Package ‘oneChannelGUI’

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Date 2 April 2015  
Title A graphical interface designed to facilitate analysis of microarrays and miRNA/RNA-seq data on laptops  
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Suggests annotate, genefilter, maSigPro, pamr, pdmclass, ChIPpeakAnno, chipseq, BSGenome, Rgraphviz, affy, annaffy, affyPLM, multtest, ssize, sizepower, RankProd, org.Hs.ot.db, org.Mm.ot.db, org.Rn.ot.db, edgeR, metaArray, MergeMaid, biomaRt, GenomeGraphs, AffyCompatible, rtracklayer, Genominator, EDASeq, limma, DESeq, DESeq2, goseq, hguene10stranscriptcluster.db, mogen10stranscriptcluster.db, ragen10stranscriptcluster.db, G0stats, AnnotationDbi, preprocessCore, baySeq, HuExExonProbesetLocation, MoExExonProbesetLocation, RaExExonProbesetLocation, snow, RmiR, RmiR.Hs.miRNA, BSGenome.Hsapiens.UCSC.hhv19, R.utils, cummeRbund, BSGenome.Mmusculus.UCSC.mm9, BSGenome.Rnorvegicus.UCSC.rn4, DESeq2, GenomicAlignments, BiocParallel, KEGG.db, miRNAPath, miRNAtap, miRNAtap.db  
Description This package was developed to simplify the use of Bioconductor tools for beginners having limited or no experience in writing R code. This library provides a graphical interface for microarray gene and exon level analysis as well as miRNA/mRNA-seq data analysis  
License Artistic-2.0  
URL http://www.bioinformatica.unito.it/oneChannelGUI/  
biocViews Sequencing, RNASeq, Microarray, OneChannel, DataImport, QualityControl, Preprocessing, StatisticalMethod, DifferentialExpression, GUI, MultipleComparison  
NeedsCompilation no
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Description

This package is directed to Bioconductor beginners that have little or no experience in writing R code. The package implements, as simple functions accessible over the affylmGUI graphical interface, some code useful for QC, data filtering, data output manipulation and identification of GO enriched classes.

Details

| Package:   | oneChannelGUI |
| Type:      | Package       |
| Version:   | 1.0           |
| Date:      | 2006-12-16    |
| License:   | GPL version 2 or newer |
**adaptorTrimm**

**Author(s)**

Author: Raffaele A Calogero
Maintainer: Raffaele A Calogero <raffaele.calogero@unito.it>

**Examples**

```r
# library(oneChannelGUI)
## To start the oneChannelGUI with the modifications
# oneChannelGUI()
```

---

**adaptorTrimm**  
**Trimming 3,5 end primers for miRNAs NGS**

**Description**

A function to trim 3, 5 end primers for miRNAs NGS fastq files. The function is not very efficient. It takes about 5 minutes for 1 million reads file.

**Usage**

```r
adaptorTrimm()
```

**Author(s)**

Raffaele A Calogero

---

**annotateNGSeset**  
**Associate annotation to NGS data loaded in oneChannelGUI**

**Description**

This function associates annotation to NGS data loaded in oneChannelGUI, using the information retrieved by genome annotation from ENSEMBL by ChIPpeakAnn package.

**Usage**

```r
annotateNGSeset()
```

**Author(s)**

Raffaele A Calogero
bayseqInterface

AptMidas

Graphical interface to APT midas

Description

This is a graphical interface to the midas program for detection of alternative splicing detection implemented in the Affymetrix APT tools

Usage

AptMidas()

Note

For more information see Affymetrix Alternative Transcript Analysis Methods for Exon Arrays whitepaper. Before using MiDAS is strongly recomanded to filter out gene level probe sets with low intensity values to avoid searching alternative splicing for probe sets which are not expressed. This can be done using filtering method implemented in oneChannelGUI which define a background intensity threshold on the basis of the intron exon signals of a set of housekeeping genes present in the exon arrays. However it is also possible to use a filter based on the dabg p-value calculated using Affymetrix APT tools. This function will also calculate Splice Index

Author(s)

Raffaele A Calogero

See Also

erankProdAltSpl

bayseqInterface

graphical interface to baySeq library.

Description

BaySeq provides statistical routines for determining differential expression in digital gene expression data. The routines can be applied equally to two groups experiments only.

Usage

bayseqInterface()

Details

This interface reorganizes NGS data loaded in oneChannelGUI to be analysed by baySeq

Author(s)

Raffaele A Calogero
**biomartFilter**

*Filtering only gene-level probe sets with multiple ensembl transcripts*

**Description**

This function allows to filter exon array data to selected only those gene-level probe sets associated to multiple mRNAs annotated in ensembl data base.

**Usage**

```r
biomartFilter()
```

**Author(s)**

Raffaele A Calogero

---

**bofa**

*Bovine microRNA data set*

**Description**

microRNA data set fasta produced using miRbase B taurus precursors reformated to have only miR name as descriptor of the fasta file.

**Usage**

```r
bofa
```

**Format**

Fasta file

---

**bowtieBuilt**

*Downloading Bowtie prebuilt reference sets*

**Description**

A function to download built reference sets for Human Mouse and Rat, available as precompiled at sourceforge oneChannelGUI.extras project.

**Usage**

```r
bowtieBuilt()
```

**Author(s)**

Raffaele A Calogero
bowtieDownload

*Description*

A function to download Bowtie and Picard tools for primary mapping of short reads and conversion of SAM to BAM format. Reference mapping for Bowtie are also generated using miRBase precursors sets for Human Mouse and Rat. Human, mouse and Rat gene-level scaffolds are also downloaded by sourceforge

*Usage*

```r
bowtieDownload()
```

*Author(s)*

Raffaele A Calogero

bowtieGenomes

*Description*

A function to build Bowtie reference sets for short reads primary mapping.

*Usage*

```r
bowtieGenomes()
```

*Author(s)*

Raffaele A Calogero

buildingLocalAnnotation

*Description*

Updates local gene-level annotation data for gene and exon arrays using the netaffx database.

*Usage*

```r
buildingLocalAnnotation()
```
chrLength

Author(s)

Raffaele A Calogero

Description

A list containing chromosome lengths for hg19, mm9 and rn4 genome releases

Usage

chrLength

Format

A list with three observations: chromosome lengths for hg19, mm9 and rn4 genome releases

Author(s)

Raffaele A Calogero

colExtract

Description

Extract a column from a tab delimited file with header

Usage

colExtract()

Author(s)

Raffaele A Calogero
**Description**

The function combines in a unique ExpressionSet the data derived from multiple Matrix Series Files belonging to a GEO experiment containing more than 255 arrays.

**Usage**

combineGeoMSF()

**Note**

see oneChannelGUI vignette for more info

**Author(s)**

Raffaele A Calogero

---

**Description**

The function allows to combine two matrices extracted from oneChannelGUI.

**Usage**

combining2eSet()

**Note**

see oneChannelGUI vignette for more info

**Author(s)**

Raffaele A Calogero
consistentFilters

This function allows filtering using the combination of multiple parameters, e.g. MiDAS p-values and Rank Product p-values

Description

This filter can be used to moderate multiple tests errors. E.g. finding the intersection between MiDAS p-values and Rank Product p-values user will remove some of the false positive produced by the two methods. A filter on the size of delta Splice Index associated to MiDAS p-values filter will allow to remove statistical significant splicing events which are characterized by a very limited variation.

Usage

consistentFilters()

Note

This function needs the presence of Splice Index data, MiDAS p-values and RP p-values. It works for two groups only

Author(s)

Raffaele A Calogero

See Also

erankProdAltSpl, AptMidas

---

cosieWrapper

A wrapper to cosie, Corrected Splicing Indices for Exon arrays, that for any given set of new exon array experiments corrects for the observed bias and improves the detection of alternative splicing

Description

COSIE is a function that for a given set of exon arrays corrects for the observed bias and improves the detection of alternative splicing. It adjusts splicing indices for exons, especially for those that belong to differentially expressed genes. For this adjustment, COSIE uses parameters that are specific for each probeset which were trained from a large number of published exon arrays. The downside of this approach is that such parameters cannot be estimated for all probesets on the microarray. Based on our training set, COSIE corrects 95.1 percent of the probesets. Separate parameter files are provided for both the full and core sets, including all probesets that are linked to transcripts. We recommend the use of the core set that was also used in the cited study below. The full set is not as well characterized.

Usage

cosieWrapper()
createGeoTarget  

*Creating a affylmGUI Target starting from a GEO Matrix series file*

**Description**

The function extracts from GEO series matrix file all the information to create a Target file, that can be used to load the GEO series matrix file into oneChannelGUI.

**Usage**

```r
createGeoTarget()
```

**Note**

see oneChannelGUI vignette for Target file description

**Author(s)**

Raffaele A Calogero

crosshybFilter  

*Removing from exon array gene/exon level probe sets characterized by cross hybridization with other transcripts*

**Description**

XHYB field is mainly an indicator of weak assignment between a transcript cluster and the assigned mRNA, suggesting a potential crosshyb, CRSSHYB is a measure of the promiscuity of the probes within a probe set among transcribed sequences.

1. 1 unique. All probes in the probe set perfectly match only one sequence in the putatively transcribed array design content. The vast majority of probe sets are unique.
2. 2 similar. All the probes in the probe set perfectly match more than one sequence in the putatively transcribed array design content.
3. 3 mixed. The probes in the probe set either perfectly match or partially match more than one sequence in the putatively transcribed array design content.

XHYB and CRSSHYB are used to remove probe sets characterized by multiple hybridization of exon probes

**Usage**

```r
crosshybFilter()
```
**ctrtrtHeatmap**

**Author(s)**

Raffaele A Calogero

---

**Description**

This function needs a maSigPro target encompassing only one treatment condition. This function is ideal to visualize the expression data for putative mRNA targets upon miRNA expression perturbation with antagonir or mimics. Expression data reorganized as the average expression differences between treatment and untreated condition over the time course.

**Usage**

```r
ctrtrtHeatmap()
```

**Author(s)**

Raffaele A Calogero

---

**desGenes**

**Description**

This function calculates differential gene expression for an RNA-seq experiment.

**Usage**

```r
desGenes()
```

**Details**

This function uses DESeq package to calculate differential expression at gene level starting from a gene table generated by DEXseq package. It returns an rda object called res.genes that can be used for further analysis.

**Author(s)**

Raffaele A Calogero
**dexCounts**

**Description**
This function summarises counts using DEXseq python scripts.

**Usage**

dexCounts()

**Details**
This function uses the python scripts present in DEXseq package to summarise at the level of exons the reads derived by a sam file. The function needs to be run for each sample. It runs in background. It is necessary the installation of HTSeq python library

**Author(s)**
Raffaele A Calogero

---

**dexExons**

**Description**
This function calculates differential expression at exon level for an RNA-seq experiment.

**Usage**

dexExons()

**Details**
This function uses DEXseq package to calculate differential expression at exon level starting from a set of counts files located in different folders and generated by dexCounts function. The target file is the one used to run tophat in oneChannelGUI. it executes in background a batch file that returns two rda object called res.exons and ecs that can be used for further analysis. res.exons is contains all results describing differential expression at exon level and ecs is the DEXseq object containing all info for the exons.

**Author(s)**
Raffaele A Calogero
**Description**

MA and Volcano plots can be generated starting from limma results summarized in a topTable. Specific subsets of the topTable defined by p-value below an user-defined threshold and/or log2 fold changes over an user-defined threshold can be saved. The subset of data can be saved as a tab delimited file

**Usage**

```r
dfMAplot(table1)
```

**Arguments**

- `table1`: topTable data.frame generate by affyImGUI

**Note**

To know more about topTable see limma help

**Author(s)**

Raffaele A Calogero

---

**Description**

Plot the Nts frequency for mapped reads extracted from a BAM file

**Usage**

```r
EDANtFreq()
```

**Details**

Interface to the plotNtFrequency of EDASeq package

**Author(s)**

Raffaele A Calogero

**See Also**

EDAplotQuality, EDAreadNumber
**EDAreadNumber**

Plotting the number of mapped reads extracted from a BAM file

**Usage**

EDAreadNumber()

**Details**

Interface to the barplot of EDASeq package

**Author(s)**

Raffaele A Calogero

**See Also**

EDAplotQuality, EDANtFreq

---

**EDAplotQuality**

Plotting Quality scores for mapped reads extracted from BAM files

**Description**

Plotting the quality scores of mapped reads extracted from a BAM file

**Usage**

EDAplotQuality()

**Details**

Interface to the plotQuality of EDASeq package

**Author(s)**

Raffaele A Calogero

**See Also**

EDAreadNumber, EDANtFreq
edgerInterface

Description

edgerInterface is a graphical interface to edgeR library.

edgeR provides statistical routines for determining differential expression in digital gene expression data. The routines can be applied equally to SAGE, CAGE, Illumina, Solexa, 454 or ABI SOLiD experiments.

Usage

edgerInterface()

Details

This interface reorganizes NGS data loaded in oneChannelGUI to be analysed by edgeR.

Author(s)

Raffaele A Calogero

edgeRNorm

Description

This function scale-normalize NGS raw data as described by Robinson and Oshlack Genome Biology 2010, 11:R25

Usage

dgeRNorm()

Author(s)

Raffaele A Calogero
### EG2probeset

This function allows to link GeneBank and Entrez Gene ids to gene-level probe set ids.

**Description**

This function allows to link oneChannelGUI embedded Affymetrix annotated accession numbers to gene-level probe set ids. Using the ACC EG are linked using the Bioconductor human, mouse or rat LLMappings annotation library.

**Usage**

```r
EG2probeset()
```

**Author(s)**

Raffaele A Calogero

---

### erankProdAltSpl

Implementation of the Rank Product method for the detection of alternative splicing events.

**Description**

This is a graphical interface to the RP function from RankProd package applied to detection of alternative splicing.

**Usage**

```r
erankProdAltSpl()
```

**Details**

Before using this method it is strongly suggested to perform a filter on the basis of DABG p-values using the filtering function available in the filtering menu. DABG values can be calculated if exon array probe set data are generated using the oneChannelGUI graphical implementation to APT tools. Affymetrix suggests to calculate probe set intensity at gene level using iterPlier and at exon level using plier. Subsequently SpliceIndex need to be calculated using the function available in the exon menu. Finally the Rank Product method could be applied exon by exon. For more details on the method see RankProd package. Selection of putative alternative splicing could be done using the filtering function available in the filtering menu of oneChannelGUI.

**Note**

IMPORTANT we are still evaluating the efficacy of this method for detection of alternative splicing events. Use it being conscious of this.

**Author(s)**

Raffaele A Calogero
**erankProdAltSplFilter**

*Filtering Rank Product results for the detection of alternative splicing events*

**Description**

This is a graphical interface to filter data on the basis of p-value generated by rank product analysis applied for the detection of alternative splicing.

**Usage**

`erankProdAltSpl()`

**Author(s)**

Raffaele A Calogero

**See Also**

`erankProdAltSpl`, `AptMidas`
exonContrasts  
*Defining t-test regularized p-values*

**Description**

This function constructs the contrasts as affylmGUI but applied to exon-level. It also performs Bayes regularization. Raw p-values are plotted to see if BH or BY type I error correction can be applied. Corrected p-values are saved and used for extraction of alternative spliced exons.

**Usage**

```r
exonContrasts()
```

**Details**

It is important to note that if multiple contrasts should be considered after the calculation of each of them it is essential to extract the alternative spliced exons with exonTopTableExtract, because everytime exonContrasts is run it overwrites the previous results.

**Author(s)**

Raffaele A Calogero

---

exonScaffold  
*stand alone function to generate a scaffold containing only exon location from ENSEMBL*

**Description**

The scaffold contains chr stand start and end position of exon. Ensembl geneID and exonID together with exon rank are also provided.

**Usage**

```r
exonScaffold(genome)
```

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>genome</td>
<td>The genome code for H sapiens is hg19, for M musculus is mm9 and for R norvegicus is rn4</td>
</tr>
</tbody>
</table>

**Details**

The scaffold is used to map reads data into regions only associated to exons

**Author(s)**

Raffaele A Calogero
**exonsSpecific2as**

*Defining the exons associated to the various alternative isoforms*

**Description**

This function uses the output derived from the function mapping2ensembl and produces a list of 1 and 0 for each of the alternative transcripts associated to a specific Entrez Gene. This function is useful to define which splicing events are not associated to exons conserved over all the possible isoforms.

**Usage**

```r
exonsSpecific2as()
```

**Author(s)**

Raffaele A Calogero

---

**exonTopTableExtract**

*Extracts data on the basis of a defined t-test regularized p-value*

**Description**

This function filters the data produced by exonContrasts to extract a list of alternative spliced exons that are saved in a file and they can be used for further analysis, i.e. extracting only variant exons. The function also filter the data present in the onechannelGUI project bot at gene and exon-level.

**Usage**

```r
exonTopTableExtract()
```

**Details**

It is important to note that if multiple contrasts should be considered after the calculation of each of them it is essential to extract the alternative spliced exons with exonTopTableExtract, because everytime exonContrasts is run it overwrites the previous results.

**Author(s)**

Raffaele A Calogero
extractAffyids

Extracting probe ids associated to a specific Gene Ontology term

**Description**

It is possible to identify the affy ids associated to a specific GO term using the extractAffyids function.

**Usage**

```r
extractAffyids()
```

**Details**

The function asks to the user to select a file containing probe set ids separated by carriage return. The file should contain only one column and no header. The user is also asked to select a specific GO term. The probe sets associated to the specific GO term will be annotated and saved in a HTML file.

**Note**

For the annotation the annotation library associated to the raw data loaded in the affylmGUI environment is used.

**Author(s)**

Raffaele A. Calogero

**See Also**

GOenrichment, plotGO

extractmirTargets

Extract miRNA targets showing a opposite regulation with respect to miRNA

**Description**

This function extract from a RmiR output the subset of genes showing a fold change variation inversely correlated to that of the miRNA selected by the user.

**Usage**

```r
extractmirTargets()
```

**Details**

The output has the same structure of an RmiR output

**Author(s)**

Raffaele A Calogero
filteringmiRtargets  
*Subsetting an expression set using a list of gene which are putative targets for a miRNA*

**Description**

This function subsets the normalized expression set present in the affyLMGUI environment on the basis of a list of probe set ids extracted on the basis of the predicted gene targets for a microRNA. Extraction is based on predicted targets for miRNA in human.

**Usage**

filteringmiRtargets()

**Details**

The function asks to the user to select a file containing probe set ids separated by carriage return. The file should contain only one column and no header.

**Note**

In transcriptional studies focusing on genes characterized by specific feature (i.e. transcription factor elements in promoters) the best filtering approach is selecting only those genes linked to the interesting biological feature.

**Author(s)**

Raffaele A Calogero

**See Also**

IPAlistFilter, iqrFilter, intensityFilter

filteringTable  
*Filtering a tab delimited file*

**Description**

This function allows to filter a tab delimited file using a vector of data present in an other file. The two files should have an header and the column name to be used for the filtering should be equal in both files

**Usage**

filteringTable()

**Author(s)**

Raffaele A Calogero
geneExonLibs

**Description**

Affymetrix Gene/Exon library files are necessary to APT tools to calculate probe set summaries. The versions downloaded from [www.bioinformatica.unito.it](http://www.bioinformatica.unito.it), with this function, contain all informations needed to analyze gene exon arrays.

**Usage**

```r
geneExonLibs()
```

**Author(s)**

Raffaele A Calogero

---

genomePlot

**Description**

This function plots average intensity signals over the genes and transcripts structure mapped on ENSEMBL to identify a specific splicing event.

**Usage**

```r
genomePlot()
```

**Author(s)**

Raffaele A Calogero

**See Also**

variantSI, variantExons, makeBED15
genomeStudioReformat

The present function reformat an expression file exported by Genome Studio in a format that is compatible with oneChannelGUI

Description

The present function reformat an expression file exported by Genome Studio in a format that is compatible with oneChannelGUI

Usage

genomeStudioReformat()

Author(s)

Raffaele A Calogero

geoVSbioc

linking GEO platforms to available BioC annotations libraries

Description

This data file gives the linke between GEO platforms and BioC annotation libraries. If the GEO BioC link exists the Bioconductor annotation lib is directly loaded in the annotation fild of the SexpressionSet

Usage

geoVSbioc

Format

A data frame with 4 observations: GEOAcc, Organisms, Title, BiocAnLIb

References

GEO and Bioconductor
getNGSAnnotation  Retrieved genome annotation from ENSEMBL

Description
This function retrieves genome annotation from ENSEMBL using ChIPpeakAnn package

Usage
getNGSAnnotation()

Author(s)
Raffaele A Calogero

GOenrichment  Searching for Gene Ontology enriched terms within a set of differentially expressed genes

Description
In Bioconductor is available a library called GOstats, which allows the calculation of enriched GO terms within a set of differentially expressed probe sets. This is a graphical implementation of a function allowing the extraction of GO enriched term in a sub set of differentially expressed probe sets. To know more about it see GOstat library

Usage
GOenrichment()

Details
The function asks to the user to select a file containing probe set ids separated by carriage return. The file should contain only one column and no header. The set of enriched terms are plotted in red over the graph of all GO term associated to the differentially expressed genes. GO enriched terms can be also saved in a tab delimited file.

Author(s)
Raffaele A Calogero

References
Robert Gentleman GOstat package

See Also
extractAffyids, plotGO
hsfa

**Description**

microRNA data set fasta produced using miRbase H sapiens precursors reformated to have only miR name as descriptor of the fasta file

**Usage**

hsfa

**Format**

Fasta file

---

inspecting.one.splice.index

*Plotting on the profiles of splice indexes for a transcript cluster ID*

**Description**

This function plots the splice index profiles for one transcript cluster ID

**Usage**

inspecting.one.splice.index()

**Author(s)**

Raffaele A Calogero

**See Also**

spliceIndex
inspecting.splice.index

Plotting on a pdf file the profiles of splice indexes

Description

This function prints in a pdf file the splice index profiles of the available genes.

Usage

inspecting.splice.index()

Author(s)

Raffaele A Calogero

See Also

spliceIndex

intensityFilter

intensity filtering with a mouse click

Description

This function removes all probe sets in which a certain percentage of experiments is below a user defined intensity threshold.

Usage

intensityFilter()

Details

The aim of non specific filtering is to remove the genes that are unlikely to carry information about the phenotypes under investigation. This filtering remove genes that do not have a certain level of, user defined, intensities in a set of, user defined, experiments.

Note

Factor analysis will be limited by the problem of having fewer samples than genes. Therefore, preselecting a smaller set of genes is definitively helpful.

Author(s)

Raffaele A Calogero

See Also

iqrFilter, listFilter, IPAlistFilter


**IPAlistFilter**

Filtering an expression set using a set of Entrez genes extracted from Ingenuity Pathways analysis (IPA)

---

**Description**

It is possible to sub set an expression set loaded in the affylmGUI environment starting form a list of Entrez genes derived by IPA search tool.

**Usage**

IPAlistFilter()

**Details**

The function asks to the user to select a file containing Entrez genes separated by carriage return. The file should contain only one column and no header.

**Author(s)**

Raffaele A Calogero

**See Also**

iqrFilter, listFilter, intensityFilter

---

**iqrFilter**

Interquantile filtering with a mouse click

---

**Description**

This function implements the interquantile filtering proposed by Heydebreck in 2004

**Usage**

iqrFilter()

**Details**

The aim of non specific filtering is to remove the genes that are unlikely to carry information about the phenotypes under investigation. This filtering remove genes that show little changes within the experimental points.

**Note**

Factor analysis will be limited by the problem of having fewer samples than genes. Therefore, preselecting a smaller set of genes is definitively helpful.

**Author(s)**

Raffaele A Calogero
References

Heydebreck et al. Bioconductor project Papers 2004

See Also

IPAlistFilter, listFilter, intensityFilter

limma2paired

graphical interface apply linear model to two sample groups with batch effect.

Description

This function allows to run Bayes regularized t-test on two groups with batch effects. Derived from the mailing list question: Paired samples and origin question, Limma 2006-05-11 13:45:41 GMT

Usage

limma2paired()

Details

This function allows to run Bayes regularized t-test on two groups with batch effects.

Author(s)

Raffaele A Calogero

limmaExons

graphical interface to limma for alternative splicing detection

Description

Applying the limma model fitting to exon-level data. Same implementation of AffylmGUI but applied to exon-level data. The first indication of alternative splicing detection using limma was proposed by Shah and Pallas in BMC Bioinformatics. 2009 Jan 20;10:26

Usage

limmaExons()

Details

The function fits the limma linear model to exon-level data

Author(s)

Raffaele A Calogero
listFilter

Subsetting an expression set using a list of Affymetrix ids

Description
This function subsets the normalized expression set present in the affylmGUI environment on the basis of a list of probe set ids passed via flat file.

Usage
listFilter()

Details
The function asks to the user to select a file containing probe set ids separated by carriage return. The file should contain only one column and no header.

Note
In transcriptional studies focusing on genes characterized by specific feature (i.e. transcription factor elements in promoters) the best filtering approach is selecting only those genes linked to the interesting biological feature.

Author(s)
Raffaele A Calogero

See Also
IPAlistFilter, iqrFilter, intensityFilter

makeBED15

This function creates files in BED15 format to be loaded on the UCSC browser

Description
This function creates files in BED15 format to be loaded on UCSC genome browser. The function uses the data derived by variantSI filter on the basis of chromosome annotation

Usage
makeBED15()

Author(s)
Raffaele A Calogero

See Also
variantSI, variantExons, plotVariantSI
makeGCcontent  
*Building a list of gene-level GC frequencies*

**Description**
A function to build a gene-level GC frequencies associated to the genes generated by the makeGeneScaffold function. Built scaffolds with GC content are available as precompiled at sourceforge oneChannelGUI.extras project.

**Usage**
makeGCcontent(scaffold, whichRef)

**Arguments**
- **scaffold**  The GRange object created by makeGeneScaffold
- **whichRef**  The genome of interest to be used to create a gene-level scaffold

**Author(s)**
Raffaele A Calogero

**See Also**
makeGeneScaffold, wrapScaffold

makeGeneScaffold  
*Building a GRange object at chromosome level on UCSC genome data*

**Description**
A function to build a gene level scaffold to associate reads derived from bowtie mapping. The function is embedded in bowtieGenome function that build, at chromosome level, the reference genome for primary mapping. Built scaffolds are available as precompiled at sourceforge oneChannelGUI.extras project.

**Usage**
makeGeneScaffold(whichRef)

**Arguments**
- **whichRef**  The genome of interest to be used to create a gene-level scaffold

**Author(s)**
Raffaele A Calogero

**See Also**
makeGCcontent, wrapScaffold
makeSam

Description

This function uses samtools to convert a bam file in a sam file. It assumes that the bam file was generated by TopHat. Therefore it searches for accepted_hits.bam in the folder.

Usage

makeSam()

Details

This function uses samtools to convert a bam file in a sam file. It assumes that the bam file was generated by TopHat. Therefore it searches for accepted_hits.bam in the folder. It produces a file named accepted_hits.sam.

Author(s)

Raffaele A Calogero

mapping2ensembl

Associating e-level probe sets to entrez gene exonic structure

Description

This function associates the statistical and expression data produced by a oneChannelGUI exon-level analysis to the exonic structure of Entrez Gene ID. This function uses biomaRt to retrieve the sequence of EG exons. RRE database is instead used to retrieve the exon-level target sequences. Any exon-level probe set id to be associated to the EG exonic sequence need to be a perfect matching substring of the exon. In the other case no exon is associated to the probe set.

Usage

mapping2ensembl()

Author(s)

Raffaele A Calogero
mapping2exon

This function maps on exon-level Probe Selection Region (PSR) starting for the file produced by function oneChannelGUI: Mapping exon level probe sets to Reference Sequences

Description
This function retrieve from RRE the PSR sequences associated to the exon-level probe sets and all exons associated to the gene associated to PRS. Subsequently identify the exon where PSR maps and produces a fasta file were are located exon-level PSR and target exon. The mapping is done using the countPattern function of the Biostrings package. Up to three mismathces are allowed in PSR mapping on exonic sequence.

Usage
mapping2exon()

Author(s)
Raffaele A Calogero

mapping2RefSeq

This function maps on NCBI Reference sequences spliced exons detected by the function oneChannelGUI: Inspecting splice indexes

Description
This function retrieve from RRE the PSR sequences associated to the exon-level probe sets using blastn detects the best refseq associated to any of the exon-level probe sets retrieve from org.XX,eg.db the EG associated to any of the detected refseq and retrieves all the refseqs associated to the EG. Subsequently check if PSR maps on all the refseqs associated to the eg (conserved exon) or only some of them (isoform specific exon)

Usage
mapping2RefSeq()

Author(s)
Raffaele A Calogero
The function executes maSigPro analysis

Description
The function creates: 1. Create a regression matrix for the full regression model (make.design.matrix function). 2. Computes the p-value associated to the F-Statistic of the model, which is used to select significant genes (p.vector function). 3. Applies a variable selection procedure to find significant variables for each gene (T.fit function). This will ultimately be used to find which are the profile differences between experimental groups. 4. Finally, it generates lists of significant genes according to R-squared of the models (get.siggenes function). To know more about the various steps see maSigPro help.

Usage
masigpro()

Author(s)
Raffaele A Calogero

See Also
masigpro.edesign, masigpro.view

The function creates an edesign object needed to run maSigPro

Description
The function creates an edesign object needed to run maSigPro. To know more about edesign object see maSigPro help. This function uses a specific configuration of Target column of the affylmGUI target file. To know more about target file see affylmGUI help. Each row of the column named Target, in the affylmGUI target file, describes the array on the basis of the experimental design. Each element needed for the construction of edesign is separated from the others by an underscore. The first three elements of the row are fixed and represent Time Replicate Control all separated by an underscore: Time_Replicate_Control. All the other elements refer to various experimental conditions. Considering two different conditions to be evaluated each row is made of 5 elements: Time_Replicate_Control_cond1_cond2 all separated by an underscore. Having an experiment made of 9 arrays, with two time points, 0h and 24h, in triplicate, and two different experimental conditions to be evaluated, the affylmGUI target file will look like:

<table>
<thead>
<tr>
<th>Name</th>
<th>FileName</th>
<th>Target</th>
</tr>
</thead>
<tbody>
<tr>
<td>mC1</td>
<td>M1.CEL</td>
<td>0_1_1_0_0</td>
</tr>
<tr>
<td>mC2</td>
<td>M4.CEL</td>
<td>0_1_1_0_0</td>
</tr>
<tr>
<td>mC3</td>
<td>M7.CEL</td>
<td>0_1_1_0_0</td>
</tr>
<tr>
<td>mE1</td>
<td>M3.CEL</td>
<td>24_2_0_1_0</td>
</tr>
<tr>
<td>mE2</td>
<td>M6.CEL</td>
<td>24_2_0_1_0</td>
</tr>
<tr>
<td>mE3</td>
<td>M9.CEL</td>
<td>24_2_0_1_0</td>
</tr>
<tr>
<td>mI1</td>
<td>M2.CEL</td>
<td>24_3_0_0_1</td>
</tr>
<tr>
<td>mI2</td>
<td>M5.CEL</td>
<td>24_3_0_0_1</td>
</tr>
<tr>
<td>mI3</td>
<td>M8.CEL</td>
<td>24_3_0_0_1</td>
</tr>
</tbody>
</table>
Usage
masigpro.edesign()

Author(s)
Raffaele A Calogero

See Also
masigpro, masigpro.view

The function allows the visualization of maSigPro results

Description
The function is a graphical implementation of the maSigPro PlotGroups function. To know more about it see maSigPro help.

Usage
masigpro.view()

Author(s)
Raffaele A Calogero

See Also
masigpro.edesign, masigpro

metaArrayIC
Graphical interface to metaArray Integrative Correlation function

Description
The integrative correlation analysis (Parmigiani et al., 2004) is a convenient tool to monitor the interstudy concordance of within-study correlations of gene expression. The gene-specific reproducibility score takes the correlation between each gene and all other genes within individual study and calculate the average correlation of these correlations across all pairs of studies.

Usage
metaArrayIC()

Author(s)
Raffaele A Calogero

References
MergeMaid package and metaArray Package
**metaArrayMerge**

Tool to create a merge object for metaArray package

**Description**
This function will create an ExpressionSet from a study starting from a tab delimited file and a target file this ExpressionSet will be merged with the NormalizedAffyData if they contain the same number of row and rownames in the same order. Data generated with this function could be analyzed using metaArrayIC function.

**Usage**

```r
metaArrayIC()
```

**Author(s)**
Raffaele A Calogero

**See Also**

mataArrayIC

---

**miRNAbowtieRun**

Primary mapping of short reads with Bowtie/Picard for miRNA

**Description**
A function for primary mapping of short reads with bowtie and the conversion of the SAM output into BAM format. In the actual implementation it is possible to run bowtie using single-end fastq files produced using Illumina platform. The available reference sets are derived by miRbase precursors and are available for human, hs, mouse, mm, rat, rn, and bovine, bo. It is strongly suggested to run a trimming of the 3 and 5 end linkers before performing the mapping with bowtie. In the present implementation bowtie runs with the following parameters: -a –best -k 1 -q -v 3 -S. Therefore only the first best alignment is shown, input files are in fastq format, alignment up to three mismatches are considered and the output is in SAM format. At the end of the mapping SAM files are converted in BAM files using picard tools.

**Usage**

```r
miRNAbowtieRun()
```

**Author(s)**
Raffaele A Calogero
ML.edesign

The function creates an data frame containing the parameters useful for class prediction.

Description

This function uses a specific configuration of Target column of the affylmGUI target file. To know more about target file see affylmGUI help. Each row of the column named Target, in the affylmGUI target file, describes the array on the basis of the experimental design. Each element needed for the construction of the data frame is separated from the others by an underscore. All the other elements refer to experimental conditions or clinical parameters. The absence of a parameter NEEDS to be described in the Target file by NA Considering two different conditions to be evaluated each row is made of 5 elements: Time_Replicate_Control_cond1_cond2 all separated by an underscore.

Having an experiment made of 9 arrays with 4 different experimental parameters the affylmGUI target file will look like:

<table>
<thead>
<tr>
<th>Name</th>
<th>FileName</th>
<th>Target</th>
</tr>
</thead>
<tbody>
<tr>
<td>mC1</td>
<td>M1.CEL</td>
<td>0_1_pos_0 NA</td>
</tr>
<tr>
<td>mC2</td>
<td>M4.CEL</td>
<td>0_1_pos_0 yes</td>
</tr>
<tr>
<td>mC3</td>
<td>M7.CEL</td>
<td>0_1_neg_0 no</td>
</tr>
<tr>
<td>mE1</td>
<td>M3.CEL</td>
<td>24_2_neg_1 NA</td>
</tr>
<tr>
<td>mE2</td>
<td>M6.CEL</td>
<td>24_2 NA_1 yes</td>
</tr>
<tr>
<td>mE3</td>
<td>M9.CEL</td>
<td>24_2_neg_1 yes</td>
</tr>
<tr>
<td>mI1</td>
<td>M2.CEL</td>
<td>12_3_0 pos yes</td>
</tr>
<tr>
<td>mI2</td>
<td>M5.CEL</td>
<td>12_3_0 pos no</td>
</tr>
<tr>
<td>mI3</td>
<td>M8.CEL</td>
<td>12_3_0 pos no</td>
</tr>
</tbody>
</table>

Usage

ML.edesign()

Author(s)

Raffaele A Calogero

mmfa

Mouse microRNA data set

Description

microRNA data set fasta produced using miRbase M musculus precursors reformatted to have only miR name as descriptor of the fasta file.

Usage

mmfa

Format

Fasta file
**mRNAbowtieRun**

*Primary mapping of short reads with Bowtie/Picard for mRNA-seq*

**Description**

A function for primary mapping of short reads with bowtie and the conversion of the SAM output into BAM format. In the actual implementation it is possible to run bowtie using pair-end fastq files produced using Illumina platform. The available reference sets are built for human, hs, mouse, mm and rat, rn. Analysis is performed chromosome by chromosome to limit RAM consumption. In the present implementation bowtie runs with the following parameters: `-a –best -k 1 -q -v 3 -S`. Therefore only the first best alignment is shown, input files are in fastq format, alignment up to three mismatches are considered and the output is in SAM format. At the end of the mapping SAM files are converted in BAM files using picard tools.

**Usage**

mRNAbowtieRun()

**Author(s)**

Raffaele A Calogero

---

**mRNAmiRCor**

*Filtering by mean of correlation between expression data and miRNA expression changes*

**Description**

This function extract the subset of gene ids which are associated to the changes of expression on putative miRNA targets upon perturbation of miR expression.

**Usage**

mRNAmiRCor()

**Details**

To run this analysis a time course experiment suitable for maSigPro analysis is needed. Expression data need to have at least 3 time points for samples transfected with a scrambled and with an Antagomir or a Mimic. The expression changes at the same time points used for microarray data need to be measured also for miRNA. miRNA expression data need to be saved as table delimited file with two columns: column Name, which has to be organized exactly as the target file Name column, and a column named Value with expression of miRNA, e.g. deltaCt. Target file for mRNA data need to be organized as for maSigPro time course analysis. Output is a three column (gene level id, miRNA-mRNA correlation, mRNA untreated - mRNA treated correlation) tab delimited file with gene IDs and correlation coefficient between mRNA profile and miRNA profile upon treatment with antagonir, i.e. mRNA expression should rise due to the reduction of miRNA expression, or mimics, i.e. mRNA expression should be reduced due to the over-expression of the specific miRNA.
Author(s)
Raffaele A Calogero

See Also
IPAlistFilter, iqrFilter, intensityFilter

---

**myExpresso**

*Running the affy expresso function with the widget*

---

**Description**

Various probe set intensity summary and normalization can be customized using the expresso function.

**Usage**

```r
myExpresso()
```

**Details**

This function run expresso with the graphical interface for parameters selection. It is important to note that expresso is more slow than the C coded rma)

Author(s)
Raffaele A Calogero

---

**ncHs.data**

*Subset of genomic regions with ncRNA of Homo sapiens*

---

**Description**

Two IRanges object referring to the plus and minus strand of genomic region encoding for ncRNAs. The ncRNAs taken in consideration are: miRNA, Mitochondrial rRNA, Mitochondrial tRNA, rRNA, snoRNA, snRNA

**Usage**

```r
ncHs.data
```

**Format**

A list with two observations: IRanges object referring to the plus and minus strand of genomic region encoding of ncRNAs

Author(s)
Raffaele A Calogero
ncMm.data

Subset of genomic regions with ncRNA of Mus musculus

Description

Two IRanges object referring to the plus and minus strand of genomic region encoding ncRNAs. The ncRNAs taken in consideration are: miRNA, Mitochondrial rRNA, Mitochondrial tRNA, rRNA, snoRNA, snRNA

Usage

ncMm.data

Format

A list with two observations: IRanges object referring to the plus and minus strand of genomic regions encoding ncRNAs

Author(s)

Raffaele A Calogero

ncRn.data

Subset of genomic regions with ncRNA in Rattus norvegicus

Description

Two IRanges object referring to the plus and minus strand of genomic region encoding ncRNAs. The ncRNAs taken in consideration are: miRNA, Mitochondrial rRNA, Mitochondrial tRNA, rRNA, snoRNA, snRNA

Usage

ncRn.data

Format

A list with two observations: IRanges object referring to the plus and minus strand of genomic region encoding ncRNAs

Author(s)

Raffaele A Calogero
ncScaffold

*stand alone function to generate a scaffold containing only ncRNA location from ENSEMBL*

Description

The scaffold contains chr stand start and end position of ncRNA. ENSEMBL retrieved species are miRNA, Mt_rRNA, Mt_tRNA, rRNA, snoRNA, snRNA

Usage

ncScaffold(genome, fasta)

Arguments

genome The genome code for H sapiens is hg19, for M musculus is mm9 and for R norvegicus is rn4
fasta If fasta is TRUE a fasta file with the sequences retrieved by ncScaffold will be produced

Details

The scaffold is used to map reads data into regions only associated to ncRNAs

Author(s)

Raffaele A Calogero

NGSreformat

*reformatting output of NGS primary tools software*

Description

This function allows to reorganize the output files produced by SHRIMP, MicroRazerS in two files with the extension: bed and logos used by oneChannelGUI

Usage

NGSreformat()

Author(s)

Raffaele A Calogero
normBoxplot

Box plot of the arrays data available in NormalizeAffyData slot

Description
Box plot visualization of normalized array data

Usage
normBoxplot()

Author(s)
Raffaele A Calogero

ocPlotHist
Gene/Exon level density plots

Description
This function runs a modified version of the plotHist of the affycoretools to be used to check density distribution plots for gene and exon expression data generated by expression console.

Usage
ocPlotHist()

Author(s)
Raffaele A Calogero

See Also
ocPlotPCA

ocPlotPCA
Gene/Exon level density plots

Description
This function runs a modified version of the plotPCA of the affycoretools to be used to check density distribution plots for gene and exon expression data.

Usage
ocPlotPCA()
Author(s)

Raffaele A Calogero

See Also

ocPlotHist

oneChannelGUI

Starting oneChannelGUI package

Description

Starting oneChannelGUI package. oneChannelGUI contains a set of functions extending the capabilities of affylmGUI package.

Usage

goseqInterface()
DESeqInterface()
EDAwithinLane()
EDAbetweenLane()
EDAbboxplot()
EDAbiasPlot()
AboutextendedaffylmGUI()
AboutaffylmGUI()
oneChannelGUI()
oneChannelGUIHelp()
maSigProHelp()
siggenesHelp()
oneChannelGUIHelp()
initialize.extAffylmGUI()
OpenExonandTargetsfiles()
GOstatsHelp()
SaveLimmaFile()
NewLimmaFile()
NewLimmaFileNGS()
OpenLimmaFile()
OpenALimmaFile(FileName)
OpenFile(FileName)
OpenLargeFile()
changeMenu()
oneChannelGUI.start()
libraryFilesDir()
whichKindOfArray()
intronicBg()
ExportNormalizedExpressionValues1() ExportNormalizedExpressionValues()
ExportFeatureNames()
SaveAsLimmaFile()
addAnnLib()}
OpenCDFandTargetsfiles()
ComputeLinearModelFit()
GetNormalizationMethod()
NormalizeNow()
ComputeContrasts()
midasFilter()
dabgFilter()
largedatasetNorm()
RankProdHelp()
affyPLMHelp()
geneFilterHelp()
pamrHelp()
pdmclassHelp()
sizelowerHelp()
ssizeHelp()
OpenAGeoFile(FileName)
OpenGeoFile()
OpenGeoFiles()
OpenNgsfiles()
recoverUnfiltered()
delete.ML()
affyPlotMA()
changes()
aptFolder()
javaFolder()
deleteLocalData()
SetED()
chooseEDir()
.annotation(eset)
.myfindOverlaps(x,y)
log2Conversion()
.myrk(x,df)
exportFASTA()
OpenmiRanalyserfiles()
OpenmiRprofiles()
OpenmiRExpress()
OpenmiRShrimp()
OpenmiRBam()
gsper1Download()
meVDownload()
extPckInfo()
startmeV()
tuxedoMenu()
setBinDir()
fastQC()
openPdf()
closePdf()
plotDE.exons()
topTable.genes()
importfusions()
fusionNS()
fusionfilter()
chimeraseq()
tophatrun()
addga()
fusionpeptides()
plotcoverage()
desCounts()
miRNAde()
downloadShrimp()
wrapperMirnaCounts(working.dir, out.dir, org = "hsa", threads = 1, cutadapt.path = "/usr/local/bin/cutadapt", parallel = FALSE, ...)
deseq2Processing()
enrich(group, permutations = permutations)
loadmirnaphi(mirnafile = "mirna_input.txt", mirnacol = "miRNA.id",
assayidcol = "ASSAYID", groupcol = "GROUP",
filterflagcol = "FILTERFLAG", expressioncol = NA,
foldchangecol = NA, pvaluecol = NA)
mirPath()
mirnaTable(mirnaobj, groups = NULL, format = "Tall", Significance = 0.2, na.char = NA, pvalueTypes = c("pvalues", "permpvalues"), maxStringLength = NA, pathwaysNames = hs.path.name)
permuting(perm)
runEnrichment(Composite = TRUE, groups = NULL, permutations = 0)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>FileName</td>
<td>Internal argument not to be set by the user</td>
</tr>
<tr>
<td>eset</td>
<td>Internal argument not to be set by the user</td>
</tr>
<tr>
<td>x</td>
<td>Internal argument not to be set by the user</td>
</tr>
<tr>
<td>y</td>
<td>Internal argument not to be set by the user</td>
</tr>
<tr>
<td>df</td>
<td>Internal argument not to be set by the user</td>
</tr>
<tr>
<td>group</td>
<td>Internal argument not to be set by the user</td>
</tr>
<tr>
<td>working.dir</td>
<td>Internal argument not to be set by the user</td>
</tr>
<tr>
<td>out.dir</td>
<td>Internal argument not to be set by the user</td>
</tr>
<tr>
<td>org</td>
<td>Internal argument not to be set by the user</td>
</tr>
<tr>
<td>threads</td>
<td>Internal argument not to be set by the user</td>
</tr>
<tr>
<td>cutadapt.path</td>
<td>Internal argument not to be set by the user</td>
</tr>
<tr>
<td>parallel</td>
<td>Internal argument not to be set by the user</td>
</tr>
<tr>
<td>mirnafile</td>
<td>Internal argument not to be set by the user</td>
</tr>
<tr>
<td>mirnacol</td>
<td>Internal argument not to be set by the user</td>
</tr>
<tr>
<td>assayidcol</td>
<td>Internal argument not to be set by the user</td>
</tr>
<tr>
<td>groupcol</td>
<td>Internal argument not to be set by the user</td>
</tr>
<tr>
<td>filterflagcol</td>
<td>Internal argument not to be set by the user</td>
</tr>
<tr>
<td>foldchangecol</td>
<td>Internal argument not to be set by the user</td>
</tr>
<tr>
<td>pvaluecol</td>
<td>Internal argument not to be set by the user</td>
</tr>
<tr>
<td>mirnaobj</td>
<td>Internal argument not to be set by the user</td>
</tr>
</tbody>
</table>
### Description

Read BeadStudio expression data file

### Usage

```r
OpenBeadStudioFiles()
```

### Details

Reads an Illumina intensity data file produced by BeadStudio. Using BeadStudio version 'One' the file will have a 'gene profile.csv' extension and using version 'Two' the file will have a .txt extension. See package vignette for more information. Multiple filenames can be specified as a vector and the data are then combined into one output file. This function should only really be used for custom analysis as the beadAnalysis() function provides easier, flexible use.
OpenmRNABam

Author(s)

Derived from readBead by Gareth Elvidge <gareth.elvidge@well.ox.ac.uk>

OpenLargefiles

This function loads large data set made from tab delimited files

Description

The function creates and expressionSet starting from all file containing the expression data in a tab delimited format. This file is loaded together with the description of the clinical parameters present in Target. This function uses a specific configuration of Target column of the affylmGUI target file. To know more about target file see affylmGUI help. Each row of the column named Target, in the affylmGUI target file, describes the clinical parameters. Each clinical parameter is separated from the others by an underscore. The affylmGUI target file will look like:

<table>
<thead>
<tr>
<th>Name</th>
<th>FileName</th>
<th>Target</th>
</tr>
</thead>
<tbody>
<tr>
<td>mC1</td>
<td>M1.CEL</td>
<td>pos_yes_1_NA_0</td>
</tr>
<tr>
<td>mC2</td>
<td>M4.CEL</td>
<td>pos_no_2_NA_0</td>
</tr>
<tr>
<td>mC3</td>
<td>M7.CEL</td>
<td>neg_no_3_pos_0</td>
</tr>
<tr>
<td>mE1</td>
<td>M3.CEL</td>
<td>neg_yes_3_neg_0</td>
</tr>
<tr>
<td>mE2</td>
<td>M6.CEL</td>
<td>neg_no_NA_1_0</td>
</tr>
<tr>
<td>mE3</td>
<td>M9.CEL</td>
<td>neg_yes_3_pos_0</td>
</tr>
<tr>
<td>mI1</td>
<td>M2.CEL</td>
<td>pos_no_2_neg_1</td>
</tr>
<tr>
<td>mI2</td>
<td>M5.CEL</td>
<td>pos_yes_2_pos_1</td>
</tr>
<tr>
<td>mI3</td>
<td>M8.CEL</td>
<td>pos_no_2_pos_1</td>
</tr>
</tbody>
</table>

Usage

OpenLargefiles()

Author(s)

Raffaele A Calogero

OpenmRNABam

This function loads Bam files generated by local mapping with bowtie in oneChannelGUI

Description

The function creates and expressionSet starting from the association of reads mapped by Bowtie using chromosome oriented mapping controlled by oneChannelGUI and associate those mapping to gene-level regions extracted by UCSC browser. IMPORTANT at the present time the expression set is base on gene-level data and not on exon-level data.

Usage

OpenmRNABam()
plierToZero

Author(s)
Raffaele A Calogero

Description
The calculation of log2 of probe set intensity by mean of plier generates a set of intensities very low
this function will set to 0 all the log2 intensities below 1 produced by iter-plier or plier alghoritm

Usage
plierToZero()

plotGO

Plotting parents of a GO term with few mouse click

Description
To know more on the parents of a specific GO term you can use the plotGO function

Usage
plotGO()

Details
A GO term to be investigated for its parents has to be placed in the graphical window.

Author(s)
Raffaele A Calogero

See Also
GOenrichment, extractAffyids
**PlotOptionsv1**  
*A modified version of the function used in affyPLM library*

**Description**
As default the plots are generated on the R GUI to reduce RAM consumption.

**Usage**

```r
PlotOptionsv1()
```

**Author(s)**
Raffaele A Calogero

---

**plotVariantSI**  
*This function plots on UCSC genome browser data derived by variantSI filter*

**Description**
This function plots on UCSC genome browser data derived by variantSI filter on the basis of chromosome annotation.

**Usage**

```r
plotVariantSI()
```

**Author(s)**
Raffaele A Calogero

**See Also**
variantSI, variantExons, makeBED15
qcMDS

This function plots the sample relations based on multidimensional scaling using the plotMDS.dge edgeR function

Description

This function is a variation on the usual multidimensional scaling (or principle coordinate) plot, in that a distance measure particularly appropriate for the digital gene expression (DGE) context is used. The distance between each pair of samples (columns) is the square root of the common dispersion for the top genes which best distinguish that pair of samples. These top genes are selected according to the tagwise dispersion of all the samples.

Usage

qcMDS()

Author(s)

Raffaele A Calogero

rankingConversion

This function transforms intensity data in normalized ranks

Description

This function transforms intensity data in normalized ranks, i.e. high intensity genes will have a value near to 0 as instead low intensity genes a normalized rank near to 1.

Usage

rankingConversion()

Author(s)

Raffaele A Calogero
rankProd

graphical interface to rank product method implemented in RankProd Bioconductor library.

Description

To know more about rank product method see RankProd help.

Usage

rankProd()

Details

The target file for the RankProd implementation contain the origin of the data as a number separated by an under score from the corresponding covariate. If all data are from the same origin the origin definition is not needed. Therefore target will contain only the covariates.

<table>
<thead>
<tr>
<th>Name</th>
<th>FileName</th>
<th>Target</th>
</tr>
</thead>
<tbody>
<tr>
<td>mC1</td>
<td>M1.CEL</td>
<td>CTRL_1</td>
</tr>
<tr>
<td>mC2</td>
<td>M4.CEL</td>
<td>CTRL_1</td>
</tr>
<tr>
<td>mC3</td>
<td>M7.CEL</td>
<td>CTRL_2</td>
</tr>
<tr>
<td>mE1</td>
<td>M3.CEL</td>
<td>CTRL_2</td>
</tr>
<tr>
<td>mE2</td>
<td>M6.CEL</td>
<td>CTRL_2</td>
</tr>
<tr>
<td>mE3</td>
<td>M9.CEL</td>
<td>TRT_1</td>
</tr>
<tr>
<td>mI1</td>
<td>M2.CEL</td>
<td>TRT_1</td>
</tr>
<tr>
<td>mI2</td>
<td>M5.CEL</td>
<td>TRT_2</td>
</tr>
<tr>
<td>mI3</td>
<td>M8.CEL</td>
<td>TRT_2</td>
</tr>
</tbody>
</table>

Author(s)

Raffaele A Calogero

---

rawBoxplotPN

Plotting raw log2 intensities from controls

Description

This function produces a box plot of the log2 raw intensities, extracted directly from CEL files, for positive and negative controls presente in XXXX.control.ps Affymetrix library file. Positive and negative controls are made of housekeeping exon and introns. It gives an idea of signal behaviour before data normalization both in the high and low intensity range

Usage

rawBoxplotPN()

Author(s)

Raffaele A Calogero
rawpCheck

**Raw p-value distribution from limma analysis by a mouse click**

**Description**
This function allow to visualize the histogram of raw p-value distribution generated by limma analysis.

**Usage**
```
rawpCheck()
```

**Details**
The histogram of raw p-value distribution will show if raw p-values are uniform in the non significant range and therefore the BH correction can be applied.

**Note**
BH is the most used method for the correction of type I errors in microarray analysis. However, it has some limitation due to the initial hypotheses: The gene expressions are independent from each other. The raw distribution of p values should be uniform in the non significant range.

**Author(s)**
Raffaele A Calogero

**References**
To know more see limma package help

---

Reads2logos

**Extracting info on the counts associated to a differentially expressed ncRNA gene**

**Description**
This function allow to plot the raw counts associated to samples or experimental groups associated to a ncRNA found differentially expressed. Furthermore, it saves the count matrix for the mapped positions on the differentially expressed ncRNA

**Usage**
```
Reads2logos()
```

**Author(s)**
Raffaele A Calogero
refiningPeaks

This function refines the structure of the genomics peaks, checking for the presence nearby peaks that can be combined.

Description

This function allows the user to define a threshold for peaks merging. If threshold is set to 0 non merging is provided.

Usage

refiningPeaks()

Author(s)

Raffaele A Calogero

reformatGdl

This function reorganizes single NGS data in a matrix to be used for statistical analysis.

Description

This function reorganizes the raw NGS data saved in a GenomeDataList object in a matrix. Matrix rows are peaks containing multiple reads. For peak detection, user has to indicate the amount of extension of each read, e.g. 200 nt in chipseq experiments, and the number of reads due by random events, e.g. 8.

Usage

reformatGdl()

Author(s)

Raffaele A Calogero

refseqDownload

Retrieving Reference Sequences from NCBI ftp

Description

This function retrieves reference sequences from NCBI ftp. RefSeq are used for mapping exon-level probe sets to refseq specific isoforms.

Usage

refseqDownload()

Author(s)

Raffaele A Calogero
**retrieveMirTargets**

This function retrieves putative targets from RmiR.Hs.miRNA given a specific microRNA.

**Usage**

retrieveMirTargets()

**Details**

This interface provides a graphical GUI to RmiR.Hs.miRNA.

**Author(s)**

Raffaele A Calogero

---

**retrievePSRseq**

This function, given a file containing exon-level probesets retrieves Probe selection Regions, PRS, from RRE db

**Description**

This function retrieve from RRE the PSR sequences associated to the exon-level probe sets

**Usage**

retrievePSRseq()

**Author(s)**

Raffaele A Calogero

---

**retrievePSRseq1gid**

This function, given a gene-level probeset id, retrieves Probe selection Regions, PRS, from RRE db

**Description**

This function retrieves from RRE the PSR sequences associated to exon-level probe sets

**Usage**

retrievePSRseq1gid()

**Author(s)**

Raffaele A Calogero
reviqrFilter

Reverse interquantile filtering with a mouse click

Description

This function implements a reverse version of the interquantile filtering proposed by Heydebreck
in 2004 to select low variance genelevel probe set. To be used to remove putative differentially
expressed genes that will make more difficult the detection of alternative splicing events.

Usage

reviqrFilter()

Details

This function can be used in a analysis focused to the detection of alternative splicing events. The
aim of this non specific filtering is to remove the genes that are likely to carry information about the
phenotypes under investigation at gene level. This filtering remove genes that show strong changes
within the experimental points at the gene level.

Author(s)

Raffaele A Calogero

References

Heydebreck et al. Bioconducotor project Papers 2004

See Also

dabgFilter, crosshybFilter

RmiRInterface
graphical interface to RmiR library.

Description

RmiR package is to couple microRNA and gene expression data (coming from the same RNA). In-
put data are obtainable with limma interface (gene-level differential expression) in oneChannelGUI
and baySeq interface (NGS microRNA differential expression).

Usage

RmiRInterface()

Details

This interfase provides a graphical GUI to RmiR package. The output structure is described in
RmiR package.
Author(s)
Raffaele A Calogero

rnfa

Rat microRNA data set

Description
microRNA data set fasta produced using miRbase R norvegicus precursors reformatted to have only miR name as descriptor of the fasta file

Usage
rnfa

Format
Fasta file

runningJetta

graphical interface to MADS/jetta R library.

Description
MADS, which stands for Microarray Analysis of Differential Splicing, is a tool to identify differential alternative splicing by exon array. The principle of MADS is to increase the precision of exon-level and gene-level expression estimates by correcting, as much as possible, noise in observed probe intensities due to background and cross-hybridization. MADS incorporates a series of novel algorithms motivated by the probe-rich design of exon-tiling arrays, such as background correction, iterative probe selection and removal of sequence-specific cross-hybridization to off-target transcripts. MADS was published in RNA, 2008, 14(8): 1470-1479. Junction and Exon array Toolkit for Transcriptome Analysis (JETTA) is compacted version of MADS.

Usage
runningJetta()

Details
Expression indexes are calculated as the order of Background Correction, Normalization and Summarization. In the Summarization step, background corrected and normalized probe intensities of a meta probeset are summarized to expression of the meta probeset. Meta probesets can be defined as gene/transcript clust/exon level.

Background Correction JETTA estimates background signal using background probes and subtracts it from the probe intensity. If the probe intensity is less than the estimated background signal, the background subtracted signal is truncated to 1. Estimation of background signal is based on several models: Median GC: median of background probe signal of the same GC counts MAT: linear model of probe sequence with 80 parameters. see Kapur et al, 2007
Normalization Normalization of JETTA is done for core probes defined in probeset annotation file. If the PSA file is not specified, it considers all probes in the MPS files as core probes. Median scaling: scaling each array so that its median is 100 Quantile: all probes of the same signal quantile have the same signal

Summarization LiWong model: multiplication model of expression and probe effect, see Li and Wong, 2001 Probe selection: select probes based on cross-array correlation of signal. see Xing et al, 2006 Median-polish

Alternative Splicing Detection Detecting alternatively expressed PSR/Exon between two sample groups based on background corrected and normalized probe intensities. It has several criteria to filter out transcript clusts and probes from the analysis. TC expression level: excluding low-expressed transcript clusts TC expression fold change: excluding transcript clusts which have big fold change between two groups Extreme probe signal: excluding probes of which signal is extremely high Cross-hybridized probes: excluding cross hybridized probes, currently pre-calculated results are needed

Author(s)

jseok@stanford.edu

---

**sample.size.evaluation**

The function executes and plots results from ssize and delta function from the ssize package

**Description**

This function represents a visual tool for helping users to understand the trade off between sample size and statistical power. To know more about see ssize help.

**Usage**

`sample.size.evaluation()`

**Details**

Both ssize and delta outputs are calculated using the BH type I error correction instead of the Bonferroni used as default in the ssize package. Furthermore, instead using the control group variance, this implementation uses the common variance described in Wei et al. BMC Genomics. 2004, 5:87

Main assumptions: A microarray experiment is set up to compare gene expressions between one treatment group and one control group. Microarray data has been normalized and transformed so that the data for each gene is sufficiently close to a normal distribution that a standard 2-sample pooled-variance t-test will reliably detect differentially expressed genes.

**Author(s)**

Raffaele A Calogero
**Description**

This function represents a tool for helping users to understand the trade off between sample size and statistical power. To know more about see ssize help.

**Usage**

```r
sample.size.evaluation1()
```

**Details**

see sizepower help

**Author(s)**

Raffaele A Calogero

---

**showDataset**

*Grabbing info about the available expression set*

**Description**

The size of the normalized expression set can change upon filtering. This function show info about the exact size of the data set.

**Usage**

```r
showDataset()
```

**Author(s)**

Raffaele A Calogero
showTopTable

*Modification of the function implemented in affylmGUI to generate a topTable*

**Description**

Modification of the function implemented in affylmGUI to generate a topTable. To know more about topTable see limma package help.

**Usage**

`showTopTable(..., export=FALSE)`

**Arguments**

- `export` defining the possibility to export data
- `...` Arguments to be passed to methods

**Author(s)**

Raffaele A Calogero

---

siggenes

*The function executes SAM analysis implemented in siggenes bioconductor library*

**Description**

To know more about SAM in Bioconductor see siggenes help.

**Usage**

`siggenes()`

**Author(s)**

Raffaele A Calogero
**simFilter**

This function allows filtering on the basis of the average splice index mean or min difference between two groups.

**Description**

Filtering out gene/exon level probe sets associated to average splice index mean or min difference between two groups lower than user defined value.

**Usage**

```r
simFilter()
```

**Note**

This function needs the presence of Splice Index.

**Author(s)**

Raffaele A Calogero

**See Also**

`simFilter`

---

**spliceIndex**

This function converts the exon intensity data into a slice index.

**Description**

Exons intensities are divided for the expression of the corresponding gene, as described by Clark et al. Science 2002 May 3;296(5569):907-10.

**Usage**

```r
spliceIndex()
```

**Details**

The function is not yet optimized, therefore it could take quite a long time to compute `spliceIndex` if more than 1000 genes are used.

**Author(s)**

Raffaele A Calogero

**See Also**

`inspecting.splice.index`
standAloneAddingAnnotation

Attach to a data frame containing gene-level data derived from Affymetrix exon arrays the annotations derived by netaffx

Description
Standalone oneChannelGUI function attaches gene-level annotation to a data frame.

Usage
standAloneAddingAnnotation(annotationdf, df.tobe.annotated, ids.column)

Arguments
annotationdf An annotation data frame generated with standAloneBuildingLocalAnnotation function
df.tobe.annotated A data frame containing a gene-level data of any type. It is mandatory that one of the column contains gene-level ids
ids.column the column of the df.tobe.annotated containing gene-level ids

Value
A data frame.

Author(s)
Raffaele A Calogero

standAloneBuildingLocalAnnotation

Creates a data frame with gene-level annotation data for exon arrays using the netaffx database

Description
Standalone oneChannelGUI function to create gene-level annotation data using netaffx database.

Usage
standAloneBuildingLocalAnnotation(libDirLocation = getwd(), netaffxUser = "myemail@somewhere.org", netaffxUserPw = "yourpassword", whichAnnotation = c("HuEx", "MoEx", "RaEx"))

Arguments
libDirLocation Folder where to save the annotation object
netaffxUser The email registered to Affymetrix netaffx web site
netaffxUserPw The password to access to netaffx
whichAnnotation Which annotation table should be used
**Value**

Location of the annotation data frame.

**Author(s)**

Raffaele A Calogero

---

**targetWidget**

*Widget to create a target file to load .CEL files*

---

**Description**

Widget to create a target file to load .CEL files to be used with NewLimmaFile function.

**Usage**

targetWidget()

**Author(s)**

Raffaele A Calogero

---

**templA**

*Generating a template A to be uploaded in Ingenuity Pathways analysis (IPA)*

---

**Description**

A template A file to be used in Ingenuity can be generated starting from a topTable containing the full array data.

**Usage**

templA()

**Note**

Template A file will contain a column with the gene ID, a column with fold change, a column with true p-value and a column with p-values for discriminating between the set of differentially expressed probe sets and the background. This column is needed to allow IPA to identify the set of enriched functional classes associated to the differentially expressed probe sets.

**Author(s)**

Raffaele A Calogero

**See Also**

IPAlistFilter
Description

This function generates a table of differentially expressed exons.

Usage

topTable.exons()

Details

topTable.exons function uses the results produced by dexExons function to generate a table of differentially expressed exons

Author(s)

Raffaele A Calogero

Description

This function allows the creation of a training set and a test set to be used for classification purposes.

Usage

trainTest()

Details

User will be asked to assign names to the available classification parameters. User will be asked to select the number associated to one of the available classification parameters. The training set will be made, using the selected classification parameter and it will be made of 2/3 of the original data set. The test set will be the remaining 1/3.

Author(s)

Raffaele A Calogero
**tuxedoSetup**

**Description**

To be able to run Tuxedo Suite with oneChannelGUI it is necessary to make soft links for samtools, bowtie, tophat, cufflinks programs

**Usage**

`tuxedoSetup()`

**Details**

samtools need to be installed by the user, as instead bowtie, tophat and cufflinks are installed via oneChannelGUI

**Author(s)**

Raffaele A Calogero

---

**TuxedoSuite**

**Description**

TuxedoSuite contains a set of functions to run tophat, cufflinks, cuffmerge, cuffdiff and to extract gene, geneDiff, isoforms, isoDiff, and CDS, cdsDiff, differentially expressed between two experimental conditions. The present implementation works only on MAC and Linux, due to the absence of a windows implementation of the tuxedo suite. tophat, cufflinks, cuffmerge and cuffdiff create the line command to execute the corresponding program and all programs are run in background, since they might require quite a lot of computation time. One differential expression is obtained geneDiff, isoDiff and cdsDiff can be used to extract respectively differentially expressed genes, isoforms and CDSs

**Usage**

`tophat()`
`createDir()`
`cufflinks()`
`cuffmerge()`
`cuffdiff()`
`geneDiff()`
`isoDiff()`
`cdsDiff()`
`chooseInDir()`
`chooseOutDir()`
Details
tophat, cufflinks, samtools and bowtie need to be installed in the system and their folder installation inserted in the PATH variable. Also the genome of interest, as fasta file and as bowtie indexed files need to be previously installed together with a GTF file containing known annotations for the genome of interest. To run the pipeline a target file is needed. createDir generates folders, with the name extracted from the Name column in the target file. In each of these folders tophat will create the files used by cufflinks, cuffmerge, cuffdiff. cuffdiff will create a folder in which differentially expressed data are saved. Access to those data is done using the cummeRbund package.

Author(s)
Raffaele A Calogero

updateLibs

This function allows to update the present installation of Bioconductor

Description

The function allows the updating of local installation of Bioconductor. It might be quite long depending on the internet connection speed.

Usage
updateLibs()

Author(s)
Raffaele A Calogero

variantExons

This function is used to generate a table containing exon-level probe set data linked to variant exons

Description

Internal oneChannelGUI annotation data linking exon-level probesets to variant exons, i.e. those exons that are specific for a subgroup of the isoforms associated to a specific gene, can be upgraded using this function. Annotation RDA files need to be saved in the data subdir of oneChannelGUI dir. Windows users need to exchange the older copies present in Rdata.zip, simply dragging them in the zip file.

Usage
variantExons()

Author(s)
Raffaele A Calogero
**variantSI**  

*This function allows filtering on the basis of variant exons*

**Description**

The function interesects a list of alternative splice exon-level probe sets detected by oneChannelGUI analysis and intersects it to the list of exon-level probe sets associated to variant exons, i.e. these exons that are associated only to a subset of all isoforms associated to a gene. The table of variant Exon is stored in RRE database and is retrieved by the function updating the UCSC tables linking probe set ids with variant exons, located in the General Tools menu. The variantSI function also attaches to the variant exons table the deltaSI associated exon-level probe set.

**Usage**

```r
variantSI()
```

**Note**

This function needs the presence of Splice Index

**Author(s)**

Raffaele A Calogero

**See Also**

variantExons, plotVariantSI

---

**VennDiagram**  

*Venn diagrams using two or three lists*

**Description**

Venn diagrams can be generated using probe sets ids or Entrez gene ids saved in flat files.

**Usage**

```r
VennDiagram()
```

**Details**

The function asks to the user to select two/three files containing probe set ids or EGs separated by carriage return. Each file should contain only one column and no header.

**Author(s)**

Raffaele A Calogero
**wrapNGS**  
*Downloading software for NGS*

**Description**

A function to download all external software needed for NGS data analysis

**Usage**

```r
wrapNGS()
```

**Author(s)**

Raffaele A Calogero

---

**wrapScaffold**  
*A wrapper for makeGCcontent, makeGeneScaffold*

**Description**

A function to wrap the makeGCcontent, makeGeneScaffold functions. Built scaffolds are available as precompiled at sourceforge oneChannelGUIextras project.

**Usage**

```r
wrapScaffold(whichRef)
```

**Arguments**

- `whichRef`  
The genome of interest to be used to create a gene-level scaffold

**Author(s)**

Raffaele A Calogero

**See Also**

makeGCcontent, makeGeneScaffold
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