## Package: ccml (via r-universe)

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Type Package

Title Consensus Clustering for Different Sample Coverage Data

Version 1.4.0

Description Consensus clustering, also called meta-clustering or cluster ensembles, has been increasingly used in clinical data. Current consensus clustering methods tend to ensemble a number of different clusters from mathematical replicates with similar sample coverage. As the fact of common variety of sample coverage in the real-world data, a new consensus clustering strategy dealing with such biological replicates is required. This is a two-step consensus clustering package, which is used to input multiple predictive labels with different sample coverage (missing labels).

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**Suggests** spelling, testthat (>= 3.0.0)

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## Description

Calculate normalized consensus weight(NCW) matrix based on permutation.

## Usage

```
callNCW(
  title = "",
  label,
  nperm = 10,
  ncore = 1,
  seedn = 100,
  stability = TRUE,
  plot = NULL
)
```

#### Arguments

title	A character value for output directory. Directory is created only if not existed. This title can be an abosulte or relative path.
label	A matrix or data frame of input labels, columns=different clustering results and rows are samples.
nperm	A integer value of the permutation numbers, or nperm=10(default), which means nperm*1000 times of permutation.
ncore	A integer value of cores to use, or ncore=1 (default). It's the input core numbers for the parallel computation in this package parallel.

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seedn	A numerical value to set the start random seed for reproducible results, or seedn=100 (default). For every 1000 iteration, the seed will +1 to get repeat results.
stability	A logical value. Should estimate the stability of normalized consensus weight based on permutation numbers (default stability=TRUE), or not?
plot	character value. NULL(default) - print to screen, 'pdf', 'png', 'pngBMP' for bitmap png, helpful for large datasets, or 'pdf'. Input for randConsensusMatrix.

#### Value

A matrix of normalized consensus weights.

#### **Examples**

```
# load data
data(example_data)
label=example_data

# if plot is not NULL, results will be saved in "result_output" directory
title="result_output"

# run ncw
ncw<-callNCW(title=title,label=label,stability=TRUE,nperm=4,ncore=1)</pre>
```

ccml

A two-step consensus clustering inputing multiple predictive labels with different sample coverages (missing labels)

### Description

A two-step consensus clustering inputing multiple predictive labels with different sample coverages (missing labels)

#### Usage

```
ccml(
  title,
  label,
  output = "rdata",
  nperm = 10,
  ncore = 1,
  seedn = 100,
  stability = TRUE,
  maxK = 15,
  reps = 1000,
  pItem = 0.9,
```

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```
plot = NULL,
  clusterAlg = "spectralClusteringAffinity",
  innerLinkage = "complete",
  ...
)
```

#### **Arguments**

A character value for output directory. Directory is created only if not existed.

This title can be an abosulte or relative path. Input for callNCW, plotCompareCW,

ConsensusClusterPlus::ConsensusClusterPlus, ConsensusClusterPlus::calcICL

label A matrix or data frame of input labels or a character value of input file name,

columns=different clustering results and rows are samples. label could be im-

port as '.rdata', '.rda', or '.csv'. Input for callNCW, plotCompareCW

output A character value for output format, or "rdata" (default) as save to .rdata when

both output and plot are not NULL, others will return to workspace.

nperm A integer value of the permutation numbers, or nperm=10(default), which means

nperm\*1000 times of permutation. Input for callNCW

ncore A integer value of cores to use, or ncore=1 (default). It's the input core numbers

for the parallel computation in this package parallel. Input for callNCW

seedn A numerical value to set the start random seed for reproducible results, or seedn=100

(default). For every 1000 iteration, the seed will +1 to get repeat results. Input

for callNCW, ConsensusClusterPlus::ConsensusClusterPlus

stability A logical value. Should estimate the stability of normalized consensus weight

based on permutation numbers (default stability=TRUE), or not? Input for

callNCW

maxK integer value. maximum cluster number to evaluate. Input for ConsensusClusterPlus::ConsensusClus

for the consensus clustering based on normalized consensus weights.

reps integer value. number of subsamples. Input for ConsensusClusterPlus::ConsensusClusterPlus

pItem numerical value. proportion of items to sample. Input for ConsensusClusterPlus::ConsensusCluster

plot character value. NULL(default) - print to screen, 'pdf', 'png', 'pngBMP' for

 $bitmap\ png, helpful\ for\ large\ datasets.\ Input\ for\ Consensus\ Cluster Plus:: Consensus\ Cluster\ Plus, helpful\ for\ large\ datasets.$ 

 ${\tt ConsensusClusterPlus::calcICL,callNCW,plotCompareCW}$ 

clusterAlg character value. cluster algorithm. 'spectralClusteringAffinity' for spectral clus-

tering of similarity/affinity matrix(default), other methods for clustering of distance matrix, 'hc' heirarchical (hclust), 'pam' for paritioning around medoids, 'km' for k-means upon data matrix, 'kmdist' for k-means upon distance matrices (former km option), or a function that returns a clustering. Input for

 ${\tt ConsensusClusterPlus::ConsensusClusterPlus.}$ 

innerLinkage heirarchical linkage method for subsampling, or "complete" (default). Input for

ConsensusClusterPlus::ConsensusClusterPlus

... Other input arguments for ConsensusClusterPlus::ConsensusClusterPlus

example\_data 5

#### Value

A list of three items

- ncw A matrix of normalized consensus weights. Output from callNCW.
- fcluster A list of length maxK. Each element is a list containing consensusMatrix (numerical matrix), consensusTree (hclust), consensusClass (consensus class asssignments). ConsensusClusterPlus also produces images. Output from ConsensusClusterPlus::ConsensusClusterPlus
- icl a list of two elements clusterConsensus and itemConsensus corresponding to cluster-consensus and item-consensus. Output from ConsensusClusterPlus::ConsensusClusterPlus

#### **Examples**

example\_data

The input data for example

#### **Description**

In this matrix, columns represent the results of different clustering results and rows are samples.

#### Usage

```
example_data
```

#### **Format**

A matrix with 10 rows and 5 columns.

6 plotCompareCW

plotCompareCW	Plot of original consensus weights vs. normalized consensus weights grouping by the number of co-appeared percent of clustering(non-missing).
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#### Description

Plot of original consensus weights vs. normalized consensus weights grouping by the number of co-appeared percent of clustering(non-missing).

#### Usage

```
plotCompareCW(title, label, ncw, plot = NULL)
```

#### Arguments

title	A character value for output directory.
label	A matrix or data frame of input labels, columns=different clustering results and rows are samples.
ncw	A matrix of normalized consensus weights with sample-by-sample as the order of sample(rows) in label.
plot	character value. NULL(default) - print to screen, 'pdf', 'png', 'pngBMP' for bitmap png, helpful for large datasets, or 'pdf'.

#### Value

A ggplot point in PDF format with x-axis: original consensus weights; y-axis: normalized consensus weights; color: percent of co-appeared of clustering; size: number of duplicates sample

#### **Examples**

```
# load data
data(example_data)
label=example_data

# if plot is not NULL, results will be saved in "result_output" directory
title="result_output"

ncw<-callNCW(title=title,label=label,stability=TRUE)
plotCompareCW(title=title,label=label,ncw=ncw)</pre>
```

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randConsensusMatrix Calculate consensus weight matrix based on the permuted input label matrix. Internal function used by callNCW

#### Description

Calculate consensus weight matrix based on the permuted input label matrix. Internal function used by callNCW

#### Usage

```
randConsensusMatrix(
    l.seed,
    l.label = label,
    l.ns = ns,
    l.nc = nc,
    l.nv = nv,
    l.index = index,
    l.pair.ind = pair.ind,
    l.ppath = ppath,
    l.plot = plot
)
```

#### Arguments

1.seed	A numerical value to set the random seed for reproducible results, 1000 random label matrix will be generated based on this seed number.
l.label	A matrix or data frame of input labels, columns=different clustering results and rows are samples.
l.ns	A integer value of number of samples, =nrow(1.label)
l.nc	A integer value of number of samples, =ncol(1.label)
l.nv	A integer vector of the number of non missing values for each column in 1.1abe1
1.index	A list of index with length of 1.nc of non missing values for each column in 1.label
l.pair.ind	A n-by-2 index matrix of array indices of upper triangular of 1.1abe1 with non missing values
l.ppath	A character value for output directory.
1.plot	character value. NULL(default) - print to screen, 'pdf', 'png', 'pngBMP' for bitmap png, helpful for large datasets, or 'pdf'.

## Value

A character of finished seed.

Write a binary file of 1000 random consensus weight matrix(as a vector n-by-1, n= nrow(1.pair.ind)) with the seed 1.seed, output file name: paste0("s",1.seed,"rcw").

 ${\tt spectralClusteringAffinity}$ 

Perform spectral clustering algorithms for an affinity matrix, using SNFtool::spectralClustering.

#### Description

Perform spectral clustering algorithms for an affinity matrix, using SNFtool::spectralClustering.

## Usage

```
spectralClusteringAffinity(affi_matrix, k, type = 3)
```

#### Arguments

affi\_matrix A numerical similarity or affinity matrix.

k A number value of clusters

type The variants of spectral clustering to use. See SNFtool::spectralClustering

#### Value

A vector consisting of cluster labels of each sample.

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