

# Package ‘EDDA’

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**Type** Package

**Title** Experimental Design in Differential Abundance analysis

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**Author** Li Juntao, Luo Huaien, Chia Kuan Hui Burton, Niranjana Nagarajan

**Maintainer** Chia Kuan Hui Burton <chiakhb@gis.a-star.edu.sg>, Niranjana Nagarajan <nagarajann@gis.a-star.edu.sg>

**Description** EDDA can aid in the design of a range of common experiments such as RNA-seq, Nanos-tring assays, RIP-seq and Metagenomic sequencing, and enables researchers to comprehensively investigate the impact of experimental decisions on the ability to detect differential abundance. This work was published on 3 December 2014 at Genome Biology under the title “The importance of study design for detecting differentially abundant features in high-throughput experiments” (<http://genomebiology.com/2014/15/12/527>).

**License** GPL (>= 2)

**Depends** Rcpp (>= 0.10.4),parallel,methods,ROCR,DESeq,baySeq,snow,edgeR

**Imports** graphics, stats, utils, parallel, methods, ROCR, DESeq, baySeq, snow, edgeR

**LinkingTo** Rcpp

**biocViews** Sequencing, ExperimentalDesign, Normalization, RNASEq, ChIPSeq

**URL** <http://edda.gis.a-star.edu.sg/>,  
<http://genomebiology.com/2014/15/12/527>

**NeedsCompilation** yes

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EDDA-package

*Experimental Design in Differential Abundance analysis***Description**

EDDA aids in the design of a range of common experiments including RNA-seq, Nanostring assays, RIP-seq and Metagenomic sequencing, and enables researchers to comprehensively investigate the impact of experimental decisions on the ability to detect differential abundance.

**Details**

Package: EDDA  
 Type: Package  
 Version: 0.99.2  
 Date: 2014-02-12  
 License: GPL (>= 2)

generateData() testDATs() computeAUC() plotROC() plotPRC()

**Author(s)**

Li Juntao, Luo Huaien, Chia Kuan Hui Burton, Niranjan Nagarajan

Maintainer: Li Juntao<lij9@gis.a-star.edu.sg>, Luo Huaien<luoh2@gis.a-star.edu.sg>, Niranjan Nagarajan <nagarajann@gis.a-star.edu.sg>

**References**

Luo Huaien, Li Juntao, Chia Kuan Hui Burton, Shyam Prabhakar, Paul Robson, Niranjan Nagarajan, The importance of study design for detecting differentially abundant features in high-throughput experiments, under review.

**Examples**

```
data <- generateData(EntityCount=500)
test.obj <- testDATs(data,DE.methods=c("DESeq","edgeR"),nor.methods="default")
auc.obj <- computeAUC(test.obj)
plotROC(auc.obj)
plotPRC(auc.obj)
```

---

BP *BaySeq Profile used in simulations by Hardcastle et al*

---

### Description

RNA-seq profile datasets.

### Usage

`data(BP)`

### Format

Data frames with 8647 observations on the following 2 variables.

`gene` a character vector

`expression` a numeric vector

### References

Hardcastle, T.J. & Kelly, K.a. baySeq: empirical Bayesian methods for identifying differential expression in sequence count data. BMC bioinformatics 11, 422-422.

---

`computeAUC` *compute AUC values.*

---

### Description

compute AUC values for each test.

### Usage

```
computeAUC(obj, cutoff=1, numCores=10,
DE.methods=c("Cuffdiff", "DESeq", "baySeq", "edgeR", "MetaStats", "NOISEq"),
nor.methods=c("default", "Mode", "UQN", "NDE"))
```

### Arguments

<code>obj</code>	Object from <code>testDATs()</code> .
<code>cutoff</code>	cutoff for ROC curve. Default is 1.
<code>numCores</code>	Number of cores for parallelization. Default is 10.
<code>DE.methods</code>	Method list for differential abundance tests. Methods currently available include "Cuffdiff", "DESeq", "baySeq", "edgeR", "MetaStats", "NOISEq".
<code>nor.methods</code>	Normalization method list. Methods currently available include "default"(default normalization for each DE method), "Mode"(Mode normalization), "UQN"(Upper quartile normalization), "NDE"(non-differential expression normalization).

**Author(s)**

Li Juntao, and Luo Huaien

**References**

Luo Huaien, Li Juntao, Chia Kuan Hui Burton, Shyam Prabhakar, Paul Robson, Niranjana Nagarajan, The importance of study design for detecting differentially abundant features in high-throughput experiments, under review.

**Examples**

```
data <- generateData(EntityCount=200)
test.obj <- testDATs(data,DE.methods="DESeq",nor.methods="default")
auc.obj <- computeAUC(test.obj)
```

---

generateData

*generate count data*

---

**Description**

Simulate count data using different models and settings.

**Usage**

```
generateData(SimulModel="Full", SampleVar="medium",
ControlRep=5, CaseRep=ControlRep, EntityCount=1000, FC="Norm(2,1)",
perDiffAbund=0.1, upPDA=perDiffAbund/2, downPDA=perDiffAbund/2,
numDataPoints=100, AbundProfile = "HBR", modelFile = NULL, minAbund=10, varLibsizes=0.1,
outlier=FALSE, perOutlier=0.15, factorOutlier=100,
inputCount=NULL, inputLabel=NULL, SimulType="auto")
```

**Arguments**

SimulModel	Simulation model used. Default is "Full". SimulModel="NegBinomial" is negative binomial model which generates data using negative binomial distribution. SimulModel="Multinom" is multinomial model which generates data mimicking the multinomial sampling process. SimulModel="Full" is a model which combines "NegBinomial" and "Multinom". SimulModel="ModelFree" uses model free approach to generate data by sub-sampling counts from modelFile (if modelFile != NULL) or from input File (if modelFile == NULL)
SampleVar	Sample variation: Default is "medium". It could be "low", "medium" and "high" or a real number.
ControlRep	Number of replicates for control group. Default is 5.
CaseRep	Number of replicates for case group. Default is same as ControlRep.
EntityCount	Entity count. Default is 1000.

FC	Fold change type. It can be "Norm(mu,sigma)", "logNorm(mu,sigma)", "log2Norm(mu,sigma)" or "Unif(a,b)". mu,sigma and a,b need be predefined. Default is "Norm(2,1)".
perDiffAbund	Percentage of differential abundance. Default is 0.1 (i.e "10 percent").
upPDA	Percentage of up-regulated differential abundance. Default is perDiffAbund/2.
downPDA	Percentage of down-regulated differential abundance. Default is perDiffAbund/2.
numDataPoints	Number of data points. Default is 100.
AbundProfile	AbundProfile for average abundance profile. It can be either the different profiles used in the paper ("HBR", "BP" and "Wu") or it can be location of the abundance profile. Default is "HBR".
modelFile	Sample data file for model free approach. Default is NULL. If modelFile = NULL, Model Free approach will subsample from Input file. If modelFile = "SingleCell", Model Free approach will subsample from the available single cell RNA-seq data. if modelFile is the name of a count file, this count file will be used as sample file for sub-sampling.
minAbund	Minimum abundance cutoff. Default is 10.
varLibsizes	Variability between library sizes. Default is 0.1.
outlier	Outlier model. Default is FALSE, i.e. outlier model is turned off.
perOutlier	Percentage of added outliers. Default is 0.15.
factorOutlier	Scaling factor to generate outliers. Default is 100.
inputCount	Input count file. Default is NULL. If not NULL, it learns the parameters (modelFile, SampleVar, perDiffAbund, upPDA, and downPDA) from count data.
inputLabel	Label of input count file. The label should be sequence of 0 or 1. Default is NULL.
SimulType	Simulation type. It is used only when user has pilot data. Default is "auto". SimulType = "auto", all the parameters (EntityCount, ControlRep, CaseRep, numDataPoint and others) are learned from user's pilot data. SimulType = "auto1", ControlRep, CaseRep, numDataPoint are specified by user input; while EntityCount and all others are learned from user's pilot data. SimulType = "auto2", EntityCount, ControlRep, CaseRep, numDataPoint are specified by user input; while all others are learned from user's pilot data.

**Value**

count	Count matrix.
DiffAbundList	Differential abundance list.
dataLabel	Data label.

**Author(s)**

Li Juntao, Luo Huaien, Chia Kuan Hui Burton, Niranjan Nagarajan

**References**

Luo Huaien, Li Juntao, Chia Kuan Hui Burton, Shyam Prabhakar, Paul Robson, Niranjan Nagarajan, The importance of study design for detecting differentially abundant features in high-throughput experiments, under review.

**Examples**

```

# generate data with all default options.
data <- generateData()
dim(data$count)
dim(data$DiffAbundList)
data$dataLabel

# generate data with input count.
x <- matrix(rnbinom(1000*15,size=1,mu=10), nrow=1000, ncol=15);
x.lable=c(rep(0,10),rep(1,5))
x[1:50,11:15] <- x[1:50,11:15]*10
x.name=paste("g",1:1000,sep="");
write.table(cbind(x.name,x),"count.txt",row.names =FALSE, sep ='\t')

data <- generateData(inputCount="count.txt",inputLabel=x.lable)
dim(data$count)
dim(data$DiffAbundList)
data$dataLabel

# or generate data with input count and redefined parameters.
data <- generateData(inputCount="count.txt",inputLabel=x.lable,
                    ControlRep=10,CaseRep=10,EntityCount=3000,SimulType="auto2")
dim(data$count)
dim(data$DiffAbundList)
data$dataLabel

```

---

HBR

*Average abundance for RNA-seq data from Human Brain Reference.*


---

**Description**

RNA-seq profile datasets.

**Usage**

```
data(HBR)
```

**Format**

Data frames with 17597 observations on the following 2 variables.

GeneName a character vector

Count a numeric vector

**References**

Au, K.F., Jiang, H., Lin, L., Xing, Y. & Wong, W.H. Detection of splice junctions from paired end RNA-seq data by SpliceMap. *Nucleic Acids Res* 38, 4570-4578.

---

plotPRC *plot precision-recall curves*

---

**Description**

plot precision-recall curves for each test.

**Usage**

```
plotPRC(obj, DE.methods=c("Cuffdiff", "DESeq", "baySeq", "edgeR", "MetaStats", "NOISEq"),
  nor.methods=c("default", "Mode", "UQN", "NDE"),
  plot_type = "o", plot_pch = 20, plot_lwd = 1.75, plot_cex = 1)
```

**Arguments**

obj	Object from testDATs().
DE.methods	Method list for differential expression tests. Methods currently available include "Cuffdiff", "DESeq", "baySeq", "edgeR", "MetaStats", "NOISEq".
nor.methods	Normalization method list. Methods currently available include "default"(default normalization for each DE method), "Mode"(Mode normalization), "UQN"(Upper quartile normalization), "NDE"(non-differential expression normalization).
plot_type	type option in plot.
plot_pch	pch option in plot.
plot_lwd	lwd option in plot.
plot_cex	cex option in plot.

**Author(s)**

Li Juntao and Chia Kuan Hui Burton

**References**

Luo Huaïen, Li Juntao, Chia Kuan Hui Burton, Shyam Prabhakar, Paul Robson, Niranjan Nagarajan, The importance of study design for detecting differentially abundant features in high-throughput experiments, under review.

**Examples**

```
data <- generateData(EntityCount=500)
test.obj <- testDATs(data, DE.methods=c("DESeq", "edgeR"), nor.methods="default")
auc.obj <- computeAUC(test.obj)
plotPRC(auc.obj)
```

---

plotROC

*plot Receiver Operating Characteristic curve*


---

**Description**

plot Receiver Operating Characteristic curve for each test.

**Usage**

```
plotROC(obj,DE.methods=c("Cuffdiff","DESeq","baySeq","edgeR","MetaStats","NOISeq"),
nor.methods=c("default","Mode","UQN","NDE"),
plot_type = "o",plot_pch = 20,plot_lwd = 1.75,plot_cex = 1)
```

**Arguments**

obj	Object from testDATs().
DE.methods	Method list for differential expression tests. Methods currently available include "Cuffdiff","DESeq","baySeq","edgeR","MetaStats","NOISeq".
nor.methods	Normalization method list. Methods currently available include "default"(default normalization for each DE method),"Mode"(Mode normalization),"UQN"(Upper quartile normalization),"NDE"(non-differential expression normalization).
plot_type	type option in plot.
plot_pch	pch option in plot.
plot_lwd	lwd option in plot.
plot_cex	cex option in plot.

**Author(s)**

Li Juntao and Chia Kuan Hui Burton

**References**

Luo Huaïen, Li Juntao, Chia Kuan Hui Burton, Shyam Prabhakar, Paul Robson, Niranjan Nagarajan, The importance of study design for detecting differentially abundant features in high-throughput experiments, under review.

**Examples**

```
data <- generateData(EntityCount=500)
test.obj <- testDATs(data,DE.methods=c("DESeq","edgeR"),nor.methods="default")
auc.obj <- computeAUC(test.obj)
plotROC(auc.obj)
```



---

 SingleCell

*Single-cell RNA-seq data for model free simulation*


---

**Description**

Single-cell RNA-seq.

**Usage**

```
data(SingleCell)
```

**Format**

Data frames with 51516 rows and 96 columns.

**Details** This is single-cell RNA-seq data from which counts were generated when using RNA-seq model free approach.

**References**

In-house data.

---

 testDATs

*Run differential abundance testings*


---

**Description**

Perform differential abundance testing on simulated count data.

**Usage**

```
testDATs(data, numCores=10, minCountsThreshold=0,
DE.methods=c("Cuffdiff", "DESeq", "baySeq", "edgeR", "MetaStats", "NOISeq"),
nor.methods=c("default", "Mode", "UQN", "NDE"), method.list=NULL)
```

**Arguments**

data	Data object from generateData() function or predefined data object similar to the output of generateData().
numCores	Number of cores for parallelization. Default is 10.
minCountsThreshold	Minimum counts threshold for filtering. Default is 0.
DE.methods	Method list for differential expression tests. Methods currently available include "Cuffdiff", "DESeq", "baySeq", "edgeR", "MetaStats", "NOISeq".

<code>nor.methods</code>	Normalization method list. Methods currently available include "default" (default normalization for each DE method), "Mode"(Mode normalization), "UQN"(Upper quartile normalization), "NDE"(non-differential expression normalization).
<code>method.list</code>	The method list for the combination of <code>DE.methods</code> and <code>nor.methods</code> . Default is NULL.

**Value**

<code>data</code>	Data object from <code>generateData()</code> function.
<code>filterCounts</code>	filtered count data.
<code>Cuffdiff</code>	Result form Cuffdiff with default normalization.
<code>Cuffdiff_uqn</code>	Result form Cuffdiff with Upper quartile normalization normalization.
<code>Cuffdiff_Mode</code>	Result form Cuffdiff with Mode normalization.
<code>Cuffdiff_nde</code>	Result form Cuffdiff with non-differential expression normalization.
<code>DESeq</code>	Result form DESeq with default normalization.
<code>DESeq_uqn</code>	Result form DESeq with Upper quartile normalization normalization.
<code>DESeq_Mode</code>	Result form DESeq with Mode normalization.
<code>DESeq_nde</code>	Result form DESeq with non-differential expression normalization.
<code>baySeq</code>	Result form baySeq with default normalization.
<code>baySeq_uqn</code>	Result form baySeq with Upper quartile normalization normalization.
<code>baySeq_Mode</code>	Result form baySeq with Mode normalization.
<code>baySeq_nde</code>	Result form baySeq with non-differential expression normalization.
<code>edgeR</code>	Result form edgeR with default normalization.
<code>edgeR_uqn</code>	Result form edgeR with Upper quartile normalization normalization.
<code>edgeR_Mode</code>	Result form edgeR with Mode normalization.
<code>edgeR_nde</code>	Result form edgeR with non-differential expression normalization.
<code>MetaStats</code>	Result form MetaStats with default normalization.
<code>MetaStats_uqn</code>	Result form MetaStats with Upper quartile normalization normalization.
<code>MetaStats_Mode</code>	Result form MetaStats with Mode normalization.
<code>MetaStats_nde</code>	Result form MetaStats with non-differential expression normalization.
<code>NOISeq</code>	Result form NOISeq with default normalization.
<code>NOISeq_uqn</code>	Result form NOISeq with Upper quartile normalization normalization.
<code>NOISeq_Mode</code>	Result form NOISeq with Mode normalization.
<code>NOISeq_nde</code>	Result form NOISeq with non-differential expression normalization.

**Author(s)**

Li Juntao, Luo Huaien, Chia Kuan Hui Burton, Niranjan Nagarajan

## References

Luo Huaïen, Li Juntao, Chia Kuan Hui Burton, Shyam Prabhakar, Paul Robson, Niranjan Nagarajan, The importance of study design for detecting differentially abundant features in high-throughput experiments, under review.

## Examples

```
data <- generateData(EntityCount=100)
test.obj <- testDATs(data,nor.methods="default")
test.obj <- testDATs(data,DE.methods="DESeq")

# test data with input count.
x <- matrix(rnbinom(1000*15,size=1,mu=10), nrow=1000, ncol=15);
x[1:50,11:15] <- x[1:50,11:15]*10
x.name=paste("g",1:1000,sep="");
write.table(cbind(x.name,x),"count.txt",row.names =FALSE, sep ='\t')

x <- read.table("count.txt",head=TRUE,sep='\t')
x.count <- x[,2:16]
x.lable=c(rep(0,10),rep(1,5))
row.names(x.count) <- x[,1]
data <- list(count=x.count,dataLabel=x.lable)
test.obj <- testDATs(data,DE.methods=c("DESeq","edgeR"),nor.methods="default")
```

---

Wu

*Average abundance for RNA-seq data from schizophrenia.*

---

## Description

RNA-seq profile datasets.

## Usage

```
data(Wu)
```

## Format

Data frames with 18982 observations on the following 2 variables.

gene a character vector

expression a numeric vector

## References

Wu, J.Q. et al. Transcriptome sequencing revealed significant alteration of cortical promoter usage and splicing in schizophrenia. PloS One 7, e36351-e36351.

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