# Bioconductor's **edge** package Version 0.99.0

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# Contents

1	Introduction	2
2	Citing this package	2
3	Getting help	3
4	Quick start guide	3
5	Objects in edge           5.1 edgeSet            5.2 edgeFit	<b>4</b> 4
	Examples 6.1 Independent time course study	<b>5</b>

#### 1 Introduction

edge is a package for significance analysis of DNA micro-array experiments and is able to identify genes that are deferentially expressed between two or more different biological conditions (e.g., healthy versus diseased tissue). There are a number of existing software packages that perform significance analysis but edge can use the odp-statistic from the Optimal Discovery Procedure (ODP) for significance testing. Whereas previously existing methods employ statistics that are essentially designed for testing one gene at a time (e.g., t-statistics and F-statistics), the ODP uses information across all genes to test for differential expression.

The improvements in power are substantial; Figure 1 shows a comparison between edge and five leading software packages, based on a well-known breast cancer expression study (Hedenfalk et al. 2001). In addition to identifying deferentially expressed genes, edge includes implementations of popular packages such as snm, sva and qvalue.

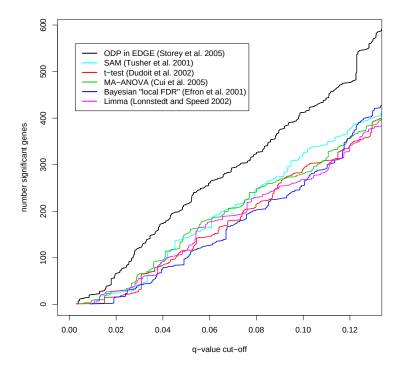


Figure 1: Comparison of EDGE to various other leading methods for identifying differential expressed genes in the Hedenfalk et al., 2001 study. Figure retrieved from Leek et al. (2005).

# 2 Citing this package

STOREY, J. D. The optimal discovery procedure for large-scale significance testing, with applications to comparative microarray experiments. *Journal of the Royal Statistical Society, Series B* 69 (2007), 347–368.

STOREY, J. D., DAI, J., AND LEEK, J. T. The optimal discovery procedure for large-scale significance

testing, with applications to comparative microarray experiments. Biostatistics 8 (2007), 414-432.

Woo, S., Leek, J. T., and Storey, J. D. A computationally efficient modular optimal discovery procedure. *Bioinformatics* 27 (2011), 509–515.

### 3 Getting help

Hopefully most questions relating to the package will be answered in the vignette but to get a more detailed account of how to use the functions simply type within R:

```
help(package = "edge")
```

Please contact the authors directly with any issues regarding bugs. Otherwise, any questions or problems implementing edge will most efficiently be addressed on the Bioconductor mailing list, http://stat.ethz.ch/mailman/listinfo/bioconductor.

# 4 Quick start guide

In order to get started using edge, it is important to first put all of the experimental data in an ExpressionSet object:

Once the ExpressionSet is created, the alternative and null hypothesis must be formulated based on the experiment. Given an ExpressionSet object and the alternative and the null hypothesis, the odp or 1rt function can be used as follows:

```
# Create Models
nullMod <- ~sex
altMod <- ~sex + ns(age, df = 3, intercept = FALSE)
# Create edgeSet object from ExpressionSet object
edgeObj <- edgeSet(expSet, full.model = fModel, null.model = nModel)
# Optimal Discovery Procedure
edgeODP <- odp(edgeObj)
# Likelihood Ratio Test
edgeLRT <- lrt(edgeObj)</pre>
```

In the above models, altMod is the alternative hypothesis and nullMod is the null hypothesis. The object edgeODP and edgeLRT contain a qvalue object which is the main slot of interest that can accessed by the qvalue.obj function. The following sections of the manual go through various case studies to show additional features of the edge package.

## 5 Objects in edge

#### 5.1 edgeSet

The main object in edge is the edgeSet object and it contains the following slots:

- full.model: the alternative model of the experiment
- null.model: the null model of the experiment
- full.matrix: the alternative model in matrix form
- null.matrix: the null model in matrix form
- individual: containing information on individuals in the experiment
- qvalue.obj: qvalue list
- ExpressionSet: inherits the slots from ExpressionSet object

As an example of how to access and set the slots of an edgeObj lets say we are interested in changing the alternative model to include the variable cov. The current models can be accessed by:

```
nullModel(object)
fullModel(object)
```

A new alternative model can be set by

```
fullModel(object) <- ~1 + cov</pre>
```

To get a summary of the object and all the slots use the summary function:

```
summary(object)
```

The package offers great flexibility in model choice and experimentation. See ?edgeSet for more details on how to extract and set each slot in the object.

#### 5.2 edgeFit

The function edgeFit fits a linear model to each gene and returns that information in an edgeFit object that contains the following slots:

- fit.full: fitted values for alternative model
- fit.null: fitted values for null model
- res.full: residuals for the alternative model
- res.null: residuals for the null models
- dH.full: diagonal elements in the projection matrix for the full model

- beta.coef: the coefficients for the full model
- stat.type: statistic type used, either "odp" or "lrt"

To access the coefficients in the edgeFit object:

```
betaCoef(object)
```

Similarly, to access the full and null residuals:

```
resFull(object)
resNull(object)
```

See ?edgeFit for more details on how to extract and set each slot of the object.

### 6 Examples

Three examples will be used to show the functionality of edge. In each example there will be a static, longitudinal and independent case study. It will become evident that in each case, the analysis procedure is similar and the only step that differs is the model setup.

There are three main steps when using edge:

- Write the alternative and null models of the experiment. In this manual they will be called altMod and nullMod.
- Create an ExpressionSet object and input the object along with the models in the edgeSet function.
- Use functions odp or lrt to obtain the q-value object which is the slot of interest. The edgeFit function can be used to extract information regarding the model fits.

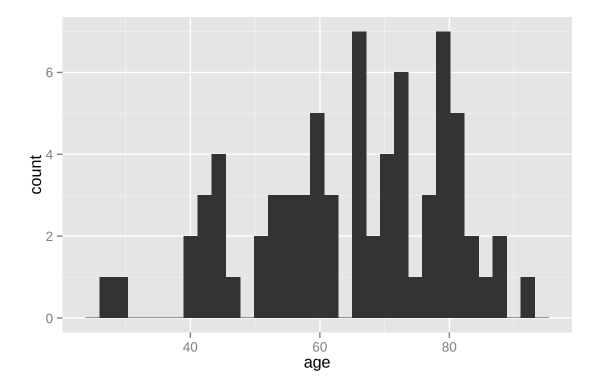
#### 6.1 Independent time course study

Gene expression measurements from kidney samples were obtained from 72 human subjects ranging in age from 27 to 92 years. Only one array was obtained per sample and the age and tissue type of each subject was recorded. To import the data:

```
data(kidney)
names(kidney)

## [1] "age" "tissue" "sex" "kidexpr"
## [5] "kidcov"
```

There are a few covariates in this data set: sex, age, tissue, kidexpr and kidcov. The two main covariates of interest for this example are the sex and age covariates. The sex variable is whether the subject was male or female and the age variable is the age of the patients. Lets view a histogram of the age to get a better idea of data:



From the histogram, it is important to keep in mind that a majority of the patients appear to be older than 50.

In the kidney experiment they were interested in finding the effect of age on gene expression. In this case, we handle the time variable, age, by fitting a natural spline curve [3].

Step 1: Create the alternative and null models:

```
library(splines)
nullMod <- ~-1 + sex
altMod <- ~-1 + sex + ns(age, intercept = FALSE,
    df = 4)</pre>
```

Where nullMod is the null model and altMod is the alternative model. The sex covariate is an adjustment variable while age is the biological variable of interest. It is important to note that it is necessary to include the adjustment variables in the formulation of the alternative models as done above.

Step 2: Once the alternative and null models have been decided, create an ExpressionSet object:

expSet contains the expression measurements and the covariates of the experiment. To access the expression values, one can use the function exprs(expSet) or to access the covariates, pData(expSet). The ExpressionSet class is a widely used object in Bioconductor and more information can be found http://www.bioconductor.org/packages/2.14/bioc/html/Biobase.html.

The function edgeSet can be used to create an edgeSet object from an ExpressionSet object:

```
edgeObj <- edgeSet(expSet, full.model = altMod, null.model = nullMod)
slotNames(edgeObj)
   [1] "null.model"
                            "full.model"
##
##
   [3] "null.matrix"
                            "full.matrix"
                            "qvalue.obj"
##
   [5] "individual"
   [7] "experimentData"
##
                            "assayData"
                            "featureData"
   [9] "phenoData"
## [11] "annotation"
                            "protocolData"
## [13] ".__classVersion__"
```

The key slot in the object is the qvalue.obj which should be the only empty slot. The other slots are directly related to the ExpressionSet object, full.model and null.model. See the section on edgeSet object for more details on these slots.

The summary function summarizes the slots in the edgeSet object:

```
summary(edgeObj)
##
## ExpressionSet Summary
##
## ExpressionSet (storageMode: lockedEnvironment)
## assayData: 1500 features, 72 samples
    element names: exprs
## protocolData: none
## phenoData
     sampleNames: 1 2 ... 72 (72 total)
##
##
     varLabels: age sex
##
    varMetadata: labelDescription
## featureData: none
## experimentData: use 'experimentData(object)'
## Annotation:
##
## edge Analysis Summary
##
## Total number of arrays: 72
## Total number of probes: 1500
##
## Biological variables:
   Null Model: ~-1 + sex
##
   Full Model: ~-1 + sex + ns(age, intercept = FALSE, df = 4)
##
##
##
  . . . . . . .
##
```

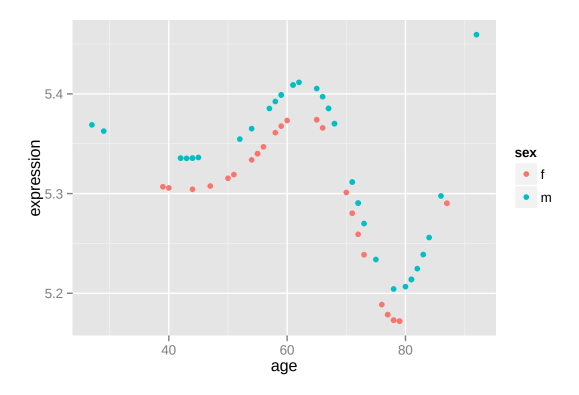
**Step 3:** Before running any significance analysis, lets view our model fits on the data. The edgeFit function can be used to extract residuals and fitted values from the alternative and null models:

```
efObj <- edgeFit(edgeObj, stat.type = "lrt")</pre>
```

The stat.type argument specifies whether you want the odp or lrt fitted values. To access the alternative model fitted values:

```
fitVals <- fitFull(ef0bj)</pre>
```

The fitted values of the first gene are shown below:



We can see that with our alternative model chosen, the expression goes up and down as the kidney ages for this particular gene. Once other interesting genes have been inspected, the user can either use the function odp or lrt to get the qvalue object. The lrt function performs a likelihood ratio test to determine p-values. If the null distribution, nullDistn, is calculated using "bootstrap" then residuals from the alternative model are re-sampled and added to the null model to simulate a distribution where there is no differential expression. Otherwise, the default input is "normal" and the assumption is that the data set follows an F-distribution.

To use 1rt on the edgeSet object:

```
edgeLRT <- lrt(edgeObj, nullDistn = "normal")</pre>
```

To view a summary of the object:

```
summary(edgeLRT)
##
## ExpressionSet Summary
```

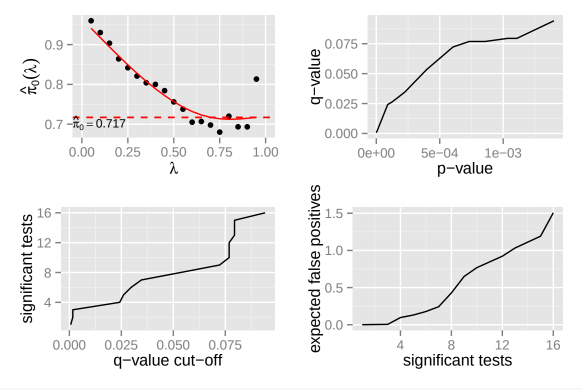
```
## ExpressionSet (storageMode: lockedEnvironment)
## assayData: 1500 features, 72 samples
## element names: exprs
## protocolData: none
## phenoData
##
   sampleNames: 1 2 ... 72 (72 total)
##
   varLabels: age sex
##
   varMetadata: labelDescription
## featureData: none
## experimentData: use 'experimentData(object)'
## Annotation:
##
## edge Analysis Summary
##
## Total number of arrays: 72
## Total number of probes: 1500
## Biological variables:
## Null Model:~-1 + sex
##
## Full Model: ~-1 + sex + ns(age, intercept = FALSE, df = 4)
##
## .....
##
##
## Statistical significance summary:
## pi0: 0.7172
##
## Cumulative number of significant calls:
##
##
           <1e-04 <0.001 <0.01 <0.025 <0.05 <0.1
## p-value 4 12 39 72 132 244
## q-value 0 1 3 4 7 16 ## local fdr 0 0 3 3 3 7
##
             <1
## p-value 1500
## q-value
           1500
## local fdr 1253
```

The slot of interest for significance analysis is the qvalue.obj slot. To access the slot:

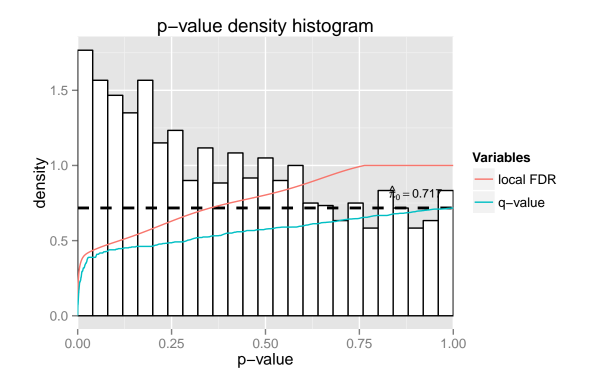
```
qval <- qvalue.obj(edgeLRT)</pre>
```

To visualize the results, plot or hist functions can be used on the qval:

```
plot(qval)
```







The odp function uses information across all tests when formulating the test statistic. In order to improve the speed of the algorithm, we utilize a k-means clustering algorithm where genes are assigned to a cluster based on the Kullback-Leiber distance. Each gene is assigned a module-average parameter to calculate the odp-statistic. The number of clusters can be adjusted by n.mods. Type ?odp for more details on the algorithm.

To use odp on an edgeSet object:

```
edgeODP <- odp(edgeObj, bs.its = 10, verbose = FALSE,
    n.mods = 20)</pre>
```

The argument bs.its controls the number of bootstrap iterations, verbose prints the iteration step and n.mods is the number of clusters formed.

To summarize the object using the odp method:

```
summary(edgeODP)
##
## ExpressionSet Summary
##
## ExpressionSet (storageMode: lockedEnvironment)
## assayData: 1500 features, 72 samples
   element names: exprs
## protocolData: none
## phenoData
    sampleNames: 1 2 ... 72 (72 total)
##
##
    varLabels: age sex
    varMetadata: labelDescription
## featureData: none
## experimentData: use 'experimentData(object)'
## Annotation:
##
## edge Analysis Summary
##
## Total number of arrays: 72
## Total number of probes: 1500
##
## Biological variables:
  Null Model: ~-1 + sex
##
   Full Model: ~-1 + sex + ns(age, intercept = FALSE, df = 4)
##
##
## .....
##
##
## Statistical significance summary:
## pi0: 0.575
##
## Cumulative number of significant calls:
##
##
             <1e-04 <0.001 <0.01 <0.025 <0.05 <0.1
                              65 138
## p-value
                  9
                      18
                                          223 354
```

```
## q-value 0 0 9 14 21 33

## local fdr 0 0 0 10 14 21

## <1

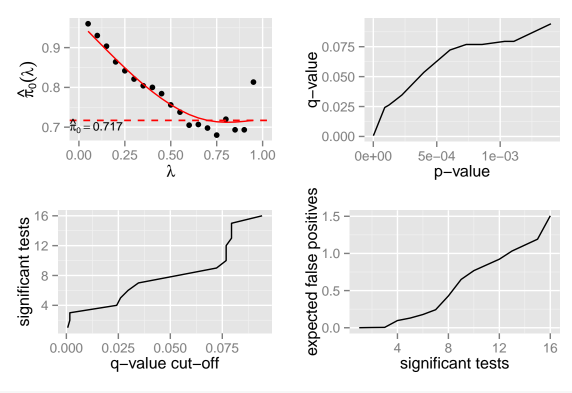
## p-value 1500

## q-value 1500

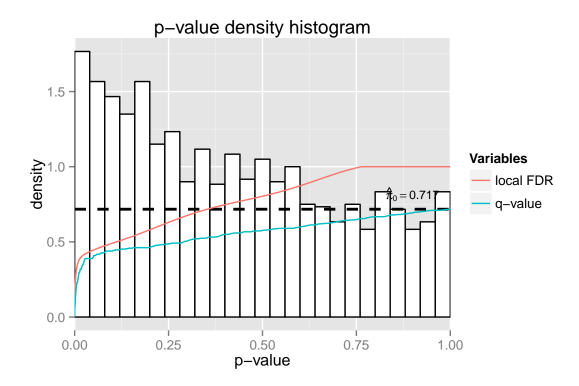
## local fdr 1500
```

Using plot and hist functions on the qvalue object:

```
qval <- qvalue.obj(edgeLRT)
plot(qval)</pre>
```



hist(qval)



We notice that the optimal discovery method finds more significant genes. As shown in previous research by Storey [1], the ODP finds more significant genes for a fixed FDR when compared to the likelihood ratio test and other popular statistical methods.

# Acknowledgements

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#### References

- [1] Storey, J. D. The optimal discovery procedure for large-scale significance testing, with applications to comparative microarray experiments. *Journal of the Royal Statistical Society, Series B* 69 (2007), 347–368.
- [2] Storey, J. D., Dai, J., and Leek, J. T. The optimal discovery procedure for large-scale significance testing, with applications to comparative microarray experiments. *Biostatistics* 8 (2007), 414–432.
- [3] Storey, J. D., Xiao, W., Leek, J. T., Tompkins, R. G., and Davis, R. W. Significance analysis of time course microarray experiments. *Proceedings of the National Academy of Sciences of the United States of America (PNAS)* 102 (2005), 12837–12842.

[4] Woo, S., Leek, J. T., and Storey, J. D. A computationally efficient modular optimal discovery procedure. Bioinformatics 27 (2011), 509-515.