Package 'coseq'

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Description Co-expression analysis for expression profiles arising from high-throughput sequencing data. Feature (e.g., gene) profiles are clustered using adapted transformations and mixture models or a K-means algorithm, and model selection criteria (to choose an appropriate number of clusters) are provided.

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coseq-package	•	•		•		•		•		•	•		•		•	•			•		•	•	•						2
clusterEntropy	•	•		•		•		•		•	•		•		•	•			•		•	•	•						4
clusterInertia	•	•		•		•	•	•	•	•	•	 •	•		•	•	•	•	•	•	•	•	•				•		4

coseq-package

30

compareARI	5
compareICL	7
convertLegacyCoseq	8
coseq	9
coseqFullResults	11
coseqResults-class	14
coseqRun	15
fietz	17
kmeansProbaPost	18
logclr	18
matchContTable	19
NormMixClus	19
NormMixClusK	21
NormMixParam	22
plot	23
summary,coseqResults-method	26
transformRNAseq	28

Index

```
coseq-package
```

Co-expression and co-abundance analysis of high-throughput sequencing data

Description

Co-expression analysis for expression profiles arising from high-throughput sequencing data. Feature (e.g., gene) profiles are clustered using adapted transformations and mixture models or a Kmeans algorithm, and model selection criteria (to choose an appropriate number of clusters) are provided.

Details

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Author(s)

Andrea Rau, Cathy Maugis-Rabusseau, Antoine Godichon-Baggioni

Maintainer: Andrea Rau <andrea.rau@inra.fr>

coseq-package

References

Godichon-Baggioni, A., Maugis-Rabusseau, C. and Rau, A. (2018) Clustering transformed compositional data using K-means, with applications in gene expression and bicycle sharing system data. Journal of Applied Statistics, doi:10.1080/02664763.2018.1454894.

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Rau, A., Maugis-Rabusseau, C., Martin-Magniette, M.-L., Celeux, G. (2015) Co-expression analysis of high-throughput transcriptome sequencing data with Poisson mixture models. Bioinformatics, doi: 10.1093/bioinformatics/btu845.

Rau, A., Celeux, G., Martin-Magniette, M.-L., Maugis-Rabusseau, C. (2011) Clustering high-throughput sequencing data with Poisson mixture models. Inria Research Report 7786. Available at http://hal.inria.fr/inria-00638082.

```
## Simulate toy data, n = 300 observations
set.seed(12345)
countmat <- matrix(runif(300*4, min=0, max=500), nrow=300, ncol=4)</pre>
countmat <- countmat[which(rowSums(countmat) > 0),]
conds <- rep(c("A", "B", "C", "D"), each=2)</pre>
## Run the Normal mixture model for K = 2,3,4
run_arcsin <- coseq(object=countmat, K=2:4, iter=5, transformation="arcsin",</pre>
                    model="Normal")
run_arcsin
## Plot and summarize results
plot(run_arcsin)
summary(run_arcsin)
## Compare ARI values for all models (no plot generated here)
ARI <- compareARI(run_arcsin, plot=FALSE)</pre>
## Compare ICL values for models with arcsin and logit transformations
run_logit <- coseq(object=countmat, K=2:4, iter=5, transformation="logit",</pre>
                   model="Normal")
compareICL(list(run_arcsin, run_logit))
## Use accessor functions to explore results
clusters(run_arcsin)
likelihood(run_arcsin)
nbCluster(run_arcsin)
ICL(run_arcsin)
## Examine transformed counts and profiles used for graphing
tcounts(run_arcsin)
profiles(run_arcsin)
## Run the K-means algorithm for logclr profiles for K = 2, \ldots, 20
run_kmeans <- coseq(object=countmat, K=2:20, transformation="logclr",</pre>
                     model="kmeans")
run_kmeans
```

clusterEntropy

Description

Provides the calculation of per-cluster entropy, equivalent to

$$Entropy(k) = \sum_{i \in C_k} \log(\tau_{ik})$$

where τ_{ik} is the conditional probability of gene *i* belonging to cluster *k* and C_k corresponds to the set of indices of genes attributed to cluster *k*.

Usage

```
clusterEntropy(probaPost)
```

Arguments

probaPost Matrix containing the conditional probabilities of belonging to each cluster for all observations

Value

Entropy per cluster

Author(s)

Cathy Maugis-Rabusseau

Examples

```
## Generate artificial matrix of conditional probabilities for K=5 clusters
tmp <- matrix(runif(100*5), nrow=100, ncol=5)
probaPost <- tmp / rowSums(tmp)
clusterEntropy(probaPost)</pre>
```

clusterInertia Calculation of within-cluster inertia

Description

Provides the calculation of within-cluster inertia, equivalent to

$$Inertia(k) = \sum_{i \in C_k} (y_{ik} - \mu_k)^2$$

where μ_k is the mean of cluster k and C_k corresponds to the set of indices of genes attributed to cluster k.

compareARI

Usage

clusterInertia(profiles, clusters)

Arguments

profiles	Matrix, data.frame, or DataFrame containing the (transformed) profiles used for
	the clustering
clusters	Vector of cluster labels corresponding to the observations in profiles

Value

Within cluster inertia

Author(s)

Andrea Rau, Antoine Godichon-Baggioni

Examples

```
## Simulate toy data, n = 300 observations
set.seed(12345)
countmat <- matrix(runif(300*4, min=0, max=500), nrow=300, ncol=4)
countmat <- countmat[which(rowSums(countmat) > 0),]
conds <- rep(c("A","B","C","D"), each=2)</pre>
```

```
## Run the K-means algorithm for logclr profiles for K = 2,..., 20
run_kmeans <- coseq(object=countmat, K=2:20, transformation="logclr",
model="kmeans")
clusterInertia(profiles=tcounts(run_kmeans), clusters=clusters(run_kmeans))</pre>
```

compareARI	Pairwise comparisons of ARI values among a set of clustering parti-
	tions

Description

Provides the adjusted rand index (ARI) between pairs of clustering paritions.

Usage

```
compareARI(object, ...)
## S4 method for signature 'coseqResults'
compareARI(object, K = NULL, parallel = FALSE,
BPPARAM = bpparam(), plot = TRUE, ...)
## S4 method for signature 'matrix'
compareARI(object, parallel = FALSE,
BPPARAM = bpparam(), plot = TRUE, ...)
## S4 method for signature 'data.frame'
compareARI(object, parallel = FALSE,
BPPARAM = bpparam(), plot = TRUE, ...)
```

Arguments

object	Object of class coseqResults or RangedSummarizedExperiment, or alternatively a $n \ge M$ data.frame or matrix containing the clustering partitions for M different models
	Additional optional parameters for corrplot
К	If NULL, pairwise ARI values will be calculated among every model in object x. Otherwise, K provides a vector of cluster numbers identifying a subset of models in x.
parallel	If FALSE, no parallelization. If TRUE, parallel execution using BiocParallel (see next argument BPPARAM). Note that parallelization is unlikely to be helpful unless the number of observations n in the clustering partitions or the number of models M are very large.
BPPARAM	Optional parameter object passed internally to bplapply when parallel=TRUE. If not specified, the parameters last registered with register will be used.
plot	If TRUE, provide a heatmap using corrplot to visualize the calculated pairwise ARI values.

Value

Matrix of adjusted rand index values calculated between each pair of models.

Author(s)

Andrea Rau

```
## Simulate toy data, n = 300 observations
set.seed(12345)
countmat <- matrix(runif(300*4, min=0, max=500), nrow=300, ncol=4)</pre>
countmat <- countmat[which(rowSums(countmat) > 0),]
conds <- rep(c("A", "B", "C", "D"), each=2)</pre>
## Run the Normal mixture model for K = 2,3,4
run_arcsin <- coseq(object=countmat, K=2:4, iter=5, transformation="arcsin",</pre>
                     model="Normal")
run_arcsin
## Plot and summarize results
plot(run_arcsin)
summary(run_arcsin)
## Compare ARI values for all models (no plot generated here)
ARI <- compareARI(run_arcsin, plot=FALSE)</pre>
\ensuremath{\texttt{#\#}} Compare ICL values for models with arcsin and logit transformations
run_logit <- coseq(object=countmat, K=2:4, iter=5, transformation="logit",</pre>
                    model="Normal")
compareICL(list(run_arcsin, run_logit))
## Use accessor functions to explore results
clusters(run_arcsin)
likelihood(run_arcsin)
```

compareICL

compareICL

Compare corrected ICL values after data transformation

Description

Compare the corrected ICL values after applying the arcsin, logit, and logMedianRef transformations in a coseq analysis

Usage

compareICL(x)

Arguments

Х

A list made up of coseqResults objects. At the current time, this function only supports the comparison of coseqResults objects using model="Normal" and transformation = c("arcsin", "logit", "logMedianRef")

Value

A plot of corrected ICL values for the models included in x (the list of coseqResults objects)

Author(s)

Andrea Rau, Cathy Maugis-Rabusseau

Examples

summary(run_arcsin)

```
## Compare ARI values for all models (no plot generated here)
ARI <- compareARI(run_arcsin, plot=FALSE)</pre>
## Compare ICL values for models with arcsin and logit transformations
run_logit <- coseq(object=countmat, K=2:4, iter=5, transformation="logit",</pre>
                   model="Normal")
compareICL(list(run_arcsin, run_logit))
## Use accessor functions to explore results
clusters(run_arcsin)
likelihood(run_arcsin)
nbCluster(run_arcsin)
ICL(run_arcsin)
## Examine transformed counts and profiles used for graphing
tcounts(run_arcsin)
profiles(run_arcsin)
## Run the K-means algorithm for logclr profiles for K = 2, ..., 20
run_kmeans <- coseq(object=countmat, K=2:20, transformation="logclr",</pre>
                    model="kmeans")
run_kmeans
```

convertLegacyCoseq Convert legacy coseq objects

Description

Convert legacy coseq S3 class objects to coseqResults S4 class objects

Usage

```
convertLegacyCoseq(object, digits = 3)
```

Arguments

object	Object of S3 class coseq arising from a call to previous versions of coseq (< 0.99.1)
digits	integer indicating the number of decimal places (round) to retain in results.

Value

Converted object of S4 class coseqResults compatible with recent versions of coseq (>= 0.99.1)

coseq

Co-expression or co-abudance analysis of high-throughput sequencing data

Description

This is the primary user interface for the coseq package. Generic S4 methods are implemented to perform co-expression or co-abudance analysis of high-throughput sequencing data, with or without data transformation, using K-means or mixture models. The supported classes are matrix, data.frame, and DESeqDataSet. The output of coseq is an S4 object of class coseqResults.

Usage

```
coseq(object, ...)
## S4 method for signature 'matrix'
coseq(object, K, subset = NULL, model = "kmeans",
 transformation = "logclr", normFactors = "TMM",
 meanFilterCutoff = NULL, modelChoice = ifelse(model == "kmeans",
  "DDSE", "ICL"), parallel = FALSE, BPPARAM = bpparam(), ...)
## S4 method for signature 'data.frame'
coseq(object, K, subset = NULL,
 model = "kmeans", transformation = "logclr", normFactors = "TMM",
 meanFilterCutoff = NULL, modelChoice = ifelse(model == "kmeans",
 "DDSE", "ICL"), parallel = FALSE, BPPARAM = bpparam(), ...)
## S4 method for signature 'DESeqDataSet'
coseq(object, K, model = "kmeans",
  transformation = "logclr", normFactors = "TMM",
 meanFilterCutoff = NULL, modelChoice = ifelse(model == "kmeans",
  "DDSE", "ICL"), parallel = FALSE, BPPARAM = bpparam(), ...)
```

Arguments

object	Data to be clustered. May be provided as a y $(n \ge q)$ matrix or data.frame of observed counts for <i>n</i> observations and <i>q</i> variables, or an object of class DESeqDataSet arising from a differential analysis via DESeq2.
	Additional optional parameters.
К	Number of clusters (a single value or a vector of values)
subset	Optional vector providing the indices of a subset of genes that should be used for the co-expression analysis (i.e., row indices of the data matrix y. For the generic function coseq, the results of a previously run differential analysis may be used to select a subset of genes on which to perform the co-expression analysis. If this is desired, subset.index can also be an object of class DESeqResults (from the results function in DESeq2).
model	Type of mixture model to use ("Poisson" or "Normal"), or alternatively "kmeans" for a K-means algorithm

transformation	Transformation type to be used: "voom", "logRPKM" (if geneLength is provided by user), "arcsin", "logit", "logMedianRef", "profile", "logclr", "clr", "alr", "ilr", or "none"
normFactors	The type of estimator to be used to normalize for differences in library size: ("TC" for total count, "UQ" for upper quantile, "Med" for median, "DESeq" for the normalization method in the DESeq package, and "TMM" for the TMM normalization method (Robinson and Oshlack, 2010). Can also be a vector (of length q) containing pre-estimated library size estimates for each sample, or "none" if no normalization is required.

meanFilterCutoff

Value used to filter low mean normalized counts if desired (by default, set to a value of 50)

- modelChoice Criterion used to select the best model. For Gaussian mixture models, "ICL" (integrated completed likelihood criterion) is currently supported. For Poisson mixture models, "ICL", "BIC" (Bayesian information criterion), and a nonasymptotic criterion calibrated via the slope heuristics using either the "DDSE" (data-driven slope estimation) or "Djump" (dimension jump) approaches may be used. See the HTSCluster package documentation for more details about the slope heuristics approaches.
- parallel If FALSE, no parallelization. If TRUE, parallel execution using BiocParallel (see next argument BPPARAM). A note on running in parallel using BiocParallel: it may be advantageous to remove large, unneeded objects from the current R environment before calling the function, as it is possible that R's internal garbage collection will copy these files while running on worker nodes.
- BPPARAMOptional parameter object passed internally to bplapply when parallel=TRUE.If not specified, the parameters last registered with register will be used.

Value

An S4 object of class coseqResults, where conditional probabilities of cluster membership for each gene in each model is stored as a SimpleList of assay data, and the corresponding log likelihood, ICL value, number of clusters, and form of Gaussian model for each model are stored as metadata.

Author(s)

Andrea Rau

coseqFullResults

```
summary(run_arcsin)
## Compare ARI values for all models (no plot generated here)
ARI <- compareARI(run_arcsin, plot=FALSE)</pre>
## Compare ICL values for models with arcsin and logit transformations
run_logit <- coseq(object=countmat, K=2:4, iter=5, transformation="logit",</pre>
                   model="Normal")
compareICL(list(run_arcsin, run_logit))
## Use accessor functions to explore results
clusters(run_arcsin)
likelihood(run_arcsin)
nbCluster(run_arcsin)
ICL(run_arcsin)
## Examine transformed counts and profiles used for graphing
tcounts(run_arcsin)
profiles(run_arcsin)
## Run the K-means algorithm for logclr profiles for K = 2,..., 20
run_kmeans <- coseq(object=countmat, K=2:20, transformation="logclr",</pre>
                    model="kmeans")
run_kmeans
```

coseqFullResults Accessors for the assigned cluster labels of a coseqResults object.

Description

The counts slot holds the count data as a matrix of non-negative integer count values, one row for each observational unit (gene or the like), and one column for each sample.

Usage

```
coseqFullResults(object, ...)
clusters(object, ...)
likelihood(object, ...)
nbCluster(object, ...)
proba(object, ...)
ICL(object, ...)
profiles(object, ...)
tcounts(object, ...)
transformationType(object, ...)
```

coseqFullResults

```
model(object, ...)
DDSEextract(object, ...)
Djumpextract(object, ...)
## S4 method for signature 'coseqResults'
clusters(object, K)
## S4 method for signature 'RangedSummarizedExperiment'
clusters(object, ...)
## S4 method for signature 'matrix'
clusters(object, ...)
## S4 method for signature 'data.frame'
clusters(object, ...)
## S4 method for signature 'MixmodCluster'
likelihood(object)
## S4 method for signature 'RangedSummarizedExperiment'
likelihood(object)
## S4 method for signature 'coseqResults'
likelihood(object)
## S4 method for signature '`NULL`'
likelihood(object)
## S4 method for signature 'MixmodCluster'
nbCluster(object)
## S4 method for signature 'RangedSummarizedExperiment'
nbCluster(object)
## S4 method for signature 'coseqResults'
nbCluster(object)
## S4 method for signature '`NULL`'
nbCluster(object)
## S4 method for signature 'RangedSummarizedExperiment'
ICL(object)
## S4 method for signature 'MixmodCluster'
ICL(object)
## S4 method for signature 'coseqResults'
ICL(object)
## S4 method for signature '`NULL`'
```

coseqFullResults

ICL(object)

S4 method for signature 'coseqResults'
profiles(object)

S4 method for signature 'coseqResults'
tcounts(object)

S4 method for signature 'coseqResults'
transformationType(object)

S4 method for signature 'coseqResults'
model(object)

S4 method for signature 'coseqResults'
coseqFullResults(object)

S4 method for signature 'coseqResults'
show(object)

S4 method for signature 'MixmodCluster'
proba(object)

S4 method for signature 'Capushe'
DDSEextract(object)

```
## S4 method for signature 'Capushe'
Djumpextract(object)
```

Arguments

object	a coseqResults, RangedSummarizedExperiment, or MixmodCluster object.
	Additional optional parameters
К	numeric indicating the model to be used (if NULL of missing, the model chosen by ICL is used by default)

Value

Output varies depending on the method. clusters returns a vector of cluster labels for each gene for the desired model.

Author(s)

Andrea Rau

```
## Simulate toy data, n = 300 observations
set.seed(12345)
countmat <- matrix(runif(300*4, min=0, max=500), nrow=300, ncol=4)
countmat <- countmat[which(rowSums(countmat) > 0),]
conds <- rep(c("A","B","C","D"), each=2)</pre>
```

```
## Run the Normal mixture model for K = 2,3,4
run_arcsin <- coseq(object=countmat, K=2:4, iter=5, transformation="arcsin",</pre>
                    model="Normal")
run_arcsin
## Plot and summarize results
plot(run_arcsin)
summary(run_arcsin)
## Compare ARI values for all models (no plot generated here)
ARI <- compareARI(run_arcsin, plot=FALSE)</pre>
## Compare ICL values for models with arcsin and logit transformations
run_logit <- coseq(object=countmat, K=2:4, iter=5, transformation="logit",</pre>
                   model="Normal")
compareICL(list(run_arcsin, run_logit))
## Use accessor functions to explore results
clusters(run_arcsin)
likelihood(run_arcsin)
nbCluster(run_arcsin)
ICL(run_arcsin)
## Examine transformed counts and profiles used for graphing
tcounts(run_arcsin)
profiles(run_arcsin)
## Run the K-means algorithm for logclr profiles for K = 2,..., 20
run_kmeans <- coseq(object=countmat, K=2:20, transformation="logclr",</pre>
                    model="kmeans")
run_kmeans
```

coseqResults-class coseqResults object and constructor

Description

coseqResults is a subclass of RangedSummarizedExperiment, used to store the co-expression results as well as some additional information useful for plotting (tcounts, y_profiles) and meta-information about the co-expression analysis (transformation, normFactors).

Usage

```
coseqResults(SummarizedExperiment, allResults, model = NULL,
transformation = NULL, tcounts = NULL, y_profiles = NULL,
normFactors = NULL)
```

Arguments

SummarizedExperiment a RangedSummarizedExperiment of coseq results allResults List of conditional probabilities of cluster membership for each gene, in all models fit

coseqRun

model	"Normal" or "Poisson", the mixture model used for co-expression
transformation	Transformation applied to counts to obtain tcounts
tcounts	Transformed counts used for mixture model fitting
y_profiles	y profiles used for coseq plotting
normFactors	Scaling factors used for normalization

Details

This constructor function would not typically be used by "end users". This simple class extends the RangedSummarizedExperiment class of the SummarizedExperiment package to allow other packages to write methods for results objects from the coseq package. It is used by coseqRun to wrap up the results table.

Value

a coseqResults object

coseqRun

Co-expression analysis

Description

Function for primary code to perform co-expression analysis, with or without data transformation, using mixture models. The output of coseqRun is an S4 object of class coseqResults.

Usage

```
coseqRun(y, K, conds = NULL, normFactors = "TMM", model = "kmeans",
  transformation = "logclr", subset = NULL, meanFilterCutoff = 50,
  modelChoice = ifelse(model == "kmeans", "DDSE", "ICL"),
  parallel = FALSE, BPPARAM = bpparam(), ...)
```

Arguments

У	$(n \ge q)$ matrix of observed counts for <i>n</i> observations (genes) and <i>q</i> variables (samples). In nearly all cases, $n > q$.
К	Number of clusters (a single value or a vector of values)
conds	Vector of length q defining the condition (treatment group) for each variable (column) in y
normFactors	The type of estimator to be used to normalize for differences in library size: ("TC" for total count, "UQ" for upper quantile, "Med" for median, "DESeq" for the normalization method in the DESeq package, and "TMM" for the TMM normalization method (Robinson and Oshlack, 2010). Can also be a vector (of length q) containing pre-estimated library size estimates for each sample, or "none" if no normalization is required.
model	Type of mixture model to use ("Poisson" or "Normal"), or alternatively "kmeans" for a K-means algorithm

transformation	Transformation type to be used: "voom", "logRPKM" (if geneLength is provided by user), "arcsin", "logit", "logMedianRef", "profile", "logclr", "clr", "alr", "ilr", or "none"
subset	Optional vector providing the indices of a subset of genes that should be used for the co-expression analysis (i.e., row indices of the data matrix y. For the generic function coseq, the results of a previously run differential analysis may be used to select a subset of genes on which to perform the co-expression analysis. If this is desired, subset.index can also be an object of class DESeqResults (from the results function in DESeq2).
meanFilterCutof	f
	Value used to filter low mean normalized counts if desired (by default, set to a value of 50)
modelChoice	Criterion used to select the best model. For Gaussian mixture models, "ICL" (integrated completed likelihood criterion) is currently supported. For Poisson mixture models, "ICL", "BIC" (Bayesian information criterion), and a non-asymptotic criterion calibrated via the slope heuristics using either the "DDSE" (data-driven slope estimation) or "Djump" (dimension jump) approaches may be used. See the HTSCluster package documentation for more details about the slope heuristics approaches.
parallel	If FALSE, no parallelization. If TRUE, parallel execution using BiocParallel (see next argument BPPARAM). A note on running in parallel using BiocParallel: it may be advantageous to remove large, unneeded objects from the current R environment before calling the function, as it is possible that R's internal garbage collection will copy these files while running on worker nodes.
BPPARAM	Optional parameter object passed internally to bplapply when parallel=TRUE. If not specified, the parameters last registered with register will be used.
	Additional optional parameters.

Value

An S4 object of class coseqResults whose assays contain a SimpleList object, where each element in the list corresponds to the conditional probabilities of cluster membership for each gene in each model. Meta data (accessible via metatdata include the model used (either Normal or Poisson), the transformation used on the data, the transformed data using to estimate model (tcounts), the normalized profiles for use in plotting (y_profiles), and the normalization factors used in the analysis (normFactors).

Author(s)

Andrea Rau

```
## Simulate toy data, n = 300 observations
set.seed(12345)
countmat <- matrix(runif(300*4, min=0, max=500), nrow=300, ncol=4)
countmat <- countmat[which(rowSums(countmat) > 0),]
conds <- rep(c("A", "B", "C", "D"), each=2)
## Run the K-means for K = 2,3,4 with logCLR transformation
## The following are equivalent:
run <- coseqRun(y=countmat, K=2:15)</pre>
```

fietz

```
run <- coseq(object=countmat, K=2:15, transformation="logclr", model="kmeans")
## Run the Normal mixture model for K = 2,3,4 with arcsine transformation
## The following are equivalent:
run <- coseqRun(y=countmat, K=2:4, iter=5, transformation="arcsin", model="Normal")
run <- coseq(object=countmat, K=2:4, iter=5, transformation="arcsin", model="Normal")</pre>
```

fietz

RNA-seq data from the mouse neocortex in Fietz et al. (2012)

Description

This dataset represents RNA-seq data from mouse neocortex RNA-seq data in five embryonic (day 14.5) mice by analyzing the transcriptome of three regions: the ventricular zone (VZ), subventricular zone (SVZ) and cortical place (CP).

Usage

data(fietz)

Format

An ExpressionSet named fietz.eset containing the phenotype data and expression data for the Fietz et al. (2012) experiment. Phenotype data may be accessed using the pData function, and expression data may be accessed using the exprs function.

Value

Object of class 'ExpressionSet'. Matrix of counts can be accessed after loading the 'Biobase' package and calling exprs(fietz)).

Source

Digital Expression Explorer (http://dee.bakeridi.edu.au/).

References

https://perso.math.univ-toulouse.fr/maugis/mixstatseq/packages

Fietz, S. A., et al. (2012). Transcriptomes of germinal zones of human and mouse fetal neocortex suggest a role of extracellular matrix in progenitor self-renewal. Proceedings of the National Academy of Sciences, 109(29):11836-11841. kmeansProbaPost

Description

Calculate conditional probabilities of cluster membership for K-means clustering

Usage

kmeansProbaPost(clusters, tcounts)

Arguments

clusters	Cluster labels arising from K-means clustering
tcounts	Transformed counts clustered using K-means

Value

Conditional probabilities of cluster membership for each observation in each cluster

Examples

```
logclr
```

Calculate the Log Centered Log Ratio (logCLR) transformation

Description

Calculate the Log Centered Log Ratio (logCLR) transformation

Usage

```
logclr(profiles)
```

Arguments

profiles Matrix of profiles. Note that the presence of 0 values causes an error message to be produced.

Value

logCLR-transformed profiles

matchContTable

Description

Permute the columns of a contingency table comparing two clusterings to load the diagonal as much as possible.

Usage

matchContTable(table_1, table_2)

Arguments

table_1	Partition from a first data clustering
table_2	Partition from a second data clustering

Value

Permuted contingency table

Examples

```
## Generate arbitrary labels from two separate clustering results
labels_1 <- sample(1:10, 1000, replace=TRUE) ## K=10 clusters
labels_2 <- sample(1:8, 1000, replace=TRUE) ## K=8 clusters
matchContTable(labels_1, labels_2)
```

NormMixClus

Normal mixture model estimation and selection for a series of cluster numbers

Description

Perform co-expression and co-abudance analysis of high-throughput sequencing data, with or without data transformation, using a Normal mixture models. The output of NormMixClus is an S4 object of class RangedSummarizedExperiment.

Usage

```
NormMixClus(y_profiles, K, subset = NULL, parallel = TRUE,
BPPARAM = bpparam(), ...)
```

Arguments

y_profiles	$(n \ge q)$ matrix of observed profiles for n observations and q variables
К	Number of clusters (a single value or a sequence of values).
subset	Optional vector providing the indices of a subset of genes that should be used for the co-expression analysis (i.e., row indices of the data matrix y.
parallel	If FALSE, no parallelization. If TRUE, parallel execution using BiocParallel (see next argument BPPARAM). A note on running in parallel using BiocParallel: it may be advantageous to remove large, unneeded objects from the current R environment before calling the function, as it is possible that R's internal garbage collection will copy these files while running on worker nodes.
BPPARAM	Optional parameter object passed internally to bplapply when parallel=TRUE. If not specified, the parameters last registered with register will be used.
	Additional optional parameters to be passed to NormMixClusK.

Value

An S4 object of class coseqResults, with conditional probabilities of cluster membership for each gene in each model stored as a list of assay data, and corresponding log likelihood, ICL value, number of clusters, and form of Gaussian model for each model stored as metadata.

Author(s)

Andrea Rau, Cathy Maugis-Rabusseau

```
conds <- rep(c("A", "B", "C", "D"), each=2)</pre>
```

```
## Run the Normal mixture model for K = 2,3
## Object of class coseqResults
run <- NormMixClus(y=profiles, K=2:3, iter=5)
run</pre>
```

```
## Run the Normal mixture model for K=2
## Object of class SummarizedExperiment0
run2 <- NormMixClusK(y=profiles, K=2, iter=5)</pre>
```

```
## Summary of results
summary(run)
```

```
## Re-estimate mixture parameters for the model with K=2 clusters
param <- NormMixParam(run, y_profiles=profiles)</pre>
```

NormMixClusK

Description

Perform co-expression and co-abudance analysis of high-throughput sequencing data, with or without data transformation, using a Normal mixture models for single number of clusters K. The output of NormMixClusK is an S4 object of class RangedSummarizedExperiment.

Usage

```
NormMixClusK(y_profiles, K, alg.type = "EM", init.runs = 50,
init.type = "small-em", GaussianModel = "Gaussian_pk_Lk_Ck",
init.iter = 20, iter = 1000, cutoff = 0.001, verbose = TRUE,
digits = 3)
```

Arguments

y_profiles	y $(n \ge q)$ matrix of observed profiles for n observations and q variables
К	Number of clusters (a single value).
alg.type	Algorithm to be used for parameter estimation: "EM", "CEM", "SEM"
init.runs	Number of runs to be used for the Small-EM strategy, with a default value of 50
init.type	Type of initialization strategy to be used: "small-em" for the Small-EM strat- egy, "random", "CEM", or "SEMMax"
GaussianModel	One of the 28 forms of Gaussian models defined in Rmixmod, by default equal to the "Gaussian_pk_Lk_Ck" (i.e., a general family model with free proportions, free volume, free shape, and free orientation)
init.iter	Number of iterations to be used within each run for the Small-EM strategry, with a default value of 20
iter	Maximum number of iterations to be run for the chosen algorithm
cutoff	Cutoff to declare algorithm convergence
verbose	If TRUE, verbose output is created
digits	Integer indicating the number of decimal places to be used for the probaPost output

Value

An S4 object of class RangedSummarizedExperiment, with conditional probabilities of cluster membership for each gene stored as assay data, and log likelihood, ICL value, number of clusters, and form of Gaussian model stored as metadata.

Author(s)

Cathy Maugis-Rabusseau, Andrea Rau

Examples

```
## Simulate toy data, n = 300 observations
set.seed(12345)
countmat <- matrix(runif(300*4, min=0, max=500), nrow=300, ncol=4)</pre>
countmat <- countmat[which(rowSums(countmat) > 0),]
profiles <- transformRNAseq(countmat, norm="none",</pre>
                              transformation="arcsin")$tcounts
conds <- rep(c("A", "B", "C", "D"), each=2)</pre>
## Run the Normal mixture model for K = 2,3
## Object of class coseqResults
run <- NormMixClus(y=profiles, K=2:3, iter=5)</pre>
run
## Run the Normal mixture model for K=2
## Object of class SummarizedExperiment0
run2 <- NormMixClusK(y=profiles, K=2, iter=5)</pre>
## Summary of results
summary(run)
## Re-estimate mixture parameters for the model with K=2 clusters
param <- NormMixParam(run, y_profiles=profiles)</pre>
```

NormMixParam Calc	culate the mean and covariance	for a Normal mixture model
-------------------	--------------------------------	----------------------------

Description

Calculates the mean and covariance parameters for a normal mixture model of the form pK_Lk_Ck

Usage

```
NormMixParam(coseqResults, y_profiles = NULL, K = NULL, digits = 3,
plot = FALSE, ...)
```

Arguments

coseqResults	Object of class coseqResults or RangedSummarizedExperiment (as output from the NormMixClus or NormMixClusK functions)
y_profiles	y $(n \ge q)$ matrix of observed profiles for n observations and q variables, required for ≥ 0 of class RangedSummarizedExperiment
К	The model used for parameter estimation for objects x of class coseq or NormMixClus. When NULL, the model selected by the ICL criterion is used; otherwise, K should designate the number of clusters in the desired model
digits	Integer indicating the number of decimal places to be used for output
plot	If true, produce heatmaps to visualize the estimated per-cluster correlation ma- trices
	Additional optional parameters to pass to corrplot, if desired

plot

Value

pi	Vector of dimension K with the estimated cluster proportions from the Gaussian mixture model, where K is the number of clusters
mu	Matrix of dimension $K \ge d$ containing the estimated mean vector from the Gaussian mixture model, where d is the number of samples in the data y_profiles and K is the number of clusters
Sigma	Array of dimension $d \ge d \ge K$ containing the estimated covariance matrices from the Gaussian mixture model, where d is the number of samples in the data y_profiles and K is the number of clusters
rho	Array of dimension $d \ge d \ge K$ containing the estimated correlation matrices from the Gaussian mixture model, where d is the number of samples in the data y_profiles and K is the number of clusters

Author(s)

Andrea Rau, Cathy Maugis-Rabusseau

Examples

```
## Simulate toy data, n = 300 observations
set.seed(12345)
countmat <- matrix(runif(300*4, min=0, max=500), nrow=300, ncol=4)</pre>
countmat <- countmat[which(rowSums(countmat) > 0),]
profiles <- transformRNAseq(countmat, norm="none",</pre>
                             transformation="arcsin")$tcounts
conds <- rep(c("A", "B", "C", "D"), each=2)</pre>
## Run the Normal mixture model for K = 2,3
## Object of class coseqResults
run <- NormMixClus(y=profiles, K=2:3, iter=5)</pre>
run
## Run the Normal mixture model for K=2
## Object of class SummarizedExperiment0
run2 <- NormMixClusK(y=profiles, K=2, iter=5)</pre>
## Summary of results
summary(run)
## Re-estimate mixture parameters for the model with K=2 clusters
param <- NormMixParam(run, y_profiles=profiles)</pre>
```

plot

Visualize results from coseq clustering

Description

Plot a coseqResults object.

Usage

plot(x, ...)

```
## S4 method for signature 'coseqResults'
plot(x, y_profiles = NULL, K = NULL,
   threshold = 0.8, conds = NULL, average_over_conds = FALSE,
   collapse_reps = "none", graphs = c("logLike", "ICL", "profiles",
    "boxplots", "probapost_boxplots", "probapost_barplots",
    "probapost_histogram"), order = FALSE, profiles_order = NULL,
   n_row = NULL, n_col = NULL, ...)
coseqGlobalPlots(object, graphs = c("logLike", "ICL"), ...)
coseqModelPlots(probaPost, y_profiles, K = NULL, threshold = 0.8,
   conds = NULL, collapse_reps = "none", graphs = c("profiles",
    "boxplots", "probapost_boxplots", "probapost_barplots",
    "probapost_histogram"), order = FALSE, profiles_order = NULL,
    n_row = NULL, n_col = NULL, ...)
```

Arguments

х	An object of class "coseqResults"
	Additional optional plotting arguments (e.g., xlab, ylab, use_sample_names, facet_labels)
y_profiles	y ($n \ge q$) matrix of observed profiles for n observations and q variables to be used for graphing results (optional for logLike, ICL, probapost_boxplots, and probapost_barplots, and by default takes value x to use to use the statement of the statement o
К	If desired, the specific model to use for plotting (or the specific cluster number(s) to use for plotting in the case of coseqModelPlots). If NULL, all clusters will be visualized, and the model chosen by ICL will be plotted
threshold	Threshold used for maximum conditional probability; only observations with maximum conditional probability greater than this threshold are visualized
conds	Condition labels, if desired
average_over_co	onds
	If TRUE, average values of y_profiles within each condition identified by conds for the profiles and boxplots plots. This argument is redundant to collapse_reps = "sum", and collapse_reps should be used instead.
collapse_reps	If "none", display all replicates. If "sum", collapse replicates within each con- dition by summing their profiles If "average", collapse replicates within each condition by averaging their profiles. For highly unbalanced experimental de- signs, using "average" will likely provide more easily interpretable plots.
graphs	Graphs to be produced, one (or more) of the following: "logLike" (log-likelihood plotted versus number of clusters), "ICL" (ICL plotted versus number of clusters), "profiles" (line plots of profiles in each cluster), "boxplots" (boxplots of profiles in each cluster), "probapost_boxplots" (boxplots of maximum conditional probabilities per cluster), "probapost_barplots" (number of observations with a maximum conditional probability greater than threshold per cluster), "probapost_histogram" (histogram of maximum conditional probabilities over all clusters)

order	If TRUE, order clusters in probapost_boxplot by median and probapost_barplot by number of observations with maximum conditional probability greater than threshold
profiles_order	If NULL or FALSE, line plots and boxplots of profiles are plotted sequentially by cluster number (K=1, K=2,). If TRUE, line plots and boxplots of profiles are plotted in an automatically calculated order (according to the Euclidean distance between cluster means) to plot clusters with similar mean profiles next to one another. Otherwise, the user may provide a vector (of length equal to the number of clusters in the given model) providing the desired order of plots.
n_row	Number of rows for plotting layout of line plots and boxplots of profiles. Note that if n_row x n_col is less than the total number of clusters plotted, plots will be divided over multiple pages.
n_col	Number of columns for plotting layout of line plots and boxplots of profiles. Note that if n_row x n_col is less than the total number of clusters plotted, plots will be divided over multiple pages.
object	An object of class "RangedSummarizedExperiment" arising from a call to NormMixClus
probaPost	Matrix or data.frame of dimension $(n \ge K)$ containing the conditional probilities of cluster membership for n genes in K clusters arising from a mixture model

Value

Named list of plots of the coseqResults object.

Author(s)

Andrea Rau, Cathy Maugis-Rabusseau

Examples

run_arcsin

```
## Plot and summarize results
plot(run_arcsin)
summary(run_arcsin)
```

```
## Compare ARI values for all models (no plot generated here)
ARI <- compareARI(run_arcsin, plot=FALSE)</pre>
```

```
## Use accessor functions to explore results
clusters(run_arcsin)
```

 ${\tt summary, coseq Results-method}$

Summarize results from coseq clustering

Description

A function to summarize the clustering results obtained from a Poisson or Gaussian mixture model estimated using coseq. In particular, the function provides the number of clusters selected for the ICL model selection approach (or alternatively, for the capushe non-asymptotic approach if K-means clustering is used), number of genes assigned to each cluster, and if desired the per-gene cluster means.

Usage

S4 method for signature 'coseqResults'
summary(object, y_profiles, digits = 3, ...)

Arguments

object	An object of class "coseqResults"
y_profiles	Data used for clustering if per-cluster means are desired
digits	Integer indicating the number of decimal places to be used for mixture model parameters
	Additional arguments

Details

Provides the following summary of results:

1) Number of clusters and model selection criterion used, if applicable.

2) Number of observations across all clusters with a maximum conditional probability greater than 90 observations) for the selected model.

3) Number of observations per cluster with a maximum conditional probability greater than 90 cluster) for the selected model.

4) If desired, the μ values and π values for the selected model in the case of a Gaussian mixture model.

Value

Summary of the coseqResults object.

Author(s)

Andrea Rau

References

Rau, A. and Maugis-Rabusseau, C. (2017) Transformation and model choice for co-expression analayis of RNA-seq data. Briefings in Bioinformatics, doi: http://dx.doi.org/10.1101/065607.

Godichon-Baggioni, A., Maugis-Rabusseau, C. and Rau, A. (2017) Clustering transformed compositional data using K-means, with applications in gene expression and bicycle sharing system data. arXiv:1704.06150.

See Also

coseq

```
## Simulate toy data, n = 300 observations
set.seed(12345)
countmat <- matrix(runif(300*4, min=0, max=500), nrow=300, ncol=4)</pre>
countmat <- countmat[which(rowSums(countmat) > 0),]
conds <- rep(c("A", "B", "C", "D"), each=2)</pre>
## Run the Normal mixture model for K = 2,3,4
run_arcsin <- coseq(object=countmat, K=2:4, iter=5, transformation="arcsin",</pre>
                    model="Normal")
run_arcsin
## Plot and summarize results
plot(run_arcsin)
summary(run_arcsin)
## Compare ARI values for all models (no plot generated here)
ARI <- compareARI(run_arcsin, plot=FALSE)</pre>
## Compare ICL values for models with arcsin and logit transformations
run_logit <- coseq(object=countmat, K=2:4, iter=5, transformation="logit",</pre>
                   model="Normal")
compareICL(list(run_arcsin, run_logit))
## Use accessor functions to explore results
clusters(run_arcsin)
likelihood(run_arcsin)
nbCluster(run_arcsin)
ICL(run_arcsin)
## Examine transformed counts and profiles used for graphing
tcounts(run_arcsin)
profiles(run_arcsin)
```

```
## Run the K-means algorithm for logclr profiles for K = 2,..., 20
```

transformRNAseq Transform RNA-seq data using common transformations

Description

Application of common transformations for RNA-seq data prior to fitting a normal mixture model

Usage

```
transformRNAseq(y, normFactors = "TMM", transformation = "arcsin",
  geneLength = NA, meanFilterCutoff = NULL, verbose = TRUE)
```

Arguments

У	$(n \ge q)$ matrix or data.frame of observed counts for n observations and q variables	
normFactors	The type of estimator to be used to normalize for differences in library size: "TC" for total count, "DESeq" for the normalization method in the DESeq package, and "TMM" for the TMM normalization method (Robinson and Oshlack, 2010). Can also be a vector (of length q) containing pre-estimated library size estimates for each sample.	
transformation	Transformation type to be used: "arcsin", "logit", "logMedianRef", "profile", "voom", "logRPKM" (if geneLength is provided by user), "logclr", "clr", "alr", "ilr", "none",	
geneLength	Vector of length equal to the number of rows in "y" providing the gene length (bp) for RPKM calculation	
meanFilterCutoff		
	Value used to filter low mean normalized counts	
verbose	If TRUE, include verbose output	

Value

tcounts	Transformed counts
normCounts	Normalized counts
snorm	Per-sample normalization factors divided by mean normalization factor
ellnorm	Per-sample normalization factors

Examples

```
set.seed(12345)
countmat <- matrix(runif(300*4, min=0, max=500), nrow=300, ncol=4)
countmat <- countmat[which(rowSums(countmat) > 0),]
conds <- rep(c("A", "B", "C", "D"), each=2)
## Arcsin transformation, TMM normalization</pre>
```

arcsin <- transformRNAseq(countmat, normFactors="TMM", transformation="arcsin")\$tcounts</pre>

transformRNAseq

Logit transformation, TMM normalization
logit <- transformRNAseq(countmat, normFactors="TMM", transformation="logit")\$tcounts
logCLR transformation, TMM normalization
logclr <- transformRNAseq(countmat, normFactors="TMM", transformation="logclr")\$tcounts</pre>

Index

*Topic cluster coseq-package, 2 *Topic **datasets** fietz, 17 *Topic **methods** coseq, 9 summary, coseqResults-method, 26 *Topic models coseq-package, 2 clusterEntropy, 4 clusterInertia, 4 clusters (coseqFullResults), 11 clusters, coseqResults-method (coseqFullResults), 11 clusters, data.frame-method (coseqFullResults), 11 clusters, matrix-method (coseqFullResults), 11 clusters,RangedSummarizedExperiment-method (coseqFullResults), 11 compareARI, 5 compareARI, coseqResults-method (compareARI), 5 compareARI, data.frame-method (compareARI), 5 compareARI, matrix-method (compareARI), 5 compareARI, RangedSummarizedExperiment-method (compareARI), 5 compareARI-methods (compareARI), 5 compareICL, 7 convertLegacyCoseq, 8 coseq, 9, 27 coseq, data.frame-method (coseq), 9 coseq,DESeqDataSet-method(coseq),9 coseq, matrix-method (coseq), 9 coseq-methods (coseq), 9 coseq-package, 2 coseqFullResults, 11 coseqFullResults, coseqResults-method (coseqFullResults), 11 coseqGlobalPlots (plot), 23 coseqModelPlots(plot), 23 coseqResults (coseqResults-class), 14

coseqResults-class, 14 coseqRun, 15, 15 DDSEextract(coseqFullResults), 11 DDSEextract,Capushe-method (coseqFullResults), 11 Djumpextract(coseqFullResults), 11 Djumpextract,Capushe-method (coseqFullResults), 11

fietz, 17

ICL (coseqFullResults), 11 ICL, coseqResults-method (coseqFullResults), 11 ICL,MixmodCluster-method (coseqFullResults), 11 ICL,mixmodCluster-method (coseqFullResults), 11 ICL,NULL-method (coseqFullResults), 11 ICL,RangedSummarizedExperiment-method (coseqFullResults), 11

 ${\tt kmeansProbaPost, 18}$

INDEX

```
nbCluster,MixmodCluster-method
        (coseqFullResults), 11
nbCluster,NULL-method
        (coseqFullResults), 11
nbCluster,RangedSummarizedExperiment-method
        (coseqFullResults), 11
NormMixClus, 19
NormMixClusK, 20, 21
NormMixParam, 22
```

plot, 23

```
show(coseqFullResults), 11
show,coseqResults-method
        (coseqFullResults), 11
summary(summary,coseqResults-method),
        26
summary,coseqResults-method, 26
summary-methods
        (summary,coseqResults-method),
        26
```

```
tcounts(coseqFullResults), 11
tcounts,coseqResults-method
        (coseqFullResults), 11
transformationType(coseqFullResults),
        11
transformationType,coseqResults-method
        (coseqFullResults), 11
```

```
transformRNAseq, 28
```