Package ‘LUCIDus’

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boot.lucid

*Deprecated function boot.lucid*

Description

This function deprecates. Please use boot_lucid instead.

Usage

```
boot.lucid(G, Z, Y, CoG = NULL, CoY = NULL, model, conf = 0.95, R = 100)
```

Arguments

- **G**
  - Exposures, a numeric vector, matrix, or data frame. Categorical variable should be transformed into dummy variables. If a matrix or data frame, rows represent observations and columns correspond to variables.

- **Z**
  - Omics data, a numeric matrix or data frame. Rows correspond to observations and columns correspond to variables.

- **Y**
  - Outcome, a numeric vector. Categorical variable is not allowed. Binary outcome should be coded as 0 and 1.

- **CoG**
  - Optional, covariates to be adjusted for estimating the latent cluster. A numeric vector, matrix or data frame. Categorical variable should be transformed into dummy variables.
**boot_lucid**

Optional, covariates to be adjusted for estimating the association between latent cluster and the outcome. A numeric vector, matrix or data frame. Categorical variable should be transformed into dummy variables.

**model**

A LUCID model fitted by `est.lucid`.

**conf**

A numeric scalar between 0 and 1 to specify confidence level(s) of the required interval(s).

**R**

An integer to specify number of bootstrap replicates for LUCID model. If feasible, it is recommended to set R ≥ 1000.

---

**Inference of LUCID model based on bootstrap resampling**

**Description**

Generate R bootstrap replicates of LUCID parameters and derive confidence interval (CI) base on bootstrap. Bootstrap replicates are generated based on nonparameteric resampling, implemented by ordinary method of codeboot::boot function.

**Usage**

```r
boot_lucid(G, Z, Y, CoG = NULL, CoY = NULL, model, conf = 0.95, R = 100)
```

**Arguments**

- **G**
  - Exposures, a numeric vector, matrix, or data frame. Categorical variable should be transformed into dummy variables. If a matrix or data frame, rows represent observations and columns correspond to variables.

- **Z**
  - Omics data, a numeric matrix or data frame. Rows correspond to observations and columns correspond to variables.

- **Y**
  - Outcome, a numeric vector. Categorical variable is not allowed. Binary outcome should be coded as 0 and 1.

- **CoG**
  - Optional, covariates to be adjusted for estimating the latent cluster. A numeric vector, matrix or data frame. Categorical variable should be transformed into dummy variables.

- **CoY**
  - Optional, covariates to be adjusted for estimating the association between latent cluster and the outcome. A numeric vector, matrix or data frame. Categorical variable should be transformed into dummy variables.

- **model**
  - A LUCID model fitted by `est.lucid`.

- **conf**
  - A numeric scalar between 0 and 1 to specify confidence level(s) of the required interval(s).

- **R**
  - An integer to specify number of bootstrap replicates for LUCID model. If feasible, it is recommended to set R ≥ 1000.
check_na

Check missing patterns in omics data Z

Description

Check missing patterns in omics data Z.

Usage

check_na(Z)

Arguments

Z A data matrix representing omics data
est.lucid

Value

1. index: indices for missing values in omics data
2. indicator_na: missing pattern for each observation
3. impute_flag: - flag to initialize imputation. Only happens when sporadic missing pattern is observed

est.lucid  Deprecated function est.lucid

Description

This function deprecates. Please use est_lucid instead.

Usage

est.lucid(
  G,
  Z,
  Y,
  CoG = NULL,
  CoY = NULL,
  K = 2,
  family = c("normal", "binary"),
  useY = TRUE,
  tol = 0.001,
  max_itr = 1000,
  max_tot.itr = 10000,
  Rho_G = 0,
  Rho_Z_Mu = 0,
  Rho_Z_Cov = 0,
  modelName = "VVV",
  seed = 123,
  init_impute = c("mclust", "lod"),
  init_par = c("mclust", "random"),
  verbose = FALSE
)

Arguments

G  Exposures, a numeric vector, matrix, or data frame. Categorical variable should be transformed into dummy variables. If a matrix or data frame, rows represent observations and columns correspond to variables.

Z  Omics data, a numeric matrix or data frame. Rows correspond to observations and columns correspond to variables.

Y  Outcome, a numeric vector. Categorical variable is not allowed. Binary outcome should be coded as 0 and 1.
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>CoG</td>
<td>Optional, covariates to be adjusted for estimating the latent cluster. A numeric vector, matrix or data frame. Categorical variable should be transformed into dummy variables.</td>
</tr>
<tr>
<td>CoY</td>
<td>Optional, covariates to be adjusted for estimating the association between latent cluster and the outcome. A numeric vector, matrix or data frame. Categorical variable should be transformed into dummy variables.</td>
</tr>
<tr>
<td>K</td>
<td>Number of latent clusters. An integer greater or equal to 2. User can use <code>lucid</code> to determine the optimal number of latent clusters.</td>
</tr>
<tr>
<td>family</td>
<td>Distribution of outcome. For continuous outcome, use &quot;normal&quot;; for binary outcome, use &quot;binary&quot;. Default is &quot;normal&quot;.</td>
</tr>
<tr>
<td>useY</td>
<td>Flag to include information of outcome when estimating the latent cluster. Default is TRUE.</td>
</tr>
<tr>
<td>tol</td>
<td>Tolerance for convergence of EM algorithm. Default is 1e-3.</td>
</tr>
<tr>
<td>max_itr</td>
<td>Max number of iterations for EM algorithm.</td>
</tr>
<tr>
<td>max_tot_itr</td>
<td>Max number of total iterations for <code>est_lucid</code> function. <code>est_lucid</code> may conduct EM algorithm for multiple times if the algorithm fails to converge.</td>
</tr>
<tr>
<td>Rho_G</td>
<td>A scalar. This parameter is the LASSO penalty to regularize exposures. If user wants to tune the penalty, use the wrapper function <code>lucid</code>.</td>
</tr>
<tr>
<td>Rho_Z_Mu</td>
<td>A scalar. This parameter is the LASSO penalty to regularize cluster-specific means for omics data (Z). If user wants to tune the penalty, use the wrapper function <code>lucid</code>.</td>
</tr>
<tr>
<td>Rho_Z_Cov</td>
<td>A scalar. This parameter is the graphical LASSO penalty to estimate sparse cluster-specific variance-covariance matrices for omics data (Z). If user wants to tune the penalty, use the wrapper function <code>lucid</code>.</td>
</tr>
<tr>
<td>modelName</td>
<td>The variance-covariance structure for omics data. See <code>mclust::mclustModelNames</code> for details.</td>
</tr>
<tr>
<td>seed</td>
<td>An integer to initialize the EM algorithm or imputing missing values. Default is 123.</td>
</tr>
<tr>
<td>init_impute</td>
<td>Method to initialize the imputation of missing values in LUCID. &quot;mclust&quot; will use <code>mclust::imputeData</code> to implement EM Algorithm for Unrestricted General Location Model to impute the missing values in omics data; <code>lod</code> will initialize the imputation via replacing missing values by ( \text{LOD} / \sqrt{2} ). LOD is determined by the minimum of each variable in omics data.</td>
</tr>
<tr>
<td>init_par</td>
<td>Method to initialize the EM algorithm. &quot;mclust&quot; will use <code>mclust</code> model to initialize parameters; &quot;random&quot; initialize parameters from uniform distribution.</td>
</tr>
<tr>
<td>verbose</td>
<td>A flag indicates whether detailed information for each iteration of EM algorithm is printed in console. Default is FALSE.</td>
</tr>
</tbody>
</table>
Fit LUCID model to conduct integrated clustering

Description

The Latent Unknown Clustering with Integrated Data (LUCID) performs integrative clustering using multi-view data. LUCID model is estimated via EM algorithm for model-based clustering. It also features variable selection, integrated imputation, bootstrap inference and visualization via Sankey diagram.

Usage

```r
est_lucid(
  G,
  Z,
  Y,
  CoG = NULL,
  CoY = NULL,
  K = 2,
  family = c("normal", "binary"),
  useY = TRUE,
  tol = 0.001,
  max_itr = 1000,
  max_tot.itr = 10000,
  Rho_G = 0,
  Rho_Z_Mu = 0,
  Rho_Z_Cov = 0,
  modelName = NULL,
  seed = 123,
  init_impute = c("mclust", "lod"),
  init_par = c("mclust", "random"),
  verbose = FALSE
)
```

Arguments

- **G**: Exposures, a numeric vector, matrix, or data frame. Categorical variable should be transformed into dummy variables. If a matrix or data frame, rows represent observations and columns correspond to variables.

- **Z**: Omics data, a numeric matrix or data frame. Rows correspond to observations and columns correspond to variables.

- **Y**: Outcome, a numeric vector. Categorical variable is not allowed. Binary outcome should be coded as 0 and 1.

- **CoG**: Optional, covariates to be adjusted for estimating the latent cluster. A numeric vector, matrix or data frame. Categorical variable should be transformed into dummy variables.
est_lucid

CoY Optional, covariates to be adjusted for estimating the association between latent cluster and the outcome. A numeric vector, matrix or data frame. Categorical variable should be transformed into dummy variables.

K Number of latent clusters. An integer greater or equal to 2. User can use lucid to determine the optimal number of latent clusters.

family Distribution of outcome. For continuous outcome, use "normal"; for binary outcome, use "binary". Default is "normal".

useY Flag to include information of outcome when estimating the latent cluster. Default is TRUE.

tol Tolerance for convergence of EM algorithm. Default is 1e-3.

max_itr Max number of iterations for EM algorithm.

max_tot.itr Max number of total iterations for est_lucid function. est_lucid may conduct EM algorithm for multiple times if the algorithm fails to converge.

Rho_G A scalar. This parameter is the LASSO penalty to regularize exposures. If user wants to tune the penalty, use the wrapper function lucid

Rho_Z_Mu A scalar. This parameter is the LASSO penalty to regularize cluster-specific means for omics data (Z). If user wants to tune the penalty, use the wrapper function lucid

Rho_Z_Cov A scalar. This parameter is the graphical LASSO penalty to estimate sparse cluster-specific variance-covariance matrices for omics data (Z). If user wants to tune the penalty, use the wrapper function lucid

modelName The variance-covariance structure for omics data. See mclust::mclustModelNames for details.

seed An integer to initialize the EM algorithm or imputing missing values. Default is 123.

init_impute Method to initialize the imputation of missing values in LUCID. "mclust" will use mclust::imputeData to implement EM Algorithm for Unrestricted General Location Model to impute the missing values in omics data; lod will initialize the imputation via relacing missing values by LOD / sqrt(2). LOD is determined by the minimum of each variable in omics data.

init_par Method to initialize the EM algorithm. "mclust" will use mclust model to initialize parameters; "random" initialize parameters from uniform distribution.

verbose A flag indicates whether detailed information for each iteration of EM algorithm is printed in console. Default is FALSE.

Value

A list which contains the several features of LUCID, including:

pars Estimates of parameters of LUCID, including beta (effect of exposure), mu (cluster-specific mean for omics data), sigma (cluster-specific variance-covariance matrix for omics data) and gamma (effect estimate of association between latent cluster and outcome)

K Number of latent cluster
modelName Geometric model to estimate variance-covariance matrix for omics data
likelihood The log likelihood of the LUCID model
post.p Posterior inclusion probability (PIP) for assigning observation i to latent cluster j
Z If missing values are observed, this is the complete dataset for omics data with missing values imputed by LUCID

References


Examples

```r
## Not run:
# use simulated data
G <- sim_data$G
Z <- sim_data$Z
Y_normal <- sim_data$Y_normal
Y_binary <- sim_data$Y_binary
cov <- sim_data$Covariate

# fit LUCID model with continuous outcome
fit1 <- est_lucid(G = G, Z = Z, Y = Y_normal, family = "normal", K = 2, seed = 1008)

# fit LUCID model with block-wise missing pattern in omics data
Z_miss_1 <- Z
Z_miss_1[sample(1:nrow(Z), 0.3 * nrow(Z)), ] <- NA
fit2 <- est_lucid(G = G, Z = Z_miss_1, Y = Y_normal, family = "normal", K = 2)

# fit LUCID model with sporadic missing pattern in omics data
Z_miss_2 <- Z
index <- arrayInd(sample(length(Z_miss_2), 0.3 * length(Z_miss_2)), dim(Z_miss_2))
Z_miss_2[index] <- NA
# initialize imputation by imputing
fit3 <- est_lucid(G = G, Z = Z_miss_2, Y = Y_normal, family = "normal", K = 2, seed = 1008, init_impute = "lod")

# initialize imputation by mclust
fit4 <- est_lucid(G = G, Z = Z_miss_2, Y = Y, family = "normal", K = 2, seed = 123, init_impute = "mclust")

# fit LUCID model with binary outcome
fit5 <- est_lucid(G = G, Z = Z, Y = Y_binary, family = "binary", K = 2, seed = 1008)

# fit LUCID model with covariates
fit6 <- est_lucid(G = G, Z = Z, Y = Y_binary, CoY = cov, family = "binary", K = 2, seed = 1008)
```
# use LUCID model to conduct integrated variable selection
# select exposure
fit6 <- est_lucid(G = G, Z = Z, Y = Y_normal, CoY = NULL, family = "normal",
K = 2, seed = 1008, Rho_G = 0.1)
# select omics data
fit7 <- est_lucid(G = G, Z = Z, Y = Y_normal, CoY = NULL, family = "normal",
K = 2, seed = 1008, Rho_Z_Mu = 90, Rho_Z_Cov = 0.1, init_par = "random")

## End(Not run)

### fill_data

**Impute missing data by optimizing the likelihood function**

**Description**

Impute missing data by optimizing the likelihood function

**Usage**

`fill_data(obs, mu, sigma, p, index)`

**Arguments**

- `obs` a vector of length M
- `mu` a matrix of size M x K
- `sigma` a matrix of size M x M x K
- `p` a vector of length K
- `index` a vector of length M, indicating whether a value is missing or not in the raw data

**Value**

an observation with updated imputed value

### gen_ci

**generate bootstrap ci (normal, basic and percentile)**

**Description**

generate bootstrap ci (normal, basic and percentile)

**Usage**

`gen_ci(x, conf = 0.95)`
### Arguments

- **x**: an object return by boot function
- **conf**: A numeric scalar between 0 and 1 to specify confidence level(s) of the required interval(s).

### Value

A matrix, the first column is t0 statistic from original model

<table>
<thead>
<tr>
<th>helix_data</th>
<th>HELIX data</th>
</tr>
</thead>
</table>

### Description

The Human Early-Life Exposome (HELIX) project is multi-center research project that aims to characterize early-life environmental exposures and associate these with omics biomarkers and child health outcomes (Vrijheid, 2014. doi: 10.1289/ehp.1307204). We used a subset of HELIX data from Exposome Data Challenge 2021 (hold by ISGlobal) as an example to illustrate LUCID model.

### Usage

`helix_data`

### Format

A list with 4 matrices corresponding to exposures (G), omics data (Z), outcome (Y) and covariates (CoY)

- **exposure**: 8 exposures to environmental pollutants. Variables end with m represent maternal exposures; end with c represent children exposures
- **omics**: 10 proteins
- **outcome**: A continuous outcome for BMI-z score based on WHO standard, A binary outcome for body mass index categories at 6-11 years old based on WHO reference (0: Thinness or Normal; 1: Overweight or Obese)
- **covariate**: 3 covariates including mother’s bmi, child sex, maternal age
**Istep_Z**

*I-step of LUCID*

**Description**

Impute missing data in Z by maximizing the likelihood given fixed parameters of LUCID

**Usage**

```
Istep_Z(Z, p, mu, sigma, index)
```

**Arguments**

- **Z**: an N by P matrix representing the omics data
- **p**: an N by K matrix representing posterior inclusion probability for each latent cluster
- **mu**: an M by K matrix representing cluster-specific means
- **sigma**: an M by M by K array representing cluster-specific covariance
- **index**: an N by M matrix representing missing values in Z

**Value**

a complete dataset of Z

---

**lucid**

*Fit a lucid model for integrated analysis on exposure, outcome and multi-omics data*

**Description**

Fit a lucid model for integrated analysis on exposure, outcome and multi-omics data

**Usage**

```
lucid(G, Z, Y, CoG = NULL, CoY = NULL, family = "normal", K = 2, Rho_G = 0, Rho_Z_Mu = 0,
```


```r
Rho_Z_Cov = 0,
verbose_tune = FALSE,
...
)
```

### Arguments

**G**  
Exposures, a numeric vector, matrix, or data frame. Categorical variable should be transformed into dummy variables. If a matrix or data frame, rows represent observations and columns correspond to variables.

**Z**  
Oмисs data, a numeric matrix or data frame. Rows correspond to observations and columns correspond to variables.

**Y**  
Outcome, a numeric vector. Categorical variable is not allowed. Binary outcome should be coded as 0 and 1.

**CoG**  
Optional, covariates to be adjusted for estimating the latent cluster. A numeric vector, matrix or data frame. Categorical variable should be transformed into dummy variables.

**CoY**  
Optional, covariates to be adjusted for estimating the association between latent cluster and the outcome. A numeric vector, matrix or data frame. Categorical variable should be transformed into dummy variables.

**family**  
Distribution of outcome. For continuous outcome, use "normal"; for binary outcome, use "binary". Default is "normal".

**K**  
Number of latent clusters (should be greater or equal than 2). Either an integer or a vector of integer. If K is a vector, model selection on K is performed.

**Rho_G**  
A scalar or a vector. This parameter is the LASSO penalty to regularize exposures. If it is a vector, `lucid` will call `tune_lucid` to conduct model selection and variable selection. User can try penalties from 0 to 1.

**Rho_Z_Mu**  
A scalar or a vector. This parameter is the LASSO penalty to regularize cluster-specific means for omics data (Z). If it is a vector, `lucid` will call `tune_lucid` to conduct model selection and variable selection. User can try penalties from 1 to 100.

**Rho_Z_Cov**  
A scalar or a vector. This parameter is the graphical LASSO penalty to estimate sparse cluster-specific variance-covariance matrices for omics data (Z). If it is a vector, `lucid` will call `tune_lucid` to conduct model selection and variable selection. User can try penalties from 0 to 1.

**verbose_tune**  
A flag to print details of tuning process.

...  
Other parameters passed to `est_lucid`

### Value

An optimal lucid model

### Examples

```r
## Not run:
G <- sim_data$G
```
Z <- sim_data$Z
Y_normal <- sim_data$Y_normal
Y_binary <- sim_data$Y_binary
cov <- sim_data$Covariate

# fit lucid model
fit1 <- lucid(G = G, Z = Z, Y = Y_normal, family = "normal")
fit2 <- lucid(G = G, Z = Z, Y = Y_binary, family = "binary", useY = FALSE)

# including covariates
fit3 <- lucid(G = G, Z = Z, Y = Y_binary, family = "binary", CoG = cov)
fit4 <- lucid(G = G, Z = Z, Y = Y_binary, family = "binary", CoY = cov)

# tune K
fit5 <- lucid(G = G, Z = Z, Y = Y_binary, family = "binary", K = 2:5)

# variable selection
fit6 <- lucid(G = G, Z = Z, Y = Y_binary, family = "binary", Rho_G = seq(0.01, 0.1, by = 0.01))
fit7 <- lucid(G = G, Z = Z, Y = Y_binary, family = "binary", Rho_Z_Mu = seq(10, 100, by = 10), Rho_Z_Cov = 0.5,
             init_par = "random", verbose_tune = TRUE)

## End(Not run)

---

plot_lucid

Visualize LUCID model through a Sankey diagram

---

Description

In the Sankey diagram, each node either represents a variable (exposure, omics or outcome) or a latent cluster. Each line represents an association. The color of the node represents variable type, either exposure, omics or outcome. The width of the line represents the effect size of a certain association; the color of the line represents the direction of a certain association.

Usage

plot_lucid(
  x,
  G_color = "dimgray",
  X_color = "#eb8c30",
  Z_color = "#2fa4da",
  Y_color = "#afa58e",
  pos_link_color = "#67928b",
  neg_link_color = "#d1e5eb",
  fontsize = 7
)
predict_lucid

Arguments

- `x`: A LUCID model fitted by `est_lucid`
- `G_color`: Color of node for exposure
- `X_color`: Color of node for latent cluster
- `Z_color`: Color of node for omics data
- `Y_color`: Color of node for outcome
- `pos_link_color`: Color of link corresponds to positive association
- `neg_link_color`: Color of link corresponds to negative association
- `fontsize`: Font size for annotation

Value

A DAG graph created by `sankeyNetwork`

Examples

```r
## Not run:
# prepare data
G <- sim_data$G
Z <- sim_data$Z
Y_normal <- sim_data$Y_normal
Y_binary <- sim_data$Y_binary
cov <- sim_data$Covariate

# plot lucid model
fit1 <- est_lucid(G = G, Z = Z, Y = Y_normal, CoY = NULL, family = "normal", K = 2, seed = 1008)
plot_lucid(fit1)

# change node color
plot_lucid(fit1, G_color = "yellow")
plot_lucid(fit1, Z_color = "red")

# change link color
plot_lucid(fit1, pos_link_color = "red", neg_link_color = "green")

## End(Not run)
```

predict_lucid  Predict cluster assignment and outcome based on LUCID model

Description

Predict cluster assignment and outcome based on LUCID model
predict_lucid

Usage

predict_lucid(model, G, Z, Y = NULL, CoG = NULL, CoY = NULL, response = TRUE)

Arguments

model  
A model fitted and returned by `est_lucid`

G  
Exposures, a numeric vector, matrix, or data frame. Categorical variable should be transformed into dummy variables. If a matrix or data frame, rows represent observations and columns correspond to variables.

Z  
Omnics data, a numeric matrix or data frame. Rows correspond to observations and columns correspond to variables.

Y  
Outcome, a numeric vector. Categorical variable is not allowed. Binary outcome should be coded as 0 and 1.

CoG  
Optional, covariates to be adjusted for estimating the latent cluster. A numeric vector, matrix or data frame. Categorical variable should be transformed into dummy variables.

CoY  
Optional, covariates to be adjusted for estimating the association between latent cluster and the outcome. A numeric vector, matrix or data frame. Categorical variable should be transformed into dummy variables.

response  
If TRUE, when predicting binary outcome, the response will be returned. If FALSE, the linear predictor is returned.

Value

A list contains predicted latent cluster and outcome for each observation

Examples

```r
## Not run:
# prepare data
G <- sim_data$G
Z <- sim_data$Z
Y_normal <- sim_data$Y_normal

# fit lucid model
fit1 <- est_lucid(G = G, Z = Z, Y = Y_normal, K = 2, family = "normal")

# prediction on training set
pred1 <- predict_lucid(model = fit1, G = G, Z = Z, Y = Y_normal)
pred2 <- predict_lucid(model = fit1, G = G, Z = Z)
## End(Not run)
```
print.lucid

Print the output of est_lucid

Description

Print the output of est_lucid

Usage

## S3 method for class 'lucid'
print(x, ...)

Arguments

x   An object of LUCID model, returned by est_lucid
...
   Other arguments to be passed to print

print.sumlucid

Print the output of LUCID in a nicer table

Description

Print the output of LUCID in a nicer table

Usage

## S3 method for class 'sumlucid'
print(x, ...)

Arguments

x   An object returned by summary_lucid
...
   Other parameters to be passed to print
**sim_data**  
*A simulated dataset for LUCID*

**Description**

This is an example dataset to illustrate LUCID model. It is simulated by assuming there are 2 latent clusters in the data. We assume the exposures are associated with latent cluster which ultimately affects the PFAS concentration and liver injury in children. The latent clusters are also characterized by differential levels of metabolites.

**Usage**

sim_data

**Format**

A list with 5 matrices corresponding to exposures (G), omics data (Z), a continuous outcome, a binary outcome and 2 covariates (can be used either as CoX or CoY). Each matrix contains 2000 observations.

- **G** 10 exposures
- **Z** 10 metabolites
- **Y_normal** Outcome, PFAS concentration in children
- **Y_binary** Binary outcome, liver injury status
- **Covariates** 2 continous covariates, can be treated as either CoX or CoY
- **X** Latent clusters

**summary_lucid**  
*Summarize results of LUCID model*

**Description**

Summarize results of LUCID model

**Usage**

summary_lucid(object, boot.se = NULL)

**Arguments**

- **object**  
  A LUCID model fitted by `est_lucid`

- **boot.se**  
  An object returned by `boot_lucid`, which contains the bootstrap confidence intervals
Examples

## Not run:
# use simulated data
G <- sim_data$G
Z <- sim_data$Z
Y_normal <- sim_data$Y_normal

# fit lucid model
fit1 <- est_lucid(G = G, Z = Z, Y = Y_normal, family = "normal", K = 2,
seed = 1008)

# conduct bootstrap resampling
boot1 <- boot_lucid(G = G, Z = Z, Y = Y_normal, model = fit1, R = 100)

# summarize lucid model
summary_lucid(fit1)

# summarize lucid model with bootstrap CIs
summary_lucid(fit1, boot.se = boot1)

## End(Not run)

tune_lucid

A wrapper function to perform model selection for LUCID

Description

Given a grid of K and L1 penalties (including Rho_G, Rho_Z_mu and Rho_Z_Cov), fit LUCID model over all combinations of K and L1 penalties to determine the optimal penalty.

Usage

tune_lucid(
  G,
  Z,
  Y,
  CoG = NULL,
  CoY = NULL,
  family = "normal",
  K = 2:5,
  Rho_G = 0,
  Rho_Z_Mu = 0,
  Rho_Z_Cov = 0,
  ...
)
Arguments

G  Exposures, a numeric vector, matrix, or data frame. Categorical variable should be transformed into dummy variables. If a matrix or data frame, rows represent observations and columns correspond to variables.

Z  Omics data, a numeric matrix or data frame. Rows correspond to observations and columns correspond to variables.

Y  Outcome, a numeric vector. Categorical variable is not allowed. Binary outcome should be coded as 0 and 1.

CoG  Optional, covariates to be adjusted for estimating the latent cluster. A numeric vector, matrix or data frame. Categorical variable should be transformed into dummy variables.

CoY  Optional, covariates to be adjusted for estimating the association between latent cluster and the outcome. A numeric vector, matrix or data frame. Categorical variable should be transformed into dummy variables.

family  Distribution of outcome. For continuous outcome, use "normal"; for binary outcome, use "binary". Default is "normal".

K  Number of latent clusters. An integer greater or equal to 2. If K is a vector, model selection on K is performed

Rho_G  A scalar or a vector. This parameter is the LASSO penalty to regularize exposures. If it is a vector, tune_lucid will conduct model selection and variable selection. User can try penalties from 0 to 1.

Rho_Z_Mu  A scalar or a vector. This parameter is the LASSO penalty to regularize cluster-specific means for omics data (Z). If it is a vector, tune_lucid will conduct model selection and variable selection. User can try penalties from 1 to 100.

Rho_Z_Cov  A scalar or a vector. This parameter is the graphical LASSO penalty to estimate sparse cluster-specific variance-covariance matrices for omics data (Z). If it is a vector, tune_lucid will conduct model selection and variable selection. User can try penalties from 0 to 1.

...  Other parameters passed to est_lucid

Value

A list:

- best_model  the best model over different combination of tuning parameters
- tune_list  a data frame contains combination of tuning parameters and corresponding BIC
- res_model  a list of LUCID models corresponding to each combination of tuning parameters

Examples

```r
## Not run:
# use simulated data
G <- sim_data$G
Z <- sim_data$Z
Y_normal <- sim_data$Y_normal
```
# find the optimal model over the grid of K
```r
tune_K <- tune_lucid(G = G, Z = Z, Y = Y_normal, useY = FALSE, tol = 1e-3,
seed = 1, K = 2:5)
```

# tune penalties
```r
tune_Rho_G <- tune_lucid(G = G, Z = Z, Y = Y_normal, useY = FALSE, tol = 1e-3,
seed = 1, K = 2, Rho_G = c(0.1, 0.2, 0.3, 0.4))
tune_Rho_Z_Mu <- tune_lucid(G = G, Z = Z, Y = Y_normal, useY = FALSE, tol = 1e-3,
seed = 1, K = 2, Rho_Z_Mu = c(10, 20, 30, 40))
tune_Rho_Z_Cov <- tune_lucid(G = G, Z = Z, Y = Y_normal, useY = FALSE, tol = 1e-3,
seed = 1, K = 2, Rho_Z_Cov = c(0.1, 0.2, 0.3))
```

## End(Not run)
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