

ConsensusClustering Documentation

Module name: ConsensusClustering

Description: Resampling-based clustering

Author: Stefano Monti (WIGR) gp-help@broad.mit.edu

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Summary: Given a set of items to be clustered (items can be either genes or chips/experiments), ConsensusClustering (CC) provides for a method to represent the consensus across multiple runs of a clustering algorithm and to assess the stability of the discovered clusters. To this end, perturbations of the original data are simulated by resampling techniques. The clustering algorithm of choice is applied to each of the perturbed data sets, and the agreement, or *consensus*, among the multiple runs is assessed and summarized in a *consensus matrix*. Each matrix entry is indexed by an item pair, and it measures the proportion of times the pair's items are clustered together across the resampling iterations (ideally, always, or never). A distinct consensus matrix is generated for each of the number of clusters considered (e.g., if kmax=5, consensus matrices corresponding to 2, 3, 4, and 5 clusters will be generated). Visual inspection of the consensus matrices, and of the corresponding summary statistics can be used to determine the best number of clusters (see reference for more details).

References:

• S. Monti, et al. "Consensus Clustering: A resampling-based method for class discovery and visualization of gene expression microarray data", *Machine Learning Journal*, 52(1-2):91-118, 2003.

Usage/Example: CC.out <- ConsensusClustering("ALB_ALT_AML.gct", kmax=5,</pre>

niter=100, normalize.type=3, norm.iter=0,

algo="hierarchical", resample="subsample0.8",

merge.type="average", pink.size=2, out.stub="CCout")

Parameters:

Name	Description	Choices
input.filename	The data based on which to carry	It can be a '.gct', a '.res', or a '.odf'
	out the clustering	file.
niter	Number of resampling iterations	Positive integer
kmax	Try K=2, 3,, kmax clusters	1 < kmax < number of items
normalize.type	Type of normalization to perform on	1=row normalize; 2=column
	data	normalize; 3=both (default);
norm.iter	Number of row/column normalization	A non-negative integer
	iterations to perform. It supercedes	
	normalize.type.	
algo	Clustering algorithm to use	hierarchical; SOM; NMF
resample	resampling scheme to use	'subsample[ratio]', 'features[nfeat]',
		'nosample'
merge.type	How to update the distance measure	"average", "single", "complete".
	(ignored if algo is other than	



'hierarchical')

descent.iter Number of SOM/NMF iterations

(ignored when algo is hierarchical):

pink.size point size of a consensus matrix's

heat map

out.stub Stub pre-pended to all the output

files

positive integer (default: 2000)

 $1 \le pink.size \le 20$

Return Value: An R list whose components have the following names (with <k>=2,...,kmax):

- 1. <out.stub>.<sampleid>.<k>.clu, is a text file listing the items belonging to each cluster. (<sampleid> indicates the type of resampling scheme, and <k> denotes the number of clusters).
- 2. <out.stub>.<sampleid>.<k>.gct is the consensus matrix for <k> clusters, with the entries sorted as in the input data.
- <out.stub>.<sampleid>.srt.<k>.gct is the consensus matrix for <k> clusters, with the entries sorted so as to have items clustering together adjacent to each other.
- 4. <out.stub>.<sampleid>.srt.<k>.gif is the heat map corresponding to the sorted consensus matrix.
- 5. <out.stub>.<sampleid>.statistics.pdf includes a series of plots of statistics (Lorenz curve, Gini index, Consensus CDF) that can be used to determine the best number of clusters.

Platform dependencies:

Task type: clustering

CPU type: any OS: any Java JVM level: 1.4

Language: Java, R (including the R library 'ineg')

Support files: none

Native command line: java –jar <libdir>geneweaver-gp.jar <input.filename> <kmax>

<niter> -o <out.stub> -p -d -s