Package 'SamSPECTRAL'

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Title Identifies cell population in flow cytometry data.
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Imports methods
Description Given a matrix of coordinates as input, SamSPECTRAL first builds the communities to sample the data points. Then, it builds a graph and after weighting the edges by conductance computation, the graph is passed to a classic spectral clustering algorithm to find the spectral clusters. The last stage of SamSPECTRAL is to combine the spectral clusters. The resulting "connected components" estimate biological cell populations in the data sample. For instructions on manual installation, refer to the PDF file provided in the following documentation.
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LazyLoad yes
R topics documented:
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SamSPECTRAL-package

Identifying cell populations in flow cytometry data.

Description

Using a faithful sampling procedure, SamSPECTRAL reduces the size of data points such that applying spectral clustering algorithm on large data such as flow cytometry is possible. Before running the spectral clustering algorithm, it uses potential theory to define similarity between sampled points.

Details

Package: SamSPECTRAL

Type: Package Version: 1.0

Date: 2009-08-31 License: GPL-2 LazyLoad: yes

The main function is SamSPECTRAL. It can be loaded using the command library(SamSPECTRAL) in R. Some parameters should be set properly including: dimensions, normal.sigma and separation.factor. These parameters can be adjusted for a data set by running the algorithm on some samples of that data set. (Normally, 2 or 3 samples are sufficient). Then the function SamSPECTRAL() can be applied to all samples in the data set to identify cell populations in each sample data.

Author(s)

Habil Zare and Parisa Shooshtari

Maintainer: Habil Zare <hzare@bccrc.ca>

References

Zare, H. and Shooshtari, P. and Gupta, A. and Brinkman R.B: Data Reduction for Spectral Clustering to Analyse High Throughput Flow Cytometry Data. BMC Bioinformatics, 2010, 11:403.

See Also

SamSPECTRAL, Building_Communities, Conductance_Calculation, Civilized_Spectral_Clustering, Connecting

```
## Not run:
library(SamSPECTRAL)
# Reading data file which has been transformed using log transform
data(small data)
```

```
\label{eq:local_continuity} $L <- small $$ L <- smsPECTRAL(data.points=full,dimensions=c(1,2,3), normal.sigma = 200, separation.factor = 0.39) $$ plot(full, pch='.', col= L) $$ \# End(Not run) $$
```

Building Communities Builds the communities from the set of all data points.

Description

Some sample points are picked up and the points close to each sample point are considered as members of that community.

Usage

Building Communities(full, m=3000, space.length=1, community.weakness.threshold=1, talk=TRUE, do.sam

Arguments

full The matrix containing the coordinates of all data points.

m Determines an upper bound on the final number of sample points which will be

in range m and $\mathrm{2}\ \mathrm{m}$

space.length An estimate for the length of a cube that is assumed to contain all data points.

community.weakness.threshold

The communities with number of members less than this threshold will be ig-

nored. Normally, setting it to 1 is reasonable.

talk A boolean flag with default value TRUE. Setting it to FALSE will keep running

the procedure quite with no messages.

do.sampling A boolean flag with default value TRUE. If set to FALSE, the sampling stage

will be ignored by picking up all the data points.

Value

Returns a society which is a list of communities.

Author(s)

Parisa Shooshtari and Habil Zare

References

Zare, H. and Shooshtari, P. and Gupta, A. and Brinkman R.B: Data Reduction for Spectral Clustering to Analyse High Throughput Flow Cytometry Data. BMC Bioinformatics, 2010, 11:403.

See Also

SamSPECTRAL

Examples

```
## Not run:
library(SamSPECTRAL)

# Reading data file which has been transformed using log transform
data(small_data)
full <- small

# Parameters:
m <- 3000; ns <- 200; sl <- 3; cwt <-1

# Sample the data and build the communities
society <- Building_Communities(full=full,m=m, space.length=sl, community.weakness.threshold=cwt)

# Ploting the representatives:
plot(full[society$representatives,])

## End(Not run)

Civilized Spectral Clustering
```

Description

The representatives of communities are considered as the vertices of a graph. Assuming the edges have been weighted according to the equivalent conductance between them, this function runs the classic spectral clustering on the graph.

Usage

 $\label{lem:civilized_Spectral_Clustering} Clustering (full, maximum.number.of.clusters, society, conductance, iterations=200, number.of.clusters, society, conductance, iterations=200, number.of.clusters=200, number.of.clus$

Runs the spectral clustering algorithm on the sample points.

Arguments

full The matrix containing the coordinates of all data points.

maximum.number.of.clusters

This parameter is used for fitting the regression line.

number.of.clusters

The default value is NA which leads to computating the number of spectral clusters automatically, otherwise this number will determine the number of spectral

clusters.

society The list of communities.

conductance A matrix in which each entry is the conductance between two communities.

Number of iterations for the k-means algorithm used by the spectral procedure.

200 is an appropriate value.

talk A boolean flag with default value TRUE. Setting it to FALSE will keep running

the procedure quite with no messages.

eigenvalues.num An integer with default value NA which prevents ploting the curve of eigenval-

ues. Otherwise, they will be ploted upto this number.

stabilizer The larger this integer is, the final results will be more stable because the under-

lying kmeans will restart many more times.

Value

A ClusteringResult class object with the following slots,

The k'th element of this list is a vector containing the labels as result of clustering to k parts.

labels.for_num.of.clusters A list containing the desired cluster numbers.

eigen.space The eigen vectors and eigen values of the normalized adjacency matrix computed for spectral clustering.

Author(s)

Parisa Shooshtari and Habil Zare

L <- labels.for num.of.clusters[[number.of.clusters]]

plot(full, pch='.', col= L)

References

Zare, H. and Shooshtari, P. and Gupta, A. and Brinkman R.B: Data Reduction for Spectral Clustering to Analyse High Throughput Flow Cytometry Data. BMC Bioinformatics, 2010, 11:403.

See Also

SamSPECTRAL

Not run:

```
library(SamSPECTRAL)

# Reading data file which has been transformed using log transform
data(small_data)
full <- small

# Parameters:
m <- 3000; ns <- 200; sl <- 3; cwt <-1; precision <- 6; mnc <-30

# Sample the data and build the communities
society <- Building_Communities(full=full,m=m, space.length=sl, community.weakness.threshold=cwt)

# Compute conductance between communities
conductance <- Conductance_Calculation(full=full, normal.sigma=ns, space.length=sl, society=society, precision=p

# Use spectral clustering to cluster the data
# First example:
clust_result <- Civilized_Spectral_Clustering(full=full, maximum.number.of.clusters=mnc, society=society, conduct number.of.clusters <- clust_result@labels.for num.of.clusters <- clust_result@labels.for num.of.clusters
```

```
# Second example:
number.of.clusters <- c(35,20)
# This is faster than runnig Civilized_Spectral_Clustering() twice because the eigen space is not needed to be computed clust_result.not.automatic <-
Civilized_Spectral_Clustering(full=full, society=society, conductance=conductance, number.of.clusters =number.of.clusters |
labels.for_num.of.clusters <- clust_result.not.automatic@labels.for_num.of.clusters
L35 <- labels.for_num.of.clusters[[35]]
L20 <- labels.for_num.of.clusters[[20]]
# plot(full, pch='.', col= L35)

## End(Not run)
```

 ${\bf Conductance_Calculation}$

Computes the conductance between communities.

Description

For each two communities, the conductance between their members is summed up and the result is returned as the conductance between the two communities.

Usage

Conductance_Calculation(full, normal.sigma, space.length, society, precision, talk=TRUE, beta=4)

Arguments

full The matrix containing the coordinates of all data points.

normal.sigma The scaling parameter, the larger it is the algorithm will find smaller clusters. space.length An estimate for the length of a cube that is assumed to contain all data points.

society The list of communities.

precision Determines the precision of computations. Setting it to 6 will work and increas-

ing it does not improve results.

talk A boolean flag with default value TRUE. Setting it to FALSE will keep running

the procedure quite with no messages.

beta A parameter with default value 4 which must NOT be changed except for huge

samples with more than 100,000 data points or for developmental purposes. Setting beta to zero will reduce computational time by applying the following approximation to the conductance calculation step. For each two community, the conductance will be the conductance between their representatives times their

sizes.

Value

Returns a matrix in which each entry is the conductance between two communities.

Author(s)

Parisa Shooshtari and Habil Zare

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References

Zare, H. and Shooshtari, P. and Gupta, A. and Brinkman R.B: Data Reduction for Spectral Clustering to Analyse High Throughput Flow Cytometry Data. BMC Bioinformatics, 2010, 11:403.

See Also

SamSPECTRAL

Examples

```
## Not run:
library(SamSPECTRAL)

# Reading data file which has been transformed using log transform
data(small_data)
full <- small

# Parameters:
m <- 3000; ns <- 200; sl <- 3; cwt <-1; precision <- 6

# Sample the data and build the communities
society <- Building_Communities(full=full,m=m, space.length=sl, community.weakness.threshold=cwt)

# Compute conductance between communities
conductance <- Conductance_Calculation(full=full, normal.sigma=ns, space.length=sl, society=society, precision=p

## End(Not run)
```

Connecting

Combines the spectral clusters to build the connected components.

Description

Considering some biological criterion based on density, the clusters which are identified by spectral clustering are combined to estimate biological populations.

Usage

Connecting(full, society, conductance, number.of.clusters, labels.for num.of.clusters, separation.factor, talk=Tl

Arguments

full The matrix containing the coordinates of all data points.

society The list of communities.

conductance A matrix in which each entry is the conductance between two communities.

number.of.clusters

A list containing the desired cluster numbers.

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labels.for num.of.clusters

The k'th element of this list, is a vector containing the labels as result of clustering to k parts.

separation.factor

This threshold controls to what extend clusters should be combined or kept separate

talk

A boolean flag with default value TRUE. Setting it to FALSE will keep running the procedure quite with no messages.

Details

A hint for setting separation.factor: While separation.factor=0.7 is normally an appropriate value for many datasets, for others some value in range 0.3 to 1.2 may produce better results depending on what populations are of particular interest.

Value

Returns two objects: 1) label, a vector containing the labels that determines to which component each data point belongs. 2) clusters.graph, the max.conductance matrix that describes the original graph based on clusters.

Author(s)

Parisa Shooshtari and Habil Zare

References

Zare, H. and Shooshtari, P. and Gupta, A. and Brinkman R.B: Data Reduction for Spectral Clustering to Analyse High Throughput Flow Cytometry Data. BMC Bioinformatics, 2010, 11:403.

See Also

SamSPECTRAL

```
## Not run:
library(SamSPECTRAL)

# Reading data file which has been transformed using log transform
data(small_data)
full <- small

# Parameters:
m <- 3000; ns <- 200; sl <- 3; cwt <-1; precision <- 6; mnc <-30

# Sample the data and build the communities
society <- Building_Communities(full=full,m=m, space.length=sl, community.weakness.threshold=cwt)

# Compute conductance between communities
conductance <- Conductance_Calculation(full=full, normal.sigma=ns, space.length=sl, society=society, precision=p

# Use spectral clustering to cluster the data
```

eigen.values.10

```
clust_result <- Civilized_Spectral_Clustering(full=full, maximum.number.of.clusters=mnc, society=society, conduct number.of.clusters <- clust_result@number.of.clusters labels.for_num.of.clusters <- clust_result@labels.for_num.of.clusters L <- labels.for_num.of.clusters[[number.of.clusters]] # plot(full, pch='.', col= L)

# Connect components
L <- Connecting(full=full, society=society, conductance=conductance, number.of.clusters=number.of.clusters, labels.for_num.of.clusters=labels.for_num.of.clusters, separation.factor=0.39)

plot(full, pch='.', col= L)

## End(Not run)
```

eigen.values.10

Eigenvalues for building the SamSPECTRAL vignette.

Description

This file contains a vector that represents the eigenvalues of the small example if normal.sigma=10.

Usage

```
data(eigen.values.10)
```

Format

This RData contains a vector.

References

Zare, H. and Shooshtari, P. and Gupta, A. and Brinkman R.B: Data Reduction for Spectral Clustering to Analyse High Throughput Flow Cytometry Data. BMC Bioinformatics, 2010, 11:403.

```
data(eigen.values.10)
plot(eigen.values.10)
```

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eigen.values.1000

Eigenvalues for building the SamSPECTRAL vignette.

Description

This file contains a vector that represents the eigenvalues of the small example if normal.sigma=1000.

Usage

```
data(eigen.values.1000)
```

Format

This RData contains a vector.

References

Zare, H. and Shooshtari, P. and Gupta, A. and Brinkman R.B: Data Reduction for Spectral Clustering to Analyse High Throughput Flow Cytometry Data. BMC Bioinformatics, 2010, 11:403.

Examples

```
data(eigen.values.1000)
plot(eigen.values.1000)
```

 ${\bf SamSPECTRAL}$

Identifies the cell populations in flow cytometry data.

Description

Given an FCS file as input, SamSPECTRAL first builds the communities to sample the data points. Then, it builds a graph and after weighting the edges of the graph by conductance computation, it is passed to a classic spectral clustering algorithm to find the spectral clusters. The last stage of SamSPECTRAL is to combine the spectral clusters. The resulting "connected components" estimate biological cell populations in the data sample.

Usage

SamSPECTRAL(data.points, dimensions=1:dim(data.points)[2], normal.sigma, separation.factor,number.of.clutalk = TRUE, precision = 6, eigenvalues.num = NA, return only.labels=TRUE, do.sampling=TRUE, beta=4, s

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Arguments

data.points A matrix that contains coordinates of the data points.

dimensions A vector that determines which dimension of the data point matrix are chosen

for investigation.

normal.sigma A scaling parameter that determines the "resolution" in the spectral clustering

stage. By increasing it, more spectral clusters are identified. This can be useful when "small" population are aimed. See the user manual for a suggestion on

how to set this parameter using the eigenvalue curve.

separation.factor

This threshold controls to what extend clusters should be combined or kept separate. Normally, an appropriate value will fall in range 0.3-2.

number.of.clusters

The default value is NA which leads to computing the number of spectral clusters automatically, otherwise this number will determine the number of spectral

clusters.

talk A boolean flag with default value TRUE. Setting it to FALSE will keep running

the procedure quite with no messages.

precision Determines the precision of computations. Setting it to 6 will work and increas-

ing it does not improve results.

eigenvalues.num An integer with default value NA which prevents ploting the curve of eigenval-

ues. Otherwise, they will be ploted upto this number.

return only.labels

A boolean flag with default value TRUE. If the user set it to FALSE, SamSPEC-TRAL function will return all the intermediate objects that are computed during

the sampling, similarity calculation, spectral clustering and combining stages.

do.sampling A boolean flag with default value TRUE. If set to FALSE, the sampling stage

will be ignored by picking up all the data points.

beta A parameter with default value 4 which must NOT be changed except for huge

samples with more than 100,000 data points or for developmental purposes. Setting beta to zero will reduce computational time by applying the following approximation to the conductance calculation step. For each two community, the conductance will be the conductance between their representatives times their

sizes.

scale A vector the length of which is equal to the number of dimensions. The coordi-

nates in each dimension are multiplied by the corresponding scaling factor. So, the bigger this factor is for a dimension, SamSPECTRAL will consider that dimension to be "more significant" and consequently, that dimension will be more

effective in clustering.

stabilizer The larger this integer is, the final results will be more stable because the under-

lying kmeans will restart many more times.

Details

Hints for setting separation.factor and normal.sigma: While separation.factor=0.7 is normally an appropriate value for many datasets, for others some value in range 0.3 to 1.2 may produce better results depending on what populations are of particular interest. The larger normal.sigma is the algorithm will find smaller clusters. It can be adjusted best by considering the plot of eigenvalues as explained in the vignette.

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Value

Returns a vector of labels for data points. If the input parameter return_only.labels is set to FALSE, all the objects that are computed during the intermediate will be returned including: society for sampling stage, conductance for similarity calculation, and clustering_result.

Author(s)

Habil Zare and Parisa Shooshtari

References

Zare, H. and Shooshtari, P. and Gupta, A. and Brinkman R.B: Data Reduction for Spectral Clustering to Analyse High Throughput Flow Cytometry Data. BMC Bioinformatics, 2010, 11:403.

See Also

SamSPECTRAL, Building_Communities, Conductance_Calculation, Civilized_Spectral_Clustering, Connecting

Examples

```
## Not run:
library(SamSPECTRAL)

# Reading data file which has been transformed using log transform
data(small_data)
full <- small

L <- SamSPECTRAL(data.points=full,dimensions=c(1,2,3), normal.sigma = 200, separation.factor = 0.39)
plot(full, pch='.', col= L)

## End(Not run)
```

small

Flow cytometry data to test SamSPECTRAL algorithm.

Description

This FCS file is a small one used to show how to set SamSPECTRAL parameters.

Usage

```
data(small_data)
```

Format

This is an FCS file.

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References

Zare, H. and Shooshtari, P. and Gupta, A. and Brinkman R.B: Data Reduction for Spectral Clustering to Analyse High Throughput Flow Cytometry Data. BMC Bioinformatics, 2010, 11:403.

Examples

```
data(small_data)
full <- small
    plot(full, pch='.')</pre>
```

stmFSC

Flow cytometry data to test SamSPECTRAL algorithm.

Description

This FCS file is used as demo data to illustrate SamSPECTRAL capabilities in identifying cell populations.

Usage

```
data(stm)
```

Format

The is an FCS file.

References

Zare, H. and Shooshtari, P. and Gupta, A. and Brinkman R.B: Data Reduction for Spectral Clustering to Analyse High Throughput Flow Cytometry Data. BMC Bioinformatics, 2010, 11:403.

```
 \begin{array}{l} {\rm data(stm)} \\ {\rm \# \,Read \,\,data \,\,files \,\,and \,\,transform \,\,them \,\,using \,\,log \,\,transform \,\,data.points <- \,\,stmFSC@exprs \,\,dimensions <- \,\,c(3,4,7) \,\,full <- \,\,log10({\rm data.points[,dimensions]}) \,\,\\ {\rm plot(full, \,\,pch='.')} \end{array}
```

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