Title  Monte Carlo Consensus Clustering

Version  1.2.0

Description  Genome-wide data is used to stratify patients into classes using class discovery algorithms. However, we have observed systematic bias present in current state-of-the-art methods. This arises from not considering reference distributions while selecting the number of classes (K). As a solution, we developed a consensus clustering-based algorithm with a hypothesis testing framework called Monte Carlo consensus clustering (M3C). M3C uses a multi-core enabled Monte Carlo simulation to generate null distributions along the range of K which are used to calculate p values to select its value. P values beyond the limits of the simulation are estimated using a beta distribution. M3C can quantify structural relationships between clusters and uses spectral clustering to deal with non-gaussian and imbalanced structures.

Depends  R (>= 3.4.0)

License  AGPL-3

Encoding  UTF-8

LazyData  true

Imports  ggplot2, Matrix, doSNOW, NMF, RColorBrewer, cluster, parallel, foreach, doParallel, matrixcalc, dendextend, sigclust

Suggests  knitr, rmarkdown

RoxygenNote  6.0.1

biocViews  Clustering, GeneExpression, Transcription, RNASeq, Sequencing

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git_last_commit  3bfc010

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clustersim

clustersim: A cluster simulator for testing clustering algorithms

Description

clustersim: A cluster simulator for testing clustering algorithms

Usage

clustersim(n, n2, r, K, alpha, wobble, redp = NULL, print = FALSE, seed = NULL)

Arguments

n
  Numerical value: The number of samples, it must be square rootable
n2
  Numerical value: The number of features
r
  Numerical value: The radius to define the initial circle (use approx n/100)
K
  Numerical value: How many clusters to simulate
alpha
  Numerical value: How far to pull apart the clusters
wobble
  Numerical value: The degree of noise to add to the sample co ordinates
redp
  Numerical value: The fraction of samples to remove from one cluster
print
  Logical flag: whether to print the PCA into current directory
seed
  Numerical value: fixes the seed if you want to repeat results

Value

A list: containing 1) matrix with simulated data in it

Examples

res <- clustersim(225, 900, 8, 4, 0.75, 0.025, redp = NULL, print = TRUE, seed=123)
**GBM clinical annotation data**

**Description**

This is the clinical annotation data from the GBM dataset, it contains the class of the tumour which is one of: classical, mesenchymal, neural, proneural. It is a data frame with 2 columns and 50 rows.

**Author(s)**

Chris John <chris.r.john86@gmail.com>

**References**


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**M3C: Monte Carlo Consensus Clustering**

**Description**

This function runs M3C, which is a consensus clustering tool with hypothesis testing. The basic idea is to use a multi-core enabled Monte Carlo simulation to drive the creation of a null distribution of stability scores. The Monte Carlo simulation maintains the correlation structure of the input data. Then the null distribution is used to compare the reference scores with the real scores and an empirical p value is calculated for every value of K. We also use the relative cluster stability index as an alternative metric which is just based on a comparison against the reference mean, the advantage being it requires fewer iterations. Small p values are estimated cheaply using a beta distribution that is inferred using parameter estimates from the Monte Carlo simulation.

**Usage**

```r
M3C(mydata, montecarlo = TRUE, cores = 1, iters = 100, maxK = 10, des = NULL, ref_method = c("reverse-pca", "chol"), repsref = 100, repsreal = 100, clusteralg = c("pam", "km", "spectral"), distance = "euclidean", pacx1 = 0.1, pacx2 = 0.9, printres = FALSE, printheatmaps = FALSE, showheatmaps = FALSE, seed = NULL, removeplots = FALSE, dend = FALSE)
```

**Arguments**

- `mydata`: Data frame or matrix: Contains the data, with samples as columns and rows as features.
- `montecarlo`: Logical flag: whether to run the Monte Carlo simulation or not (recommended: TRUE).
- `cores`: Numerical value: how many cores to split the Monte Carlo simulation over.
**mydata**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>iters</td>
<td>Numerical value: how many Monte Carlo iterations to perform (default: 100, recommended: 100-1000)</td>
</tr>
<tr>
<td>maxK</td>
<td>Numerical value: the maximum number of clusters to test for, K (default: 10)</td>
</tr>
<tr>
<td>des</td>
<td>Data frame: contains annotation data for the input data for automatic reordering (optional)</td>
</tr>
<tr>
<td>ref_method</td>
<td>Character string: refers to which reference method to use (recommended: leaving as default)</td>
</tr>
<tr>
<td>repsref</td>
<td>Numerical value: how many reps to use for the Monte Carlo reference data (suggest 100)</td>
</tr>
<tr>
<td>repsreal</td>
<td>Numerical value: how many reps to use for the real data (recommended: 100)</td>
</tr>
<tr>
<td>clusteralg</td>
<td>String: dictates which algorithm to use for M3C (recommended: leaving as default)</td>
</tr>
<tr>
<td>distance</td>
<td>String: dictates which distance metric to use for M3C (recommended: leaving as default)</td>
</tr>
<tr>
<td>pacx1</td>
<td>Numerical value: The 1st x co-ordinate for calculating the pac score from the CDF (default: 0.1)</td>
</tr>
<tr>
<td>pacx2</td>
<td>Numerical value: The 2nd x co-ordinate for calculating the pac score from the CDF (default: 0.9)</td>
</tr>
<tr>
<td>printres</td>
<td>Logical flag: whether to print all results into current directory</td>
</tr>
<tr>
<td>printheatmaps</td>
<td>Logical flag: whether to print all the heatmaps into current directory</td>
</tr>
<tr>
<td>showheatmaps</td>
<td>Logical flag: whether to show the heatmaps on screen (can be slow)</td>
</tr>
<tr>
<td>seed</td>
<td>Numerical value: fixes the seed if you want to repeat results, set the seed to 123 for example here</td>
</tr>
<tr>
<td>removeplots</td>
<td>Logical flag: whether to remove all plots (recommended: leaving as default)</td>
</tr>
<tr>
<td>dend</td>
<td>Logical flag: whether to compute the dendrogram and p values for the optimal K or not</td>
</tr>
</tbody>
</table>

**Value**

A list, containing: 1) the stability results and 2) all the output data (another list) 3) reference stability scores (see vignette for more details on how to easily access)

**Examples**

```r
res <- M3C(mydata, cores=1, iters=100, ref_method = 'reverse-pca', montecarlo = TRUE,printres = FALSE, maxK = 10, showheatmaps = FALSE, repsreal = 100, repsref = 100, printheatmaps = FALSE, seed = 123, des = desx)
```

**mydata**

*GBM expression data*

**Description**

This is the expression data from the GBM dataset. It is a data frame with 50 columns and 1740 rows.

**Author(s)**

Chris John <chris.r.john86@gmail.com>
References


pca: A principal component analysis function

Description

pca: A principal component analysis function

Usage

pca(mydata, K = FALSE, printres = FALSE, labels = FALSE)

Arguments

mydata  Data frame or matrix or M3C results object: if dataframe/matrix should have samples as columns and rows as features
K  Numerical value: if running on the M3C results object, which value was the optimal K?
printres  Logical flag: whether to print the PCA into current directory
labels  Factor: if we want to just display gender for example

Value

A PCA plot object

Examples

PCA <- pca(mydata)
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