Package ‘econet’

November 8, 2018

Type Package

Title Estimation of Parameter-Dependent Network Centrality Measures

Version 0.1.81

Description

Provides methods for estimating parameter-dependent network centrality measures with linear-in-means models. Both non-linear least squares and maximum likelihood estimators are implemented. The methods allow for both link and node heterogeneity in network effects, endogenous network formation and the presence of unconnected nodes. The routines also compare the explanatory power of parameter-dependent network centrality measures with those of standard measures of network centrality. Benefits and features of the ‘econet’ package are illustrated using data from Battaglini and Patacchini (2018), which examine the determinants of US campaign contributions when legislators care about the behavior of other legislators to whom they are socially connected. For additional details, see the vignette.

Depends R (>= 3.5.0)

Imports bbmle, igraph, intergraph, Matrix, MASS, minpack.lm, sna, spatstat.utils, stats, tnet, utils, plyr, dplyr

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Encoding UTF-8

LazyData true

RoxygenNote 6.1.1

Suggests testthat, R.rsp

VignetteBuilder R.rsp

NeedsCompilation no

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Repository CRAN

Date/Publication 2018-11-08 11:00:03 UTC
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```
a_db_alumni                  Dataset from Battaglini, Patacchini (2018)
```

**Description**

Dataset from Battaglini, Patacchini (2018)

**Usage**

```r
data("a_db_alumni")
```

**Format**

an object of class `data.frame` with 2176 rows and 48 columns

- `id` unique congressman id.
- `PAC` PAC contributions to a member of Congress (in US dollars).
- `isolate` dummy variable taking value of 1 if the congressman did not graduate from the same institution within eight years with any other congressman.
- `S1-S65` list of school dummies. Each one refers to a different school. The dummy takes value one if the congressman attended the given school.
- `S_other` dummy variable taking value of 1 if the congressman attended any other school not previously listed.
- `party` dummy variable taking value 1 if the congressman is a Democrat, and 0 if the congressman is a Republican.
- `gender` dummy variable taking value of 1 if the congressman is female.
- `nchair` dummy variable taking value of 1 if the congressman is a chair of at least one committee.
- `sen` number of consecutive years in the House of Representatives.
- `margin_b` dummy variable taking value of 1 if the election margin of victory is less than 5%.
**dw** distance to the center in terms of ideology measured using the absolute value of the first dimension of the dw-nominate score created by McCarty et al. (1997).

**majority** dummy variable taking value of 1 if the member of Congress is a member of the party holding the majority of the seats in the House of Representatives.

**relcom** dummy variable taking value of 1 if the congressman is member of a powerful committee (Appropriations, Budget, Rules and Ways and Means).

**time** categorical variable. It takes 1 if the record refers to the 109th congress, 2 if the record refers to the 110th congress, 3 if the record refers to the 111th congress, 4 if the record refers to the 112th congress, 5 if the record refers to the 113th congress.

**weights** for each congressman $i$, the value is equal to the inverse of the variance of PAC contributions received by all congressmen in $i$'s State of election.

### References


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**a_G_alumni_111**

Column-normalized adjacency matrix representing the alumni network of the 111th U.S. Congress.

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**Description**

Column-normalized adjacency matrix representing the alumni network of the 111th U.S. Congress.

**Usage**

data("a_G_alumni_111")

**Format**

an object of class Matrix with 426 rows and 426 columns

**References**

boot

boot: Bootstrap residuals with cross-sectional dependence

Description

boot: Bootstrap residuals with cross-sectional dependence

Usage

```r
## S3 method for class 'econet'
boot(object, hypothesis = c("lim", "het", "het_l", 
                         "het_r", "par"), group = NULL, niter, weights, ...)
```

Arguments

- `object`: an object of class `econet` obtained using 'NLLS' estimation.
- `hypothesis`: string. One of c("lim", "het", "het_l", "het_r", "par").
- `group`: NULL or vector of positive integers specifying the indices for resampling within groups.
- `niter`: number of required iterations.
- `weights`: logical. It is TRUE if the object `object` was estimated using weights, and FALSE otherwise.
- `...`: additional parameters

Details

For additional details, see the vignette

Warning: This function is available only when `net_dep` is run with `estimation == "NLLS"`

Value

A numeric vector containing bootstrapped standard errors (see Anselin, 1990). If the procedure is not feasible, it returns a vector of NAs.

References


See Also

`net_dep`
Examples

```r
# Load data
data("db_cosponsor")
data("G.alumni_111")
db_model_B <- db_cosponsor
G_model_B <- G.cosponsor_111
G_exclusion_restriction <- G.alumni_111
are_factors <- c("party", "gender", "nchair")
db_model_B[are_factors] <- lapply(db_model_B[are_factors], factor)

# Specify formula
f_model_B <- formula("les ~ gender + party + nchair")

# Specify starting values
starting <- c(alpha = 0.23952,
               beta_gender1 = -0.22024,
               beta_party1 = 0.42947,
               beta_nchair1 = 3.09615,
               phi = 0.40038,
               unobservables = 0.07714)

# object Linear-in-means model
lim_model_B <- net_dep(formula = f_model_B, data = db_model_B,
                       G = G_model_B, model = "model_B", estimation = "NLLS",
                       hypothesis = "lim", endogeneity = TRUE, correction = "heckman",
                       first_step = "standard",
                       exclusion_restriction = G_exclusion_restriction,
                       start.val = starting)

# Bootstrap
# Warning: this may take a very long time to run.
# Decrease the number of iterations to reduce runtime.
boot_lim_estimate <- boot(object = lim_model_B, hypothesis = "lim",
                           group = NULL, niter = 1000, weights = FALSE)

boot_lim_estimate
```

Description

Dataset from Battaglini, Patachini (2018)

Usage

```r
data("db.alumni_test")
```
Format

Data set for runtime executions. An object of class `data.frame` with 42 rows and 48 columns

- **id** unique congressman id.
- **PAC** PAC contributions to a member of Congress (in US dollars).
- **isolate** dummy variable taking value of 1 if the congressman did not graduate from the same institution within eight years with any other congressman.
- **S1-S65** list of school dummies. Each one refers to a different school. The dummy takes value one if the congressman attended the given school.
- **S_other** dummy variable taking value of 1 if the congressman attended any other school not previously listed.
- **party** dummy variable taking value 1 if the congressman is a Democrat, and 0 if the congressman is a Republican.
- **gender** dummy variable taking value of 1 if the congressman is female.
- **nchair** dummy variable taking value of 1 if the congressman is a chair of at least one committee.
- **sen** number of consecutive years in the House of Representatives.
- **margin_b** dummy variable taking value of 1 if the election margin of victory is less than 5%.
- **dw** distance to the center in terms of ideology measured using the absolute value of the first dimension of the dw-nominate score created by McCarty et al. (1997).
- **majority** dummy variable taking value of 1 if the member of Congress is a member of the party holding the majority of the seats in the House of Representatives.
- **recom** dummy variable taking value of 1 if the congressman is member of a powerful committee (Appropriations, Budget, Rules and Ways and Means).
- **time** categorical variable. It takes 1 if the record refers to the 109th congress, 2 if the record refers to the 110th congress, 3 if the record refers to the 111th congress, 4 if the record refers to the 112th congress, 5 if the record refers to the 113th congress.
- **weights** for each congressman $i$, the value is equal to the inverse of the variance of PAC contributions received by all congressmen in $i$'s State of election.

---

**db_cosponsor**  
*Dataset from Battaglini, Leone Sciabolazza, Patacchini (2018)*

Description

Dataset from Battaglini, Leone Sciabolazza, Patacchini (2018)

Usage

`data("db_cosponsor")`
Format

an object of class data.frame with 2176 rows and 48 columns

id unique congressman id.

time categorical variable. It takes 1 if the record refers to the 109th congress, 2 if the record refers to the 110th congress, 3 if the record refers to the 111th congress, 4 if the record refers to the 112th congress, 5 if the record refers to the 113th congress.

party dummy variable taking value 1 if the congressman is a Democrat, and 0 if the congressman is a Republican.

gender dummy variable taking value of 1 if the congressman is female.

nchair dummy variable taking value of 1 if the congressman is a chair of at least one committee.

margin election Margin of Victory of the member of Congress.

dw distance to the center in terms of ideology measured using the absolute value of the first dimension of the dw-nominate score created by McCarty et al. (1997).

les weighted average of the following the number of a congressman’s sponsored bill that were introduced, received any action in committee and beyond committee, passed the House, and became law. It differentially weights commemorative, substantive and significant legislation. Created by Volden C. and Wiseman A. E. (2014).

majority dummy variable taking value of 1 if the member of Congress is a member of the party holding the majority of the seats in the House of Representatives.

state_leg dummy variable taking value of 1 if the member of Congress served in state legislature.

state_leg_prof Squire’s (1992) index of state professionalism relative to Congress.

speaker dummy variable taking value of 1 if the member of Congress is speaker of the House.

seniority number of consecutive years in the House of Representatives.

maj_leader dummy variable taking value of 1 if the member of Congress is member of the majority party leadership, as reported by the Almanac of American Politics.

min_leader dummy variable taking value of 1 if the member of Congress is member of the minority party leadership, as reported by the Almanac of American Politics.

nowhite dummy variable taking value of 1 if the member of Congress is Afro-American or Latino, and 0 otherwise.

H.102-251 list of committee dummies. Each one refers to a different committee. The dummy takes value of 1 if the congressman served in the given committee.

weights for each congressman \( i \), the value is equal to the inverse of the variance of LES of all congressmen in \( i \)'s State of election.

References


Squire, P. (1992), "Legislative professionalization and membership diversity in state legislatures", Legislative Studies Quarterly, 17(1) 69-79.

\begin{itemize}
  \item \textit{G_alumni_111} \hspace{1cm} \textit{Column-normalized adjacency matrix representing the alumni network of the 111th U.S. Congress.}
  
  \textbf{Description} \\
  Column-normalized adjacency matrix representing the alumni network of the 111th U.S. Congress.
  
  \textbf{Usage} \\
  \texttt{data("G_alumni_111")}
  
  \textbf{Format} \\
  \texttt{an object of class Matrix with 426 rows and 426 columns}
  
  \textbf{References} \\

  \item \textit{G_cosponsor_111} \hspace{1cm} \textit{Column-normalized adjacency matrix representing the cosponsorship network of the 111th U.S. Congress.}
  
  \textbf{Description} \\
  Column-normalized adjacency matrix representing the cosponsorship network of the 111th U.S. Congress.
  
  \textbf{Usage} \\
  \texttt{data("G_cosponsor_111")}
  
  \textbf{Format} \\
  \texttt{an object of class Matrix with 439 rows and 439 columns}
  
  \textbf{References} \\
\end{itemize}
G_model_A_test

Column-normalized adjacency matrix representing a subsample of the alumni network of the 111th U.S. Congress.

Description

Column-normalized adjacency matrix representing a subsample of the alumni network of the 111th U.S. Congress.

Usage

data("G_model_A_test")

Format

Data set for runtime executions. An object of class Matrix with 42 rows and 42 columns

References


horse_race

Compare the explanatory power of parameter-dependent network centrality measures with those of standard measures of network centrality.

Description

Compare the explanatory power of parameter-dependent network centrality measures with those of standard measures of network centrality.

Usage

horse_race(formula = formula(), centralities = c("indegree", "outdegree", "degree", "betweenness", "incloseness", "outcloseness", "closeness", "eigenvector"), directed = FALSE, weighted = FALSE, normalization = FALSE, data = list(), unobservables = list(), G = list(), model = c("model_A", "model_B"), estimation = c("NLLS", "MLE"), endogeneity = FALSE, first_step = NULL, data_first_step = NULL, exclusion_restriction = NULL, start.val = NULL, to_weight = NULL, time_fixed_effect = NULL, mle_controls = NULL)
Arguments

formula an object of class `formula`: a symbolic description of the model to be fitted. The constant (i.e. intercept) and the autogressive parameter needs not to be specified.

centralities at least one of c("indegree", "outdegree", "degree", "betweenness", "incloseness", "outcloseness", "closeness", "eigenvector").

directed logical. TRUE if the social network is directed, FALSE otherwise.

weighted logical. TRUE if the social network is weighted, FALSE otherwise.

normalization Default is NULL. Alternatively, it can be set to c("bygraph", "bycomponent", "bymaxcomponent", "bymaxgraph"). See details.

data an object of class `data.frame` containing the variables in the model. If data are longitudinal, observations must be ordered by time period and then by individual.

unobservables a numeric vector used to obtain an unbiased estimate of the parameter-dependent centrality when the network is endogenous. See details.

G an object of class `Matrix` representing the social network. Row and column names must be specified and match the order of the observations in data.

model string. One of c("model_A", "model_B"). See details.

estimation string. One of c("NLLS", "MLE"). They are used to implement respectively a non-linear least square and a maximum likelihood estimator.

endogeneity logical. Default is FALSE. If TRUE, `net_dep` implements a two-step correction procedure to control for the endogeneity of the network.

first_step Default is NULL. If `endogeneity` = TRUE, it requires to specify one of c("standard", "fe", "shortest", "coauthors", "degree"). See details.

data_first_step an optional object of class `data.frame`. If provided, it is used to implement the first step of the estimation when `endogeneity` = TRUE.

exclusion_restriction an object of class `Matrix` representing the exogenous matrix used to instrument the endogenous social network, if `unobservables` is non-NULL. Row and column names must be specified and match the order of the observations in data.

start.val an optional list containing the starting values for the estimations. Object names must match the names provided in `formula`. It is also required to specify the value of both the constant and the decay parameter(s).

to_weight an optional vector of weights to be used in the fitting process to indicate that different observations have different variances. Should be NULL or a numeric vector. If non-NULL, weighted non-linear least squares (if `estimation = "NLLS"`) or weighted maximum likelihood (if `estimation = "MLE"`) is estimated.

time_fixed_effect an optional string. It indicates the name of the time index used in formula. It is used for models with longitudinal data.

mle_controls a list allowing the user to set upper and lower bounds for control variables in MLE estimation and the variance for the ML estimator.
Details

A number of different normalization are available to the user:

- bygraph and bycomponent are used to divide degree and closeness centrality by \( n - 1 \), and betweenness centrality by \((n - 1) \times (n - 2)\) if directed = TRUE, or by \((n - 1) \times (n - 2) / 2\) if directed = FALSE. In the former case (i.e. bygraph), \( n \) is equal to the number of nodes in the network. In the latter case (i.e. bycomponent), \( n \) is equal to the number of nodes of the component in which the node is embedded.
- bymaxgraph and bymaxcomponent are used to divide degree, betweenness and closeness centrality by the maximum value of the centrality of the network (bymaxgraph) or component (bymaxcomponent) in which the node is embedded.

If the network is endogenous, the user is required to run separately net_dep and extract from the resulting object the vector of unobservables necessary for obtaining an unbiased estimate of the parameter.dependent centrality. This vector can be passed through the argument unobservables. If endogeneity = TRUE, a two-step estimation is implemented to control for network endogeneity. The argument first_step is used to control for the specification of the first-step model, e.g.:

- first_step = "standard" is used when agents’ connection are predicted by the differences in their characteristics (i.e. those on the right hand side of formula), and an exclusion_restriction: i.e., their connections in a different network.
- first_step = "fe" adds individual fixed effects to the standard model, as in Graham (2017).
- first_step = "shortest" adds to the standard model, the shortest distance between \( i \) and \( j \), excluding the link between \( i \) and \( j \) itself, as in Fafchamps et al (2010).
- first_step = "coauthor" adds to the standard model, the number of shared connections between \( i \) and \( j \), as in Graham (2015).
- first_step = "degree" adds to the standard model, the difference in the degree centrality of \( i \) and \( j \).

For additional details, see the vignette.

Value

A list of two objects:

- A list of estimates, each one setting the decay parameter to zero, and adding one of the centralities to the specification of formula. The last object adds to formula all the selected centralities and the decay parameter is set different from zero.
- An object of class data.frame containing the computed centrality measures.
- A list of first-step estimations used to correct the effect of centrality measures when the network is endogenous.

References

See Also

net_dep

Examples

```r
# Load data
data("db_cosponsor")
data("G_alumni_111")
db_model_B <- db_cosponsor
G_model_B <- G_cosponsor_111
G_exclusion_restriction <- G_alumni_111
are_factors <- c("party", "gender", "nchair")
db_model_B[are_factors] <- lapply(db_model_B[are_factors], factor)

# Specify formula
f_model_B <- formula("les ~ gender + party + nchair")

# Specify starting values
starting <- c(alpha = 0.214094,
  beta_gender1 = -0.212706,
  beta_party1 = 0.478518,
  beta_nchair1 = 3.09234,
  beta_betweenness = 7.06287e-05,
  phi = 0.344787)

# Fit model
horse_model_B <- horse_race(formula = f_model_B,
  centralities = "betweenness",
  directed = TRUE, weighted = TRUE,
  data = db_model_B, G = G_model_B,
  model = "model_b", estimation = "NLLS",
  start.val = starting)

# Store and print results
summary(horse_model_B)
summary(horse_model_B, centrality = "betweenness")
horse_model_B$centrality

# WARNING, This toy example is provided only for runtime execution.
# Please refer to previous examples for sensible calculations.
data("db_alumni_test")
data("G_model_A_test")
db_model <- db_alumni_test
G_model <- G_model_A_test
f_model <- formula("les ~ dw")
horse_model_test <- horse_race(formula = f_model,
  centralities = "betweenness",
  directed = TRUE, weighted = FALSE, normalization = NULL,
  data = db_model, unobservables = NULL, G = G_model,
  model = "model_A", estimation = "NLLS",
  start.val = c(alpha = -0.31055275,
  beta_dw = 1.50666982,
```
Description

Implement a number of modifications to the linear-in-means model to obtain different weighted versions of Katz-Bonacich centrality.

Usage

```
net_dep(formula = formula(), data = list(), G = list(),
model = c("model_A", "model_B"), estimation = c("NLLS", "MLE"),
hypothesis = c("lim", "het", "het_l", "het_r", "par"),
endogeneity = FALSE, correction = NULL, first_step = NULL,
z = NULL, data_first_step = NULL, exclusion_restriction = NULL,
start.val = NULL, to_weight = NULL, time_fixed_effect = NULL,
mle_controls = NULL, kappa = NULL)
```

Arguments

- **formula**: an object of class formula: a symbolic description of the model to be fitted. The constant (i.e. intercept) and the autogressive parameter needs not to be specified.
- **data**: an object of class data.frame containing the variables in the model. If data are longitudinal, observations must be ordered by time period and then by individual.
- **G**: an object of class Matrix representing the social network. Row and column names must be specified and match the order of the observations in data.
- **model**: string. One of c("model_A", "model_B"). See details.
- **estimation**: string. One of c("NLLS", "MLE"). They are used to implement respectively a non-linear least square and a Maximum Likelihood estimator.
- **hypothesis**: string. One of c("lim", "het", "het_l", "het_r", "par"). See details.
- **endogeneity**: logical. Default is FALSE. If TRUE, net_dep implements a two-step correction procedure to control for the endogeneity of the network.
- **correction**: Default is NULL. If endogeneity = TRUE, it is required to specify if the main regression should use an instrumental variable ("iv") or Heckman ("heckman") approach.
- **first_step**: Default is NULL. If endogeneity = TRUE, it requires to specify one of c("standard" "fe", "shortest", "coauthors", "degree"). See details.
Agent's parameter-dependent centrality is obtained as a function of
• the agent’s characteristics and the performance of its socially connected peers, as in Battaglini, Leone Sciabolazza, Patacchini (2018), if model = "model_B";
• the performance of its socially connected peers, as in Battaglini, Patacchini (2018), if model = "model_A".

Peer effects are assumed to be homogenous if hypothesis = "lim". They are assumed to be heterogenous by setting:
• hypothesis = "het", when peers’ performance is susceptible to agent’s characteristics and model = "model_A".
• hypothesis = "het_1", when peers’ performance is susceptible to agent’s characteristics and model = "model_B".
• hypothesis = "het_r", when agent’s performance is susceptible to peers’ characteristics and model = "model_B".
• hypothesis = "par", when model = "model_B", if the network is formed by interactions between and within two different groups.
When hypothesis is equal to "het", "het_l", or "het_r", the argument z is used to specify the source of heterogeneity; i.e., the attribute affecting the ability of the agent to influence or be influenced by peers. When hypothesis is equal to "par", the argument "z" is used to partition observations in two groups; e.g. the generic element i of vector z takes the value 1 if agent i is member of the first group, and it takes 2 otherwise.

If endogeneity = TRUE, a two-step estimation is implemented to control for network endogeneity. The argument first_step is used to control for the specification of the first-step model, e.g.:

- first_step = "standard" is used when agents' connection are predicted by the differences in their characteristics (i.e. those on the right hand side of formula), and an exclusion_restriction: i.e., their connections in a different network.
- first_step = "fe" adds to the standard model, individual fixed effects, as in Graham (2017).
- first_step = "shortest" adds to the standard model, the shortest distance between i and j, excluding the link between i and j itself, as in Fafchamps et al (2010).
- first_step = "coauthor" adds to the standard model, the number of shared connections between i and j, as in Graham (2015).
- first_step = "degree" adds to the standard model, the difference in the degree centrality of i and j.

The argument start.val is used to specify starting estimates. This can be done with a named list. If a factor is present, a value for each treatment contrast must be provided. Labels of treatment contrasts must be assigned following R model design standards: e.g., a number is appended to contrast names as in contrasts().

The starting value referring to the intercept (constant) must be labelled as "alpha". The label(s) for decay parameter(s) must be:

- "phi", if hypothesis="lim" or hypothesis="het"
- "theta_0", "theta_1", if hypothesis="het_l"
- "eta_0", "eta_1", if hypothesis="het_r"
- "phi_within", "phi_between", if hypothesis="par"

The interaction term when hypothesis="het" must be labelled "gamma". The label to be used for unobservables when endogeneity = TRUE is "unobservables". When estimation = "MLE", it is required to set also the starting value for the variance of the ML estimator. This should be labelled as "sigma".

The argument mle_controls takes a list of two objects. The first is a named numeric vector used to set upper and lower bounds for control variables. The second object is a vector used to set upper (first value) and lower (second value) bounds for the variance of the Maximum Likelihood estimator.

Names in mle_controls must be equal to those used in start.val. For additional details, see the vignette.

Value

A list of three objects: i) Estimates of the main regression; ii) The vector of agents' parameter-dependent centrality; iii) Estimates of the first-step regression (if endogeneity = TRUE)
References


Examples

# Model A

# Load data
data("a_db_alumni") data("a_G_alumni_111")
db_model_A <- a_db_alumni G_model_A <- a_G_alumni_111
are_factors <- c("party", "gender", "nchair", "isolate")
db_model_A[are_factors] <- lapply(db_model_A[are_factors], factor)
db_model_A$PAC <- db_model_A$PAC/1e+06

# Specify formula
f_model_A <- formula("PAC ~ gender + party + nchair + isolate")

# Specify starting values
starting <- c(alpha = 0.47325,
            beta_gender1 = -0.26991,
            beta_party1 = 0.55883,
            beta_nchair1 = -0.17409,
            beta_isolate1 = 0.18813,
            phi = 0.21440)

# Fit Linear-in-means model
lim_model_A <- net_dep(formula = f_model_A, data = db_model_A,
            G = G_model_A, model = "model_A", estimation = "NLLS",
            hypothesis = "lim", start.val = starting)

summary(lim_model_A)
lm_model_A$centrality

# Test Heterogeneity

# Heterogeneous factor
```
z <- as.numeric(as.character(db_model_A$gender))

# Specify formula
f_het_model_A <- formula("PAC ~ party + nchair + isolate")

# Specify starting values
starting <- c(alpha = 0.44835,
              beta_party1 = 0.56004,
              beta_nchair1 = -0.16349,
              beta_isolate1 = 0.21011,
              beta_z = -0.26015,
              phi = 0.34212,
              gamma = -0.49960)

# Fit model
het_model_A <- net_dep(formula = f_het_model_A, data = db_model_A,
                        G = G_model_A, model = "model_A", estimation = "NLLS",
                        hypothesis = "het", z = z, start.val = starting)

summary(het_model_A)
het_model_A$centrality

# Model B

# Load data
data("db_cosponsor")
data("G_alumni_111")
db_model_B <- db_cosponsor
G_model_B <- G_cosponsor_111
G_exclusion_restriction <- G_alumni_111
are_factors <- c("party", "gender", "nchair")
db_model_B[are_factors] <- lapply(db_model_B[are_factors], factor)

# Specify formula
f_model_B <- formula("les ~gender + party + nchair")

# Specify starting values
starting <- c(alpha = 0.23952,
              beta_gender1 = -0.22024,
              beta_party1 = 0.42947,
              beta_nchair1 = 3.09615,
              phi = 0.40038,
              unobservables = 0.07714)

# Fit Linear-in-means model
lim_model_B <- net_dep(formula = f_model_B, data = db_model_B,
                      G = G_model_B, model = "model_B", estimation = "NLLS",
                      hypothesis = "lim", endogeneity = TRUE,
                      correction = "heckman", first_step = "standard",
                      exclusion_restriction = G_exclusion_restriction,
                      start.val = starting)

summary(lim_model_B)
```
```r
lim_model_B$centrality
summary(lim_model_B, print = "first.step")

# Test Heterogeneity

# Heterogeneous factor (node -level)
z <- as.numeric(as.character(db_model_B$gender))

# Specify formula
f_het_model_B <- formula("les ~ party + nchair")

# Specify starting values
starting <- c(alpha = 0.23952,
       beta_party1 = 0.42947,
       beta_nchair1 = 3.09615,
       beta_z = -0.12749,
       theta_0 = 0.42588,
       theta_1 = 0.08007)

# Fit model
het_model_B_l <- net_dep(formula = f_het_model_B,
data = db_model_B,
      G = G_model_B, model = "model_B", estimation = "NLLS",
      hypothesis = "het_l", z = z, start.val = starting)

# Store and print results
summary(het_model_B_l)
het_model_B_l$centrality

# Specify starting values
starting <- c(alpha = 0.04717,
       beta_party1 = 0.51713,
       beta_nchair1 = 3.12683,
       beta_z = 0.01975,
       eta_0 = 1.02789,
       eta_1 = 2.71825)

# Fit model
het_model_B_r <- net_dep(formula = f_het_model_B,
data = db_model_B,
      G = G_model_B, model = "model_B", estimation = "NLLS",
      hypothesis = "het_r", z = z, start.val = starting)

# Store and print results
summary(het_model_B_r)
het_model_B_r$centrality

# Heterogeneous factor (edge -level)
z <- as.numeric(as.character(db_model_B$party))

# Specify starting values
starting <- c(alpha = 0.242486,
       beta_gender1 = -0.229895,
```
quantify

beta_party1 = 0.42848,
beta_nchair1 = 3.0959,
phi_within = 0.396371,
phi_between = 0.414135)

# Fit model
party_model_B <- net_dep(formula = f_model_B, data = db_model_B,
                          G = G_model_B, model = "model_B",
estimation = "NLSS", hypothesis = "par",
z = z, start.val = starting)

# Store and print results
summary(party_estimate_model_B)
party_estimate_model_B$centrality

# WARNING, This toy example is provided only for runtime execution.
# Please refer to previous examples for sensible calculations.
data("db_alumni_test")
data("G_model_A_test")
db_model_A <- db_alumni_test
G_model_A <- G_model_A_test
f_model_A <- formula(les ~ dw)
lim_model_A_test <- net_dep(formula = f_model_A, data = db_model_A,
                           G = G_model_A, model = "model_A", estimation = "NLSS",
hypothesis = "lim", start.val = c(alpha = 0.09030594,
beta_dw = 1.21401940,
phi = 1.47140647))

summary(lim_model_A_test)

quantify

quantify: quantification of marginal effects in linear-in-means models.

Description

quantify: quantification of marginal effects in linear-in-means models.

Usage

## S3 method for class 'econet'
quantify(object, ...)

Arguments

object               first object in the list of outcomes returned by net_dep (available only if the argument model is set to "model_B").
...                  other arguments
Details

`quantify` returns marginal effects for `net_dep` objects when `model = "model_B"` and `hypothesis = "lim"`. For additional details, see the vignette.

Value

an object of class `data.frame` listing direct and indirect variable effects (mean, standard deviation, max, min).

See Also

`net_dep`

Examples

```r
# Load data
data("db_cosponsor")
data("G_alumni_111")
db_model_B <- db_cosponsor
G_model_B <- G_cosponsor_111
G_exclusion_restriction <- G_alumni_111
are_factors <- c("party", "gender", "nchair")
db_model_B[are_factors] <- lapply(db_model_B[are_factors], factor)

# Specify formula
f_model_B <- formula("les ~ gender + party + nchair")

# Specify starting values
starting <- c(alpha = 0.23952,
        beta_gender1 = -0.22024,
        beta_party1 = 0.42947,
        beta_nchair1 = 3.09615,
        phi = 0.40038,
        unobservables = 0.07714)

# Fit Linear-in-means model
lim_model_B <- net_dep(formula = f_model_B, data = db_model_B,
        G = G_model_B, model = "model_B", estimation = "NLLS",
        hypothesis = "lim", endogeneity = TRUE, correction = "heckman",
        first_step = "standard",
        exclusion_restriction = G_exclusion_restriction,
        start.val = starting)
quantify(fit = lim_model_B)

# WARNING, This toy example is provided only for runtime execution. # Please refer to previous examples for sensible calculations.
data("db_alumni_test")
data("G_model_A_test")
db_model <- db_alumni_test
G_model <- G_model_A_test
f_model <- formula("les ~ dw")
```
```r
lim_model_test <- net_dep(formula = f_model, data = db_model,
                          G = G_model, model = "model_B", estimation = "NLLS",
                          hypothesis = "lim", start.val = c(alpha = 0.4553839,
                          beta_dw = -0.7514903,
                          phi = 1.6170539))

quantify(lim_model_test)
```
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