

Package ‘EGAnet’

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Title Exploratory Graph Analysis – a Framework for Estimating the Number of Dimensions in Multivariate Data using Network Psychometrics

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Description Implements the Exploratory Graph Analysis (EGA) framework for dimensionality and psychometric assessment. EGA is part of a new area called network psychometrics that uses undirected network models for the assessment of psychometric properties. EGA estimates the number of dimensions (or factors) using graphical lasso or Triangulated Maximally Filtered Graph (TMFG) and a weighted network community detection algorithm. A bootstrap method for verifying the stability of the dimensions and items in those dimensions is available. The fit of the structure suggested by EGA can be verified using Entropy Fit Indices. A novel approach called Unique Variable Analysis (UVA) can be used to identify and reduce redundant variables in multivariate data. Network loadings, which are roughly equivalent to factor loadings when the data generating model is a factor model, are available. Network scores can also be computed using the network loadings. Dynamic EGA (dynEGA) will estimate dimensions from time series data for individual, group, and sample levels. Golino, H., & Epskamp, S. (2017) <doi:10.1371/journal.pone.0174035>. Golino, H., Shi, D., Christensen, A. P., Garrido, L. E., Nieto, M. D., Sadana, R., & Thiyagarajan, J. A. (2020) <doi:10.31234/osf.io/gzcre>. Christensen, A. P., & Golino, H. (under review) <doi:10.31234/osf.io/hz89e>. Golino, H., Moulder, R. G., Shi, D., Christensen, A. P., Garrido, L. E., Nieto, M. D., Nesselroade, J., Sadana, R., Thiyagarajan, J. A., & Boker, S. M. (2020) <doi:10.31234/osf.io/mtka2>. Christensen, A. P. & Golino, H. (2021) <doi:10.3390/psych3030032>. Christensen, A. P., Garrido, L. E., & Golino, H. (under review) <doi:10.31234/osf.io/4kra2>. Golino, H., Christensen, A. P., Moulder, R. G., Kim, S., & Boker, S. M. (under review) <doi:10.31234/osf.io/tfs7c>.

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EGAnet-package

EGAnet-package

Description

Implements the Exploratory Graph Analysis (EGA; Golino & Epskamp, 2017; Golino, Shi et al., 2020) framework for dimensionality and psychometric assessment. EGA is part of a new area called *network psychometrics* that uses undirected network models for the assessment of psychometric properties. EGA estimates the number of dimensions (or factors) using graphical lasso [EBICglasso](#) or Triangulated Maximally Filtered Graph (TMFG) and a weighted network community detection algorithm (Christensen, Garrido, Golino, under review A). A bootstrap method for verifying the stability of the dimensions and items in those dimensions is available ([bootEGA](#); Christensen & Golino, 2021a). The fit of the structure suggested by EGA can be verified using Entropy Fit Indices ([entropyFit](#), [tefi](#); Golino, Moulder et al., 2020). A novel approach called Unique Variable Analysis (UVA) can be used to identify and reduce redundant variables in multivariate data (Christensen, Garrido, & Golino, under review B). Network loadings ([net.loads](#)), which are roughly equivalent to factor loadings when the data generating model is a factor model, are available (Christensen & Golino, 2021b, 2021c). Network scores ([net.scores](#)) can also be computed using the network loadings. Finally, dynamic EGA ([dynEGA](#)) will estimate dimensions from time series data for individual, group, and sample levels (Golino, Christensen et al., 2021).

Author(s)

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References

- Christensen, A. P., Garrido, L. E., & Golino, H. (under review A). Comparing community detection algorithms in psychological data: A Monte Carlo simulation. *PsyArXiv*.
Related functions: [EGA](#)
- Christensen, A. P., Garrido, L. E., & Golino, H. (under review B). Unique Variable Analysis: A novel approach to detect redundant variables in multivariate data. *PsyArXiv*.
Related functions: [UVA](#)
- Christensen, A. P., & Golino, H. (2021a). Estimating the stability of the number of factors via Bootstrap Exploratory Graph Analysis: A tutorial. *Psych*, 3(3), 479-500.
Related functions: [bootEGA](#), [dimensionStability](#), # and [itemStability](#)
- Christensen, A. P., & Golino, H. (2021b). Factor or network model? Predictions from neural networks. *Journal of Behavioral Data Science*, 1(1), 85-126.
Related functions: [LCT](#)
- Christensen, A. P., & Golino, H. (2021c). On the equivalency of factor and network loadings. *Behavior Research Methods*, 53, 1563-1580.
Related functions: [LCT](#) and [net.loads](#)
- Christensen, A. P., Golino, H., & Silvia, P. J. (2020). A psychometric network perspective on the validity and validation of personality trait questionnaires. *European Journal of Personality*, 34, 1095-1108.
Related functions: [bootEGA](#), [dimensionStability](#), # [EGA](#), [itemStability](#), and [UVA](#)
- Golino, H., Christensen, A. P., Moulder, R., Kim, S., & Boker, S. M. (2021). Modeling latent topics in social media using Dynamic Exploratory Graph Analysis: The case of the right-wing and left-wing trolls in the 2016 US elections. *Psychometrika*.
Related functions: [dynEGA](#) and [simDFM](#)
- Golino, H., & Demetriou, A. (2017). Estimating the dimensionality of intelligence like data using Exploratory Graph Analysis. *Intelligence*, 62, 54-70.
Related functions: [EGA](#)
- Golino, H., & Epskamp, S. (2017). Exploratory graph analysis: A new approach for estimating the number of dimensions in psychological research. *PLoS ONE*, 12, e0174035.
Related functions: [EGA](#)
- Golino, H., Moulder, R., Shi, D., Christensen, A. P., Garrido, L. E., Nieto, M. D., Nesselroade, J., Sadana, R., Thiyagarajan, J. A., & Boker, S. M. (2020). Entropy fit indices: New fit measures for assessing the structure and dimensionality of multiple latent variables. *Multivariate Behavioral Research*.
Related functions: [entropyFit](#), [tefi](#), and [vn.entropy](#)
- Golino, H., Shi, D., Christensen, A. P., Garrido, L. E., Nieto, M. D., Sadana, R., Thiyagarajan, J. A., & Martinez-Molina, A. (2020). Investigating the performance of exploratory graph analysis and traditional techniques to identify the number of latent factors: A simulation and tutorial. *Psychological Methods*, 25, 292-320.
Related functions: [EGA](#)
- Golino, H., Thiyagarajan, J. A., Sadana, M., Teles, M., Christensen, A. P., & Boker, S. M. (under review). Investigating the broad domains of intrinsic capacity, functional ability, and environment: An exploratory graph analysis approach for improving analytical methodologies for measuring healthy aging. *PsyArXiv*.
Related functions: [EGA.fit](#) and [tefi](#)

Jamison, L., Christensen, A. P., & Golino, H. (under review). Optimizing Walktrap's community detection in networks using the Total Entropy Fit Index. *PsyArXiv*.

Related functions: [EGA.fit](#) and [tefi](#)

 boot.ergoInfo

Bootstrap Test for the Ergodicity Information Index

Description

Computes a parametric Bootstrap Test for the Ergodicity Information Index, comparing the empirical Ergodicity Information index to values obtained in data generated using N parametric bootstraps of the correlation matrix estimated using the [dynEGA](#) function, for the population structure. The p-values in the bootstrap test can be calculated as $(\text{sum}(\text{EII} \geq \text{boot.EII}) + 1) / (\text{iter} + 1)$ and as $(\text{sum}(\text{EII} \leq \text{boot.EII}) + 1) / (\text{iter} + 1)$, where EII is the empirical Ergodicity Information Index, boot.EII is the values of the Ergodicity Information Index obtained in the bootstrapped samples, and iter is the number of random samples generated in the simulation. The two-sided p-value is computed as two times the lowest p-value. In the bootstrap Test for the Ergodicity Information Index, the null hypothesis is that the empirical value of EII is equal to the values of EII obtained in multiple individuals with the same structure as the population structure estimated via [dynEGA](#). Small values of p indicate that is very unlikely to obtain an EII as large as the one obtained in the empirical sample if the null hypothesis is true (i.e. all individuals have the same structure as the population structure), thus there is convincing evidence that the empirical Ergodicity Information Index is different than it could be expected if all individuals had a similar latent structure.

Usage

```
boot.ergoInfo(
  dynEGA.pop,
  iter,
  EII,
  use,
  embed,
  tau,
  delta,
  derivatives,
  model,
  model.args = list(),
  algorithm = c("walktrap", "louvain"),
  algorithm.args = list(),
  corr,
  ncores,
  ...
)
```

Arguments

`dynEGA.pop` A `dynEGA` or a `dynEGA.pop.ind` object.

iter	Numeric integer. Number of random samples to generate in the Monte-Carlo simulation. At least 500 is recommended
EII	Numeric. Empirical Ergodicity Information Index obtained via the ergoInfo function.
use	Character. A string indicating what network element will be used to compute the algorithm complexity in the ergoInfo function, the list of edges or the weights of the network. Defaults to use = "edge.list". Current options are: <ul style="list-style-type: none"> • edge.list Calculates the algorithm complexity using the list of edges. • weights Calculates the algorithm complexity using the weights of the network.
embed	Integer. Number of embedded dimensions (the number of observations to be used in the Embed function). For example, an "embed = 5" will use five consecutive observations to estimate a single derivative. Default is "embed = 5".
tau	Integer. Number of observations to offset successive embeddings in the Embed function. A tau of one uses adjacent observations. Default is "tau = 1".
delta	Integer. The time between successive observations in the time series. Default is "delta = 1".
derivatives	Integer. The order of the derivative to be used in the EGA procedure. Default to 1.
model	Character. A string indicating the method to use. Defaults to glasso. Current options are: <ul style="list-style-type: none"> • glasso Estimates the Gaussian graphical model using graphical LASSO with extended Bayesian information criterion to select optimal regularization parameter. This is the default method • TMFG Estimates a Triangulated Maximally Filtered Graph
model.args	List. A list of additional arguments for EBICglasso.qgraph or TMFG
algorithm	A string indicating the algorithm to use or a function from igraph Current options are: <ul style="list-style-type: none"> • walktrap Computes the Walktrap algorithm using cluster_walktrap • louvain Computes the Walktrap algorithm using cluster_louvain
algorithm.args	List. A list of additional arguments for cluster_walktrap , cluster_louvain , or some other community detection algorithm function (see examples)
corr	Type of correlation matrix to compute. The default uses cor_auto . Current options are: <ul style="list-style-type: none"> • cor_auto Computes the correlation matrix using the cor_auto function from qgraph. • pearson Computes Pearson's correlation coefficient using the pairwise complete observations via the cor function. • spearman Computes Spearman's correlation coefficient using the pairwise complete observations via the cor function.

ncores	Numeric. Number of cores to use in computing results. Defaults to <code>parallel::detectCores() / 2</code> or half of your computer's processing power. Set to 1 to not use parallel computing. Recommended to use maximum number of cores minus one If you're unsure how many cores your computer has, then use the following code: <code>parallel::detectCores()</code>
...	Additional arguments. Used for deprecated arguments from previous versions of EGA

Value

Returns a list containing:

boot.ergoInfo	The values of the Ergodicity Information Index obtained in the Monte-Carlo Simulation
p.value.twosided	The p-value of the Monte-Carlo test for the Ergodicity Information Index. The null hypothesis is that the empirical Ergodicity Information index is equal to the expected value of the EII if the all individuals had similar latent structures.
effect	Indicates wheter the empirical EII is greater or less then the Monte-Carlo obtained EII.
plot.dist	Histogram of the bootstrapped ergodicity information index

Author(s)

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Examples

```
## Not run:
\donttest{
dyn1 <- dynEGA.ind.pop(data = sim.dynEGA[,-c(22)], n.embed = 5, tau = 1,
                      delta = 1, id = 21, use.derivatives = 1,
                      model = "glasso", ncores = 2, corr = "pearson")

eii1 <- ergoInfo(data = dyn1)$EII

testing.ergoInfo <- boot.ergoInfo(dynEGA.pop = dyn1, iter = 10,EII = eii1,
embed = 5, tau = 1, delta = 1, derivatives = 1,
model = "glasso", ncores = 2, corr = "pearson")
}
## End(Not run)
```

`boot.wmt`*bootEGA Results of wmt2Data*

Description

[bootEGA](#) results using the "glasso" model and 500 iterations of the Wiener Matrizen-Test 2 (WMT-2)

[bootEGA](#) Results of [wmt2Data](#)

Usage

```
data(boot.wmt)
```

```
data(boot.wmt)
```

Format

A list with 9 objects (see [bootEGA](#))

A list with 8 objects (see [bootEGA](#))

Details

[bootEGA](#) results using the "glasso" model and 500 iterations of the Wiener Matrizen-Test 2 (WMT-2)

Examples

```
data("boot.wmt")
```

```
data("boot.wmt")
```

`bootEGA`*Dimension Stability Analysis of [EGA](#)*

Description

[bootEGA](#) Estimates the number of dimensions of n bootstraps using the empirical (partial) correlation matrix (parametric) or resampling from the empirical dataset (non-parametric). It also estimates a typical median network structure, which is formed by the median or mean pairwise (partial) correlations over the n bootstraps.

Usage

```
bootEGA(
  data,
  n = NULL,
  uni.method = c("expand", "LE"),
  iter,
  type = c("parametric", "resampling"),
  seed = 1234,
  corr = c("cor_auto", "pearson", "spearman"),
  model = c("glasso", "TMFG"),
  model.args = list(),
  algorithm = c("walktrap", "louvain"),
  algorithm.args = list(),
  typicalStructure = TRUE,
  plot.typicalStructure = TRUE,
  plot.type = c("GGally", "qgraph"),
  plot.args = list(),
  ncores,
  ...
)
```

Arguments

<code>data</code>	Matrix or data frame. Includes the variables to be used in the bootEGA analysis
<code>n</code>	Integer. Sample size if data provided is a correlation matrix
<code>uni.method</code>	Character. What unidimensionality method should be used? Defaults to "LE". Current options are: <ul style="list-style-type: none"> • <code>expand</code> Expands the correlation matrix with four variables correlated .50. If number of dimension returns 2 or less in check, then the data are unidimensional; otherwise, regular EGA with no matrix expansion is used. This is the method used in the Golino et al. (2020) <i>Psychological Methods</i> simulation. • <code>LE</code> Applies the leading eigenvalue algorithm (cluster_leading_eigen) on the empirical correlation matrix. If the number of dimensions is 1, then the leading eigenvalue solution is used; otherwise, regular EGA is used. This is the final method used in the Christensen, Garrido, and Golino (2021) simulation.
<code>iter</code>	Numeric integer. Number of replica samples to generate from the bootstrap analysis. At least 500 is recommended
<code>type</code>	Character. A string indicating the type of bootstrap to use. Current options are: <ul style="list-style-type: none"> • <code>"parametric"</code> Generates n new datasets (multivariate normal random distributions) based on the original dataset, via the mvrnorm function • <code>"resampling"</code> Generates n random subsamples of the original data
<code>seed</code>	Numeric. Seed to reproduce results. Defaults to 1234. For random results, set to NULL

corr	Type of correlation matrix to compute. The default uses <code>cor_auto</code> . Current options are: <ul style="list-style-type: none"> • <code>cor_auto</code> Computes the correlation matrix using the <code>cor_auto</code> function from <code>qgraph</code>. • <code>pearson</code> Computes Pearson's correlation coefficient using the pairwise complete observations via the <code>cor</code> function. • <code>spearman</code> Computes Spearman's correlation coefficient using the pairwise complete observations via the <code>cor</code> function.
model	Character. A string indicating the method to use. Current options are: <ul style="list-style-type: none"> • <code>glasso</code> Estimates the Gaussian graphical model using graphical LASSO with extended Bayesian information criterion to select optimal regularization parameter. This is the default method • <code>TMFG</code> Estimates a Triangulated Maximally Filtered Graph
model.args	List. A list of additional arguments for <code>EBICglasso.qgraph</code> or <code>TMFG</code>
algorithm	A string indicating the algorithm to use or a function from <code>igraph</code> Current options are: <ul style="list-style-type: none"> • <code>walktrap</code> Computes the Walktrap algorithm using <code>cluster_walktrap</code> • <code>louvain</code> Computes the Walktrap algorithm using <code>cluster_louvain</code>
algorithm.args	List. A list of additional arguments for <code>cluster_walktrap</code> , <code>cluster_louvain</code> , or some other community detection algorithm function (see examples)
typicalStructure	Boolean. If TRUE, returns the typical network of partial correlations (estimated via graphical lasso or via TMFG) and estimates its dimensions. The "typical network" is the median of all pairwise correlations over the n bootstraps. Defaults to TRUE
plot.typicalStructure	Boolean. If TRUE, returns a plot of the typical network (partial correlations), which is the median of all pairwise correlations over the n bootstraps, and its estimated dimensions. Defaults to TRUE
plot.type	Character. Plot system to use. Current options are <code>qgraph</code> and <code>GGally</code> . Defaults to "GGally".
plot.args	List. A list of additional arguments for the network plot. For <code>plot.type = "qgraph"</code> : <ul style="list-style-type: none"> • <code>vsize</code> Size of the nodes. Defaults to 6. For <code>plot.type = "GGally"</code> (see <code>ggnet2</code> for full list of arguments): <ul style="list-style-type: none"> • <code>vsize</code> Size of the nodes. Defaults to 6. • <code>label.size</code> Size of the labels. Defaults to 5. • <code>alpha</code> The level of transparency of the nodes, which might be a single value or a vector of values. Defaults to 0.7. • <code>edge.alpha</code> The level of transparency of the edges, which might be a single value or a vector of values. Defaults to 0.4.

- `legend.names` A vector with names for each dimension
- `color.palette` The color palette for the nodes. For custom colors, enter HEX codes for each dimension in a vector. See [color_palette_EGA](#) for more details and examples

`ncores` Numeric. Number of cores to use in computing results. Defaults to `parallel::detectCores() / 2` or half of your computer's processing power. Set to 1 to not use parallel computing

If you're unsure how many cores your computer has, then use the following code: `parallel::detectCores()`

`...` Additional arguments. Used for deprecated arguments from previous versions of [EGA](#)

Value

Returns a list containing:

<code>iter</code>	Number of replica samples in bootstrap
<code>boot.ndim</code>	Number of dimensions identified in each replica sample
<code>boot.wc</code>	Item allocation for each replica sample
<code>bootGraphs</code>	Networks of each replica sample
<code>summary.table</code>	Summary table containing number of replica samples, median, standard deviation, standard error, 95% confidence intervals, and quantiles (lower = 2.5% and upper = 97.5%)
<code>frequency</code>	Proportion of times the number of dimensions was identified (e.g., .85 of 1,000 = 850 times that specific number of dimensions was found)
<code>EGA</code>	Output of the original EGA results
<code>typicalGraph</code>	A list containing: <ul style="list-style-type: none"> • <code>graph</code> Network matrix of the median network structure • <code>typical.dim.variables</code> An ordered matrix of item allocation • <code>wc</code> Item allocation of the median network

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References

- # Original implementation of bootEGA
Christensen, A. P., & Golino, H. (2021). Estimating the stability of the number of factors via Bootstrap Exploratory Graph Analysis: A tutorial. *Psych*, 3(3), 479-500.
- # Structural consistency (see [dimensionStability](#))
Christensen, A. P., Golino, H., & Silvia, P. J. (2020). A psychometric network perspective on the validity and validation of personality trait questionnaires. *European Journal of Personality*, 34(6), 1095-1108.

See Also

[EGA](#) to estimate the number of dimensions of an instrument using EGA and [CFA](#) to verify the fit of the structure suggested by EGA using confirmatory factor analysis.

Examples

```
# Load data
wmt <- wmt2[,7:24]

# bootEGA glasso example
## plot.type = "qgraph" used for CRAN checks
## plot.type = "GGally" is the default
boot.wmt <- bootEGA(data = wmt, iter = 500, plot.type = "qgraph",
  type = "parametric", ncores = 2)

# bootEGA TMFG example
boot.wmt <- bootEGA(data = wmt, iter = 500, model = "TMFG",
  plot.type = "qgraph", type = "parametric", ncores = 2, seed = 1234)

# bootEGA Louvain example
boot.wmt <- bootEGA(data = wmt, iter = 500, algorithm = "louvain",
  plot.type = "qgraph", type = "parametric", ncores = 2, seed = 1234)

# bootEGA Spinglass example
boot.wmt <- bootEGA(data = wmt, iter = 500, model = "TMFG", plot.type = "qgraph",
  algorithm = igraph::cluster_spinglass, type = "parametric", ncores = 2)
```

CFA

CFA Fit of [EGA](#) Structure

Description

Verifies the fit of the structure suggested by [EGA](#) using confirmatory factor analysis

Usage

```
CFA(ega.obj, data, estimator, plot.CFA = TRUE, layout = "spring", ...)
```

Arguments

ega.obj	An EGA object
data	A dataframe with the variables to be used in the analysis
estimator	The estimator used in the confirmatory factor analysis. 'WLSMV' is the estimator of choice for ordinal variables. 'ML' or 'WLS' for interval variables. See lavOptions for more details

<code>plot.CFA</code>	Logical. Should the CFA structure with its standardized loadings be plot? Defaults to TRUE
<code>layout</code>	Layout of plot (see semPaths). Defaults to "spring"
<code>...</code>	Arguments passed to cfa

Value

Returns a list containing:

<code>fit</code>	Output from cfa
<code>summary</code>	Summary output from lavaan-class
<code>fit.measures</code>	Fit measures: chi-squared, degrees of freedom, p-value, CFI, RMSEA, GFI, and NFI. Additional fit measures can be applied using the fitMeasures function (see examples)

Author(s)

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References

- Christensen, A. P., Gross, G. M., Golino, H., Silvia, P. J., & Kwapil, T. R. (2019). Exploratory graph analysis of the Multidimensional Schizotypy Scale. *Schizophrenia Research*, *206*, 43-51.
- Golino, H., & Epskamp, S. (2017). Exploratory graph analysis: A new approach for estimating the number of dimensions in psychological research. *PLoS ONE*, *12*, e0174035.

See Also

[EGA](#) to estimate the number of dimensions of an instrument using EGA and [bootEGA](#) to investigate the stability of EGA's estimation via bootstrap.

Examples

```
# Load data
wmt <- wmt2[,7:24]

# Estimate EGA
ega.wmt <- EGA(data = wmt, plot.EGA = FALSE)

# Fit CFA model to EGA results
cfa.wmt <- CFA(ega.obj = ega.wmt, estimator = 'WLSMV', plot.CFA = TRUE, data = wmt)

# Additional fit measures
lavaan::fitMeasures(cfa.wmt$fit, fit.measures = "all")

# Load data
```

```

intel <- intelligenceBattery[,8:66]

# Estimate EGA
ega.intel <- EGA(data = intel, plot.EGA = FALSE)

# Fit CFA model to EGA results
cfa.intel <- CFA(ega.obj = ega.intel, estimator = 'WLSMV', plot.CFA = TRUE,
data = intel)

```

color_palette_EGA [EGA Color Palettes](#)

Description

Color palettes for plotting [ggnet2 EGA](#) network plots

Usage

```
color_palette_EGA(name, wc, sorted = FALSE)
```

Arguments

name	<p>Character. Name of color scheme (see RColorBrewer). Defaults to "polychrome". EGA palettes:</p> <ul style="list-style-type: none"> • "polychrome" Default 20 color palette • "grayscale" "grayscale", "greyscale", or "colorblind" will produce plots suitable for publication purposes • "blue.ridge1" Palette inspired by the Blue Ridge Mountains • "blue.ridge2" Second palette inspired by the Blue Ridge Mountains • "rainbow" Rainbow colors. Default for qgraph • "rio" Palette inspired by Rio de Janeiro, Brazil • "itacare" Palette inspired by Itacare, Brazil <p>For custom colors, enter HEX codes for each dimension in a vector</p>
wc	<p>Vector. A vector representing the community (dimension) membership of each node in the network. NA values mean that the node was disconnected from the network</p>
sorted	<p>Boolean. Should colors be sorted by wc? Defaults to TRUE</p>

Value

Vector of colors for community memberships

Author(s)

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Examples

```
# Default
color_palette_EGA(name = "polychrome", wc = ega.wmt$wc)

# Blue Ridge Moutains 1
color_palette_EGA(name = "blue.ridge1", wc = ega.wmt$wc)

# Custom
color_palette_EGA(name = "#7FD1B9", wc = ega.wmt$wc)
```

compare.EGA.plots *Visually Compares [EGAnet](#) plots*

Description

Organizes EGA plots for comparison. Ensures that nodes are placed in the same layout to maximize comparison. Community memberships are also homogenized across EGA outputs to enhance interpretation

Usage

```
compare.EGA.plots(
  ...,
  input_list = NULL,
  base_plot = 1,
  labels,
  rows,
  columns,
  plot.type = c("GGally", "qgraph"),
  plot.args = list()
)
```

Arguments

...	EGAnet objects
input_list	List. Bypasses ... argument in favor of using a list as an input
base_plot	Numeric. Plot to be used as the base for the configuration of the networks. Uses the number of the order in which the plots are input. Defaults to 1 or the first plot
labels	Character vector. Labels for each EGAnet object
rows	Numeric. Number of rows to spread plots across

<code>columns</code>	Numeric. Number of columns to spread plots down
<code>plot.type</code>	Character. Plot system to use. Current options are qgraph and GGally . Defaults to "GGally"
<code>plot.args</code>	List. A list of additional arguments for the network plot. For <code>plot.type = "qgraph"</code> : <ul style="list-style-type: none"> • <code>vsize</code> Size of the nodes. Defaults to 6. For <code>plot.type = "GGally"</code> (see ggnet2 for full list of arguments): <ul style="list-style-type: none"> • <code>vsize</code> Size of the nodes. Defaults to 6. • <code>label.size</code> Size of the labels. Defaults to 5. • <code>alpha</code> The level of transparency of the nodes, which might be a single value or a vector of values. Defaults to 0.7. • <code>edge.alpha</code> The level of transparency of the edges, which might be a single value or a vector of values. Defaults to 0.4. • <code>legend.names</code> A vector with names for each dimension • <code>color.palette</code> The color palette for the nodes. For custom colors, enter HEX codes for each dimension in a vector. See color_palette_EGA for more details and examples

Value

Visual comparison of [EGAnet](#) objects

Author(s)

Alexander Christensen <alexpaulchristensen@gmail.com>

Examples

```
# obtain SAPA items
items <- psychTools::spi[,c(11:20)]
```

depression

Depression Data

Description

A response matrix (n = 574) of the Beck Depression Inventory, Beck Anxiety Inventory and the Athens Insomnia Scale.

A response matrix (n = 574) of the Beck Depression Inventory, Beck Anxiety Inventory and the Athens Insomnia Scale.

Usage

```
data(depression)
```

```
data(depression)
```


Format

A 574x78 response matrix

A 574x78 response matrix

Examples

```
data("depression")
```

```
data("depression")
```

dimensionStability *Dimension Stability Statistics from* [bootEGA](#)

Description

Based on the [bootEGA](#) results, this function computes the stability of dimensions. This is computed by assessing the proportion of times the original dimension is exactly replicated in across bootstrap samples

Usage

```
dimensionStability(bootega.obj, ...)
```

Arguments

bootega.obj A [bootEGA](#) object

... Additional arguments. Used for deprecated arguments from previous versions of dimStability

Value

Returns a list containing:

dimension.stability

A list containing:

- structural.consistency The proportion of times that each empirical [EGA](#) dimension *exactly* replicates across the [bootEGA](#) samples
- average.item.stability The average item stability in each empirical [EGA](#) dimension

item.stability Results from [itemStability](#)

Author(s)

Hudson Golino <hfg9s at virginia.edu> and Alexander P. Christensen <alexpaulchristensen@gmail.com>

References

Christensen, A. P., & Golino, H. (2021). Estimating the stability of the number of factors via Bootstrap Exploratory Graph Analysis: A tutorial. *Psych*, 3(3), 479-500.

Christensen, A. P., Golino, H., & Silvia, P. J. (2020). A psychometric network perspective on the validity and validation of personality trait questionnaires. *European Journal of Personality*, 34(6), 1095-1108.

See Also

[EGA](#) to estimate the number of dimensions of an instrument using EGA and [CFA](#) to verify the fit of the structure suggested by EGA using confirmatory factor analysis.

Examples

```
# Load data
wmt <- wmt2[,7:24]

## Not run: # Estimate EGA network
## plot.type = "qgraph" used for CRAN checks
## plot.type = "GGally" is the default
ega.wmt <- EGA(data = wmt, model = "glasso", plot.type = "qgraph")

# Estimate bootstrap EGA
boot.wmt <- bootEGA(data = wmt, iter = 500, typicalStructure = TRUE,
plot.typicalStructure = TRUE, model = "glasso", plot.type = "qgraph",
type = "parametric", ncores = 2)

## End(Not run)

# Estimate stability statistics
res <- dimensionStability(boot.wmt)
res$dimension.stability

# Changing plot features (ggplot2)
## Changing colors (ignore warnings)
### qgraph Defaults
res$item.stability$plot +
  ggplot2::scale_color_manual(values = rainbow(length(
res$dimension.stability$structural.consistency)))

### Pastel
res$item.stability$plot +
  ggplot2::scale_color_brewer(palette = "Pastel1")

## Changing Legend (ignore warnings)
res$item.stability$plot +
  ggplot2::scale_color_discrete(labels = "Intelligence")
```

dnn.weights

Loadings Comparison Test Deep Learning Neural Network Weights

Description

A list of weights from four different neural network models: random vs. non-random model (r_nr_weights), low correlation factor vs. network model (lf_n_weights), high correlation with variables less than or equal to factors vs. network model (hlf_n_weights), and high correlation with variables greater than factors vs. network model (hgf_n_weights)

A list of weights from four different neural network models: random vs. non-random model (r_nr_weights), low correlation factor vs. network model (lf_n_weights), high correlation with variables less than or equal to factors vs. network model (hlf_n_weights), and high correlation with variables greater than factors vs. network model (hgf_n_weights)

Usage

```
data(dnn.weights)
```

```
data(dnn.weights)
```

Format

A list of with a length of 4

A list of with a length of 4

Examples

```
data("dnn.weights")
```

```
data("dnn.weights")
```

dynEGA

Dynamic Exploratory Graph Analysis

Description

Estimates dynamic factors in multivariate time series (i.e. longitudinal data, panel data, intensive longitudinal data) at multiple time scales, in different levels of analysis: individuals (intraindividual structure), groups or population (structure of the population). Exploratory graph analysis is applied in the derivatives estimated using generalized local linear approximation ([glla](#)). Instead of estimating factors by modeling how variables are covarying, as in traditional EGA, dynEGA is a dynamic model that estimates the factor structure by modeling how variables are changing together. GLLA is a filtering method for estimating derivatives from data that uses time delay embedding and a variant of Savitzky-Golay filtering to accomplish the task.

Usage

```

dynEGA(
  data,
  n.embed,
  tau = 1,
  delta = 1,
  level = c("individual", "group", "population"),
  id = NULL,
  group = NULL,
  use.derivatives = 1,
  model = c("glasso", "TMFG"),
  model.args = list(),
  algorithm = c("walktrap", "louvain"),
  algorithm.args = list(),
  corr = c("cor_auto", "pearson", "spearman"),
  ncores,
  ...
)

```

Arguments

<code>data</code>	A dataframe with the variables to be used in the analysis. The dataframe should be in a long format (i.e. observations for the same individual (for example, individual 1) are placed in order, from time 1 to time t, followed by the observations from individual 2, also ordered from time 1 to time t.)
<code>n.embed</code>	Integer. Number of embedded dimensions (the number of observations to be used in the Embed function). For example, an "n.embed = 5" will use five consecutive observations to estimate a single derivative.
<code>tau</code>	Integer. Number of observations to offset successive embeddings in the Embed function. A tau of one uses adjacent observations. Default is "tau = 1".
<code>delta</code>	Integer. The time between successive observations in the time series. Default is "delta = 1".
<code>level</code>	Character. A string indicating the level of analysis. If the interest is in modeling the intraindividual structure only (one dimensionality structure per individual), then <code>level</code> should be set to "individual". If the interest is in the structure of a group of individuals, then <code>level</code> should be set to "group". Finally, if the interest is in the population structure, then <code>level</code> should be set to "population". Current options are: <ul style="list-style-type: none"> • <code>individual</code> Estimates the dynamic factors per individual. This should be the preferred method if one is interested in the factor structure of individuals. An additional parameter ("<code>id</code>") needs to be provided identifying each individual. • <code>group</code> Estimates the dynamic factors for each group. An additional parameter ("<code>group</code>") needs to be provided identifying the group membership. • <code>population</code> Estimates the dynamic factors of the population
<code>id</code>	Numeric. Number of the column identifying each individual.

group	Numeric or character. Number of the column identifying group membership. Must be specified only if level = "group".
use.derivatives	Integer. The order of the derivative to be used in the EGA procedure. Default to 1.
model	Character. A string indicating the method to use. Current options are: <ul style="list-style-type: none"> • <code>glasso</code> Estimates the Gaussian graphical model using graphical LASSO with extended Bayesian information criterion to select optimal regularization parameter. This is the default method • <code>TMFG</code> Estimates a Triangulated Maximally Filtered Graph
model.args	List. A list of additional arguments for <code>EBICglasso.qgraph</code> or <code>TMFG</code>
algorithm	A string indicating the algorithm to use or a function from <code>igraph</code> Current options are: <ul style="list-style-type: none"> • <code>walktrap</code> Computes the Walktrap algorithm using <code>cluster_walktrap</code> • <code>louvain</code> Computes the Walktrap algorithm using <code>cluster_louvain</code>
algorithm.args	List. A list of additional arguments for <code>cluster_walktrap</code> , <code>cluster_louvain</code> , or some other community detection algorithm function (see examples)
corr	Type of correlation matrix to compute. The default uses "pearson". Current options are: <ul style="list-style-type: none"> • <code>cor_auto</code> Computes the correlation matrix using the <code>cor_auto</code> function from <code>qgraph</code>. • <code>pearson</code> Computes Pearson's correlation coefficient using the pairwise complete observations via the <code>cor</code> function. • <code>spearman</code> Computes Spearman's correlation coefficient using the pairwise complete observations via the <code>cor</code> function.
ncores	Numeric. Number of cores to use in computing results. Defaults to <code>parallel::detectCores() / 2</code> or half of your computer's processing power. Set to 1 to not use parallel computing. Recommended to use maximum number of cores minus one If you're unsure how many cores your computer has, then use the following code: <code>parallel::detectCores()</code>
...	Additional arguments. Used for deprecated arguments from previous versions of <code>EGA</code>

Author(s)

Hudson Golino <hfg9s at virginia.edu>

References

Boker, S. M., Deboeck, P. R., Edler, C., & Keel, P. K. (2010) Generalized local linear approximation of derivatives from time series. In S.-M. Chow, E. Ferrer, & F. Hsieh (Eds.), *The Notre Dame series on quantitative methodology. Statistical methods for modeling human dynamics: An interdisciplinary dialogue*, (p. 161-178). *Routledge/Taylor & Francis Group*.

Deboeck, P. R., Montpetit, M. A., Bergeman, C. S., & Boker, S. M. (2009) Using derivative estimates to describe intraindividual variability at multiple time scales. *Psychological Methods*, *14*(4), 367-386.

Golino, H., Christensen, A. P., Moulder, R. G., Kim, S., & Boker, S. M. (2021). Modeling latent topics in social media using Dynamic Exploratory Graph Analysis: The case of the right-wing and left-wing trolls in the 2016 US elections. *Psychometrika*.

Savitzky, A., & Golay, M. J. (1964). Smoothing and differentiation of data by simplified least squares procedures. *Analytical Chemistry*, *36*(8), 1627-1639.

Examples

```
# Population structure:
## plot.type = "qgraph" used for CRAN checks
## plot.type = "GGally" is the default
dyn.random <- dynEGA(data = sim.dynEGA, n.embed = 5, tau = 1,
delta = 1, id = 21, group = 22, use.derivatives = 1,
level = "population", model = "glasso", ncores = 2)

plot(dyn.random, plot.type = "qgraph")

# Group structure:
dyn.group <- dynEGA(data = sim.dynEGA, n.embed = 5, tau = 1,
delta = 1, id = 21, group = 22, use.derivatives = 1,
level = "group", model = "glasso", ncores = 2)

plot(dyn.group, ncol = 2, nrow = 1, plot.type = "qgraph")

# Intraindividual structure (commented out for CRAN tests):
# dyn.individual <- dynEGA(data = sim.dynEGA, n.embed = 5, tau = 1,
# delta = 1, id = 21, group = 22, use.derivatives = 1,
# level = "individual", model = "glasso", ncores = 2)
```

dynEGA.ind.pop

Dynamic EGA used in the mctest.ergoInfo function

Description

Dynamic EGA used in the `mctest.ergoInfo` function. DynEGA estimates dynamic factors in multivariate time series (i.e. longitudinal data, panel data, intensive longitudinal data) at multiple time scales, in different levels of analysis: individuals (intraindividual structure) and population (structure of the population). Exploratory graph analysis is applied in the derivatives estimated using generalized local linear approximation ([glla](#)). Instead of estimating factors by modeling how variables are covarying, as in traditional EGA, dynEGA is a dynamic model that estimates the factor structure by modeling how variables are changing together. GLLA is a filtering method for estimating derivatives from data that uses time delay embedding and a variant of Savitzky-Golay filtering to accomplish the task.

Usage

```

dynEGA.ind.pop(
  data,
  n.embed,
  tau = 1,
  delta = 1,
  id = NULL,
  use.derivatives = 1,
  model = c("glasso", "TMFG"),
  model.args = list(),
  algorithm = c("walktrap", "louvain"),
  algorithm.args = list(),
  corr = c("cor_auto", "pearson", "spearman"),
  ncores,
  ...
)

```

Arguments

data	A dataframe with the variables to be used in the analysis. The dataframe should be in a long format (i.e. observations for the same individual (for example, individual 1) are placed in order, from time 1 to time t, followed by the observations from individual 2, also ordered from time 1 to time t.)
n.embed	Integer. Number of embedded dimensions (the number of observations to be used in the Embed function). For example, an "n.embed = 5" will use five consecutive observations to estimate a single derivative.
tau	Integer. Number of observations to offset successive embeddings in the Embed function. A tau of one uses adjacent observations. Default is "tau = 1".
delta	Integer. The time between successive observations in the time series. Default is "delta = 1".
id	Numeric. Number of the column identifying each individual.
use.derivatives	Integer. The order of the derivative to be used in the EGA procedure. Default to 1.
model	Character. A string indicating the method to use. Defaults to glasso. Current options are: <ul style="list-style-type: none"> • glasso Estimates the Gaussian graphical model using graphical LASSO with extended Bayesian information criterion to select optimal regularization parameter. This is the default method • TMFG Estimates a Triangulated Maximally Filtered Graph
model.args	List. A list of additional arguments for EBICglasso.qgraph or TMFG
algorithm	A string indicating the algorithm to use or a function from igraph Current options are: <ul style="list-style-type: none"> • walktrap Computes the Walktrap algorithm using cluster_walktrap • louvain Computes the Walktrap algorithm using cluster_louvain

algorithm.args	List. A list of additional arguments for <code>cluster_walktrap</code> , <code>cluster_louvain</code> , or some other community detection algorithm function (see examples)
corr	Type of correlation matrix to compute. The default uses <code>cor_auto</code> . Current options are: <ul style="list-style-type: none"> • <code>cor_auto</code> Computes the correlation matrix using the <code>cor_auto</code> function from <code>qgraph</code>. • <code>pearson</code> Computes Pearson's correlation coefficient using the pairwise complete observations via the <code>cor</code> function. • <code>spearman</code> Computes Spearman's correlation coefficient using the pairwise complete observations via the <code>cor</code> function.
ncores	Numeric. Number of cores to use in computing results. Defaults to <code>parallel::detectCores() / 2</code> or half of your computer's processing power. Set to 1 to not use parallel computing. Recommended to use maximum number of cores minus one If you're unsure how many cores your computer has, then use the following code: <code>parallel::detectCores()</code>
...	Additional arguments. Used for deprecated arguments from previous versions of EGA

Author(s)

Hudson Golino <hfg9s at virginia.edu>

Examples

```
## Not run:
\donttest{# Population structure:
dyn.ega1 <- dynEGA.ind.pop(data = sim.dynEGA, n.embed = 5, tau = 1,
delta = 1, id = 21, use.derivatives = 1, model = "glasso", ncores = 2,
cor = "pearson")
}

## End(Not run)
```

EBICglasso.qgraph [EBICglasso from qgraph 1.4.4](#)

Description

This function uses the [glasso](#) package (Friedman, Hastie and Tibshirani, 2011) to compute a sparse gaussian graphical model with the graphical lasso (Friedman, Hastie & Tibshirani, 2008). The tuning parameter is chosen using the Extended Bayesian Information criterium (EBIC) described by Foygel & Drton (2010).

Usage

```
EBICglasso.qgraph(
  data,
  n = NULL,
  gamma = 0.5,
  penalize.diagonal = FALSE,
  nlambda = 100,
  lambda.min.ratio = 0.01,
  returnAllResults = FALSE,
  penalizeMatrix,
  countDiagonal = FALSE,
  refit = FALSE,
  ...
)
```

Arguments

<code>data</code>	Data matrix
<code>n</code>	Number of participants
<code>gamma</code>	EBIC tuning parameter. 0.5 is generally a good choice. Setting to zero will cause regular BIC to be used.
<code>penalize.diagonal</code>	Should the diagonal be penalized?
<code>nlambda</code>	Number of lambda values to test.
<code>lambda.min.ratio</code>	Ratio of lowest lambda value compared to maximal lambda
<code>returnAllResults</code>	If TRUE this function does not return a network but the results of the entire glasso path.
<code>penalizeMatrix</code>	Optional logical matrix to indicate which elements are penalized
<code>countDiagonal</code>	Should diagonal be counted in EBIC computation? Defaults to FALSE. Set to TRUE to mimic qgraph < 1.3 behavior (not recommended!).
<code>refit</code>	Logical, should the optimal graph be refitted without LASSO regularization? Defaults to FALSE.
<code>...</code>	Arguments sent to glasso

Details

The glasso is run for 100 values of the tuning parameter logarithmically spaced between the maximal value of the tuning parameter at which all edges are zero, `lambda_max`, and `lambda_max/100`. For each of these graphs the EBIC is computed and the graph with the best EBIC is selected. The partial correlation matrix is computed using [wi2net](#) and returned.

Value

A partial correlation matrix

Author(s)

Sacha Epskamp <mail@sachaepskamp.com>

References

Friedman, J., Hastie, T., & Tibshirani, R. (2008). Sparse inverse covariance estimation with the graphical lasso. *Biostatistics*, 9, 432-441.

#glasso package Jerome Friedman, Trevor Hastie and Rob Tibshirani (2011). glasso: Graphical lasso-estimation of Gaussian graphical models. R package version 1.7.

Foygel, R., & Drton, M. (2010). Extended Bayesian information criteria for Gaussian graphical models. In *Advances in neural information processing systems* (pp. 604-612).

#psych package Revelle, W. (2014) psych: Procedures for Personality and Psychological Research, Northwestern University, Evanston, Illinois, USA. R package version 1.4.4.

#Matrix package Douglas Bates and Martin Maechler (2014). Matrix: Sparse and Dense Matrix Classes and Methods. R package version 1.1-3.

Examples

```
### Using wmt2 dataset from EGAnet ###
data(wmt2)

# Compute correlations:
CorMat <- qgraph::cor_auto(wmt2[,7:24])

# Compute graph with tuning = 0 (BIC):
BICgraph <- EBICglasso.qgraph(CorMat, n = nrow(wmt2), gamma = 0)

# Compute graph with tuning = 0.5 (EBIC)
EBICgraph <- EBICglasso.qgraph(CorMat, n = nrow(wmt2), gamma = 0.5)
```

EGA

Applies the Exploratory Graph Analysis technique

Description

Estimates the number of dimensions of a given dataset or correlation matrix using the graphical lasso ([EBICglasso.qgraph](#)) or the Triangulated Maximally Filtered Graph ([TMFG](#)) network estimation methods.

Usage

```
EGA(
  data,
  n = NULL,
  uni.method = c("expand", "LE"),
  corr = c("cor_auto", "pearson", "spearman"),
  model = c("glasso", "TMFG"),
  model.args = list(),
  algorithm = c("walktrap", "louvain"),
  algorithm.args = list(),
  plot.EGA = TRUE,
  plot.type = c("GGally", "qgraph"),
  plot.args = list(),
  verbose = TRUE,
  ...
)
```

Arguments

data	Matrix or data frame. Variables (down columns) or correlation matrix. If the input is a correlation matrix, then argument n (number of cases) is required
n	Integer. Sample size if data provided is a correlation matrix
uni.method	Character. What unidimensionality method should be used? Defaults to "LE". Current options are: <ul style="list-style-type: none"> • expand Expands the correlation matrix with four variables correlated .50. If number of dimension returns 2 or less in check, then the data are unidimensional; otherwise, regular EGA with no matrix expansion is used. This is the method used in the Golino et al. (2020) <i>Psychological Methods</i> simulation. • LE Applies the leading eigenvalue algorithm (cluster_leading_eigen) on the empirical correlation matrix. If the number of dimensions is 1, then the leading eigenvalue solution is used; otherwise, regular EGA is used. This is the final method used in the Christensen, Garrido, and Golino (2021) simulation.
corr	Type of correlation matrix to compute. The default uses cor_auto . Current options are: <ul style="list-style-type: none"> • cor_auto Computes the correlation matrix using the cor_auto function from qgraph. • pearson Computes Pearson's correlation coefficient using the pairwise complete observations via the cor function. • spearman Computes Spearman's correlation coefficient using the pairwise complete observations via the cor function.
model	Character. A string indicating the method to use. Defaults to "glasso". Current options are:

	<ul style="list-style-type: none"> • <code>glasso</code> Estimates the Gaussian graphical model using graphical LASSO with extended Bayesian information criterion to select optimal regularization parameter • <code>TMFG</code> Estimates a Triangulated Maximally Filtered Graph
<code>model.args</code>	List. A list of additional arguments for <code>EBICglasso.qgraph</code> or <code>TMFG</code>
<code>algorithm</code>	A string indicating the algorithm to use or a function from <code>igraph</code> Defaults to "walktrap". Current options are: <ul style="list-style-type: none"> • <code>walktrap</code> Computes the Walktrap algorithm using <code>cluster_walktrap</code> • <code>louvain</code> Computes the Louvain algorithm using <code>cluster_louvain</code>
<code>algorithm.args</code>	List. A list of additional arguments for <code>cluster_walktrap</code> , <code>cluster_louvain</code> , or some other community detection algorithm function (see examples)
<code>plot.EGA</code>	Boolean. If TRUE, returns a plot of the network and its estimated dimensions. Defaults to TRUE
<code>plot.type</code>	Character. Plot system to use. Current options are <code>qgraph</code> and <code>GGally</code> . Defaults to "GGally"
<code>plot.args</code>	List. A list of additional arguments for the network plot. For <code>plot.type = "qgraph"</code> : <ul style="list-style-type: none"> • <code>vsize</code> Size of the nodes. Defaults to 6. For <code>plot.type = "GGally"</code> (see <code>ggnet2</code> for full list of arguments): <ul style="list-style-type: none"> • <code>vsize</code> Size of the nodes. Defaults to 6. • <code>label.size</code> Size of the labels. Defaults to 5. • <code>alpha</code> The level of transparency of the nodes, which might be a single value or a vector of values. Defaults to 0.7. • <code>edge.alpha</code> The level of transparency of the edges, which might be a single value or a vector of values. Defaults to 0.4. • <code>legend.names</code> A vector with names for each dimension • <code>color.palette</code> The color palette for the nodes. For custom colors, enter HEX codes for each dimension in a vector. See <code>color_palette_EGA</code> for more details and examples
<code>verbose</code>	Boolean. Should network estimation parameters be printed? Defaults to TRUE. Set to FALSE for no print out
<code>...</code>	Additional arguments. Used for deprecated arguments from previous versions of <code>EGA</code>

Details

Two community detection algorithms, Walktrap (Pons & Latapy, 2006) and Louvain (Blondel et al., 2008), are pre-programmed because of their superior performance in simulation studies on psychological data generated from factor models (Christensen & Golino; 2020; Golino et al., 2020). Notably, any community detection algorithm from the `igraph` can be used to estimate the number of communities (see examples).

Value

Returns a list containing:

network	A symmetric network estimated using either the EBICglasso.qgraph or TMFG
wc	A vector representing the community (dimension) membership of each node in the network. NA values mean that the node was disconnected from the network
n.dim	A scalar of how many total dimensions were identified in the network
cor.data	The zero-order correlation matrix

Author(s)

Hudson Golino <hfg9s at virginia.edu>, Alexander P. Christensen <alexpaulchristensen at gmail.com>, Maria Dolores Nieto <acinodam at gmail.com> and Luis E. Garrido <garrido.luiseduardo at gmail.com>

References

- # Louvain algorithm
Blondel, V. D., Guillaume, J.-L., Lambiotte, R., & Lefebvre, E. (2008). Fast unfolding of communities in large networks. *Journal of Statistical Mechanics: Theory and Experiment*, 2008, P10008.
- # Compared all *igraph* community detections algorithms, introduced Louvain algorithm, simulation with continuous and polytomous data
Also implements the Leading Eigenvalue unidimensional method
Christensen, A. P., Garrido, L. E., & Golino, H. (2021). Comparing community detection algorithms in psychological data: A Monte Carlo simulation. *PsyArXiv*.
- # Original simulation and implementation of EGA
Golino, H. F., & Epskamp, S. (2017). Exploratory graph analysis: A new approach for estimating the number of dimensions in psychological research. *PLoS ONE*, 12, e0174035.
- Golino, H. F., & Demetriou, A. (2017). Estimating the dimensionality of intelligence like data using Exploratory Graph Analysis. *Intelligence*, 62, 54-70.
- # Current implementation of EGA, introduced unidimensional checks, continuous and dichotomous data
Golino, H., Shi, D., Christensen, A. P., Garrido, L. E., Nieto, M. D., Sadana, R., & Thiyagarajan, J. A. (2020). Investigating the performance of Exploratory Graph Analysis and traditional techniques to identify the number of latent factors: A simulation and tutorial. *Psychological Methods*, 25, 292-320.
- # Walktrap algorithm
Pons, P., & Latapy, M. (2006). Computing communities in large networks using random walks. *Journal of Graph Algorithms and Applications*, 10, 191-218.

See Also

[bootEGA](#) to investigate the stability of EGA's estimation via bootstrap and [CFA](#) to verify the fit of the structure suggested by EGA using confirmatory factor analysis.

Examples

```
# Estimate EGA
## plot.type = "qgraph" used for CRAN checks
## plot.type = "GGally" is the default
ega.wmt <- EGA(data = wmt2[,7:24], plot.type = "qgraph")

# Summary statistics
summary(ega.wmt)

# Estimate EGAtmfg
ega.wmt <- EGA(data = wmt2[,7:24], model = "TMFG", plot.type = "qgraph")

# Estimate EGA with Louvain algorithm
ega.wmt <- EGA(data = wmt2[,7:24], algorithm = "louvain", plot.type = "qgraph")

# Estimate EGA with Spinglass algorithm
ega.wmt <- EGA(data = wmt2[,7:24],
algorithm = igraph::cluster_spinglass, plot.type = "qgraph")

# Estimate EGA
ega.intel <- EGA(data = intelligenceBattery[,8:66], model = "glasso", plot.EGA = FALSE)

# Summary statistics
summary(ega.intel)
```

EGA.estimate

A Sub-routine Function for EGA

Description

Estimates the number of dimensions of a given dataset or correlation matrix using the graphical lasso ([EBICglasso.qgraph](#)) or the Triangulated Maximally Filtered Graph ([TMFG](#)) network estimation methods.

Usage

```
EGA.estimate(
  data,
  n = NULL,
  model = c("glasso", "TMFG"),
  model.args = list(),
  algorithm = c("walktrap", "louvain"),
  algorithm.args = list(),
  corr = c("cor_auto", "pearson", "spearman"),
  verbose = TRUE,
```

...
)

Arguments

data	Matrix or data frame. Variables (down columns) or correlation matrix. If the input is a correlation matrix, then argument n (number of cases) is required
n	Integer. Sample size if data provided is a correlation matrix
model	Character. A string indicating the method to use. Current options are: <ul style="list-style-type: none"> • <code>glasso</code> Estimates the Gaussian graphical model using graphical LASSO with extended Bayesian information criterion to select optimal regularization parameter. This is the default method • <code>TMFG</code> Estimates a Triangulated Maximally Filtered Graph
model.args	List. A list of additional arguments for <code>EBICglasso.qgraph</code> or <code>TMFG</code>
algorithm	A string indicating the algorithm to use or a function from <code>igraph</code> Current options are: <ul style="list-style-type: none"> • <code>walktrap</code> Computes the Walktrap algorithm using <code>cluster_walktrap</code> • <code>louvain</code> Computes the Walktrap algorithm using <code>cluster_louvain</code>
algorithm.args	List. A list of additional arguments for <code>cluster_walktrap</code> , <code>cluster_louvain</code> , or some other community detection algorithm function (see examples)
corr	Type of correlation matrix to compute. The default uses <code>cor_auto</code> . Current options are: <ul style="list-style-type: none"> • <code>cor_auto</code> Computes the correlation matrix using the <code>cor_auto</code> function from <code>qgraph</code>. • <code>pearson</code> Computes Pearson's correlation coefficient using the pairwise complete observations via the <code>cor</code> function. • <code>spearman</code> Computes Spearman's correlation coefficient using the pairwise complete observations via the <code>cor</code> function.
verbose	Boolean. Should network estimation parameters be printed? Defaults to TRUE. Set to FALSE for no print out
...	Additional arguments. Used for deprecated arguments from previous versions of <code>EGA</code>

Details

Two community detection algorithms, Walktrap (Pons & Latapy, 2006) and Louvain (Blondel et al., 2008), are pre-programmed because of their superior performance in simulation studies on psychological data generated from factor models (Christensen & Golino; 2020; Golino et al., 2020). Notably, any community detection algorithm from the `igraph` can be used to estimate the number of communities (see examples).

Value

Returns a list containing:

<code>estimated.network</code>	A symmetric network estimated using either the EBICglasso.qgraph or TMFG
<code>wc</code>	A vector representing the community (dimension) membership of each node in the network. NA values mean that the node was disconnected from the network
<code>n.dim</code>	A scalar of how many total dimensions were identified in the network
<code>cor.data</code>	The zero-order correlation matrix

Author(s)

Alexander P. Christensen <alexpaulchristensen at gmail.com> and Hudson Golino <hfg9s at virginia.edu>

References

- # Louvain algorithm
Blondel, V. D., Guillaume, J.-L., Lambiotte, R., & Lefebvre, E. (2008). Fast unfolding of communities in large networks. *Journal of Statistical Mechanics: Theory and Experiment*, 2008, P10008.
- # Compared all *igraph* community detections algorithms, introduced Louvain algorithm, simulation with continuous and polytomous data
Christensen, A. P., & Golino, H. (under review). Estimating factors with psychometric networks: A Monte Carlo simulation comparing community detection algorithms. *PsyArXiv*.
- # Original simulation and implementation of EGA
Golino, H. F., & Epskamp, S. (2017). Exploratory graph analysis: A new approach for estimating the number of dimensions in psychological research. *PLoS ONE*, 12, e0174035.
- Golino, H. F., & Demetriou, A. (2017). Estimating the dimensionality of intelligence like data using Exploratory Graph Analysis. *Intelligence*, 62, 54-70.
- # Current implementation of EGA, introduced unidimensional checks, continuous and dichotomous data
Golino, H., Shi, D., Christensen, A. P., Garrido, L. E., Nieto, M. D., Sadana, R., & Thiyagarajan, J. A. (2020). Investigating the performance of Exploratory Graph Analysis and traditional techniques to identify the number of latent factors: A simulation and tutorial. *Psychological Methods*, 25, 292-320.
- # Walktrap algorithm
Pons, P., & Latapy, M. (2006). Computing communities in large networks using random walks. *Journal of Graph Algorithms and Applications*, 10, 191-218.

See Also

[bootEGA](#) to investigate the stability of EGA's estimation via bootstrap and [CFA](#) to verify the fit of the structure suggested by EGA using confirmatory factor analysis.

Examples

```
# Estimate EGA
ega.wmt <- EGA.estimate(data = wmt2[,7:24], model = "glasso")

# Estimate EGAtmfg
ega.wmt <- EGA.estimate(data = wmt2[,7:24], model = "TMFG")

# Estimate EGA with Spinglass
ega.wmt <- EGA.estimate(data = wmt2[,7:24], model = "glasso",
algorithm = igraph::cluster_spinglass)
```

EGA.fit

[EGA Optimal Model Fit using the Total Entropy Fit Index \(tefi\)](#)

Description

Estimates the best fitting model using [EGA](#). The number of steps in the [cluster_walktrap](#) detection algorithm is varied and unique community solutions are compared using [tefi](#).

Usage

```
EGA.fit(
  data,
  n = NULL,
  uni.method = c("expand", "LE"),
  corr = c("cor_auto", "pearson", "spearman"),
  model = c("glasso", "TMFG"),
  algorithm = c("leiden", "walktrap"),
  algorithm.args = list(steps = c(3:8), resolution_parameter = seq(0, 2, 0.001))
)
```

Arguments

data	Matrix or data frame. Dataset or correlation matrix
n	Integer. Sample size (if the data provided is a correlation matrix)
uni.method	Character. What unidimensionality method should be used? Defaults to "LE". Current options are: <ul style="list-style-type: none"> • expand Expands the correlation matrix with four variables correlated .50. If number of dimension returns 2 or less in check, then the data are unidimensional; otherwise, regular EGA with no matrix expansion is used. This is the method used in the Golino et al. (2020) <i>Psychological Methods</i> simulation.

	<ul style="list-style-type: none"> • LE Applies the leading eigenvalue algorithm (<code>cluster_leading_eigen</code>) on the empirical correlation matrix. If the number of dimensions is 1, then the leading eigenvalue solution is used; otherwise, regular EGA is used. This is the final method used in the Christensen, Garrido, and Golino (2021) simulation.
<code>corr</code>	<p>Type of correlation matrix to compute. The default uses <code>cor_auto</code>. Current options are:</p> <ul style="list-style-type: none"> • <code>cor_auto</code> Computes the correlation matrix using the <code>cor_auto</code> function from <code>qgraph</code>. • <code>pearson</code> Computes Pearson's correlation coefficient using the pairwise complete observations via the <code>cor</code> function. • <code>spearman</code> Computes Spearman's correlation coefficient using the pairwise complete observations via the <code>cor</code> function.
<code>model</code>	<p>Character. A string indicating the method to use. Defaults to "glasso" Current options are:</p> <ul style="list-style-type: none"> • "glasso" Estimates the Gaussian graphical model using graphical LASSO with extended Bayesian information criterion to select optimal regularization parameter. See <code>EBICglasso.qgraph</code> • "TMFG" Estimates a Triangulated Maximally Filtered Graph. See <code>TMFG</code>
<code>algorithm</code>	<p>A string indicating the algorithm to use or a function from <code>igraph</code> Defaults to "walktrap". Current options are:</p> <ul style="list-style-type: none"> • <code>walktrap</code> Computes the Walktrap algorithm using <code>cluster_walktrap</code> • <code>leiden</code> Computes the Leiden algorithm using <code>cluster_louvain</code>
<code>algorithm.args</code>	<p>List. A list of additional arguments for <code>cluster_walktrap</code> or <code>cluster_leiden</code>. Options are:</p> <ul style="list-style-type: none"> • <code>steps</code> Number of steps used in the Walktrap algorithm. Defaults to <code>c(3:8)</code> • <code>leiden</code> Resolution parameter used in the Leiden algorithm. Defaults to <code>seq(0, 2, .001)</code>. Higher values lead to smaller communities, lower values lead to larger communities

Value

Returns a list containing:

<code>EGA</code>	The <code>EGA</code> output for the best fitting model
<code>steps</code>	The number of steps used in the best fitting model from the <code>cluster_walktrap</code> algorithm
<code>resolution_parameter</code>	The resolution parameter used in the best fitting model from the <code>cluster_leiden</code> algorithm
<code>EntropyFit</code>	The <code>tefi</code> Index for the unique solutions given the range of steps (vector names represent the number of steps)
<code>Lowest.EntropyFit</code>	The lowest value for the <code>tefi</code> Index

Author(s)

Hudson Golino <hfg9s at virginia.edu> and Alexander P. Christensen <alexpaulchristensen@gmail.com>

References

Entropy fit measures

Golino, H., Moulder, R. G., Shi, D., Christensen, A. P., Garrido, L. E., Neito, M. D., Nesselroade, J., Sadana, R., Thiyagarajan, J. A., & Boker, S. M. (in press). Entropy fit indices: New fit measures for assessing the structure and dimensionality of multiple latent variables. *Multivariate Behavioral Research*.

Simulation for EGA.fit

Jamison, L., Christensen, A. P., & Golino, H. (under review). Optimizing Walktrap's community detection in networks using the Total Entropy Fit Index. *PsyArXiv*.

Leiden algorithm

Traag, V. A., Waltman, L., & Van Eck, N. J. (2019). From Louvain to Leiden: guaranteeing well-connected communities. *Scientific Reports*, 9(1), 1-12.

Walktrap algorithm

Pons, P., & Latapy, M. (2006). Computing communities in large networks using random walks. *Journal of Graph Algorithms and Applications*, 10, 191-218.

See Also

[bootEGA](#) to investigate the stability of EGA's estimation via bootstrap, [EGA](#) to estimate the number of dimensions of an instrument using EGA, and [CFA](#) to verify the fit of the structure suggested by EGA using confirmatory factor analysis.

Examples

```
# Load data
wmt <- wmt2[,7:24]

# Estimate EGA
## plot.type = "qgraph" used for CRAN checks
## plot.type = "GGally" is the default
ega.wmt <- EGA(data = wmt, plot.type = "qgraph")

# Estimate optimal EGA
fit.wmt <- EGA.fit(data = wmt)

# Plot optimal fit
plot(fit.wmt$EGA, plot.type = "qgraph")

# Compare with CFA
cfa.ega <- CFA(ega.wmt, estimator = "WLSMV", data = wmt)
cfa.fit <- CFA(fit.wmt$EGA, estimator = "WLSMV", data = wmt)

lavaan::lavTestLRT(cfa.ega$fit, cfa.fit$fit, method = "satorra.bentler.2001")
```

`ega.wmt`*EGA WMT-2 Data*

Description

[EGA](#) Network of `wmt2Data`

An [EGA](#) using the "glasso" model of the Wiener Matrizen-Test 2 (WMT-2)

Usage

```
data(ega.wmt)
```

```
data(ega.wmt)
```

Format

A 17 x 17 adjacency matrix

A 17 x 17 adjacency matrix

Details

An [EGA](#) using the "glasso" model of the Wiener Matrizen-Test 2 (WMT-2)

Examples

```
data("ega.wmt")
```

```
data("ega.wmt")
```

`Embed`*Time-delay Embedding*

Description

Reorganizes an individual's observed time series into an embedded matrix. The embedded matrix is constructed with replicates of an individual time series that are offset from each other in time. The function requires two parameters, one that specifies the number of observations to be used (i.e. the number of embedded dimensions) and the other that specifies the number of observations to offset successive embeddings.

Usage

```
Embed(x, E, tau)
```

Arguments

x	Vector. An observed time series to be reorganized into a time-delayed embedded matrix.
E	Integer. Number of embedded dimensions or the number of observations to be used. For example, an "E = 5" will generate a matrix with five columns, meaning that five consecutive observations are used to create each row of the embedded matrix.
tau	Integer. Number of observations to offset successive embeddings. A tau of one uses adjacent observations. Default is "tau = 1".

Value

Returns a matrix containing the embedded matrix.

Author(s)

Pascal Deboeck <pascal.deboeck at psych.utah.edu>

References

Deboeck, P. R., Montpetit, M. A., Bergeman, C. S., & Boker, S. M. (2009) Using derivative estimates to describe intraindividual variability at multiple time scales. *Psychological Methods, 14*, 367-386.

Examples

```
# A time series with 8 time points
tseries <- 49:56
embed.tseries <- Embed(tseries, E = 4, tau = 1)
```

entropyFit

Entropy Fit Index

Description

Computes the fit of a dimensionality structure using empirical entropy. Lower values suggest better fit of a structure to the data.

Usage

```
entropyFit(data, structure)
```

Arguments

data	Matrix or data frame. Contains variables to be used in the analysis
structure	A vector representing the structure (numbers or labels for each item). Can be theoretical factors or the structure detected by EGA

Value

Returns a list containing:

Total.Correlation	The total correlation of the dataset
Total.Correlation.MM	Miller-Madow correction for the total correlation of the dataset
Entropy.Fit	The Entropy Fit Index
Entropy.Fit.MM	Miller-Madow correction for the Entropy Fit Index
Average.Entropy	The average entropy of the dataset

Author(s)

Hudson F. Golino <hfg9s@virginia.edu>, Alexander P. Christensen <alexpaulchristensen@gmail.com> and Robert Moulder <rgm4fd@virginia.edu>

References

Golino, H., Moulder, R. G., Shi, D., Christensen, A. P., Garrido, L. E., Nieto, M. D., Nesselroade, J., Sadana, R., Thiyagarajan, J. A., & Boker, S. M. (2020). Entropy fit indices: New fit measures for assessing the structure and dimensionality of multiple latent variables. *Multivariate Behavioral Research*.

See Also

[EGA](#) to estimate the number of dimensions of an instrument using EGA and [CFA](#) to verify the fit of the structure suggested by EGA using confirmatory factor analysis.

Examples

```
# Load data
wmt <- wmt2[,7:24]

# Estimate EGA model
ega.wmt <- EGA(data = wmt, model = "glasso", plot.EGA = FALSE)

# Compute entropy indices
entropyFit(data = wmt, structure = ega.wmt$wc)
```

ergoInfo	<i>Ergodicity Information Index</i>
----------	-------------------------------------

Description

Computes the Ergodicity Information Index

Usage

```
ergoInfo(data, use = c("edge.list", "weights"))
```

Arguments

data	A dynEGA.ind.pop object
use	Character. A string indicating what network element will be used to compute the algorithm complexity, the list of edges or the weights of the network. Defaults to use = "edge.list". Current options are: <ul style="list-style-type: none">• <code>edge.list</code> Calculates the algorithm complexity using the list of edges.• <code>weights</code> Calculates the algorithm complexity using the weights of the network.

Value

Returns a list containing:

PrimeWeight	The prime-weight encoding of the individual networks
PrimeWeight.pop	The prime-weight encoding of the population network
Kcomp	The Kolmogorov complexity of the prime-weight encoded individual networks
Kcomp.pop	The Kolmogorov complexity of the prime-weight encoded population network
EII	The Ergodicity Information Index

Author(s)

Hudson Golino <hfg9s at virginia.edu>

glla

*Generalized Local Linear Approximation***Description**

Estimates the derivatives of a time series using generalized local linear approximation (GLLA). GLLA is a filtering method for estimating derivatives from data that uses time delay embedding and a variant of Savitzky-Golay filtering to accomplish the task.

Usage

```
glla(x, n.embed, tau, delta, order)
```

Arguments

x	Vector. An observed time series.
n.embed	Integer. Number of embedded dimensions (the number of observations to be used in the Embed function).
tau	Integer. Number of observations to offset successive embeddings in the Embed function. A tau of one uses adjacent observations. Default is "tau = 1".
delta	Integer. The time between successive observations in the time series. Default is "delta = 1".
order	Integer. The maximum order of the derivative to be estimated. For example, "order = 2" will return a matrix with three columns with the estimates of the observed scores and the first and second derivative for each row of the embedded matrix (i.e. the reorganization of the time series implemented via the Embed function).

Value

Returns a matrix containing n columns, in which n is one plus the maximum order of the derivatives to be estimated via generalized local linear approximation.

Author(s)

Hudson Golino <hfg9s at virginia.edu>

References

Boker, S. M., Deboeck, P. R., Edler, C., & Keel, P. K. (2010) Generalized local linear approximation of derivatives from time series. In S.-M. Chow, E. Ferrer, & F. Hsieh (Eds.), *The Notre Dame series on quantitative methodology. Statistical methods for modeling human dynamics: An interdisciplinary dialogue*, (p. 161-178). *Routledge/Taylor & Francis Group*.

Deboeck, P. R., Montpetit, M. A., Bergeman, C. S., & Boker, S. M. (2009) Using derivative estimates to describe intraindividual variability at multiple time scales. *Psychological Methods*, *14*(4), 367-386.

Savitzky, A., & Golay, M. J. (1964). Smoothing and differentiation of data by simplified least squares procedures. *Analytical Chemistry*, 36(8), 1627-1639.

Examples

```
# A time series with 8 time points
tseries <- 49:56
deriv.tseries <- glla(tseries, n.embed = 4, tau = 1, delta = 1, order = 2)
```

hierEGA

Hierarchical [EGA](#)

Description

Estimates EGA using the lower-order solution of [cluster_louvain](#) to identify the lower-order dimensions and then uses factor or network loadings to estimate factor or network scores, which are used to estimate the higher-order dimensions

Usage

```
hierEGA(
  data,
  scores = c("factor", "network"),
  consensus.iter = 1000,
  consensus.method = c("highest_modularity", "most_common", "iterative", "lowest_tefi"),
  uni.method = c("expand", "LE"),
  corr = c("cor_auto", "pearson", "spearman"),
  model = c("glasso", "TMFG"),
  model.args = list(),
  algorithm = c("walktrap", "louvain"),
  algorithm.args = list(),
  plot.EGA = TRUE,
  plot.type = c("GGally", "qgraph"),
  plot.args = list(),
  verbose = TRUE
)
```

Arguments

data	Matrix or data frame. Variables (down columns) only. Does not accept correlation matrices
scores	Character. How should scores for the higher-order structure be estimated? Defaults to "network" for network scores computed using the net.scores function. Set to "factor" for factor scores computed using fa . Factors are assumed to be correlated using the "oblimin" rotation. <i>NOTE</i> : Factor scores

use the number of communities from [EGA](#). Estimated factor may not align with these communities. The plots using factor scores with have higher order factors that may not completely map onto the lower order communities. Look at the `$hierarchical$higher_order$lower_loadings` to determine the composition of the lower order factors.

By default, both factor and network scores are computed and stored in the output. The selected option only appears in the main output (`$hierarchical`)

<code>consensus.iter</code>	Numeric. Number of iterations to perform in consensus clustering (see Lancichinetti & Fortunato, 2012). Defaults to 1000
<code>consensus.method</code>	<p>Character. What consensus clustering method should be used? Defaults to "highest_modularity". Current options are:</p> <ul style="list-style-type: none"> • <code>highest_modularity</code> Uses the community solution that achieves the highest modularity across iterations • <code>most_common</code> Uses the community solution that is found the most across iterations • <code>iterative</code> Identifies the most common community solutions across iterations and determines how often nodes appear in the same community together. A threshold of 0.30 is used to set low proportions to zero. This process repeats iteratively until all nodes have a proportion of 1 in the community solution. • <code>lowest_tefi</code> Uses the community solution that achieves the lowest tefi across iterations <p>By default, all <code>consensus.method</code> options are computed and stored in the output. The selected method will be used to plot and appear in the main output (<code>\$hierarchical</code>)</p>
<code>uni.method</code>	<p>Character. What unidimensionality method should be used? Defaults to "LE". Current options are:</p> <ul style="list-style-type: none"> • <code>expand</code> Expands the correlation matrix with four variables correlated .50. If number of dimension returns 2 or less in check, then the data are unidimensional; otherwise, regular EGA with no matrix expansion is used. This is the method used in the Golino et al. (2020) <i>Psychological Methods</i> simulation. • <code>LE</code> Applies the leading eigenvalue algorithm (cluster_leading_eigen) on the empirical correlation matrix. If the number of dimensions is 1, then the leading eigenvalue solution is used; otherwise, regular EGA is used. This is the final method used in the Christensen, Garrido, and Golino (2021) simulation.
<code>corr</code>	<p>Type of correlation matrix to compute. The default uses <code>cor_auto</code>. Current options are:</p> <ul style="list-style-type: none"> • <code>cor_auto</code> Computes the correlation matrix using the <code>cor_auto</code> function from qgraph. • <code>pearson</code> Computes Pearson's correlation coefficient using the pairwise complete observations via the <code>cor</code> function.

	<ul style="list-style-type: none"> • <code>spearman</code> Computes Spearman's correlation coefficient using the pairwise complete observations via the <code>cor</code> function.
<code>model</code>	<p>Character. A string indicating the method to use. Defaults to "glasso". Current options are:</p> <ul style="list-style-type: none"> • <code>glasso</code> Estimates the Gaussian graphical model using graphical LASSO with extended Bayesian information criterion to select optimal regularization parameter • <code>TMFG</code> Estimates a Triangulated Maximally Filtered Graph
<code>model.args</code>	List. A list of additional arguments for <code>EBICglasso.qgraph</code> or <code>TMFG</code>
<code>algorithm</code>	<p>A string indicating the algorithm to use or a function from <code>igraph</code> Defaults to "louvain". Current options are:</p> <ul style="list-style-type: none"> • <code>walktrap</code> Computes the Walktrap algorithm using <code>cluster_walktrap</code> • <code>louvain</code> Computes the Louvain algorithm using <code>cluster_louvain</code>
<code>algorithm.args</code>	List. A list of additional arguments for <code>cluster_walktrap</code> , <code>cluster_louvain</code> , or some other community detection algorithm function (see examples)
<code>plot.EGA</code>	Boolean. If TRUE, returns a plot of the network and its estimated dimensions. Defaults to TRUE
<code>plot.type</code>	Character. Plot system to use. Current options are <code>qgraph</code> and <code>GGally</code> . Defaults to "GGally"
<code>plot.args</code>	<p>List. A list of additional arguments for the network plot. For <code>plot.type = "qgraph"</code>:</p> <ul style="list-style-type: none"> • <code>vsize</code> Size of the nodes. Defaults to 6. <p>For <code>plot.type = "GGally"</code> (see <code>ggnet2</code> for full list of arguments):</p> <ul style="list-style-type: none"> • <code>vsize</code> Size of the nodes. Defaults to 6. • <code>label.size</code> Size of the labels. Defaults to 5. • <code>alpha</code> The level of transparency of the nodes, which might be a single value or a vector of values. Defaults to 0.7. • <code>edge.alpha</code> The level of transparency of the edges, which might be a single value or a vector of values. Defaults to 0.4. • <code>legend.names</code> A vector with names for each dimension • <code>color.palette</code> The color palette for the nodes. For custom colors, enter HEX codes for each dimension in a vector. See <code>color_palette_EGA</code> for more details and examples
<code>verbose</code>	Boolean. Should network estimation parameters be printed? Defaults to TRUE. Set to FALSE for no print out

Value

Returns a list of lists containing:

Main Results

- hierarhical The main results list containing:
- lower_order Lower order EGA results for the selected methods
 - higher_order Higher order EGA results for the selected methods
- If plot.EGA = TRUE, then:
- lower_plot Plot of the lower order results
 - higher_plot Plot of the higher order results
 - hier_plot Plot of the lower and higher order results together, side-by-side

Secondary Results

- lower_ega A list containing the lower order EGA results. The \$wc does not contain valid results. Do not use its output.
- lower_wc A list containing consensus clustering results:
- highest_modularity Community memberships based on the highest modularity across the cluster_louvain applications
 - most_common Community memberships based on the most commonly found memberships across the cluster_louvain applications
 - iterative Community memberships based on consensus clustering described by Lancichinetti & Fortunato (2012)
 - lowest_tefi Community memberships based on the lowest tefi across the cluster_louvain applications
 - summary_table A data frame summarizing the unique community solutions across the iterations. Down the columns indicate: number of dimensions (N_Dimensions), proportion of times each community solution was identified (Proportion), modularity of each community solution (Modularity), total entropy fit index of each community solution (tefi), and the memberships for each item. Across the rows indicate each unique community solution
- factor_results A list containing higher order results based on factor scores. A list for each consensus.method is provided with their EGA results
- network_results A list containing higher order results based on network scores. A list for each consensus.method is provided with their EGA results

Author(s)

Marcos Jimenez <marcosjnezhquez@gmail.com>, Francisco J. Abad <fjose.abad@uam.es>, Eduardo Garcia-Garzon <egarcia@ucjc.edu>, Luis Eduardo Garrido <luisgarrido@pucmm.edu.do>, Alexander P. Christensen <alexpaulchristensen@gmail.com>, and Hudson Golino <hfg9s@virginia.edu>

References

Lancichinetti, A., & Fortunato, S. (2012). Consensus clustering in complex networks. *Scientific Reports*, 2(1), 1-7.

Examples

```
# Obtain example data
data <- optimism

# hierEGA example (no plots)
opt.hier<- hierEGA(
  data = optimism,
  algorithm = "louvain",
  plot.EGA = FALSE # no plots for CRAN check
)

## Not run:

# hierEGA example (plots)
opt.hier <- hierEGA(
  data = optimism,
  algorithm = "louvain"
)

# Save plots
ggplot2::ggsave(
  filename = "hierEGA_plot.png", # name of plot
  plot = opt.res$hierarchical$hier_plot, # plot to save
  height = 8, # figure height
  width = 10, # figure width
  dpi = 600 # dots per inch
)

## End(Not run)
```

intelligenceBattery *Intelligence Data*

Description

A response matrix (n = 1152) of the International Cognitive Ability Resource (ICAR) intelligence battery developed by Condon and Revelle (2016).

A response matrix (n = 1152) of the International Cognitive Ability Resource (ICAR) intelligence battery developed by Condon and Revelle (2016).

Usage

```
data(intelligenceBattery)
```

```
data(intelligenceBattery)
```

Format

A 1185x125 response matrix

A 1185x125 response matrix

Examples

```
data("intelligenceBattery")
```

```
data("intelligenceBattery")
```

invariance

Measurement Invariance of [EGA](#) Structure

Description

Estimates metric invariance of [EGA](#) or specified structure

Usage

```
invariance(
  data,
  groups,
  memberships = NULL,
  type = c("loadings"),
  iter = 500,
  ncores,
  ...
)
```

Arguments

<code>data</code>	Matrix or data frame. Variables to be used in the analysis
<code>groups</code>	Vector. Group membership corresponding to each case in data
<code>memberships</code>	Vector. Node membership for each community or factor. Defaults to NULL. When NULL, EGA is used to compute node memberships
<code>type</code>	Character. Type of measurement invariance to estimate. Only includes "loadings" at the moment
<code>iter</code>	Numeric. Number of iterations to perform for the permutation. Defaults to 500
<code>ncores</code>	Numeric. Number of cores to use in computing results. Defaults to <code>parallel::detectCores() / 2</code> or half of your computer's processing power. Set to 1 to not use parallel computing If you're unsure how many cores your computer has, then use the following code: <code>parallel::detectCores()</code>
<code>...</code>	Arguments passed to EGA

Value

Returns a list containing:

memberships	Original memberships provided in memberships or from EGA if NULL
EGA	Original EGA results for the sample
groups	<ul style="list-style-type: none"> • EGA EGA results for each group • loadings Network loadings for each group • loadingsDifference Difference between the dominant loadings of each group
permutation	<ul style="list-style-type: none"> • groups Permuted groups across iterations • loadings Loadings for each group for each permutation • loadingsDifference Difference between the dominant loadings of each group for each permutation
results	Data frame of the results (which are printed)

Author(s)

Laura Jamison <lj5yn@virginia.edu>, Alexander P. Christensen <alexpaulchristensen@gmail.com>, and Hudson F. Golino <hfg9s@virginia.edu>

Examples

```
# Load data
wmt <- wmt2[-1,7:24]

# Groups
groups <- rep(1:2, each = nrow(wmt) / 2)

# Measurement invariance
results <- invariance(wmt, groups, ncores = 2)
```

itemStability

Item Stability Statistics from [bootEGA](#)

Description

Based on the [bootEGA](#) results, this function computes and plots the number of times an item (variable) is estimated in the same factor/dimension as originally estimated by [EGA](#) (`item.replication`). The output also contains each item's replication frequency (i.e., proportion of bootstraps that an item appeared in each dimension; `item.dim.rep`) as well as the average network loading for each item in each dimension (`item.loadings`).

Usage

```
itemStability(bootega.obj, IS.plot = TRUE, structure = NULL, ...)
```

Arguments

bootega.obj	A bootEGA object
IS.plot	Should the plot be produced for <code>item.replication</code> ? If TRUE, then a plot for the <code>item.replication</code> output will be produced. Defaults to TRUE
structure	User specified dimensionality structure.
...	Additional arguments. Used for deprecated arguments from previous versions of itemStability

Value

Returns a list containing:

membership	A list containing: <ul style="list-style-type: none"> • <code>empirical</code> The empirical memberships from the empirical EGA result • <code>unique</code> The unique dimensions from the empirical EGA result • <code>bootstrap</code> The memberships from the replicate samples in the bootEGA results
item.stability	A list containing: <ul style="list-style-type: none"> • <code>empirical.dimensions</code> The proportion of times each item replicated within the empirical EGA defined dimension. This EGA result is defined using the input from bootEGA • <code>all.dimensions</code> The proportion of times each item replicated in each of the empirical EGA defined dimensions. This EGA result is defined using the input from bootEGA
plot	A plot of the number of times each item replicated within the empirical EGA defined dimension.
mean.loadings	Matrix of the average standardized network loading (computed using net.loads) for each item in each dimension

Author(s)

Hudson Golino <hfg9s at virginia.edu> and Alexander P. Christensen <alexpaulchristensen@gmail.com>

References

- Christensen, A. P., & Golino, H. (2021). Estimating the stability of the number of factors via Bootstrap Exploratory Graph Analysis: A tutorial. *Psych*, 3(3), 479-500.
- Christensen, A. P., Golino, H., & Silvia, P. J. (2020). A psychometric network perspective on the validity and validation of personality trait questionnaires. *European Journal of Personality*, 34(6), 1095-1108.

See Also

[EGA](#) to estimate the number of dimensions of an instrument using EGA and [CFA](#) to verify the fit of the structure suggested by EGA using confirmatory factor analysis.

Examples

```
# Load data
wmt <- wmt2[,7:24]

## Not run: # Estimate EGA network
## plot.type = "qgraph" used for CRAN checks
## plot.type = "GGally" is the default
ega.wmt <- EGA(data = wmt, model = "glasso", plot.type = "qgraph")

# Estimate dimension stability
boot.wmt <- bootEGA(data = wmt, iter = 100, typicalStructure = TRUE,
plot.typicalStructure = TRUE, model = "glasso", plot.type = "qgraph",
type = "parametric", ncores = 2)

## End(Not run)

# Estimate item stability statistics
res <- itemStability(boot.wmt)

# Changing plot features (ggplot2)
## Changing colors (ignore warnings)
### qgraph Defaults
res$plot +
  ggplot2::scale_color_manual(values = rainbow(max(res$membership$unique)))

### Pastel
res$plot +
  ggplot2::scale_color_brewer(palette = "Pastel1")

## Changing Legend (ignore warnings)
res$plot +
  ggplot2::scale_color_discrete(labels = "Intelligence")
```

Description

An algorithm to identify whether data were generated from a factor or network model using factor and network loadings. The algorithm uses heuristics based on theory and simulation. These heuristics were then submitted to several deep learning neural networks with 240,000 samples per model with varying parameters.

Usage

```
LCT(
  data,
  n,
  iter = 100,
  dynamic = FALSE,
  dynamic.args = list(n.embed = 4, tau = 1, delta = 1, use.derivatives = 1)
)
```

Arguments

<code>data</code>	Matrix or data frame. A data frame with the variables to be used in the test or a correlation matrix. If the data used is a correlation matrix, the argument <code>n</code> will need to be specified
<code>n</code>	Integer. Sample size (if the data provided is a correlation matrix)
<code>iter</code>	Integer. Number of replicate samples to be drawn from a multivariate normal distribution (uses <code>mvtnorm::mvrnorm</code>). Defaults to 100
<code>dynamic</code>	Boolean. Is the dataset a time series where rows are time points and columns are variables? Defaults to <code>FASLE</code> .
<code>dynamic.args</code>	List. Arguments to be used in <code>dynEGA</code> . Defaults: <ul style="list-style-type: none"> • <code>n.embed</code> Number of embeddings: 4 • <code>tau</code> Lag: 1 • <code>delta</code> Delta: 1 • <code>use.derivatives</code> Derivatives: 1

Value

Returns a list containing:

<code>empirical</code>	Prediction of model based on empirical dataset only
<code>bootstrap</code>	Prediction of model based on means of the loadings across the bootstrap replicate samples
<code>proportion</code>	Proportions of models suggested across bootstraps

Author(s)

Hudson F. Golino <hfg9s at virginia.edu> and Alexander P. Christensen <alexpaulchristensen at gmail.com>

References

Christensen, A. P., & Golino, H. (2021). Factor or network model? Predictions from neural networks. *Journal of Behavioral Data Science*, 1(1), 85-126.

Examples

```
# Compute LCT
## Network model
LCT(data = wmt2[,7:24])

## Factor model
LCT(data = psychTools::bfi[,1:25])

# Dynamic LCT
LCT(sim.dynEGA[sim.dynEGA$ID == 1,1:20], dynamic = TRUE)
```

louvain

Louvain Community Detection Algorithm

Description

Computes the Louvain community detection algorithm (Blondel et al., 2008)

Usage

```
louvain(A, method = c("modularity", "tefi"), resolution = 1, corr = NULL)
```

Arguments

A	Matrix or data frame. A network adjacency matrix
method	Character. Whether modularity or <code>tefi</code> should be used to optimize communities. Defaults to "modularity"
resolution	Numeric. Resolution parameter for computing modularity. Defaults to 1. Values smaller than 1 favor larger communities; values larger than 1 favor smaller communities
corr	Matrix or data frame. Correlation matrix to be used when method = "tefi"

Details

This version was adapted from the Matlab code available here: <https://perso.uclouvain.be/vincent.blondel/research/louvain.htm>. The code was adjusted to mirror the results of `cluster_louvain`. The Louvain algorithm's results can vary depending on node ordering. In this version, nodes are **not** shuffled so that consistent results can be achieved with the same node ordering. Results from `cluster_louvain` will shuffle nodes **within** the function and therefore will sometimes produce similar results and sometimes produce slightly different results. This version is based all in R and therefore is slower than the version in `igraph`.

Value

Returns a list containing:

wc	A matrix of lower to higher order community membership detected in the network
modularity	A vector of modularity values corresponding the rows of the wc matrix

Author(s)

Alexander P. Christensen <alexpaulchristensen@gmail.com> and Hudson Golino <hfg9s@virginia.edu>

References

Blondel, V. D., Guillaume, J.-L., Lambiotte, R., & Lefebvre, E. (2008). Fast unfolding of communities in large networks. *Journal of Statistical Mechanics: Theory and Experiment*, 2008, P10008.

Examples

```
# Load data
dep <- depression[,24:44]

# Estimate correlations
corr <- qgraph::cor_auto(dep)

# Estimate network
net <- EBICglasso.qgraph(corr, n = nrow(dep))

# Estimate communities using modularity
louvain(net, method = "modularity")

# Estimate communities using tefi
louvain(net, method = "tefi", corr = corr)
```

Description

Computes a Monte-Carlo Test for the Ergodicity Information Index, comparing the empirical Ergodicity Information index to values obtained in a Monte-Carlo simulation in which all individuals have a similar latent structure. The p-values in the Monte-Carlo test can be calculated as $(\text{sum}(\text{EII} \geq \text{MC.EII}) + 1) / (\text{iter} + 1)$ and as $(\text{sum}(\text{EII} \leq \text{MC.EII}) + 1) / (\text{iter} + 1)$, where EII is the empirical Ergodicity Information Index, MC.EII is the values of the Ergodicity Information Index obtained in the simulation, and *iter* is the number of random samples generated in the simulation. The two-sided p-value is computed as two times the lowest p-value. In the Monte-Carlo Test for the Ergodicity Information Index, the null hypothesis is that the empirical value of EII is equal to the Monte-Carlo value of EII obtained in multiple individuals with a similar latent structure. Small

values of p indicate that is very unlikely to obtain an EII as large as the one obtained in the empirical sample if the null hypothesis is true, thus there is convincing evidence that the empirical Ergodicity Information Index is different than it could be expected if all individuals had a similar latent structure, conditioned on the parameters used to simulate the data.

Usage

```
mctest.ergoInfo(
  iter,
  N,
  EII,
  use,
  variab,
  timep,
  nfact,
  error,
  dfm,
  loadings,
  autoreg,
  crossreg,
  var.shock,
  cov.shock,
  embed,
  tau,
  delta,
  derivatives,
  model,
  model.args = list(),
  algorithm = c("walktrap", "louvain"),
  algorithm.args = list(),
  corr,
  ncores,
  ...
)
```

Arguments

<code>iter</code>	Numeric integer. Number of random samples to generate in the Monte-Carlo simulation. At least 500 is recommended
<code>N</code>	Numeric integer. Number of individuals to simulate data from, using the simDFM function.
<code>EII</code>	Numeric. Empirical Ergodicity Information Index obtained via the ergoInfo function.
<code>use</code>	Character. A string indicating what network element will be used to compute the algorithm complexity in the ergoInfo function, the list of edges or the weights of the network. Defaults to <code>use = "edge.list"</code> . Current options are: <ul style="list-style-type: none"> <code>edge.list</code> Calculates the algorithm complexity using the list of edges.

	<ul style="list-style-type: none"> • <code>weights</code> Calculates the algorithm complexity using the weights of the network.
<code>variab</code>	Number of variables per factor.
<code>timep</code>	Number of time points.
<code>nfact</code>	Number of factors.
<code>error</code>	Value to be used to construct a diagonal matrix Q . This matrix is $p \times p$ covariance matrix Q that will generate random errors following a multivariate normal distribution with mean zeros. The value provided is squared before constructing Q .
<code>dfm</code>	<p>A string indicating the dynamical factor model to use. Defaults to "DAFS". Current options are:</p> <ul style="list-style-type: none"> • <code>DAFS</code> Simulates data using the direct autoregressive factor score model. This is the default method • <code>RandomWalk</code> Simulates data using a dynamic factor model with random walk factor scores.
<code>loadings</code>	Magnitude of the loadings.
<code>autoreg</code>	Magnitude of the autoregression coefficients. Default is " <code>autoreg = 0.8</code> ".
<code>crossreg</code>	Magnitude of the cross-regression coefficients. Default is " <code>crossreg = 0.1</code> ".
<code>var.shock</code>	Magnitude of the random shock variance. Default is " <code>var.shock = 0.18</code> ".
<code>cov.shock</code>	Magnitude of the random shock covariance Default is " <code>cov.shock = 0.36</code> ".
<code>embed</code>	Integer. Number of embedded dimensions (the number of observations to be used in the <code>Embed</code> function). For example, an " <code>embed = 5</code> " will use five observations to estimate a single derivative. Defaults to <code>embed = 5</code> .
<code>tau</code>	Integer. Number of observations to offset successive embeddings in the <code>Embed</code> function. A tau of one uses adjacent observations. Default is " <code>tau = 1</code> ".
<code>delta</code>	Integer. The time between successive observations in the time series. Default is " <code>delta = 1</code> ".
<code>derivatives</code>	Integer. The order of the derivative to be used in the EGA procedure. Default to 1.
<code>model</code>	<p>Character. A string indicating the method to use. Defaults to <code>glasso</code>. Current options are:</p> <ul style="list-style-type: none"> • <code>glasso</code> Estimates the Gaussian graphical model using graphical LASSO with extended Bayesian information criterion to select optimal regularization parameter. This is the default method • <code>TMFG</code> Estimates a Triangulated Maximally Filtered Graph
<code>model.args</code>	List. A list of additional arguments for <code>EBICglasso.qgraph</code> or <code>TMFG</code>
<code>algorithm</code>	<p>A string indicating the algorithm to use or a function from <code>igraph</code></p> <p>Current options are:</p> <ul style="list-style-type: none"> • <code>walktrap</code> Computes the Walktrap algorithm using <code>cluster_walktrap</code> • <code>louvain</code> Computes the Walktrap algorithm using <code>cluster_louvain</code>

algorithm.args	List. A list of additional arguments for <code>cluster_walktrap</code> , <code>cluster_louvain</code> , or some other community detection algorithm function (see examples)
corr	Type of correlation matrix to compute. The default uses <code>cor_auto</code> . Current options are: <ul style="list-style-type: none"> • <code>cor_auto</code> Computes the correlation matrix using the <code>cor_auto</code> function from <code>qgraph</code>. • <code>pearson</code> Computes Pearson's correlation coefficient using the pairwise complete observations via the <code>cor</code> function. • <code>spearman</code> Computes Spearman's correlation coefficient using the pairwise complete observations via the <code>cor</code> function.
ncores	Numeric. Number of cores to use in computing results. Defaults to <code>parallel::detectCores() / 2</code> or half of your computer's processing power. Set to 1 to not use parallel computing. Recommended to use maximum number of cores minus one If you're unsure how many cores your computer has, then use the following code: <code>parallel::detectCores()</code>
...	Additional arguments. Used for deprecated arguments from previous versions of EGA

Value

Returns a list containing:

mc.ergoInfo	The values of the Ergodicity Information Index obtained in the Monte-Carlo Simulation
p.value.twosided	The p-value of the Monte-Carlo test for the Ergodicity Information Index. The null hypothesis is that the empirical Ergodicity Information index is equal to the expected value of the EII if the all individuals had similar latent structures.
effect	Indicates wheter the empirical EII is greater or less then the Monte-Carlo obtained EII.
plot.dist	Histogram of the bootstrapped ergodicity information index

Author(s)

Hudson Golino <hfg9s at virginia.edu>

Examples

```
## Not run:
\donttest{
dyn1 <- dynEGA.ind.pop(data = sim.dynEGA, n.embed = 5, tau = 1,
                      delta = 1, id = 21, group = 22, use.derivatives = 1,
                      model = "glasso", ncores = 2, corr = "pearson")

eii1 <- ergoInfo(data = dyn1)$EII

dist.ergoinfo <- mctest.ergoInfo(iter = 10, N = 10, EII = eii1,
```

```

variab = 4,
timep = 100, nfact = 2, error = 0.05, dfm = "DAFS", loadings = 0.55, autoreg = 0.8,
crossreg = 0.1, var.shock = 0.18, cov.shock = 0.36, embed = 5, tau=1, delta=1, derivatives=1,
model = "glasso", ncores = 2, corr = "pearson")
}
## End(Not run)

```

methods.section *Automated Methods Section for EGA_{net} Objects*

Description

This function accepts [EGA](#) objects and generates a Methods section for your analysis. The output is an HTML page containing the descriptions of the methods and parameters as well as a Reference section for appropriate citation.

Usage

```

methods.section(
  ...,
  stats = c("net.loads", "net.scores", "dimensionStability", "itemStability")
)

```

Arguments

... [EGA_{net}](#) objects. Available methods (more methods will be added soon!):

- [EGA](#) Exploratory graph analysis
- [bootEGA](#) Bootstrap exploratory graph analysis
- [UVA](#) Unique variable analysis

stats Methods section for statistics in [EGA_{net}](#). Multiple statistics can be input. Available statistics:

- [net.loads](#) Network loadings. Requires [EGA](#) object to be input
- [net.scores](#) Network scores. Requires [EGA](#) object to be input
- [dimensionStability](#) Structural consistency. Requires [bootEGA](#) object to be input
- [itemStability](#) Item stability. Requires [bootEGA](#) object to be input

Value

Automated HTML Methods section in your default browser

Examples

```

# Estimate EGA
## plot.type = "qgraph" used for CRAN checks
## plot.type = "GGally" is the default
ega.wmt <- EGA(data = wmt2[,7:24], plot.type = "qgraph")

# EGA Methods section
if(interactive()){
  methods.section(ega.wmt)
}

# Estimate standardized network loadings
wmt.loads <- net.loads(ega.wmt)$std

# EGA Methods section with network loadings
if(interactive()){
  methods.section(ega.wmt, stats = "net.loads")
}

## Not run: # bootEGA example
## plot.type = "qgraph" used for CRAN checks
## plot.type = "GGally" is the default
boot.wmt <- bootEGA(data = wmt2[,7:24], iter = 500, plot.type = "qgraph",
  type = "parametric", ncores = 2)

## End(Not run)
# EGA and bootEGA Methods section
if(interactive()){
  methods.section(ega.wmt, boot.wmt)
}

# Estimate structural consistency
sc.wmt <- dimensionStability(boot.wmt)

# EGA and bootEGA Methods section with structural consistency and item stability
if(interactive()){
  methods.section(boot.wmt, stats = c("dimensionStability", "itemStability"))
}

# EGA with network loadings and
# bootEGA Methods section with structural consistency and item stability
if(interactive()){
  methods.section(ega.wmt, boot.wmt, stats = c("net.loads", "dimensionStability", "itemStability"))
}

```

Description

Computes the between- and within-community strength of each item for each community. This function uses the `comcat` and `stable` functions to calculate the between- and within-community strength of each item, respectively.

Usage

```
net.loads(A, wc, pos.manifold = FALSE, min.load = 0, plot.NL = FALSE)
```

Arguments

<code>A</code>	Matrix, data frame, or EGA object. A network adjacency matrix
<code>wc</code>	Numeric or character vector. A vector of community assignments. If input into <code>A</code> is an EGA object, then <code>wc</code> is automatically detected
<code>pos.manifold</code>	Boolean. Should a positive manifold be applied (i.e., should all dimensions be positively correlated)? Defaults to <code>FALSE</code> . Set to <code>TRUE</code> for a positive manifold
<code>min.load</code>	Numeric. Sets the minimum loading allowed in the standardized network loading matrix. Values equal or greater than the minimum loading are kept in the output. Values less than the minimum loading are removed. This matrix can be viewed using <code>print()</code> or <code>summary()</code> Defaults to <code>0</code>
<code>plot.NL</code>	Boolean. Should proportional loadings be plotted? Defaults to <code>FALSE</code> . Set to <code>TRUE</code> for plot with pie charts visualizing the proportion of loading associated with each dimension

Details

Simulation studies have demonstrated that a node's strength centrality is roughly equivalent to factor loadings (Christensen, Golino, & Silvia, 2019; Hallquist, Wright, & Molenaar, in press). Hallquist and colleagues (in press) found that node strength represented a combination of dominant and cross-factor loadings. This function computes each node's strength within each specified dimension, providing a rough equivalent to factor loadings (including cross-loadings).

For more details, type `vignette("Network_Scores")`

Value

Returns a list containing:

<code>unstd</code>	A matrix of the unstandardized within- and between-community strength values for each node
<code>std</code>	A matrix of the standardized within- and between-community strength values for each node
<code>minLoad</code>	The minimum loading to appear in summary of network loadings. Use <code>print()</code> or <code>summary()</code> to view
<code>plot</code>	A qgraph plot of the network loadings. Use <code>plot</code> to view

Author(s)

Alexander P. Christensen <alexpaulchristensen@gmail.com> and Hudson Golino <hfg9s at virginia.edu>

References

Christensen, A. P., & Golino, H. (2021). On the equivalency of factor and network loadings. *Behavior Research Methods*, 53, 1563-1580.

Christensen, A. P., Golino, H., & Silvia, P. J. (2020). A psychometric network perspective on the validity and validation of personality trait questionnaires. *European Journal of Personality*, 34, 1095-1108.

Hallquist, M., Wright, A. C. G., & Molenaar, P. C. M. (2019). Problems with centrality measures in psychopathology symptom networks: Why network psychometrics cannot escape psychometric theory. *Multivariate Behavioral Research*, 1-25.

Examples

```
# Load data
wmt <- wmt2[,7:24]

## Not run:
# Estimate EGA
ega.wmt <- EGA(wmt)

## End(Not run)

# Network loadings
net.loads(ega.wmt)
```

net.scores

Network Scores

Description

This function computes network scores computed based on each node's strength within each community (i.e., factor) in the network (see [net.loads](#)). These values are used as network "factor loadings" for the weights of each item. Notably, network analysis allows nodes to contribute to more than one community. These loadings are considered in the network scores. In addition, if the construct is a hierarchy (e.g., personality questionnaire; items in facet scales in a trait domain), then an overall score can be computed (see argument `global`). An important difference is that the network scores account for cross-loadings in their estimation of scores

Usage

```
net.scores(data, A, wc, global = FALSE, impute, ...)
```

Arguments

data	Matrix or data frame. Must be a dataset
A	Matrix, data frame, or EGA object. An adjacency matrix of network data
wc	Numeric. A vector of community assignments. Not necessary if an EGA object is input for argument A
global	Boolean. Should general network loadings be computed in scores? Defaults to FALSE. If there is more than one dimension and there is theoretically one global dimension, then general loadings of the dimensions onto the global dimension can be included in the weighted scores
impute	Character. In the presence of missing data, imputation can be implemented. Currently, three options are available: <ul style="list-style-type: none"> • none No imputation is performed. This is the default. • mean The "mean" value of the columns are used to replace the missing data. • median The "median" value of the columns are used to replace the missing data.
...	Additional arguments for EGA

Details

For more details, type `vignette("Network_Scores")`

Value

Returns a list containing:

unstd.scores	The unstandardized network scores for each participant and community (including the overall score)
std.scores	The standardized network scores for each participant and community (including the overall score)
commCor	Partial correlations between the specified or identified communities
loads	Standardized network loadings for each item in each dimension (computed using net.loads)

Author(s)

Alexander P. Christensen <alexpaulchristensen@gmail.com> and Hudson F. Golino <hfg9s@virginia.edu>

References

- Christensen, A. P., & Golino, H. (2021). On the equivalency of factor and network loadings. *Behavior Research Methods*, *53*, 1563-1580.
- Christensen, A. P., Golino, H., & Silvia, P. J. (2020). A psychometric network perspective on the validity and validation of personality trait questionnaires. *European Journal of Personality*, *34*, 1095-1108.

Golino, H., Christensen, A. P., Moulder, R., Kim, S., & Boker, S. M. (2021). Modeling latent topics in social media using Dynamic Exploratory Graph Analysis: The case of the right-wing and left-wing trolls in the 2016 US elections. *Psychometrika*.

Examples

```
# Load data
wmt <- wmt2[,7:24]

## Not run:
# Estimate EGA
ega.wmt <- EGA(wmt)

## End(Not run)

# Network scores
net.scores(data = wmt, A = ega.wmt)
```

network.descriptives *Descriptive Statistics for Networks*

Description

Computes descriptive statistics for network models

Usage

```
network.descriptives(network)
```

Arguments

network Matrix, data frame, [qgraph](#), or [EGA](#) object

Value

Numeric vector including:

Mean_weight	The average of the edge weights in the network
SD_weight	The standard deviation of the edge weights in the network
Min_weight	The minimum of the edge weights in the network
Max_weight	The maximum of the edge weights in the network
Density	The density of the network
ASPL	The average shortest path length (ASPL) of the network (computed as unweighted)
CC	The clustering coefficient (CC) of the network (computed as unweighted)

swn.rand	Small-worldness measure based on random networks: $swn.rand = (ASPL/ASPL_{r,andom})/(CC/CC_{r,andom})$ swn.rand > 1 suggests the network is small-world
swn.HG	Small-worldness measure based on Humphries & Gurney (2008): $swn.HG = (transitivity/transitivity_{r,andom})/(ASPL/ASPL_{r,andom})$ swn.HG > 1 suggests the network is small-world
swn.TJHBL	Small-worldness measure based on Telesford, Joyce, Hayasaka, Burdette, & Laurienti (2011): $swn.TJHBL = (ASPL_{r,andom}/ASPL) - (CC/CC_{lattice})$ swn.TJHBL near 0 suggests the network is small-world, positive values suggest more random network characteristics, negative values suggest more lattice network characteristics
scale-free_R-sq	The R-squared fit of whether the degree distribution follows the power-law (many small degrees, few large degrees)

Author(s)

Hudson Golino <hfg9s at virginia.edu> and Alexander P. Christensen <alexpaulchristensen@gmail.com>

References

```
# swn.HG
Humphries, M. D., & Gurney, K. (2008). Network 'small-world-ness': A quantitative method for determining canonical network equivalence. PLoS one, 3, e0002051

# swn.TJHBL
Telesford, Q. K., Joyce, K. E., Hayasaka, S., Burdette, J. H., & Laurienti, P. J. (2011). The ubiquity of small-world networks. Brain Connectivity, 1(5), 367-375

# scale-free_R-sq
Langfelder, P., & Horvath, S. (2008). WGCNA: an R package for weighted correlation network analysis. BMC Bioinformatics, 9, 559
```

Examples

```
# Load data
wmt <- wmt2[,7:24]

# EGA example
## plot.type = "qgraph" used for CRAN checks
## plot.type = "GGally" is the default
ega.wmt <- EGA(data = wmt, plot.type = "qgraph")

# Compute descriptives
network.descriptives(ega.wmt)
```

optimism

Optimism Data

Description

A response matrix (n = 282) containing responses to 10 items of the Revised Life Orientation Test (LOT-R), developed by Scheier, Carver, & Bridges (1994).

A response matrix (n = 282) containing responses to 10 items of the Revised Life Orientation Test (LOT-R), developed by Scheier, Carver, & Bridges (1994).

Usage

```
data(optimism)
```

```
data(optimism)
```

Format

A 282x10 response matrix

A 282x10 response matrix

References

Scheier, M. F., Carver, C. S., & Bridges, M. W. (1994). Distinguishing optimism from neuroticism (and trait anxiety, self-mastery, and self-esteem): a reevaluation of the Life Orientation Test. *Journal of Personality and Social Psychology*, *67*, 1063-1078.

Scheier, M. F., Carver, C. S., & Bridges, M. W. (1994). Distinguishing optimism from neuroticism (and trait anxiety, self-mastery, and self-esteem): a reevaluation of the Life Orientation Test. *Journal of Personality and Social Psychology*, *67*, 1063-1078.

Examples

```
data("optimism")
```

```
data("optimism")
```

Description

Plots for EGAnet objects

Usage

```
## S3 method for class 'bootEGA'
plot(x, plot.type = c("GGally", "qgraph"),
     plot.args = list(), produce = TRUE, ...)

## S3 method for class 'CFA'
plot(x, layout = "spring", vsize = 6, ...)

## S3 method for class 'dynEGA'
plot(x, title = "", plot.type = c("GGally", "qgraph"),
     plot.args = list(), produce = TRUE, ...)

## S3 method for class 'dynEGA.Groups'
plot(x, ncol, nrow, title = "",
     plot.type = c("GGally", "qgraph"), plot.args = list(), produce = TRUE, ...)

## S3 method for class 'dynEGA.Individuals'
plot(x, title = "", id = NULL,
     plot.type = c("GGally", "qgraph"), plot.args = list(), produce = TRUE, ...)

## S3 method for class 'EGA'
plot(x, title = "", plot.type = c("GGally", "qgraph"),
     plot.args = list(), produce = TRUE, ...)

## S3 method for class 'NetLoads'
plot(x, ...)
```

Arguments

<code>x</code>	Object from EGAnet package
<code>plot.type</code>	Character. Plot system to use. Current options are qgraph and GGally . Defaults to "GGally".
<code>plot.args</code>	List. A list of additional arguments for the network plot. For <code>plot.type = "qgraph"</code> : <ul style="list-style-type: none"> • <code>vsize</code> Size of the nodes. Defaults to 6. For <code>plot.type = "GGally"</code> (see ggnet2 for full list of arguments): <ul style="list-style-type: none"> • <code>vsize</code> Size of the nodes. Defaults to 6.

- `label.size` Size of the labels. Defaults to 5.
- `alpha` The level of transparency of the nodes, which might be a single value or a vector of values. Defaults to 0.7.
- `edge.alpha` The level of transparency of the edges, which might be a single value or a vector of values. Defaults to 0.4.
- `legend.names` A vector with names for each dimension
- `color.palette` The color palette for the nodes. For custom colors, enter HEX codes for each dimension in a vector. See [color_palette_EGA](#) for more details and examples

<code>produce</code>	Boolean. This argument is used internally. Should plot be produced? Defaults to TRUE
<code>...</code>	Arguments passed on to <ul style="list-style-type: none"> • qgraph Functions: <code>bootEGA</code>, <code>dynEGA</code>, <code>dynEGA.Groups</code>, <code>dynEGA.Individuals</code>, <code>EGA</code>, and <code>net.loads</code> • semPaths Functions: <code>CFA</code>
<code>vsiz</code>	Numeric. Size of vertices in CFA plots. Defaults to 6
<code>layout</code>	Character. Layout of plot (see semPaths). Defaults to "spring"
<code>ncol</code>	Numeric. Number of columns
<code>nrow</code>	Numeric. Number of rows
<code>title</code>	Character. Title of the plot. Defaults to ""
<code>id</code>	Numeric. An integer or character indicating the ID of the individual to plot

Value

Plots of EGAnet object

Author(s)

Hudson Golino <hfg9s at virginia.edu> and Alexander P. Christensen <alexpaulchristensen@gmail.com>

<code>prime.num</code>	<i>Prime Numbers through 100,000</i>
------------------------	--------------------------------------

Description

Numeric vector of primes generated from the primes package. Used in the function `[EGAnet]{ergoInfo}`.
Not for general use

Numeric vector of primes generated from the primes package. Used in the function `[EGAnet]{ergoInfo}`.
Not for general use

Usage

```
data(prime.num)
```

```
data(prime.num)
```

Format

A 1185x24 response matrix

A 1185x24 response matrix

Examples

```
data("prime.num")
```

```
data("prime.num")
```

prints	<i>S3Methods for Printing</i>
--------	-------------------------------

Description

Prints for EGAnet objects

Usage

```
## S3 method for class 'dynEGA'  
print(x, ...)  
  
## S3 method for class 'dynEGA.Groups'  
print(x, ...)  
  
## S3 method for class 'dynEGA.Individuals'  
print(x, ...)  
  
## S3 method for class 'EGA'  
print(x, ...)  
  
## S3 method for class 'NetLoads'  
print(x, ...)  
  
## S3 method for class 'invariance'  
print(x, ...)  
  
## S3 method for class 'hierEGA'  
print(x, ...)
```

Arguments

x	Object from EGAnet package
...	Additional arguments

Value

Prints EGAnet object

Author(s)

Hudson Golino <hfg9s at virginia.edu> and Alexander P. Christensen <alexpaulchristensen@gmail.com>

residualEGA

Residualized EGA

Description

residualEGA Estimates the number of dimensions after controlling for wording effects. EGA is applied in the residual of a random intercept item factor model (RIIFA) with one method factor and one substantive factor.

Usage

```
residualEGA(data, manifests, lat, negative.items, plot = TRUE)
```

Arguments

data	Matrix or data frame. Includes the variables to be used in the residualEGA analysis
manifests	Character vector. Vector indicating the names of the variables (items) to be used in the analysis.
lat	Numeric integer. Number of latent factors to be estimated. Only one substantive latent factor is recommended in the current version of the function.
negative.items	Numeric vector A numeric vector indicating the column of the negative items.
plot	Boolean. If TRUE, returns a plot of the residualized network and its estimated dimensions. Defaults to TRUE

Value

Returns a list containing:

openMx.model	OpenMX model
openMx.result	OpenMX results
openMx.std.par	OpenMX standardized parameters
ResidualMatrix	Residual matrix
EGA.Residuals	Results of the residualized EGA
Fit	Fit metrics of the network structure, calculated using the ggmlfit function of the qgraph package
WordLoads	Loadings of the wording effects

Author(s)

Hudson F. Golino <hfg9s at virginia.edu> and Robert Moulder <rgm4fd@virginia.edu>

See Also

[EGA](#) to estimate the number of dimensions of an instrument using EGA and [CFA](#) to verify the fit of the structure suggested by EGA using confirmatory factor analysis.

Examples

```
data <- optimism

## Not run:
# resEGA example
opt.res <- residualEGA(data = data, manifests = colnames(optimism),
  lat = 1, negative.items = c(3,7,9), plot = TRUE)

# Fit:
opt.res$Fit

## End(Not run)
```

riEGA

Random-Intercept EGA

Description

Estimates the number of substantive dimensions after controlling for wording effects. EGA is applied to a residual correlation matrix after subtracting and random intercept factor with equal unstandardized loadings from all the regular and unrecoded reversed items in the database

Usage

```
riEGA(
  data,
  n = NULL,
  uni.method = c("expand", "LE"),
  corr = c("cor_auto", "pearson", "spearman"),
  model = c("glasso", "TMFG"),
  model.args = list(),
  algorithm = c("walktrap", "louvain"),
  algorithm.args = list(),
  plot.EGA = TRUE,
  plot.type = c("GGally", "qgraph"),
  plot.args = list(),
  estimator = c("auto", "WLSMV", "MLR"),
```

```

lavaan.args = list(),
verbose = TRUE
)

```

Arguments

data	Matrix or data frame. Variables (down columns) or correlation matrix. If the input is a correlation matrix, then argument n (number of cases) is required . Variables MUST be unrecoded – reversed items should remain reversed
n	Integer. Sample size if data provided is a correlation matrix
uni.method	Character. What unidimensionality method should be used? Defaults to "LE". Current options are: <ul style="list-style-type: none"> • expand Expands the correlation matrix with four variables correlated .50. If number of dimension returns 2 or less in check, then the data are unidimensional; otherwise, regular EGA with no matrix expansion is used. This is the method used in the Golino et al. (2020) <i>Psychological Methods</i> simulation. • LE Applies the leading eigenvalue algorithm (cluster_leading_eigen) on the empirical correlation matrix. If the number of dimensions is 1, then the leading eigenvalue solution is used; otherwise, regular EGA is used. This is the final method used in the Christensen, Garrido, and Golino (2021) simulation.
corr	Type of correlation matrix to compute. The default uses cor_auto . Current options are: <ul style="list-style-type: none"> • cor_auto Computes the correlation matrix using the cor_auto function from qgraph. • pearson Computes Pearson's correlation coefficient using the pairwise complete observations via the cor function. • spearman Computes Spearman's correlation coefficient using the pairwise complete observations via the cor function.
model	Character. A string indicating the method to use. Defaults to "glasso". Current options are: <ul style="list-style-type: none"> • glasso Estimates the Gaussian graphical model using graphical LASSO with extended Bayesian information criterion to select optimal regularization parameter • TMFG Estimates a Triangulated Maximally Filtered Graph
model.args	List. A list of additional arguments for EBICglasso , qgraph or TMFG
algorithm	A string indicating the algorithm to use or a function from igraph Defaults to "walktrap". Current options are: <ul style="list-style-type: none"> • walktrap Computes the Walktrap algorithm using cluster_walktrap • louvain Computes the Louvain algorithm using cluster_louvain
algorithm.args	List. A list of additional arguments for cluster_walktrap , cluster_louvain , or some other community detection algorithm function (see examples)

<code>plot.EGA</code>	Boolean. If TRUE, returns a plot of the network and its estimated dimensions. Defaults to TRUE
<code>plot.type</code>	Character. Plot system to use. Current options are qgraph and GGally . Defaults to "GGally"
<code>plot.args</code>	List. A list of additional arguments for the network plot. For <code>plot.type = "qgraph"</code> : <ul style="list-style-type: none"> • <code>vsize</code> Size of the nodes. Defaults to 6. For <code>plot.type = "GGally"</code> (see ggnet2 for full list of arguments): <ul style="list-style-type: none"> • <code>vsize</code> Size of the nodes. Defaults to 6. • <code>label.size</code> Size of the labels. Defaults to 5. • <code>alpha</code> The level of transparency of the nodes, which might be a single value or a vector of values. Defaults to 0.7. • <code>edge.alpha</code> The level of transparency of the edges, which might be a single value or a vector of values. Defaults to 0.4. • <code>legend.names</code> A vector with names for each dimension • <code>color.palette</code> The color palette for the nodes. For custom colors, enter HEX codes for each dimension in a vector. See color_palette_EGA for more details and examples
<code>estimator</code>	Character. Estimator to use for random-intercept model (see Estimators for more details). Defaults to "auto", which selects "MLR" for continuous data and "WLSMV" for mixed and categorical data. Data are considered continuous data if they have 6 or more categories (see Rhemtulla, Brosseau-Liard, & Savalei, 2012)
<code>lavaan.args</code>	List. If <code>reduce.method = "latent"</code> , then lavaan 's <code>cfa</code> function will be used to create latent variables to reduce variables. Arguments should be input as a list. Some example arguments (see lavOptions for full details)
<code>verbose</code>	Boolean. Should network estimation parameters be printed? Defaults to TRUE. Set to FALSE for no print out

Value

Returns a list containing:

EGA	Results from EGA
RI	A list containing information about the random-intercept model (if the model converged): <ul style="list-style-type: none"> • <code>fit</code> The fit object for the random-intercept model using cfa • <code>lavaan.args</code> The arguments used in cfa • <code>loadings</code> Standardized loadings from the random-intercept model • <code>correlation</code> Residual correlations after accounting for the random-intercept model

Author(s)

Alejandro Garcia-Pardina <alejandrop97@gmail.com>, Francisco J. Abad <fjose.abad@uam.es>, Alexander P. Christensen <alexpaulchristensen@gmail.com>, Hudson Golino <hfg9s@virginia.edu>, Luis Eduardo Garrido <luisgarrido@pucmm.edu.do>, and Robert Moulder <rgm4fd@virginia.edu>

References

Selection of CFA Estimator
Rhemtulla, M., Brosseau-Liard, P. E., & Savalei, V. (2012). When can categorical variables be treated as continuous? A comparison of robust continuous and categorical SEM estimation methods under suboptimal conditions. *Psychological Methods, 17*, 354-373.

Examples

```
# Obtain example data
data <- optimism

## Not run:
# riEGA example
opt.res <- riEGA(data = optimism)

## End(Not run)
```

sim.dynEGA

sim.dynEGA Data

Description

A simulated (multivariate time series) data with 20 variables, 200 individual observations, 50 time points per individual and 2 groups of individuals.

A simulated (multivariate time series) data with 20 variables, 200 individual observations, 50 time points per individual and 2 groups of individuals.

Usage

```
data(sim.dynEGA)
```

```
data(sim.dynEGA)
```

Format

A 10000x22 multivariate time series

A 10000x22 multivariate time series

Examples

```
data("sim.dynEGA")
```

```
data("sim.dynEGA")
```

simDFM

*Simulate data following a Dynamic Factor Model***Description**

Function to simulate data following a dynamic factor model (DFM). Two DFMs are currently available: the direct autoregressive factor score model (Engle & Watson, 1981; Nesselroade, McArdle, Aggen, and Meyers, 2002) and the dynamic factor model with random walk factor scores.

Usage

```
simDFM(
  variab,
  timep,
  nfact,
  error,
  dfm = c("DAFS", "RandomWalk"),
  loadings,
  autoreg,
  crossreg,
  var.shock,
  cov.shock,
  burnin = 1000
)
```

Arguments

variab	Number of variables per factor.
timep	Number of time points.
nfact	Number of factors.
error	Value to be used to construct a diagonal matrix Q. This matrix is p x p covariance matrix Q that will generate random errors following a multivariate normal distribution with mean zeros. The value provided is squared before constructing Q.
dfm	A string indicating the dynamical factor model to use. Current options are: <ul style="list-style-type: none"> • DAFS Simulates data using the direct autoregressive factor score model. This is the default method • RandomWalk Simulates data using a dynamic factor model with random walk factor scores.
loadings	Magnitude of the loadings.
autoreg	Magnitude of the autoregression coefficients.
crossreg	Magnitude of the cross-regression coefficients.
var.shock	Magnitude of the random shock variance.
cov.shock	Magnitude of the random shock covariance
burnin	Number of n first samples to discard when computing the factor scores. Defaults to 1000.

Author(s)

Hudson F. Golino <hfg9s at virginia.edu>

References

Engle, R., & Watson, M. (1981). A one-factor multivariate time series model of metropolitan wage rates. *Journal of the American Statistical Association*, 76(376), 774-781.

Nesselroade, J. R., McArdle, J. J., Aggen, S. H., & Meyers, J. M. (2002). Dynamic factor analysis models for representing process in multivariate time-series. In D. S. Moskowitz & S. L. Hershberger (Eds.), *Multivariate applications book series. Modeling intraindividual variability with repeated measures data: Methods and applications*, 235-265.

Examples

```
## Not run:
\donttest{
# Estimate EGA network
data1 <- simDFM(variab = 5, timep = 50, nfact = 3, error = 0.05,
dfm = "DAFS", loadings = 0.7, autoreg = 0.8,
crossreg = 0.1, var.shock = 0.18,
cov.shock = 0.36, burnin = 1000)
}

## End(Not run)
```

summaries

S3Methods for Summaries

Description

Summaries for EGAnet objects

Usage

```
## S3 method for class 'CFA'
summary(object, ...)

## S3 method for class 'dynEGA'
summary(object, ...)

## S3 method for class 'dynEGA.Groups'
summary(object, ...)

## S3 method for class 'dynEGA.Individuals'
summary(object, ...)
```

```
## S3 method for class 'EGA'
summary(object, ...)

## S3 method for class 'NetLoads'
summary(object, ...)
```

Arguments

object Object from EGAnet package
 ... Additional arguments

Value

Summarizes EGAnet object

Author(s)

Hudson Golino <hfg9s at virginia.edu> and Alexander P. Christensen <alexpaulchristensen@gmail.com>

tefi *Total Entropy Fit Index using Von Neumman's entropy (Quantum Information Theory) for correlation matrices*

Description

Computes the fit (TEFI) of a dimensionality structure using Von Neumman's entropy when the input is a correlation matrix. Lower values suggest better fit of a structure to the data.

Usage

```
tefi(data, structure)
```

Arguments

data A dataframe or correlation matrix
 structure A vector representing the structure (numbers or labels for each item). Can be theoretical factors or the structure detected by [EGA](#)

Value

Returns a list containing:

VN.Entropy.Fit The Entropy Fit Index using Von Neumman's entropy
 Total.Correlation The total correlation of the dataset
 Average.Entropy The average entropy of the dataset

Author(s)

Hudson Golino <hfg9s at virginia.edu>, Alexander P. Christensen <alexpaulchristensen@gmail.com>, and Robert Moulder <rgm4fd@virginia.edu>

References

Golino, H., Moulder, R. G., Shi, D., Christensen, A. P., Garrido, L. E., Nieto, M. D., Nesselroade, J., Sadana, R., Thiyagarajan, J. A., & Boker, S. M. (2020). Entropy fit indices: New fit measures for assessing the structure and dimensionality of multiple latent variables. *Multivariate Behavioral Research*.

See Also

[EGA](#) to estimate the number of dimensions of an instrument using EGA and [CFA](#) to verify the fit of the structure suggested by EGA using confirmatory factor analysis.

Examples

```
# Load data
wmt <- wmt2[,7:24]

## Not run:
# Estimate EGA model
ega.wmt <- EGA(data = wmt, model = "glasso")

## End(Not run)

# Compute entropy indices
tefi(data = ega.wmt$correlation, structure = ega.wmt$wc)
```

 TMFG

Triangulated Maximally Filtered Graph

Description

Applies the Triangulated Maximally Filtered Graph (TMFG) filtering method (**please see and cite Massara et al., 2016**). The TMFG method uses a structural constraint that limits the number of zero-order correlations included in the network ($3n - 6$; where n is the number of variables). The TMFG algorithm begins by identifying four variables which have the largest sum of correlations to all other variables. Then, it iteratively adds each variable with the largest sum of three correlations to nodes already in the network until all variables have been added to the network. This structure can be associated with the inverse correlation matrix (i.e., precision matrix) to be turned into a GGM (i.e., partial correlation network) by using Local-Global Inversion Method (see Barfuss et al., 2016 for more details). See Details for more information on this network estimation method.

Usage

```
TMFG(cormat)
```

Arguments

```
cormat      A correlation matrix
```

Details

The TMFG method applies a structural constraint on the network, which restrains the network to retain a certain number of edges ($3n-6$, where n is the number of nodes; Massara et al., 2016). The network is also composed of 3- and 4-node cliques (i.e., sets of connected nodes; a triangle and tetrahedron, respectively). The TMFG method constructs a network using zero-order correlations and the resulting network can be associated with the inverse covariance matrix (yielding a GGM; Barfuss, Massara, Di Matteo, & Aste, 2016). Notably, the TMFG can use any association measure and thus does not assume the data is multivariate normal.

Construction begins by forming a tetrahedron of the four nodes that have the highest sum of correlations that are greater than the average correlation in the correlation matrix. Next, the algorithm iteratively identifies the node that maximizes its sum of correlations to a connected set of three nodes (triangles) already included in the network and then adds that node to the network. The process is completed once every node is connected in the network. In this process, the network automatically generates what's called a planar network. A planar network is a network that could be drawn on a sphere with no edges crossing (often, however, the networks are depicted with edges crossing; Tumminello, Aste, Di Matteo, & Mantegna, 2005).

Value

Returns a list containing:

```
A          The filtered adjacency matrix
separators The separators (3-cliques) in the network
cliques    The cliques (4-cliques) in the network
```

Author(s)

Alexander Christensen <alexpaulchristensen@gmail.com>

References

- Barfuss, W., Massara, G. P., Di Matteo, T., & Aste, T. (2016). Parsimonious modeling with information filtering networks. *Physical Review E*, *94*, 062306.
- Christensen, A. P., Kenett, Y. N., Aste, T., Silvia, P. J., & Kwapił, T. R. (2018). Network structure of the Wisconsin Schizotypy Scales-Short Forms: Examining psychometric network filtering approaches. *Behavior Research Methods*, *50*, 2531-2550.
- Massara, G. P., Di Matteo, T., & Aste, T. (2016). Network filtering for big data: Triangulated maximally filtered graph. *Journal of Complex Networks*, *5*, 161-178.

Examples

```
# Pearson's correlation only for CRAN checks
A <- TMFG(cor(wmt2[,7:24]))$A
```

totalCor	<i>Total Correlation</i>
----------	--------------------------

Description

Computes the total correlation of a dataset

Usage

```
totalCor(data)
```

Arguments

data Matrix or data frame. Variables to be used in the analysis

Value

Returns a list containing:

Ind.Entropies	Individual entropies for each variable
Joint.Entropy	The joint entropy of the dataset
Total.Cor	The total correlation of the dataset

Author(s)

Hudson F. Golino <hfg9s at virginia.edu>

References

Watanabe, S. (1960). Information theoretical analysis of multivariate correlation. *IBM Journal of Research and Development* 4, 66-82.

Implementation Felix, L. M., Mansur-Alves, M., Teles, M., Jamison, L., & Golino, H. (2021). Longitudinal impact and effects of booster sessions in a cognitive training program for healthy older adults. *Archives of Gerontology and Geriatrics*, 94, 104337.

Examples

```
# Compute total correlation
totalCor(wmt2[,7:24])
```

totalCorMat	<i>Total Correlation Matrix</i>
-------------	---------------------------------

Description

Computes the pairwise total correlation for a dataset

Usage

```
totalCorMat(data)
```

Arguments

data Matrix or data frame. Variables to be used in the analysis

Value

Returns a square matrix with pairwise total correlations

Author(s)

Hudson F. Golino <hfg9s at virginia.edu>

References

Watanabe, S. (1960). Information theoretical analysis of multivariate correlation. *IBM Journal of Research and Development* 4, 66-82.

Implementation Felix, L. M., Mansur-Alves, M., Teles, M., Jamison, L., & Golino, H. (2021). Longitudinal impact and effects of booster sessions in a cognitive training program for healthy older adults. *Archives of Gerontology and Geriatrics*, 94, 104337.

Examples

```
## Not run:  
\donttest{  
# Compute total correlation  
totalCorMat(wmt2[,7:24])  
}  
## End(Not run)
```

`toy.example`*Toy Example Data*

Description

A simulated dataset with 2 factors, three items per factor and $n = 500$.

Usage

```
data(toy.example)
```

Format

A 500x6 response matrix

Examples

```
data("toy.example")
```

`UVA`*Unique Variable Analysis*

Description

Identifies redundant variables in a multivariate dataset using a number of different association methods and types of significance values (see Christensen, Garrido, & Golino, 2020 for more details)

Usage

```
UVA(  
  data,  
  n = NULL,  
  model = c("glasso", "TMFG"),  
  corr = c("cor_auto", "pearson", "spearman"),  
  method = c("cor", "pcor", "wTO"),  
  type = c("adapt", "alpha", "threshold"),  
  sig,  
  key = NULL,  
  reduce = TRUE,  
  auto = TRUE,  
  label_latent = TRUE,  
  reduce.method = c("latent", "remove", "sum"),  
  lavaan.args = list(),  
  adhoc = TRUE,  
  plot.redundancy = FALSE,  
  plot.args = list()  
)
```

Arguments

<code>data</code>	Matrix or data frame. Input can either be data or a correlation matrix
<code>n</code>	Numeric. If input in <code>data</code> is a correlation matrix, then sample size is required. Defaults to NULL
<code>model</code>	Character. A string indicating the method to use. Current options are: <ul style="list-style-type: none"> • <code>glasso</code> Estimates the Gaussian graphical model using graphical LASSO with extended Bayesian information criterion to select optimal regularization parameter. This is the default method • <code>TMFG</code> Estimates a Triangulated Maximally Filtered Graph
<code>corr</code>	Type of correlation matrix to compute. The default uses <code>cor_auto</code> . Current options are: <ul style="list-style-type: none"> • <code>cor_auto</code> Computes the correlation matrix using the <code>cor_auto</code> function from <code>qgraph</code>. • <code>pearson</code> Computes Pearson's correlation coefficient using the pairwise complete observations via the <code>cor</code> function. • <code>spearman</code> Computes Spearman's correlation coefficient using the pairwise complete observations via the <code>cor</code> function.
<code>method</code>	Character. Computes weighted topological overlap (" <code>wT0</code> " using <code>EBICglasso</code>), partial correlations (" <code>pcor</code> "), or correlations (" <code>cor</code> ") Defaults to " <code>wT0</code> "
<code>type</code>	Character. Type of significance. Computes significance using the standard p -value (" <code>alpha</code> "), adaptive alpha p -value (<code>adapt.a</code>), or some threshold " <code>threshold</code> ". Defaults to " <code>threshold</code> "
<code>sig</code>	Numeric. p -value for significance of overlap (defaults to <code>.05</code>). Defaults for " <code>threshold</code> " for each method: <ul style="list-style-type: none"> • "<code>wT0</code>" <code>.25</code> • "<code>pcor</code>" <code>.35</code> • "<code>cor</code>" <code>.50</code>
<code>key</code>	Character vector. A vector with variable descriptions that correspond to the order of variables input into data. Defaults to NULL or the column names of data
<code>reduce</code>	Boolean. Should redundancy reduction be performed? Defaults to TRUE. Set to FALSE for redundancy analysis only
<code>auto</code>	Boolean. Should redundancy reduction be automated? Defaults to TRUE. Set to FALSE for manual selection
<code>label_latent</code>	Boolean. Should latent variables be labelled? Defaults to TRUE. Set to FALSE for arbitrary labelling (i.e., " <code>LV_</code> ")
<code>reduce.method</code>	Character. How should data be reduced? Defaults to " <code>latent</code> " <ul style="list-style-type: none"> • "<code>latent</code>" Redundant variables will be combined into a latent variable • "<code>remove</code>" All but one redundant variable will be removed • "<code>sum</code>" Redundant variables are combined by summing across cases (rows)

lavaan.args	<p>List. If <code>reduce.method = "latent"</code>, then <code>lavaan</code>'s <code>cfa</code> function will be used to create latent variables to reduce variables. Arguments should be input as a list. Some example arguments (see lavOptions for full details):</p> <ul style="list-style-type: none"> • <code>estimator</code> Estimator to use for latent variables (see Estimators) for more details. Defaults to "MLR" for continuous data and "WLSMV" for mixed and categorical data. Data are considered continuous data if they have 6 or more categories (see Rhemtulla, Brosseau-Liard, & Savalei, 2012) • <code>missing</code> How missing data should be handled. Defaults to "fiml" • <code>std.lv</code> If TRUE, the metric of each latent variable is determined by fixing their (residual) variances to 1.0. If FALSE, the metric of each latent variable is determined by fixing the factor loading of the first indicator to 1.0. If there are multiple groups, <code>std.lv = TRUE</code> and "loadings" is included in the <code>group.label</code> argument, then only the latent variances i of the first group will be fixed to 1.0, while the latent variances of other groups are set free. Defaults to TRUE
adhoc	<p>Boolean. Should adhoc check of redundancies be performed? Defaults to TRUE. If TRUE, adhoc check will run the redundancy analysis on the reduced variable set to determine if there are any remaining redundancies. This check is performed with the arguments: <code>method = "wTO"</code>, <code>type = "threshold"</code>, and <code>sig = .20</code>. This check is based on Christensen, Garrido, and Golino's (2020) simulation where these parameters were found to be the most conservative, demonstrating few false positives and false negatives</p>
plot.redundancy	<p>Boolean. Should redundancies be plotted in a network plot? Defaults to FALSE</p>
plot.args	<p>List. Arguments to be passed onto ggnet2. Defaults:</p> <ul style="list-style-type: none"> • <code>vsize = 6</code> Changes node size • <code>alpha = 0.4</code> Changes transparency • <code>label.size = 5</code> Changes label size • <code>edge.alpha = 0.7</code> Changes edge transparency

Value

Returns a list:

redundancy	<p>A list containing several objects:</p> <ul style="list-style-type: none"> • <code>redudant</code> Vectors nested within the list corresponding to redundant nodes with the name of object in the list • <code>data</code> Original data • <code>correlation</code> Correlation matrix of original data • <code>weights</code> Weights determine by weighted topological overlap, partial correlation, or zero-order correlation • <code>network</code> If <code>method = "wTO"</code>, then the network computed following EGA with EBICglasso network estimation • <code>plot</code> If <code>redundancy.plot = TRUE</code>, then a plot of all redundancies found • <code>descriptives</code>
------------	---

- basic A vector containing the mean, standard deviation, median, median absolute deviation (MAD), 3 times the MAD, 6 times the MAD, minimum, maximum, and critical value for the overlap measure (i.e., weighted topological overlap, partial correlation, or threshold)
 - centralTendency A matrix for all (absolute) non-zero values and their respective standard deviation from the mean and median absolute deviation from the median
 - method Returns method argument
 - type Returns type argument
 - distribution If type != "threshold", then distribution that was used to determine significance
- reduced If reduce = TRUE, then a list containing:
- data New data with redundant variables merged or removed
 - mergedA matrix containing the variables that were decided to be redundant with one another
 - method Method used to perform redundancy reduction
- adhoc If adhoc = TRUE, then the adhoc check containing the same objects as in the redundancy list object in the output

Author(s)

Alexander Christensen <alexpaulchristensen@gmail.com>

References

Simulation using UVA

Christensen, A. P., Garrido, L. E., & Golino, H. (under review). Unique Variable Analysis: A novel approach for detecting redundant variables in multivariate data. *PsyArXiv*.

Implementation of UVA (formally node .redundant)

Christensen, A. P., Golino, H., & Silvia, P. J. (2020). A psychometric network perspective on the validity and validation of personality trait questionnaires. *European Journal of Personality*, *34*, 1095-1108.

wTO measure

Nowick, K., Gernat, T., Almaas, E., & Stubbs, L. (2009). Differences in human and chimpanzee gene expression patterns define an evolving network of transcription factors in brain. *Proceedings of the National Academy of Sciences*, *106*, 22358-22363.

Selection of CFA Estimator

Rhemtulla, M., Brosseau-Liard, P. E., & Savalei, V. (2012). When can categorical variables be treated as continuous? A comparison of robust continuous and categorical SEM estimation methods under suboptimal conditions. *Psychological Methods*, *17*, 354-373.

Examples

```
# Select Five Factor Model personality items only
idx <- na.omit(match(gsub("-", ""), unlist(psychTools::spi.keys[1:5])), colnames(psychTools::spi)))
items <- psychTools::spi[,idx]
```

```

# Change names in redundancy output to each item's description
key.ind <- match(colnames(items), as.character(psychTools::spi.dictionary$item_id))
key <- as.character(psychTools::spi.dictionary$item[key.ind])

# Automated selection of redundant variables (default)

uva.results <- UVA(data = items, key = key)

# Manual selection of redundant variables
if(interactive()){
uva.results <- UVA(data = items, key = key, type = "adapt")
}

```

vn.entropy	<i>Entropy Fit Index using Von Neumman's entropy (Quantum Information Theory) for correlation matrices</i>
------------	--

Description

Computes the fit of a dimensionality structure using Von Neumman's entropy when the input is a correlation matrix. Lower values suggest better fit of a structure to the data.

Usage

```
vn.entropy(data, structure)
```

Arguments

data	A dataframe or a correlation matrix
structure	A vector representing the structure (numbers or labels for each item). Can be theoretical factors or the structure detected by EGA

Value

Returns a list containing:

VN.Entropy.Fit	The Entropy Fit Index using Von Neumman's entropy
Total.Correlation	The total correlation of the dataset
Average.Entropy	The average entropy of the dataset

Author(s)

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References

Golino, H., Moulder, R. G., Shi, D., Christensen, A. P., Garrido, L. E., Nieto, M. D., Nesselroade, J., Sadana, R., Thiyagarajan, J. A., & Boker, S. M. (2020). Entropy fit indices: New fit measures for assessing the structure and dimensionality of multiple latent variables. *Multivariate Behavioral Research*.

See Also

[EGA](#) to estimate the number of dimensions of an instrument using EGA and [CFA](#) to verify the fit of the structure suggested by EGA using confirmatory factor analysis.

Examples

```
# Load data
dep <- depression[,24:44]

# Estimate EGA
## plot.type = "qqgraph" used for CRAN checks
## plot.type = "GGally" is the default
ega.dep <- EGA(data = dep, model = "glasso", plot.type = "qgraph")

# Compute entropy indices
vn.entropy(data = ega.dep$correlation, structure = ega.dep$wc)
```

wmt2

WMT-2 Data

Description

A response matrix (n = 1185) of the Wiener Matrizen-Test 2 (WMT-2).

A response matrix (n = 1185) of the Wiener Matrizen-Test 2 (WMT-2).

Usage

```
data(wmt2)
```

```
data(wmt2)
```

Format

A 1185x24 response matrix

A 1185x24 response matrix

Examples

```
data("wmt2")
```

```
data("wmt2")
```

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