# Package: eclust (via r-universe)

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Type Package

**Title** Environment Based Clustering for Interpretable Predictive Models in High Dimensional Data

Version 0.1.0

Description Companion package to the paper: An analytic approach for interpretable predictive models in high dimensional data, in the presence of interactions with exposures. Bhatnagar, Yang, Khundrakpam, Evans, Blanchette, Bouchard, Greenwood (2017) <DOI:10.1101/102475>. This package includes an algorithm for clustering high dimensional data that can be affected by an environmental factor.

**Depends** R (>= 3.3.1)

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URL https://github.com/sahirbhatnagar/eclust/,
 http://sahirbhatnagar.com/eclust/

BugReports https://github.com/sahirbhatnagar/eclust/issues

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Repository https://sahirbhatnagar.r-universe.dev

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plot.eclust

Plot Heatmap of Cluster Summaries by Exposure Status

# Description

Plots cluster summaries such as the 1st principal component or average by exposure status. This is a plot method for object of class eclust returned by the r\_cluster\_data function. Two heatmaps, side-by-side are returned, where the first heatmap corresponds to the unexposed subjects and the second heatmap corresponds to the exposed subjects.

### Usage

```
## S3 method for class 'eclust'
plot(x, type = c("ECLUST", "CLUST"), summary = c("pc",
   "avg"), sample = c("training", "test"), unexposed_title = "E=0",
   exposed_title = "E=1", ...)
```

### **Arguments**

x object of class eclust, which is returned by the r\_cluster\_data function

type show results from the "ECLUST" (which considers the environment) or "CLUST"

(which ignores the environment) methods. Default is "ECLUST". See r\_cluster\_data

for details. This function uses the clustersAddon object for "ECLUST" and the

clustersAll for "CLUST"

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```
summary show the 1st principal component or the average of each cluster. Default is "pc".

which sample to show, the "training" or the "test" set. Default is "training". This is determined by the train_index and test_index arguments in the r_cluster_data function. If you want to show all subjects, then provide the numeric vector 1:n to either argument, where n is the entire sample size.

unexposed_title

The title for the unexposed subjects heatmap. Default is "E=0".

exposed_title

The title for the exposed subjects heatmap. Default is "E=1".

other arguments passed to the Heatmap function
```

#### **Details**

Rows are the cluster summaries and columns are the subjects. This function determines the minimum and maximum value for the whole dataset and then creates a color scale using those values with the colorRamp2. This is so that both heatmaps are on the same color scale, i.e., each color represents the same value in both heatmaps. This is done for being able to visually compare the results.

### Value

a plot of two Heatmaps, side-by-side, of the cluster summaries by exposure status

```
## Not run:
data("tcgaov")
tcgaov[1:5,1:6, with = FALSE]
Y <- log(tcgaov[["OS"]])</pre>
E <- tcgaov[["E"]]</pre>
genes <- as.matrix(tcgaov[,-c("OS","rn","subtype","E","status"),with = FALSE])</pre>
trainIndex <- drop(caret::createDataPartition(Y, p = 1, list = FALSE, times = 1))</pre>
testIndex <- setdiff(seq_len(length(Y)),trainIndex)</pre>
cluster_res <- r_cluster_data(data = genes,</pre>
                                response = Y,
                                exposure = E,
                                train_index = trainIndex,
                                test_index = testIndex,
                                cluster_distance = "tom",
                                eclust_distance = "difftom",
                                measure_distance = "euclidean",
                                clustMethod = "hclust",
                                cutMethod = "dynamic",
                                method = "average",
                                nPC = 1,
                                minimum_cluster_size = 60)
class(cluster_res)
plot(cluster_res, show_column_names = FALSE)
```

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```
## End(Not run)
```

plot.similarity

Function to generate heatmap

# **Description**

Plots a heatmap of a similarity matrix such as a correlation matrix or a TOM matrix. This function is a plotting method for an object of class similarity. These objects are returned by the s\_generate\_data and s\_generate\_data\_mars functions

## Usage

```
## S3 method for class 'similarity'
plot(x, color = viridis::viridis(100), truemodule,
   active, ...)
```

# **Arguments**

X	an object of class similarity. This is a p x p symmetric matix such as a correlation matrix or a TOM matrix, where p is the number of genes
color	colors for the heatmap. By default it uses the viridis color scheme. The viridis package needs to be installed.
truemodule	a numeric vector of length p where p is the number of genes, giving the module membership. By default, $0 = \text{Grey}$ , $1 = \text{Turquoise}$ , $2 = \text{Blue}$ , $3 = \text{Red}$ , $4 = \text{Green}$ , and $5 = \text{Yellow}$ . This information is used for annotating the heatmap
active	a binary vector of length p (where p is the number of genes) where 0 means that gene is not related to the response, and 1 means that the gene is associated to the response.
	other arguments passed to the pheatmap function

### Value

a heatmap of a similarity matrix

### Note

this function is only meant to be used with output from the s\_generate\_data and s\_generate\_data\_mars functions, since it assumes a fixed number of modules.

```
## Not run:
corrX <- cor(simdata[,c(-1,-2)])
class(corrX) <- append(class(corrX), "similarity")
plot(corrX, truemodule = c(rep(1:5, each=150), rep(0, 250)))
## End(Not run)</pre>
```

r\_cluster\_data 5

	-		
r	$\mathcal{C}$	luster	data

Cluster data using environmental exposure

# **Description**

This is one of the functions for real data analysis, which will cluster the data based on the environment, as well as ignoring the environment

### Usage

```
r_cluster_data(data, response, exposure, train_index, test_index,
   cluster_distance = c("corr", "corr0", "corr1", "tom", "tom0", "tom1",
   "diffcorr", "difftom", "fisherScore"), eclust_distance = c("fisherScore",
   "corScor", "diffcorr", "difftom"), measure_distance = c("euclidean",
   "maximum", "manhattan", "canberra", "binary", "minkowski"),
   minimum_cluster_size = 50, ...)
```

### **Arguments**

data n x p matrix of data. rows are samples, columns are genes or cpg sites. Should

not contain the environment variable

response numeric vector of length n

exposure binary (0,1) numeric vector of length n for the exposure status of the n samples

train\_index numeric vector indicating the indices of response and the rows of data that are

in the training set

test\_index numeric vector indcating the indices of response and the rows of data that are

in the test set

cluster\_distance

character representing which matrix from the training set that you want to use to cluster the genes. Must be one of the following

 corr, corr0, corr1, tom, tom0, tom1, diffcorr, difftom, corScor, tomScor, fisherScore

### eclust\_distance

character representing which matrix from the training set that you want to use to cluster the genes based on the environment. See cluster\_distance for avaiable options. Should be different from cluster\_distance. For example, if cluster\_distance=corr and EclustDistance=fisherScore. That is, one should be based on correlations ignoring the environment, and the other should be based on correlations accounting for the environment. This function will always return this add on

#### measure\_distance

one of "euclidean", "maximum", "manhattan", "canberra", "binary", "minkowski" to be passed to dist function for calculating the distance for the clusters based on the corr,corr1,corr0, tom, tom0, tom1 matrices

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```
minimum_cluster_size
```

The minimum cluster size. Only applicable if cutMethod='dynamic'. This argument is passed to the cutreeDynamic function through the u\_cluster\_similarity function. Default is 50.

. . arguments passed to the u\_cluster\_similarity function

#### **Details**

This function clusters the data. The results of this function should then be passed to the r\_prepare\_data function which output the appropriate X and Y matrices in the right format for regression packages such as mgcv, caret and glmnet

#### Value

a list of length 8:

clustersAddon clustering results based on the environment and not the environment. see u\_cluster\_similarity for details

**clustersAll** clustering results ignoring the environment. See u\_cluster\_similarity for details **etrain** vector of the exposure variable for the training set

**cluster\_distance\_similarity** the similarity matrix based on the argument specified in cluster\_distance **eclust\_distance\_similarity** the similarity matrix based on the argument specified in eclust\_distance

**clustersAddonMembership** a data.frame and data.table of the clustering membership for clustering results based on the environment and not the environment. As a result, each gene will show up twice in this table

**clustersAllMembership** a data.frame and data.table of the clustering membership for clustering results based on all subjects i.e. ignoring the environment. Each gene will only show up once in this table

**clustersEclustMembership** a data.frame and data.table of the clustering membership for clustering results accounting for the environment. Each gene will only show up once in this table

# See Also

```
u_cluster_similarity
```

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```
exposure = E,
                              train_index = trainIndex,
                              test_index = testIndex,
                              cluster_distance = "tom",
                              eclust_distance = "difftom",
                              measure_distance = "euclidean",
                              clustMethod = "hclust",
                              cutMethod = "dynamic",
                              method = "average",
                              nPC = 1,
                              minimum_cluster_size = 60)
# the number of clusters determined by the similarity matrices specified
# in the cluster_distance and eclust_distance arguments. This will always be larger
# than cluster_res$clustersAll$nclusters which is based on the similarity matrix
# specified in the cluster_distance argument
cluster_res$clustersAddon$nclusters
# the number of clusters determined by the similarity matrices specified
# in the cluster_distance argument only
cluster_res$clustersAll$nclusters
## End(Not run)
```

r\_prepare\_data

Prepare data for regression routines

# Description

This function will output the appropriate X and Y matrices in the right format for regression packages such as mgcv, caret and glmnet

# Usage

```
r_prepare_data(data, response = "Y", exposure = "E", probe_names)
```

# Arguments

data	the data frame which contains the response, exposure, and genes or cpgs or covariates. the columns should be labelled.
response	the column name of the response in the data argument
exposure	the column name of the exposure in the data argument
probe_names	the column names of the genes, or cpg sites or covariates

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### Value

a list of length 5:

X the X matrix

Y the response vector

E the exposure vector

main\_effect\_names the names of the main effects including the exposure

interaction\_names the names of the interaction effects

```
data("tcgaov")
tcgaov[1:5,1:6, with = FALSE]
Y <- log(tcgaov[["OS"]])
E <- tcgaov[["E"]]</pre>
genes <- as.matrix(tcgaov[,-c("OS","rn","subtype","E","status"),with = FALSE])</pre>
trainIndex <- drop(caret::createDataPartition(Y, p = 0.5, list = FALSE, times = 1))</pre>
testIndex <- setdiff(seq_len(length(Y)),trainIndex)</pre>
## Not run:
cluster_res <- r_cluster_data(data = genes,</pre>
                               response = Y,
                               exposure = E,
                               train_index = trainIndex,
                               test_index = testIndex,
                               cluster_distance = "tom",
                               eclust_distance = "difftom",
                               measure_distance = "euclidean",
                               clustMethod = "hclust",
                               cutMethod = "dynamic",
                               method = "average",
                               nPC = 1,
                               minimum_cluster_size = 50)
pc_eclust_interaction <- r_prepare_data(data = cbind(cluster_res$clustersAddon$PC,</pre>
                                                       survival = Y[trainIndex],
                                                       subtype = E[trainIndex]),
                                          response = "survival", exposure = "subtype")
names(pc_eclust_interaction)
dim(pc_eclust_interaction$X)
pc_eclust_interaction$main_effect_names
pc_eclust_interaction$interaction_names
## End(Not run)
```

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simdata

Simulated Data with Environment Dependent Correlations

# Description

A dataset containing simulated data for example use of the eclust package functions. This data was generated using the s\_modules and s\_generate\_data

# Usage

simdata

### **Format**

A matrix with 100 rows and 502 variables:

Y continuous response vector

E binary environment variable for ECLUST method. E = 0 for unexposed (n=50) and E = 1 for exposed (n=50)

columns 3:502 gene expression data for 1000 genes. column names are the gene names

### Note

Code used to generate this data can be found on the GitHub page for this package. See URL below.

#### **Source**

 $https://raw.githubusercontent.com/sahirbhatnagar/eclust/master/data-raw/simulated-data-processing. \\ R$ 

### References

Bhatnagar, SR., Yang, Y., Blanchette, M., Bouchard, L., Khundrakpam, B., Evans, A., Greenwood, CMT. (2016+). An analytic approach for interpretable predictive models in high dimensional data, in the presence of interactions with exposures *Preprint* 

```
simdata[1:5, 1:10]
table(simdata[,"E"])
```

s_generate_data	Generate linear response data and test and training sets for simulation study
	stuay

### **Description**

create a function that takes as input, the number of genes, the true beta vector, the gene expression matrix created from the generate\_blocks function and returns a list of data matrix, as well as correlation matrices, TOM matrices, cluster information, training and test data

# Usage

```
s_generate_data(p, X, beta, binary_outcome = FALSE,
   cluster_distance = c("corr", "corr0", "corr1", "tom", "tom0", "tom1",
   "diffcorr", "difftom", "corScor", "tomScor", "fisherScore"), n, n0,
   include_interaction = F, signal_to_noise_ratio = 1,
   eclust_distance = c("fisherScore", "corScor", "diffcorr", "difftom"),
   cluster_method = c("hclust", "protoclust"), cut_method = c("dynamic",
   "gap", "fixed"), distance_method = c("euclidean", "maximum", "manhattan",
   "canberra", "binary", "minkowski"), n_clusters,
   agglomeration_method = c("complete", "average", "ward.D2", "single",
   "ward.D", "mcquitty", "median", "centroid"), nPC = 1, K.max = 10,
   B = 10)
```

# **Arguments**

p number of genes in design matrix Χ gene expression matrix of size n x p using the generate\_blocks function true beta coefficient vector beta binary\_outcome Logical. Should a binary outcome be generated. Default is FALSE. See details on how a binary outcome is generated cluster\_distance character representing which matrix from the training set that you want to use to cluster the genes. Must be one of the following • corr, corr0, corr1, tom, tom0, tom1, diffcorr, difftom, corScor, tomScor, fisherScore total number of subjects n total number of subjects with E=0 include\_interaction Should an interaction with the environment be generated as part of the response.

signal\_to\_noise\_ratio

signal to noise ratio, default is 1

Default is FALSE.

eclust\_distance

character representing which matrix from the training set that you want to use to cluster the genes based on the environment. See cluster\_distance for avaiable options. Should be different from cluster\_distance. For example, if cluster\_distance=corr and EclustDistance=fisherScore. That is, one should be based on correlations ignoring the environment, and the other should be based on correlations accounting for the environment. This function will always return this add on

cluster\_method Cluster the data using hierarchical clustering or prototype clustering. Defaults

cluster\_method="hclust". Other option is protoclust, however this pack-

age must be installed before proceeding with this option

cut\_method what method to use to cut the dendrogram. 'dynamic' refers to dynamicTreeCut

library. 'gap' is Tibshirani's gap statistic clusGap using the 'Tibs2001SEmax' rule. 'fixed' is a fixed number specified by the n\_clusters argument

distance\_method

one of "euclidean", "maximum", "manhattan", "canberra", "binary", "minkowski" to be passed to dist function.

agglomeration\_method

the agglomeration method to be used. This should be (an unambiguous abbreviation of) one of "ward.D", "ward.D2", "single", "complete", "average" (= UPGMA), "mcquitty" (= WPGMA), "median" (= WPGMC) or "centroid" (=

UPGMC).

nPC number of principal components to extract from each cluster. Default is 1. Only

1 or 2 is allowed.

K.max the maximum number of clusters to consider, must be at least two. Only used if

cutMethod='gap'

B integer, number of Monte Carlo ("bootstrap") samples. Only used if cutMethod='gap'

# **Details**

To generate a binary outcome we first generate a continuous outcome Y which is  $X^T\beta$ , defined p=1/(1+exp(-Y)) and used this to generate a two-class outcome z with Pr(z=1)=p and Pr(z=0)=1-p.

# Value

list of (in the following order)

beta\_truth a 1 column matrix containing the true beta coefficient vector

**similarity** an object of class similarity which is the similarity matrix specified by the cluster\_distance argument

similarityEclust an object of class similarity which is the similarity matrix specified by the eclust\_distance argument

**DT** data.table of simulated data from the s\_response function

Y The simulated response

**X0** the n0 x p design matrix for the unexposed subjects

X1 the n1 x p design matrix for the exposed subjects

**X\_train** the training design matrix for all subjects

**X\_test** the test set design matrix for all subjects

Y train the training set response

**Y\_test** the test set response

**DT** train the training response and training design matrix in a single data.frame object

DT\_test the test response and training design matrix in a single data.frame object

S0 a character vector of the active genes i.e. the ones that are associated with the response

**n\_clusters\_All** the number of clusters identified by using the similarity matrix specified by the cluster\_distance argument

n\_clusters\_Eclust the number of clusters identified by using the similarity matrix specified by the eclust\_distance argument

n\_clusters\_Addon the sum of n\_clusters\_All and n\_clusters\_Eclust

clustersAll the cluster membership of each gene based on the cluster\_distance matrix

**clustersAddon** the cluster membership of each gene based on both the cluster\_distance matrix and the eclust\_distance matrix. Note that each gene will appear twice here

**clustersEclust** the cluster membership of each gene based on the eclust\_distance matrix

gene\_groups\_inter cluster membership of each gene with a penalty factor used for the group lasso

gene\_groups\_inter\_Addon cluster membership of each gene with a penalty factor used for the group lasso

tom train all the TOM matrix based on all training subjects

tom\_train\_diff the absolute difference of the exposed and unexposed TOM matrices:  $|TOM_{E=1} - TOM_{E=0}|$ 

tom\_train\_e1 the TOM matrix based on training exposed subjects only

tom train e0 the TOM matrix based on training unexposed subjects only

corr train all the Pearson correlation matrix based on all training subjects

**corr\_train\_diff** the absolute difference of the exposed and unexposed Pearson correlation matrices:  $|Cor_{E=1} - Cor_{E=0}|$ 

corr\_train\_e1 the Pearson correlation matrix based on training exposed subjects only

corr\_train\_e0 the Pearson correlation matrix based on training unexposed subjects only

fisherScore The fisher scoring matrix. see u\_fisherZ for details

**corScor** The correlation scoring matrix:  $|Cor_{E=1} + Cor_{E=0} - 2|$ 

mse\_null The MSE for the null model

**DT\_train\_folds** The 10 training folds used for the stability measures

**X\_train\_folds** The 10 X training folds (the same as in DT\_train\_folds)

**Y\_train\_folds** The 10 Y training folds (the same as in DT\_train\_folds)

#### Note

this function calls the s\_response to generate phenotype as a function of the gene expression data. This function also returns other information derived from the simulated data including the test and training sets, the correlation and TOM matrices and the clusters.

the PCs and averages need to be calculated in the fitting functions, because these will change based on the CV fold

```
library(magrittr)
# simulation parameters
rho = 0.90; p = 500; SNR = 1; n = 200; n0 = n1 = 100; nActive = p*0.10; cluster_distance = "tom";
Ecluster_distance = "difftom"; rhoOther = 0.6; betaMean = 2;
alphaMean = 1; betaE = 3; distanceMethod = "euclidean"; clustMethod = "hclust";
cutMethod = "dynamic"; agglomerationMethod = "average"
#in this simulation its blocks 3 and 4 that are important
#leaveOut: optional specification of modules that should be left out
#of the simulation, that is their genes will be simulated as unrelated
#("grey"). This can be useful when simulating several sets, in some which a module
#is present while in others it is absent.
d0 <- s_modules(n = n0, p = p, rho = 0, exposed = FALSE,
                modProportions = c(0.15, 0.15, 0.15, 0.15, 0.15, 0.25),
                minCor = 0.01,
                maxCor = 1,
                corPower = 1,
                propNegativeCor = 0.3,
                backgroundNoise = 0.5,
                signed = FALSE,
                leaveOut = 1:4)
d1 <- s_{modules}(n = n1, p = p, rho = rho, exposed = TRUE,
                modProportions = c(0.15, 0.15, 0.15, 0.15, 0.15, 0.25),
                minCor = 0.4,
                maxCor = 1,
                corPower = 0.3,
                propNegativeCor = 0.3,
                backgroundNoise = 0.5,
                signed = FALSE)
truemodule1 <- d1$setLabels
X <- rbind(d0$datExpr, d1$datExpr) %>%
  magrittr::set_colnames(paste0("Gene", 1:p)) %>%
  magrittr::set_rownames(paste0("Subject",1:n))
betaMainEffect <- vector("double", length = p)</pre>
betaMainInteractions <- vector("double", length = p)</pre>
# the first nActive/2 in the 3rd block are active
```

```
betaMainEffect[which(truemodule1 %in% 3)[1:(nActive/2)]] <- runif(</pre>
 nActive/2, betaMean - 0.1, betaMean + 0.1)
# the first nActive/2 in the 4th block are active
betaMainEffect[which(truemodule1 %in% 4)[1:(nActive/2)]] <- runif(</pre>
 nActive/2, betaMean+2 - 0.1, betaMean+2 + 0.1)
betaMainInteractions[which(betaMainEffect!=0)] <- runif(nActive, alphaMean - 0.1, alphaMean + 0.1)
beta <- c(betaMainEffect, betaE, betaMainInteractions)</pre>
## Not run:
result <- s_generate_data(p = p, X = X,
                           beta = beta,
                           include_interaction = TRUE,
                           cluster_distance = cluster_distance,
                           n = n, n0 = n0,
                           eclust_distance = Ecluster_distance,
                           signal_to_noise_ratio = SNR,
                           distance_method = distanceMethod,
                           cluster_method = clustMethod,
                           cut_method = cutMethod,
                           agglomeration_method = agglomerationMethod,
                           nPC = 1
names(result)
## End(Not run)
```

 ${\tt s\_generate\_data\_mars} \quad \textit{Generate non linear response and test and training sets for non-linear} \\ simulation \textit{study}$ 

# **Description**

create a function that takes as input, the number of genes, the true beta vector, the gene expression matrix created from the generate\_blocks function and returns a list of data matrix, as well as correlation matrices, TOM matrices, cluster information, training and test data

### **Usage**

```
s_generate_data_mars(p, X, beta, binary_outcome = FALSE, truemodule, nActive,
  cluster_distance = c("corr", "corr0", "corr1", "tom", "tom0", "tom1",
  "diffcorr", "difftom", "corScor", "tomScor", "fisherScore"), n, n0,
  include_interaction = F, signal_to_noise_ratio = 1,
  eclust_distance = c("fisherScore", "corScor", "diffcorr", "difftom"),
  cluster_method = c("hclust", "protoclust"), cut_method = c("dynamic",
  "gap", "fixed"), distance_method = c("euclidean", "maximum", "manhattan",
  "canberra", "binary", "minkowski"), n_clusters,
  agglomeration_method = c("complete", "average", "ward.D2", "single",
  "ward.D", "mcquitty", "median", "centroid"), nPC = 1, K.max = 10,
  B = 10)
```

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#### **Arguments**

number of genes in design matrix р

Χ gene expression matrix of size n x p using the generate\_blocks function

beta true beta coefficient vector

Logical. Should a binary outcome be generated. Default is FALSE. See details binary\_outcome

on how a binary outcome is generated

truemodule numeric vector of the true module membership used in the s\_response\_mars

function. Modules 3 and 4 are active in the response. See s\_response\_mars

function for details.

nActive number of active genes in the response used in the s\_response\_mars

cluster\_distance

character representing which matrix from the training set that you want to use to cluster the genes. Must be one of the following

• corr, corr0, corr1, tom, tom0, tom1, diffcorr, difftom, corScor, tomScor, fisherScore

total number of subjects n

total number of subjects with E=0 n0

include\_interaction

Should an interaction with the environment be generated as part of the response. Default is FALSE.

signal\_to\_noise\_ratio

signal to noise ratio, default is 1

eclust\_distance

character representing which matrix from the training set that you want to use to cluster the genes based on the environment. See cluster\_distance for avaialble options. Should be different from cluster\_distance. For example, if cluster\_distance=corr and EclustDistance=fisherScore. That is, one should be based on correlations ignoring the environment, and the other should be based on correlations accounting for the environment. This function will always return this add on

cluster\_method Cluster the data using hierarchical clustering or prototype clustering. Defaults cluster\_method="hclust". Other option is protoclust, however this package must be installed before proceeding with this option

cut\_method

what method to use to cut the dendrogram. 'dynamic' refers to dynamicTreeCut library. 'gap' is Tibshirani's gap statistic clusGap using the 'Tibs2001SEmax' rule. 'fixed' is a fixed number specified by the n\_clusters argument

distance\_method

one of "euclidean", "maximum", "manhattan", "canberra", "binary", "minkowski" to be passed to dist function.

n clusters

Number of clusters specified by the user. Only applicable when cut\_method="fixed" agglomeration\_method

> the agglomeration method to be used. This should be (an unambiguous abbreviation of) one of "ward.D", "ward.D2", "single", "complete", "average" (= UPGMA), "mcquitty" (= WPGMA), "median" (= WPGMC) or "centroid" (= UPGMC).

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nPC number of principal components. Can be 1 or 2.

K. max the maximum number of clusters to consider, must be at least two. Only used if

cutMethod='gap'

B integer, number of Monte Carlo ("bootstrap") samples. Only used if cutMethod='gap'

#### Value

list of (in the following order)

beta\_truth a 1 column matrix containing the true beta coefficient vector

similarity an object of class similarity which is the similarity matrix specified by the cluster\_distance argument

**similarityEclust** an object of class similarity which is the similarity matrix specified by the eclust\_distance argument

DT data.table of simulated data from the s\_response function

Y The simulated response

**X0** the n0 x p design matrix for the unexposed subjects

**X1** the n1 x p design matrix for the exposed subjects

**X\_train** the training design matrix for all subjects

X test the test set design matrix for all subjects

**Y\_train** the training set response

**Y\_test** the test set response

**DT\_train** the training response and training design matrix in a single data.frame object

DT\_test the test response and training design matrix in a single data.frame object

**S0** a character vector of the active genes i.e. the ones that are associated with the response

n\_clusters\_All the number of clusters identified by using the similarity matrix specified by the cluster\_distance argument

n\_clusters\_Eclust the number of clusters identified by using the similarity matrix specified by the eclust\_distance argument

n\_clusters\_Addon the sum of n\_clusters\_All and n\_clusters\_Eclust

clustersAll the cluster membership of each gene based on the cluster\_distance matrix

**clustersAddon** the cluster membership of each gene based on both the cluster\_distance matrix and the eclust\_distance matrix. Note that each gene will appear twice here

**clustersEclust** the cluster membership of each gene based on the eclust\_distance matrix

gene\_groups\_inter cluster membership of each gene with a penalty factor used for the group lasso

**gene\_groups\_inter\_Addon** cluster membership of each gene with a penalty factor used for the group lasso

tom\_train\_all the TOM matrix based on all training subjects

tom\_train\_diff the absolute difference of the exposed and unexposed TOM matrices:  $|TOM_{E=1} - TOM_{E=0}|$ 

tom\_train\_e1 the TOM matrix based on training exposed subjects only

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```
tom_train_e0 the TOM matrix based on training unexposed subjects only
corr_train_all the Pearson correlation matrix based on all training subjects
corr_train_diff the absolute difference of the exposed and unexposed Pearson correlation matrices: |Cor_{E=1} - Cor_{E=0}|
corr_train_e1 the Pearson correlation matrix based on training exposed subjects only
corr_train_e0 the Pearson correlation matrix based on training unexposed subjects only
fisherScore The fisher scoring matrix. see u_fisherZ for details
corScor The correlation scoring matrix: |Cor_{E=1} + Cor_{E=0} - 2|
mse_null The MSE for the null model
DT_train_folds The 10 training folds used for the stability measures
X_train_folds The 10 X training folds (the same as in DT_train_folds)
Y_train_folds The 10 Y training folds (the same as in DT_train_folds)
```

### **Examples**

truemodule1 <- d1\$setLabels

```
library(magrittr)
# simulation parameters
rho = 0.90; p = 500; SNR = 1; n = 200; n0 = n1 = 100; nActive = p*0.10; cluster_distance = "tom";
Ecluster_distance = "difftom"; rhoOther = 0.6; betaMean = 2;
alphaMean = 1; betaE = 3; distanceMethod = "euclidean"; clustMethod = "hclust";
cutMethod = "dynamic"; agglomerationMethod = "average"
#in this simulation its blocks 3 and 4 that are important
#leaveOut: optional specification of modules that should be left out
#of the simulation, that is their genes will be simulated as unrelated
#("grey"). This can be useful when simulating several sets, in some which a module
#is present while in others it is absent.
d0 < -s_{modules}(n = n0, p = p, rho = 0, exposed = FALSE,
                modProportions = c(0.15, 0.15, 0.15, 0.15, 0.15, 0.25),
                minCor = 0.01,
                maxCor = 1,
                corPower = 1,
                propNegativeCor = 0.3,
                backgroundNoise = 0.5,
                signed = FALSE,
                leaveOut = 1:4)
d1 <- s_modules(n = n1, p = p, rho = rho, exposed = TRUE,</pre>
                modProportions = c(0.15, 0.15, 0.15, 0.15, 0.15, 0.25),
                minCor = 0.4,
                maxCor = 1,
                corPower = 0.3,
                propNegativeCor = 0.3,
                backgroundNoise = 0.5,
                signed = FALSE)
```

```
X <- rbind(d0$datExpr, d1$datExpr) %>%
  magrittr::set_colnames(paste0("Gene", 1:p)) %>%
  magrittr::set_rownames(paste0("Subject",1:n))
betaMainEffect <- vector("double", length = p)</pre>
# the first nActive/2 in the 3rd block are active
betaMainEffect[which(truemodule1 %in% 3)[1:(nActive/2)]] <- runif(</pre>
  nActive/2, betaMean - 0.1, betaMean + 0.1)
# the first nActive/2 in the 4th block are active
betaMainEffect[which(truemodule1 %in% 4)[1:(nActive/2)]] <- runif(</pre>
  nActive/2, betaMean+2 - 0.1, betaMean+2 + 0.1)
beta <- c(betaMainEffect, betaE)</pre>
result <- s_generate_data_mars(p = p, X = X,
                                beta = beta,
                                binary_outcome = FALSE,
                                truemodule = truemodule1,
                                nActive = nActive,
                                include_interaction = FALSE,
                                cluster_distance = cluster_distance,
                                n = n, n0 = n0,
                                eclust_distance = Ecluster_distance,
                                signal_to_noise_ratio = SNR,
                                distance_method = distanceMethod,
                                cluster_method = clustMethod,
                                cut_method = cutMethod,
                                agglomeration_method = agglomerationMethod,
                                nPC = 1)
names(result)
```

 $s_mars_clust$ 

Fit MARS Models on Simulated Cluster Summaries

### **Description**

This function creates summaries of the given clusters (e.g. 1st PC or average), and then runs Friedman's MARS on those summaries. To be used with simulated data where the 'truth' is known i.e., you know which features are associated with the response. This function was used to produce the simulation results in Bhatnagar et al. 2016.

# Usage

```
s_mars_clust(x_train, x_test, y_train, y_test, s0, summary = c("pc", "avg"),
model = c("MARS"), exp_family = c("gaussian", "binomial"), gene_groups,
true_beta = NULL, topgenes = NULL, stability = F, filter = F,
include_E = T, include_interaction = F, clust_type = c("CLUST",
    "ECLUST"), nPC = 1)
```

#### **Arguments**

ntrain x p matrix of simulated training set where ntrain is the number of trainx\_train ing observations and p is total number of predictors. This matrix needs to have named columns representing the feature names or the gene names ntest x p matrix of simulated training set where ntest is the number of training  $x_{test}$ observations and p is total number of predictors. This matrix needs to have named columns representing the feature names or the gene names y\_train numeric vector of length ntrain representing the responses for the training subjects. If continuous then you must set exp\_family = "gaussion". For exp\_family="binomial" should be either a factor with two levels, or a twocolumn matrix of counts or proportions (the second column is treated as the target class; for a factor, the last level in alphabetical order is the target class) numeric vector of length ntest representing the responses for the test subjects. y\_test If continuous then you must set exp\_family = "gaussion". For exp\_family="binomial" should be either a factor with two levels, or a two-column matrix of counts or proportions (the second column is treated as the target class; for a factor, the last level in alphabetical order is the target class). s0 chracter vector of the active feature names, i.e., the features in x\_train that are truly associated with the response. the summary of each cluster. Can be the principal component or average. Desummary fault is summary = "pc" which takes the first number\_pc principal components. Currently a maximum of 2 principal components can be chosen. mode1 Type of non-linear model to be fit. Currently only Friedman's MARS is supported. exp\_family Response type. See details for y\_train argument above. gene\_groups data.frame that contains the group membership for each feature. The first column is called 'gene' and the second column should be called 'cluster'. The 'gene' column identifies the features and must be the same identifiers in the x\_train, x\_test matrices. The 'cluster' column is a numeric integer indicating the cluster group membership. A cluster group membership of 0 implies the feature did not cluster into any group. true\_beta numeric vector of true beta coefficients topgenes List of features to keep if filter=TRUE. Default is topgenes = NULL which means all features are kept for the analysis Should stability measures be calculated. Default is stability=FALSE. See destability tails filter Should analysis be run on a subset of features. Default is filter = FALSE include\_E Should the environment variable be included in the regression analysis. Default is include\_E = TRUE include\_interaction

Should interaction effects between the features in x\_train and the environment

variable be fit. Default is include\_interaction=TRUE

clust\_type Method used to cluster the features. This is used for naming the output only

and has no consequence for the results. clust\_type = "CLUST" is the default which means that the environment varible was not used in the clustering step. clust\_type = "ECLUST" means that the environment variable was used in the

clustering aspect.

nPC Number of principal components if summary = "pc". Default is nPC = 1. Can be

either 1 or 2.

#### **Details**

This function first does 10 fold cross-validation to tune the degree (either 1 or 2) using the train function with method="earth" and nprune is fixed at 1000. Then the earth function is used, with nk = 1000 and pmethod = "backward" to fit the MARS model using the optimal degree from the 10-fold CV.

#### Value

This function has two different outputs depending on whether stability = TRUE or Stability = FALSE

If stability = TRUE then this function returns a p x 2 data.frame or data.table of regression coefficients without the intercept. The output of this is used for subsequent calculations of stability.

If stability = FALSE then returns a vector with the following elements (See Table 3: Measures of Performance in Bhatnagar et al (2016+) for definitions of each measure of performance):

mse or AUC Test set mean squared error if exp\_family = "gaussion" or test set Area under

the curve if exp\_family = "binomial" calculated using the roc function

RMSE Square root of the mse. Only applicable if exp\_family = "gaussion"

Shat Number of non-zero estimated regression coefficients. The non-zero estimated

regression coefficients are referred to as being selected by the model

TPR true positive rate

FPR false positive rate

Correct Sparsity

Correct true positives + correct true negative coefficients divided by the total

number of features

CorrectZeroMain

Proportion of correct true negative main effects

CorrectZeroInter

Proportion of correct true negative interactions

IncorrectZeroMain

Proportion of incorrect true negative main effects

IncorrectZeroInter

Proportion of incorrect true negative interaction effects

```
## Not run:
library(magrittr)
# simulation parameters
rho = 0.90; p = 500; snr = 1; rn = 200; rnr = 100; rnr = 1
Ecluster_distance = "difftom"; rhoOther = 0.6; betaMean = 2;
alphaMean = 1; betaE = 3; distanceMethod = "euclidean"; clustMethod = "hclust";
cutMethod = "dynamic"; agglomerationMethod = "average"
#in this simulation its blocks 3 and 4 that are important
#leaveOut: optional specification of modules that should be left out
#of the simulation, that is their genes will be simulated as unrelated
#("grey"). This can be useful when simulating several sets, in some which a module
#is present while in others it is absent.
d0 < -s_{modules}(n = n0, p = p, rho = 0, exposed = FALSE,
                               modProportions = c(0.15, 0.15, 0.15, 0.15, 0.15, 0.25),
                                minCor = 0.01,
                               maxCor = 1,
                                corPower = 1,
                                propNegativeCor = 0.3,
                                backgroundNoise = 0.5,
                                signed = FALSE,
                                leaveOut = 1:4)
d1 <- s_modules(n = n1, p = p, rho = rho, exposed = TRUE,
                                modProportions = c(0.15, 0.15, 0.15, 0.15, 0.15, 0.25),
                                minCor = 0.4,
                                maxCor = 1.
                                corPower = 0.3,
                                propNegativeCor = 0.3,
                                backgroundNoise = 0.5,
                                signed = FALSE)
truemodule1 <- d1$setLabels
X <- rbind(d0$datExpr, d1$datExpr) %>%
    magrittr::set_colnames(paste0("Gene", 1:p)) %>%
    magrittr::set_rownames(paste0("Subject",1:n))
betaMainEffect <- vector("double", length = p)</pre>
# the first nActive/2 in the 3rd block are active
betaMainEffect[which(truemodule1 %in% 3)[1:(nActive/2)]] <- runif(</pre>
    nActive/2, betaMean - 0.1, betaMean + 0.1)
# the first nActive/2 in the 4th block are active
betaMainEffect[which(truemodule1 %in% 4)[1:(nActive/2)]] <- runif(</pre>
    nActive/2, betaMean+2 - 0.1, betaMean+2 + 0.1)
beta <- c(betaMainEffect, betaE)</pre>
result <- s_generate_data_mars(p = p, X = X,
```

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```
beta = beta,
                                binary_outcome = FALSE,
                                truemodule = truemodule1,
                                nActive = nActive,
                                include_interaction = FALSE,
                                cluster_distance = cluster_distance,
                                n = n, n0 = n0,
                                eclust_distance = Ecluster_distance,
                                signal_to_noise_ratio = SNR,
                                distance_method = distanceMethod,
                                cluster_method = clustMethod,
                                cut_method = cutMethod,
                                agglomeration_method = agglomerationMethod,
                                nPC = 1)
mars_res <- s_mars_clust(x_train = result[["X_train"]],</pre>
                         x_test = result[["X_test"]],
                         y_train = result[["Y_train"]],
                         y_test = result[["Y_test"]],
                          s0 = result[["S0"]],
                          summary = "pc",
                          exp_family = "gaussian",
                          gene_groups = result[["clustersAddon"]],
                          clust_type = "ECLUST")
unlist(mars_res)
## End(Not run)
```

s\_mars\_separate

Fit Multivariate Adaptive Regression Splines on Simulated Data

# **Description**

This function can run Friedman's MARS models on the untransformed design matrix. To be used with simulated data where the 'truth' is known i.e., you know which features are associated with the response. This function was used to produce the simulation results in Bhatnagar et al. 2016. Uses caret functions to tune the degree and the nprune parameters

# Usage

```
s_mars_separate(x_train, x_test, y_train, y_test, s0, model = c("MARS"),
  exp_family = c("gaussian", "binomial"), topgenes = NULL, stability = F,
  filter = F, include_E = T, include_interaction = F, ...)
```

### **Arguments**

x\_train

ntrain x p matrix of simulated training set where ntrain is the number of training observations and p is total number of predictors. This matrix needs to have named columns representing the feature names or the gene names

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x_test	ntest x p matrix of simulated training set where ntest is the number of training observations and p is total number of predictors. This matrix needs to have named columns representing the feature names or the gene names
y_train	numeric vector of length ntrain representing the responses for the training subjects. If continuous then you must set exp_family = "gaussion". For exp_family="binomial" should be either a factor with two levels, or a two-column matrix of counts or proportions (the second column is treated as the target class; for a factor, the last level in alphabetical order is the target class)
y_test	numeric vector of length ntest representing the responses for the test subjects. If continuous then you must set exp_family = "gaussion". For exp_family="binomial" should be either a factor with two levels, or a two-column matrix of counts or proportions (the second column is treated as the target class; for a factor, the last level in alphabetical order is the target class).
s0	chracter vector of the active feature names, i.e., the features in x_train that are truly associated with the response.
model	Type of non-linear model to be fit. Currently only Friedman's MARS is supported.
exp_family	Response type. See details for y_train argument above.
topgenes	List of features to keep if filter=TRUE. Default is topgenes = NULL which means all features are kept for the analysis
stability	Should stability measures be calculated. Default is stability=FALSE. See details
filter	Should analysis be run on a subset of features. Default is filter = FALSE
include_E	Should the environment variable be included in the regression analysis. Default is include_E = TRUE
include_interac	
	Should interaction effects between the features in x_train and the environment variable be fit. Default is include_interaction=TRUE
	other parameters passed to trainControl function

# **Details**

This function first does 10 fold cross-validation to tune the degree (either 1 or 2) using the train function with method="earth" and nprune is fixed at 1000. Then the earth function is used, with nk = 1000 and pmethod = "backward" to fit the MARS model using the optimal degree from the 10-fold CV.

# Value

This function has two different outputs depending on whether stability = TRUE or Stability = TRUE or Stability = TRUE

If stability = TRUE then this function returns a p x 2 data.frame or data.table of regression coefficients without the intercept. The output of this is used for subsequent calculations of stability.

If stability = FALSE then returns a vector with the following elements (See Table 3: Measures of Performance in Bhatnagar et al (2016+) for definitions of each measure of performance):

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mse or AUC Test set mean squared error if exp\_family = "gaussion" or test set Area under

the curve if exp\_family = "binomial" calculated using the roc function

RMSE Square root of the mse. Only applicable if exp\_family = "gaussion"

Shat Number of non-zero estimated regression coefficients. The non-zero estimated

regression coefficients are referred to as being selected by the model

TPR true positive rate
FPR false positive rate

Correct Sparsity

Correct true positives + correct true negative coefficients divided by the total

number of features

CorrectZeroMain

Proportion of correct true negative main effects

CorrectZeroInter

Proportion of correct true negative interactions

IncorrectZeroMain

Proportion of incorrect true negative main effects

IncorrectZeroInter

Proportion of incorrect true negative interaction effects

```
## Not run:
library(magrittr)
# simulation parameters
rho = 0.90; p = 500; SNR = 1; n = 200; n0 = n1 = 100; nActive = p*0.10; cluster_distance = "tom";
Ecluster_distance = "difftom"; rhoOther = 0.6; betaMean = 2;
alphaMean = 1; betaE = 3; distanceMethod = "euclidean"; clustMethod = "hclust";
cutMethod = "dynamic"; agglomerationMethod = "average"
#in this simulation its blocks 3 and 4 that are important
#leaveOut: optional specification of modules that should be left out
#of the simulation, that is their genes will be simulated as unrelated
#("grey"). This can be useful when simulating several sets, in some which a module
#is present while in others it is absent.
d0 <- s_modules(n = n0, p = p, rho = 0, exposed = FALSE,
                modProportions = c(0.15, 0.15, 0.15, 0.15, 0.15, 0.25),
                minCor = 0.01,
                maxCor = 1,
                corPower = 1,
                propNegativeCor = 0.3,
                backgroundNoise = 0.5,
                signed = FALSE,
                leaveOut = 1:4)
d1 <- s_modules(n = n1, p = p, rho = rho, exposed = TRUE,</pre>
                modProportions = c(0.15, 0.15, 0.15, 0.15, 0.15, 0.25),
                minCor = 0.4,
                maxCor = 1,
```

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```
corPower = 0.3,
                propNegativeCor = 0.3,
                backgroundNoise = 0.5,
                signed = FALSE)
truemodule1 <- d1$setLabels
X <- rbind(d0$datExpr, d1$datExpr) %>%
  magrittr::set_colnames(paste0("Gene", 1:p)) %>%
  magrittr::set_rownames(paste0("Subject",1:n))
betaMainEffect <- vector("double", length = p)</pre>
# the first nActive/2 in the 3rd block are active
betaMainEffect[which(truemodule1 %in% 3)[1:(nActive/2)]] <- runif(</pre>
  nActive/2, betaMean - 0.1, betaMean + 0.1)
# the first nActive/2 in the 4th block are active
betaMainEffect[which(truemodule1 %in% 4)[1:(nActive/2)]] <- runif(</pre>
  nActive/2, betaMean+2 - 0.1, betaMean+2 + 0.1)
beta <- c(betaMainEffect, betaE)</pre>
result <- s_generate_data_mars(p = p, X = X,
                                beta = beta,
                                binary_outcome = FALSE,
                                truemodule = truemodule1,
                                nActive = nActive,
                                include_interaction = FALSE,
                                cluster_distance = cluster_distance,
                                n = n, n0 = n0,
                                eclust_distance = Ecluster_distance,
                                signal_to_noise_ratio = SNR,
                                distance_method = distanceMethod,
                                cluster_method = clustMethod,
                                cut_method = cutMethod,
                                agglomeration_method = agglomerationMethod,
                                nPC = 1)
mars_res <- s_mars_separate(x_train = result[["X_train"]],</pre>
                             x_test = result[["X_test"]],
                             y_train = result[["Y_train"]],
                             y_test = result[["Y_test"]],
                             s0 = result[["S0"]],
                             exp_family = "gaussian")
unlist(mars_res)
## End(Not run)
```

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# **Description**

This is a wrapper of the simulateDatExpr function which simulates data in a modular structure (i.e. in blocks). This function simulates data in 5 blocks referred to as Turquoise, Blue, Red, Green and Yellow, separately for exposed (E=1) and unexposed (E=0) observations.

# Usage

```
s_modules(n, p, rho, exposed, ...)
```

### **Arguments**

n	number of observations
р	total number of predictors to simulate
rho	numeric value representing the expected correlation between green module and red module
exposed	binary numeric vector of length n with 0 for unexposed and 1 for exposed
	arguments passed to the simulateDatExpr function

#### Value

n x p matrix of simulated data

```
library(magrittr)
p <- 1000
n <- 200
d\theta <- s_{modules}(n = 100, p = 1000, rho = 0, exposed = FALSE,
                modProportions = c(0.15, 0.15, 0.15, 0.15, 0.15, 0.25),
                minCor = 0.01,
                maxCor = 1,
                corPower = 1,
                propNegativeCor = 0.3,
                backgroundNoise = 0.5,
                signed = FALSE,
                leaveOut = 1:4)
d1 <- s_{modules}(n = 100, p = 1000, rho = 0.90, exposed = TRUE,
                modProportions = c(0.15, 0.15, 0.15, 0.15, 0.15, 0.25),
                minCor = 0.4,
                maxCor = 1,
                corPower = 0.3,
                propNegativeCor = 0.3,
                backgroundNoise = 0.5,
                 signed = FALSE)
X <- rbind(d0$datExpr, d1$datExpr) %>%
magrittr::set_colnames(paste0("Gene", 1:p)) %>%
magrittr::set_rownames(paste0("Subject",1:n))
dim(X)
```

s\_pen\_clust

Fit Penalized Regression Models on Simulated Cluster Summaries

### **Description**

This function creates summaries of the given clusters (e.g. 1st PC or average), and then fits a penalized regression model on those summaries. To be used with simulated data where the 'truth' is known i.e., you know which features are associated with the response. This function was used to produce the simulation results in Bhatnagar et al. 2016. Can run lasso, elasticnet, SCAD or MCP models

# Usage

```
s_pen_clust(x_train, x_test, y_train, y_test, s0, gene_groups,
  summary = c("pc", "avg"), model = c("lasso", "elasticnet", "scad", "mcp"),
  exp_family = c("gaussian", "binomial"), filter = F, topgenes = NULL,
  stability = F, include_E = T, include_interaction = F,
  clust_type = c("CLUST", "ECLUST"), number_pc = 1)
```

# **Arguments**

x_train	ntrain x p matrix of simulated training set where ntrain is the number of training observations and p is total number of predictors. This matrix needs to have named columns representing the feature names or the gene names
x_test	ntest x p matrix of simulated training set where ntest is the number of training observations and p is total number of predictors. This matrix needs to have named columns representing the feature names or the gene names
y_train	numeric vector of length ntrain representing the responses for the training subjects. If continuous then you must set exp_family = "gaussion". For exp_family="binomial" should be either a factor with two levels, or a two-column matrix of counts or proportions (the second column is treated as the target class; for a factor, the last level in alphabetical order is the target class)
y_test	numeric vector of length ntest representing the responses for the test subjects. If continuous then you must set exp_family = "gaussion". For exp_family="binomial" should be either a factor with two levels, or a two-column matrix of counts or proportions (the second column is treated as the target class; for a factor, the last level in alphabetical order is the target class).
s0	chracter vector of the active feature names, i.e., the features in x_train that are truly associated with the response.
gene_groups	data.frame that contains the group membership for each feature. The first column is called 'gene' and the second column should be called 'cluster'. The 'gene' column identifies the features and must be the same identifiers in the x_train,x_test matrices. The 'cluster' column is a numeric integer indicating the cluster group membership. A cluster group membership of 0 implies the feature did not cluster into any group.

summary the summary of each cluster. Can be the principal component or average. De-

fault is summary = "pc" which takes the first number\_pc principal components.

Currently a maximum of 2 principal components can be chosen.

model Regression model to be fit on cluster summaries. Default is model="lasso"

which corresponds to glmnet mixing parameter alpha=1. model="elasticnet"

corresponds to glmnet mixing parameter alpha=0.5, model="mcp" and model="scad"

are the non-convex models from the novreg package

exp\_family Response type. See details for y\_train argument above.

filter Should analysis be run on a subset of features. Default is filter = FALSE

topgenes List of features to keep if filter=TRUE. Default is topgenes = NULL which

means all features are kept for the analysis

stability Should stability measures be calculated. Default is stability=FALSE. See de-

tails

include\_E Should the environment variable be included in the regression analysis. Default

is include\_E = TRUE

include\_interaction

Should interaction effects between the features in x\_train and the environment

variable be fit. Default is include\_interaction=TRUE

clust\_type Method used to cluster the features. This is used for naming the output only

and has no consequence for the results. clust\_type = "CLUST" is the default which means that the environment varible was not used in the clustering step. clust\_type = "ECLUST" means that the environment variable was used in the

clustering aspect.

number\_pc Number of principal components if summary = "pc". Default is number\_pc = 1.

Can be either 1 or 2.

#### **Details**

The stability of feature importance is defined as the variability of feature weights under perturbations of the training set, i.e., small modifications in the training set should not lead to considerable changes in the set of important covariates (Toloşi, L., & Lengauer, T. (2011)). A feature selection algorithm produces a weight, a ranking, and a subset of features. In the CLUST and ECLUST methods, we defined a predictor to be non-zero if its corresponding cluster representative weight was non-zero. Using 10-fold cross validation (CV), we evaluated the similarity between two features and their rankings using Pearson and Spearman correlation, respectively. For each CV fold we re-ran the models and took the average Pearson/Spearman correlation of the 10 choose 2 combinations of estimated coefficients vectors. To measure the similarity between two subsets of features we took the average of the Jaccard distance in each fold. A Jaccard distance of 1 indicates perfect agreement between two sets while no agreement will result in a distance of 0.

### Value

This function has two different outputs depending on whether stability = TRUE or STADISE

If stability = TRUE then this function returns a p x 2 data.frame or data.table of regression coefficients without the intercept. The output of this is used for subsequent calculations of stability.

If stability = FALSE then returns a vector with the following elements (See Table 3: Measures of Performance in Bhatnagar et al (2016+) for definitions of each measure of performance):

mse or AUC Test set mean squared error if exp\_family = "gaussion" or test set Area under

the curve if exp\_family = "binomial" calculated using the roc function

RMSE Square root of the mse. Only applicable if exp\_family = "gaussion"

Shat Number of non-zero estimated regression coefficients. The non-zero estimated

regression coefficients are referred to as being selected by the model

TPR true positive rate
FPR false positive rate

Correct Sparsity

Correct true positives + correct true negative coefficients divided by the total

number of features

CorrectZeroMain

Proportion of correct true negative main effects

CorrectZeroInter

Proportion of correct true negative interactions

IncorrectZeroMain

Proportion of incorrect true negative main effects

IncorrectZeroInter

Proportion of incorrect true negative interaction effects

nclusters number of estimated clusters by the cutreeDynamic function

# Note

number\_pc=2 will not work if there is only one feature in an estimated cluster

### References

Toloşi, L., & Lengauer, T. (2011). Classification with correlated features: unreliability of feature ranking and solutions. Bioinformatics, 27(14), 1986-1994.

Bhatnagar, SR., Yang, Y., Blanchette, M., Bouchard, L., Khundrakpam, B., Evans, A., Greenwood, CMT. (2016+). An analytic approach for interpretable predictive models in high dimensional data, in the presence of interactions with exposures *Preprint* 

Langfelder, P., Zhang, B., & Horvath, S. (2008). *Defining clusters from a hierarchical cluster tree:* the Dynamic Tree Cut package for R. Bioinformatics, 24(5), 719-720.

Friedman, J., Hastie, T. and Tibshirani, R. (2008) *Regularization Paths for Generalized Linear Models via Coordinate Descent*, http://www.stanford.edu/~hastie/Papers/glmnet.pdf

Breheny, P. and Huang, J. (2011) Coordinate descent algorithms for nonconvex penalized regression, with applications to biological feature selection. Ann. Appl. Statist., 5: 232-253.

### **Examples**

library(magrittr)

# simulation parameters

```
rho = 0.90; p = 500; SNR = 1; n = 200; n0 = n1 = 100; nActive = p*0.10; cluster_distance = "tom";
Ecluster_distance = "difftom"; rhoOther = 0.6; betaMean = 2;
alphaMean = 1; betaE = 3; distanceMethod = "euclidean"; clustMethod = "hclust";
cutMethod = "dynamic"; agglomerationMethod = "average"
#in this simulation its blocks 3 and 4 that are important
#leaveOut: optional specification of modules that should be left out
#of the simulation, that is their genes will be simulated as unrelated
#("grey"). This can be useful when simulating several sets, in some which a module
#is present while in others it is absent.
d0 \leftarrow s_modules(n = n0, p = p, rho = 0, exposed = FALSE,
                modProportions = c(0.15, 0.15, 0.15, 0.15, 0.15, 0.25),
                minCor = 0.01,
                maxCor = 1,
                 corPower = 1,
                 propNegativeCor = 0.3,
                 backgroundNoise = 0.5,
                 signed = FALSE,
                leaveOut = 1:4)
d1 <- s_modules(n = n1, p = p, rho = rho, exposed = TRUE,</pre>
                modProportions = c(0.15, 0.15, 0.15, 0.15, 0.15, 0.25),
                minCor = 0.4,
                maxCor = 1,
                corPower = 0.3,
                 propNegativeCor = 0.3,
                 backgroundNoise = 0.5,
                 signed = FALSE)
truemodule1 <- d1$setLabels
X <- rbind(d0$datExpr, d1$datExpr) %>%
  magrittr::set_colnames(paste0("Gene", 1:p)) %>%
  magrittr::set_rownames(paste0("Subject",1:n))
betaMainEffect <- vector("double", length = p)</pre>
betaMainInteractions <- vector("double", length = p)</pre>
# the first nActive/2 in the 3rd block are active
betaMainEffect[which(truemodule1 %in% 3)[1:(nActive/2)]] <- runif(</pre>
  nActive/2, betaMean - 0.1, betaMean + 0.1)
# the first nActive/2 in the 4th block are active
betaMainEffect[which(truemodule1 %in% 4)[1:(nActive/2)]] <- runif(</pre>
  nActive/2, betaMean+2 - 0.1, betaMean+2 + 0.1)
betaMainInteractions[which(betaMainEffect!=0)] <- runif(nActive, alphaMean - 0.1, alphaMean + 0.1)
beta <- c(betaMainEffect, betaE, betaMainInteractions)</pre>
result <- s_generate_data(p = p, X = X,</pre>
                           beta = beta,
                           include_interaction = TRUE,
                           cluster_distance = cluster_distance,
                           n = n, n0 = n0,
```

```
eclust_distance = Ecluster_distance,
                           signal_to_noise_ratio = SNR,
                           distance_method = distanceMethod,
                           cluster_method = clustMethod,
                           cut_method = cutMethod,
                           agglomeration_method = agglomerationMethod,
                           nPC = 1)
pen_res <- s_pen_clust(x_train = result[["X_train"]],</pre>
                        x_test = result[["X_test"]],
                        y_train = result[["Y_train"]],
                        y_test = result[["Y_test"]],
                        s0 = result[["S0"]],
                        gene_groups = result[["clustersAddon"]],
                        summary = "pc",
                        model = "lasso",
                        exp_family = "gaussian",
                        clust_type = "ECLUST",
                        include_interaction = TRUE)
unlist(pen_res)
```

s\_pen\_separate

Fit Penalized Regression Models on Simulated Data

# Description

This function can run penalized regression models on the untransformed design matrix. To be used with simulated data where the 'truth' is known i.e., you know which features are associated with the response. This function was used to produce the simulation results in Bhatnagar et al. 2016. Can run lasso, elasticnet, SCAD or MCP models

# Usage

```
s_pen_separate(x_train, x_test, y_train, y_test, s0,
  exp_family = c("gaussian", "binomial"), model = c("lasso", "elasticnet",
  "scad", "mcp"), topgenes = NULL, stability = F, filter = F,
  include_E = T, include_interaction = F)
```

# **Arguments**

 $x_{train}$ 

ntrain x p matrix of simulated training set where ntrain is the number of training observations and p is total number of predictors. This matrix needs to have named columns representing the feature names or the gene names

x\_test

ntest x p matrix of simulated training set where ntest is the number of training observations and p is total number of predictors. This matrix needs to have named columns representing the feature names or the gene names

y\_train numeric vector of length ntrain representing the responses for the training subjects. If continuous then you must set exp\_family = "gaussion". For exp\_family="binomial" should be either a factor with two levels, or a twocolumn matrix of counts or proportions (the second column is treated as the target class; for a factor, the last level in alphabetical order is the target class) y\_test numeric vector of length ntest representing the responses for the test subjects. If continuous then you must set exp\_family = "gaussion". For exp\_family="binomial" should be either a factor with two levels, or a two-column matrix of counts or proportions (the second column is treated as the target class; for a factor, the last level in alphabetical order is the target class). s0 chracter vector of the active feature names, i.e., the features in x\_train that are truly associated with the response. exp\_family Response type. See details for y\_train argument above. model Regression model to be fit on cluster summaries. Default is model="lasso" which corresponds to glmnet mixing parameter alpha=1. model="elasticnet" corresponds to glmnet mixing parameter alpha=0.5, model="mcp" and model="scad" are the non-convex models from the nevreg package List of features to keep if filter=TRUE. Default is topgenes = NULL which topgenes means all features are kept for the analysis Should stability measures be calculated. Default is stability=FALSE. See destability filter Should analysis be run on a subset of features. Default is filter = FALSE include\_E Should the environment variable be included in the regression analysis. Default is include\_E = TRUE include\_interaction

Should interaction effects between the features in x\_train and the environment variable be fit. Default is include\_interaction=TRUE

#### **Details**

The stability of feature importance is defined as the variability of feature weights under perturbations of the training set, i.e., small modifications in the training set should not lead to considerable changes in the set of important covariates (Toloşi, L., & Lengauer, T. (2011)). A feature selection algorithm produces a weight, a ranking, and a subset of features. In the CLUST and ECLUST methods, we defined a predictor to be non-zero if its corresponding cluster representative weight was non-zero. Using 10-fold cross validation (CV), we evaluated the similarity between two features and their rankings using Pearson and Spearman correlation, respectively. For each CV fold we re-ran the models and took the average Pearson/Spearman correlation of the 10 choose 2 combinations of estimated coefficients vectors. To measure the similarity between two subsets of features we took the average of the Jaccard distance in each fold. A Jaccard distance of 1 indicates perfect agreement between two sets while no agreement will result in a distance of 0.

#### Value

This function has two different outputs depending on whether stability = TRUE or Stability = FALSE

If stability = TRUE then this function returns a p x 2 data.frame or data.table of regression coefficients without the intercept. The output of this is used for subsequent calculations of stability.

If stability = FALSE then returns a vector with the following elements (See Table 3: Measures of Performance in Bhatnagar et al (2016+) for definitions of each measure of performance):

mse or AUC Test set mean squared error if exp\_family = "gaussion" or test set Area under

the curve if exp\_family = "binomial" calculated using the roc function

RMSE Square root of the mse. Only applicable if exp\_family = "gaussion"

Shat Number of non-zero estimated regression coefficients. The non-zero estimated

regression coefficients are referred to as being selected by the model

TPR true positive rate
FPR false positive rate

Correct Sparsity

Correct true positives + correct true negative coefficients divided by the total

number of features

CorrectZeroMain

Proportion of correct true negative main effects

CorrectZeroInter

Proportion of correct true negative interactions

IncorrectZeroMain

Proportion of incorrect true negative main effects

IncorrectZeroInter

Proportion of incorrect true negative interaction effects

# References

Toloşi, L., & Lengauer, T. (2011). Classification with correlated features: unreliability of feature ranking and solutions. Bioinformatics, 27(14), 1986-1994.

Bhatnagar, SR., Yang, Y., Blanchette, M., Bouchard, L., Khundrakpam, B., Evans, A., Greenwood, CMT. (2016+). An analytic approach for interpretable predictive models in high dimensional data, in the presence of interactions with exposures *Preprint* 

Langfelder, P., Zhang, B., & Horvath, S. (2008). *Defining clusters from a hierarchical cluster tree:* the Dynamic Tree Cut package for R. Bioinformatics, 24(5), 719-720.

Friedman, J., Hastie, T. and Tibshirani, R. (2008) *Regularization Paths for Generalized Linear Models via Coordinate Descent*, http://www.stanford.edu/~hastie/Papers/glmnet.pdf

Breheny, P. and Huang, J. (2011) Coordinate descent algorithms for nonconvex penalized regression, with applications to biological feature selection. Ann. Appl. Statist., 5: 232-253.

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

# simulation parameters
rho = 0.90; p = 500 ;SNR = 1 ; n = 200; n0 = n1 = 100 ; nActive = p*0.10 ; cluster_distance = "tom";
Ecluster_distance = "difftom"; rhoOther = 0.6; betaMean = 2;
```

```
alphaMean = 1; betaE = 3; distanceMethod = "euclidean"; clustMethod = "hclust";
cutMethod = "dynamic"; agglomerationMethod = "average"
#in this simulation its blocks 3 and 4 that are important
#leaveOut: optional specification of modules that should be left out
#of the simulation, that is their genes will be simulated as unrelated
#("grey"). This can be useful when simulating several sets, in some which a module
#is present while in others it is absent.
d\theta <- s_modules(n = n\theta, p = p, rho = \theta, exposed = FALSE,
                modProportions = c(0.15, 0.15, 0.15, 0.15, 0.15, 0.25),
                minCor = 0.01,
                maxCor = 1,
                 corPower = 1,
                 propNegativeCor = 0.3,
                 backgroundNoise = 0.5,
                 signed = FALSE,
                leaveOut = 1:4)
d1 \leftarrow s_{modules}(n = n1, p = p, rho = rho, exposed = TRUE,
                modProportions = c(0.15, 0.15, 0.15, 0.15, 0.15, 0.25),
                minCor = 0.4,
                maxCor = 1,
                corPower = 0.3,
                 propNegativeCor = 0.3,
                 backgroundNoise = 0.5,
                 signed = FALSE)
truemodule1 <- d1$setLabels
X <- rbind(d0$datExpr, d1$datExpr) %>%
  magrittr::set_colnames(paste0("Gene", 1:p)) %>%
  magrittr::set_rownames(paste0("Subject",1:n))
betaMainEffect <- vector("double", length = p)</pre>
betaMainInteractions <- vector("double", length = p)</pre>
# the first nActive/2 in the 3rd block are active
betaMainEffect[which(truemodule1 %in% 3)[1:(nActive/2)]] <- runif(</pre>
  nActive/2, betaMean - 0.1, betaMean + 0.1)
# the first nActive/2 in the 4th block are active
betaMainEffect[which(truemodule1 %in% 4)[1:(nActive/2)]] <- runif(</pre>
  nActive/2, betaMean+2 - 0.1, betaMean+2 + 0.1)
betaMainInteractions[which(betaMainEffect!=0)] <- runif(nActive, alphaMean - 0.1, alphaMean + 0.1)
beta <- c(betaMainEffect, betaE, betaMainInteractions)</pre>
result <- s_generate_data(p = p, X = X,
                           beta = beta,
                           include_interaction = TRUE,
                           cluster_distance = cluster_distance,
                           n = n, n0 = n0,
                           eclust_distance = Ecluster_distance,
                           signal_to_noise_ratio = SNR,
```

s\_response 35

s\_response

Generate True Response vector for Linear Simulation

# **Description**

Given the true beta vector, covariates and environment variable this function generates the linear response with specified signal to noise ratio.

# Usage

```
s_response(n, n0, p, genes, binary_outcome = FALSE, E,
   signal_to_noise_ratio = 1, include_interaction = FALSE, beta = NULL)
```

# Arguments

n	Total number of subjects
n0	Total number of unexposed subjects
р	Total number of genes (or covariates)
genes	nxp matrix of the genes or covariates
binary_outcome	Logical. Should a binary outcome be generated. Default is FALSE. See details on how a binary outcome is generated
Е	binary 0,1, vector of the exposure/environment variable
signal_to_noise	e_ratio
	a numeric variable for the signal to noise ratio
include_interac	tion
	$Logical. \ Should \ the \ response \ include \ the \ interaction \ between \ E \ and \ the \ genes \ (for \ the \ non-zero \ beta \ coefficient \ vector)$
beta	true beta coefficient vector. Assumes this vector is in the same order as the genes.

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#### Value

a data.frame/data.table containing the response and the design matrix. Also an object of class expression

```
library(magrittr)
# simulation parameters
rho = 0.90; p = 500; snr = 1; rn = 200; rno = 100; rno = 1
Ecluster_distance = "difftom"; rhoOther = 0.6; betaMean = 2;
alphaMean = 1; betaE = 3; distanceMethod = "euclidean"; clustMethod = "hclust";
cutMethod = "dynamic"; agglomerationMethod = "average"
#in this simulation its blocks 3 and 4 that are important
#leaveOut: optional specification of modules that should be left out
#of the simulation, that is their genes will be simulated as unrelated
#("grey"). This can be useful when simulating several sets, in some which a module
#is present while in others it is absent.
d0 < -s_{modules}(n = n0, p = p, rho = 0, exposed = FALSE,
                                 modProportions = c(0.15, 0.15, 0.15, 0.15, 0.15, 0.25),
                                 minCor = 0.01,
                                 maxCor = 1,
                                 corPower = 1,
                                 propNegativeCor = 0.3,
                                 backgroundNoise = 0.5,
                                 signed = FALSE,
                                 leaveOut = 1:4)
d1 <- s_modules(n = n1, p = p, rho = rho, exposed = TRUE,</pre>
                                 modProportions = c(0.15, 0.15, 0.15, 0.15, 0.15, 0.25),
                                 minCor = 0.4,
                                 maxCor = 1,
                                 corPower = 0.3,
                                 propNegativeCor = 0.3,
                                 backgroundNoise = 0.5,
                                 signed = FALSE)
truemodule1 <- d1$setLabels
X <- rbind(d0$datExpr, d1$datExpr) %>%
    magrittr::set_colnames(paste0("Gene", 1:p)) %>%
    magrittr::set_rownames(paste0("Subject",1:n))
betaMainEffect <- vector("double", length = p)</pre>
# the first nActive/2 in the 3rd block are active
betaMainEffect[which(truemodule1 %in% 3)[1:(nActive/2)]] <- runif(</pre>
    nActive/2, betaMean - 0.1, betaMean + 0.1)
# the first nActive/2 in the 4th block are active
betaMainEffect[which(truemodule1 %in% 4)[1:(nActive/2)]] <- runif(</pre>
```

s\_response\_mars 37

s\_response\_mars

Generate True Response vector for Non-Linear Simulation

# **Description**

Given the covariates and environment variable this function generates the nonlinear response with specified signal to noise ratio.

### Usage

```
s_response_mars(n, n0, p, genes, beta, binary_outcome = FALSE, E,
    signal_to_noise_ratio = 1, truemodule, nActive)
```

### **Arguments**

total number of subjects n n0 total number of subjects with E=0 number of genes in design matrix genes nxp matrix of the genes or covariates true beta coefficient vector beta binary\_outcome Logical. Should a binary outcome be generated. Default is FALSE. See details on how a binary outcome is generated F binary 0,1, vector of the exposure/environment variable signal\_to\_noise\_ratio signal to noise ratio, default is 1 truemodule numeric vector of the true module membership used in the s\_response\_mars function. Modules 3 and 4 are active in the response. See s\_response\_mars function for details. number of active genes in the response used in the s\_response\_mars nActive

# Value

a data.frame/data.table containing the response and the design matrix. Also an object of class expression

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#### Note

See Bhatnagar et al (2017+) for details on how the response is simulated.

```
library(magrittr)
# simulation parameters
rho = 0.90; p = 500; SNR = 1; n = 200; n0 = n1 = 100; nActive = p*0.10; cluster_distance = "tom";
Ecluster_distance = "difftom"; rhoOther = 0.6; betaMean = 2;
alphaMean = 1; betaE = 3; distanceMethod = "euclidean"; clustMethod = "hclust";
cutMethod = "dynamic"; agglomerationMethod = "average"
#in this simulation its blocks 3 and 4 that are important
#leaveOut: optional specification of modules that should be left out
#of the simulation, that is their genes will be simulated as unrelated
#("grey"). This can be useful when simulating several sets, in some which a module
#is present while in others it is absent.
d0 <- s_modules(n = n0, p = p, rho = 0, exposed = FALSE,
                modProportions = c(0.15, 0.15, 0.15, 0.15, 0.15, 0.25),
                minCor = 0.01,
                maxCor = 1,
                corPower = 1,
                propNegativeCor = 0.3,
                backgroundNoise = 0.5,
                signed = FALSE,
                leaveOut = 1:4)
d1 <- s_modules(n = n1, p = p, rho = rho, exposed = TRUE,</pre>
                modProportions = c(0.15, 0.15, 0.15, 0.15, 0.15, 0.25),
                minCor = 0.4,
                maxCor = 1,
                corPower = 0.3,
                propNegativeCor = 0.3,
                backgroundNoise = 0.5,
                signed = FALSE)
truemodule1 <- d1$setLabels
X <- rbind(d0$datExpr, d1$datExpr) %>%
  magrittr::set_colnames(paste0("Gene", 1:p)) %>%
  magrittr::set_rownames(paste0("Subject",1:n))
betaMainEffect <- vector("double", length = p)</pre>
# the first nActive/2 in the 3rd block are active
betaMainEffect[which(truemodule1 %in% 3)[1:(nActive/2)]] <- runif(</pre>
  nActive/2, betaMean - 0.1, betaMean + 0.1)
# the first nActive/2 in the 4th block are active
betaMainEffect[which(truemodule1 %in% 4)[1:(nActive/2)]] <- runif(</pre>
  nActive/2, betaMean+2 - 0.1, betaMean+2 + 0.1)
```

tcgaov 39

tcgaov

Subset of TCGA mRNA Ovarian serous cystadenocarcinoma data

### **Description**

A dataset containing a subset of the TCGA mRNA Ovarian serous cystadenocarcinoma data generated using Affymetrix HTHGU133a arrays. Differences in gene expression profiles have led to the identification of robust molecular subtypes of ovarian cancer; these are of biological and clinical importance because they have been shown to correlate with overall survival (Tothill et al., 2008). Improving prediction of survival time based on gene expression signatures can lead to targeted therapeutic interventions (Helland et al., 2011). The proposed ECLUST algorithm was applied to gene expression data from 511 ovarian cancer patients profiled by the Affymetrix Human Genome U133A 2.0 Array. The data were obtained from the TCGA Research Network: http://cancergenome.nih.gov/ and downloaded via the TCGA2STAT R library (Wanet al., 2015). Using the 881 signature genes from Helland et al. (2011) we grouped subjects into two groups based on the results in this paper, to create a "positive control" environmental variable expected to have a strong effect. Specifically, we defined an environment variable in our framework as: E = 0 for subtypes C1 and C2 (E = 0), and E = 1 for subtypes C4 and C5 (E = 0).

# Usage

tcgaov

# Format

A data.table and data.frame with 511 rows and 886 variables:

```
rn unique patient identifier (character)
subtype cancer subtype (1,2,3 or 4) as per Helland et al. 2011 (integer)
E binary environment variable for ECLUST method. E = 0 for subtypes 1 and 2 (n = 253), and E = 1 for subtypes 4 and 5 (n = 258) (numeric)
status vital status, 0 = alive, 1 = dead (numeric)
OS overall survival time (numeric)
columns 6:886 gene expression data for 881 genes, column names are the gene names (numeric)
```

40 u\_cluster\_similarity

### **Source**

```
http://www.liuzlab.org/TCGA2STAT/#import-gene-expression
http://gdac.broadinstitute.org/
http://journals.plos.org/plosone/article/asset?unique&id=info:doi/10.1371/journal.
pone.0018064.s015
```

#### References

Richard W Tothill, Anna V Tinker, Joshy George, Robert Brown, Stephen B Fox, Stephen Lade, Daryl S Johnson, Melanie K Trivett, Dariush Etemadmoghadam, Bianca Locandro, et al. Novel molecular subtypes of serous and endometrioid ovarian cancer linked to clinical outcome. Clinical Cancer Research, 14(16):5198–5208, 2008.

Aslaug Helland, Michael S Anglesio, Joshy George, Prue A Cowin, Cameron N Johnstone, Colin M House, Karen E Sheppard, Dariush Etemadmoghadam, Nataliya Melnyk, Anil K Rustgi, et al. Deregulation of mycn, lin28b and let7 in a molecular subtype of aggressive high-grade serous ovarian cancers. PloS one, 6(4):e18064, 2011.

# **Examples**

```
# using data.table syntax from the data.table package
tcgaov[1:5, 1:10, with = FALSE]
tcgaov[,table(subtype, E, useNA = "always")]
```

```
u_cluster_similarity Cluster similarity matrix
```

Description

Return cluster membership of each predictor. This function is called internally by the s\_generate\_data and s\_generate\_data\_mars functions. Is also used by the r\_clust function for real data analysis.

# Usage

```
u_cluster_similarity(x, expr, exprTest, distanceMethod,
  clustMethod = c("hclust", "protoclust"), cutMethod = c("dynamic", "gap",
  "fixed"), nClusters, method = c("complete", "average", "ward.D2", "single",
  "ward.D", "mcquitty", "median", "centroid"), K.max = 10, B = 50, nPC,
  minimum_cluster_size = 50)
```

#### **Arguments**

X	similarity matrix. must have non-NULL dimnames i.e., the rows and columns should be labelled, e.g. "Gene1, Gene2,"
expr	gene expression data (training set). rows are people, columns are genes
exprTest	gene expression test set. If using real data, and you dont have enough samples for a test set then just supply the same data supplied to the expr argument

u\_cluster\_similarity 41

distanceMethod one of "euclidean", "maximum", "manhattan", "canberra", "binary", "minkowski"

to be passed to dist function. If missing, then this function will take 1-x as the dissimilarity measure. This functionality is for diffCorr,diffTOM, fisherScore

matrices which need to be converted to a distance type matrix.

clustMethod Cluster the data using hierarchical clustering or prototype clustering. Defaults

clustMethod="hclust". Other option is protoclust, however this package

must be installed before proceeding with this option

cutMethod what method to use to cut the dendrogram. 'dynamic' refers to cutreeDynamicTree

library. 'gap' is Tibshirani's gap statistic clusGap using the 'Tibs2001SEmax'

rule. 'fixed' is a fixed number specified by the nClusters argument

nClusters number of clusters. Only used if cutMethod = fixed

method the agglomeration method to be used. This should be (an unambiguous ab-

breviation of) one of "ward.D", "ward.D2", "single", "complete", "average" (= UPGMA), "mcquitty" (= WPGMA), "median" (= WPGMC) or "centroid" (=

UPGMC).

K.max the maximum number of clusters to consider, must be at least two. Only used if

cutMethod='gap'

B integer, number of Monte Carlo ("bootstrap") samples. Only used if cutMethod='gap'

nPC number of principal components. Can be 1 or 2.

minimum\_cluster\_size

The minimum cluster size. Only applicable if cutMethod='dynamic'. This argument is passed to the cutreeDynamic function. Default is 50.

# Value

a list of length 2:

**clusters** a p x 3 data.frame or data.table which give the cluster membership of each gene, where p is the number of genes. The first column is the gene name, the second column is the cluster number (numeric) and the third column is the cluster membership as a character vector of color names (these will match up exactly with the cluster number)

**pcInfo** a list of length 9:

**eigengenes** a list of the eigengenes i.e. the 1st (and 2nd if nPC=2) principal component of each module

averageExpr a data.frame of the average expression for each module for the training set
 averageExprTest a data.frame of the average expression for each module for the test set
 varExplained percentage of variance explained by each 1st (and 2nd if nPC=2) principal component of each module

validColors cluster membership of each gene

**PC** a data.frame of the 1st (and 2nd if nPC=2) PC for each module for the training set **PCTest** a data.frame of the 1st (and 2nd if nPC=2) PC for each module for the test set **prcompObj** the prcomp object

nclusters a numeric value for the total number of clusters

### **Examples**

```
u_extract_selected_earth
```

Get selected terms from an earth object

# **Description**

function to extract the selected terms from an earth object

### Usage

```
u_extract_selected_earth(obj)
```

# **Arguments**

obj

object of class earth returned by the earth function

#### **Details**

called internally by the s\_mars\_separate and s\_mars\_clust functions

#### Value

character vector of selected terms from the MARS model

u\_extract\_summary 43

u\_extract\_summary

Calculates cluster summaries

# Description

This is a modified version of moduleEigengenes. It can extract (1st and 2nd principal component) of modules in a given single dataset. It can also return the average, the variance explained This function is more flexible and the nPC argument is used. currently only nPC = 1 and nPC = 2 are supported

# Usage

```
u_extract_summary(x_train, colors, x_test, y_train, y_test, impute = TRUE,
nPC, excludeGrey = FALSE, grey = if (is.numeric(colors)) 0 else "grey",
subHubs = TRUE, trapErrors = FALSE, returnValidOnly = trapErrors,
softPower = 6, scale = TRUE, verbose = 0, indent = 0)
```

# **Arguments**

x_train	Training data for a single set in the form of a data frame where rows are samples and columns are genes (probes, cpgs, covariates).
colors	A vector of the same length as the number of probes in expr, giving module color for all probes (genes). Color "grey" is reserved for unassigned genes.
x_test	Test set in the form of a data frame where rows are samples and columns are genes (probes, cpgs, covariates).
y_train	Training response numeric vector
y_test	Test response numeric vector
impute	If TRUE, expression data will be checked for the presence of NA entries and if the latter are present, numerical data will be imputed, using function impute.knn and probes from the same module as the missing datum. The function impute.knn uses a fixed random seed giving repeatable results.
nPC	Number of principal components and variance explained entries to be calculated. Note that only 1 or 2 is possible.
excludeGrey	Should the improper module consisting of 'grey' genes be excluded from the eigengenes?
grey	Value of colors designating the improper module. Note that if colors is a factor of numbers, the default value will be incorrect.
subHubs	Controls whether hub genes should be substituted for missing eigengenes. If TRUE, each missing eigengene (i.e., eigengene whose calculation failed and the error was trapped) will be replaced by a weighted average of the most connected hub genes in the corresponding module. If this calculation fails, or if subHubs==FALSE, the value of trapErrors will determine whether the offending module will be removed or whether the function will issue an error and stop.

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trapErrors

Controls handling of errors from that may arise when there are too many NA entries in expression data. If TRUE, errors from calling these functions will be trapped without abnormal exit. If FALSE, errors will cause the function to stop. Note, however, that subHubs takes precedence in the sense that if subHubs==TRUE and trapErrors==FALSE, an error will be issued only if both the principal component and the hubgene calculations have failed.

returnValidOnly

logical; controls whether the returned data frame of module eigengenes contains columns corresponding only to modules whose eigengenes or hub genes could be calculated correctly (TRUE), or whether the data frame should have columns for each of the input color labels (FALSE).

softPower The power u

The power used in soft-thresholding the adjacency matrix. Only used when the hubgene approximation is necessary because the principal component calculation failed. It must be non-negative. The default value should only be changed if there is a clear indication that it leads to incorrect results.

scale logical; can be used to turn off scaling of the expression data before calculating

the singular value decomposition. The scaling should only be turned off if the data has been scaled previously, in which case the function can run a bit faster. Note however that the function first imputes, then scales the expression data in each module. If the expression contain missing data, scaling outside of the function and letting the function impute missing data may lead to slightly different

results than if the data is scaled within the function.

verbose Controls verbosity of printed progress messages. 0 means silent, up to (about) 5

the verbosity gradually increases.

indent A single non-negative integer controlling indentation of printed messages. 0

means no indentation, each unit above that adds two spaces.

# Details

This function is called internally by the u\_cluster\_similarity function

### Value

A list with the following components:

eigengenes Module eigengenes in a dataframe, with each column corresponding to one eigengene

averageExpr the average expression per module in the training set

averageExprTest the average expression per module in the training set

varExplained The variance explained by the first PC in each module

**validColors** A copy of the input colors with entries corresponding to invalid modules set to grey if given, otherwise 0 if colors is numeric and "grey" otherwise.

PC The 1st or 1st and 2nd PC from each module in the training set

PCTest The 1st or 1st and 2nd PC from each module in the test set

prcompObj The prcomp object returned by prcomp

**nclusters** the number of modules (clusters)

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### References

Zhang, B. and Horvath, S. (2005), "A General Framework for Weighted Gene Co-Expression Network Analysis", Statistical Applications in Genetics and Molecular Biology: Vol. 4: No. 1, Article 17

# **Examples**

```
## Not run:
#see u_cluster_similarity for examples
## End(Not run)
```

u\_fisherZ

Calculate Fisher's Z Transformation for Correlations

# Description

Calculate Fisher's Z transformation for correlations. This can be used as an alternative measure of similarity. Used in the s\_generate\_data function

# Usage

```
u_fisherZ(n0, cor0, n1, cor1)
fisherTransform(n_1, r1, n_2, r2)
```

# Arguments

n0	number of unexposed subjects
cor0	correlation matrix of unexposed covariate values. Should be dimension pxp
n1	number of exposed subjects
cor1	correlation matrix of exposed covariate values. Should be dimension pxp
n_1	number of unexposed subjects
r1	correlation for unexposed
n_2	number of exposed subjects
r2	correlation for exposed

# Value

a pxp matrix of Fisher's Z transformation of correlations

# Note

fisherTransform is called internally by u\_fisherZ function

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# References

https://en.wikipedia.org/wiki/Fisher\_transformation

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