Bioconductor's **edge** package Version 0.99.0

John D. Storey and Andrew J. Bass Princeton University http://genomine.org/contact.html

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1 Introduction

edge is a package for significance analysis of DNA micro-array experiments and is able to identify genes that are differentially expressed between two or more different biological conditions (e.g., healthy versus diseased tissue). edge performs significance analysis and uses 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.

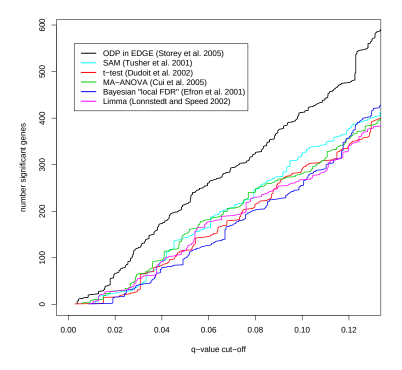


Figure 1: Comparison of EDGE to various other leading methods for identifying differential expressed genes in the Hedenfalk et al., 2001 study. Figure retrieved from Leek et al. (2005).

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

To get started let's first load the kidney dataset that is included in the package:

```
library(edge)
data(kidney)
kidexpr <- kidney$kidexpr
age <- kidney$age
sex <- kidney$sex</pre>
```

The kidney study was interested in finding out what happens as the kidney ages. The covariate age is the age of the subject and the covariate sex is whether the subject was male or female. The expression values for the genes are contained in the kidexpr variable which is a 1500 by 72 matrix. The kidney dataset included in the package is a subset of the full dataset.

Once the data has been loaded, the user can proceed to use the function edgeModel to create the alternative and null hypothesis of the experiment:

```
edgeObj <- edgeModel(data = kidexpr, adj.var = data.frame(sex),
   tme = age, sampling = "timecourse", basis.type = "ncs",
   basis.df = 4)</pre>
```

Since the study examines the kidney tissue over time, the sampling method is "timecourse" where the tme argument is age. The sex variable is an adjustment variable and basis.type and basis.df describe the spline fit on the tme argument.

The alternative and null hypothesis formulated by edgeModel can be accessed by fullModel and nullModel, respectively:

```
fullModel(edgeObj)
## ~sex + ns(tme, df = 4, intercept = FALSE)
## <environment: 0x898e370>
```

```
nullModel(edgeObj)
## ~sex
## <environment: 0x898e370>
```

adj.var is the adjustment variable sex and time.basis is the spline fit of age created by edgeModel. The alternative and null models created by edgeModel are fitted to the data by least squares and test statistics are formed by using either odp or lrt:

```
# Optimal Discovery Procedure
edgeODP <- odp(edgeObj, verbose = FALSE)
# Likelihood Ratio Test
edgeLRT <- lrt(edgeObj)</pre>
```

The summary function can be used to summarize the objects:

```
summary(edgeODP)
```

The objects edgeODP and edgeLRT contain a qvalue object which provides p-values, q-values and local false discovery rate values:

```
qval <- qvalueObj(edgeODP)</pre>
summary(qval)
##
## Call:
## qvalue(p = pval)
##
## pi0: 0.5017
##
## Cumulative number of significant calls:
##
##
            <1e-04 <0.001 <0.01 <0.025 <0.05 <0.1
## p-value
              9
                       24
                             86 163 254 399
## q-value
                 0
                       0
                             10
                                   21
                                         34 106
                        0
                            5
                                    13
                                          21
## local FDR
                 0
                                             65
##
              <1
## p-value
            1500
## q-value
           1500
## local FDR 1500
```

The following sections of the manual go through various case studies to show additional features of the edge package.

5 Examples

Three examples will be used to show the functionality of edge. The examples will cover static, longitudinal and independent case studies. 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:

- Load experimental data. Optionally, create an ExpressionSet object.
- Use edgeModel to create an edgeSet object. If an ExpressionSet object is created use 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.

5.1 Static study

Step 1: The gibson dataset provides gene expression measurements in peripheral blood leukocyte samples from three Moroccan Amazigh groups leading distinct ways of life: desert nomadic (DESERT), mountain agrarian (VILLAGE), and coastal urban (AGADIR). Suppose we are interested in finding the genes that differentiate the Moroccan Amazigh groups the most.

To import the data:

```
data(gibson)
names(gibson)
## [1] "batch" "gibexpr" "gender" "location"
```

There are a few variables in this data set: batch, gibexpr, gender, and location. The three covariates of interest are gender, batch and location. There are three locations where individuals were sampled (location): "VILLAGE", "DESERT" and "AGADIR". At each location there were either "males" or "females" (gender) and there were different batches. The gibexpr variable contains the expression matrix of the experiment.

Step 2: Use the function edgeModel to create an edgeSet object:

The gibson study is a static experiment so the sampling argument will be "static". The grp argument is for the location variable and the adj.var argument is the adjustment variables. The adj.var must be in model.matrix form. A brief overview of the arguments of edgeModel:

- data Matrix of expression values
- tme A vector of time measurements for a time-course study
- ind A factor that assigns each observations to an individual in the experiment
- basis.df Degree of freedom of spline fit in a time-course study
- basis.type A spline curve is fitted to the tme variable in a "timecourse" study. The type can be "ncs" (B-spline for a natural cubic spline) or "ps" (polynomial spline)
- bio.var Biological variables
- adj.var Adjustment variables (matrix)

- grp Numerical vector describing which group each observation belong (i.e "DESERT", "VILLAGE" or "AGADIR")
- sampling Can either be "timecourse" or "static" depending on the experiment

Other examples in the vignette will show when to use the additional arguments in edgeModel. edgeSet is the main object in the package:

```
slotNames(edgeObj)
                           "full.model"
   [1] "null.model"
##
   [3] "null.matrix"
                           "full.matrix"
  [5] "individual"
                          "qvalueObj"
  [7] "experimentData" "assayData"
##
                         "featureData"
   [9] "phenoData"
##
                           "protocolData"
## [11] "annotation"
## [13] ".__classVersion__"
```

The edgeObj is an edgeSet object that extends an ExpressionSet object. It contains the covariates and expression values along with the alternative and null models. The alternative and null models generated by edgeModel can be accessed using

```
fullModel(edgeObj)

## ~gibson.gender + gibson.batch + gibson.location
## <environment: 0x87e6278>

nullModel(edgeObj)

## ~gibson.gender + gibson.batch
## <environment: 0x87e6278>
```

The key slot in edgeObj is the qvalueObj which should be the only empty slot. The other slots are directly related to the input data and hypothesis models. See the section 6.1 for more details on the edgeSet object.

The summary function summarizes the slots in the edgeSet object:

```
summary(edgeObj)
##
## ExpressionSet Summary
##
## ExpressionSet (storageMode: lockedEnvironment)
## assayData: 2000 features, 46 samples
## element names: exprs
## protocolData: none
## phenoData
    sampleNames: 1 2 ... 46 (46 total)
##
    varLabels: gibson.gender gibson.batch
##
##
       gibson.location
   varMetadata: labelDescription
## featureData: none
## experimentData: use 'experimentData(object)'
```

```
## Annotation:
##
## edge Analysis Summary
##
## Total number of arrays: 46
## Total number of probes: 2000
##
## Biological variables:
## Null Model: "gibson.gender + gibson.batch
## <environment: 0x87e6278>
##
  Full Model: "gibson.gender + gibson.batch + gibson.location
## <environment: 0x87e6278>
##
##
##
```

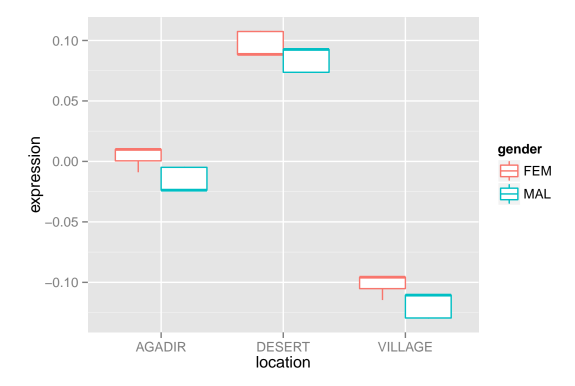
Step 3: Before running any significance analysis, lets view our model fits of the data. The edgeFit function can be used to extract residuals and fitted values from the alternative and null models:

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

edgeFit is simply an implementation of least squares using the alternative and null models. The stat.type argument specifies whether you want the odp or lrt fitted values. The difference between choosing "odp" and "lrt" is that "odp" centers the data by the null model fit. To access the alternative model fitted values:

```
fitVals <- fitFull(efObj)
```

The fitted values of the first gene are shown below:



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.

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 or lrt on the edgeSet object:

The argument bs.its controls the number of bootstrap iterations, verbose prints the iteration step and n.mods is the number of clusters formed. If n.mods is equal to the number of genes than the full Optimal Discovery Procedures is used.

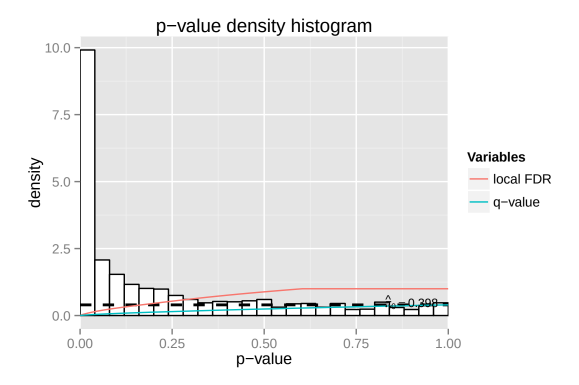
The slot of interest for significance analysis is the qvalueObj slot. To access the slot:

```
qval <- qvalueObj(edgeODP)
summary(qval)
##</pre>
```

```
## Call:
## qvalue(p = pval)
## pi0: 0.3981
##
## Cumulative number of significant calls:
##
             <1e-04 <0.001 <0.01 <0.025 <0.05 <0.1
##
## p-value
                102
                        240
                              531
                                     702
                                            841 1020
                  0
                        163
                              450
                                            854 1141
## q-value
                                     661
                   0
                        102
## local FDR
                              256
                                     405
                                            531 695
                <1
## p-value
             2000
## q-value
             2000
## local FDR 1716
```

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

```
hist(qval)
```



5.2 Independent time course study

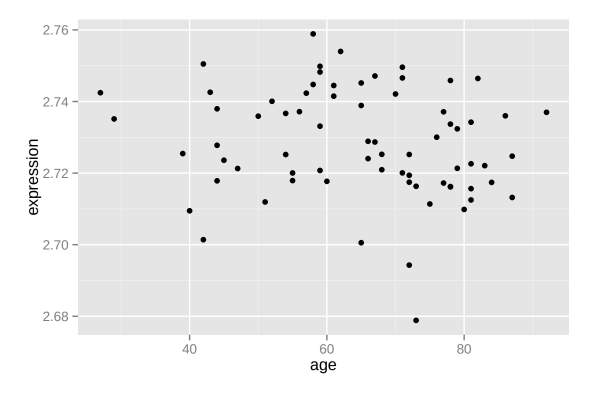
Step 1: Gene expression measurements from kidney samples were obtained from 72 human subjects ranging in age from 27 to 92 years. Only one array was obtained per sample and the age and tissue type of each subject was recorded. To import the data:

```
data(kidney)
names(kidney)

## [1] "age" "sex" "kidexpr"

kidexpr <- kidney$kidexpr
age <- kidney$age
sex <- kidney$sex</pre>
```

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 the first gene to get a better idea of data:



For this particular gene, it seems that there is a slight decrease in expression past 60 and an increase past 80.

Step 2: Use the function edgeModel to create an edgeSet object:

```
edgeObj <- edgeModel(data = kidexpr, adj.var = data.frame(sex),
   tme = age, sampling = "timecourse", basis.type = "ncs",
   basis.df = 4)</pre>
```

Since the kidney study is a time-course study the sampling method will be "timecourse". The adjustment variable in the study is sex while the time variable is age. A brief overview of the arguments can be found in section 5.1.

The alternative and null models can be accessed using

```
fullModel(edgeObj)

## ~sex + ns(tme, df = 4, intercept = FALSE)

## <environment: 0x9f8a590>

nullModel(edgeObj)

## ~sex

## <environment: 0x9f8a590>
```

The adj.var corresponds to the adjustment variables and time.basis corresponds to the time variable inputed in edgeModel. See section 6.1 for more details.

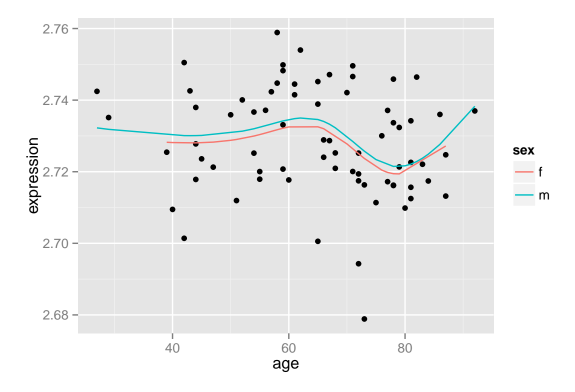
Step 3: Before running any significance analysis, lets view our model fits of the data. The edgeFit function can be used to extract residuals and fitted values from the alternative and null models. edgeFit performs a least squares regression on the alternative and null models. In the endotoxin dataset since there is a ind factor, the data is centered by individual in order to remove the effects from various individuals:

```
ef0bj <- edgeFit(edgeObj, stat.type = "lrt")</pre>
```

The stat.type argument specifies whether you want the odp or lrt fitted values. As mentioned in section 5.1, the only difference between the fitted values of "odp" and "lrt" is that the "odp" method centers the data by the null model fit. To access the alternative model fitted values:

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

The fitted values for the gene shown in the first step is shown below:



Our initial intuition was correct: A slight decrease in expression follow by an increase as time goes on. As mentioned in section 5.1, the lrt or odp function can be used for differential analysis:

To use odp or lrt on the edgeSet object:

Next, lets see if the gene mentioned above is significant:

```
qval <- qvalueObj(edgeODP)
qval$pvalues[1]

## [1] 0.05645

qval$qvalue[1]

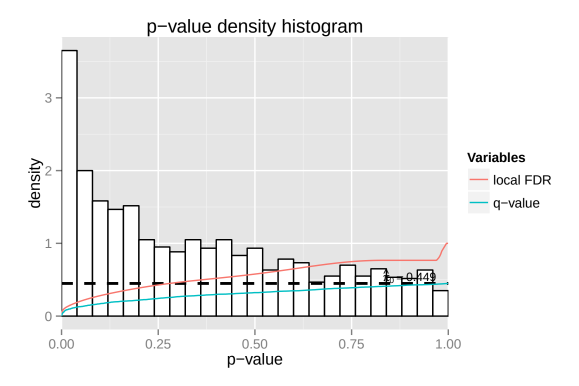
## [1] 0.1359

qval$lfdr[1]

## [1] 0.2181</pre>
```

If we assume a p-value threshold of 0.5 then the gene does not show any significant changes over time. The high q-value and local FDR value confirm that the gene is not significant.

Using the hist function on the qvalue object:



5.3 Longitudinal time course study

Step 1: The endotoxin dataset provide gene expression measurements in an endotoxin study where four subjects were given endotoxin and four subjects were given a placebo. Blood samples were collected and leukocytes were isolated from the samples before infusion and measurement were recorded at times 2, 4, 6, 9, 24 hours. We are interested in identifying genes that vary over time between the endotoxin and control groups.

To import the data:

```
data(endotoxin)
names(endotoxin)
## [1] "class" "endoexpr" "ind" "time"
```

There are a few covariates in this data set: expr, class, individual, and time. There are 8 individuals in the experiment (ind) that were sampled at multiple time points (time) that were either "endotoxin" or "control" (class). The expr variable contains the expression values of the experiment.

Step 2: Use the function edgeModel to create an edgeSet object:

```
edgeObj <- edgeModel(data = endotoxin$endoexpr, ind = endotoxin$ind,
    tme = endotoxin$time, grp = data.frame(endotoxin$class),</pre>
```

```
sampling = "timecourse", basis.type = "ncs",
basis.df = 4)
```

The endotoxin experiment is a time-course study so the sampling argument will be "timecourse". The tme argument is for the time variable in the experiment and the ind argument is to identify which observations corresponds what individuals. Since the sampling method is "timecourse", we fit a spline curve described by variables basis.type and basis.df. Additional arguments can be viewed by typing ?edgeModel.

The alternative and null models can be accessed using

The adj.var corresponds to the adjustment variables, the time.basis corresponds to the time variable and the grp corresponds to the treatment variable inputed in edgeModel. See the section 5.1 object for more details on accessing and setting edgeSet slots.

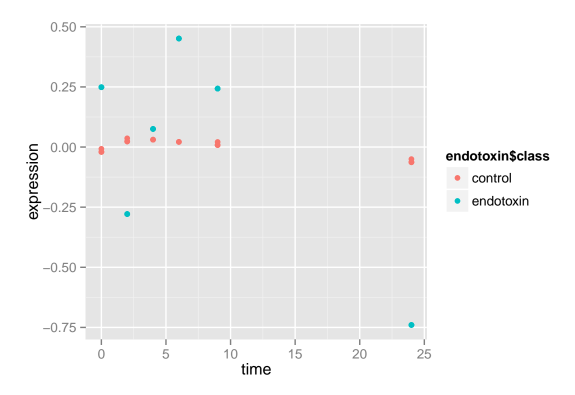
Step 3: Before running any significance analysis, lets view our model fits of the data. The edgeFit function can be used to extract residuals and fitted values from the alternative and null models:

```
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:



Next, the user can either use the function odp or lrt to get the qvalue object. See section 5.1 for more details on the odp and lrt method. To use odp or lrt:

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

```
qval <- qvalueObj(edgeODP)</pre>
```

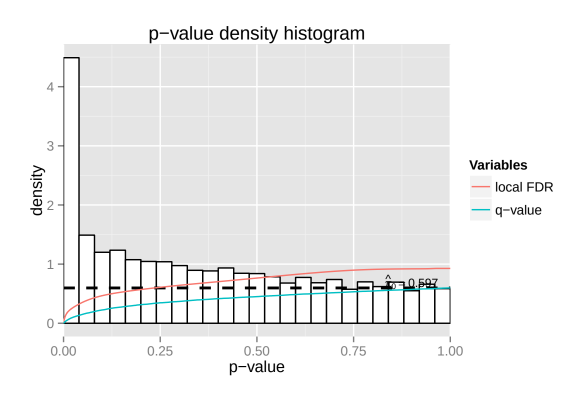
To summarize the object using the odp method:

```
summary(qval)
##
## Call:
## qvalue(p = pval)
## pi0: 0.597
##
## Cumulative number of significant calls:
##
##
              <1e-04 <0.001 <0.01 <0.025 <0.05 <0.1
## p-value
                191
                        324
                              551
                                      731
                                            978 1327
                              339
## q-value
                  0
                        189
                                      424
                                            533
                                                718
                   0
                              203
## local FDR
                        167
                                      288
                                            347
                                                419
##
                <1
```

```
## p-value 5000
## q-value 5000
## local FDR 5000
```

Using the hist function on the qvalue object:

hist(qval)



6 Objects in edge

6.1 edgeSet

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

- ullet 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
- qvalueObj: qvalue list

• ExpressionSet: inherits the slots from ExpressionSet object

ExpressionSet contains the expression measurements and the covariates of the experiment. To access the expression values, one can use the function exprs or to access the covariates, pData. 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.

As an example of how to access the slots of an edgeObj lets say we are interested in viewing the alternative model. The model can be accessed by:

```
fullModel(edgeObj)

## ~endotoxin.class + ns(tme, df = 4, intercept = FALSE) + (endotoxin.class):ns(tme,
## df = 4, intercept = FALSE)
## <environment: 0x835c5c8>
```

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

```
summary(edgeObj)
##
## ExpressionSet Summary
##
## ExpressionSet (storageMode: lockedEnvironment)
## assayData: 5000 features, 46 samples
   element names: exprs
## protocolData: none
## phenoData
    sampleNames: 1 2 ... 46 (46 total)
##
   varLabels: endotoxin.class tme
##
   varMetadata: labelDescription
## featureData: none
## experimentData: use 'experimentData(object)'
## Annotation:
## edge Analysis Summary
##
## Total number of arrays: 46
## Total number of probes: 5000
##
## Biological variables:
## Null Model: endotoxin.class + ns(tme, df = 4, intercept = FALSE)
## <environment: 0x835c5c8>
##
## Full Model: endotoxin.class + ns(tme, df = 4, intercept = FALSE) + (endotoxin.class):ns(tme,
      df = 4, intercept = FALSE)
## <environment: 0x835c5c8>
## Individuals:
## [1] 1 1 1 1 1 1 2 2 2 2 2 2 3 3 3 3 3 3 4 4 4 4
## [23] 4 4 5 5 5 5 5 5 6 6 6 6 7 7 7 7 7 7 8 8 8 8
## [45] 8 8
## Levels: 1 2 3 4 5 6 7 8
```

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.

6.2 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 from the alternative model
- fit.null: fitted values from null model
- res.full: residuals from the alternative model
- res.null: residuals from the null model
- 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 of the edgeFit object in section 5.2:

```
betaCoef(ef0bj)
```

Similarly, to access the full and null residuals:

```
resFull(ef0bj)
resNull(ef0bj)
```

A summary of the object can be displayed by:

```
summary(ef0bj)
```

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

7 Using the sva package

The sva package is useful for removing batch effects or any unwanted variation in an experiment. It does this by forming surrogate variables to adjust for sources of unknown variation. edge uses the sva package in the function edgeSVA. An example of how to use this on the kidney dataset:

```
newEdgeObj <- edgeSVA(edgeObj, n.sv = 5, B = 10)
## Number of significant surrogate variables is: 5</pre>
```

```
## Iteration (out of 10 ):1 2 3 4 5 6 7 8 9 10
```

A new edgeObj is created that includes the surrogate variables in the null and full matrices from sva. See ?sva for additional input parameters in edgeSVA.

Now odp or lrt can simply be used as before:

```
edgeODP <- odp(newEdgeObj, verbose = FALSE)
edgeLRT <- lrt(newEdgeObj)</pre>
```

Acknowledgements

This software development has been supported in part by funding from the National Institutes of Health and the Office of Naval Research.

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