# MAMA: an R package for Meta-Analysis of MicroArray

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April 29, 2013

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# Introduction

This paper provides a user guide to R-package MAMA. The package implements eight different methods that have been proposed in meta-analysis of microarray and are designed to identify differentially expressed genes.

In here, we will demonstrate the features of the package with an example of meta-analysis in cancer microarray data, the comparison of expression profiles in MSI (microsatelite instable) and MSS (microsatelite stable) colon cancer. We gathered three microarray data from public databases. The data are stored in ColonData. It is an object of new S4 class - MetaArray, in which separate slots for gene expression data matrices (expression profiles), clinical sample characteristics and datanames were defined.

We start with package and sample data loading.

```
> rm(list=ls(all=TRUE))
> options(width=60)
> library(MAMA)
> data(ColonData)
> ColonData
Dataset Denmark containing 500 probes and 77 samples.
Sumarization of samples:
MSIstatus
MSI MSS
 39
    38
age
   Min. 1st Qu. Median
                           Mean 3rd Qu.
                                           Max.
        45.00
                54.00
                          54.92
                                  62.00
                                          80.00
Dataset Australia containing 500 probes and 36 samples.
Sumarization of samples:
MSIstatus
MSI MSS
  5 31
age
```

```
Min. 1st Qu.
                 Median
                            Mean 3rd Qu.
                                            Max.
  50.00
          57.75
                  66.00
                           66.36
                                   73.75
                                           85.00
Dataset Japan containing 500 probes and 41 samples.
Sumarization of samples:
location
  distal proximal
                   unknown
      28
               11
MSIstatus
MSI MSS
 16
    25
age
                            Mean 3rd Qu.
  Min. 1st Qu.
                 Median
                                            Max.
  40.00
          47.00
                  53.00
                           53.61
                                   60.00
                                           65.00
```

The original data sets have been preprocessed and subsampled in order to reduce the computational complexity of the meta-analysis. All data sets have been normalized and are in  $log_2$ -scale. The corresponding sample sizes for the three datasets can be seen in outprint above. The same set of 500 gene has been selected in each dataset.

A different method is described in each of the parts below and the descriptions are independed from each other, so you can directly move to the method of your interest. Meta-analysis usually consist of three steps: Data preparation, Detection of differentially expressed genes and Extraction and visualization of results. There is also a wrapper function for each method that outperforms all necessary step as one-line command.

We select the significant genes at significance level of 0.05 in all methods.

# Methods that combine p-values

#### Introduction

In this part we will focus on methods that combine p-values [4], [5]. These methods are inspired by Fisher's S-statistic published in 1925 [6].

Two measurements of significance of change in gene expression are the most common. These are: value of test-statistic and p-value. In here the p-values from study-specific analysis are combined into one p-value in sense of sum of logs. Methods differ in test statistic used to calculate the study-specific p-values.

## Usage

#### Data preparation

One needs a list of gene expression data matrices esets and a list of vectors with class labels classes. The former is GEDM slot of MetaArray object, the latter can be obtained by function selectClass.

```
> esets <- GEDM(ColonData)
> classes <- selectClass(ColonData, "MSIstatus", "factor")</pre>
```

#### Detecting differentially expressed genes

Functions pvalcombination and pvalcombination.paired provide meta-analysis based on combination of p-values. The first one is designed for unpaired data and the second for paired design of microarray experiments. Because, our data sets are unpaired, we will use pvalcombination. The function requires: a list of gene expression data matrices (esets), a list of vectors of class labels (classes), type of test statistics (moderated) and threshold for significance (BHth). It returns list of indices of selected genes. Three possible values for argument moderated are available: "t" for common t-test, "limma" for moderated t-test used in limma package [7] and "SMVar" for moderated t-test defined in SMVar package [8].

> pvalt<-pvalcombination(esets, classes, moderated = "t", BHth = 0.05)

```
DE IDD Loss IDR IRR
215.00 43.00 20.00 20.00 10.42
```

Several characteristics, which have been defined in meta-analysis of microarray (especially for methods which combine p-values or effect sizes), are outprinted by the function. DE denotes number of significant genes in meta-analysis. IDD represents Integration Driven Discoveries, it means genes which are significant in meta-analysis but not in any of the individual studies. Other way round, if a gene is significant only in individual data sets but not in meta-analysis, it is called Integration Driven Revision and Loss is a number of such genes. IDR and IRR are percentages of Integration Driven Discoveries and Integration Driven Revisions in identified differentially expressed genes (DE).

#### Results

> summary(pvalt)

	Length	Class	Mode
study1	162	-none-	numeric
study2	14	-none-	numeric
study3	99	-none-	numeric
${\tt AllIndStudies}$	192	-none-	numeric
Meta	215	-none-	numeric
${\tt TestStatistic}$	500	-none-	numeric

This object is a list with six slots. Study1 to Study3 are numeric vectors with indices of differentially expressed genes in data sets 1 to 3. AllIndStudies is a vector of indices of differentially expressed genes in at least one data set. Differentially expressed genes found by meta-analysis have their indices stored in Meta. And finally, a slot called TestStatistic is a vector with test statistics in meta-analysis.

#### Wrapper function

Function metaMA is a wrapper function for this method. It does not require the data preparation step and returns the same output as pvalcombination. By

```
> pval<- metaMA(ColonData, "MSIstatus", which ="pval")
```

```
DE IDD Loss IDR IRR
212.00 43.00 22.00 20.28 11.52
```

we combine study specific p-values(pval) in ColonData dataset using column "MSIstatus" from clinical data as class labels. The only difference between pval and pvalt is the *gene.names* slot in the wrapper function.

# Methods that combine effect sizes

#### Introduction

Methods that combine effect size use hierarchical model:

$$y_i = \theta_i + \epsilon_i, \epsilon_i \sim N(0, \sigma_i^2)$$

$$\theta_i = \mu + \delta_i, \delta_i \sim N(0, \tau_i^2),$$

where  $\mu$  is true difference in mean expression between two classes,  $y_i$  denotes the measure effect for study i, with  $i=1,..,k,\,\tau^2$  represents the between study variability,  $\sigma_i^2$  denotes the within study variability. The analysis is different depending on whether a fixed-effect model (FEM) or a random-effect model (REM) is deemed appropriate. Under a FEM,  $\tau=0$  is assumed, otherwise a REM need to be fit. The estimates of the overall effect  $\mu$  are different depending on which model is used.

Two papers dealing with effect size combination as method for meta analysis of microarray have been published [4] and [9]. They differ in effect size definition and implementation.

Method presented in [4] offers three variants of effect sizes (classical and moderated T-test) and uses explicitly random-effect model. It is implemented as two functions EScombination for unpaired data and EScombination.paired for paired data.

On the other hand, in [9] the effect size is defined as Hedge's and Olkin's g and both random-effect and fixed-effect are available. Package GeneMeta [10] implements this method.

## Algorithm

- 1. Data recoding.
- 2. Effect size calculation in each data set.
- 3. Decision between random-effect model (REM) and fixed-effect model (FEM).

- 4. Model application.
- 5. Identification of differentialy expressed genes.

### Usage

Because there are two different ways of implementation of this method, we will discuss them separately.

#### Implementation from metaMA package

**Data preparation** This method requires two lists, one containing the data matrices and the other one the corresponding vectors of group labels.

```
> esets <- GEDM(ColonData)
> classes <- selectClass(ColonData, "MSIstatus", "factor")</pre>
```

Detecting differentially expressed genes As we have unpaired data, we are going to use function EScombination. This function has four arguments: a list of gene expression data matrices (esets), a list of class labels vectors (classes), effect size definition (moderated) and a threshold for false discovery rate (FDR) (BHth). Three possible values for moderated are available: "t" for common t-test, "limma" for moderated t-test used in limma package [7] and "SMVar" for moderated t-test defined in SMVar package [8].

```
> ESt<-EScombination(esets, classes, moderated = "t" , BHth = 0.01)
```

```
DE IDD Loss IDR IRR
109.00 28.00 51.00 25.69 38.64
```

Function EScombination prints several measures defined in meta-analysis of microarray. DE denotes number of significant genes in meta-analysis. IDD represents Integration Driven Discoveries, it means genes which are significant in meta-analysis but not in any of the individual studies. Other way round, if a gene is significant only in individual data sets but not in meta-analysis, it is called Integration Driven Revision and Loss is a number of such genes. IDR and IRR are percentages of Integration Driven Discoveries and Integration Driven Revisions in identified differentially expressed genes (DE).

#### > summary(ESt)

	Length	Class	Mode
study1	113	-none-	${\tt numeric}$
study2	8	-none-	${\tt numeric}$
study3	59	-none-	${\tt numeric}$
${\tt AllIndStudies}$	132	-none-	${\tt numeric}$
Meta	109	-none-	${\tt numeric}$
${\tt TestStatistic}$	500	-none-	numeric

This object is a list with six slots. Study1 to Study3 are indices of differentially expressed genes in data sets 1 to 3. AllIndStudies is a vector of indices of

differentially expressed genes in at least one data set. Differentially expressed genes found by meta-analysis have their indices stored in *Meta*. And finally, a slot called *TestStatistic* is a vector with test statistics ("combined effect size") in meta-analysis.

Wrapper function The same results can be obtained via

```
> es <- metaMA(ColonData, "MSIstatus", which = "ES")

DE IDD Loss IDR IRR

166.00 27.00 52.00 16.27 27.23
```

#### Implementation from GeneMeta package

**Data preparation** Before calculating effect sizes we have to create numeric vectors with class labels in form of 1's and 0's. 1 is supposed to be for diseased samples and 0 for normal samples. In data sets used as example in this document 1 refers to MSI samples and 0 to MSS.

```
> ph1<-clinical(ColonData)[[1]][,1]</pre>
> levels(ph1) <- c(1, 0)
> clinical(ColonData)[[1]][,1]
[71] MSS MSS MSS MSS MSS MSS MSS
Levels: MSI MSS
> ph1
Levels: 1 0
> ph2<-clinical(ColonData)[[2]][,1]</pre>
> levels(ph2) <- c(1, 0)
> ph3<-clinical(ColonData)[[3]][,2]</pre>
levels(ph3) \leftarrow c(1, 0)
```

ph1, ph2 and ph3 are numeric vectors containing class labels for data sets denmark, australia and japan. These vectors are needed as arguments for functions which provide effect size and its variability estimates.

Detecting differentially expressed genes Functions getdF, dstar and sigmad estimate effect size and its variability for a individual data set, therefore we have to use them three-times. For denmark data set

```
> d.den <- getdF(GEDM(ColonData)[[1]], ph1)</pre>
> d.adj.den <- dstar(d.den, length(ph1))</pre>
> var.d.adj.den <- sigmad(d.adj.den, sum(ph1 == 0), sum(ph1 == 1))
> head(d.adj.den)
[1] 0.4835090 -0.1882238 -0.2740332 -0.4466879 -0.9850491
[6] -1.1694108
> head(var.d.adj.den)
[1] 0.05347487 0.05218687 0.05244444 0.05325247 0.05825761
[6] 0.06083683
and for other two data sets
> d.aus <- getdF(GEDM(ColonData)[[2]], ph2)</pre>
> d.adj.aus <- dstar(d.aus, length(ph2))</pre>
> var.d.adj.aus <- sigmad(d.adj.aus, sum(ph2 == 0), sum(ph2 == 1))</pre>
> d.jap <- getdF(GEDM(ColonData)[[3]], ph3)</pre>
> d.adj.jap <- dstar(d.jap, length(ph3))</pre>
> var.d.adj.jap <- sigmad(d.adj.jap, sum(ph3 == 0), sum(ph3 == 1))</pre>
```

Function getdF has two arguments: the data set (a ExpressionSet object or a matrix) and class labels (a factor or numeric vector with 1 and 0) and computes estimates of standardized mean difference, found in Hedge and Olkin's [11]. Function dstar corrects the estimates for sample size bias, therefore its second argument is sample size of the data set. Function sigmad calculates the estimate of variance of unbiased effect size. For calculation, the user has to provide effect size estimates and sample size of each class.

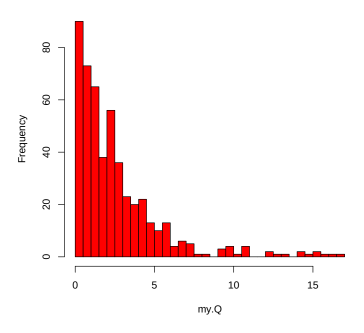
Now, we are going to use Chochran's Q statistic [12] to test between-study variability, so we can decide whether we should be considering random-effect (REM) or fixed-effect model(FEM) for the data.

Function f. Q provides a straightforward calculation of Cochran's Q statistic. If the null hypothesis that the between-study variance is equal to zero (data are well modeled by a fixed effects design) then the estimated Q values will have approximately a chi-squared distribution with degrees of freedom equal to the number of studies minus one. We are going to look at mean and histogram of Q statistics. Later we will compare quantiles of Q to quantiles of chi-square distribution.

```
> mymns <- cbind(d.adj.den, d.adj.aus,d.adj.jap)
> myvars <- cbind(var.d.adj.den, var.d.adj.aus,var.d.adj.jap)
> my.Q <- f.Q(mymns, myvars)
> mean(my.Q)

[1] 2.576469
> hist(my.Q, breaks = 50, col = "red")
```

## Histogram of my.Q



> num.studies <- 3
> plotQvsChi(my.Q, num.studies)

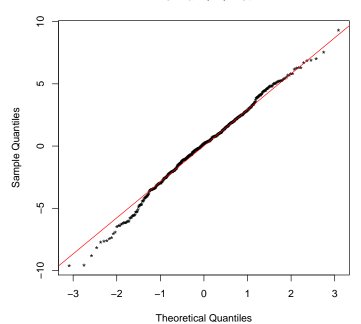
# 

According to Q-Q plot the hypothesis seems to be valid and fixed-effect model (FEM) should be used. However, we are going to use random-effect model (REM) too, so we can see if there is any difference in estimates of combined effect size.

The computation is simpler for FEM than for REM. Functions mu.tau2 and var.tau2 estimate combined effect size (mu.tau2) and variance (var.tau2). Each effect size is a weighted average of the effects for the individual data sets divided by its standard error. The weights are the reciprocal of the estimated variances.

```
> #FEM
> muFEM = mu.tau2(mymns, myvars)
> sdFEM = var.tau2(myvars)
> ZFEM = muFEM/sqrt(sdFEM)
> qqnorm(ZFEM, pch = "*")
> qqline(ZFEM, col = "red")
```

#### Normal Q-Q Plot



Plotting the quantiles of the effects we can see that the presumption of approximate Normality seems to be appropriate.

In REM we have to account between-study variability  $(\tau^2)$ . Function tau2.DL provides DerSimonian's and Laird's [13] estimates of  $\tau^2$  from Cochran's Q. It has two addional arguments: number of studies (num.studies) and weights (my.weights=1/myvars). We add between-study variability to estimated variance (myvars) and calculate the combined effect size like in FEM.

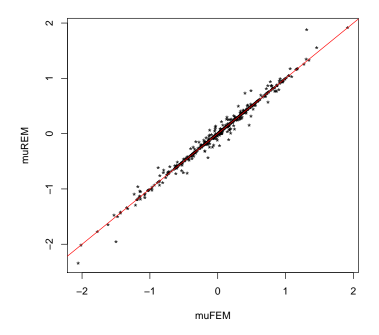
```
> #REM
> num.studies <- 3</pre>
```

```
> my.tau2.DL <- tau2.DL(my.Q, num.studies, my.weights=1/myvars)
> myvarsDL <- myvars + my.tau2.DL
> muREM <- mu.tau2(mymns, myvarsDL)
> varREM <- var.tau2(myvarsDL)
> ZREM <- muREM/sqrt(varREM)</pre>
```

muFEM or muREM are numeric vectors with estimated combined (overall) effect size for a gene in FEM or REM. The estimated standard error of overall effect size for each gene is stored in numeric vectors: varFEM or varREM. We will test significance of overall effect size by Z-score (ZFEM or ZREM) defined as mean divided by standard error.

We can easily compare FEM estimates and REM estimates

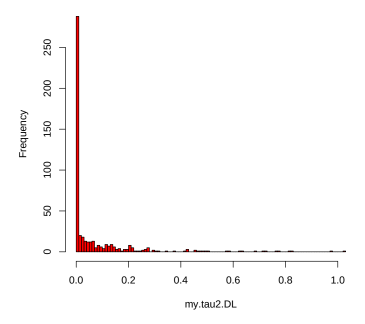
```
> plot(muFEM, muREM, pch = "*")
> abline(0, 1, col = "red")
```



We do not see much difference here. Actually, for most of the genes the  $\tau^2$  is estimated as zero.

```
> hist.tau<-hist(my.tau2.DL, col = "red", breaks = 100, main = "Histogram of tau")
```

#### Histogram of tau



**Results** The procedure described in details above is also implemented in function zScores. The arguments of this function are a list of expression sets (esets) and a list of classes (classes). Argument useREM chooses between REM and FEM.

```
> esets <- GEDM(ColonData)</pre>
> classes <- selectClass(ColonData, "MSIstatus", "binary")</pre>
> theScores <- zScores(esets, classes, useREM = FALSE)
 round(theScores[1:2, ],3)
            zSco_Ex_1 zSco_Ex_2 zSco_Ex_3
                                              zSco MUvals
217562_at
                                      0.881
                                             1.865
                 2.091
                          -0.542
                                                     0.326
203766_s_at
               -0.824
                          -0.085
                                      0.855 -0.196 -0.034
            MUsds Qvals df Qpvalues Chisq Effect_Ex_1
            0.175 1.964
                          2
                                0.375 0.062
                                                   0.484
217562_at
203766_s_at 0.174 1.379
                          2
                                0.502 0.845
                                                  -0.188
            Effect_Ex_2 Effect_Ex_3 EffectVar_Ex_1
217562_at
                  -0.262
                                0.283
                                                0.053
                                                0.052
203766_s_at
                  -0.041
                                0.275
            EffectVar_Ex_2 EffectVar_Ex_3
217562_at
                      0.233
                                      0.103
                                      0.103
203766_s_at
                      0.232
```

We get a matrix (theScores) with the following columns:

•  $Effect\_Ex\_$  are the unbiased estimates of the effect (d.adj.)

- EffectVar\_Ex\_ are the estimated variances of the unbiased effects (var.d.adj.
- zSco\_Ex\_ are the unbiased estimates of the effects divided by their standard deviation
- Qvals are the Q statistics (my.Q) and df is the number of combined experiments minus one
- MUvals and MUsds are equal to muFEM and sdFEM (the overall mean effect size and its standard deviation)
- zSco are the z scores (ZFEM)
- *Qpvalues* is for each gene the probability that a chi-square distribution with *df* degree of freedom has a higher value than its *Q* statistic
- Chisq is the probability that a chi-square distribution with 1 degree of freedom has a higher value than zSco2

Function zScoresFDR implements SAM [?] type analysis to estimate the false discovery rate (FDR).

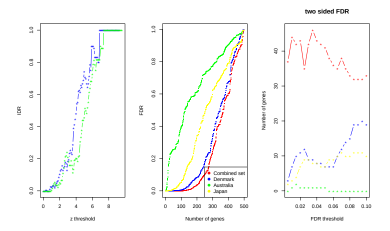
> round(ScoresFDR\$pos[1:2, ],3)

```
zSco_Ex_1 FDR_Ex_1 zSco_Ex_2 FDR_Ex_2 zSco_Ex_3
217562_at
                2.091
                         0.083
                                   -0.542
                                             1.073
                                                        0.881
                                   -0.085
203766 s at
               -0.824
                         1.204
                                             1.006
                                                       0.855
            FDR_Ex_3
                       zSco
                              FDR MUvals MUsds Qvals df
217562_at
               0.585 1.865 0.115 0.326 0.175 1.964
203766_s_at
               0.585 -0.196 1.044 -0.034 0.174 1.379
            Qpvalues Chisq
217562_at
               0.375 0.062
               0.502 0.845
203766_s_at
```

Function plotES provides several visualizations of the results. Specifying which=1 will plot so called *IDRplot*. This plot shows the fraction of the genes that have a higher effect size than the threshold for the combined Z-score, but not for any of the data set specific Z-scores. Genes with combined Z-score > 0 and < 0 are plotted separately. Selection which=2 will plot the number of genes and the corresponding FDR for the two sided situation. If the user is more interested in the number of genes that are below a given threshold for the FDR, one decides for which=3. It shows for each study (indicated by different colors) and various thresholds for the FDR (x axis) the number of genes that are below this threshold in the given study but above in all other studies are shown (y axis). If numeric vector is used that all figure specified in the vectors are plot.

Argument legend.names is a character vector with names of the date set used in legends and colors is a vector of colors to be used for plotting.

- > plotES(theScores, ScoresFDR, num.studies=3, legend.names=c("Combined set",
- + "Denmark", "Australia", "Japan"), colors=c("red", "blue",
- + "green", "yellow"), which=1:3)



Wrapper function There is also another function (ES.GeneMeta) that wrapes data preparation and functions zScore, ScoresFDR.

> es2<-ES.GeneMeta(ColonData, "MSIstatus", nperm = 100)

 ${\tt nperm} = 100$  is used only for computational complexity. A thousand of permutation is more appropriate.

# Similarity of Ordered Gene Lists (SOGL)

#### Introduction

Similarity of Ordered Gene Lists is another method for meta-analysis of microarray. It is call as "comparison of comparisons" by its authors [?].

Briefly, it assigns a similarity score to a comparison of ranked (ordered) gene lists. The score is based on the number of overlapping genes in the top ranks. It computes the size of overlap for each rank. The final score is a weighted sum of these values, with more weight put on the top ranks.

## Algorithm

- 1. Required data sets data sets with same set of genes are required.
- 2. Ranking of genes The genes are then ranked based on gene-wise test on difference of class mean. There is only one assumption about test result: a large positive test score corresponds to up-regulation and a large negative value to down-regulation.
- 3. Computing the overlap for each rank (from 1 to number of genes) we count the number of genes that appear in both ordered lists up to that position. It is denoted as  $O_n(G_A, G_B)$ , where  $G_A$  and  $G_B$  refer to ordered gene lists.
- 4. Similarity score First we compute a total overlap  $A_n$  at position n given as  $O_n(G_A, G_B) + O_n(f(G_A), f(G_B))$ , where f() means flipped list (down-regulated genes on top). Later we add weights to it and we sum it up to preliminary score. Weights  $w_{\alpha} = e^{-\alpha \cdot n}$  are used as default and tunning of parameter  $\alpha$  is needed. The weights are used to put more importance on top genes.

The algorithm above is valid for meta-analysis in which expression data are not available. However, in this situation we can not use same approach for tunning  $\alpha$ .

## Usage

#### Data preparation

As long as we have data stored in MetaArray object, there is no need for data transformation of any kind. We can move directly to order the genes as the first step in detecting differentially expressed genes.

### Detecting differentially expressed genes

Detection of differentially expressed gene can be performed by a sequence of function or by one wrapper function. We will discuss both approaches below.

We start with ordering the genes from the most up-regulated to the most down-regulated ones. Function computeOrdering provides such functionality. It has three arguments: the dataset, name of clinical parameter with class labels and the test statistic (fold-change or t-test with unequal variance).

```
> ordering<- computeOrdering(ColonData, "MSIstatus", "T")</pre>
```

Now we have to tune the  $\alpha$  parameter so we could obtain the similarity score.

Function computeAlpha returns vector of possible values for  $\alpha$ . These depend on the number of genes in analysis (argument ngenes) or minimal weight (argument min.weight) to be considered in score calculation. One can also select a particular number of top genes to be analyzed (argument n).

```
    A<-computeAlpha(ngenes=500)</li>
    A
    [1] 0.11512925 0.07675284 0.05756463 0.03837642 0.02878231
    [6] 0.02302585
    When n is used only one value of α is returned.
```

```
> a<-computeAlpha(n=25, ngenes=500)</pre>
```

> a

[1] 0.460517

Random permutation of class labels and subsampling are necessary to gather the optimal value for  $\alpha$ . This is the most computationally complex step of the analysis.

```
> random.score<-RandomScore(data = ColonData, varname = "MSIstatus",
+ B = 10, alpha = A, test = "T", two.sided = TRUE)</pre>
```

Function RandomScore performs random permutation of class labels and subsampling and returns a list with three slots. A matrix of smiliarity scores in which rows refer to  $\alpha$ 's and column to permutations is stored in random slot. Similar results from subsampling are stored in subsample slot. If "empirical" is part of which argument then also 95% empirical confidence intervals and medians of expected number of overlapping genes for each position are provided in empiricial.ci. The argument two.sided is TRUE if both sides of ordered gene list are to be compared and FALSE if only top genes have to be considered.

Now, optimal value for  $\alpha$  can be selected according to pAUC as a measure of the separability of two distributions: scores after random permutation of class labels and scores after subsampling. The  $\alpha$  with the highest pAUC is selected.

```
> A.opt <- selectAlpha(A, random.score$subsample, random.score$random)
```

#### Results

The final similarity score score, its significance sig and vector of genes which contribute to the overall similarity score genes can be obtained via:

```
> score<-prelimScore(ordering, A.opt$alpha)
> sig<-sigScore(ordering, A.opt$alpha, 100)
> genes<-selectGenes(ordering, A.opt$alpha, 0.95)</pre>
```

#### Wrapper function

Function performSOGL is a wrapper function for Similarity of Ordered Gene Lists.

```
> SOGL.res <- performSOGL(ColonData, varname = "MSIstatus", test = "FCH", + B = 100, which=c("score", "empirical"), min.weight = 1e-05, two.sided = TRUE )
```

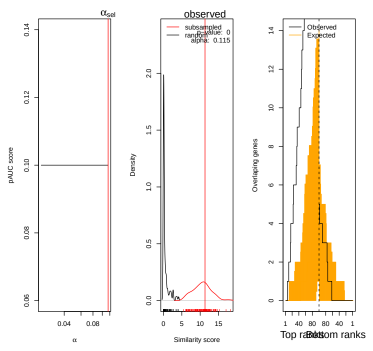
Processing data...Tuning alpha..Significance and genes...

SOGL.res is a list that contains:

- ordering ordered gene lists as a data.frame where columns refer to datasets
- alpha.selected selected value of alpha parameter
- alpha.considered vector of alpha considered for selection
- $\bullet$  pAUC pAUC values related to all alphas considered
- random random scores (permutations of class labels)
- subsample scores after subsampling from each class and dataset
- emp.ci empirical confidence intervals for number of overlapping genes
- ullet common.genes vector of number of overlapping genes
- ullet score observed similarity score
- significance significance of the observed score in form of p-value
- ullet genes genes that account for observed similarity score

The SOGL.res is an object of class SOGLresult for which plot function exist.

```
> par(mfrow = c(1, 3))
> plot(SOGL.res, "alpha selection")
> plot(SOGL.res, "density")
> plot(SOGL.res, "empirical CI")
```



The left graph shows pAUC of all  $\alpha$ 's considered. The selected  $\alpha$  is singed by red vertical line. In the center, one can see estimated densities of random (in black) and subsampled score (in red) for selected  $\alpha$ . The observed similarity score is marked by vertical line. The bottom rugs are scores actually achieved in permutations. The right graph displays the empirical confidence intervals for number of overlapping genes to each position. Observed overlaps are drawn as black step-function.

# RankProduct

#### Introduction

RankProduct is a non-parametric statistic that detects up-regulated and down-regulated genes under one condition against another condition. In our sample data set we look for difference in expression between MSI and MSS colon cancer.

It focuses on genes which are consistently highly ranked in a number of lists, for example genes that are regularly found among top up-regulated genes in many microarray studies. It assumes that under the null hypothesis the order of all items is random then the probability of finding a certain item among the top r of n items in a list is p = r/n. Rank product is defined by multiplying these probabilities  $RP = \prod_i \frac{r_i}{n_i}$ , where  $r_i$  is the rank of the item in the i-th list and  $n_i$  is the total number of the items on i-th list. The smaller the RP value the smaller the probability that the observation of the item at the top of the lists is due to chance. It is equivalent to calculating the geometric mean rank. A list of up- or down-regulated genes are selected based on the estimated percentage of false positive prediction (pfp) - known as false discovery rate (FDR), too.

## Algorithm

Algorithm of the method has five steps:

- $1.\ \,$  Fold-change ratio is calculated in each data set.
- 2. Ranks are assigned (1 for the highest value) according to fold-change ratio.  $r_{gi}$  is rank of gene g in comparison i, where i is from 1 to K, where K is sum of products of number of slides in groups.
- 3. Rank Product for a gene  $(RP_g)$  is calculated as  $\prod_i r_{gi}^{1/K}$
- 4. l permutations of expression values at each microarray slide is performed and all previous steps repeated. We obtain  $RP_g^{(l)}$
- 5. Step 4 is repeated L times to estimate the distribution of  $RP_g^{(l)}$ . This distribution is used to calculate p-value and pfp for each gene.

## Usage

This section depends on the RankProd package, which is free only for non-commercial users. Non-academic users MUST confirm with the authors.

> library(RankProd)

#### Data preparation

In order to run a rank product meta-analysis, user needs to call function RPadvance. It requires three arguments: data, cl and origin. The first one data is the matrix (or data frame) containing the gene expression data that should be analyzed. Each of its rows corresponds to a gene, and each column corresponds to a sample. Second and third argument, cl and origin, are vectors of length ncol(data) containing the class labels of the samples or the origin labels of the samples. Function mergedata returns a list with three slots corresponding to arguments described above. varname argument is a string indicating which columns of the clinical data matrices should be used as class labels.

In cl all 1's refer to MSI samples and all 2's to MSS samples. Similarly in origin, 1 belongs to samples from first data set (denmark), 2 from second data set (australia) and 3 from japan study. You can choose different numbers for labels, but same numbers are always treated like same samples from same class or with same origin.

#### Detecting differentially expressed genes

In this section, we show how the rank product method can be applied to detect differentially expressed gene in our data sets in sence of meta-analysis. It means we will get two separate lists (up- and down-regulated genes separately) not two such lists for each data set. For each gene, one pfp (percentage of false prediction) is computed and used to select significant genes. We can run meta-analysis by

- > RP.out<-RPadvance(rankdata\$dat,rankdata\$cl,rankdata\$origin,num.perm=10,
- + logged=TRUE,na.rm=FALSE,gene.names=rownames(GEDM(ColonData)[[1]]), plot=FALSE)

The data is from 3 different origins

Rank Product analysis for two-class case

Warning: Expected classlabels are 0 and 1. cl will thus be set to 0 and 1.

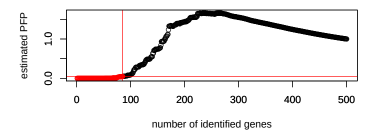
Starting 10 permutations... Computing pfp...

The data are log-transformed, therefore we set logged=TRUE. The number of permutations is default set to 100, you can change it to higher number, if you wish more precise estimates of the pfp. The argument plot=FALSE will prevent the graphical display of the estimated pfp vs. number of identified genes. We will use function plotRP for a such display.

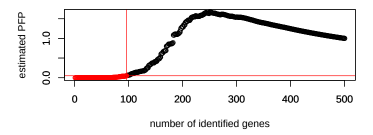
#### Results

> plotRP(RP.out, cutoff=0.05)

#### Identification of Up-regulated genes under class 2



#### Identification of down-regulated genes under class 2



The function plotRP graphically displays the estimated pfp vs. number of identified genes using the output from RPadvance. If cutoff (the maximum accepted pfp) is specified, identified genes are marked in red.

> RankRes<-topGene(RP.out, cutoff=0.05)

Table1: Genes called significant under class1 < class2

Table2: Genes called significant under class1 > class2

#### > head(round(RankRes\$Table1,3))

	gene.index	RP/Rsum	<pre>FC:(class1/class2)</pre>	pfp
228030_at	254	15.955	0.234	0
228915_at	462	26.810	0.407	0
206239_s_at	77	28.268	0.344	0
243669_s_at	237	30.786	0.465	0
213880_at	258	40.144	0.520	0
213385_at	213	43.146	0.456	0
	P.value			
228030_at	0			
228915_at	0			
206239_s_at	0			
243669_s_at	0			
213880_at	0			
213385_at	0			

#### > head(round(RankRes\$Table2,3))

	gene.index	RP/Rsum	FC:(class1/class2)	pfp	P.value
205242_at	257	31.645	3.092	0	0
37145_at	154	34.843	2.678	0	0
209301_at	164	37.957	2.276	0	0
206442_at	280	42.928	2.927	0	0
206391_at	168	49.136	1.836	0	0
204818_at	277	50.370	2.216	0	0

The function topGene is used to output a table of the identified genes from the RP.out. Table contains genes according to other arguments. It is obligatory to specify either the cutoff (the desired significance of the identification) or num.gene (the number of top genes identified), otherwise a error message will be printed and the function will be stopped. If cutoff is selected, user needs to choose between pfp (percentage of false prediction) or pval (p-value). pfp is the default setting, which is selected when no selection is made.

Two tables are output, listing identified up- (Table1: class 1 < class 2) and down- (Table2: class 1 > class 2) regulated genes. There are 5 columns in the table

- 1. gene.index is the gene index in the original data set
- 2. RP/Rsum is the computed rank product for each gene
- 3. FC:(class1/class2) is the computed fold change of the average expression levels under two conditions, which would be converted to the original scale using input logbase (default value is 2) if logged=TRUE is specified
- 4. pfp is the estimated pfp value for each gene in the list if that gene serves as the cutoff point
- 5. P.value is the associated P-values for each gene

### Wrapper function

Function RankProduct needs no data preparation and provides the tables of identified up- and down- regulated genes.

```
> rp <- RankProduct(ColonData, "MSIstatus", num.perm = 10)
The data is from 3 different origins
Rank Product analysis for two-class case
Starting 10 permutations...
Computing pfp...
Table1: Genes called significant under class1 < class2</pre>
Table2: Genes called significant under class1 > class2
```

# Z-statistic - posterior mean differential expression

#### Introduction

The main idea of this method is that one can use data from one study to construct a prior distribution of differential expression and thus utilize the posterior mean differential expression, weighted by variances, whose distribution is standard normal distribution due to classic Bayesian probability calculation.

It is based on assumption that gene expression is normally distributed with mean  $\mu_g$  and SD  $\sigma_g^2$  and that we can estimate  $\sigma_g^2$  by pooling together all genes with similar levels of mean intensity. The difference in gene expression is tested by

$$Z = \frac{D}{\sigma_D} = \frac{\overline{X}_1 - \overline{X}_2}{\sqrt{\frac{\sigma_1^2}{n_1} + \frac{\sigma_2^2}{n_2}}} \sim N(0, 1),$$

where  $\overline{X}_1$  and  $\overline{X}_2$  denotes mean gene expression values in classes,  $\sigma_1^2$  and  $\sigma_2^2$  denotes the estimated SD in classes and  $n_1$  and  $n_2$  denotes the number of samples in classes.

## Usage

#### Data preparation

Because the same number of samples in each class and study is used in primary publication of the method [16], we will first look at number of samples in our data.

> ColonData

Dataset Denmark containing 500 probes and 77 samples. Sumarization of samples:
MSIstatus
----MSI MSS

```
39
    38
age
  Min. 1st Qu. Median
                           Mean 3rd Qu.
                                           Max.
  36.00
          45.00
                 54.00
                          54.92
                                  62.00
                                          80.00
Dataset Australia containing 500 probes and 36 samples.
Sumarization of samples:
MSIstatus
MSI MSS
  5 31
age
   Min. 1st Qu. Median
                           Mean 3rd Qu.
                                           Max.
                  66.00
  50.00
         57.75
                          66.36
                                  73.75
                                          85.00
Dataset Japan containing 500 probes and 41 samples.
Sumarization of samples:
location
  distal proximal unknown
               11
MSIstatus
_____
MSI MSS
 16
    25
age
   Min. 1st Qu.
                 Median
                           Mean 3rd Qu.
                                           Max.
  40.00
          47.00
                  53.00
                          53.61
                                  60.00
                                          65.00
```

The smallest value in the tables above is 5, therefore we will randomly choose 5 samples in each class and data set. Function posterior.mean performs such data reduction and Z-statistic calculation. It has three required arguments: a data set as MetaArray object (data), name of clinical parameter with class labels (varname) and number of samples to be selected (nsamp).

#### Detecting differentially expressed genes

```
We apply this method by

> z.stat<-posterior.mean(ColonData, "MSIstatus", 5)

0 marked as 0

1 marked as 1

Contrast will be 1 - 0
```

#### Results

> head(round(z.stat,3))

```
Zscore Pvalue
1552281_at 3.483 0.000
1552365_at -4.342 0.000
1552485_at -2.926 0.003
1552502_s_at -0.984 0.325
1552546_a_at -0.483 0.629
1552553_a_at -1.012 0.311
```

Only values of Z-statistic ( ${\tt Zscore}$ ) and their p-values ( ${\tt Pvalue}$ ) are outputs from this method.

## Notes and discussion

This implementation expects either same microarray platform or same scale of expression values (like after POE transformation [17]) in all data sets.

# VennMapping

#### Introduction

VennMapping [20] is a method based on Venn diagrams and contingency tables. It looks for number of common genes in pairs of gene lists, statistical significance of observed match and returns also names of the common genes.

## Algorithm

Algorithm of this method consists of three steps:

- 1. Calculation of fold-change in each data set.
- 2. Selection of significant (interesting) genes.
- 3. Comparison of gene lists pairs.

## Usage

#### Data preparation

Function fold.change calculates mean fold-change in MetaArray object. It has two arguments: data set (e.g. ColonData) and name of the column with class labels to be used (e.g. "MSIstatus"). It assumes data are on  $log_2$  scale.

```
> fc<-fold.change(ColonData, "MSIstatus")
```

Function gene.select.FC selects significant/interesting genes from mean fold-change matrix with rows referring to genes and columns to data sets. The user has to specify (apart from mean fold-change matrix) a cutoff for selection. The cutoff is on  $log_2$  scale, too. We chose 1 for genes with at least 2-fold change in expression.

```
> list<-gene.select.FC(fc,1)
> summary(list)
```

```
Length Class Mode

Denmark 33 -none- character

Australia 27 -none- character

Japan 35 -none- character
```

Object list is a list in which each slot contains names of selected genes in one study. For example from the print above 33 genes have been selected in denmark data set.

#### Detecting differentially expressed genes

Now, we can move on comparison of selected gene lists in pairs of data sets. There are three functions to perform such a analysis: conting.tab, Z and gene.list. conting.tab returns contingency table with number of common genes. Z provides Z statistic to measure significance of observed number of common genes and gene.list outputs table with names of common genes. All of them have one argument same - it is a list object with names of selected genes in individual data sets. For function Z one additional argument is necessary - the number of genes involved in meta-analysis.

#### > conting.tab(list)

	Denmark	Australia	Japan
Denmark	NA	12	16
${\tt Australia}$	12	NA	7
Japan	16	7	NA
> Z(list,	n = 500	)	

```
        Denmark
        Australia
        Japan

        Denmark
        NA
        7.920245
        9.320174

        Australia
        7.869850
        NA
        3.821593

        Japan
        9.340196
        3.854327
        NA
```

#### > gene.list(list)

```
Denmark

NA

Australia "205009_at;206239_s_at;37145_at;205044_at;213385_at;228030_at;205242_at;204818_a

Japan "202803_s_at;230964_at;213915_at;206239_s_at;1556055_at;1552281_at;37145_at;2093

Australia

Denmark "205009_at;206239_s_at;37145_at;205044_at;213385_at;228030_at;205242_at;204818_a

Australia NA
```

```
Japan "206239_s_at;209583_s_at;37145_at;228030_at;206442_at;210143_at;230793_at"
```

Denmark "202803\_s\_at;230964\_at;213915\_at;206239\_s\_at;1556055\_at;1552281\_at;37145\_at;2093 Australia "206239\_s\_at;209583\_s\_at;37145\_at;228030\_at;206442\_at;210143\_at;230793\_at"

ustrarra ~200259\_s\_at;209505\_s\_at;5/145\_at;220050\_at;200442\_at;210145\_at;250/95\_at

Japan NA

#### Wrapper function

Function VennMapper provides both gene selection in each dataset and all comparisons of selected gene lists.

```
> vm <- VennMapper(ColonData, "MSIstatus", 1)</pre>
```

#### \$conting.tab

	Denmark	Australia	Japan
Denmark	NA	12	16
Australia	12	NA	7
Japan	16	7	NA

#### \$z.score

Denmark Australia Japan NA 14.964793 17.550323 Denmark NA 8.098665 Australia 14.93428 17.56230 8.120747 Japan NA

#### \$genes

Denmark

Denmark

Denmark

Australia "205009\_at;206239\_s\_at;37145\_at;205044\_at;213385\_at;228030\_at;205242\_at;204818\_a Japan "202803\_s\_at;230964\_at;213915\_at;206239\_s\_at;1556055\_at;1552281\_at;37145\_at;2093

"205009\_at;206239\_s\_at;37145\_at;205044\_at;213385\_at;228030\_at;205242\_at;204818\_a

Australia

Australia NA

"206239\_s\_at;209583\_s\_at;37145\_at;228030\_at;206442\_at;210143\_at;230793\_at" Japan

Denmark "202803\_s\_at;230964\_at;213915\_at;206239\_s\_at;1556055\_at;1552281\_at;37145\_at;2093

Australia "206239\_s\_at;209583\_s\_at;37145\_at;228030\_at;206442\_at;210143\_at;230793\_at"

Japan

attr(,"class")

[1] "VennMapper.res"

# **MAP-Matches**

#### Introduction

Meta-Analysis Pattern Matches (MAP-Matches) [21] is a method that extends VennMapping [20] and meta-profiling [22]. It is designed to analyze more distinct microarray data (search for common molecular mechanism in all types of cancer). It assumes same gene set in all data sets.

## Algorithm

Algorithm of this method has five steps:

- 1. Calculation of T-statistic for each two classes in each data set.
- 2. Building matrix of T-statistics (T-matrix) with rows referring to genes and columns to pairs of classes and data set.
- 3. Selection of threshold for T-statistic.
- 4. Transformation of T-matrix into a binary matrix: 1 for T-statistics above threshold, 0 for T-statistics below threshold.
- 5. Statistical analysis of transformed T-matrix (more details in Usage section).

## Usage

#### Data preparation

The analysis starts with calculation of T-statistics. Function meta.test returns a list with two slots: matrix of test statistics (test) and matrix of p-values (p). In each of the matrices rows correspond to genes and columns to data sets. We need only test slot for this method. Argument varname is a string indicating which column of clinical data should be used as class labels.

> stat.real<-meta.test(ColonData, varname = "MSIstatus")\$test

#### Detecting differentially expressed genes

The do not select significant genes in each study we only set threshold for T-statistics. We decided for 95 % quantile (500 genes in 3 datasets with 95% quantile leaves top 75 significance tests for the analysis). If there is a lot of genes and datasets in the meta-analysis one should consider higher quantiles.

```
> stat <- c(stat.real)
> quan <- quantile(abs(stat), seq(0.00, 1.00, 0.0001))
> T.default <- quan["95.00%"]</pre>
```

Now, we transform stat.real (T-matrix) into a binary matrix. We replace T-statistics above threshold with 1 and below with 0.

```
> value.dis <- apply(stat.real,MARGIN=c(1,2), function(x) ifelse(abs(x)>T.default,1,0))
> rownames(value.dis)<-rownames(GEDM(ColonData)[[1]])</pre>
```

>	head	(value,	dis
_	Heau	lvarue.	· uıbı

	Denmark	Australia	Japan
217562_at	0	0	0
203766_s_at	0	0	0
1554394_at	0	0	0
212662_at	0	0	0
1555370_a_at	0	0	0
240574_at	1	0	0

Each row value.dis is called a meta-analysis pattern. We are going to analyze their occurrence, significance and genes they occur at. Function ratio provides basic summarization of value.dis.

```
> results<-ratio(value.dis)
> summary(results)
```

```
Length Class Mode
n 3 -none- numeric
X.string 54 -none- character
p.strong 7 -none- numeric
p.soft 7 -none- numeric
```

In **results** we can find: number of genes with T-statistic sufficiently high in each study n, patterns observed in data (X.String), probability of observing strong match (p.strong) and probability of observing soft match (p.soft). We say two patterns match strongly if they are equal. The rule for soft match is weaker as only 1's in patterns must match.

Function MAPmatrix calculates a matrix with rows corresponding to patterns and four columns: unique patterns that are being observed in our data (uniqe.pat), number of observed soft matches with the pattern (n.soft), number of observed strong matches (n.strong) and number of 1's in the pattern n.sig).

```
> MAPmat<-MAPmatrix(value.dis)
```

<sup>&</sup>gt; MAPmat

	unique.pat	${\tt n.soft}$	n.strong	n.sig
100	100	38	22	1
110	110	7	4	2
010	010	15	6	1
101	101	12	9	2
001	001	22	8	1
011	011	5	2	2
111	111	3	3	3

Only pattern with multiply 1's are connected with common molecular mechanism and we will focus on them in the rest of analysis.

```
> MAPmat2<-MAPmat[MAPmat$n.sig>1,]
> unique.pat<-as.character(MAPmat2[,1])</pre>
```

We assume that sufficiently high number of strong matches may provides evidence of common molecular mechanism. Functions MAPsig1 and MAPsig2 perform statistical analysis to answer whether we observe significant number of matches or not. The statistical analysis can be done in two ways (both based on permutation testing): we either permute columns of T-matrix (in binary form) or permute class labels in data sets and repeat the whole procedure with same threshold for T-statistics. The former is implemented in MAPsig1 and the latter in MAPsig2. Function test.group.shuffle calculates T-statistics with permuted class label repeatedly.

```
> p1<-MAPsig1(unique.pat, value.dis, iter=100)
```

```
Permutation:
100
> p1
  p.soft p.strong
             0.00
       0
2
             0.00
       0
3
       0
             0.12
             0.00
> out<-test.group.shuffle(ColonData, "MSIstatus", B = 100)</pre>
> p2 <- MAPsig2(out, value.dis, unique.pat, B = 100)
> p2
  permu.soft permu.strong
```

	F	F
1	0	0.03
2	0	0.00
3	0	0.17
4	0	0.00

Both p1 and p2 have same structure: rows refer to patterns and columns to statistical signifficance of observed soft (p.soft or permu.soft) or strong (p.strong or permu.strong) matches.

#### Results

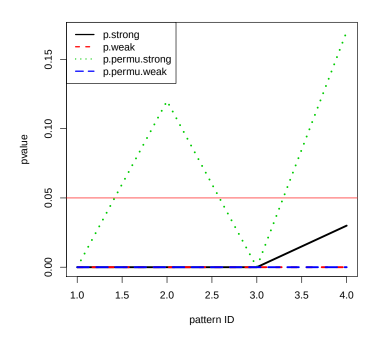
Finally, we will bind all necessary outputs together.

```
> resx <- cbind(MAPmat2, p1, p2)
> colnames(resx) <-c(colnames(MAPmat2), "p.strong",
+ "p.weak", "p.permu.strong", "p.permu.weak")
> intx <- t(as.matrix(resx[which(resx[,4]<0.06),]))
> t(resx)
```

	110	101	011	111
unique.pat	"110"	"101"	"011"	"111"
n.soft	" 7"	"12"	" 5"	" 3"
n.strong	"4"	"9"	"2"	"3"
n.sig	"2"	"2"	"2"	"3"
p.strong	"0"	"0"	"0"	"0"
p.weak	"0.00"	"0.00"	"0.12"	"0.00"
p.permu.strong	"0"	"0"	"0"	"0"
p.permu.weak	"0.03"	"0.00"	"0.17"	"0.00"

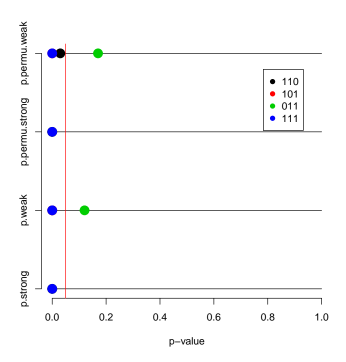
We can plot p-values by

> plotpattern(resx, method=1)



or

> plotpattern(resx, method=2)



Until now, we have only found out that for some patterns there is significantly high count of strong or soft matches being observed. Obviously we want to know expression of which genes is changed in these patterns. Function MAP.genes returns a list in which each slot contains list of genes involved in one pattern. If argument files is set to TRUE a files with gene names are also saved.

```
> probs<-MAP.genes(resx,value.dis, files=FALSE)
> names(probs)<-rownames(resx)
> summary(probs)

Length Class Mode
110 7 -none- character
101 12 -none- character
011 5 -none- character
111 3 -none- character
```

probs is a list with each slot referring to one pattern and list of gene names is stored there. The pattern has been observed at these genes.

#### Wrapper function

Function MAP.Matches provides all steps of MAP-Matches method in one-line command. In

```
> map <- MAP.Matches(ColonData, "MSIstatus",
+ t.cutoff = "95.00%", nperm = 100, sig.col = "p.col.strong")</pre>
```

```
Examining the data...
Statistical analysis...
Permutation:
50
100
Permuting class labels in dataset:
Denmark
Australia
Japan
Permutation:
50
100
```

the first argument is the dataset (MetaArray object), the second one is a column name of the class labels, nperm denotes the number of permutations and the last one sig.col is column name for selection of significant patterns - "p.col.strong" refer to the p-value of the strong matches when permutation of columns are used.

### Chapter 9

## **METRADISC**

#### Introduction

METRADISC [23] is unique among rank-based methods because it provides an estimate of heterogeneity as one of its outputs. Additionally the method can deal with genes which are being measured in only some of the studies. The implementation available in MAMA package is restricted to genes common in all microarray studies analyzed.

#### Algorithm

- 1. Gene Ranking In microarray analysis we usually test samples for a large number of genes. The results provide for each gene a test statistic and its statistical significance (p-value). Therefore we can rank the tested genes in each study based on direction in expression change and statistical significance. If there are n genes being tested, the highest rank n is given to the gene that shows the lowest p-value and it is up-regulated in diseased samples. Then follow all other up-regulated genes ranked according to increasing p-value. These are followed by down-regulated genes and the lowest rank (1) is given to gene that shows the lowest p-value and is down-regulated in diseased samples. Genes with equal p-values are assigned tied ranks.
- 2. The Average Rank and Heterogeneity metrics In this step we compute a average rank and heterogeneity metrics. The average rank  $R^*$  is defined as  $R^* = \frac{\sum_{i=1}^s R_i}{s}$ , where  $R_i$  is the rank of the gene in study i and s is total number of studies (i=1,2,...,s). The heterogeneity metrics  $Q^*$  is given by formula  $Q^* = \sum_{i=1}^s (R_i R^*)^2$ , it is actually generalization of Cochran's Q statistic.
- 3. Monte Carlo permutation test To obtain statistical significance for average rank and heterogeneity metrics we randomly permute the ranks of each study and the stimulated metrics are calculated. Then we repeat the procedure to generate null distribution for the metrics. Each variable is then tested against the corresponding null distribution. We are interested

genuinely in four statistical significances: for high average rank, for low average rank, for high heterogeneity and for low heterogeneity. Distinction between high and low average rank is important as we want to keep the direction of effect in mind. Ignoring it can lead to spurious results that a gene is consistently significant even if it is up-regulated in one study and down-regulated in second one. On the other hand, statistically low heterogeneity may suggest consistent results among different studies. The statistical significance for high average rank  $(R^*)$  is defined as the percentage of simulated metrics that exceed or are equal to the observed  $(R^*)$ . The statistical significance for low average rank  $(R^*)$  is defined as the percentage of simulated metrics that are below or equal to the observed  $(R^*)$ . Significance of heterogeneity is defined analogously.

#### Usage

#### Data preparation

We will start with computing test statistic and p-value for each gene and data set. Function meta.test returns a list with two slots: data frame of test statistics and data frame of p-values. In each of the matrices rows correspond to genes and columns to data sets. Argument varname is a string indicating which column of clinical data should be used as class labels.

```
> metra<-meta.test(ColonData, varname = "MSIstatus")
> head(metra$test)
```

	Denmark	Australia	Japan
217562_at	-2.1666481	1.0669144	-0.9286918
203766_s_at	0.8318955	0.1014991	-0.9459710
1554394_at	1.2176000	4.0282590	1.1763280
212662_at	1.9755557	1.3046695	2.5983263
1555370_a_at	4.3678119	-0.6042763	2.7174563
240574_at	5.1746999	1.6404196	1.8830312

#### Detecting differentially expressed genes

Now, we can proceed to ranking genes. Function rank.genes.adv ranks the genes as described in Algorithm section above.

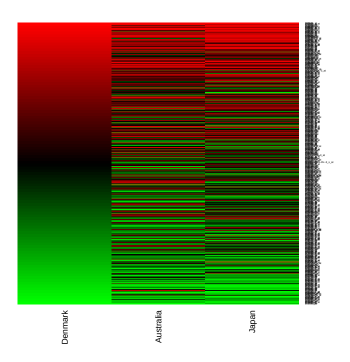
```
> RANK<-rank.genes.adv(metra)
> head(RANK)
```

	Denmark	Australia	Japan
217562_at	105	368	156
203766_s_at	326	257	153
1554394_at	348	488	380
212662_at	390	397	442
1555370_a_at	473	170	445
240574_at	484	424	419

The genes ranks can be visualized by

```
> RANK2<-RANK[order(RANK[,1]),]</pre>
```

- > colnames(RANK2)<-c("Denmark","Australia", "Japan")</pre>
- > heatcol<-colorRampPalette(c("Green", "Black", "Red"))(100)</pre>
- > metaheat(as.matrix(RANK2), col=heatcol, c.cex = 0.8)



The next step is to compute average rank  $R^*$  and heterogeneity metric  $Q^*$  for each gene.

```
> RQ<-compute.RQ(RANK)
```

> head(round(RQ,1))

```
r.star q.star
217562_at 209.7 38904.7
203766_s_at 245.3 15168.7
1554394_at 405.3 10762.7
212662_at 409.7 1592.7
1555370_a_at 362.7 56072.7
240574_at 442.3 2616.7
```

And finally we use function MCtest to perform Monte Carlo permutation test. Function requires the observed ranks (RANK), observed average rank and heterogeneity metric (RQ) and number of permutations (nper) as arguments. Number of permutations depends on the required accuracy for the final p-values. 1/nper is the accuracy for the final p-values. For example with 1000 permutations the p-values are calculated with three decimal places.

> MC<-MCtest(RANK,RQ, nper=1000)

```
100
      200
            300
                   400
                         500
                               600
                                     700
                                            800
                                                  900
                                                         1000
> head(MC)
             R.high R.low Q.high Q.low
217562_at
              0.667 0.333
                            0.468 0.532
203766_s_at
              0.523 0.477
                            0.747 0.253
1554394_at
              0.041 0.959
                            0.801 0.199
212662_at
              0.033 0.967
                            0.950 0.050
1555370_a_at
              0.080 0.920
                            0.303 0.697
              0.011 0.989
240574_at
                            0.942 0.058
```

#### Results

The command below creates a character vector of genes with significant average ranks and low heterogeneity. The selected threshold for statistical significance is 0.05.

```
> METRA<-c(rownames(MC)[MC[,1]<0.05 & MC[,4]<0.05],
+ rownames(MC)[MC[,2]<0.05 & MC[,4]<0.05])
> METRA[1:10]

[1] "239442_at" "234207_at" "225802_at" "230964_at"

[5] "1563364_at" "206239_s_at" "236223_s_at" "222551_s_at"

[9] "1560119_at" "201279_s_at"
```

#### Wrapper function

Function METRADISC performs all steps of this method in one command.

```
> metra<-METRADISC(ColonData,
                              "MSIstatus", nperm = 1000)
100
      200
            300
                  400
                        500
                              600
                                    700
                                          800
                                                 900
                                                       1000
> str(metra)
List of 3
 $ ranks :'data.frame':
                               500 obs. of 3 variables:
  ..$ Denmark : num [1:500] 105 326 348 390 473 484 83 126 498 482 ...
  ..$ Australia: num [1:500] 368 257 488 397 170 424 179 187 490 495 ...
               : num [1:500] 156 153 380 442 445 419 88 187 390 469 ...
         : num [1:500, 1:2] 210 245 405 410 363 ...
 $ RQ
  ..- attr(*, "dimnames")=List of 2
  ....$ : chr [1:500] "217562_at" "203766_s_at" "1554394_at" "212662_at" ...
  .. ..$ : chr [1:2] "r.star" "q.star"
 $ MCtest: num [1:500, 1:4] 0.683 0.524 0.031 0.024 0.111 0.006 0.934 0.845 0.002 0 ...
  ..- attr(*, "dimnames")=List of 2
  ....$ : chr [1:500] "217562_at" "203766_s_at" "1554394_at" "212662_at" ...
  ....$ : chr [1:4] "R.high" "R.low" "Q.high" "Q.low"
 - attr(*, "class")= chr "METRADISC.res"
```

It returns a list with ranks for each gene and dataset ranks, average rank and heterogeneity for each gene RQ and results of the Monte Carlo test MCtest.

# Part I Results combination

In this part we are going to compare and combine outputs from all methods so we can look and changes in gene expression in various ways.

We are going to start with lists of differentially expressed genes, because this is the only one output common for all methods mentioned in this vignette. We will merge all lists into one variable via function join.DEG. The function requires a complete list of genes involved in meta-analysis so it can map indices to gene names like for example function pvalcombination provides. Because the same set of genes was measured in each data set we can arbitrarily choose one data set.

```
> lists<-join.DEG(pvalt, ESt, ScoresFDR, SOGL.res, RankRes, z.stat,
+ probs,genenames=rownames(GEDM(ColonData)[[1]]),
+ type=c(1,1,3,4,5,6,7), cutoff=0.02)
> #or
> lists2<-join.DEG(pval, es, es2, SOGL.res, rp, z.stat, map,
+ genenames=rownames(GEDM(ColonData)[[1]]), cutoff=0.05)
> names(lists)<-c("PvalCom", "ESCom", "ESCom2", "OrderedList", "RankProduct",
+ "Z-stat", "MAP")
> summary(lists)
```

	Length	Class	Mode
PvalCom	215	-none-	character
ESCom	109	-none-	character
ESCom2	178	-none-	character
OrderedList	11	-none-	character
RankProduct	178	-none-	character
Z-stat	149	-none-	${\tt character}$
MAP	18	-none-	character

The argument type is a numeric identification of the method the object originates. When wrapper function are used there is no need for this argument.

Now, we will transform this list to a binary matrix where rows refer to genes and columns to method and 1 means that the gene was identified as a differentially expressed gene in the method. Function make.matrix provides such transformation.

> MAT<-make.matrix(lists)
> MAT[1:5,1:5]

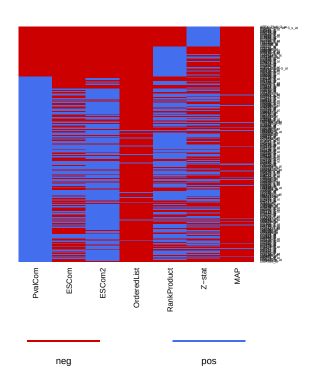
	PvalCom	ESCom	ESCom2	OrderedList	RankProduct
1554394_at	1	0	0	0	0
212662_at	1	0	1	0	0
1555370_a_at	1	0	1	0	1
240574_at	1	1	1	0	1
203553_s_at	1	1	1	0	0

It is very popular to visualize results of microarray analysis as a heatmap. A heatmap is a graphical representation for a numeric matrix where values are presented as colors. Gene expression values are usually used in microarray analysis. In these pictures colors go continuously from green (for down-regulation) through black (for no change in gene expression) to red (for up-regulation).

There are several R-packages which implement plotting heatmaps in slightly different way. Functions metaheat and metaheat2 are modification of two of them, so a discrete set of colors (only two in metaheat but even several in metaheat2) can be used with an appropriate legend.

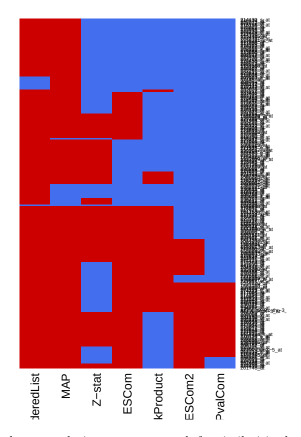
Function metaheat has three arguments: a data matrix (MAT), a number defining position of legend (legend=1 is legend drawn below the picture) and vector of colors (col).

> metaheat(MAT, legend=1, col=c("red3", "royalblue2"))



Function metaheat2 has as many arguments as heatmap.2 form gplots package and two more. Argument legend.names is a character vector with labels to be used in legend. Setting discret=TRUE will indicate that legend for discrete values should be drawn.

```
> metaheat2(MAT, col=c("red3", "royalblue2"), legend.names=c("DEG","noDEG"),
+ discrete=TRUE, trace="none", dendrogram="none")
```



The user can perform cluster analysis on MAT to search for similarities between methods or genes.

We can look at number of genes found by number of methods by

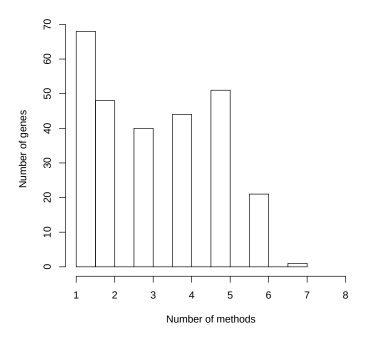
#### > dim(MAT)

#### [1] 273 7

According to the outsprint above, eight different methods have found 217 differentially expressed genes.

The histogram below shows that the most of the genes have been selected in only one method.

- > n.met<-apply(MAT,1,sum)</pre>
- > hist(n.met, main="", xlab="Number of methods", ylab="Number of genes", xlim=c(1,8))



n.met is a numeric vector of number of methods that identified the gene as differentially expressed.

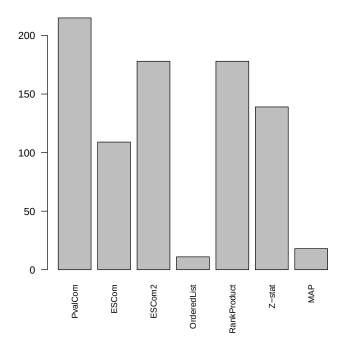
Next, we can look for example how many genes have been found as differentially expressed in at least 6 methods.

> dim(MAT[n.met>5,])

#### [1] 22 7

On the other hand, we can find out how many genes have been found by a method.

- > n.gen<-apply(MAT,2,sum)</pre>
- > barplot(n.gen, cex.names=0.8, las=2)



Function contig.tab provides a number of genes common in two gene lists. It can be applied to lists, too.

- > TAB<-conting.tab(lists)
- > TAB[1:5,1:5]

	PvalCom	ESCom	ESCom2	${\tt OrderedList}$	${\tt RankProduct}$
PvalCom	NA	109	178	11	143
ESCom	109	NA	109	11	97
ESCom2	178	109	NA	11	134
OrderedList	11	11	11	NA	11
RankProduct	143	97	134	11	NA

#### Expression of one gene

In this section we are going to focus on one gene and to look at its expression change from different points of view. The different points of view are represented by different approaches used in the methods.

First we will join all the available results to one list and then select only rows for one gene.

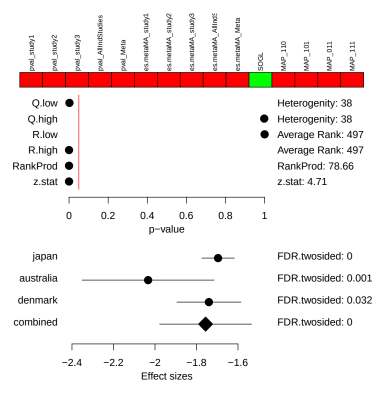
- > results<-join.results(pvalt, ESt, theScores, ScoresFDR\$two.sided, SOGL.res,
- + RankRes, z.stat, probs, MC, RQ, type=c(1,1,5,5,2,3,5,4,5,5),
- + genenames=rownames(GEDM(ColonData)[[1]]))
- > #or
- > results2<-join.results(pval, es, es2, SOGL.res, rp, z.stat, map, metra)

```
> gene<-metagene("203008_x_at",results2)</pre>
> names(gene)<-c("pval", "es.metaMA", "es.GeneMeta", "SOGL", "RP", "z.stat",
   "MAP", "METRADISC")
> gene
$pval
       study1
                     study2
                                    study3 AllIndStudies
     1.000000
                   1.000000
                                  1.000000
                                                1.000000
         Meta TestStatistic
     1.000000
                   8.732214
attr(,"class")
[1] "metaMA"
                  "metaMA.gene"
$es.metaMA
       study1
                     study2
                                    study3 AllIndStudies
     1.000000
                   1.000000
                                  1.000000
                                                1.000000
         Meta TestStatistic
     1.000000
                  -8.492688
attr(,"class")
[1] "metaMA"
                  "metaMA.gene"
$es.GeneMeta
                                    zSco_Ex_3
     zSco_Ex_1
                    zSco_Ex_2
                                                        zSco
 -6.4432924942 -3.7652548865 -4.6965956199 -8.8027593869
                        MUsds
        MUvals
                                        Qvals
                                                          df
 -1.7730928661
                 0.2014246656
                                0.2626001205
                                                2.0000000000
      Qpvalues
                        Chisq
                                 Effect_Ex_1
                                                 Effect_Ex_2
                 0.000000000 -1.7184785951 -2.0248658800
  0.8769545957
   Effect_Ex_3 EffectVar_Ex_1 EffectVar_Ex_2 EffectVar_Ex_3
 -1.7586784853
                 0.0711332351
                                 0.2892036455
                                                0.1402189026
                                    zSco_Ex_2
                                                    FDR_Ex_2
     zSco_Ex_1
                     FDR_Ex_1
 -6.4432924942
                 0.000000000 -3.7652548865
                                                0.0325000000
     zSco_Ex_3
                     FDR_Ex_3
                                         zSco
                                                         FDR
 -4.6965956199
                 0.0007142857
                                -8.8027593869
                                                0.000000000
        {\tt MUvals}
                        MUsds
                                                          df
                                        Qvals
 -1.7730928661
                 0.2014246656
                                 0.2626001205
                                                2.0000000000
      Qpvalues
                        Chisq
  0.8769545957
                 0.000000000
attr(,"class")
[1] "ES.GeneMeta"
                       "ES.GeneMeta.gene"
$SOGL
NULL
<0 rows> (or 0-length row.names)
$RP
                              RP/Rsum FC:(class1/class2)
        gene.index
          415.0000
                              78.6607
                                                   0.5803
               pfp
                              P.value
            0.0000
                               0.0000
```

```
attr(,"class")
[1] "RankProduct"
                       "RankProduct.gene"
$z.stat
                           Pvalue
              Zscore
203008_x_at 4.708621 2.493978e-06
$MAP
110 101 011 111
TRUE TRUE TRUE TRUE
attr(,"class")
[1] "MAP.Matches"
                       "MAP.Matches.gene"
$METRADISC
r.star q.star R.high
                          R.low
                                 Q.high
                                          Q.low
497.000 38.000
                  0.000
                          1.000
                                  0.998
                                          0.002
attr(,"class")
[1] "METRADISC"
                     "METRADISC.gene"
attr(,"class")
[1] "metagene"
```

This output provides much of the information available on the gene through all the described methods. It is a rather complicated structure, so we will try to represent it graphically in comprehensible form.

```
> plotgene(gene, type=c(0,0,1,4,5,6,7,10), datalabels=c("denmark", "australia", + "japan", "combined"))
```



The picture above shows in top part occurrence of gene in lists of selected genes in some methods. The dark box means that the gene is present in the list. Values from objects: pvalt, ESt, x.z and probs are used in here.

The middle part is dedicated to p-values available in meta-analysis. Specific values of the statistics can be found on the right side of the chart. The vertical dashed line denotes the signifficance threshold 5%. P-values from MC, RankRes and z.stat are drawn in here.

Combination of effect size is plotted in the bottom graph. The point marks the effect size. Horizontal lines denote the variance of effect size. Statistical significance of the difference in gene expression (FDR adjusted) can be found on the right side of the chart. This graph uses values from the Scores and ScoresFDR.

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