics converge to eachother, as expected.

4 Additional functionality

4.1 Ferreira's π_0 estimate

Ferreira *et al.* propose a semi-parametric method to estimate the proportion of non-differentially expressed genes[1, 2]. Use sampleSize(pd, method="Ferreira",

pi0=seq(0.05, 0.5, 0.05), doplot=TRUE).

4.2 Rupperts estimation method

For using the method proposed by Ruppert [6] two additional packages need to be installed for using this method namely quadprog and splines. Using sampleSize(pd, method="Ruppert", nKnots = 11, bDegree = 3) both π_0 and the density of effect sizes is estimated by Ruppert's method.

5 Details

This document was written using:

- R version 2.15.0 (2012-03-30), x86_64-unknown-linux-gnu
- Locale: LC_CTYPE=en_US.UTF-8, LC_NUMERIC=C, LC_TIME=en_US.UTF-8, LC_COLLATE=C, LC_MONETARY=en_US.UTF-8, LC_MESSAGES=en_US.UTF-8, LC_PAPER=C, LC_NAME=C, LC_ADDRESS=C, LC_TELEPHONE=C, LC_MEASUREMENT=en_US.UTF-8, LC_IDENTIFICATION=C
- Base packages: base, datasets, grDevices, graphics, methods, stats, utils
- Other packages: Biobase 2.16.0, BiocGenerics 0.2.0, SSPA 1.12.0, convert 1.32.0, limma 3.12.0, marray 1.34.0, multtest 2.12.0, qvalue 1.30.0
- Loaded via a namespace (and not attached): MASS 7.3-17, splines 2.15.0, stats4 2.15.0, survival 2.36-12, tcltk 2.15.0, tools 2.15.0

References

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