Package 'graphclass'

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COBRE.data

fMRI brain networks of the COBRE dataset

Description

This dataset contains functional connectivity brain networks of 124 subjects and corresponding class labels indicating schizophrenia status (70 healthy controls and 54 schizophrenic subjects) obtained from the COBRE dataset.

Usage

```
data(COBRE.data)
```

Format

COBRE. data is a list with three elements

X.cobre A matrix containing the upper triangular part of the networks in column major order and without the diagonal entries. The networks are composed of 263 nodes obtained from the Power parcellation (Power et al. 2011) and 34453 different edges. Each row corresponds to a subject in the data, and the columns correspond to edges in the networks. The edge weights are the Fisher-transformed correlation between the fMRI time series of the nodes. Nuisance covariates like age, gender, motion (meanFD and meanFDquad) and handedness have been regressed out. For a description of the preprocessing steps to obtain the network edge weights, see Arroyo Relion et al. (2019).

Y.cobre Class labels of the subjects in the dataset. Y=1 represents schizophrenia status. **subject.label** Subject ID number in the COBRE dataset.

Source

```
http://fcon_1000.projects.nitrc.org/indi/retro/cobre.html
```

References

C. J. Aine, H. J. Bockholt, J. R. Bustillo, J. M. CaClive, A. Caprihan, C. Gasparovic, F. M. Hanlon, J. M. Houck, R. E. Jung, J. Lauriello, J. Liu, A. R. Mayer, N. I. Perrone-Bizzozero, S. Posse, J. M. Stephen, J. A. Turner, V. P. Clark, Vince D. Calhoun (2017). "Multimodal Neuroimaging in Schizophrenia: Description and Dissemination." *Neuroinformatics*, **15**(4), 343–364. ISSN 1559-0089, doi: 10.1007/s1202101793389, https://doi.org/10.1007/s12021-017-9338-9.

J.D. Arroyo Relion, D. Kessler, E. Levina, S.F. Taylor (2019). "Network classification with applications to brain connectomics." *Ann Appl Stat.*, **13**(3), 1648.

Jonathan D Power, Alexander L Cohen, Steven M Nelson, Gagan S Wig, Kelly Anne Barnes, Jessica A Church, Alecia C Vogel, Timothy O Laumann, Fran M Miezin, Bradley L Schlaggar, others (2011). "Functional network organization of the human brain." *Neuron*, **72**(4), 665–678.

```
data(COBRE.data)
plot_adjmatrix(COBRE.data$X.cobre[1,])
```

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construct_D

Constructor for penalty matrix

Description

This function constructs an auxiliary matrix to compute the node penalty in the regularized classifier, which is internally used by the optimization algorithm. This matrix can be passed to the function graphclass in order to avoid creating it every time this function is called. Use it for efficiency when running the classifier multiple times (for example, in a cross-validation routine).

Usage

```
construct_D(nodes = 264)
```

Arguments

nodes

Number of nodes in the network, by default is 264 (Power parcellation).

Value

A sparse matrix

```
D263 = construct_D(263)
# Load COBRE data
data(COBRE.data)
X <- COBRE.data$X.cobre</pre>
Y <- COBRE.data$Y.cobre
# 5-fold cross validation of the subgraph selection penalty
fold_index <- (1:length(Y) %% 5) + 1</pre>
gclist <- list()</pre>
for(fold in 1:5) {
    foldout <- which(fold_index == fold)</pre>
    gclist[[fold]] <- graphclass(X = X[-foldout,], Y = factor(Y[-foldout]),</pre>
                      Xtest = X[foldout,], Ytest = factor(Y[foldout]),
                      type = "intersection",
                      lambda = 1e-4, rho = 1, gamma = 1e-5,
                      D = D263)
# test error on each fold
lapply(gclist, function(gc) gc$test_error)
```

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σ <u>e</u> t	matrix	

Converts a vector into an adjacency matrix

Description

Given a vector that encodes an adjacency matrix, returns the matrix representation.

Usage

```
get_matrix(beta, type = c("undirected", "directed"))
```

Arguments

beta Vectorized adjacency matrix. If the network is undirected, the vector is assumed

to represent the upper triangular part of the adjacency matrix in column major order. For undirected network, the vector contains the entries ordered by column

and excluding the diagonal elements.

type Specifies whether the vector represents an undirected or directed network.

Default is undirected.

Value

Adjacency matrix.

Examples

```
# Obtain the adjacency matrix of a COBRE data subject
data(COBRE.data)
A <- get_matrix(COBRE.data$X.cobre[1,], type = "undirected")
plot_adjmatrix(A)</pre>
```

graphclass

Regularized logistic regression classifier for networks.

Description

graphclass fits a regularized logistic regression to a set of network adjacency matrices with responses, and returns an object with the classifier.

Plots the coefficients matrix obtained with the function graphclass.

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Usage

```
graphclass(
  X = NULL,
  Y = NULL
  type = c("intersection", "union", "groups", "fusion"),
)
## Default S3 method:
graphclass(
  X = NULL
  Y = NULL
  Xtest = NULL,
  Ytest = NULL,
  Adj_list = NULL,
  type = c("intersection", "union", "groups", "fusion"),
  lambda = 0,
  rho = 0,
  gamma = 1e-05,
  params = NULL,
  id = "",
  verbose = F,
  D = NULL,
  Groups = NULL,
  G_penalty_factors = NULL,
)
## S3 method for class 'graphclass'
plot(object, ...)
```

Arguments

Χ

D

	(by column order) upper triangular part of a network adjacency matrix.
Υ	A vector containing the class labels of the training samples (only 2 classes are supported for now).
type	Type of penalty function. Default is "intersection". See details.
Xtest	Optional argument for providing a matrix containing the test samples, with each row representing an upper-triangular vectorized adjacency matrix.
Ytest	Optional argument containing the labels of test samples.
Adj_list	A training list of of symmetric adjacency matrices with zeros in the diagonal
lambda	penalty parameter $lambda$, by default is set to 0.
rho	penalty parameter ${\it rho}$ controlling sparsity, by default is set to 0.
gamma	ridge parameter (for numerical purposes). Default is gamma = 1e-5.

params

A list containing internal parameters for the optimization algorithm. See details.

verbose whether output is printed

matrix D used by the penalty to define the groups. This optional argument can be used to pass a precomputed matrix D, which can be time saving if the method is fitted multiple times. See the function construct_D.

A matrix with the training samples, in wich each row represents the vectorized

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Groups list of lists, where each list correspond to a grouping and each sublist to sets of

indexes in X. Each sublist should be a non-overlapping group.

G_penalty_factors

For type "groups", each group is penalized by this factor. Should sum to 1.

object trained graphclass object

Details

The function graphclass fits a regularized logistic regression to classify a set of network adjacency matrices with N labeled nodes and corresponding responses. The classifier fits a matrix of coefficients $B \in \mathbb{R}^{N \times N}$, in which B_{ij} indicates the coefficient corresponding to the edge (i, j).

The argument type provides options to choose the penalty function. If type = "intersection" or "union", the penalty corresponds to the node selection penalty defined as

$$\Omega(B) = \lambda \left(\sum_{i=1}^{N} \sqrt{\sum_{j=1}^{N} B_{ij}^2} + \rho \sum_{i=1}^{N} \sum_{j=1}^{N} |B_{ij}| \right).$$

When type = "intersection", a symmetric restriction on B is enforced, and the penalty promotes subgraph selection. If type = "union", the penalty promotes individual node selection. See Arroyo Relion et al. (2019) for more details.

The value type = "groups" corresponds to a generic group lasso penalty. The groups of edges have to be specified using the argument Groups with a list of arrays, in which each element of the list corresponds to a group, and the array indicates the indexes of the variables in that group. The optional argument G_penalty_factors is an array of size equal to the number of groups, and can be used to specify different weights for each group on the penalty (for example, when groups have different sizes).

The optional argument params is a list that allows to control some internal parameters of the optimization algorithm. The elements beta_start and b_start are initial values for the optimization algorithm. The value of beta_start is a vector that indicates the weights of the upper triangular part of B, and b_start is the initial value of the threshold in the logistic regression. By default, these parameters are set to zero. The elements MAX_ITER and CONV_CRIT can be used to change the maximum number of iterations and the convergence criterion in the proximal algorithm for fitting the node selection penalty (see Arroyo Relion et al. (2019)). By default, these values are set to MAX_ITER=300 and CONV_CRIT = 1e-5.

Value

An object containing the trained graph classifier.

beta Edge coefficients vector of the regularized logistic regression solution.

b Intercept value.

Yfit Fitted logistic regression probabilities in the train data.

Ypred Predicted class for the test samples (if available).
train_error Percentage of train samples that are misclassified.

test_error Percentage of test samples that are misclassified (if available).

References

J.D. Arroyo Relion, D. Kessler, E. Levina, S.F. Taylor (2019). "Network classification with applications to brain connectomics." *Ann Appl Stat.*, **13**(3), 1648.

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See Also

```
plot.graphclass, predict.graphclass
```

Examples

```
# Load COBRE data
data(COBRE.data)
X <- COBRE.data$X.cobre</pre>
Y <- COBRE.data$Y.cobre
# An example of the subgraph selection penalty
gc = graphclass(X = X, Y = factor(Y), type = "intersection",
               lambda = 1e-4, rho = 1, gamma = 1e-5)
plot(gc)
# 5-fold cross validation
fold_index <- (1:length(Y) \%\% 5) + 1
# Make penalty matrix in advance to save time
D263 <- construct_D(nodes = 263)</pre>
gclist <- list()</pre>
for(fold in 1:5) {
    foldout <- which(fold_index == fold)</pre>
    gclist[[fold]] <- graphclass(X = X[-foldout,], Y = factor(Y[-foldout]),</pre>
                      Xtest = X[foldout,], Ytest = factor(Y[foldout]),
                      type = "intersection",
                      lambda = 1e-4, rho = 1, gamma = 1e-5,
                      D = D263)
}
# test error on each fold
lapply(gclist, function(gc) gc$test_error)
data(COBRE.data)
X <- COBRE.data$X.cobre</pre>
Y <- COBRE.data$Y.cobre
# An example of the subgraph selection penalty
gc = graphclass(X, Y = factor(Y), lambda = 1e-5, rho = 1)
plot(gc)
```

matrix_to_vec

Convert matrix to vector.

Description

The function encodes an adjacency matrix into a vector.

Usage

```
matrix_to_vec(A, type = c("undirected", "directed"))
```

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Arguments

A Adjacency matrix of a network.

type Parameter to specify whether the adjacency matrix represents an undirected or

directed network. Default is undirected.

Value

A vector containing the upper triangular part of an adjacency matrix (if the graph is undirected) or adjacency entry values ordered by column and excluding the diagonal entries (if the graph is directed).

node_sparsity

Percentage of inactive nodes in a network

Description

Calculates the node sparsity of a vectorized adjacency matrix, defined as the average number of nodes with no edges.

Usage

```
node_sparsity(beta, type = c("undirected", "directed"))
```

Arguments

beta Vectorized adjacency matrix.

type Specifies whether the vector represents an undirected or directed network.

Default is undirected.

Value

Percentage of inactive nodes in the graph

```
A <- matrix(0, ncol = 4, nrow = 4)
A[2, 1] <- 1
A[1, 2] <- 1
A[2, 3] <- 1
A[3, 2] <- 1
vec <- matrix_to_vec(A)
node_sparsity(vec)</pre>
```

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plot_adjmatrix	Plots a vectorized adjacency matrix.

Description

Draws a plot of the adjacency matrix represented by a vector using the function levelplot of the lattice package.

Usage

```
plot_adjmatrix(
  edgevalues,
  type = c("undirected", "directed"),
  edgetype = c("real", "prob", "binary"),
  communities = NULL,
  community_labels = NULL,
  main = "",
  axislabel = "Nodes",
  colorlims = NULL
)
```

Arguments

edgevalues	Edge values of th	ne adiacency matrix	. The argument can	be either a square ad-

jacency matrix, or a vectorized adjacency matrix. For undirected networks, the vector should contain the upper triangular part in column-major order. For di-

rected networks use both

type If edgevalues is a vector, this parameter specifies whether the vector represents

an undirected or directed network. Default is undirected.

edgetype Specify the type of edge values. For real-valued edges, type = "real". If the

edges are between 0 and 1, tipe = "prob". For binary edges, type = "binary".

communities Optional argument to specify a list in which each element contains an array

indicating the indexes of the nodes on each community.

community_labels

Labels for each community. The array should have the same length than communities.

main Title of the plot axislabel Label for the axes.

colorlims An array with two elements indicating the minimum and maximum value in the

color bar

Value

An objected returned by the function levelplot of the lattice package.

```
# Plot the adjacency matrix of a COBRE data subject
data(COBRE.data)
X1 <- COBRE.data$X.cobre[1,]</pre>
```

10 power.parcellation

power.parcellation

Node assignments of the Power parcellation

Description

Table containing the node assignments to brain systems of the 264 regions of interest from Power et al. (2011). These nodes were used to construct COBRE.data and UMich.data. Note that node 75 is missing in the COBRE data.

Usage

```
power.parcellation
```

Format

power.parcellation is a data frame in which the nodes represent nodes. The data frame has three columns:

Master.Assignment Brain system number

Color Color representing the system.

Suggested.System Suggested brain system.

Source

```
http://www.nil.wustl.edu/labs/petersen/Resources_files/Consensus264.xls
```

References

Jonathan D Power, Alexander L Cohen, Steven M Nelson, Gagan S Wig, Kelly Anne Barnes, Jessica A Church, Alecia C Vogel, Timothy O Laumann, Fran M Miezin, Bradley L Schlaggar, others (2011). "Functional network organization of the human brain." *Neuron*, **72**(4), 665–678.

```
data(power.parcellation)
```

predict.graphclass 11

r grapn ciassiпer	
1	r graph classifier

Description

Given an object of type graphclass, this function obtains the fitted classes for a new data.

Usage

```
## S3 method for class 'graphclass'
predict(object, newdata, type = c("class", "prob", "error"), Ytest, ...)
```

Arguments

type	Indicates the type of response: class for class predictions, prob for predicted probabilities, error for misclassification error (if Ytest is provided).
Ytest	If type = "error", true classes to compare.
gc	A trained graph classifier object
X	A matrix containing rows with vectorized upper triangular adjacency matrices (column-major order)

Value

A vector containing the predicted classes.

Examples

UMich.data

fMRI brain networks of the UMich dataset

Description

This dataset contains functional connectivity brain networks of 79 subjects and corresponding class labels indicating schizophrenia status (40 healthy controls and 39 schizophrenic subjects). The fMRI data used to obtain these networks were collected by Professor Stephen F. Taylor's lab at the University of Michigan.

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Usage

```
data(UMich.data)
```

Format

UMich. data is a list with two elements

X.cobre A matrix containing the upper triangular part of the networks in column major order and without the diagonal entries. The networks are composed of 264 nodes obtained from the Power parcellation (Power et al. 2011) and 34716 different edges. Each row corresponds to a subject in the data, and the columns correspond to edges in the networks. The edge weights are the Fisher-transformed correlations between the fMRI time series of the nodes. Nuisance covariates like age, gender, motion (meanFD and meanFDquad) and handedness have been regressed out. For a description of the preprocessing steps to obtain the network edge weights, see Arroyo Relion et al. (2019).

Y.cobre Class labels of the subjects in the dataset. Y = 1 represents schizophrenia status.

References

J.D. Arroyo Relion, D. Kessler, E. Levina, S.F. Taylor (2019). "Network classification with applications to brain connectomics." *Ann Appl Stat.*, **13**(3), 1648.

Jonathan D Power, Alexander L Cohen, Steven M Nelson, Gagan S Wig, Kelly Anne Barnes, Jessica A Church, Alecia C Vogel, Timothy O Laumann, Fran M Miezin, Bradley L Schlaggar, others (2011). "Functional network organization of the human brain." *Neuron*, **72**(4), 665–678.

```
data(UMich.data)
plot_adjmatrix(UMich.data$X.umich[1,])
```

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