# Package 'ISoLDE'

May 15, 2024

Type Package

Title Integrative Statistics of alleLe Dependent Expression

Version 1.32.0

Date 2015-10-29

**Description** This package provides ISoLDE a new method for identifying imprinted genes. This method is dedicated to data arising from RNA sequencing technologies. The ISoLDE package implements original statistical methodology described in the publication below.

**Encoding** UTF-8

License GPL (>= 2.0)

**Depends** R (>= 3.3.0),graphics,grDevices,stats,utils

URL www.r-project.org

LazyData no

NeedsCompilation yes

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biocViews ImmunoOncology, GeneExpression, Transcription, GeneSetEnrichment, Genetics, Sequencing, RNASeq, MultipleComparison, SNP, GeneticVariability, Epigenetics, MathematicalBiology, GeneRegulation

git\_url https://git.bioconductor.org/packages/ISoLDE

git\_branch RELEASE\_3\_19

git\_last\_commit a4f5743

git\_last\_commit\_date 2024-04-30

Repository Bioconductor 3.19

**Date/Publication** 2024-05-15

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# Description

This package provides a new method for identifying genes with allelic bias. This method is dedicated to data arising from RNA sequencing technologies. The ISoLDE package implements original statistical methodology described in the publication below.

# **Details**

ISoLDE method has been motivated by several literature limitations in taking into account the data specificities and in making the most of biological replicates. It is based on the definition of a new criterion using robust estimation of the data variability. Variability estimation is of high importance in statistical testing procedures because a difference significance can only be assessed with regards to the intrinsic data variability.

Two methods are available to identify allele specific expression: one is based on bootstrap resampling while the second one uses an empirical threshold. The first one is much more satisfying and is likely to give the most reliable results but it can only be applied to data with at least three biological replicates for each reciprocal cross. While strongly recommending to use at least three replicates, the second method implements a robust solution when only two replicates are available.

# Author(s)

```
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Marine Rohmer <marine.rohmer@mgx.cnrs.fr>,
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```

#### References

Reynès, C. et al. (2016): ISoLDE: a new method for identification of allelic imbalance. Submitted

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| 0.17              | WORLD AND THEFTED AND THE SPECIAL PLANTS           |
|-------------------|----------------------------------------------------|
| filteredASRcounts | NORMALIZED AND FILTERED ALLELE SPECIFIC READ (ASR) |
|                   | COUNTS FOR AN EXPERIENCE WITH MORE THAN TWO REPLI- |
|                   | CATES                                              |

# Description

A data.frame containing the normalized and filtered values of allele specific read counts, for an experiment with more than two replicates. This data.frame is obtained with the filterT function run on the normASRcounts data.frame.

### **Format**

A data.frame.

### **Details**

This data. frame is obtained with the filterT function. Each line represents a feature (e.g. a gene, transcript). Each column represents the number of allele specific sens reads from either the paternal or maternal parent for a given biological replicate, so that you expect to have two columns per biological replicate. Values in the matrix are filtered and normalized (RLE method) ASR counts.

# Source

Extract from Bouschet, T. et al. (2016): In vitro corticogenesis from embryonic stem cells recapitulates the in vivo epigenetic control of imprinted gene expression. *Submitted* Subset of 6062 genes (after filtering).

# References

Bouschet, T. et al. (2016): In vitro corticogenesis from embryonic stem cells recapitulates the in vivo epigenetic control of imprinted gene expression. *Submitted* 

# See Also

```
filterT: a function to filter the ASR counts and produce the filteredASRcounts
data.frame.
isolde_test: example of which uses filteredASRcounts.
```

# Examples

data(filteredASRcounts)

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filterT

# FILTERING GENES BEFORE STATISTICAL ANALYSIS

## **Description**

Filter lowly expressed genes (or transcripts) according to a data driven threshold, before any statistical analysis. This step is not mandatory but strongly recommended.

# Usage

```
filterT(rawASRcounts, normASRcounts, target, tol_filter = 0,
bias)
```

## **Arguments**

rawASRcounts the data. frame containing raw counts (obtained with the readRawInput func-

tion or any data.frame following rawASRcounts format specifications).

Raw count data. frame is required when filtering on raw or on normalized data when the normalized data do not contain 0 counts. (For simplicity purpose, we

call '0 count' any value of zero in a count file).

normASRcounts the data. frame containing normalized counts (obtained with the

readNormInput function or any data.frame following normASRcounts format

specifications).

We strongly recommend to filter on normalized ASR counts.

the data. frame containing the target meta data (obtained with the

readTarget function or any data.frame following target format specifications).

tol\_filter a value between 0 and 100 allowing to introduce tolerance rate into filtering step:

if tol\_filter = 25 all genes having less than 25% of their counts from at least one parental (or strain) origin below the threshold are selected (the default value 0 means all raw counts from at least one parental (or strain) origin must be above

threshold, 100 means that no filtering is applied).

bias The kind of allele expression bias you want to study. It must be one of "parental"

or "strain".

### **Details**

Filtering in statistical analysis is recommended to avoid considering genes (or transcript) without enough information, and thus to avoid a too strong effect of multiple test correction.

The aim of our filtering method is to eliminate from analysis not enough quantified genes, that is genes having mostly counts of 0 or near 0 for each replicate in at least one condition (parent, strain). In this purpose, the filterT function searches for the distribution of counts of a gene in a condition when most of read counts are 0 for this condition. This distribution allows to define a threshold. Hence, genes having less counts than this threshold are eliminated.

The filtering step is not mandatory but strongly recommended.

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### Value

A list of two data. frame:

filteredASRcounts

This data.frame contains ASR counts that have successfully passed the filtering step.

removedASRcounts

This data. frame contains ASR counts that have been removed by the filtering step.

Each line represents a feature (e.g. a gene, transcript). Each column represents the number of allele-specific sens reads from either the paternal or maternal parent for a given biological replicate, so that you expect to have two columns per biological replicate.

#### Note

filterT output on normalized data is the typical input for isolde\_test.

### Note

A minimal filtering step will always be performed while applying the isolde\_test function. It consists of eliminating all genes not satisfying these two conditions:

- At least one of the two medians (of paternal or maternal ASR counts) is different from 0;
- There is at least one ASR count (different from 0) in each cross.

# Author(s)

```
Marine Rohmer <marine.rohmer@mgx.cnrs.fr>,
Christelle Reynès <christelle.reynes@igf.cnrs.fr>
```

## References

Reynès, C. et al. (2016): ISoLDE: a new method for identification of allelic imbalance. Submitted

# **Examples**

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| isolde_test | Statistical analysis of Allele specific read (ASR) counts |
|-------------|-----------------------------------------------------------|
|             |                                                           |

# Description

The main function of the ISoLDE package. Performs statistical test to identify genes with allelic bias and produces both graphical and textual outputs.

# Usage

# Arguments

| bias        | The kind of bias you want to study. It must be one of "parental" or "strain".                                                                                                                                                                                                                                                                                                                                                                                                                     |
|-------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| method      | or "threshold". Default behaviour is to adapt to the number of replicates: when at least three biological replicates for each reciprocal cross are available the bootstrap resampling method is used, else the threshold method is applied. It is possible to force isolde_test to use the threshold method even when more than three replicates are available. In this case method must be set to "threshold". It is *not possible* to force a bootstrap method with less than three replicates. |
| asr_counts  | the data.frame containing the ASR counts to be tested. These data should be normalized and filtered (see the filterT function), although the function can run with non-normalized and non-filtered data (not recommended).                                                                                                                                                                                                                                                                        |
| target      | the target data.frame (obtained by the readTarget function).                                                                                                                                                                                                                                                                                                                                                                                                                                      |
| nboot       | specifies how many resampling steps to do for the bootstrap method. This option is not considered if "threshold" value is set for method. Low values of nboot leads to less relevent results (default to 5000).                                                                                                                                                                                                                                                                                   |
| pcore       | a value between 0 and 100 (default to 75) which specifies the proportion of cores (in percent) to be used for the bootstrap method.                                                                                                                                                                                                                                                                                                                                                               |
| graph       | if TRUE (default) graphical outputs are produced (both on device and file).                                                                                                                                                                                                                                                                                                                                                                                                                       |
| ext         | specifies the extension of the graphical file output (does not work if graph = FALSE). It must be one of "pdf" (default), "png" or "eps".                                                                                                                                                                                                                                                                                                                                                         |
| text        | if TRUE (default) textual output files are produced.                                                                                                                                                                                                                                                                                                                                                                                                                                              |
| split_files | if text = TRUE, specifies if your want to have all genes in one same output file (FALSE, default) or four separate files according to the result: ASE, biallelic, undetermined or filtered (TRUE).                                                                                                                                                                                                                                                                                                |
| prefix      | specifies the prefix for all output file names (default to "ISoLDE_result").                                                                                                                                                                                                                                                                                                                                                                                                                      |
| outdir      | specifies the path where to write the output file(s) (default to current directory).                                                                                                                                                                                                                                                                                                                                                                                                              |
|             |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   |

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### **Details**

Before using this function, your data should be normalized and filtered (see the filterT function for filtering) although the function can run with non-normalized and/or non-filtered data.

The method depends on your minimum number of replicates for each reciprocal cross.

If only one replicate is found, the test can not be achieved and exits.

method="default": If more than two replicates per cross, the method takes advantage of having enough information by using bootstrap resampling to identify genes with allelic bias.

If only two replicates are found in at least one cross, there is too few information to obtain reliable distributions from resampling. Genes with allelic bias are identified thanks to empirically defined thresholds.

method="threshold": The empirical method will be processed instead of the bootstrap one, even if more than two replicates per cross are found.

Note that in differential RNA-seq analysis, at least three replicates are strongly recommended, as variability estimation quality is a key factor in statistical analysis.

More details in Reynès, C. et al. (2016) ISoLDE: a new method for identification of allelic imbalance. *Submitted* 

# Value

listASE

a data. frame with one row per gene (or transcript) identified as having an allelic bias and five columns:

- "names" contains gene (or transcript) names such as asr\_counts row names,
- "criterion" contains the criterion value (see vignette or Reynès et al. (2016)),
- "diff\_prop" the criterion numerator which contains the difference between proportions of either parents or strain origins,
- "variability" the criterion denominator which quantifies the gene (or transcript) variability between replicates,
- "origin" specifies the bias direction either "P" or "M" for parental bias or one of specified strain names for strain bias.

listBA

a data.frame with one row per gene (or transcript) identified as biallelically expressed and four columns corresponding to the first four ones in listASE.

listUN

- a data.frame with one row per gene (or transcript) with undetermined status and six columns. The first five columns are the same as listASE, the last one may take three values:
- "FLAG\_consistency" for genes no statistical evidence of neither bias nor biallelic expression but whose parental or strain bias is always in the same direction across replicates,
- "FLAG\_significance" for genes with statistical evidence of bias but with dis-

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crepancies in bias direction across replicates,

- "NO\_FLAG" for other undetermined genes.

listFILT a data.frame containing names of genes that have failed the minimal filtering

step and thus that have not been considered during the statistical test.

ASE, BA and UN lists are sorted according to their criterion value.

### Note

The bootstrap resampling step is performed many times (default to 5000). Hence, the function may run for a long time if performing the bootstrap method (until several minutes).

### Note

A minimal filtering step will always be performed while applying the isolde\_test function. It consists of eliminating all genes not satisfying these two conditions:

- At least one of the two medians (of paternal or maternal ASR counts) is different from 0;
- There is at least one ASR count (different from 0) in each cross.

# Author(s)

```
Christelle Reynès <christelle.reynes@igf.cnrs.fr>,
Marine Rohmer <marine.rohmer@mgx.cnrs.fr>
```

#### References

Reynès, C. et al. (2016): ISoLDE: a new method for identification of allelic imbalance. Submitted

# **Examples**

```
# Loading all required data.frames
data(filteredASRcounts)
data(target)
# Statistical analysis (forcing the threshold option)
isolde_res <- isolde_test(bias = "parental", method = "threshold",
asr_counts = filteredASRcounts, target = target, ext = "pdf",
prefix = "ISoLDE_test")</pre>
```

normASRcounts

NORMALIZED ALLELE SPECIFIC READ (ASR) COUNTS FOR AN EXPERIENCE WITH MORE THAN TWO REPLICATES

# **Description**

normASRcounts\_file.txt: A tab-delimited text file containing the normalized values of ASR counts for an experiment with more than two replicates.

normASRcounts.rda: the normASRcounts\_file.txt loaded into a data.frame by the readNormInput function.

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### **Format**

normASRcounts\_file.txt: A tab-delimited file. normASRcounts.rda: A data.frame.

### **Details**

Each line represents a feature (e.g. a gene or a transcript). Each column represents the number of allele-specific sens reads from either

the paternal or maternal parent for a given biological replicate, so that

you expect to have two columns per biological replicate.

Values in the matrix are normalized (RLE method) ASR counts.

In case of double input, columns must be in the same order in both raw and normalized ASR counts files.

The normASRcounts\_file.txt file should be read and checked by the readNormInput function.

### Note

A minimum of two biological replicates per cross is mandatory, however, we strongly recommend to use more than two replicates per cross. This enables a better estimation of variability and to use the bootstrap method to perform the statistical test (see the isolde\_test function).

### Source

Extract from Bouschet, T. et al. (2016): In vitro corticogenesis from embryonic stem cells recapitulates the in vivo epigenetic control of imprinted gene expression. *Submitted* Subset of 6062 genes (after filtering).

### References

Bouschet, T. et al. (2016): In vitro corticogenesis from embryonic stem cells recapitulates the in vivo epigenetic control of imprinted gene expression. *Submitted* 

#### See Also

readNormInput example of which uses the normASRcounts file.

### **Examples**

```
# normASRcounts_file.txt
normfile <- system.file("extdata", "normASRcounts_file.txt",
package = "ISoLDE")
normASRcounts <- readNormInput(norm_file = normfile, del = "tab",
rownames = TRUE, colnames = TRUE)
# normASRcounts.rda
data(normASRcounts)</pre>
```

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rawASRcounts

RAW ALLELE SPECIFIC READ (ASR) COUNTS FOR AN EXPERI-ENCE WITH MORE THAN TWO REPLICATES

# **Description**

rawASRcounts\_file.txt: A tab-delimited text file containing the raw values of ASR counts for an experiment with more than two replicates.

rawASRcounts.rda: the rawASRcounts\_file.txt loaded into a data.frame by the readRawInput function.

#### **Format**

rawASRcounts\_file.txt: A tab-delimited file.

rawASRcounts.rda: A data.frame.

### **Details**

Each line represents a feature (e.g. a gene or a transcript).

Each column represents the number of allele-specific sens reads from either

the paternal or maternal parent for a given biological replicate, so that

you expect to have two columns per biological replicate.

Values in the matrix are raw allele-specific read counts.

In case of double input, columns must be in the same order in both raw and normalized ASR counts files.

The rawASRcounts\_file.txt file should be read and checked by the readRawInput function.

# Note

A minimum of two biological replicates per cross is mandatory, however, we strongly recommend to use more than two replicates per cross. This enables a better estimation of variability and to use the bootstrap method to perform the statistical test (see the isolde\_test function).

# Source

Extract from Bouschet, T. et al. (2016): In vitro corticogenesis from embryonic stem cells recapitulates the in vivo epigenetic control of imprinted gene expression. *Submitted* Subset of 6062 genes (after filtering).

# References

Bouschet, T. et al. (2016): In vitro corticogenesis from embryonic stem cells recapitulates the in vivo epigenetic control of imprinted gene expression. *Submitted* 

### See Also

readRawInput example of which uses the rawASRcounts file.

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# **Examples**

```
# rawASRcounts_file.txt
rawfile <- system.file("extdata", "rawASRcounts_file.txt",
package = "ISoLDE")
rawASRcounts <- readRawInput(raw_file = rawfile, del = "tab",
colnames = TRUE, rownames = TRUE)
# rawASRcounts.rda
data(rawASRcounts)</pre>
```

readNormInput

READ THE NORMALIZED DATA FILE

# **Description**

Checks and loads into a data. frame the input file containing normalized allele-specific read (ASR) counts so that it can be input into filterT and isolde\_test.

# Usage

```
readNormInput(norm_file, del = "\t", rownames = TRUE, colnames =
TRUE, dec = ".")
```

# **Arguments**

| norm_file | A character-delimited input file containing normalized counts such as described in normASRcounts_file.                                                                                                                     |
|-----------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| del       | Specifies the delimiter for the input file, usually a semi-colon ";", a coma "," or a tabulation "\t". (default : "\t"). Note : None of your data values must contain this delimiter (be specially careful in gene names). |
| rownames  | Specifies if the file contains some row names to consider. Possible values: TRUE or FALSE (default: TRUE).                                                                                                                 |
| colnames  | Specifies if the file contains some column names to consider. Possible values: TRUE or FALSE (default: TRUE).                                                                                                              |
| dec       | Specifies the character used in the file for decimal mark (default: ".").                                                                                                                                                  |

# Value

A data. frame containing normalized ASR counts from your input file.

# Author(s)

```
Marine Rohmer <marine.rohmer@mgx.cnrs.fr>,
Christelle Reynès <christelle.reynes@igf.cnrs.fr>
```

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# See Also

```
normASRcounts_file.txt: the normalized ASR count file on which to run the readNormInput function.
readRawInput: a similar function for raw (non-normalized) ASR count files.
```

# **Examples**

```
# character-delimited input file containing normalized ASR counts
normfile <- system.file("extdata", "normASRcounts_file.txt",
package = "ISoLDE")
# loading it into a data.frame using the readNormInput function
nbreadnorm <- readNormInput(norm_file = normfile, del = "tab",
rownames = TRUE, colnames = TRUE, dec = ".")</pre>
```

 ${\tt readRawInput}$ 

READ THE RAW DATA FILE

# **Description**

Checks and loads into a data. frame the input file containing raw allele specific read (ASR) counts so that it can be input into filterT.

# Usage

```
readRawInput(raw_file, del = "\t", rownames = TRUE, colnames =
    TRUE)
```

# Arguments

| raw_file | A character-delimited input file containing raw ASR counts such as described in rawASRcounts_file.                                                                                                                       |
|----------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| del      | Specifies the delimiter for the input file, usually a semi-colon ";", a coma "," or a tabulation "\t". (default: "\t"). Note: None of your data values must contain this delimiter (be specially careful in gene names). |
| rownames | Specifies if the file contains some row names to consider. Possible values: TRUE or FALSE (default: TRUE).                                                                                                               |
| colnames | Specifies if the file contains some column names to consider. Possible values: TRUE or FALSE (default: TRUE)                                                                                                             |

# **Details**

Raw ASR counts are only required for the filtering step (with the filterT function) in case the normalized data do not contain 0 counts anymore. If you do not want to perform the filtering step or if you still have 0 counts in your normalized file, you do not need to load raw ASR counts. (For simplicity purpose, we call '0 count' any value of zero in a count file).

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# Value

A data. frame containing raw ASR counts from your input file.

# Author(s)

```
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Christelle Reynès <christelle.reynes@igf.cnrs.fr>
```

# See Also

rawASRcounts\_file.txt: the raw ASR count file on which to run the readRawInput function. readNormInput: a similar function for normalized ASR count file.

# **Examples**

```
# character-delimited input file containing raw ASR counts
rawfile <- system.file("extdata", "rawASRcounts_file.txt",
package = "ISoLDE")
# loading it into a data.frame using the readRawInput function
nbread <- readRawInput(raw_file = rawfile, del = "tab",
rownames = TRUE, colnames = TRUE)</pre>
```

readTarget

READ THE TARGET FILE

# Description

Checks and loads into a data. frame your target input file.

# Usage

```
readTarget(target_file, asr_counts, del = "\t")
```

# **Arguments**

| target_file | A character-delimited text input file, containing metadata about ASR counts files (see target_file.txt).                                                                                                                          |
|-------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| asr_counts  | The data.frame containing values of ASR counts (obtained either by the readRawInput or the readNormInput function). It is used to perform checks on compatibility with the target file.                                           |
| del         | Specifies the delimiter for the target input file, usually a semi-colon ";", a coma "," or a tabulation "\t". (default : "\t"). Note : None of your data values must contain this delimiter (be specially careful in gene names). |

# **Details**

See target\_file.txt for more details about the target\_file format.

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### Value

a data. frame containing the target.

#### Author(s)

```
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Christelle Reynès <christelle.reynes@igf.cnrs.fr>
```

### See Also

target\_file.txt: the metadata file on which to run the readTarget function.

# **Examples**

```
# Target input file
targetfile <- system.file("extdata", "target_file.txt",
package = "ISoLDE")
# The data.frame containing ASR counts is also required
data(rawASRcounts)
# Load into a data.frame and check the target file
target <- readTarget(target_file = targetfile,
asr_counts = rawASRcounts, del = "\t")</pre>
```

target

METADATA ABOUT THE ASR COUNT DATA.

# **Description**

target\_file.txt: A tab-delimited file describing your input data (raw and / or normalized allele specific read (ASR) count file(s)).

Each line of the target file corresponds to a column of the rawASRcounts and / or normASRcounts data. frames. Lines of target file MUST be in the same order as the columns in ASR count data. Each line contains four values, separated by a character (e.g. a tabulation): samples, parent, strain and replicate (see the Details section for more information). target.rda: The target\_file.txt file loaded into a data.frame by the link{readTarget} function.

# **Format**

```
target_file.txt: A tab-delimited text file. target.rda: A data.drame.
```

## Details

Details of the three columns:

sample: the biological sample name. A same sample name has to appear twice in the target file: one line for the maternal allele and one line for the paternal allele.

allele: the parental origin of the ASR count. Two possible values: maternal or paternal.

strain: the strain origin of the ASR count. Exactly two different values have to be provided in the

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whole file.

The first line of the target file has to contain these column names in the same order. These metadata are required for both filterT and isolde\_test functions.

Factice example: sample,parent,strain samp1,maternal,str1 samp1,paternal,str2 samp2,maternal,str1 samp2,paternal,str1 samp3,maternal,str1 samp4,maternal,str1 samp4,paternal,str2

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```

# See Also

readTarget is a function to load into a data. frame and check the input target file.

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