database releases. The biomaRt package is designed so that new BioMart databases can be automatically included, once they provide public MySQL access or have an active BioMart web service. The tight integration of large public databases with data analysis in R makes biomaRt a powerful platform for biological data integration and data mining.

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Identifying Interesting Genes with siggenes

by Holger Schwender, Andreas Krause and Katja Ickstadt

A common and important task in microarray experiments is the identification of genes whose expression values differ substantially between groups or conditions. Finding such differentially expressed genes requires methods that can deal with multiple testing problems in which thousands or even tens of thousands of hypotheses are tested simultaneously.

Usually, a statistic appropriate for testing if the expression levels are associated with a covariate of interest and the corresponding p-value are computed for each gene. Afterwards, these raw p-values are adjusted for multiplicity such that a Type I error rate is strongly controlled at a pre-specified level of significance. The classical example of such an error rate is the family-wise error rate (FWER), i.e. the probability of at least one false positive. This error rate, however, might be too conservative for a situation in which thousands of hypotheses are tested and several tens of genes should be identified. In the analysis of microarray data, another error rate has hence become very popular: The False Discovery Rate (FDR) which is loosely spoken the expected proportion of false positives among all rejected null hypotheses, i.e.

identified genes.

There are, however, other ways to adjust for multiplicity: For example, QQ plots or the Bayesian framework can be employed for this purpose. If the observed test statistics are plotted against the values of the test statistics that would be expected under the null hypothesis most of the points will approximately lie on the diagonal. Those points that differ substantially from this line correspond to genes that are most likely differentially expressed. The Significance Analysis of Microarrays (SAM) proposed by Tusher et al. (2001) can be used to specify what "differ substantially" means. While Tusher et al. (2001) base their analysis on a moderated *t* statistic, Schwender et al. (2003) compare this approach with a SAM version based on Wilcoxon rank sums.

Efron et al. (2001) use an empirical Bayes analysis (EBAM) to model the distribution of the observed test statistics as a mixture of two components, one for the differentially expressed genes and the other for the not differentially expressed genes. Following their analysis, a gene is called differentially expressed if the corresponding posterior probability is larger than 0.9.

Both SAM and EBAM are implemented in the

Bioconductor package **siggenes**. In this article, we, however, will concentrate on SAM. In the following, we briefly describe the SAM procedure, its implementation in **siggenes** (for more details, see Schwender et al., 2003) and the test statistics already available in this package. Afterwards, we show how you can write your own function for other testing situations. Finally, we will give an example of how sam can be applied to gene expression data.

Significance Analysis of Microarrays

In SAM, a statistic d appropriate for testing if there is an association between the expression levels and the covariate of interest is computed for each of the m genes. These observed test scores are sorted and plotted against the scores expected under the null hypothesis, where the expected test scores $\bar{d}_{(i)}$, $i=1,\ldots,m$, are computed as follows: If the null distribution is known, then $\bar{d}_{(i)}$ is the (i-0.5)/m quantile of this null distribution. Otherwise, $\bar{d}_{(i)}$ is assessed by

- generating *B* permutations of the group labels,
- computing the *m* test statistics and sorting them for each of the *B* permutations,
- averaging over the Bi^{th} -smallest scores.

Two lines parallel to the diagonal in a distance of Δ are then drawn into this plot called the SAM plot. Any gene that has a d value

- larger than or equal to the d value of the gene, say $d_{\rm up}$, that corresponds to the left-most point on the right side of the origin that lies above the upper Δ line,
- smaller than or equal to the d value of the gene, say d_{low} , that corresponds to the right-most point on the left side of the origin that lies below the lower Δ line

is called differentially expressed. Afterwards, the FDR is estimated by

- counting how many of the mB permuted test scores are larger than or equal to $d_{\rm up}$ or smaller than and equal to $d_{\rm low}$, and dividing this number by B,
- dividing this average by the number of identified genes,
- multiplying this ratio by the prior probability that a gene is not differentially expressed (by default, sam estimates this probability by the procedure of Storey and Tibshirani, 2003).

This procedure is repeated for several values of Δ and the value of Δ is chosen that provides the best balance between the number of identified genes and the estimated FDR.

The following test statistics can be called in sam by setting the argument 'method' to

- d.stat: Moderated *t* and *F* statistics. The "usual" *t* or *F* statistics are computed if the fudge factor 's0' is set to zero. (The fudge factor is added to the denominator of the statistics to prevent genes with very low expression levels to become differentially expressed. For details, see Tusher et al., 2001).
- wilc.stat: Wilcoxon rank sums for one and two class analyses.
- cat.stat: Pearson's χ^2 -statistic for testing categorical data such as SNP (Single Nucleotide Polymorphism) data (Schwender, 2005).

Writing Your Own Test Score Function

It is also possible to write your own function for another testing situation and use this function in sam. This function must have as input the two required arguments

- 'data': A matrix or data frame containing the data. Each row of this data set should correspond to one of the *m* variables, i.e. genes, and each column to one of the *n* observations.
- 'c1': A vector consisting of the class labels of the observations.

The function can also have additional optional arguments that can be called in sam.

The output of this function must be a list consisting of the following objects

- 'd': A numeric vector containing the test scores of the genes.
- 'd.bar': A numeric vector of length na.exclude(d) consisting of the sorted test scores expected under the null hypothesis.
- 'p.value': A numeric vector of the same length and order as 'd' containing the p-values of the genes.
- 'vec.false': A numeric vector of the same length as 'd' consisting of the one-sided numbers of falsely called genes (for more details, see Schwender et al., 2003, and the help files of sam).
- 's': A numeric vector containing the standard errors of the expression values.

's0': A numeric value specifying the fudge factor.

'mat . samp': A $B \times n$ matrix containing the permuted class labels.

'msg': A character vector containing messages that are displayed when the SAM specific S4 methods print and summary are called.

'fold': A numeric vector containing the fold changes of the genes. Should be set to numeric(0) if another analysis than a two-class analysis is performed.

Assume, e.g., that we would like to perform a SAM analysis with the "usual" t-statistic assuming equal group variances and normality. The code of a function t.stat for such an analysis is given by

```
t.stat <- function(data, cl){</pre>
    require(genefilter) ||
         stop("genefilter required.")
    row.out <- rowttests(data, cl)</pre>
    d <- row.out$statistic</pre>
    m <- length(na.exclude(d))</pre>
    d.bar \leftarrow qt(((1:m) - 0.5)/m, row.out$df)
    p.value <- row.out$p.value</pre>
    vec.false <- m * p.value/2</pre>
    s <- row.out$dm/d
    # dm: differences in group means
    msg <- paste("SAM Two-Class Analysis",</pre>
          "Assuming Normality\n\n")
    list(d=-d, d.bar=d.bar, p.value=p.value,
         vec.false=vec.false, s=s, s0=0,
        mat.samp=matrix(numeric(0)),
        msg=msg, fold=numeric(0))
}
```

Please note that in the output of t.stat 'd' is set to -d since in rowttests the mean of group 2 is subtracted from the mean of group 1, whereas in sam the difference is taken the other way around.

Now t.stat can be used in sam by setting method=t.stat.

Example: ALL Data

As example we here employ one of the data sets used in Gentleman et al. (2005). The package ALL containing this data set can be downloaded by

```
> source(
    "http://www.bioconductor.org/getBioC.R")
> getBioC("ALL")
> library(ALL)
> data(ALL)
```

Even though it is in general not a good idea to filter genes / probe sets prior to a SAM analysis since SAM assumes that most of the genes are not differentially expressed, we here follow the code of Scholtens and von Heydebreck (2005) and filter the genes and select a subset of the samples.

```
> library(genefilter)
> subALL <- filterALL()
```

(The code of filterALL can be found in the Appendix.) This leads to an exprSet object containing gene expression data of 2,391 probe sets and 79 samples.

Following Scholtens and von Heydebreck (2005) we would like to identify the probe sets whose expression values differ strongly between the samples for which

```
> mol.biol <- pData(subALL)$mol.biol</pre>
```

is equal to "BCR/ABL" and the samples for which mol.biol=="NEG". Thus, sam is applied to this data set by specifying the required arguments 'data' and 'cl', where

'data' can either be a matrix, a data frame or an exprSet object containing the gene expression data.

'cl' is a vector containing the class labels of the samples. If 'data' is an exprSet object, then 'c1' can also be a character string naming the column of pData(data) that contains the class labels.

So

```
> library(siggenes)
> clALL <- ifelse(mol.biol=="BCR/ABL", 0, 1)</pre>
> dataALL <- exprs(subALL)</pre>
> out1 <- sam(dataALL, clALL,
   var.equal = TRUE, rand = 123456)
```

leads to the same results as

```
> out2 <- sam(subALL, "mol.biol",
  var.equal = TRUE, rand = 123456)
```

where 'var.equal' is set to TRUE since we here would like to assume that the group variances are equal, and 'rand' is set to 123456 to make the results of this analysis reproducible.

By default, the number of identified genes and the estimated FDR is computed for ten values of Δ equidistantly spaced between 0.1 and $\max_i |d_{(i)}|$ $d_{(i)}$. The output of our SAM analysis is thus given by

```
> out1
```

SAM Analysis for the Two-Class Unpaired Case Assuming Equal Variances

```
Delta p0
           False Called
 0.1 0.63 1900.61 2075 0.57726
```

2	0.7 0.63	125.07	400	0.19706
3	1.4 0.63	3.86	90	0.02703
4	2.0 0.63	0.1	25	0.00252
5	2.7 0.63	0	6	0
6	3.3 0.63	0	4	0
7	4.0 0.63	0	3	0
8	4.6 0.63	0	2	0
9	5.3 0.63	0	2	0
10	5.9 0.63	0	1	0

where p0 is the estimated prior probability that a gene is not differentially expressed, False is the number of falsely called genes (see Tusher et al., 2001), Called is the number of identified genes, and FDR = p0 * False / Called is the estimated FDR.

More information, e.g., the value of the fudge factor can be obtained using summary. Both summary and print can also be used to generate the above table for other values of Δ . For example,

```
> print(out1, seq(1.4, 2, 0.1))
```

SAM Analysis for the Two-Class Unpaired Case Assuming Equal Variances

	Delta	p0	${\tt False}$	${\tt Called}$	FDR
1	1.4	0.63	3.86	90	0.02703
2	1.5	0.63	2.36	77	0.01932
3	1.6	0.63	1.45	64	0.01428
4	1.7	0.63	0.80	54	0.00934
5	1.8	0.63	0.41	45	0.00574
6	1.9	0.63	0.26	36	0.00455
7	2.0	0.63	0.10	25	0.00252

Let's say our choice of Δ is 1.5. The SAM plot for this selection shown in Figure 1 is generated by

```
> plot(out1, 1.5, sig.col = c(3,2), pch = 16,
+ pos.stats = 2, cex = 0.6)
```

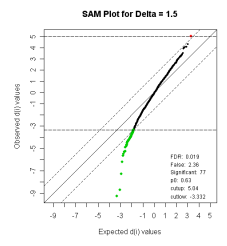


Figure 1: SAM Plot for $\Delta = 1.5$.

where

'sig.col' is a numeric value or vector specifying the color of the identified down- and up-regulated genes,

'pos.stats' indicates where the statistics are shown in the SAM plot,

'cex' specifies the relative size of the plotting symbols of the genes not identified as differentially expressed.

While the relative size of the symbols can be specified separately for the identified and the not identified genes, the symbol itself ('pch') is the same for both types of genes. For all arguments of the SAM specific method plot, see

> help.sam(plot)

Information about the identified genes such as their *d* values, the corresponding raw p-values and the q-values (see Storey and Tibshirani, 2003) can be obtained by

> summary(out1,1.5)

An excerpt from the output of summary is shown in Figure 2. This information can also be stored in a csv file via sam2excel or in an html file using sam2html. If 'data' is an exprSet object or 'chipname' is specified in sam2html, then the html file will also contain the gene symbols and links to public repositories such as Entrez, RefSeq and UniGene. If 'cdfname' is specified, links to the Affymetrix webpages of the identified probe sets will also be available. For example, the html file generated by

```
> sam2html(out1, 1.5, "out1.html", ll = TRUE,
+ cdfname = "HG-U95Av2")
```

is available at http://www.statistik.uni-dortmund.de/de/content/einrichtungen/lehrstuehle/personen/holgers/out1.html.

Finally, we would like to check if method="t.stat" (see previous section) really works.

```
> out3 <- sam(subALL, "mol.biol",
+ method = "t.stat")
> out3
```

SAM Two-Class Analysis Assuming Normality

	Delta	p0	False	${\tt Called}$	FDR
1	0.1	0.63	1832.197	2050	0.56307
2	0.7	0.63	94.609	347	0.17177
3	1.3	0.63	3.555	100	0.02240
4	1.9	0.63	0.044	19	0.00145
5	2.5	0.63	0.000567	6	5.95e-05
6	3.1	0.63	3.33e-05	4	5.24e-06
7	3.7	0.63	2.92e-07	3	6.14e-08
8	4.4	0.63	5.73e-10	2	1.80e-10
9	5.0	0.63	5.73e-10	2	1.80e-10
10	5.6	0.63	0	0	0

```
SAM Analysis for the Two-Class Unpaired Case Assuming Equal Variances
Number of permutations: 100
MEAN number of falsely called genes is computed.
Delta: 1.5
 cutlow: -3.332
cutup: 5.04
p0: 0.63
Significant Genes: 77
Falsely Called Genes: 2.36
FDR: 0.0193
Genes called significant (using Delta = 1.5):
   Row d.value stdev rawp q.value R.fold
                                              Name
   134
        -9.26 0.1188
                               0 0.457 1636_g_at
 1787
        -8.69 0.1327 0
                                0 0.442 39730_at
  133
        -7.28 0.1652 0
                                0 0.420
                                          1635_at
 1890
        -6.18 0.2878 0
                                0 0.429
                                           40202_at
        -5.65 0.2388
                                          37027_at
 1193
                                0 0.460
```

Figure 2: An excerpt from the output of summary (out1, 1.5).

Since in both analyses we have computed the t statistic assuming equal group variances, the d values in both analyses should be the same:

```
> tmp <- sum(round(out1@d, 8) ==
+ round(out3@d, 8))
> tmp == length(out1@d)
[1] TRUE
```

Summary

The package **siggenes** contains functions for performing both a Significance Analysis of Microarrays (SAM) and an Empirical Bayes Analysis of Microarrays (EBAM). The function sam provides not only a set of statistics for standard tests such as the t test and F test but also the possibility to use user-written functions for other testing situations. After identifying a list of genes, not only statistics of these genes such as their test scores and p-values can be obtained but also links to public repositories containing biological information about these genes.

The EBAM functions are currently under revision to provide more user-friendly and less memory-consuming versions of these functions having all the features that the SAM functions already have.

This article will be available as vignette in **siggenes 1.7.1** and later.

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Appendix

```
filterALL <- function(){
    pdat <- pData(ALL)</pre>
    subset<-intersect(grep("^B",</pre>
         as.character(pdat$BT)),
         which(pdat$mol %in% c("BCR/ABL",
         "NEG")))
    eset <- ALL[, subset]
    require(genefilter)
    f1 <- pOverA(0.25, log2(100))
    f2 \leftarrow function(x) IQR(x) > 0.5
    selected <- genefilter(eset,</pre>
        filterfun(f1, f2))
    esetSub <- eset[selected, ]</pre>
    pdat <- pData(esetSub)</pre>
    esetSub$mol.biol <-
         as.character(esetSub$mol.biol)
    esetSub
}
```

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Reverse Engineering Genetic Networks using the GeneNet Package

by Juliane Schäfer, Rainer Opgen-Rhein, and Korbinian Strimmer

GeneNet is a package for analyzing highdimensional (time series) data obtained from highthroughput functional genomics assays, such as expression microarrays or metabolic profiling. Specifically, GeneNet allows to infer large-scale gene association networks. These are graphical Gaussian models (GGMs) that represent multivariate dependencies in biomolecular networks by means of partial correlation. Therefore, the output of an analysis conducted by GeneNet is a graph where each gene corresponds to a node and the edges included in the graph portray direct dependencies between them.

GeneNet implements a specific learning algorithm that allows to estimate GGMs from small sample high-dimensional data that is both computationally as well as statistically efficient. This approach relies on analytic shrinkage estimation of covariance and (partial) correlation matrices and on model selection using (local) false discovery rate multiple testing. Hence, GeneNet includes a computational algorithm that decides which edges are to be included in the final network, in dependence of the *relative* values of the pairwise partial correlations.

In a recent comparative survey (Werhli et al., 2006) the **GeneNet** procedure was found to recover the topology of gene regulatory networks with similar accuracy as computationally much more demanding methods such as dynamical Bayesian networks (Friedman, 2004).

We note that the approach implemented in **GeneNet** should be regarded as an exploratory approach that may help to identify interesting genes (such as "hubs") or clusters of genes that are functionally related or co-regulated, rather than that it yields the precise network of mechanistic interactions. Therefore, the resulting network topologies need be interpreted and validated in the light of biological background information, ideally accompanied by further integrative analysis employing data from different levels of the cellular system.

Prerequisites

GeneNet is available from the CRAN repository and from the webpage http://strimmerlab.org/software/genenet/. It requires prior installation of four further R packages also found on CRAN: corpcor, longitudinal, fdrtool, and locfdr (Efron, 2004).