

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 estimates the number of dimensions in psychological data using network estimation methods and community detection algorithms. A bootstrap method is provided to assess the stability of dimensions and items. Fit is evaluated using the Entropy Fit family of indices. Unique Variable Analysis evaluates the extent to which items are locally dependent (or redundant). Network loadings provide similar information to factor loadings and can be used to compute network scores. A bootstrap and permutation approach are available to assess configural and metric invariance. Hierarchical structures can be detected using Hierarchical EGA. Time series and intensive longitudinal data can be analyzed using Dynamic EGA, supporting individual, group, and population level assessments.

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Suggests fitdistrplus, gridExtra, pbapply, progress, psych, pwr, RColorBrewer

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URL <https://r-ega.net>

BugReports <https://github.com/hfgolino/EGAnet/issues>

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EGAnet-package	<i>EGAnet-package</i>
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Description

Implements the Exploratory Graph Analysis (EGA) framework for dimensionality and psychometric assessment. EGA estimates the number of dimensions in psychological data using network estimation methods and community detection algorithms. A bootstrap method is provided to assess the stability of dimensions and items. Fit is evaluated using the Entropy Fit family of indices. Unique Variable Analysis evaluates the extent to which items are locally dependent (or redundant). Network loadings provide similar information to factor loadings and can be used to compute network scores. A bootstrap and permutation approach are available to assess configural and metric invariance. Hierarchical structures can be detected using Hierarchical EGA. Time series and intensive longitudinal data can be analyzed using Dynamic EGA, supporting individual, group, and population level assessments.

Author(s)

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References

- Christensen, A. P. (2023). Unidimensional community detection: A Monte Carlo simulation, grid search, and comparison. *PsyArXiv*.
 # Related functions: [community.unidimensional](#)
- Christensen, A. P., Garrido, L. E., & Golino, H. (2023). Unique variable analysis: A network psychometrics method to detect local dependence. *Multivariate Behavioral Research*.
 # Related functions: [UVA](#)
- Christensen, A. P., Garrido, L. E., Guerra-Pena, K., & Golino, H. (2023). Comparing community detection algorithms in psychometric networks: A Monte Carlo simulation. *Behavior Research Methods*.
 # Related functions: [EGA](#)
- 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](#)

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. #

Related functions: [CFA](#) and [EGA](#)

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: [CFA](#), [EGA](#), and [bootEGA](#)

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., Nesselroade, J. R., & Christensen, A. P. (2022). Towards a psychology of individuals: The ergodicity information index and a bottom-up approach for finding generalizations. *PsyArXiv*.

Related functions: [boot.ergoInfo](#), [ergoInfo](#), [jsd](#), and [infoCluster](#)

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. (2020). 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. (2021). Optimizing Walktrap's community detection in networks using the Total Entropy Fit Index. *PsyArXiv*.

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

Jamison, L., Golino, H., & Christensen, A. P. (2023). Metric invariance in exploratory graph analysis via permutation testing. *PsyArXiv*.

Related functions: [invariance](#)

Shi, D., Christensen, A. P., Day, E., Golino, H., & Garrido, L. E. (2023). A Bayesian approach for dimensionality assessment in psychological networks. *PsyArXiv*

Related functions: [EGA](#)

See Also

Useful links:

- <https://r-ega.net>
- Report bugs at <https://github.com/hfgolino/EGAnet/issues>

auto.correlate	<i>Automatic correlations</i>
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Description

This wrapper is similar to [cor_auto](#). There are some minor adjustments that make this function simpler and to function within [EGAnet](#). NA values are not treated as categories (this behavior differs from [cor_auto](#))

Usage

```
auto.correlate(
  data,
  corr = c("cosine", "kendall", "pearson", "spearman"),
  ordinal.categories = 7,
  forcePD = TRUE,
  na.data = c("pairwise", "listwise"),
  empty.method = c("none", "zero", "all"),
  empty.value = c("none", "point_five", "one_over"),
  forceReturn = FALSE,
  verbose = FALSE,
  ...
)
```

Arguments

<code>data</code>	Matrix or data frame. Should consist only of variables to be used in the analysis
<code>corr</code>	Character (length = 1). The standard correlation method to be used. Defaults to "pearson". Using "pearson" will compute polychoric, tetrachoric, polyserial, and biserial correlations for categorical and categorical/continuous correlations by default. To obtain "pearson" correlations regardless, use cor . Other options of "kendall" and "spearman" are provided for completeness and use cor.cosine is also available
<code>ordinal.categories</code>	Numeric (length = 1). <i>Up to</i> the number of categories <i>before</i> a variable is considered continuous. Defaults to 7 categories before 8 is considered continuous

forcePD	Boolean (length = 1). Whether positive definite matrix should be enforced. Defaults to TRUE
na.data	Character (length = 1). How should missing data be handled? Defaults to "pairwise". Available options: <ul style="list-style-type: none"> • "pairwise" — Computes correlation for all available cases between two variables • "listwise" — Computes correlation for all complete cases in the dataset
empty.method	Character (length = 1). Method for empty cell correction in <code>polychoric.matrix</code> . Defaults to "none". Available options: <ul style="list-style-type: none"> • "none" — Adds no value (<code>empty.value = "none"</code>) to the empirical joint frequency table between two variables • "zero" — Adds <code>empty.value</code> to the cells with zero in the joint frequency table between two variables • "all" — Adds <code>empty.value</code> to all in the joint frequency table between two variables
empty.value	Character (length = 1). Value to add to the joint frequency table cells in <code>polychoric.matrix</code> . Defaults to "none". Accepts numeric values between 0 and 1 or specific methods: <ul style="list-style-type: none"> • "none" — Adds no value (0) to the empirical joint frequency table between two variables • "point_five" — Adds 0.5 to the cells defined by <code>empty.method</code> • "one_over" — Adds $1/n$ where n equals the number of cells based on <code>empty.method</code>. For <code>empty.method = "zero"</code>, n equals the number of zero cells
forceReturn	Boolean (length = 1). Whether correlation matrix should be forced to return. Defaults to FALSE. Set to TRUE to receive the correlation matrix "as is"
verbose	Boolean (length = 1). Whether messages should be printed. Defaults to FALSE
...	Not actually used but makes it easier for general functionality in the package

Author(s)

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Examples

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

# Obtain correlations
wmt_corr <- auto.correlate(wmt)
```

boot.ergoInfo *Bootstrap Test for the Ergodicity Information Index*

Description

Tests the Ergodicity Information Index obtained in the empirical sample with a distribution of EII obtained by a variant of bootstrap sampling (see **Details** for the procedure)

Usage

```
boot.ergoInfo(
  dynEGA.object,
  EII,
  use = c("edge.list", "unweighted", "weighted"),
  shuffles = 5000,
  iter = 100,
  ncores,
  verbose = TRUE
)
```

Arguments

dynEGA.object	A dynEGA or a dynEGA.ind.pop object. If a dynEGA object, then level = c("individual", "population") is required
EII	A ergoInfo object used to estimate the Empirical Ergodicity Information Index or the estimated value of EII estimated using the ergoInfo function. Inherits use from ergoInfo . If no ergoInfo object is provided, then it is estimated
use	Character (length = 1). 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 = "unweighted". Current options are: <ul style="list-style-type: none"> • "edge.list" — Calculates the algorithm complexity using the list of edges • "unweighted" — Calculates the algorithm complexity using the binary weights of the encoded prime transformed network. 0 = edge absent and 1 = edge present • "weighted" — Calculates the algorithm complexity using the weights of encoded prime-weight transformed network
shuffles	Numeric. Number of shuffles used to compute the Kolmogorov complexity. Defaults to 5000
iter	Numeric (length = 1). Number of replica samples to generate from the bootstrap analysis. Defaults to 100 (1000 for robustness)
ncores	Numeric (length = 1). Number of cores to use in computing results. Defaults to ceiling(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 type: parallel::detectCores()
verbose	Boolean (length = 1). Should progress be displayed? Defaults to TRUE. Set to FALSE to not display progress

Details

In traditional bootstrap sampling, individual participants are resampled with replacement from the empirical sample. This process is time consuming when carried out across v number of variables, n number of participants, t number of time points, and i number of iterations. Instead, `boot.ergoInfo` uses the premise of an ergodic process to establish more efficient test that works directly on the sample's networks.

With an ergodic process, the expectation is that all individuals will have a systematic relationship with the population. Destroying this relationship should result in a significant loss of information. Following this conjecture, `boot.ergoInfo` shuffles a random subset of edges that exist in the **population** that is *equal* to the number of shared edges it has with an individual. An individual's unique edges remain the same, controlling for their unique information. The result is a replicate individual that contains the same total number of edges as the actual individual but its shared information with the population has been scrambled.

This process is repeated over each individual to create a replicate sample and is repeated for X iterations (e.g., 100). This approach creates a sampling distribution that represents the expected information between the population and individuals when a random process generates the shared information between them. If the shared information between the population and individuals in the empirical sample is sufficiently meaningful, then this process should result in significant information loss.

How to interpret the results: the result of `boot.ergoInfo` is a sampling distribution of EII values that would be expected if the process was random (null distribution). If the empirical EII value is *greater than* or not significantly different from the null distribution, then the empirical data can be expected to be generated from a nonergodic process and the population structure is not sufficient to describe all individuals. If the empirical EII value is significantly *lower than* the null distribution, then the empirical data can be described by the population structure – the population structure is sufficient to describe all individuals.

Value

Returns a list containing:

<code>empirical.ergoInfo</code>	Empirical Ergodicity Information Index
<code>boot.ergoInfo</code>	The values of the Ergodicity Information Index obtained in the bootstrap
<code>p.value</code>	The two-sided p -value of the bootstrap test for the Ergodicity Information Index. The null hypothesis is that the empirical Ergodicity Information index is equal to or greater than the expected value of the EII with small variation in the population structure
<code>effect</code>	Indicates whether the empirical EII is greater or less than the bootstrap distribution of EII.
<code>interpretation</code>	How you can interpret the result of the test in plain English

Author(s)

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References

Original Implementation

Golino, H., Nesselroade, J. R., & Christensen, A. P. (2022). Toward a psychology of individuals: The ergodicity information index and a bottom-up approach for finding generalizations. *PsyArXiv*.

See Also

[plot.EGAnet](#) for plot usage in [EGAnet](#)

Examples

```
# Obtain simulated data
sim.data <- sim.dynEGA

## Not run:
# Dynamic EGA individual and population structures
dyn1 <- dynEGA.ind.pop(
  data = sim.dynEGA[,-26], n.embed = 5, tau = 1,
  delta = 1, id = 25, use.derivatives = 1,
  model = "glasso", ncores = 2, corr = "pearson"
)

# Empirical Ergodicity Information Index
eii1 <- ergoInfo(dynEGA.object = dyn1, use = "unweighted")

# Bootstrap Test for Ergodicity Information Index
testing.ergoInfo <- boot.ergoInfo(
  dynEGA.object = dyn1, EII = eii1,
  ncores = 2, use = "unweighted"
)

# Plot result
plot(testing.ergoInfo)

# Example using `dynEGA`
dyn2 <- dynEGA(
  data = sim.dynEGA, n.embed = 5, tau = 1,
  delta = 1, use.derivatives = 1, ncores = 2,
  level = c("individual", "population")
)

# Empirical Ergodicity Information Index
eii2 <- ergoInfo(dynEGA.object = dyn2, use = "unweighted")

# Bootstrap Test for Ergodicity Information Index
testing.ergoInfo2 <- boot.ergoInfo(
  dynEGA.object = dyn2, EII = eii2,
  ncores = 2
)

# Plot result
plot(testing.ergoInfo2)
```

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

```
boot.wmt          bootEGA Results of wmt2Data
```

Description

`bootEGA` results from `boot.wmt <- bootEGA(wmt2[,7:24], seed = 1234)`

Usage

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

Format

A list with 12 objects (see **Value** in `bootEGA`)

Examples

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

```
bootEGA          Bootstrap Exploratory Graph Analysis
```

Description

`bootEGA` Estimates the number of dimensions of iter bootstraps using the empirical zero-order correlation matrix ("parametric") or "resampling" from the empirical dataset (non-parametric). `bootEGA` estimates a typical median network structure, which is formed by the median or mean pairwise (partial) correlations over the *iter* bootstraps (see **Details** for information about the typical median network structure).

Usage

```
bootEGA(
  data,
  n = NULL,
  corr = c("auto", "cor_auto", "cosine", "pearson", "spearman"),
  na.data = c("pairwise", "listwise"),
  model = c("BGGM", "glasso", "TMFG"),
  algorithm = c("leiden", "louvain", "walktrap"),
  uni.method = c("expand", "LE", "louvain"),
  iter = 500,
  type = c("parametric", "resampling"),
  ncores,
```

```

EGA.type = c("EGA", "EGA.fit", "hierEGA", "riEGA"),
plot.itemStability = TRUE,
typicalStructure = FALSE,
plot.typicalStructure = FALSE,
seed = NULL,
verbose = TRUE,
...
)

```

Arguments

<code>data</code>	Matrix or data frame. Should consist only of variables to be used in the analysis
<code>n</code>	Numeric (length = 1). Sample size if data provided is a correlation matrix
<code>corr</code>	Character (length = 1). Method to compute correlations. Defaults to "auto". Available options: <ul style="list-style-type: none"> • "auto" — Automatically computes appropriate correlations for the data using Pearson's for continuous, polychoric for ordinal, tetrachoric for binary, and polyserial/biserial for ordinal/binary with continuous. To change the number of categories that are considered ordinal, use <code>ordinal.categories</code> (see polychoric.matrix for more details) • "cor_auto" — Uses <code>cor_auto</code> to compute correlations. Arguments can be passed along to the function • "cosine" — Uses <code>cosine</code> to compute cosine similarity • "pearson" — Pearson's correlation is computed for all variables regardless of categories • "spearman" — Spearman's rank-order correlation is computed for all variables regardless of categories <p>For other similarity measures, compute them first and input them into data with the sample size (n)</p>
<code>na.data</code>	Character (length = 1). How should missing data be handled? Defaults to "pairwise". Available options: <ul style="list-style-type: none"> • "pairwise" — Computes correlation for all available cases between two variables • "listwise" — Computes correlation for all complete cases in the dataset
<code>model</code>	Character (length = 1). Defaults to "glasso". Available options: <ul style="list-style-type: none"> • "BGM" — Computes the Bayesian Gaussian Graphical Model. Set argument <code>ordinal.categories</code> to determine levels allowed for a variable to be considered ordinal. See <code>?BGM:estimate</code> for more details • "glasso" — Computes the GLASSO with EBIC model selection. See EBICglasso.qgraph for more details • "TMFG" — Computes the TMFG method. See TMFG for more details
<code>algorithm</code>	Character or <code>igraph</code> <code>cluster_*</code> function (length = 1). Defaults to "walktrap". Three options are listed below but all are available (see community.detection for other options):

	<ul style="list-style-type: none"> • "leiden" — See cluster_leiden for more details • "louvain" — By default, "louvain" will implement the Louvain algorithm using the consensus clustering method (see community.consensus for more information). This function will implement <code>consensus.method = "most_common"</code> and <code>consensus.iter = 1000</code> unless specified otherwise • "walktrap" — See cluster_walktrap for more details
uni.method	<p>Character (length = 1). What unidimensionality method should be used? Defaults to "louvain". Available options:</p> <ul style="list-style-type: none"> • "expand" — Expands the correlation matrix with four variables correlated 0.50. 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 method was used in the Golino et al.'s (2020) <i>Psychological Methods</i> simulation • "LE" — Applies the Leading Eigenvector algorithm (cluster_leading_eigen) on the empirical correlation matrix. If the number of dimensions is 1, then the Leading Eigenvector solution is used; otherwise, regular EGA is used. This method was used in the Christensen et al.'s (2023) <i>Behavior Research Methods</i> simulation • "louvain" — Applies the Louvain algorithm (cluster_louvain) on the empirical correlation matrix. If the number of dimensions is 1, then the Louvain solution is used; otherwise, regular EGA is used. This method was validated Christensen's (2022) <i>PsyArXiv</i> simulation. Consensus clustering can be used by specifying either "consensus.method" or "consensus.iter"
iter	Numeric (length = 1). Number of replica samples to generate from the bootstrap analysis. Defaults to 500 (recommended)
type	<p>Character (length = 1). What type of bootstrap should be performed? Defaults to "parametric". Available options:</p> <ul style="list-style-type: none"> • "parametric" — Generates <code>iter</code> new datasets from (multivariate normal random distributions) based on the original dataset using mvrnorm • "resampling" — Generates <code>iter</code> new datasets from random subsamples of the original data
ncores	<p>Numeric (length = 1). Number of cores to use in computing results. Defaults to <code>ceiling(parallel::detectCores() / 2)</code> or half of your computer's processing power. Set to 1 to not use parallel computing</p> <p>If you're unsure how many cores your computer has, then type: <code>parallel::detectCores()</code></p>
EGA.type	<p>Character (length = 1). Type of EGA model to use. Defaults to "EGA" Available options:</p> <ul style="list-style-type: none"> • "EGA" — Uses standard exploratory graph analysis • "EGA.fit" — Uses tefi to determine best fit of EGA • "hierEGA" — Uses hierarchical exploratory graph analysis • "riEGA" — Uses random-intercept exploratory graph analysis <p>Arguments for EGA.type can be added (see links for details on specific function arguments)</p>

plot.itemStability	Boolean (length = 1). Should the plot be produced for item.replication? Defaults to TRUE
typicalStructure	Boolean (length = 1). If TRUE, returns the median ("glasso" or "BGM") or mean ("TMFG") network structure and estimates its dimensions (see Details for more information). Defaults to FALSE
plot.typicalStructure	Boolean (length = 1). If TRUE, returns a plot of the typical network structure. Defaults to FALSE
seed	Numeric (length = 1). Defaults to NULL or random results. Set for reproducible results. See Reproducibility and PRNG for more details on random number generation in EGAnet
verbose	Boolean (length = 1). Should progress be displayed? Defaults to TRUE. Set to FALSE to not display progress
...	Additional arguments that can be passed on to auto.correlate , network.estimate , community.detection , community.consensus , EGA , EGA.fit , hierEGA , and riEGA

Details

The typical network structure is derived from the median (or mean) value of each pairwise relationship. These values tend to reflect the "typical" value taken by an edge across the bootstrap networks. Afterward, the same community detection algorithm is applied to the typical network as the bootstrap networks.

Because the community detection algorithm is applied to the typical network structure, there is a possibility that the community algorithm determines a different number of dimensions than the median number derived from the bootstraps. The typical network structure (and number of dimensions) may *not* match the empirical [EGA](#) number of dimensions or the median number of dimensions from the bootstrap. This result is known and *not* a bug.

Value

Returns a list containing:

iter	Number of replica samples in bootstrap
bootGraphs	A list containing the networks of each replica sample
bootCorrs	A list containing the zero-order correlations of each replica sample
boot.wc	A matrix of membership assignments for each replica network with variables down the columns and replicas across the rows
boot.ndim	Number of dimensions identified in each replica sample
summary.table	A data frame containing number of replica samples, median, standard deviation, standard error, 95% confidence intervals, and quantiles (lower = 2.5% and upper = 97.5%)
frequency	A data frame containing the proportion of times the number of dimensions was identified (e.g., .85 of 1,000 = 850 times that specific number of dimensions was found)

TEFI	tefi value for each replica sample
type	Type of bootstrap used
EGA	Output of the empirical EGA results (output will vary based on EGA.type)
EGA.type	Type of *EGA function used
typicalGraph	A list containing: <ul style="list-style-type: none"> • graph — Network matrix of the median network structure • typical.dim.variables — An ordered matrix of item allocation • wc — Membership assignments of the median network
plot.typical.ega	Plot output if plot.typicalStructure = TRUE

Author(s)

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

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.

See Also

[itemStability](#) to estimate the stability of the variables in the empirical dimensions and [dimensionStability](#) to estimate the stability of the dimensions (structural consistency)

Examples

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

## Not run:
# Standard EGA parametric example
boot.wmt <- bootEGA(
  data = wmt, iter = 500,
  type = "parametric", ncores = 2
)

# Standard resampling example
boot.wmt <- bootEGA(
  data = wmt, iter = 500,
  type = "resampling", ncores = 2
)

# Example using {igraph} `cluster_*` function
boot.wmt.spinglass <- bootEGA(
  data = wmt, iter = 500,
  algorithm = igraph::cluster_spinglass,
  # use any function from {igraph}
```

```

    type = "parametric", ncores = 2
  )

# EGA fit example
boot.wmt.fit <- bootEGA(
  data = wmt, iter = 500,
  EGA.type = "EGA.fit",
  type = "parametric", ncores = 2
)

# Hierarchical EGA example
boot.wmt.hier <- bootEGA(
  data = wmt, iter = 500,
  EGA.type = "hierEGA",
  type = "parametric", ncores = 2
)

# Random-intercept EGA example
boot.wmt.ri <- bootEGA(
  data = wmt, iter = 500,
  EGA.type = "riEGA",
  type = "parametric", ncores = 2
)
## End(Not run)

```

CFA

CFA Fit of [EGA](#) or [hierEGA](#) Structure

Description

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

Usage

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

Arguments

<code>ega.obj</code>	An EGA object or an hierEGA
<code>data</code>	Matrix or data frame. Should consist only of variables to be used in the analysis
<code>estimator</code>	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 <code>cfa</code>
<code>summary</code>	Summary output from <code>lavaan-class</code>
<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 <code>fitMeasures</code> function (see examples)

Author(s)

Hudson F. Golino <hfg9s at virginia.edu>

References**Demonstrative use**

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.

Initial implementation

Golino, H., & Epskamp, S. (2017). Exploratory graph analysis: A new approach for estimating the number of dimensions in psychological research. *PLoS ONE*, 12, e0174035.

Examples

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

## Not run:
# Estimate EGA
ega.wmt <- EGA(
  data = wmt,
  plot.EGA = FALSE # No plot for CRAN checks
)

# Fit CFA model to EGA results
cfa.wmt <- CFA(
  ega.obj = ega.wmt, estimator = "WLSMV",
  plot.CFA = FALSE, # No plot for CRAN checks
  data = wmt
)

# Additional fit measures
lavaan::fitMeasures(cfa.wmt$fit, fit.measures = "all")
## End(Not run)
```

color_palette_EGA [EGA Color Palettes](#)

Description

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

Usage

```
color_palette_EGA(
  name = c("polychrome", "blue.ridge1", "blue.ridge2", "rainbow", "rio", "itacare",
           "grayscale"),
  wc,
  sorted = FALSE
)
```

Arguments

name	Character. Name of color scheme (see RColorBrewer). Defaults to "polychrome". EGA palettes: <ul style="list-style-type: none"> • "polychrome" — Default 40 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
wc	For custom colors, enter HEX codes for each dimension in a vector Numeric 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
sorted	Boolean. Should colors be sorted by wc? Defaults to FALSE

Value

Vector of colors for community memberships

Author(s)

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

See Also

[plot.EGAnet](#) for plot usage in [EGAnet](#)

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 = c("#7FD1B9", "#24547e"), wc = ega.wmt$wc)
```

community.compare *Compares Community Detection Solutions Using Permutation*

Description

A permutation implementation to determine statistical significance of whether the community comparison measure is different from zero

Usage

```
community.compare(
  base,
  comparison,
  method = c("vi", "nmi", "split.join", "rand", "adjusted.rand"),
  iter = 1000,
  shuffle.base = TRUE,
  verbose = TRUE,
  seed = NULL
)
```

Arguments

base	Character or numeric vector. A vector of characters or numbers that are treated as the baseline communities
comparison	Character or numeric vector (length = length(base)). A vector of characters or numbers that are treated as the baseline communities
method	Character (length = 1). Comparison metrics from compare . Defaults to "adjusted.rand". Available options: <ul style="list-style-type: none"> • "vi" — Variation of information (Meila, 2003) • "nmi" — Normalized mutual information (Danon et al., 2003) • "split.join" — Split-join distance (Dongen, 2000) • "rand" — Rand index (Rand, 1971) • "adjusted.rand" — adjusted Rand index (Hubert & Arabie, 1985; Steinley, 2004)

iter	Numeric (length = 1). Number of permutations to perform. Defaults to 1000 (recommended)
shuffle.base	Boolean (length = 1). Whether the base cluster solution should be shuffled. Defaults to TRUE to remain consistent with original implementation (Qannari et al., 2014); however, from a theoretical standpoint, it might make sense to only shuffle the comparison to determine whether it is specifically different from the recognized base
verbose	Boolean (length = 1). Should progress be displayed? Defaults to TRUE. Set to FALSE to not display progress
seed	Numeric (length = 1). Defaults to NULL or random results. Set for reproducible results. See Reproducibility and PRNG for more details on random number generation in EGAnet

Value

Returns data frame containing method used (Method), empirical or observed value (Empirical), and p-value based on the permutation test (p.value)

Author(s)

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

References

Implementation of Permutation Test

Qannari, E. M., Courcoux, P., & Faye, P. (2014). Significance test of the adjusted Rand index. Application to the free sorting task. *Food Quality and Preference*, 32, 93–97.

Variation of Information

Meila, M. (2003, August). Comparing clusterings by the variation of information. In *Learning Theory and Kernel Machines: 16th Annual Conference on Learning Theory and 7th Kernel Workshop*, COLT/Kernel 2003, Washington, DC, USA, August 24-27, 2003. Proceedings (pp. 173-187). Berlin, DE: Springer Berlin Heidelberg.

Normalized Mutual Information

Danon, L., Diaz-Guilera, A., Duch, J., & Arenas, A. (2005). Comparing community structure identification. *Journal of Statistical Mechanics: Theory and Experiment*, 2005(09), P09008.

Split-join Distance

Dongen, S. (2000). Performance criteria for graph clustering and Markov cluster experiments. *CWI (Centre for Mathematics and Computer Science)*.

Rand Index

Rand, W. M. (1971). Objective criteria for the evaluation of clustering methods. *Journal of the American Statistical Association*, 66(336), 846-850.

Adjusted Rand Index

Hubert, L., & Arabie, P. (1985). Comparing partitions. *Journal of Classification*, 2, 193-218.

Steinley, D. (2004). Properties of the Hubert-Arabie adjusted rand index. *Psychological Methods*, 9(3), 386.

Examples

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

# Estimate network
network <- EBICglasso.qgraph(data = wmt)

# Compute Edge Betweenness
edge_between <- community.detection(network, algorithm = "edge_betweenness")

# Compute Fast Greedy
fast_greedy <- community.detection(network, algorithm = "fast_greedy")

# Perform permutation test
community.compare(edge_between, fast_greedy)
```

community.consensus *Applies the Consensus Clustering Method (Louvain only)*

Description

Applies the consensus clustering method introduced by (Lancichinetti & Fortunato, 2012). The original implementation of this method applies a community detection algorithm repeatedly to the same network. With stochastic networks, the algorithm is likely to identify different community solutions with many repeated applications.

Usage

```
community.consensus(
  network,
  order = c("lower", "higher"),
  resolution = 1,
  consensus.method = c("highest_modularity", "iterative", "most_common", "lowest_tefi"),
  consensus.iter = 1000,
  correlation.matrix = NULL,
  allow.singleton = FALSE,
  membership.only = TRUE,
  ...
)
```

Arguments

network	Matrix or igraph network object
order	Character (length = 1). Defaults to "higher". Whether "lower" or "higher" order memberships from the Louvain algorithm should be obtained for the consensus. The "lower" order Louvain memberships are from the first initial pass of the Louvain algorithm whereas the "higher" order Louvain memberships are from the last pass of the Louvain algorithm

resolution	Numeric (length = 1). A parameter that adjusts modularity to allow the algorithm to prefer smaller (resolution > 1) or larger (0 < resolution < 1) communities. Defaults to 1 (standard modularity computation)
consensus.method	Character (length = 1). Defaults to "most_common". Available options for arriving at a consensus (<i>Note</i> : All methods except "iterative" are considered experimental until validated): <ul style="list-style-type: none"> • "highest_modularity" — EXPERIMENTAL. Selects the community solution with the highest modularity across the applications. Modularity is a reasonable metric for identifying the number of communities in a network but it comes with limitations (e.g., resolution limit) • "iterative" — The original approach proposed by Lancichinetti & Fortunato (2012). See "Details" for more information • "most_common" — Selects the community solution that appears the most frequently across the applications. The idea behind this method is that the solution that appears most often will be the most likely solution for the algorithm as well as most reproducible. Can be less stable as the number of nodes increase requiring a larger value for consensus.iter. This method is the default • "lowest_tefi" — EXPERIMENTAL. Selects the community solution with the lowest Total Entropy Fit Index (tefi) across the applications. TEFI is a reasonable metric to identify the number of communities in a network based on Golino, Moulder et al. (2020)
consensus.iter	Numeric (length = 1). Number of algorithm applications to the network. Defaults to 1000
correlation.matrix	Symmetric matrix. Used for computation of tefi . Only needed when consensus.method = "tefi"
allow.singleton	Boolean (length = 1). Whether singleton or single node communities should be allowed. Defaults to FALSE. When FALSE, singleton communities will be set to missing (NA); otherwise, when TRUE, singleton communities will be allowed
membership.only	Boolean. Whether the memberships only should be output. Defaults to TRUE. Set to FALSE to obtain all output for the community detection algorithm
...	Not actually used but makes it easier for general functionality in the package

Details

The goal of the consensus clustering method is to identify a stable solution across algorithm applications to derive a "consensus" clustering. The standard or "iterative" approach is to apply the community detection algorithm N times. Then, a co-occurrence matrix is created representing how often each pair of nodes co-occurred across the applications. Based on some cut-off value (e.g., 0.30), co-occurrences below this value are set to zero, forming a "new" sparse network. The procedure proceeds until all nodes co-occur with all other nodes in their community (or a proportion of 1.00).

Variations of this procedure are also available in this package but are **experimental**. Use these experimental procedures with caution. More work is necessary before these experimental procedures are validated

At this time, seed setting for consensus clustering is not supported

Value

Returns either a vector with the selected solution or a list when `membership.only = FALSE`:

<code>selected_solution</code>	Resulting solution from the consensus method
<code>memberships</code>	Matrix of memberships across the consensus iterations
<code>proportion_table</code>	For methods that use frequency, a table that reports those frequencies alongside their corresponding memberships

Author(s)

Hudson Golino <hfg9s at virginia.edu> and Alexander P. Christensen <alexpaulchristensen@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(10), P10008.

Consensus clustering

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

Entropy fit indices

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*.

Examples

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

# Estimate correlation matrix
correlation.matrix <- auto.correlate(wmt)

# Estimate network
network <- EBICglasso.qgraph(data = wmt)

# Compute standard Louvain with highest modularity approach
community.consensus(
  network,
```

```

    consensus.method = "highest_modularity"
  )

# Compute standard Louvain with iterative (original) approach
community.consensus(
  network,
  consensus.method = "iterative"
)

# Compute standard Louvain with most common approach
community.consensus(
  network,
  consensus.method = "most_common"
)

# Compute standard Louvain with lowest TEFI approach
community.consensus(
  network,
  consensus.method = "lowest_tefi",
  correlation.matrix = correlation.matrix
)

```

community.detection *Apply a Community Detection Algorithm*

Description

General function to apply community detection algorithms available in igraph. Follows the EGAnet approach of setting singleton and disconnected nodes to missing (NA)

Usage

```

community.detection(
  network,
  algorithm = c("edge_betweenness", "fast_greedy", "fluid", "infomap", "label_prop",
    "leading_eigen", "leiden", "louvain", "optimal", "spinglass", "walktrap"),
  allow.singleton = FALSE,
  membership.only = TRUE,
  ...
)

```

Arguments

network	Matrix or igraph network object
algorithm	Character or igraph cluster_* function (length = 1). Available options: <ul style="list-style-type: none"> "edge_betweenness" — See cluster_edge_betweenness for more details

- "fast_greedy" — See [cluster_fast_greedy](#) for more details
- "fluid" — See [cluster_fluid_communities](#) for more details
- "infomap" — See [cluster_infomap](#) for more details
- "label_prop" — See [cluster_label_prop](#) for more details
- "leading_eigen" — See [cluster_leading_eigen](#) for more details
- "leiden" — See [cluster_leiden](#) for more details. *Note:* The Leiden algorithm will default to the modularity objective function (`objective_function = "modularity"`). Set `objective_function = "CPM"` to use the Constant Potts Model instead (see examples)
- "louvain" — See [cluster_louvain](#) for more details
- "optimal" — See [cluster_optimal](#) for more details
- "spinglass" — See [cluster_spinglass](#) for more details
- "walktrap" — See [cluster_walktrap](#) for more details

`allow.singleton`

Boolean (length = 1). Whether singleton or single node communities should be allowed. Defaults to FALSE. When FALSE, singleton communities will be set to missing (NA); otherwise, when TRUE, singleton communities will be allowed

`membership.only`

Boolean (length = 1). Whether the memberships only should be output. Defaults to TRUE. Set to FALSE to obtain all output for the community detection algorithm

...

Additional arguments to be passed on to `igraph`'s community detection functions (see `algorithm` for link to arguments of each algorithm)

Value

Returns memberships from a community detection algorithm

Author(s)

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

References

Csardi, G., & Nepusz, T. (2006). The `igraph` software package for complex network research. *InterJournal, Complex Systems*, 1695.

Examples

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

# Estimate network
network <- EBICglasso.qgraph(data = wmt)

# Compute Edge Betweenness
community.detection(network, algorithm = "edge_betweenness")

# Compute Fast Greedy
```

```
community.detection(network, algorithm = "fast_greedy")

# Compute Fluid
community.detection(
  network, algorithm = "fluid",
  no.of.communities = 2 # needs to be set
)

# Compute Infomap
community.detection(network, algorithm = "infomap")

# Compute Label Propagation
community.detection(network, algorithm = "label_prop")

# Compute Leading Eigenvector
community.detection(network, algorithm = "leading_eigen")

# Compute Leiden (with modularity)
community.detection(
  network, algorithm = "leiden",
  objective_function = "modularity"
)

# Compute Leiden (with CPM)
community.detection(
  network, algorithm = "leiden",
  objective_function = "CPM",
  resolution_parameter = 0.05 # "edge density"
)

# Compute Louvain
community.detection(network, algorithm = "louvain")

# Compute Optimal (identifies maximum modularity solution)
community.detection(network, algorithm = "optimal")

# Compute Spinglass
community.detection(network, algorithm = "spinglass")

# Compute Walktrap
community.detection(network, algorithm = "walktrap")

# Example with {igraph} network
community.detection(
  convert2igraph(network), algorithm = "walktrap"
)
```

Description

Memberships from community detection algorithms do not always align numerically. This function seeks to homogenize community memberships between a target membership (the membership to homogenize toward) and one or more other memberships. This function is the core of the [dimensionStability](#) and [itemStability](#) functions

Usage

```
community.homogenize(target.membership, convert.membership)
```

Arguments

target.membership

Vector, matrix, or data frame. The target memberships that all other memberships input into convert.membership should be homogenize **toward**

convert.membership

Vector, matrix, or data frame. Either a vector of memberships the same length as target.membership or a matrix or data frame of many membership solutions with either across rows or down columns the same length as target.membership (this function will automatically determine this orientation for you with precedence given solutions *across rows*)

Value

Returns a vector or matrix the length or size of convert.membership with memberships homogenized toward target.membership

Author(s)

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

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.

Examples

```
# Get network
network <- network. estimation(wmt2[,7:24])

# Apply Walktrap
network_walktrap <- community.detection(
  network, algorithm = "walktrap"
)

# Apply Louvain
network_louvain <- community.detection(
  network, algorithm = "louvain"
```

```
)
# Homogenize toward Walktrap
community.homogenize(network_walktrap, network_louvain)
```

```
community.unidimensional
```

Approaches to Detect Unidimensional Communities

Description

A function to apply several approaches to detect a unidimensional community in networks. There have many different approaches recently such as expanding the correlation matrix to have orthogonal correlations ("expand"), applying the Leading Eigenvalue community detection algorithm [cluster_leading_eigen](#) to the correlation matrix ("LE"), and applying the Louvain community detection algorithm [cluster_louvain](#) to the correlation matrix ("louvain"). Not necessarily intended for individual use – it's better to use [EGA](#)

Usage

```
community.unidimensional(
  data,
  n = NULL,
  corr = c("auto", "cor_auto", "pearson", "spearman"),
  na.data = c("pairwise", "listwise"),
  model = c("BGGM", "glasso", "TMFG"),
  uni.method = c("expand", "LE", "louvain"),
  verbose = FALSE,
  ...
)
```

Arguments

- | | |
|------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| data | Matrix or data frame. Should consist only of variables that are desired to be in analysis |
| n | Numeric (length = 1). Sample size if data provided is a correlation matrix |
| corr | Character (length = 1). Method to compute correlations. Defaults to "auto". Available options: <ul style="list-style-type: none"> • "auto" — Automatically computes appropriate correlations for the data using Pearson's for continuous, polychoric for ordinal, tetrachoric for binary, and polyserial/biserial for ordinal/binary with continuous. To change the number of categories that are considered ordinal, use <code>ordinal.categories</code> (see polychoric.matrix for more details) • "cor_auto" — Uses cor_auto to compute correlations. Arguments can be passed along to the function |

	<ul style="list-style-type: none"> • "pearson" — Pearson's correlation is computed for all variables regardless of categories • "spearman" — Spearman's rank-order correlation is computed for all variables regardless of categories
	For other similarity measures, compute them first and input them into data with the sample size (n)
na.data	Character (length = 1). How should missing data be handled? Defaults to "pairwise". Available options: <ul style="list-style-type: none"> • "pairwise" — Computes correlation for all available cases between two variables • "listwise" — Computes correlation for all complete cases in the dataset
model	Character (length = 1). Defaults to "glasso". Available options: <ul style="list-style-type: none"> • "BGM" — Computes the Bayesian Gaussian Graphical Model. Set argument <code>ordinal.categories</code> to determine levels allowed for a variable to be considered ordinal. See <code>?BGM::estimate</code> for more details • "glasso" — Computes the GLASSO with EBIC model selection. See EBICglasso.qgraph for more details • "TMFG" — Computes the TMFG method. See TMFG for more details
uni.method	Character (length = 1). What unidimensionality method should be used? Defaults to "louvain". Available options: <ul style="list-style-type: none"> • "expand" — Expands the correlation matrix with four variables correlated 0.50. 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 method was used in the Golino et al.'s (2020) <i>Psychological Methods</i> simulation • "LE" — Applies the Leading Eigenvector algorithm (cluster_leading_eigen) on the empirical correlation matrix. If the number of dimensions is 1, then the Leading Eigenvector solution is used; otherwise, regular EGA is used. This method was used in the Christensen et al.'s (2023) <i>Behavior Research Methods</i> simulation • "louvain" — Applies the Louvain algorithm (cluster_louvain) on the empirical correlation matrix. If the number of dimensions is 1, then the Louvain solution is used; otherwise, regular EGA is used. This method was validated Christensen's (2022) <i>PsyArXiv</i> simulation. Consensus clustering can be used by specifying either "consensus.method" or "consensus.iter"
verbose	Boolean. Whether messages and (insignificant) warnings should be output. Defaults to FALSE (silent calls). Set to TRUE to see all messages and warnings for every function call
...	Additional arguments to be passed on to auto.correlate , network.estimate , community.consensus , and community.detection

Value

Returns the memberships of the community detection algorithm. The memberships will output *regardless* of whether the network is unidimensional

Author(s)

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

References**Expand approach**

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.

Leading Eigenvector approach

Christensen, A. P., Garrido, L. E., Guerra-Pena, K., & Golino, H. (2023). Comparing community detection algorithms in psychometric networks: A Monte Carlo simulation. *Behavior Research Methods*.

Louvain approach

Christensen, A. P. (2023). Unidimensional community detection: A Monte Carlo simulation, grid search, and comparison. *PsyArXiv*.

Examples

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

# Louvain with Consensus Clustering (default)
community.unidimensional(wmt)

# Leading Eigenvector
community.unidimensional(wmt, uni.method = "LE")

# Expand
community.unidimensional(wmt, uni.method = "expand")
```

compare.EGA.plots

Visually Compare Two or More EGAnet plots

Description

Organizes EGA plots for comparison. Ensures that nodes are placed in the same layout to maximize comparison

Usage

```
compare.EGA.plots(
  ...,
  input.list = NULL,
  base = 1,
```

```

    same.layout = TRUE,
    labels = NULL,
    rows = NULL,
    columns = NULL,
    plot.all = TRUE
  )

```

Arguments

...	Handles multiple arguments: <ul style="list-style-type: none"> • *EGA objects — can be dropped in without any argument designation. The function will search across input to find necessary EGAnet objects • ggnet2 arguments — can be passed along to ggnet2 • gplot.layout — can be specified using mode = or layout = using the name of the layout (e.g., mode = "circle" will produce the circle layout from gplot.layout). By default, the layout is the same as qgraph
input.list	List. Bypasses ... argument in favor of using a list as an input
base	Numeric (length = 1). 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
same.layout	Boolean (length = 1). Whether the nodes should be in the same position for all networks. Defaults to TRUE. Set to FALSE for their individual layouts
labels	Character (same length as input). Labels for each EGAnet object
rows	Numeric (length = 1). Number of rows to spread plots across
columns	Numeric (length = 1). Number of columns to spread plots down
plot.all	Boolean (length = 1). Whether plot should be produced or just output. Defaults to TRUE. Set to FALSE to avoid plotting (but still obtain plot objects)

Value

Visual comparison of EGAnet objects

Author(s)

Alexander Christensen <alexpaulchristensen@gmail.com>

See Also

[plot.EGAnet](#) for plot usage in EGAnet

Examples

```

# Obtain WMT-2 data
wmt <- wmt2[,7:24]

# Draw random samples of 300 cases
sample1 <- wmt[sample(1:nrow(wmt), 300),]

```

```
sample2 <- wmt[sample(1:nrow(wmt), 300),]

# Estimate EGAs
ega1 <- EGA(sample1)
ega2 <- EGA(sample2)

# Compare EGAs via plot
compare.EGA.plots(
  ega1, ega2,
  base = 1, # use "ega1" as base for comparison
  labels = c("Sample 1", "Sample 2"),
  rows = 1, columns = 2
)

# Change layout to circle plots
compare.EGA.plots(
  ega1, ega2,
  labels = c("Sample 1", "Sample 2"),
  mode = "circle"
)
```

convert2igraph

Convert networks to igraph

Description

Converts networks to igraph format

Usage

```
convert2igraph(A, diagonal = 0)
```

Arguments

A	Matrix or data frame. $N \times N$ matrix where N is the number of nodes
diagonal	Numeric. Value to be placed on the diagonal of A. Defaults to 0

Value

Returns a network in the igraph format

Author(s)

Hudson Golino <hfg9s at virginia.edu> & Alexander P. Christensen <alexander.christensen at Vanderbilt.Edu>

Examples

```
convert2igraph(ega.wmt$network)
```

convert2tidygraph	<i>Convert networks to tidygraph</i>
-------------------	--------------------------------------

Description

Converts networks to tidygraph format

Usage

```
convert2tidygraph(EGA.object)
```

Arguments

EGA.object A single [EGAnet](#) object containing the outputs \$network and \$wc

Value

Returns a network in the tidygraph format

Author(s)

Dominique Makowski, Hudson Golino <hfg9s at virginia.edu>, & Alexander P. Christensen <alexander.christensen at Vanderbilt.Edu>

Examples

```
convert2tidygraph(ega.wmt)
```

cosine	<i>Cosine similarity</i>
--------	--------------------------

Description

Computes cosine similarity

Usage

```
cosine(x, y = NULL, ...)
```

Arguments

x	Numeric vector, matrix, or data frame. If $nrow(x) > 1$, then x will be treated as a matrix to compute an n by n similarity matrix (y will not be used!)
y	Numeric vector, matrix, or data frame. Only used if x is a single variable. Used to compute similarity between one variable and n other variables. Defaults to NULL
...	Not actually used but makes it easier for general functionality in the package

Details

On missing values: \emptyset will be used to replace missing values. When using (matrix) multiplication, the \emptyset value cancels out the product rendering the missing value as "not counting" in the sums

Author(s)

Alexander P. Christensen <alexpaulchristensen@gmail.com>

Examples

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

# Obtain cosines
wmt_cosine <- cosine(wmt)
```

depression

Depression Data

Description

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

Usage

```
data(depression)
```

Format

A 574x78 response matrix

Examples

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

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

Description

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

Usage

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

Arguments

bootega.obj	A bootEGA object
IS.plot	Boolean (length = 1). Should the plot be produced for item.replication? Defaults to TRUE
structure	Numeric (length = number of variables). A theoretical or pre-defined structure. Defaults to NULL or the empirical EGA result in the bootega.obj
...	Additional arguments. Used for deprecated arguments from previous versions of itemStability

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

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.

Conceptual introduction

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.

Examples

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

## Not run:
# Estimate bootstrap EGA
boot.wmt <- bootEGA(
  data = wmt, iter = 500,
  type = "parametric", ncores = 2
)
## End(Not run)

# Estimate stability statistics
dimensionStability(boot.wmt)
```

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`)

Usage

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

Format

A list of with a length of 4

Examples

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

dynamic.network.compare

Compares Dynamic Network Structures Using Permutation

Description

A permutation implementation to determine statistical significance of whether the dynamic network structures are different from one another

Usage

```
dynamic.network.compare(
  data,
  paired = FALSE,
  corr = c("auto", "cor_auto", "pearson", "spearman"),
  na.data = c("pairwise", "listwise"),
  model = c("BGGM", "glasso", "TMFG"),
  id = NULL,
  group = NULL,
  n.embed = 5,
  n.embed.optimize = FALSE,
  tau = 1,
  delta = 1,
  use.derivatives = 1,
  na.derivative = c("none", "kalman", "rowwise", "skipover"),
  zero.jitter = 0.001,
  iter = 1000,
  ncores,
  seed = NULL,
  verbose = TRUE,
  ...
)
```

Arguments

data	Matrix or data frame. Should consist only of variables to be used in the analysis as well as an ID column
paired	Boolean (length = 1). Whether groups are repeated measures representing paired samples. Defaults to FALSE
corr	Character (length = 1). Method to compute correlations. Defaults to "auto". Available options: <ul style="list-style-type: none"> "auto" — Automatically computes appropriate correlations for the data using Pearson's for continuous, polychoric for ordinal, tetrachoric for binary, and polyserial/biserial for ordinal/binary with continuous. To change the number of categories that are considered ordinal, use <code>ordinal.categories</code> (see polychoric.matrix for more details)

- "cor_auto" — Uses `cor_auto` to compute correlations. Arguments can be passed along to the function
- "cosine" — Uses `cosine` to compute cosine similarity
- "pearson" — Pearson's correlation is computed for all variables regardless of categories
- "spearman" — Spearman's rank-order correlation is computed for all variables regardless of categories

For other similarity measures, compute them first and input them into data with the sample size (n)

na.data	Character (length = 1). How should missing data be handled? Defaults to "pairwise". Available options: <ul style="list-style-type: none"> • "pairwise" — Computes correlation for all available cases between two variables • "listwise" — Computes correlation for all complete cases in the dataset
model	Character (length = 1). Defaults to "glasso". Available options: <ul style="list-style-type: none"> • "BGGM" — Computes the Bayesian Gaussian Graphical Model. Set argument <code>ordinal.categories</code> to determine levels allowed for a variable to be considered ordinal. See <code>?BGGM::estimate</code> for more details • "glasso" — Computes the GLASSO with EBIC model selection. See EBICglasso.qgraph for more details • "TMFG" — Computes the TMFG method. See TMFG for more details
id	Numeric or character (length = 1). Number or name of the column identifying each individual. Defaults to NULL
group	Numeric or character (length = 1). Number of the column identifying group membership. Defaults to NULL
n.embed	Numeric (length = 1). Defaults to 5. Number of embedded dimensions (the number of observations to be used in the <code>Embed</code> function). For example, an "n.embed = 5" will use five consecutive observations to estimate a single derivative
n.embed.optimize	Boolean (length = 1). If TRUE, performs optimization of n.embed for each individual, then constructs the population based on optimized derivatives. When TRUE, individual networks are considered of interest and will always be output. Defaults to FALSE
tau	Numeric (length = 1). Defaults to 1. Number of observations to offset successive embeddings in the <code>Embed</code> function. Generally recommended to leave "as is"
delta	Numeric (length = 1). Defaults to 1. The time between successive observations in the time series (i.e, lag). Generally recommended to leave "as is"
use.derivatives	Numeric (length = 1). Defaults to 1. The order of the derivative to be used in the analysis. Available options: <ul style="list-style-type: none"> • 0 — No derivatives; consistent with moving average • 1 — First-order derivatives; interpreted as "velocity" or rate of change over time

- 2 — Second-order derivatives; interpreted as "acceleration" or rate of the rate of change over time

Generally recommended to leave "as is"

na.derivative	Character (length = 1). How should missing data in the embeddings be handled? Available options (see Boker et al. (2018) in glla references for more details):
	<ul style="list-style-type: none"> • "none" (default) — does nothing and leaves NAs in data • "kalman" — uses Kalman smoothing (KalmanSmooth) with structural time series models (StructTS) to impute missing values. This approach models the underlying temporal dependencies (trend, seasonality, autocorrelation) to generate estimates for missing observations while preserving the original time scale. More computationally intensive than the other methods but typically provides the most accurate imputation by respecting the stochastic properties of the time series • "rowwise" — adjusts time interval with respect to each embedding ensuring time intervals are adaptive to the missing data (tends to be more accurate than "none") • "skipover" — "skips over" missing data and treats the non-missing points as continuous points in time (note that the time scale shifts to the "per mean time interval," which is different and <i>larger</i> than the original scale)
zero.jitter	Numeric (length = 1). Small amount of Gaussian noise added to zero variance derivatives to prevent estimation failures. For more than one variable, noise is generated multivariate normal distribution to ensure orthogonal noise is added. The jitter preserves the overall structure but avoids singular covariance matrices during network estimation. Defaults to 0.001
iter	Numeric (length = 1). Number of permutations to perform. Defaults to 1000 (recommended)
ncores	Numeric (length = 1). Number of cores to use in computing results. Defaults to <code>ceiling(parallel::detectCores() / 2)</code> or half of your computer's processing power. Set to 1 to not use parallel computing
seed	Numeric (length = 1). Defaults to NULL or random results. Set for reproducible results. See Reproducibility and PRNG for more details on random number generation in EGAnet
verbose	Boolean (length = 1). Should progress be displayed? Defaults to TRUE. Set to FALSE to not display progress
...	Additional arguments that can be passed on to auto.correlate , network.estimate , EGA , and jsd

Value

Returns a list:

network	Data frame with row names of each measure, empirical value (<code>statistic</code>), and p -value based on the permutation test (<code>p.value</code>)
edges	List containing matrices of values for empirical values (<code>statistic</code>), p -values (<code>p.value</code>), and Benjamini-Hochberg corrected p -values (<code>p.adjusted</code>)

Author(s)

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

References**sF**

Ulitzsch, E., Khanna, S., Rhemtulla, M., & Domingue, B. W. (2023). A graph theory based similarity metric enables comparison of subpopulation psychometric networks. *Psychological Methods*.

Jensen-Shannon Similarity (1 - Distance)

De Domenico, M., Nicosia, V., Arenas, A., & Latora, V. (2015). Structural reducibility of multilayer networks. *Nature Communications*, 6(1), 1–9.

Total Network Strength

van Borkulo, C. D., van Bork, R., Boschloo, L., Kossakowski, J. J., Tio, P., Schoevers, R. A., Borsboom, D., & Waldorp, L. J. (2023). Comparing network structures on three aspects: A permutation test. *Psychological Methods*, 28(6), 1273–1285.

Examples

```
# Three similar groups

# Set seed
set.seed(42)

# Simulate dynamic data
participants <- lapply(
  seq_len(50), function(i){

    # Get output
    output <- simDFM(
      variab = 6, timep = 15,
      nfact = 2, error = 0.100,
      dfm = "DAFS", loadings = 0.60,
      autoreg = 0.80, crossreg = 0.10,
      var.shock = 0.36, cov.shock = 0.18,
      burnin = 2000
    )

    # Add ID
    df <- data.frame(
      ID = i,
      Group = rep(1:3, each = 5),
      output$data
    )

    # Return data
    return(df)
  }
)
```

```

# Put participants into a data frame
df <- do.call(rbind.data.frame, participants)

## Not run:
# Perform comparison
dynamic.network.compare(
  data = df, paired = TRUE,
  # EGA arguments
  corr = "auto", na.data = "pairwise", model = "glasso",
  # dynEGA arguments
  id = "ID", group = "Group", n.embed = 3,
  tau = 1, delta = 1, use.derivatives = 1,
  # Permutation arguments
  iter = 1000, ncores = 2, verbose = TRUE, seed = 42
)
## End(Not run)

# Two similar groups and one different

# Simulate dynamic data
participants <- lapply(
  seq_len(50), function(i){

  # Get output
  output <- simDFM(
    variab = 4, timep = 5,
    nfact = 3, error = 0.100,
    dfm = "DAFS", loadings = 0.60,
    autoreg = 0.80, crossreg = 0.10,
    var.shock = 0.36, cov.shock = 0.18,
    burnin = 2000
  )

  # Add ID
  df <- data.frame(
    ID = i,
    Group = rep(3, each = 5),
    output$data
  )

  # Return data
  return(df)

  }
)

# Replace group 3
new_group <- do.call(rbind.data.frame, participants)
df[df$Group == 3,] <- new_group

## Not run:
# Perform comparison
dynamic.network.compare(

```

```

data = df, paired = TRUE,
# EGA arguments
corr = "auto", na.data = "pairwise", model = "glasso",
# dynEGA arguments
id = "ID", group = "Group", n.embed = 3,
tau = 1, delta = 1, use.derivatives = 1,
# Permutation arguments
iter = 1000, ncores = 2, verbose = TRUE, seed = 42
)
## End(Not run)

```

dynEGA

Dynamic Exploratory Graph Analysis

Description

Estimates dynamic communities in multivariate time series (e.g., panel data, longitudinal data, intensive longitudinal data) at multiple time scales and at different levels of analysis: individuals (intraindividual structure), groups, and population (interindividual structure)

Usage

```

dynEGA(
  data,
  id = NULL,
  group = NULL,
  n.embed = 5,
  n.embed.optimize = FALSE,
  tau = 1,
  delta = 1,
  use.derivatives = 1,
  na.derivative = c("none", "kalman", "rowwise", "skipover"),
  zero.jitter = 0.001,
  level = c("individual", "group", "population"),
  corr = c("auto", "cor_auto", "pearson", "spearman"),
  na.data = c("pairwise", "listwise"),
  model = c("BGGM", "glasso", "TMFG"),
  algorithm = c("leiden", "louvain", "walktrap"),
  uni.method = c("expand", "LE", "louvain"),
  ncores,
  seed = NULL,
  verbose = TRUE,
  ...
)

```

Arguments

data	<p>Matrix or data frame. Participants and variable should be in long format such that row t represents observations for all variables at time point t for a participant. The next row, $t + 1$, represents the next measurement occasion for that same participant. The next participant's data should immediately follow, in the same pattern, after the previous participant</p> <p>data should have an ID variable labeled "ID"; otherwise, it is assumed that the data represent the population</p> <p>For groups, data should have a Group variable labeled "Group"; otherwise, it is assumed that there are no groups in data</p> <p>Arguments <code>id</code> and <code>group</code> can be specified to tell the function which column in data it should use as the ID and Group variable, respectively</p> <p>A measurement occasion variable is not necessary and should be <i>removed</i> from the data before proceeding with the analysis</p>
id	Numeric or character (length = 1). Number or name of the column identifying each individual. Defaults to NULL
group	Numeric or character (length = 1). Number of the column identifying group membership. Defaults to NULL
n.embed	<p>Numeric (length = 1 or more). Defaults to 5. Number of embedded dimensions (the number of observations to be used in the <code>Embed</code> function). For example, an "n.embed = 5" will use five consecutive observations to estimate a single derivative.</p> <p>If more than one value is provided, then the number of embeddings will be optimized over using <code>tefi</code> to determine the optimal length of the embedding dimensions for <i>each</i> individual in the sample</p>
n.embed.optimize	<p>Boolean (length = 1). If TRUE, performs optimization of n.embed for each individual, then constructs the population based on optimized derivatives. When TRUE, individual networks are considered of interest and will always be output. Defaults to FALSE</p>
tau	Numeric (length = 1). Defaults to 1. Number of observations to offset successive embeddings in the <code>Embed</code> function. Generally recommended to leave "as is"
delta	Numeric (length = 1). Defaults to 1. The time between successive observations in the time series (i.e, lag). Generally recommended to leave "as is"
use.derivatives	<p>Numeric (length = 1). Defaults to 1. The order of the derivative to be used in the analysis. Available options:</p> <ul style="list-style-type: none"> • 0 — No derivatives; consistent with moving average • 1 — First-order derivatives; interpreted as "velocity" or rate of change over time • 2 — Second-order derivatives; interpreted as "acceleration" or rate of the rate of change over time <p>Generally recommended to leave "as is"</p>
na.derivative	<p>Character (length = 1). How should missing data in the embeddings be handled? Available options (see Boker et al. (2018) in glla references for more details):</p>

	<ul style="list-style-type: none"> • "none" (default) — does nothing and leaves NAs in data • "kalman" — uses Kalman smoothing (KalmanSmooth) with structural time series models (StructTS) to impute missing values. This approach models the underlying temporal dependencies (trend, seasonality, autocorrelation) to generate estimates for missing observations while preserving the original time scale. More computationally intensive than the other methods but typically provides the most accurate imputation by respecting the stochastic properties of the time series • "rowwise" — adjusts time interval with respect to each embedding ensuring time intervals are adaptive to the missing data (tends to be more accurate than "none") • "skipover" — "skips over" missing data and treats the non-missing points as continuous points in time (note that the time scale shifts to the "per mean time interval," which is different and <i>larger</i> than the original scale)
zero.jitter	Numeric (length = 1). Small amount of Gaussian noise added to zero variance derivatives to prevent estimation failures. For more than one variable, noise is generated multivariate normal distribution to ensure orthogonal noise is added. The jitter preserves the overall structure but avoids singular covariance matrices during network estimation. Defaults to 0.001
level	Character vector (up to length of 3). A character vector indicating which level(s) to estimate: <ul style="list-style-type: none"> • "individual" — Estimates EGA for each individual in data (intraindividual structure; requires an "ID" column, see data) • "group" — Estimates EGA for each group in data (group structure; requires a "Group" column, see data) • "population" — Estimates EGA across all data (interindividual structure)
corr	Character (length = 1). Method to compute correlations. Defaults to "auto". Available options: <ul style="list-style-type: none"> • "auto" — Automatically computes appropriate correlations for the data using Pearson's for continuous, polychoric for ordinal, tetrachoric for binary, and polyserial/biserial for ordinal/binary with continuous. To change the number of categories that are considered ordinal, use <code>ordinal.categories</code> (see polychoric.matrix for more details) • "cor_auto" — Uses <code>cor_auto</code> to compute correlations. Arguments can be passed along to the function • "pearson" — Pearson's correlation is computed for all variables regardless of categories • "spearman" — Spearman's rank-order correlation is computed for all variables regardless of categories <p>For other similarity measures, compute them first and input them into data with the sample size (n)</p>
na.data	Character (length = 1). How should missing data be handled? Defaults to "pairwise". Available options: <ul style="list-style-type: none"> • "pairwise" — Computes correlation for all available cases between two variables

	<ul style="list-style-type: none"> • "listwise" — Computes correlation for all complete cases in the dataset
model	<p>Character (length = 1). Defaults to "glasso". Available options:</p> <ul style="list-style-type: none"> • "BGGM" — Computes the Bayesian Gaussian Graphical Model. Set argument <code>ordinal.categories</code> to determine levels allowed for a variable to be considered ordinal. See <code>?BGGM::estimate</code> for more details • "glasso" — Computes the GLASSO with EBIC model selection. See EBICglasso.qgraph for more details • "TMFG" — Computes the TMFG method. See TMFG for more details
algorithm	<p>Character or <code>igraph cluster_*</code> function (length = 1). Defaults to "walktrap". Three options are listed below but all are available (see community.detection for other options):</p> <ul style="list-style-type: none"> • "leiden" — See cluster_leiden for more details • "louvain" — By default, "louvain" will implement the Louvain algorithm using the consensus clustering method (see community.consensus for more information). This function will implement <code>consensus.method = "most_common"</code> and <code>consensus.iter = 1000</code> unless specified otherwise • "walktrap" — See cluster_walktrap for more details
uni.method	<p>Character (length = 1). What unidimensionality method should be used? Defaults to "louvain". Available options:</p> <ul style="list-style-type: none"> • "expand" — Expands the correlation matrix with four variables correlated 0.50. 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 method was used in the Golino et al.'s (2020) <i>Psychological Methods</i> simulation • "LE" — Applies the Leading Eigenvector algorithm (cluster_leading_eigen) on the empirical correlation matrix. If the number of dimensions is 1, then the Leading Eigenvector solution is used; otherwise, regular EGA is used. This method was used in the Christensen et al.'s (2023) <i>Behavior Research Methods</i> simulation • "louvain" — Applies the Louvain algorithm (cluster_louvain) on the empirical correlation matrix. If the number of dimensions is 1, then the Louvain solution is used; otherwise, regular EGA is used. This method was validated Christensen's (2022) <i>PsyArXiv</i> simulation. Consensus clustering can be used by specifying either <code>"consensus.method"</code> or <code>"consensus.iter"</code>
ncores	<p>Numeric (length = 1). Number of cores to use in computing results. Defaults to <code>ceiling(parallel::detectCores() / 2)</code> or half of your computer's processing power. Set to 1 to not use parallel computing</p> <p>If you're unsure how many cores your computer has, then type: <code>parallel::detectCores()</code></p>
seed	<p>Numeric (length = 1). Defaults to NULL or random results. Set for reproducible results. See Reproducibility and PRNG for more details on random number generation in EGAnet</p>
verbose	<p>Boolean (length = 1). Should progress be displayed? Defaults to TRUE. Set to FALSE to not display progress</p>
...	<p>Additional arguments to be passed on to auto.correlate, network.estimate, community.detection, community.consensus, and EGA</p>

Details

Derivatives for each variable's time series for each participant are estimated using generalized local linear approximation (see [glla](#)). EGA is then applied to these derivatives to model how variables are changing together over time. Variables that change together over time are detected as communities

Value

A list containing:

Derivatives A list containing:

- Estimates — A list the length of the unique IDs containing data frames of zero- to second-order derivatives for each ID in data
- EstimatesDF — A data frame of derivatives across all IDs containing columns of the zero- to second-order derivatives as well as id and group variables (group is automatically set to 1 for all if no group is provided)

dynEGA A list containing:

- population — If level includes "populaton", then the EGA results for the entire sample
- group — If level includes "group", then a list containing the EGA results for each group
- individual — If level includes "individual", then a list containing the EGA results for each id

Author(s)

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

References

Generalized local linear approximation

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.

Original dynamic EGA implementation

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*.

Time delay embedding procedure

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

See Also

[plot.EGAnet](#) for plot usage in EGAnet

Examples

```
# Population structure
simulated_population <- dynEGA(
  data = sim.dynEGA, level = "population"
  # uses simulated data in package
  # useful to understand how data should be structured
)

## Not run:
# Group structure
simulated_group <- dynEGA(
  data = sim.dynEGA, level = "group"
  # uses simulated data in package
  # useful to understand how data should be structured
)

# Individual structure
simulated_individual <- dynEGA(
  data = sim.dynEGA, level = "individual",
  ncores = 2, # use more for quicker results
  verbose = TRUE # progress bar
)

# Population, group, and individual structure
simulated_all <- dynEGA(
  data = sim.dynEGA,
  level = c("individual", "group", "population"),
  ncores = 2, # use more for quicker results
  verbose = TRUE # progress bar
)

# Plot population
plot(simulated_all$dynEGA$population)

# Plot groups
plot(simulated_all$dynEGA$group)

# Plot individual
plot(simulated_all$dynEGA$individual, id = 1)

# Step through all plots
# Unless `id` is specified, 4 random IDs
# will be drawn from individuals
plot(simulated_all)

# Optimize over multiple embeddings
optimized_all <- dynEGA(
  data = sim.dynEGA,
```

```

level = c("individual", "group", "population"),
n.embed = 3:10, # set number of dimensions to search over
n.embed.optimize = TRUE, # set to TRUE to optimize
ncores = 2, # use more for quicker results
verbose = TRUE # progress bar
)
## End(Not run)

```

dynEGA.ind.pop

Intra- and Inter-individual [dynEGA](#)

Description

A wrapper function to estimate both intraindividual (level = "individual") and interindividual (level = "population") structures using [dynEGA](#)

Usage

```

dynEGA.ind.pop(
  data,
  id = NULL,
  n.embed = 5,
  n.embed.optimize = FALSE,
  tau = 1,
  delta = 1,
  use.derivatives = 1,
  na.derivative = c("none", "kalman", "rowwise", "skipover"),
  zero.jitter = 0.001,
  corr = c("auto", "cor_auto", "pearson", "spearman"),
  na.data = c("pairwise", "listwise"),
  model = c("BGGM", "glasso", "TMFG"),
  algorithm = c("leiden", "louvain", "walktrap"),
  uni.method = c("expand", "LE", "louvain"),
  ncores,
  seed = NULL,
  verbose = TRUE,
  ...
)

```

Arguments

data Matrix or data frame. Participants and variable should be in long format such that row t represents observations for all variables at time point t for a participant. The next row, $t + 1$, represents the next measurement occasion for that same participant. The next participant's data should immediately follow, in the same pattern, after the previous participant

data should have an ID variable labeled "ID"; otherwise, it is assumed that the data represent the population

For groups, data should have a Group variable labeled "Group"; otherwise, it is assumed that there are no groups in data

Arguments `id` and `group` can be specified to tell the function which column in data it should use as the ID and Group variable, respectively

A measurement occasion variable is not necessary and should be *removed* from the data before proceeding with the analysis

<code>id</code>	Numeric or character (length = 1). Number or name of the column identifying each individual. Defaults to NULL
<code>n.embed</code>	Numeric (length = 1 or more). Defaults to 5. Number of embedded dimensions (the number of observations to be used in the <code>Embed</code> function). For example, an <code>"n.embed = 5"</code> will use five consecutive observations to estimate a single derivative. If more than one value is provided, then the number of embeddings will be optimized over using <code>tefi</code> to determine the optimal length of the embedding dimensions for <i>each</i> individual in the sample
<code>n.embed.optimize</code>	Boolean (length = 1). If TRUE, performs optimization of <code>n.embed</code> for each individual, then constructs the population based on optimized derivatives. When TRUE, individual networks are considered of interest and will always be output. Defaults to FALSE
<code>tau</code>	Numeric (length = 1). Defaults to 1. Number of observations to offset successive embeddings in the <code>Embed</code> function. Generally recommended to leave "as is"
<code>delta</code>	Numeric (length = 1). Defaults to 1. The time between successive observations in the time series (i.e, lag). Generally recommended to leave "as is"
<code>use.derivatives</code>	Numeric (length = 1). Defaults to 1. The order of the derivative to be used in the analysis. Available options: <ul style="list-style-type: none"> • 0 — No derivatives; consistent with moving average • 1 — First-order derivatives; interpreted as "velocity" or rate of change over time • 2 — Second-order derivatives; interpreted as "acceleration" or rate of the rate of change over time Generally recommended to leave "as is"
<code>na.derivative</code>	Character (length = 1). How should missing data in the embeddings be handled? Available options (see Boker et al. (2018) in <code>glla</code> references for more details): <ul style="list-style-type: none"> • "none" (default) — does nothing and leaves NAs in data • "kalman" — uses Kalman smoothing (<code>KalmanSmooth</code>) with structural time series models (<code>StructTS</code>) to impute missing values. This approach models the underlying temporal dependencies (trend, seasonality, autocorrelation) to generate estimates for missing observations while preserving the original time scale. More computationally intensive than the other methods but typically provides the most accurate imputation by respecting the stochastic properties of the time series

	<ul style="list-style-type: none"> • "rowwise" — adjusts time interval with respect to each embedding ensuring time intervals are adaptive to the missing data (tends to be more accurate than "none") • "skipover" — "skips over" missing data and treats the non-missing points as continuous points in time (note that the time scale shifts to the "per mean time interval," which is different and <i>larger</i> than the original scale)
zero.jitter	Numeric (length = 1). Small amount of Gaussian noise added to zero variance derivatives to prevent estimation failures. For more than one variable, noise is generated multivariate normal distribution to ensure orthogonal noise is added. The jitter preserves the overall structure but avoids singular covariance matrices during network estimation. Defaults to 0.001
corr	<p>Character (length = 1). Method to compute correlations. Defaults to "auto". Available options:</p> <ul style="list-style-type: none"> • "auto" — Automatically computes appropriate correlations for the data using Pearson's for continuous, polychoric for ordinal, tetrachoric for binary, and polyserial/biserial for ordinal/binary with continuous. To change the number of categories that are considered ordinal, use <code>ordinal.categories</code> (see polychoric.matrix for more details) • "cor_auto" — Uses <code>cor_auto</code> to compute correlations. Arguments can be passed along to the function • "pearson" — Pearson's correlation is computed for all variables regardless of categories • "spearman" — Spearman's rank-order correlation is computed for all variables regardless of categories <p>For other similarity measures, compute them first and input them into data with the sample size (n)</p>
na.data	<p>Character (length = 1). How should missing data be handled? Defaults to "pairwise". Available options:</p> <ul style="list-style-type: none"> • "pairwise" — Computes correlation for all available cases between two variables • "listwise" — Computes correlation for all complete cases in the dataset
model	<p>Character (length = 1). Defaults to "glasso". Available options:</p> <ul style="list-style-type: none"> • "BGGM" — Computes the Bayesian Gaussian Graphical Model. Set argument <code>ordinal.categories</code> to determine levels allowed for a variable to be considered ordinal. See <code>?BGGM:estimate</code> for more details • "glasso" — Computes the GLASSO with EBIC model selection. See EBICglasso.qgraph for more details • "TMFG" — Computes the TMFG method. See TMFG for more details
algorithm	<p>Character or <code>igraph_cluster_*</code> function (length = 1). Defaults to "walktrap". Three options are listed below but all are available (see community.detection for other options):</p> <ul style="list-style-type: none"> • "leiden" — See cluster_leiden for more details

	<ul style="list-style-type: none"> • "louvain" — By default, "louvain" will implement the Louvain algorithm using the consensus clustering method (see community.consensus for more information). This function will implement <code>consensus.method = "most_common"</code> and <code>consensus.iter = 1000</code> unless specified otherwise • "walktrap" — See cluster_walktrap for more details
<code>uni.method</code>	<p>Character (length = 1). What unidimensionality method should be used? Defaults to "louvain". Available options:</p> <ul style="list-style-type: none"> • "expand" — Expands the correlation matrix with four variables correlated 0.50. 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 method was used in the Golino et al.'s (2020) <i>Psychological Methods</i> simulation • "LE" — Applies the Leading Eigenvector algorithm (cluster_leading_eigen) on the empirical correlation matrix. If the number of dimensions is 1, then the Leading Eigenvector solution is used; otherwise, regular EGA is used. This method was used in the Christensen et al.'s (2023) <i>Behavior Research Methods</i> simulation • "louvain" — Applies the Louvain algorithm (cluster_louvain) on the empirical correlation matrix. If the number of dimensions is 1, then the Louvain solution is used; otherwise, regular EGA is used. This method was validated Christensen's (2022) <i>PsyArXiv</i> simulation. Consensus clustering can be used by specifying either "consensus.method" or "consensus.iter"
<code>ncores</code>	<p>Numeric (length = 1). Number of cores to use in computing results. Defaults to <code>ceiling(parallel::detectCores() / 2)</code> or half of your computer's processing power. Set to 1 to not use parallel computing</p> <p>If you're unsure how many cores your computer has, then type: <code>parallel::detectCores()</code></p>
<code>seed</code>	<p>Numeric (length = 1). Defaults to NULL or random results. Set for reproducible results. See Reproducibility and PRNG for more details on random number generation in EGAnet</p>
<code>verbose</code>	<p>Boolean (length = 1). Should progress be displayed? Defaults to TRUE. Set to FALSE to not display progress</p>
<code>...</code>	<p>Additional arguments to be passed on to auto.correlate, network.estimate, community.detection, community.consensus, and EGA</p>

Value

Same output as `EGAnet{dynEGA}` returning list objects for `level = "individual"` and `level = "population"`

Author(s)

Hudson Golino <hfg9s at virginia.edu>

See Also

[plot.EGAnet](#) for plot usage in EGAnet

Examples

```

# Obtain data
sim.dynEGA <- sim.dynEGA # bypasses CRAN checks

## Not run:
# Dynamic EGA individual and population structure
dyn.ega1 <- dynEGA.ind.pop(
  data = sim.dynEGA, n.embed = 5, tau = 1,
  delta = 1, id = 25, use.derivatives = 1,
  ncores = 2, corr = "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 criterion (EBIC) described by Foygel & Drton (2010).

Usage

```

EBICglasso.qgraph(
  data,
  n = NULL,
  corr = c("auto", "cor_auto", "cosine", "pearson", "spearman"),
  na.data = c("pairwise", "listwise"),
  gamma = 0.5,
  penalize.diagonal = FALSE,
  nlambda = 100,
  lambda.min.ratio = 0.1,
  fast = FALSE,
  returnAllResults = FALSE,
  penalizeMatrix = NULL,
  countDiagonal = FALSE,
  refit = FALSE,
  model.selection = c("EBIC", "JSD"),
  verbose = FALSE,
  ...
)

```

Arguments

data	Matrix or data frame. Should consist only of variables to be used in the analysis
n	Numeric (length = 1). Sample size if data provided is a correlation matrix
corr	Character (length = 1). Method to compute correlations. Defaults to "auto". Available options: <ul style="list-style-type: none"> • "auto" — Automatically computes appropriate correlations for the data using Pearson's for continuous, polychoric for ordinal, tetrachoric for binary, and polyserial/biserial for ordinal/binary with continuous. To change the number of categories that are considered ordinal, use <code>ordinal.categories</code> (see polychoric.matrix for more details) • "cor_auto" — Uses <code>cor_auto</code> to compute correlations. Arguments can be passed along to the function • "cosine" — Uses <code>cosine</code> to compute cosine similarity • "pearson" — Pearson's correlation is computed for all variables regardless of categories • "spearman" — Spearman's rank-order correlation is computed for all variables regardless of categories <p>For other similarity measures, compute them first and input them into data with the sample size (n)</p>
na.data	Character (length = 1). How should missing data be handled? Defaults to "pairwise". Available options: <ul style="list-style-type: none"> • "pairwise" — Computes correlation for all available cases between two variables • "listwise" — Computes correlation for all complete cases in the dataset
gamma	Numeric (length = 1) EBIC tuning parameter. Defaults to 0.50 and is generally a good choice. Setting to 0 will cause regular BIC to be used
penalize.diagonal	Boolean (length = 1). Should the diagonal be penalized? Defaults to FALSE
nlambda	Numeric (length = 1). Number of lambda values to test. Defaults to 100
lambda.min.ratio	Numeric (length = 1). Ratio of lowest lambda value compared to maximal lambda. Defaults to 0.1. NOTE qgraph sets the default to 0.01
fast	Boolean (length = 1). Whether the <code>glassoFast</code> version should be used to estimate the GLASSO. Defaults to FALSE. The fast results <i>may</i> differ by less than floating point of the original GLASSO implemented by <code>glasso</code> and should not impact reproducibility much
returnAllResults	Boolean (length = 1). Whether all results should be returned. Defaults to FALSE (network only). Set to TRUE to access <code>glassopath</code> output
penalizeMatrix	Boolean matrix. Optional logical matrix to indicate which elements are penalized
countDiagonal	Boolean (length = 1). Should diagonal be counted in EBIC computation? Defaults to FALSE. Set to TRUE to mimic qgraph < 1.3 behavior (not recommended!)

<code>refit</code>	Boolean (length = 1). Should the optimal graph be refitted without LASSO regularization? Defaults to FALSE
<code>model.selection</code>	Character (length = 1). How lambda should be selected within GLASSO. Defaults to "EBIC". "JSD" is experimental and should not be used otherwise
<code>verbose</code>	Boolean (length = 1). Whether messages and (insignificant) warnings should be output. Defaults to FALSE (silent calls). Set to TRUE to see all messages and warnings for every function call
<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; for maintenance, Hudson Golino <hfg9s at virginia.edu> and Alexander P. Christensen <alexpaulchristensen at gmail.com>

References

Instantiation of GLASSO

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

glasso + EBIC

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

glasso package

Friedman, J., Hastie, T., & Tibshirani, R. (2011). `glasso`: Graphical lasso-estimation of Gaussian graphical models. R package version 1.7.

Tutorial on EBICglasso

Epskamp, S., & Fried, E. I. (2018). A tutorial on regularized partial correlation networks. *Psychological Methods*, 23(4), 617-634.

Examples

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

# Fast
fast <- EBICglasso.qgraph(wmt)
```

```

# Regular
regular <- EBICglasso.qgraph(wmt, fast = FALSE)

# Difference between fast and regular
sum(abs(fast - regular))

# Compute graph with tuning = 0 (BIC)
BICgraph <- EBICglasso.qgraph(data = wmt, gamma = 0)

# Compute graph with tuning = 0.5 (EBIC)
EBICgraph <- EBICglasso.qgraph(data = wmt, gamma = 0.5)

```

 EGA

Exploratory Graph Analysis

Description

Estimates the number of communities (dimensions) of a dataset or correlation matrix using a network estimation method (Golino & Epskamp, 2017; Golino et al., 2020). After, a community detection algorithm is applied (Christensen et al., 2023) for multidimensional data. A unidimensional check is also applied based on findings from Golino et al. (2020) and Christensen (2023)

Usage

```

EGA(
  data,
  n = NULL,
  corr = c("auto", "cor_auto", "cosine", "pearson", "spearman"),
  na.data = c("pairwise", "listwise"),
  model = c("BGGM", "glasso", "TMFG"),
  algorithm = c("leiden", "louvain", "walktrap"),
  uni.method = c("expand", "LE", "louvain"),
  plot.EGA = TRUE,
  verbose = FALSE,
  ...
)

```

Arguments

data	Matrix or data frame. Should consist only of variables to be used in the analysis. Can be raw data or a correlation matrix
n	Numeric (length = 1). Sample size if data provided is a correlation matrix
corr	Character (length = 1). Method to compute correlations. Defaults to "auto". Available options:

- "auto" — Automatically computes appropriate correlations for the data using Pearson's for continuous, polychoric for ordinal, tetrachoric for binary, and polyserial/biserial for ordinal/binary with continuous. To change the number of categories that are considered ordinal, use `ordinal.categories` (see `polychoric.matrix` for more details)
- "cor_auto" — Uses `cor_auto` to compute correlations. Arguments can be passed along to the function
- "cosine" — Uses `cosine` to compute cosine similarity
- "pearson" — Pearson's correlation is computed for all variables regardless of categories
- "spearman" — Spearman's rank-order correlation is computed for all variables regardless of categories

For other similarity measures, compute them first and input them into data with the sample size (n)

<code>na.data</code>	Character (length = 1). How should missing data be handled? Defaults to "pairwise". Available options: <ul style="list-style-type: none"> • "pairwise" — Computes correlation for all available cases between two variables • "listwise" — Computes correlation for all complete cases in the dataset
<code>model</code>	Character (length = 1). Defaults to "glasso". Available options: <ul style="list-style-type: none"> • "BGGM" — Computes the Bayesian Gaussian Graphical Model. Set argument <code>ordinal.categories</code> to determine levels allowed for a variable to be considered ordinal. See <code>?BGGM::estimate</code> for more details • "glasso" — Computes the GLASSO with EBIC model selection. See <code>EBICglasso.qgraph</code> for more details • "TMFG" — Computes the TMFG method. See <code>TMFG</code> for more details
<code>algorithm</code>	Character or <code>igraph_cluster_*</code> function (length = 1). Defaults to "walktrap". Three options are listed below but all are available (see <code>community.detection</code> for other options): <ul style="list-style-type: none"> • "leiden" — See <code>cluster_leiden</code> for more details • "louvain" — By default, "louvain" will implement the Louvain algorithm using the consensus clustering method (see <code>community.consensus</code> for more information). This function will implement <code>consensus.method = "most_common"</code> and <code>consensus.iter = 1000</code> unless specified otherwise • "walktrap" — See <code>cluster_walktrap</code> for more details
<code>uni.method</code>	Character (length = 1). What unidimensionality method should be used? Defaults to "louvain". Available options: <ul style="list-style-type: none"> • "expand" — Expands the correlation matrix with four variables correlated 0.50. 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 method was used in the Golino et al.'s (2020) <i>Psychological Methods</i> simulation

- "LE" — Applies the Leading Eigenvector algorithm ([cluster_leading_eigen](#)) on the empirical correlation matrix. If the number of dimensions is 1, then the Leading Eigenvector solution is used; otherwise, regular EGA is used. This method was used in the Christensen et al.'s (2023) *Behavior Research Methods* simulation
- "louvain" — Applies the Louvain algorithm ([cluster_louvain](#)) on the empirical correlation matrix. If the number of dimensions is 1, then the Louvain solution is used; otherwise, regular EGA is used. This method was validated Christensen's (2022) *PsyArXiv* simulation. Consensus clustering can be used by specifying either "consensus.method" or "consensus.iter"

plot.EGA	Boolean (length = 1). Defaults to TRUE. Whether the plot should be returned with the results. Set to FALSE for no plot
verbose	Boolean (length = 1). Whether messages and (insignificant) warnings should be output. Defaults to FALSE (silent calls). Set to TRUE to see all messages and warnings for every function call
...	Additional arguments to be passed on to auto.correlate , network.estimate , community.detection , community.consensus , and community.unidimensional

Value

Returns a list containing:

network	A matrix containing a network estimated using <code>link[EGAnet]{network.estimate}</code>
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
correlation	The zero-order correlation matrix
n	Number of cases in data
dim.variables	An ordered matrix of item allocation
TEFI	<code>link[EGAnet]{tefi}</code> for the estimated structure
plot.EGA	Plot output if <code>plot.EGA = TRUE</code>

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

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.

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., & Thiagarajan, 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.

Compared all *igraph* community detection algorithms, introduced Louvain algorithm, simulation with continuous and polytomous data

Also implements the Leading Eigenvalue unidimensional method

Christensen, A. P., Garrido, L. E., Pena, K. G., & Golino, H. (2023). Comparing community detection algorithms in psychological data: A Monte Carlo simulation. *Behavior Research Methods*.

Comprehensive unidimensionality simulation

Christensen, A. P. (2023). Unidimensional community detection: A Monte Carlo simulation, grid search, and comparison. *PsyArXiv*.

Compared all *igraph* community detection algorithms, simulation with continuous and polytomous data

Christensen, A. P., Garrido, L. E., Guerra-Pena, K., & Golino, H. (2023). Comparing community detection algorithms in psychometric networks: A Monte Carlo simulation. *Behavior Research Methods*.

See Also

[plot.EGAnet](#) for plot usage in EGAnet

Examples

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

# Estimate EGA
ega.wmt <- EGA(
  data = wmt,
  plot.EGA = FALSE # No plot for CRAN checks
)

# Print results
print(ega.wmt)

# Plot with colorblind friendly edges
print(ega.wmt, edge.color = c("#0B64AD", "red"))

# Estimate EGAtmfg
ega.wmt.tmfg <- EGA(
  data = wmt, model = "TMFG",
  plot.EGA = FALSE # No plot for CRAN checks
)

# Estimate EGA with Louvain algorithm
ega.wmt.louvain <- EGA(
  data = wmt, algorithm = "louvain",
  plot.EGA = FALSE # No plot for CRAN checks
)

# Estimate EGA with an {igraph} function (Fast-greedy)
```

```
ega.wmt.greedy <- EGA(
  data = wmt,
  algorithm = igraph::cluster_fast_greedy,
  plot.EGA = FALSE # No plot for CRAN checks
)
```

 EGA.estimate

Estimates EGA for Multidimensional Structures

Description

A basic function to estimate [EGA](#) for multidimensional structures. This function does *not* include the unidimensional check and it does not plot the results. This function can be used as a streamlined approach for quick [EGA](#) estimation when unidimensionality or visualization is not a priority

Usage

```
EGA.estimate(
  data,
  n = NULL,
  corr = c("auto", "cor_auto", "cosine", "pearson", "spearman"),
  na.data = c("pairwise", "listwise"),
  model = c("BGGM", "glasso", "TMFG"),
  algorithm = c("leiden", "louvain", "walktrap"),
  verbose = FALSE,
  ...
)
```

Arguments

- | | |
|------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| data | Matrix or data frame. Should consist only of variables to be used in the analysis |
| n | Numeric (length = 1). Sample size if data provided is a correlation matrix |
| corr | Character (length = 1). Method to compute correlations. Defaults to "auto". Available options: <ul style="list-style-type: none"> "auto" — Automatically computes appropriate correlations for the data using Pearson's for continuous, polychoric for ordinal, tetrachoric for binary, and polyserial/biserial for ordinal/binary with continuous. To change the number of categories that are considered ordinal, use <code>ordinal.categories</code> (see polychoric.matrix for more details) "cor_auto" — Uses cor_auto to compute correlations. Arguments can be passed along to the function "cosine" — Uses cosine to compute cosine similarity "pearson" — Pearson's correlation is computed for all variables regardless of categories |

- "spearman" — Spearman's rank-order correlation is computed for all variables regardless of categories

For other similarity measures, compute them first and input them into data with the sample size (n)

na.data	Character (length = 1). How should missing data be handled? Defaults to "pairwise". Available options: <ul style="list-style-type: none"> • "pairwise" — Computes correlation for all available cases between two variables • "listwise" — Computes correlation for all complete cases in the dataset
model	Character (length = 1). Defaults to "glasso". Available options: <ul style="list-style-type: none"> • "BGGM" — Computes the Bayesian Gaussian Graphical Model. Set argument <code>ordinal.categories</code> to determine levels allowed for a variable to be considered ordinal. See <code>?BGGM:estimate</code> for more details • "glasso" — Computes the GLASSO with EBIC model selection. See EBICglasso.qgraph for more details • "TMFG" — Computes the TMFG method. See TMFG for more details
algorithm	Character or <code>igraph cluster_*</code> function (length = 1). Defaults to "walktrap". Three options are listed below but all are available (see community.detection for other options): <ul style="list-style-type: none"> • "leiden" — See cluster_leiden for more details • "louvain" — By default, "louvain" will implement the Louvain algorithm using the consensus clustering method (see community.consensus for more information). This function will implement <code>consensus.method = "most_common"</code> and <code>consensus.iter = 1000</code> unless specified otherwise • "walktrap" — See cluster_walktrap for more details
verbose	Boolean (length = 1). Whether messages and (insignificant) warnings should be output. Defaults to FALSE (silent calls). Set to TRUE to see all messages and warnings for every function call
...	Additional arguments to be passed on to auto.correlate , network.estimate , community.detection , and community.consensus

Value

Returns a list containing:

network	A matrix containing a network estimated using <code>link[EGAnet]{network.estimate}</code>
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
n	Number of cases in data

Author(s)

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

References

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.

Introduced unidimensional checks, simulation with continuous and dichotomous data

Golino, H., Shi, D., Christensen, A. P., Garrido, L. E., Nieto, M. D., Sadana, R., & Thiagarajan, 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.

Compared all igraph community detection algorithms, simulation with continuous and polytomous data

Christensen, A. P., Garrido, L. E., Guerra-Pena, K., & Golino, H. (2023). Comparing community detection algorithms in psychometric networks: A Monte Carlo simulation. *Behavior Research Methods*.

See Also

[plot.EGAnet](#) for plot usage in EGAnet

Examples

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

# Estimate EGA
ega.wmt <- EGA.estimate(data = wmt)

# Estimate EGA with TMFG
ega.wmt.tmfG <- EGA.estimate(data = wmt, model = "TMFG")

# Estimate EGA with an {igraph} function (Fast-greedy)
ega.wmt.greedy <- EGA.estimate(
  data = wmt,
  algorithm = igraph::cluster_fast_greedy
)
```

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,
  corr = c("auto", "cor_auto", "cosine", "pearson", "spearman"),
  na.data = c("pairwise", "listwise"),
  model = c("BGGM", "glasso", "TMFG"),
  algorithm = c("leiden", "louvain", "walktrap"),
  plot.EGA = TRUE,
  verbose = FALSE,
  ...
)
```

Arguments

<code>data</code>	Matrix or data frame. Should consist only of variables to be used in the analysis
<code>n</code>	Numeric (length = 1). Sample size if data is a correlation matrix
<code>corr</code>	Character (length = 1). Method to compute correlations. Defaults to "auto". Available options: <ul style="list-style-type: none"> • "auto" — Automatically computes appropriate correlations for the data using Pearson's for continuous, polychoric for ordinal, tetrachoric for binary, and polyserial/biserial for ordinal/binary with continuous. To change the number of categories that are considered ordinal, use <code>ordinal.categories</code> (see polychoric.matrix for more details) • "cor_auto" — Uses <code>cor_auto</code> to compute correlations. Arguments can be passed along to the function • "cosine" — Uses <code>cosine</code> to compute cosine similarity • "pearson" — Pearson's correlation is computed for all variables regardless of categories • "spearman" — Spearman's rank-order correlation is computed for all variables regardless of categories <p>For other similarity measures, compute them first and input them into data with the sample size (n)</p>
<code>na.data</code>	Character (length = 1). How should missing data be handled? Defaults to "pairwise". Available options: <ul style="list-style-type: none"> • "pairwise" — Computes correlation for all available cases between two variables • "listwise" — Computes correlation for all complete cases in the dataset
<code>model</code>	Character (length = 1). Defaults to "glasso". Available options: <ul style="list-style-type: none"> • "BGGM" — Computes the Bayesian Gaussian Graphical Model. Set argument <code>ordinal.categories</code> to determine levels allowed for a variable to be considered ordinal. See <code>?BGGM::estimate</code> for more details • "glasso" — Computes the GLASSO with EBIC model selection. See EBICglasso.qgraph for more details

	<ul style="list-style-type: none"> • "TMFG" — Computes the TMFG method. See TMFG for more details
algorithm	<p>Character or igraph <code>cluster_*</code> function. Three options are listed below but all are available (see community.detection for other options):</p> <ul style="list-style-type: none"> • "leiden" — See cluster_leiden for more details. <i>Note:</i> The Leiden algorithm will default to the Constant Potts Model objective function (<code>objective_function = "CPM"</code>). Set <code>objective_function = "modularity"</code> to use modularity instead (see examples). By default, searches along resolutions from 0 to <code>max(abs(network))</code> or the maximum absolute edge weight in the network in 0.01 increments (<code>resolution_parameter = seq.int(0, max(abs(network)), 0.01)</code>). For modularity, searches along resolutions from 0 to 2 in 0.05 increments (<code>resolution_parameter = seq.int(0, 2, 0.05)</code>) by default. Use the argument <code>resolution_parameter</code> to change the search parameters (see examples) • "louvain" — See community.consensus for more details. By default, searches along resolutions from 0 to 2 in 0.05 increments (<code>resolution_parameter = seq.int(0, 2, 0.05)</code>). Use the argument <code>resolution_parameter</code> to change the search parameters (see examples) • "walktrap" — This algorithm is the default. See cluster_walktrap for more details. By default, searches along 3 to 8 steps (<code>steps = 3:8</code>). Use the argument <code>steps</code> to change the search parameters (see examples)
plot.EGA	Boolean. If TRUE, returns a plot of the network and its estimated dimensions. Defaults to TRUE
verbose	Boolean. Whether messages and (insignificant) warnings should be output. Defaults to FALSE (silent calls). Set to TRUE to see all messages and warnings for every function call
...	Additional arguments to be passed on to auto.correlate , network.estimate , community.detection , community.consensus , and EGA.estimate

Value

Returns a list containing:

EGA	EGA results of the best fitting solution
EntropyFit	tefi fit values for each solution
Lowest.EntropyFit	The best fitting solution based on tefi
parameter.space	Parameter values used in search space

Author(s)

Hudson Golino <hfg9s@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.

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(10), P10008.

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

[plot.EGAnet](#) for plot usage in EGAnet

Examples

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

# Estimate optimal EGA with Walktrap
fit.walktrap <- EGA.fit(
  data = wmt, algorithm = "walktrap",
  steps = 3:8, # default
  plot.EGA = FALSE # no plot for CRAN checks
)

# Estimate optimal EGA with Leiden and CPM
fit.leiden <- EGA.fit(
  data = wmt, algorithm = "leiden",
  objective_function = "CPM", # default
  # resolution_parameter = seq.int(0, max(abs(network)), 0.01),
  # For CPM, the default max resolution parameter
  # is set to the largest absolute edge in the network
  plot.EGA = FALSE # no plot for CRAN checks
)

# Estimate optimal EGA with Leiden and modularity
fit.leiden <- EGA.fit(
  data = wmt, algorithm = "leiden",
```

```

    objective_function = "modularity",
    resolution_parameter = seq.int(0, 2, 0.05),
    # default for modularity
    plot.EGA = FALSE # no plot for CRAN checks
  )

## Not run:
# Estimate optimal EGA with Louvain
fit.louvain <- EGA.fit(
  data = wmt, algorithm = "louvain",
  resolution_parameter = seq.int(0, 2, 0.05), # default
  plot.EGA = FALSE # no plot for CRAN checks
)
## End(Not run)

```

 ega.wmt

 EGA Network of [wmt2Data](#)

Description

EGA results from `ega.wmt <- EGA(wmt2[, 7:24])` for the Wiener Matrizen-Test (WMT-2)

Usage

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

Format

A list with 8 objects (see **Value** in [EGA](#))

Examples

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

 EGAnet-plot

 S3 Plot Methods for EGAnet

Description

General usage for plots created by EGAnet's S3 methods. Plots across the EGAnet package leverage GGally's [ggnnet2](#) and [ggplot2](#)'s [ggplot](#).

Most plots allow the full usage of the `gg*` series functionality and therefore plotting arguments should be referenced through those packages rather than here in EGAnet.

The sections below list the functions and their usage for the S3 plot methods. The plot methods are intended to be generic and without many arguments so that nearly all arguments are passed to [ggnnet2](#) and [ggplot](#).

There are some constraints placed on certain plots to keep the EGAnet style throughout the (network) plots in the package, so be aware that if some settings are not changing your plot output, then these settings might be fixed to maintain the EGAnet style

General Usage

```
plot(x, ...)

plot.dynEGA(x, base = 1, id = NULL, ...)

plot.dynEGA.Group(x, base = 1, ...)

plot.dynEGA.Individual(x, base = 1, id = NULL, ...)

plot.hierEGA(
  x, plot.type = c("multilevel", "separate"),
  color.match = FALSE, ...
)

plot.invariance(x, p_type = c("p", "p_BH"), p_value = 0.05, ...)

plot.TEFI.compare(x, base.name, comparison.name, base.color, comparison.color, ...)
```

General Arguments

- `x` — EGAnet object with available S3 plot method (see full list below)
- `color.palette` — Character (vector). Either a character (length = 1) from the pre-defined palettes in `color_palette_EGA` or character (length = total number of communities) using HEX codes (see **Color Palettes** and **Examples** sections)
- `layout` — Character (length = 1). Layouts can be set using `gplot.layout` and the ending layout name; for example, `gplot.layout.circle` can be set in these functions using `layout = "circle"` or `mode = "circle"` (see **Examples**)
- `base` — Numeric (length = 1). 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
- `id` — Numeric index(es) or character name(s). IDs to use when plotting `dynEGA` level = "individual". Defaults to NULL or 4 IDs drawn at random
- `plot.type` — Character (length = 1). Whether `hierEGA` networks should be plotted in a stacked, "multilevel" fashion or as "separate" plots. Defaults to "multilevel"
- `color.match` — Boolean (length = 1). Whether lower order community colors in the `hierEGA` plot should be "matched" and used as the border color for the higher order communities. Defaults to FALSE
- `p_type` — Character (length = 1). Type of p -value when plotting `invariance`. Defaults to "p" or uncorrected p -value. Set to "p_BH" for the Benjamini-Hochberg corrected p -value
- `p_value` — Numeric (length = 1). The p -value to use alongside `p_type` when plotting `invariance`. Defaults to 0.05

- `base.name` — Character (length = 1). A string to label the base structure in the plot. Defaults to "Base"
- `comparison.name` — Character (length = 1). A string to label the comparison structure in the plot. Defaults to "Comparison"
- `base.color` — Character (length = 1). A string to specifying the color of the base structure in the plot. Hex codes can be used. Defaults to "blue"
- `comparison.color` — Character (length = 1). A string to specifying the color of the comparison structure in the plot. Hex codes can be used. Defaults to "red"
- ... — Additional arguments to pass on to `ggnet2` and `gplot.layout` (see **Examples**)

*EGA Plots

`bootEGA`, `dynEGA`, `EGA`, `EGA.estimate`, `EGA.fit`, `hierEGA`, `invariance`, `riEGA`

All Available S3 Plot Methods

`boot.ergoInfo`, `bootEGA`, `dynEGA`, `dynEGA.Group`, `dynEGA.Individual`, `dynEGA.Population`, `EGA`, `EGA.estimate`, `EGA.fit`, `hierEGA`, `infoCluster`, `invariance`, `itemStability`, `riEGA`

Color Palettes

`color_palette_EGA` will implement some color palettes in EGAnet. The main EGAnet style palette is "polychrome". This palette currently has 40 colors but there will likely be a need to expand it further (e.g., `hierEGA` demands a lot of colors).

The `color.palette` argument will also accept HEX code colors that are the same length as the number of communities in the plot.

In any network plots, the `color.palette` argument can be used to select color palettes from `color_palette_EGA` as well as those in the color scheme of `RColorBrewer`

For more worked examples than below, see [Plots in {EGAnet}](#)

Examples

```
# Using different arguments in {GGally}'s `ggnet2`
plot(ega.wmt, node.size = 6, edge.size = 4)

# Using a different layout in {sna}'s `gplot.layout`
plot(ega.wmt, layout = "circle") # 'layout' argument
plot(ega.wmt, mode = "circle") # 'mode' argument

# Using different color palettes with `color_palette_EGA`

## Pre-defined palette
plot(ega.wmt, color.palette = "blue.ridge2")

## University of Virginia colors
plot(ega.wmt, color.palette = c("#232D4B", "#F84C1E"))

## Vanderbilt University colors
## (with additional {GGally} `ggnet2` argument)
```

```
plot(  
  ega.wmt, color.palette = c("#FFFFFF", "#866D4B"),  
  label.color = "#000000"  
)
```

Embed

Time-delay Embedding

Description

Reorganizes a single 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	Numeric vector. An observed time series to be reorganized into a time-delayed embedded matrix.
E	Numeric (length = 1). Number of embedded dimensions or the number of observations to be used. E = 5, for example, will generate a matrix with five columns corresponding to five consecutive observations across each row of the embedded matrix
tau	Numeric (length = 1). Number of observations to offset successive embeddings. A tau of one uses adjacent observations. Default is tau = 1

Value

Returns a numeric matrix

Author(s)

Pascal Deboeck <pascal.deboeck at psych.utah.edu> and Alexander P. Christensen <alexpaulchristensen@gmail.com>

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
time_series <- 49:56

# Time series embedding
Embed(time_series, E = 5, tau = 1)
```

entropyFit	<i>Entropy Fit Index</i>
------------	--------------------------

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	Numeric or character vector (length = ncol(data)). 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 at virginia.edu>, Alexander P. Christensen <alexpaulchristensen@gmail.com> and Robert Moulder <rgm4fd@virginia.edu>

References

Initial formalization and simulation

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*.

Examples

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

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

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

ergoInfo

Ergodicity Information Index

Description

Computes the Ergodicity Information Index

Usage

```
ergoInfo(
  dynEGA.object,
  use = c("edge.list", "unweighted", "weighted"),
  shuffles = 5000
)
```

Arguments

`dynEGA.object` A `dynEGA.ind.pop` object

`use` Character (length = 1). 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 = "unweighted"`. Current options are:

- `"edge.list"` — Calculates the algorithm complexity using the list of edges
- `"unweighted"` — Calculates the algorithm complexity using the binary weights of the encoded prime transformed network. 0 = edge absent and 1 = edge present

- "weighted" — Calculates the algorithm complexity using the weights of encoded prime-weight transformed network
- shuffles Numeric. Number of shuffles used to compute the Kolmogorov complexity. Defaults to 5000

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
complexity	The complexity metric proposed by Santora and Nicosia (2020)
EII	The Ergodicity Information Index

Author(s)

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

References**Original Implementation**

Golino, H., Nesselroade, J. R., & Christensen, A. P. (2022). Toward a psychology of individuals: The ergodicity information index and a bottom-up approach for finding generalizations. *PsyArXiv*.

Examples

```
# Obtain data
sim.dynEGA <- sim.dynEGA # bypasses CRAN checks

## Not run:
# Dynamic EGA individual and population structure
dyn.ega1 <- dynEGA.ind.pop(
  data = sim.dynEGA[,-26], n.embed = 5, tau = 1,
  delta = 1, id = 25, use.derivatives = 1,
  ncores = 2, corr = "pearson"
)

# Compute empirical ergodicity information index
eii <- ergoInfo(dyn.ega1)
## End(Not run)
```

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

Description

Computes the fit (Generalized TEFI) of a hierarchical or correlated bifactor dimensionality structure (or `hierEGA` objects) using Von Neumman's entropy when the input is a correlation matrix. Lower values suggest better fit of a structure to the data

Usage

```
genTEFI(data, structure = NULL, verbose = TRUE)
```

Arguments

<code>data</code>	Matrix, data frame, or <code>hierEGA</code> object. Can be raw data or correlation matrix
<code>structure</code>	List (length = levels). A list containing the hierarchical structure. Each list element corresponds to increasing levels (1 = first level, 2 = second level, etc.). The length of the first element (first level) should be the same as the number of variables in data. Each level after should either be the number of variables (<code>ncol(data)</code>) or the maximum number of dimensions from the preceding level (<code>max(previous_dimensions)</code>)
<code>verbose</code>	Boolean (length = 1). Whether messages and (insignificant) warnings should be output. Defaults to TRUE to see all messages and warnings for every function call. Set to FALSE to ignore messages and warnings

Value

Returns a `levels + 1` columns data frame of the Generalized Total Entropy Fit Index using Von Neumman's entropy (`VN.Entropy.Fit`) (first column) as well as each individual's levels entropy

Author(s)

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

Examples

```
# Example using network scores
opt.hier <- hierEGA(
  data = optimism, scores = "network",
  plot.EGA = FALSE # No plot for CRAN checks
)

# Compute the Generalized Total Entropy Fit Index
genTEFI(opt.hier)
```

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,
  na.derivative = c("none", "kalman", "rowwise", "skipover")
)
```

Arguments

x	Numeric vector. An observed time series
n.embed	Numeric (length = 1). Number of embedded dimensions (the number of observations to be used in the Embed function)
tau	Numeric (length = 1). Number of observations to offset successive embeddings in the Embed function. A tau of one uses adjacent observations. Default is 1
delta	Numeric (length = 1). The time between successive observations in the time series. Default is 1
order	Numeric (length = 1). 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)
na.derivative	Character (length = 1). How should missing data in the embeddings be handled? Available options (see Boker et al. (2018) for more details): <ul style="list-style-type: none"> "none" (default) — does nothing and leaves NAs in data "kalman" — uses Kalman smoothing (KalmanSmooth) with structural time series models (StructTS) to impute missing values. This approach models the underlying temporal dependencies (trend, seasonality, autocorrelation) to generate estimates for missing observations while preserving the original time scale. More computationally intensive than the other methods but typically provides the most accurate imputation by respecting the stochastic properties of the time series

- "rowwise" — adjusts time interval with respect to each embedding ensuring time intervals are adaptive to the missing data (tends to be more accurate than "none")
- "skipover" — "skips over" missing data and treats the non-missing points as continuous points in time (note that the time scale shifts to the "per mean time interval," which is different and *larger* than the original scale)

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

GLLA implementation

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.

Filtering procedure

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

Missing Data

Boker, S. M., Tiberio, S. S., & Moulder, R. G. (2018). Robustness of time delay embedding to sampling interval misspecification. In K. van Montfort, J. H. L. Oud, & M. C. Voelke (Eds.), *Continuous Time Modeling in the Behavioral and Related Sciences* (pp. 239–258). Springer International Publishing.

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 the Louvain algorithm (`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 (for more details, see Jiménez et al., 2023)

Usage

```
hierEGA(
  data,
  loading.method = c("original", "revised"),
  rotation = NULL,
  scores = c("factor", "network"),
  loading.structure = c("simple", "full"),
  impute = c("mean", "median", "none"),
  corr = c("auto", "cor_auto", "pearson", "spearman"),
  na.data = c("pairwise", "listwise"),
  model = c("BGGM", "glasso", "TMFG"),
  lower.algorithm = "louvain",
  higher.algorithm = c("leiden", "louvain", "walktrap"),
  uni.method = c("expand", "LE", "louvain"),
  plot.EGA = TRUE,
  verbose = FALSE,
  ...
)
```

Arguments

<code>data</code>	Matrix or data frame. Should consist only of variables to be used in the analysis (does not accept correlation matrices)
<code>loading.method</code>	Character (length = 1). Sets network loading calculation based on implementation described in "original" (Christensen & Golino, 2021) or the "revised" (Christensen et al., 2024) implementation. Defaults to "revised"
<code>rotation</code>	Character. A rotation to use to obtain a simpler structure. For a list of rotations, see rotations for options. Defaults to NULL or no rotation. By setting a rotation, scores estimation will be based on the rotated loadings rather than unrotated loadings
<code>scores</code>	Character (length = 1). How should scores for the higher-order structure be estimated? Defaults to "network" for network scores computed using the <code>net.scores</code> function. Set to "factor" for factor scores computed using <code>fa</code> . Factors scores will be based on EFA (as in Jiménez et al., 2023)

Factor scores use the number of communities from EGA. Estimated factor loadings may not align with these communities. The plots using factor scores will have higher order factors that may not completely map on to the lower order communities. Look at `$hierarchical$higher_order$lower_loadings` to determine the composition of the lower order factors.

loading.structure	<p>Character (length = 1). Whether simple structure or the saturated loading matrix should be used when computing scores (scores = "network" only). Defaults to "simple"</p> <p>"simple" structure more closely mirrors traditional hierarchical factor analytic methods such as CFA; "full" structure more closely mirrors EFA methods</p> <p>Simple structure is the more conservative (established) approach and is therefore the default. Treat "full" as experimental as proper vetting and validation has not been established</p>
impute	<p>Character (length = 1). If there are any missing data, then imputation can be implemented. Available options:</p> <ul style="list-style-type: none"> • "none" — Default. No imputation is performed • "mean" — The mean value of each variable is used to replace missing data for that variable • "median" — The median value of each variable is used to replace missing data for that variable
corr	<p>Character (length = 1). Method to compute correlations. Defaults to "auto". Available options:</p> <ul style="list-style-type: none"> • "auto" — Automatically computes appropriate correlations for the data using Pearson's for continuous, polychoric for ordinal, tetrachoric for binary, and polyserial/biserial for ordinal/binary with continuous. To change the number of categories that are considered ordinal, use <code>ordinal.categories</code> (see polychoric.matrix for more details) • "cor_auto" — Uses <code>cor_auto</code> to compute correlations. Arguments can be passed along to the function • "pearson" — Pearson's correlation is computed for all variables regardless of categories • "spearman" — Spearman's rank-order correlation is computed for all variables regardless of categories <p>For other similarity measures, compute them first and input them into data with the sample size (n)</p>
na.data	<p>Character (length = 1). How should missing data be handled? Defaults to "pairwise". Available options:</p> <ul style="list-style-type: none"> • "pairwise" — Computes correlation for all available cases between two variables • "listwise" — Computes correlation for all complete cases in the dataset
model	<p>Character (length = 1). Defaults to "glasso". Available options:</p> <ul style="list-style-type: none"> • "BGGM" — Computes the Bayesian Gaussian Graphical Model. Set argument <code>ordinal.categories</code> to determine levels allowed for a variable to be considered ordinal. See <code>?BGGM::estimate</code> for more details

	<ul style="list-style-type: none"> • "glasso" — Computes the GLASSO with EBIC model selection. See EBICglasso.qgraph for more details • "TMFG" — Computes the TMFG method. See TMFG for more details
lower.algorithm	<p>Character or <code>cluster_*</code> function (length = 1). Defaults to the lower order "louvain" with most common consensus clustering (1000 iterations; see community.consensus for more details)</p> <p>Louvain with consensus clustering is <i>strongly</i> recommended. Using any other algorithm is considered <i>experimental</i> as they have not been designed to capture lower order communities</p>
higher.algorithm	<p>Character or <code>cluster_*</code> function (length = 1). Defaults to "louvain". Three options are listed below but all are available (see community.detection for other options):</p> <ul style="list-style-type: none"> • "leiden" — See cluster_leiden for more details • "louvain" — By default, "louvain" will implement the higher-order (order = "higher") Louvain algorithm using the consensus clustering method (see community.consensus for more information). This function will implement <code>consensus.method = "most_common"</code> and <code>consensus.iter = 1000</code> unless specified otherwise • "walktrap" — See cluster_walktrap for more details <p>Using <code>algorithm</code> will set only <code>higher.algorithm</code> and <code>lower.algorithm</code> will default to Louvain with most common consensus clustering (1000 iterations)</p>
uni.method	<p>Character (length = 1). What unidimensionality method should be used? Defaults to "louvain". Available options:</p> <ul style="list-style-type: none"> • "expand" — Expands the correlation matrix with four variables correlated 0.50. 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 method was used in the Golino et al.'s (2020) <i>Psychological Methods</i> simulation • "LE" — Applies the Leading Eigenvector algorithm (cluster_leading_eigen) on the empirical correlation matrix. If the number of dimensions is 1, then the Leading Eigenvector solution is used; otherwise, regular EGA is used. This method was used in the Christensen et al.'s (2023) <i>Behavior Research Methods</i> simulation • "louvain" — Applies the Louvain algorithm (cluster_louvain) on the empirical correlation matrix. If the number of dimensions is 1, then the Louvain solution is used; otherwise, regular EGA is used. This method was validated Christensen's (2022) <i>PsyArXiv</i> simulation. Consensus clustering can be used by specifying either <code>"consensus.method"</code> or <code>"consensus.iter"</code>
plot.EGA	<p>Boolean. If TRUE, returns a plot of the network and its estimated dimensions. Defaults to TRUE</p>
verbose	<p>Boolean (length = 1). Whether messages and (insignificant) warnings should be output. Defaults to FALSE (silent calls). Set to TRUE to see all messages and warnings for every function call</p>
...	<p>Additional arguments to be passed on to auto.correlate, network.estimate, community.detection, community.consensus, EGA, and rotations</p>

Value

Returns a list of lists containing:

lower_order	EGA results for the lower order structure
higher_order	EGA results for the higher order structure
parameters	A list containing lower_loadings and lower_scores that were used to estimate scores and the higher order EGA results, respectively
dim.variables	A data frame with variable names and their lower and higher order assignments
TEFI	Generalized TEFI using <code>tefi</code>
plot.hierEGA	Plot output if <code>plot.EGA = TRUE</code>

Author(s)

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

References**Hierarchical EGA simulation**

Jiménez, M., Abad, F. J., Garcia-Garzon, E., Golino, H., Christensen, A. P., & Garrido, L. E. (2023). Dimensionality assessment in bifactor structures with multiple general factors: A network psychometrics approach. *Psychological Methods*.

3+ level hierarchical EGA

Samo, A., Christensen, A. P., Abad, F. J., Garrido, L. E., Garcia-Garzon, E., Golino, H. & McAbee, S. T. (2023). Building the structure of personality from the bottom-up using Hierarchical Exploratory Graph Analysis. *PsyArXiv*.

Conceptual implementation

Golino, H., Thiagarajan, J. A., Sadana, R., Teles, M., Christensen, A. P., & Boker, S. M. (2020). 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*.

Revised network loadings

Christensen, A. P., Golino, H., Abad, F. J., & Garrido, L. E. (2024). Revised network loadings. *PsyArXiv*.

See Also

[plot.EGAnet](#) for plot usage in

Examples

```
# Example using network scores
opt.hier <- hierEGA(
  data = optimism, scores = "network",
  plot.EGA = FALSE # No plot for CRAN checks
)
```

```
# Plot multilevel plot
plot(opt.hier, plot.type = "multilevel")

# Plot multilevel plot with higher order
# border color matching the corresponding
# lower order color
plot(opt.hier, color.match = TRUE)

# Plot levels separately
plot(opt.hier, plot.type = "separate")
```

igraph2matrix *Convert network to matrix*

Description

Converts network to matrix

Usage

```
igraph2matrix(igraph_network, diagonal = 0)
```

Arguments

igraph_network network object
diagonal Numeric (length = 1). Value to be placed on the diagonal of network. Defaults to 0

Value

Returns a network in the format

Author(s)

Hudson Golino <hfg9s at virginia.edu> & Alexander P. Christensen <alexander.christensen at Vanderbilt.Edu>

Examples

```
# Convert network to {igraph}
igraph_network <- convert2igraph(ega.wmt$network)

# Convert network back to matrix
igraph2matrix(igraph_network)
```

 infoCluster

Information Theoretic Mixture Clustering for dynEGA

Description

Performs hierarchical clustering using Jensen-Shannon distance followed by the Louvain algorithm with consensus clustering. The method iteratively identifies smaller and smaller clusters until there is no change in the clusters identified

Usage

```
infoCluster(dynEGA.object, plot.cluster = TRUE, ...)
```

Arguments

<code>dynEGA.object</code>	A dynEGA or a dynEGA.ind.pop object that is used to match the arguments of the EII object
<code>plot.cluster</code>	Boolean (length = 1). Should plot of optimal and hierarchical clusters be output? Defaults to TRUE. Set to FALSE to not plot
<code>...</code>	Additional arguments to be passed on to jsd

Value

Returns a list containing:

<code>clusters</code>	A vector corresponding to cluster each participant belongs to
<code>clusterTree</code>	The dendrogram from hclust the hierarhical clustering
<code>clusterPlot</code>	Plot output from results
<code>JSD</code>	Jensen-Shannon Distance

Author(s)

Hudson Golino <hfg9s at virginia.edu> & Alexander P. Christensen <alexander.christensen at Vanderbilt.Edu>

See Also

[plot.EGAnet](#) for plot usage in [EGAnet](#)

Examples

```
# Obtain data
sim.dynEGA <- sim.dynEGA # bypasses CRAN checks

## Not run:
# Dynamic EGA individual and population structure
dyn.ega1 <- dynEGA.ind.pop(
```

```

data = sim.dynEGA, n.embed = 5, tau = 1,
delta = 1, id = 25, use.derivatives = 1,
ncores = 2, corr = "pearson"
)

# Perform information-theoretic clustering
clust1 <- infoCluster(dynEGA.object = dyn.ega1)
## End(Not run)

```

information

Information Theory Metrics

Description

A general function to compute several different information theory metrics

Usage

```

information(
  data,
  base = 2.718282,
  bins = floor(sqrt(nrow(data)/5)),
  statistic = c("entropy", "joint.entropy", "conditional.entropy", "total.correlation",
    "dual.total.correlation", "o.information")
)

```

Arguments

data	Matrix or data frame. Should consist only of variables to be used in the analysis
base	Numeric (length = 1). Base of logarithm to use for entropy. Common options include: <ul style="list-style-type: none"> • 2 — bits • 2.718282 — nats • 10 — bans Defaults to $\exp(1)$ or 2.718282
bins	Numeric (length = 1). Number of bins if data are not discrete. Defaults to $\text{floor}(\sqrt{\text{nrow}(\text{data}) / 5})$
statistic	Character. Information theory statistics to compute. Available options: <ul style="list-style-type: none"> • "entropy" — Shannon's entropy (Shannon, 1948) for each variable in data. Values range from \emptyset to $\log(k)$ where k is the number of categories for the variable • "joint.entropy" — shared uncertainty over all variables in data. Values range from the maximum of the individual entropies to the sum of individual entropies

- "conditional.entropy" — uncertainty remaining after considering all other variables in data. Values range from 0 to the individual entropy of the conditioned variable
- "total.correlation" — generalization of mutual information to more than two variables (Watanabe, 1960). Quantifies the redundancy of information in data. Values range from 0 to the sum of individual entropies minus the maximum of the individual entropies
- "dual.total.correlation" — "shared randomness" or total uncertainty remaining in the data (Han, 1978). Values range from 0 to joint entropy
- "o.information" — quantifies the extent to which the data is represented by lower-order (> 0 ; redundancy) or higher-order (< 0 ; synergy) constraint (Crutchfield, 1994)

By default, all statistics are computed

Value

Returns list containing *only* requested statistic

Author(s)

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

References

Shannon's entropy

Shannon, C. E. (1948). A mathematical theory of communication. *The Bell System Technical Journal*, 27(3), 379-423.

Formalization of total correlation

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

Applied implementation of total correlation

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.

Formalization of dual total correlation

Te Sun, H. (1978). Nonnegative entropy measures of multivariate symmetric correlations. *Information and Control*, 36, 133-156.

Formalization of O-information

Crutchfield, J. P. (1994). The calculi of emergence: Computation, dynamics and induction. *Physica D: Nonlinear Phenomena*, 75(1-3), 11-54.

Applied implementation of O-information

Marinazzo, D., Van Roozendaal, J., Rosas, F. E., Stella, M., Comolatti, R., Colenbier, N., Stramaglia, S., & Rosseel, Y. (2024). An information-theoretic approach to build hypergraphs in psychometrics. *Behavior Research Methods*, 1-23.

Examples

```
# All measures
information(wmt2[,7:24])

# One measures
information(wmt2[,7:24], statistic = "joint.entropy")
```

intelligenceBattery *Intelligence Data*

Description

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

Usage

```
data(intelligenceBattery)
```

Format

A 1185x125 response matrix

Examples

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

invariance *Measurement Invariance of [EGA Structure](#)*

Description

Estimates configural invariance using [bootEGA](#) on all data (across groups) first. After configural variance is established, then metric invariance is tested using the community structure that established configural invariance (see **Details** for more information on this process)

Usage

```

invariance(
  data,
  groups,
  structure = NULL,
  iter = 500,
  configural.threshold = 0.7,
  configural.type = c("parametric", "resampling"),
  corr = c("auto", "cor_auto", "pearson", "spearman"),
  na.data = c("pairwise", "listwise"),
  model = c("BGGM", "glasso", "TMFG"),
  algorithm = c("leiden", "louvain", "walktrap"),
  uni.method = c("expand", "LE", "louvain"),
  ncores,
  seed = NULL,
  verbose = TRUE,
  ...
)

```

Arguments

<code>data</code>	Matrix or data frame. Should consist only of variables to be used in the analysis
<code>groups</code>	Numeric or character vector (length = <code>nrow(data)</code>). Group membership corresponding to each case in data
<code>structure</code>	Numeric or character vector (length = <code>ncol(data)</code>). A vector representing the structure (numbers or labels for each item). Can be theoretical factors or the structure detected by EGA . If supplied, then configural invariance check is skipped (i.e., configural invariance is assumed based on the given structure)
<code>iter</code>	Numeric (length = 1). Number of iterations to perform for the permutation. Defaults to 500 (recommended)
<code>configural.threshold</code>	Numeric (length = 1). Value to use a threshold in itemStability to determine which items should be removed during configural invariance (see Details for more information). Defaults to 0.70 (recommended)
<code>configural.type</code>	Character (length = 1). Type of bootstrap to use for configural invariance in bootEGA . Defaults to "parametric"
<code>corr</code>	Character (length = 1). Method to compute correlations. Defaults to "auto". Available options: <ul style="list-style-type: none"> "auto" — Automatically computes appropriate correlations for the data using Pearson's for continuous, polychoric for ordinal, tetrachoric for binary, and polyserial/biserial for ordinal/binary with continuous. To change the number of categories that are considered ordinal, use <code>ordinal.categories</code> (see polychoric.matrix for more details) "cor_auto" — Uses cor_auto to compute correlations. Arguments can be passed along to the function

- "pearson" — Pearson's correlation is computed for all variables regardless of categories
- "spearman" — Spearman's rank-order correlation is computed for all variables regardless of categories

For other similarity measures, compute them first and input them into data with the sample size (n)

na.data	Character (length = 1). How should missing data be handled? Defaults to "pairwise". Available options: <ul style="list-style-type: none"> • "pairwise" — Computes correlation for all available cases between two variables • "listwise" — Computes correlation for all complete cases in the dataset
model	Character (length = 1). Defaults to "glasso". Available options: <ul style="list-style-type: none"> • "BGGM" — Computes the Bayesian Gaussian Graphical Model. Set argument <code>ordinal.categories</code> to determine levels allowed for a variable to be considered ordinal. See <code>?BGGM:estimate</code> for more details • "glasso" — Computes the GLASSO with EBIC model selection. See EBICglasso.qgraph for more details • "TMFG" — Computes the TMFG method. See TMFG for more details
algorithm	Character or <code>cluster_*</code> function (length = 1). Defaults to "walktrap". Three options are listed below but all are available (see community.detection for other options): <ul style="list-style-type: none"> • "leiden" — See cluster_leiden for more details • "louvain" — By default, "louvain" will implement the Louvain algorithm using the consensus clustering method (see community.consensus for more information). This function will implement <code>consensus.method = "most_common"</code> and <code>consensus.iter = 1000</code> unless specified otherwise • "walktrap" — See cluster_walktrap for more details
uni.method	Character (length = 1). What unidimensionality method should be used? Defaults to "louvain". Available options: <ul style="list-style-type: none"> • "expand" — Expands the correlation matrix with four variables correlated 0.50. 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 method was used in the Golino et al.'s (2020) <i>Psychological Methods</i> simulation • "LE" — Applies the Leading Eigenvector algorithm (cluster_leading_eigen) on the empirical correlation matrix. If the number of dimensions is 1, then the Leading Eigenvector solution is used; otherwise, regular EGA is used. This method was used in the Christensen et al.'s (2023) <i>Behavior Research Methods</i> simulation • "louvain" — Applies the Louvain algorithm (cluster_louvain) on the empirical correlation matrix. If the number of dimensions is 1, then the Louvain solution is used; otherwise, regular EGA is used. This method was validated Christensen's (2022) <i>PsyArXiv</i> simulation. Consensus clustering can be used by specifying either <code>"consensus.method"</code> or <code>"consensus.iter"</code>

ncores	Numeric (length = 1). Number of cores to use in computing results. Defaults to <code>ceiling(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 type: <code>parallel::detectCores()</code>
seed	Numeric (length = 1). Defaults to NULL or random results. Set for reproducible results. See Reproducibility and PRNG for more details on random number generation in
verbose	Boolean (length = 1). Should progress be displayed? Defaults to TRUE. Set to FALSE to not display progress
...	Additional arguments that can be passed on to auto.correlate , network. estimation , community.detection , community.consensus , EGA , bootEGA , and net.loads

Details

In traditional psychometrics, measurement invariance is performed in sequential testing from more flexible (more free parameters) to more rigid (fewer free parameters) structures. Measurement invariance in network psychometrics is no different.

Configural Invariance

To establish configural invariance, the data are collapsed across groups and a common sample structure is identified used [bootEGA](#) and [itemStability](#). If some variables have a replication less than 0.70 in their assigned dimension, then they are considered unstable and therefore not invariant. These variables are removed and this process is repeated until all items are considered stable (replication values greater than 0.70) or there are no variables left. If configural invariance cannot be established, then the last run of results are returned and metric invariance is not tested (because configural invariance is not met). Importantly, if any variables *are* removed, then configural invariance is not met for the original structure. Any removal would suggest only partial configural invariance is met.

Metric Invariance

The variables that remain after configural invariance are submitted to metric invariance. First, each group estimates a network and then network loadings ([net.loads](#)) are computed using the assigned community memberships (determined during configural invariance). Then, the difference between the assigned loadings of the groups is computed. This difference represents the empirical values. Second, the group memberships are permuted and networks are estimated based on the these permuted groups for `iter` times. Then, network loadings are computed and the difference between the assigned loadings of the group is computed, resulting in a null distribution. The empirical difference is then compared against the null distribution using a two-tailed p -value based on the number of null distribution differences that are greater and less than the empirical differences for each variable. Both uncorrected and false discovery rate corrected p -values are returned in the results. Uncorrected p -values are flagged for significance along with the direction of group differences.

Three or More Groups

When there are 3 or more groups, the function performs metric invariance testing by comparing all possible pairs of groups. Specifically:

- *Pairwise Comparisons*: The function generates all possible unique group pairings and computes the differences in network loadings for each pair. The same community structure, derived from configural invariance or provided by the user, is used for all groups.

- *Permutation Testing*: For each group pair, permutation tests are conducted to assess the statistical significance of the observed differences in loadings. p -values are calculated based on the proportion of permuted differences that are greater than or equal to the observed difference.
- *Result Compilation*: The function compiles the results for each pair including both uncorrected (p) and FDR-corrected (Benjamini-Hochberg; p_{BH}) p -values, and the direction of differences. It returns a summary of the findings for all pairwise comparisons.

This approach allows for a detailed examination of metric invariance across multiple groups, ensuring that all potential differences are thoroughly assessed while maintaining the ability to identify specific group differences.

For more details, see Jamison, Golino, and Christensen (2023)

Value

Returns a list containing:

<code>configural.results</code>	<code>bootEGA</code> results from the final run that produced configural invariance. This output will be output on the final run of unsuccessful configural invariance runs
<code>memberships</code>	Original memberships provided in <code>structure</code> or from <code>EGA</code> if <code>structure = NULL</code>
<code>EGA</code>	Original <code>EGA</code> results for the full sample
<code>groups</code>	A list containing: <ul style="list-style-type: none"> • <code>EGA</code> — <code>EGA</code> results for each group • <code>loadings</code> — Network loadings (<code>net.loadings</code>) for each group • <code>loadingsDifference</code> — Difference between the dominant loadings of each group
<code>permutation</code>	A list containing: <ul style="list-style-type: none"> • <code>groups</code> — Permuted groups across iterations • <code>loadings</code> — Network loadings (<code>net.loadings</code>) for each group for each permutation • <code>loadingsDifference</code> — Difference between the dominant loadings of each group for each permutation
<code>results</code>	Data frame of the results (which are printed)

Author(s)

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

References

Original implementation

Jamison, L., Christensen, A. P., & Golino, H. F. (2024). Metric invariance in exploratory graph analysis via permutation testing. *Methodology*, 20(2), 144-186.

See Also

[plot.EGAnet](#) for plot usage in

Examples

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

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

## Not run:
# Measurement invariance
results <- invariance(wmt, groups, ncores = 2)

# Plot with uncorrected alpha = 0.05
plot(results, p_type = "p", p_value = 0.05)

# Plot with BH-corrected alpha = 0.10
plot(results, p_type = "p_BH", p_value = 0.10)
## End(Not run)
```

itemDiagnostics

Diagnostics Analysis for Low Stability Items

Description

Computes item diagnostics to determine whether there exist several potential psychometric issues that affect [itemStability](#). These issues include local dependence (using [UVA](#)), minor dimensions (using [cosine](#) on the stability patterns), multidimensional items (using [net.loads](#)), and items with low loadings (using [net.loads](#))

Usage

```
itemDiagnostics(data, ...)
```

Arguments

<code>data</code>	Matrix or data frame. Should consist only of variables to be used in the analysis
<code>...</code>	Additional arguments to pass on to bootEGA , net.loads , and UVA

Value

Returns a list containing:

<code>diagnostics</code>	A data frame containing the diagnostics of low item stabilities (see itemStability)
<code>boot</code>	Output from bootEGA

uva	Output from UVA
minor	A list containing suggested items to keep, remove, and a matrix for probable minor dimensions (<code>minor.matrix</code>)
loadings	Output from net.loads
suggested	Variables that are suggested to be retained to increase item stability

Author(s)

Alexander P. Christensen <alexpaulchristensen@gmail.com>, Hudson Golino <hfg9s@virginia.edu>, and Luis Eduardo Garrido <garrido.luiseduardo@gmail.com>

Examples

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

## Not run:
# Obtain diagnostics
diagnostics <- itemDiagnostics(wmt, ncores = 2)
## End(Not run)
```

itemStability *Item Stability Statistics from* [bootEGA](#)

Description

Based on the [bootEGA](#) results, this function computes and plots the number of times an variable is estimated in the same dimension as originally estimated by an empirical [EGA](#) structure or a theoretical/input structure. The output also contains each variable's replication frequency (i.e., proportion of bootstraps that a variable appeared in each dimension)

Usage

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

Arguments

bootega.obj	A bootEGA object
IS.plot	Boolean (length = 1). Should the plot be produced for item.replication? Defaults to TRUE
structure	Numeric (length = number of variables). A theoretical or pre-defined structure. Defaults to NULL or the empirical EGA result in the bootega.obj
...	Deprecated arguments from previous versions of itemStability

Value

Returns a list containing:

membership	<p>A list containing:</p> <ul style="list-style-type: none"> • <code>empirical</code> — A vector of the empirical memberships from the empirical EGA result • <code>bootstrap</code> — A matrix of the homogenized memberships from the replicate samples in the bootEGA results • <code>structure</code> — A vector of the structure used in the analysis. If <code>structure = NULL</code>, then this output will be the same as <code>empirical</code>
item.stability	<p>A list containing:</p> <ul style="list-style-type: none"> • <code>empirical.dimensions</code> — A vector of the proportion of times each item replicated within the structure defined by <code>structure</code> • <code>all.dimensions</code> — A matrix of the proportion of times each item replicated in each of the structure defined dimensions
plot	Plot output if <code>IS.plot = TRUE</code>

Author(s)

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

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.

Conceptual introduction

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

[plot.EGAnet](#) for plot usage in [EGAnet](#)

Examples

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

## Not run:
# Standard EGA example
boot.wmt <- bootEGA(
  data = wmt, iter = 500,
  type = "parametric", ncores = 2
)
## End(Not run)
```

```
# Standard item stability
wmt.is <- itemStability(boot.wmt)

## Not run:
# EGA fit example
boot.wmt.fit <- bootEGA(
  data = wmt, iter = 500,
  EGA.type = "EGA.fit",
  type = "parametric", ncores = 2
)

# EGA fit item stability
wmt.is.fit <- itemStability(boot.wmt.fit)

# Hierarchical EGA example
boot.wmt.hier <- bootEGA(
  data = wmt, iter = 500,
  EGA.type = "hierEGA",
  type = "parametric", ncores = 2
)

# Hierarchical EGA item stability
wmt.is.hier <- itemStability(boot.wmt.hier)

# Random-intercept EGA example
boot.wmt.ri <- bootEGA(
  data = wmt, iter = 500,
  EGA.type = "riEGA",
  type = "parametric", ncores = 2
)

# Random-intercept EGA item stability
wmt.is.ri <- itemStability(boot.wmt.ri)
## End(Not run)
```

jsd

Jensen-Shannon Distance

Description

Computes the Jensen-Shannon Distance between two networks

Usage

```
jsd(network1, network2, method = c("kld", "spectral"), signed = TRUE)
```

Arguments

network1	Matrix or data frame. Network to be compared
network2	Matrix or data frame. Second network to be compared
method	Character (length = 1). Method to compute Jensen-Shannon Distance. Defaults to "spectral". Available options: <ul style="list-style-type: none"> • "kld" — Uses Kullback-Leibler Divergence • "spectral" — Uses eigenvalues of combinatorial Laplacian matrix to compute Von Neumann entropy
signed	Boolean. (length = 1). Should networks be remain signed? Defaults to TRUE

Value

Returns Jensen-Shannon Distance

Author(s)

Hudson Golino <hfg9s at virginia.edu> & Alexander P. Christensen <alexander.christensen at Vanderbilt.Edu>

Examples

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

# Set seed (for reproducibility)
set.seed(1234)

# Split data
split1 <- sample(
  1:nrow(wmt), floor(nrow(wmt) / 2)
)
split2 <- setdiff(1:nrow(wmt), split1)

# Obtain split data
data1 <- wmt[split1,]
data2 <- wmt[split2,]

# Perform EBICglasso
glas1 <- EBICglasso.qgraph(data1)
glas2 <- EBICglasso.qgraph(data2)

# Spectral JSD
jsd(glas1, glas2)
# 0.1595893

# Spectral JSS (similarity)
1 - jsd(glas1, glas2)
# 0.8404107
```

```
# Jensen-Shannon Divergence
jsd(glas1, glas2, method = "kld")
# 0.1393621
```

known.graph *Re-fit Network*

Description

Refits a network with unregularized partial correlations based on some known graph (perhaps estimated with a regularization method) following Hastie, Tibshirani, and Friedman (2008)

Usage

```
known.graph(S, A, method = c("glasso", "HTF"), tol = 0.000001, max.iter = 100)
```

Arguments

S	Matrix. Covariance or correlation matrix
A	Adjacency matrix. Unweighted network structure where 1 is an edge present and 0 is an edge absent
method	Character (length = 1). Whether to use the glasso method without penalization or the HTF (Haste, Tibshirani, & Friedman, 2008) method. Defaults to "glasso", which tends to be more robust
tol	Numeric (length = 1). Tolerance for convergence. The algorithm stops when the maximum absolute change in covariance matrix elements between iterations is less than tol. Defaults to 1e-06
max.iter	Numeric (length = 100). Maximum number of iterations to achieve tolerance before stopping

Value

Returns a list containing:

network	Estimated network
W	Estimated covariance matrix
Theta	Estimated inverse covariance matrix
iterations	Number of iterations to converge (or maximum if it did not)
converged	Whether the algorithm converged

Author(s)

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

References

HTF Implementation on p. 631–634

Hastie, T., Tibshirani, R., & Friedman, J. (2008). The elements of statistical learning: Data mining, inference, and prediction (2nd ed.). New York, NY: Springer.

Examples

```
# Obtain data
wmt <- depression[,24:44]

# Obtain correlation matrix
wmt_R <- auto.correlate(wmt)

# Estimate network
wmt_network <- network.estimate(wmt_R, n = nrow(wmt))

# Obtain adjacency
wmt_A <- wmt_network
wmt_A[] <- ifelse(wmt_A != 0, 1, 0)

# Obtain unregularized estimate
wmt_unreg <- known.graph(S = wmt_R, A = wmt_A)
```

LCT

Loadings Comparison Test

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 = NULL,
  corr = c("auto", "cor_auto", "pearson", "spearman"),
  na.data = c("pairwise", "listwise"),
  model = c("BGGM", "glasso", "TMFG"),
  algorithm = c("leiden", "louvain", "walktrap"),
  uni.method = c("expand", "LE", "louvain"),
  iter = 100,
  seed = NULL,
  verbose = TRUE,
  ...
)
```

Arguments

<code>data</code>	Matrix or data frame. Should consist only of variables to be used in the analysis. Can be raw data or a correlation matrix
<code>n</code>	Numeric (length = 1). Sample size if data provided is a correlation matrix
<code>corr</code>	Character (length = 1). Method to compute correlations. Defaults to "auto". Available options: <ul style="list-style-type: none"> • "auto" — Automatically computes appropriate correlations for the data using Pearson's for continuous, polychoric for ordinal, tetrachoric for binary, and polyserial/biserial for ordinal/binary with continuous. To change the number of categories that are considered ordinal, use <code>ordinal.categories</code> (see polychoric.matrix for more details) • "cor_auto" — Uses <code>cor_auto</code> to compute correlations. Arguments can be passed along to the function • "pearson" — Pearson's correlation is computed for all variables regardless of categories • "spearman" — Spearman's rank-order correlation is computed for all variables regardless of categories <p>For other similarity measures, compute them first and input them into <code>data</code> with the sample size (<code>n</code>)</p>
<code>na.data</code>	Character (length = 1). How should missing data be handled? Defaults to "pairwise". Available options: <ul style="list-style-type: none"> • "pairwise" — Computes correlation for all available cases between two variables • "listwise" — Computes correlation for all complete cases in the dataset
<code>model</code>	Character (length = 1). Defaults to "glasso". Available options: <ul style="list-style-type: none"> • "BGGM" — Computes the Bayesian Gaussian Graphical Model. Set argument <code>ordinal.categories</code> to determine levels allowed for a variable to be considered ordinal. See <code>?BGGM::estimate</code> for more details • "glasso" — Computes the GLASSO with EBIC model selection. See EBICglasso.qgraph for more details • "TMFG" — Computes the TMFG method. See TMFG for more details
<code>algorithm</code>	Character or <code>igraph cluster_*</code> function (length = 1). Defaults to "walktrap". Three options are listed below but all are available (see community.detection for other options): <ul style="list-style-type: none"> • "leiden" — See cluster_leiden for more details • "louvain" — By default, "louvain" will implement the Louvain algorithm using the consensus clustering method (see community.consensus for more information). This function will implement <code>consensus.method = "most_common"</code> and <code>consensus.iter = 1000</code> unless specified otherwise • "walktrap" — See cluster_walktrap for more details
<code>uni.method</code>	Character (length = 1). What unidimensionality method should be used? Defaults to "louvain". Available options:

- "expand" — Expands the correlation matrix with four variables correlated 0.50. 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 method was used in the Golino et al.'s (2020) *Psychological Methods* simulation
- "LE" — Applies the Leading Eigenvector algorithm (`cluster_leading_eigen`) on the empirical correlation matrix. If the number of dimensions is 1, then the Leading Eigenvector solution is used; otherwise, regular EGA is used. This method was used in the Christensen et al.'s (2023) *Behavior Research Methods* simulation
- "louvain" — Applies the Louvain algorithm (`cluster_louvain`) on the empirical correlation matrix. If the number of dimensions is 1, then the Louvain solution is used; otherwise, regular EGA is used. This method was validated Christensen's (2022) *PsyArXiv* simulation. Consensus clustering can be used by specifying either "consensus.method" or "consensus.iter"

iter	Numeric (length = 1). Number of replicate samples to be drawn from a multivariate normal distribution (uses MASS::mvrnorm). Defaults to 100 (recommended)
seed	Numeric (length = 1). Defaults to NULL or random results. Set for reproducible results. See Reproducibility and PRNG for more details on random number generation in EGAnet
verbose	Boolean (length = 1). Should progress be displayed? Defaults to TRUE. Set to FALSE to not display progress
...	Additional arguments that can be passed on to <code>auto.correlate</code> , <code>network.estimate</code> , <code>community.detection</code> , <code>community.consensus</code> , and EGA

Value

Returns a list containing:

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

Author(s)

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

References

Model training and validation

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

Examples

```
# Get data
data <- psych::bfi[,1:25]

## Not run: # Compute LCT
## Factor model
LCT(data)
## End(Not run)
```

`modularity`*Computes the (Signed) Modularity Statistic*

Description

Computes (signed) modularity statistic given a network and community structure. Allows the resolution parameter to be set

Usage

```
modularity(network, memberships, resolution = 1, signed = FALSE)
```

Arguments

<code>network</code>	Matrix or data frame. A symmetric matrix representing a network
<code>memberships</code>	Numeric (length = <code>ncol(network)</code>). A numeric vector of integer values corresponding to each node's community membership
<code>resolution</code>	Numeric (length = 1). A parameter that adjusts modularity to prefer smaller (<code>resolution > 1</code>) or larger (<code>0 < resolution < 1</code>) communities. Defaults to 1 (standard modularity computation)
<code>signed</code>	Boolean (length = 1). Whether signed or absolute modularity should be computed. The most common modularity metric is defined by positive values only. Gomez et al. (2009) introduced a signed version of modularity that will discount modularity for edges with negative values. This property isn't always desired for psychometric networks. If TRUE, then this signed modularity metric will be computed. If FALSE, then the absolute value of the edges in the network (using <code>abs</code>) will be used to compute modularity. Defaults to FALSE

Value

Returns the modularity statistic

Author(s)

Alexander P. Christensen <alexpaulchristensen@gmail.com> with assistance from GPT-4

References

Gomez, S., Jensen, P., & Arenas, A. (2009). Analysis of community structure in networks of correlated data. *Physical Review E*, 80(1), 016114.

Examples

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

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

# Compute standard (absolute values) modularity
modularity(
  network = ega.wmt$network,
  memberships = ega.wmt$wc,
  signed = FALSE
)
# 0.1697952

# Compute signed modularity
modularity(
  network = ega.wmt$network,
  memberships = ega.wmt$wc,
  signed = TRUE
)
# 0.1701946
```

net.loads

Network Loadings

Description

Computes the between- and within-community strength of each variable for each community

Usage

```
net.loads(
  A,
  wc,
  loading.method = c("original", "revised"),
  scaling = 2,
  rotation = NULL,
  ordered = c("descending", "variable"),
  ...
)
```

Arguments

A	Network matrix, data frame, or EGA object
wc	Numeric or character vector (length = ncol(A)). A vector of community assignments. If input into A is an EGA object, then wc is automatically detected
loading.method	Character (length = 1). Sets network loading calculation based on implementation described in "original" (Christensen & Golino, 2021) or the "revised" (Christensen et al., 2024) implementation. Defaults to "revised"
scaling	Numeric (length = 1). Scaling factor for the magnitude of the "experimental" network loadings. Defaults to 2. 10 makes loadings roughly the size of factor loadings when correlations between factors are orthogonal
rotation	Character (length = 1). A rotation to use to obtain a simpler structure. For a list of rotations, see rotations for options. Defaults to NULL or no rotation. By setting a rotation, scores estimation will be based on the rotated loadings rather than unrotated loadings
ordered	Character (length = 1). How the loadings should be ordered in the output. Available options: <ul style="list-style-type: none"> • "descending" (default) — Sorts loadings in descending order on their assigned community. This option is best for interpretation • "variable" — Keeps variables in the same order as the original network. This option is best for analyses
...	Additional arguments to pass on to rotations

Details

Simulation studies have demonstrated that a node's strength centrality is roughly equivalent to factor loadings (Christensen & Golino, 2021; Hallquist, Wright, & Molenaar, 2019). Hallquist and colleagues (2019) 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; Christensen & Golino, 2021).

Value

Returns a list containing:

unstd	A matrix of the unstandardized within- and between-community strength values for each node
std	A matrix of the standardized within- and between-community strength values for each node
rotated	NULL if rotation = NULL; otherwise, a list containing the rotated standardized network loadings (loadings) and correlations between dimensions (Phi) from the rotation

Author(s)

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

References

Original implementation and simulation

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

Demonstration of node strength similarity to CFA loadings

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.

Revised network loadings

Christensen, A. P., Golino, H., Abad, F. J., & Garrido, L. E. (2025). Revised network loadings. *Behavior Research Methods*, *57*, 114.

Examples

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

# Estimate EGA
ega.wmt <- EGA(
  data = wmt,
  plot.EGA = FALSE # No plot for CRAN checks
)

# 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 in the network (see [net.loads](#)). These values are used as "network loadings" for the weights of each variable.

Network scores are computed as a formative composite rather than a reflective factor. This composite representation is consistent with no latent factors that psychometric network theory proposes.

Scores can be computed as a "simple" structure, which is equivalent to a weighted sum scores or as a "full" structure, which is equivalent to an EFA approach. Conservatively, the "simple" structure approach is recommended until further validation

Usage

```
net.scores(
  data,
  A,
```

```

    wc,
    loading.method = c("original", "revised"),
    rotation = NULL,
    scores = c("Anderson", "Bartlett", "components", "Harman", "network", "tenBerge",
              "Thurstone"),
    loading.structure = c("simple", "full"),
    impute = c("mean", "median", "none"),
    ...
  )

```

Arguments

data	Matrix or data frame. Should consist only of variables to be used in the analysis
A	Network matrix, data frame, or EGA object
wc	Numeric or character vector (length = ncol(A)). A vector of community assignments. If input into A is an EGA object, then wc is automatically detected
loading.method	Character (length = 1). Sets network loading calculation based on implementation described in "original" (Christensen & Golino, 2021) or the "revised" (Christensen et al., 2024) implementation. Defaults to "revised"
rotation	Character. A rotation to use to obtain a simpler structure. For a list of rotations, see rotations for options. Defaults to NULL or no rotation. By setting a rotation, scores estimation will be based on the rotated loadings rather than unrotated loadings
scores	Character (length = 1). How should scores be estimated? Defaults to "network" for network scores. Set to other scoring methods which will be computed using factor.scores (see link for arguments and explanations for other methods)
loading.structure	Character (length = 1). Whether simple structure or the saturated loading matrix should be used when computing scores. Defaults to "simple" "simple" structure more closely mirrors sum scores and CFA; "full" structure more closely mirrors EFA Simple structure is the more "conservative" (established) approach and is therefore the default. Treat "full" as experimental as proper vetting and validation has not been established
impute	Character (length = 1). If there are any missing data, then imputation can be implemented. Available options: <ul style="list-style-type: none"> • "none" — Default. No imputation is performed • "mean" — The mean value of each variable is used to replace missing data for that variable • "median" — The median value of each variable is used to replace missing data for that variable
...	Additional arguments to be passed on to net.loads and factor.scores

Value

Returns a list containing:

scores	A list containing the standardized (std.scores) rotated (rot.scores) scores. If rotation = NULL, then rot.scores will be NULL
loadings	Output from net.load s

Author(s)

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

References**Original implementation and simulation for loadings**

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

Preliminary simulation for scores

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*.

Revised network loadings

Christensen, A. P., Golino, H., Abad, F. J., & Garrido, L. E. (2024). Revised network loadings. *PsyArXiv*.

Examples

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

# Estimate EGA
ega.wmt <- EGA(
  data = wmt,
  plot.EGA = FALSE # No plot for CRAN checks
)

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

network.compare

Compares Network Structures Using Permutation

Description

A permutation implementation to determine statistical significance of whether the network structures are different from one another

Usage

```
network.compare(
  base,
  comparison,
  corr = c("auto", "cor_auto", "pearson", "spearman"),
  na.data = c("pairwise", "listwise"),
  model = c("BGGM", "glasso", "TMFG"),
  iter = 1000,
  ncores,
  verbose = TRUE,
  seed = NULL,
  ...
)
```

Arguments

base	Matrix or data frame. Should consist only of variables to be used in the analysis. First dataset
comparison	Matrix or data frame. Should consist only of variables to be used in the analysis. Second dataset
corr	Character (length = 1). Method to compute correlations. Defaults to "auto". Available options: <ul style="list-style-type: none"> • "auto" — Automatically computes appropriate correlations for the data using Pearson's for continuous, polychoric for ordinal, tetrachoric for binary, and polyserial/biserial for ordinal/binary with continuous. To change the number of categories that are considered ordinal, use <code>ordinal.categories</code> (see polychoric.matrix for more details) • "cor_auto" — Uses <code>cor_auto</code> to compute correlations. Arguments can be passed along to the function • "pearson" — Pearson's correlation is computed for all variables regardless of categories • "spearman" — Spearman's rank-order correlation is computed for all variables regardless of categories <p>For other similarity measures, compute them first and input them into data with the sample size (n)</p>
na.data	Character (length = 1). How should missing data be handled? Defaults to "pairwise". Available options: <ul style="list-style-type: none"> • "pairwise" — Computes correlation for all available cases between two variables • "listwise" — Computes correlation for all complete cases in the dataset
model	Character (length = 1). Defaults to "glasso". Available options: <ul style="list-style-type: none"> • "BGGM" — Computes the Bayesian Gaussian Graphical Model. Set argument <code>ordinal.categories</code> to determine levels allowed for a variable to be considered ordinal. See <code>?BGGM::estimate</code> for more details

- "glasso" — Computes the GLASSO with EBIC model selection. See [EBICglasso.qgraph](#) for more details
- "TMFG" — Computes the TMFG method. See [TMFG](#) for more details

iter	Numeric (length = 1). Number of permutations to perform. Defaults to 1000 (recommended)
ncores	Numeric (length = 1). Number of cores to use in computing results. Defaults to <code>ceiling(parallel::detectCores() / 2)</code> or half of your computer's processing power. Set to 1 to not use parallel computing
verbose	Boolean (length = 1). Should progress be displayed? Defaults to TRUE. Set to FALSE to not display progress
seed	Numeric (length = 1). Defaults to NULL or random results. Set for reproducible results. See Reproducibility and PRNG for more details on random number generation in EGAnet
...	Additional arguments that can be passed on to auto.correlate , network.estimate , community.detection , community.consensus , EGA , and jsd

Value

Returns a list:

network	Data frame with row names of each measure, empirical value (<code>statistic</code>), and p -value based on the permutation test (<code>p.value</code>)
edges	List containing matrices of values for empirical values (<code>statistic</code>), p -values (<code>p.value</code>), and Benjamini-Hochberg corrected p -values (<code>p.adjusted</code>)

Author(s)

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

References**sF**

Ulitzsch, E., Khanna, S., Rhemtulla, M., & Domingue, B. W. (2023). A graph theory based similarity metric enables comparison of subpopulation psychometric networks. *Psychological Methods*.

Jensen-Shannon Similarity (1 - Distance)

De Domenico, M., Nicosia, V., Arenas, A., & Latora, V. (2015). Structural reducibility of multilayer networks. *Nature Communications*, 6(1), 1–9.

Total Network Strength

van Borkulo, C. D., van Bork, R., Boschloo, L., Kossakowski, J. J., Tio, P., Schoevers, R. A., Borsboom, D., & Waldorp, L. J. (2023). Comparing network structures on three aspects: A permutation test. *Psychological Methods*, 28(6), 1273–1285.

Examples

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

```

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

# Groups
group1 <- wmt[groups == 1,]
group2 <- wmt[groups == 2,]

## Not run: # Perform comparison
results <- network.compare(group1, group2)

# Print results
print(results)

# Plot edge differences
plot(results)
## End(Not run)

```

network.confusion *Confusion Matrix Metrics for Edge Comparison and Recovery*

Description

Computes many commonly used confusion matrix metrics

Usage

```

network.confusion(
  base,
  comparison,
  metric = c("all", "sen", "spec", "ppv", "npv", "fdr", "fom", "ba", "f1", "csi", "mcc"),
  full.names = FALSE
)

```

Arguments

base	Matrix or data frame. Network that will be treated as the "ground truth" such that a false positive represents an edge that is present in comparison but not in this network
comparison	Matrix or data frame. Network that will be treated as the estimator such that a false positive represents an edge that is present in this network but not in base
metric	Character vector. Defaults to "all" metrics. Available options: <ul style="list-style-type: none"> "all" — All available metrics (default) "sen" — Sensitivity (True Positive Rate):

$$\frac{TP}{TP + FN}$$

- "spec" — Specificity (True Negative Rate):

$$\frac{TN}{TN + FP}$$

- "ppv" — Positive Predictive Value (Precision):

$$\frac{TP}{TP + FP}$$

- "npv" — Negative Predictive Value:

$$\frac{TN}{TN + FN}$$

- "fdr" — False Discovery Rate:

$$1 - PPV = \frac{FP}{TP + FP}$$

- "fom" — False Omission Rate:

$$1 - NPV = \frac{FN}{TN + FN}$$

- "ba" — Balanced Accuracy:

$$\frac{\textit{Sensitivity} + \textit{Specificity}}{2}$$

- "f1" — F1 Score (harmonic mean of PPV and Sensitivity):

$$\frac{2TP}{2TP + FP + FN}$$

- "csi" — Critical Success Index (Jaccard / Threat Score):

$$\frac{TP}{TP + FP + FN}$$

- "mcc" — Matthews Correlation Coefficient:

$$\frac{TP \times TN - FP \times FN}{\sqrt{(TP + FP)(TP + FN)(TN + FP)(TN + FN)}}$$

`full.names` Boolean (length = 1). Whether full or abbreviated names should be used. Defaults to FALSE. Set to TRUE for full names

Value

Returns a named vector of confusion matrix metrics

Author(s)

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

Examples

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

# Set split
split <- sample(1:nrow(wmt), round(nrow(wmt) / 2))

# Estimate networks
split1 <- network. estimation(wmt[split,])
split2 <- network. estimation(wmt[-split,])

# Estimate metrics
network. confusion(split1, split2)
```

network. estimation *Apply a Network Estimation Method*

Description

General function to apply network estimation methods in [EGAnet](#)

Usage

```
network. estimation(
  data,
  n = NULL,
  corr = c("auto", "cor_ auto", "cosine", "pearson", "spearman"),
  na. data = c("pairwise", "listwise"),
  model = c("BGGM", "glasso", "TMFG"),
  network. only = TRUE,
  verbose = FALSE,
  ...
)
```

Arguments

data	Matrix or data frame. Should consist only of variables to be used in the analysis
n	Numeric (length = 1). Sample size if data provided is a correlation matrix
corr	Character (length = 1). Method to compute correlations. Defaults to "auto". Available options: <ul style="list-style-type: none"> "auto" — Automatically computes appropriate correlations for the data using Pearson's for continuous, polychoric for ordinal, tetrachoric for binary, and polyserial/biserial for ordinal/binary with continuous. To change the number of categories that are considered ordinal, use <code>ordinal. categories</code> (see polychoric. matrix for more details)

- "cor_auto" — Uses `cor_auto` to compute correlations. Arguments can be passed along to the function
- "cosine" — Uses `cosine` to compute cosine similarity
- "pearson" — Pearson's correlation is computed for all variables regardless of categories
- "spearman" — Spearman's rank-order correlation is computed for all variables regardless of categories

For other similarity measures, compute them first and input them into data with the sample size (n)

na.data	Character (length = 1). How should missing data be handled? Defaults to "pairwise". Available options: <ul style="list-style-type: none"> • "pairwise" — Computes correlation for all available cases between two variables • "listwise" — Computes correlation for all complete cases in the dataset
model	Character (length = 1). Defaults to "glasso". Available options: <ul style="list-style-type: none"> • "BGGM" — Computes the Bayesian Gaussian Graphical Model. Set argument <code>ordinal.categories</code> to determine levels allowed for a variable to be considered ordinal. See <code>?BGGM::estimate</code> for more details • "glasso" — Computes the GLASSO with EBIC model selection. See EBICglasso.qgraph for more details • "TMFG" — Computes the TMFG method. See TMFG for more details
network.only	Boolean (length = 1). Whether the network only should be output. Defaults to TRUE. Set to FALSE to obtain all output for the network estimation method
verbose	Boolean (length = 1). Whether messages and (insignificant) warnings should be output. Defaults to FALSE (silent calls). Set to TRUE to see all messages and warnings for every function call
...	Additional arguments to be passed on to <code>auto.correlate</code> and the different network estimation methods (see <code>model</code> for model specific details)

Value

Returns a matrix populated with a network from the input data

Author(s)

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

References**Graphical Least Absolute Shrinkage and Selection Operator (GLASSO)**

Friedman, J., Hastie, T., & Tibshirani, R. (2008). Sparse inverse covariance estimation with the graphical lasso. *Biostatistics*, 9(3), 432–441.

GLASSO with Extended Bayesian Information Criterion (EBICglasso)

Epskamp, S., & Fried, E. I. (2018). A tutorial on regularized partial correlation networks. *Psychological Methods*, 23(4), 617–634.

Bayesian Gaussian Graphical Model (BGGM)

Williams, D. R. (2021). Bayesian estimation for Gaussian graphical models: Structure learning, predictability, and network comparisons. *Multivariate Behavioral Research*, 56(2), 336–352.

Triangulated Maximally Filtered Graph (TMFG)

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

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

# EBICglasso (default for EGA functions)
glasso_network <- network.estimate(
  data = wmt, model = "glasso"
)

# TMFG
tmfg_network <- network.estimate(
  data = wmt, model = "TMFG"
)
```

network.fit

Traditional Fit Metrics for Networks

Description

Computes several traditional fit metrics for networks including

- chi-square (χ^2)
- root mean square error of approximation (RMSEA) with confidence intervals
- confirmatory fit index (CFI)
- Tucker-Lewis index (TLI)
- standardized root mean residual (SRMR)
- log-likelihood
- Akaike's information criterion (AIC)
- Bayesian information criterion (BIC)

Usage

```
network.fit(network, n, S, ci = 0.95)
```

Arguments

network	Matrix or data frame. A p by p square network matrix
n	Numeric (length = 1). Sample size
S	Matrix or data frame. A p by p square zero-order correlation matrix corresponding with the input network
ci	Numeric (length = 1). Confidence interval for RMSEA

Value

Returns a named vector of fit statistics

Author(s)

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

References

Epskamp, S., Rhemtulla, M., & Borsboom, D. (2017). Generalized network psychometrics: Combining network and latent variable models. *Psychometrika*, 82(4), 904–927.

Examples

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

# Obtain correlation matrix
S <- auto.correlate(wmt)

# EBICglasso (default for EGA functions)
glasso_network <- network.estimation(
  data = wmt, model = "glasso"
)

# Obtain fit (expects continuous variables!)
network.fit(network = glasso_network, n = nrow(wmt), S = S)
# Scaled metrics are not yet available for
# dichotomous or polytomous data!
```

network.predictability

Predict New Data based on Network

Description

General function to compute a network's predictive power on new data, following Haslbeck and Waldorp (2018) and Williams and Rodriguez (2022)

This implementation is different from the `predictability` in the `mgm` package (Haslbeck), which is based on (regularized) regression. This implementation uses the network directly, converting the partial correlations into an implied precision (inverse covariance) matrix. See **Details** for more information

Usage

```
network.predictability(network, original.data, newdata, ordinal.categories = 7)
```

Arguments

<code>network</code>	Matrix or data frame. A partial correlation network
<code>original.data</code>	Matrix or data frame. Must consist only of variables to be used to estimate the network. See Examples
<code>newdata</code>	Matrix or data frame. Must consist of the same variables in the same order as <code>original.data</code> . See Examples
<code>ordinal.categories</code>	Numeric (length = 1). <i>Up to</i> the number of categories <i>before</i> a variable is considered continuous. Defaults to 7 categories before 8 is considered continuous

Details

This implementation of network predictability proceeds in several steps with important assumptions:

1. Network was estimated using (partial) correlations (not regression like the `mgm` package!)
2. Original data that was used to estimate the network in 1. is necessary to apply the original scaling to the new data
3. (Linear) regression-like coefficients are obtained by reserve engineering the inverse covariance matrix using the network's partial correlations (i.e., by setting the diagonal of the network to -1 and computing the inverse of the opposite signed partial correlation matrix; see `EGAnet:::pcor2inv`)
4. Predicted values are obtained by matrix multiplying the new data with these coefficients
5. **Dichotomous and polytomous** data are given categorical values based on the **original data's** thresholds and these thresholds are used to convert the continuous predicted values into their corresponding categorical values
6. Evaluation metrics:
 - dichotomous — "Accuracy" or the percent correctly predicted for the 0s and 1s and "Kappa" or Cohen's Kappa (see cite)
 - polytomous — "Linear Kappa" or linearly weighted Kappa and "Krippendorff's alpha" (see cite)
 - continuous — R-squared ("R2") and root mean square error ("RMSE")

Value

Returns a list containing:

predictions	Predicted values of newdata based on the network
betas	Beta coefficients derived from the network
results	Performance metrics for each variable in newdata

Author(s)

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

References**Original Implementation of Node Predictability**

Haslbeck, J. M., & Waldorp, L. J. (2018). How well do network models predict observations? On the importance of predictability in network models. *Behavior Research Methods*, 50(2), 853–861.

Derivation of Regression Coefficients Used (Formula 3)

Williams, D. R., & Rodriguez, J. E. (2022). Why overfitting is not (usually) a problem in partial correlation networks. *Psychological Methods*, 27(5), 822–840.

Examples

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

# Set seed (to reproduce results)
set.seed(42)

# Split data
training <- sample(
  1:nrow(wmt), round(nrow(wmt) * 0.80) # 80/20 split
)

# Set splits
wmt_train <- wmt[training,]
wmt_test <- wmt[-training,]

# EBICglasso (default for EGA functions)
glasso_network <- network.estimation(
  data = wmt_train, model = "glasso"
)

# Check predictability
network.predictability(
  network = glasso_network, original.data = wmt_train,
  newdata = wmt_test
)
```

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).

Usage

```
data(optimism)
```

Format

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.

Examples

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

plot_clusters

Plot Clustered Individual Networks

Description

Visualize clusters of individual networks identified by [infoCluster](#) (or another clustering method) applied to a [dynEGA](#) object

Provides visualization modes:

- "population": Cluster-level "population" networks, obtained by stacking derivatives across individuals and estimating a new EGA
- "average": Cluster-average networks, obtained by averaging adjacency matrices across individuals in the cluster

Usage

```
plot_clusters(
  dynEGA.object,
  clustering,
  include,
  type = c("population", "average"),
  node.size = 3,
  label.size = 1.2,
  ...
)
```

Arguments

<code>dynEGA.object</code>	A dynEGA or a dynEGA.ind.pop object
<code>clustering</code>	Vector (length of individuals). A vector of cluster membership for each individual in the <code>dynEGA.object</code> . A common and easy option is to use an infoCluster object containing cluster assignments for individuals (see infoCluster). Accepts any vector of memberships
<code>include</code>	Numeric vector. Specifies which cluster memberships should be explicitly included in the plot. By default, all clusters are shown. Use this argument to restrict the visualization to a subset of clusters (e.g., <code>include = c(1, 3, 5)</code> will only display clusters 1, 3, and 5)
<code>type</code>	Character (length = 1). Type of visualization to produce: <ul style="list-style-type: none"> • "population" (default) — Cluster-level population networks by stacking derivatives across individuals in the cluster and re-estimating with EGA • "average" — Averaged networks with a consensus clustering matrix (pairwise) membership similarity that is supplied to community.detection In either type, the argument setting applied in your <code>'dynEGA.object'</code> will be carried forward into the network estimation ("population") and community detection (both)
<code>node.size</code>	Numeric (length = 1 or number of variables). Node size in network visualizations. Defaults to 3
<code>label.size</code>	Numeric (length = 1 or number of nodes). Label size in network visualizations. Defaults to 1.2
<code>...</code>	Additional arguments passed to EGA and compare.EGA.plots

Details

This function provides flexible visualization of individual clusters obtained from [infoCluster](#) (or other clustering input; see examples):

- Cluster-level population networks treat each cluster as a "mini-population," stacking derivatives across individuals to re-estimate an EGA
- Average networks show the mean connectivity structure for each cluster, with consensus clustering memberships

Function automatically extracts the appropriate derivatives ("population") and networks ("average") from the `dynEGA.object` results.

Value

Plots created using [compare.EGA.plots](#):

"population" A plot combining cluster-level population network plots
"average" A plot combining cluster-average network plots

Author(s)

Hudson Golino <hfg9s at virginia.edu>

See Also

[dynEGA](#), [infoCluster](#), [EGA](#), [compare.EGA.plots](#)

Examples

```
# Load data
data <- sim.dynEGA

## Not run:
# Run dynEGA with optimization
optimized_all <- dynEGA(
  data = data, level = c("individual", "population"),
  n.embed = 3:25, n.embed.optimize = TRUE
)

# Cluster individuals
clust <- infoCluster(dynEGA.object = optimized_all)

# Cluster-level population networks
plot_clusters(
  dynEGA.object = optimized_all,
  clustering = clust,
  type = "population"
)

# Average networks per cluster
plot_clusters(
  dynEGA.object = optimized_all,
  clustering = clust,
  type = "average"
)

# Cluster-level population networks, including only Cluster 2:
plot_clusters(
  dynEGA.object = optimized_all,
  clustering = clust, include = 2,
  type = "population"
)

# Using alternative clusters
```

```

plot_clusters(
  dynEGA.object = optimized_all,
  clustering = rep(1:2, each = 50), # vector of memberships
  type = "population"
)

# Run with non-optimized dynEGA
standard_all <- dynEGA(
  data = data,
  level = c("individual", "population")
)

# Obtain clusters
clust_standard <- infoCluster(dynEGA.object = standard_all)

# Plot clusters with population
plot_clusters(
  dynEGA.object = standard_all,
  clustering = clust_standard,
  type = "population"
)
## End(Not run)

```

polychoric.matrix

Computes Polychoric Correlations

Description

A fast implementation of polychoric correlations in C. Uses the Beasley-Springer-Moro algorithm (Boro & Springer, 1977; Moro, 1995) to estimate the inverse univariate normal CDF, the Drezner-Wesolosky approximation (Drezner & Wesolosky, 1990) to estimate the bivariate normal CDF, and Brent's method (Brent, 2013) for optimization of rho

Usage

```

polychoric.matrix(
  data,
  na.data = c("pairwise", "listwise"),
  empty.method = c("none", "zero", "all"),
  empty.value = c("none", "point_five", "one_over"),
  ...
)

```

Arguments

data	Matrix or data frame. A dataset with all ordinal values (rows = cases, columns = variables). Data are required to be between 0 and 11. Proper adjustments should be made prior to analysis (e.g., scales from -3 to 3 in increments of 1 should be shifted by added 4 to all values)
------	--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------

na.data	Character (length = 1). How should missing data be handled? Defaults to "pairwise". Available options: <ul style="list-style-type: none"> • "pairwise" — Computes correlation for all available cases between two variables • "listwise" — Computes correlation for all complete cases in the dataset
empty.method	Character (length = 1). Method for empty cell correction. Available options: <ul style="list-style-type: none"> • "none" — Adds no value (<code>empty.value = "none"</code>) to the empirical joint frequency table between two variables • "zero" — Adds <code>empty.value</code> to the cells with zero in the joint frequency table between two variables • "all" — Adds <code>empty.value</code> to all in the joint frequency table between two variables
empty.value	Character (length = 1). Value to add to the joint frequency table cells. Accepts numeric values between 0 and 1 or specific methods: <ul style="list-style-type: none"> • "none" — Adds no value (0) to the empirical joint frequency table between two variables • "point_five" — Adds 0.5 to the cells defined by <code>empty.method</code> • "one_over" — Adds $1/n$ where n equals the number of cells based on <code>empty.method</code>. For <code>empty.method = "zero"</code>, n equals the number of zero cells
...	Not used but made available for easier argument passing

Value

Returns a polychoric correlation matrix

Author(s)

Alexander P. Christensen <alexpaulchristensen@gmail.com> with assistance from GPT-4

References**Beasley-Moro-Springer algorithm**

Beasley, J. D., & Springer, S. G. (1977). Algorithm AS 111: The percentage points of the normal distribution. *Journal of the Royal Statistical Society. Series C (Applied Statistics)*, 26(1), 118-121.

Moro, B. (1995). The full monte. *Risk* 8 (February), 57-58.

Brent optimization

Brent, R. P. (2013). Algorithms for minimization without derivatives. Mineola, NY: Dover Publications, Inc.

Drezner-Wesolowsky bivariate normal approximation

Drezner, Z., & Wesolowsky, G. O. (1990). On the computation of the bivariate normal integral. *Journal of Statistical Computation and Simulation*, 35(1-2), 101-107.

Examples

```
# Load data (ensure matrix for missing data example)
wmt <- as.matrix(wmt2[,7:24])

# Compute polychoric correlation matrix
correlations <- polychoric.matrix(wmt)

# Randomly assign missing data
wmt[sample(1:length(wmt), 1000)] <- NA

# Compute polychoric correlation matrix
# with pairwise missing
pairwise_correlations <- polychoric.matrix(
  wmt, na.data = "pairwise"
)

# Compute polychoric correlation matrix
# with listwise missing
pairwise_correlations <- polychoric.matrix(
  wmt, na.data = "listwise"
)
```

prime.num

Prime Numbers through 100,000

Description

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

Usage

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

Format

A 1185x24 response matrix

Examples

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

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,
  corr = c("auto", "cor_auto", "cosine", "pearson", "spearman"),
  na.data = c("pairwise", "listwise"),
  model = c("glasso", "TMFG"),
  algorithm = c("leiden", "louvain", "walktrap"),
  uni.method = c("expand", "LE", "louvain"),
  plot.EGA = TRUE,
  verbose = FALSE,
  ...
)
```

Arguments

- | | |
|------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| data | Matrix or data frame. Should consist only of variables to be used in the analysis. Can be raw data or a correlation matrix.
<i>If a correlation matrix is input, then the correlations are assumed to be computed using the original keyed data (where reverse-keyed items remain reverse-keyed). Using correlations matrices where reverse-keyed items have been recoded will result in invalid results.</i> |
| n | Numeric (length = 1). Sample size if data provided is a correlation matrix |
| corr | Character (length = 1). Method to compute correlations. Defaults to "auto". Available options: <ul style="list-style-type: none"> • "auto" — Automatically computes appropriate correlations for the data using Pearson's for continuous, polychoric for ordinal, tetrachoric for binary, and polyserial/biserial for ordinal/binary with continuous. To change the number of categories that are considered ordinal, use <code>ordinal.categories</code> (see polychoric.matrix for more details) • "cor_auto" — Uses <code>cor_auto</code> to compute correlations. Arguments can be passed along to the function • "cosine" — Uses <code>cosine</code> to compute cosine similarity • "pearson" — Pearson's correlation is computed for all variables regardless of categories |

	<ul style="list-style-type: none"> • "spearman" — Spearman's rank-order correlation is computed for all variables regardless of categories
	For other similarity measures, compute them first and input them into data with the sample size (n)
na.data	Character (length = 1). How should missing data be handled? Defaults to "pairwise". Available options: <ul style="list-style-type: none"> • "pairwise" — Computes correlation for all available cases between two variables • "listwise" — Computes correlation for all complete cases in the dataset
model	Character (length = 1). Defaults to "glasso". Available options: <ul style="list-style-type: none"> • "BGM" — Computes the Bayesian Gaussian Graphical Model. Set argument <code>ordinal.categories</code> to determine levels allowed for a variable to be considered ordinal. See <code>?BGM::estimate</code> for more details • "glasso" — Computes the GLASSO with EBIC model selection. See EBICglasso.qgraph for more details • "TMFG" — Computes the TMFG method. See TMFG for more details
algorithm	Character or <code>igraph_cluster_*</code> function (length = 1). Defaults to "walktrap". Three options are listed below but all are available (see community.detection for other options): <ul style="list-style-type: none"> • "leiden" — See cluster_leiden for more details • "louvain" — By default, "louvain" will implement the Louvain algorithm using the consensus clustering method (see community.consensus for more information). This function will implement <code>consensus.method = "most_common"</code> and <code>consensus.iter = 1000</code> unless specified otherwise • "walktrap" — See cluster_walktrap for more details
uni.method	Character (length = 1). What unidimensionality method should be used? Defaults to "louvain". Available options: <ul style="list-style-type: none"> • "expand" — Expands the correlation matrix with four variables correlated 0.50. 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 method was used in the Golino et al.'s (2020) <i>Psychological Methods</i> simulation • "LE" — Applies the Leading Eigenvector algorithm (cluster_leading_eigen) on the empirical correlation matrix. If the number of dimensions is 1, then the Leading Eigenvector solution is used; otherwise, regular EGA is used. This method was used in the Christensen et al.'s (2023) <i>Behavior Research Methods</i> simulation • "louvain" — Applies the Louvain algorithm (cluster_louvain) on the empirical correlation matrix. If the number of dimensions is 1, then the Louvain solution is used; otherwise, regular EGA is used. This method was validated Christensen's (2022) <i>PsyArXiv</i> simulation. Consensus clustering can be used by specifying either <code>"consensus.method"</code> or <code>"consensus.iter"</code>
plot.EGA	Boolean (length = 1). If TRUE, returns a plot of the network and its estimated dimensions. Defaults to TRUE

verbose	Boolean (length = 1). Whether messages and (insignificant) warnings should be output. Defaults to FALSE (silent calls). Set to TRUE to see all messages and warnings for every function call
...	Additional arguments to be passed on to auto.correlate , network.estimate , community.detection , community.consensus , and EGA

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
TEFI	<code>link[EGAnet]{tefi}</code> for the estimated structure
<code>plot.EGA</code>	Plot output if <code>plot.EGA = TRUE</code>

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.

See Also

[plot.EGAnet](#) for plot usage in EGAnet

Examples

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

# riEGA example
riEGA(data = wmt, plot.EGA = FALSE)
# no plot for CRAN checks
```

sF

sF Similarity Metric

Description

Computes the sF norm based metric, s_F (Ulitzsch et al., 2023)

Usage

```
sF(network1, network2)
```

Arguments

network1	Matrix or data frame. Network to be compared
network2	Matrix or data frame. Second network to be compared

Value

Returns sF

Author(s)

Hudson Golino <hfg9s at virginia.edu> & Alexander P. Christensen <alexander.christensen at Vanderbilt.Edu>

References

Simulation Study

Ulitzsch, E., Khanna, S., Rhemtulla, M., & Domingue, B. W. (2023). A graph theory based similarity metric enables comparison of subpopulation psychometric networks *Psychological Methods*.

Examples

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

# Set seed (for reproducibility)
set.seed(1234)

# Split data
split1 <- sample(
  1:nrow(wmt), floor(nrow(wmt) / 2)
)
split2 <- setdiff(1:nrow(wmt), split1)

# Obtain split data
data1 <- wmt[split1,]
data2 <- wmt[split2,]
```

```
# Perform EBICglasso
glas1 <- EBICglasso.qgraph(data1)
glas2 <- EBICglasso.qgraph(data2)

# sF
sF(glas1, glas2)
# 0.7070395
```

sim.dynEGA

sim.dynEGA Data

Description

A simulated (multivariate time series) data with 24 variables, 100 individual observations, 50 time points per individual and 2 groups of individuals

Usage

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

Format

A 5000 x 26 multivariate time series

Details

Data were generated using the [simDFM](#) function with the following arguments:

Group 1

```
simDFM( variab = 12, timep = 50, nfact = 2, error = 0.125, dfm = "DAFS", loadings = EGAnet:::runif_xoshiro(
1, min = 0.50, max = 0.70 ), autoreg = 0.80, crossreg = 0.00, var.shock = 0.36, cov.shock
= 0.18 )
```

Group 2

```
simDFM( variab = 8, timep = 50, nfact = 3, error = 0.125, dfm = "DAFS", loadings = EGAnet:::runif_xoshiro(
1, min = 0.50, max = 0.70 ), autoreg = 0.80, crossreg = 0.00, var.shock = 0.36, cov.shock
= 0.18 )
```

Examples

```
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:  
# 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 = 0.36,  
cov.shock = 0.18, burnin = 1000)  
## End(Not run)
```

simEGM

Simulate data following a Exploratory Graph Model (EGM)

Description

Function to simulate data based on EGM

Usage

```
simEGM(  
  communities,  
  variables,  
  loadings,  
  cross.loadings = 0.01,  
  correlations,  
  sample.size,  
  quality = c("acceptable", "robust"),  
  max.iterations = 100  
)
```

Arguments

<code>communities</code>	Numeric (length = 1). Number of communities to generate
<code>variables</code>	Numeric vector (length = 1 or <code>communities</code>). Number of variables per community
<code>loadings</code>	Numeric (length = 1, <code>communities</code> , or <code>total variables × communities</code>). Magnitude of the assigned network loadings. For reference, small (0.20), moderate (0.35), and large (0.50). Input can be a loading matrix but must have the dimensions: <code>total variables × communities</code> Uses <code>runif(n, min = value - 0.025, max = value + 0.025)</code> for some jitter in the loadings
<code>cross.loadings</code>	Numeric (length = 1). Standard deviation of a normal distribution with a mean of zero (<code>n, mean = 0, sd = value</code>). Defaults to 0.01. Not recommended to change too drastically (small increments such as 0.01 work best)
<code>correlations</code>	Numeric (length = 1 or <code>communities × communities</code> matrix). Magnitude of the community correlations. Input can be a correlations matrix but must have the dimensions: <code>communities × communities</code>
<code>sample.size</code>	Numeric (length = 1). Number of observations to generate
<code>quality</code>	Character (length = 1). Quality metrics related to the alignment of the correlations implied by the loadings and network are computed with certain standards in place to accept a solution. These metrics include: <ul style="list-style-type: none"> • SRMR (or RMSE) — standardized root mean residual where acceptable equals 0.02 and robust equals 0.01 • MAE — mean absolute error where acceptable equals 0.02 and robust equals 0.01 • sF — sF where acceptable equals 0.90 and robust equals 0.95 • jsd — Jensen-Shannon Distance where acceptable equals 0.10 and robust equals 0.05 Defaults to "acceptable". "robust" is available but most often needs <code>density.power</code> to be increased to allow for more cross-loadings to converge
<code>max.iterations</code>	Numeric (length = 1). Number of iterations to attempt to get convergence before erroring out. Defaults to 100

Author(s)

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

Examples

```
simulated <- simEGM(
  communities = 2, variables = 6,
  loadings = 0.35, # use network loading sizes
  correlations = 0.30, sample.size = 1000
)
```

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

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 = NULL, verbose = TRUE)
```

Arguments

data	Matrix, data frame, or *EGA class object. Matrix or data frame can be raw data or a correlation matrix. All *EGA objects are accepted. hierEGA input will produced the Generalized TEFI (see genTEFI)
structure	Numeric or character vector (length = ncol(data)). Can be theoretical factors or the structure detected by EGA
verbose	Boolean (length = 1). Whether messages and (insignificant) warnings should be output. Defaults to TRUE to see all messages and warnings for every function call. Set to FALSE to ignore messages and warnings

Value

Returns a data frame with columns:

Non-hierarchical Structure

VN.Entropy.Fit The Total Entropy Fit Index using Von Neumman's entropy

Total.Correlation

The total correlation of the dataset

Average.Entropy

The average entropy of the dataset

Hierarchical Structure

VN.Entropy.Fit The Generalized Total Entropy Fit Index using Von Neumman's entropy

Level_#_VN An individual level's Von Neumann's entropy

Author(s)

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

References

Initial formalization and simulation

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*.

Examples

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

# Estimate EGA model
ega.wmt <- EGA(
  data = wmt, model = "glasso",
  plot.EGA = FALSE # no plot for CRAN checks
)

# Compute entropy indices for empirical EGA
tefi(ega.wmt)

# User-defined structure (with `EGA` object)
tefi(ega.wmt, structure = c(rep(1, 5), rep(2, 5), rep(3, 8)))
```

tefi.compare

Compare Total Entropy Fit Index (tefi) Between Two Structures

Description

This function computes the [tefi](#) values for two different structures using bootstrapped correlation matrices from [bootEGA](#) and compares them using a non-parametric bootstrap test. It also visualizes the distributions of [tefi](#) values for both structures.

Usage

```
tefi.compare(bootega.obj, base, comparison, plot.TEFI = TRUE, ...)
```

Arguments

<code>bootega.obj</code>	A bootEGA object
<code>base</code>	Numeric (length = columns in original dataset). A vector representing the base structure to be tested
<code>comparison</code>	Numeric (length = columns in original dataset). A vector representing the structure to be compared against the base structure
<code>plot.TEFI</code>	Boolean (length = 1). Whether the TEFI comparison and the p-value should be plotted. Defaults to TRUE
<code>...</code>	Additional arguments that can be passed on to plot.EGAnet . See Examples for plotting arguments

Details

The null hypothesis is that the TEFI values obtained in the bootstrapped correlation matrices for the base structure are than the TEFI values obtained in the bootstrapped correlation matrices for the comparison structure. Therefore, the p -value in this bootstrap test can be interpreted as follows:

- If the p -value less than 0.05: TEFI values for the base structure tend to be lower than the comparison structure, indicating that the former provides a better fit (lower entropy) than the latter
- If the p -value is greater than 0.05: TEFI values for the base structure are not significantly lower than the comparison structure, suggesting that both structures may provide similar fits or that comparison might fit better

Value

A list containing:

TEFI.df	A data frame containing the TEFI values for both structures
p.value	The p -value from the non-parametric bootstrap hypothesis test

Author(s)

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

Examples

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

## Not run:
# Perform bootstrap EGA
boot.wmt <- bootEGA(
  data = wmt, iter = 500,
  type = "parametric", ncores = 2
)
## End(Not run)

# Perform comparison
comparing_tefi <- tefi.compare(
  boot.wmt,
  base = boot.wmt$EGA$wc, # Compare Walktrap
  comparison = community.detection(
    boot.wmt$EGA$network, algorithm = "louvain"
  ) # With Louvain
)

# Plot options (UVa colors)
plot(
  comparing_tefi,
  base.name = "Walktrap", base.color = "#232D4B",
  comparison.name = "Louvain", comparison.color = "#E57200"
)
```

Description

Applies the Triangulated Maximally Filtered Graph (TMFG) filtering method (see 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 (LoGo; see Barfuss et al., 2016 for more details). See **Details** for more information

Usage

```
TMFG(
  data,
  n = NULL,
  corr = c("auto", "cor_auto", "cosine", "pearson", "spearman"),
  na.data = c("pairwise", "listwise"),
  partial = FALSE,
  returnAllResults = FALSE,
  verbose = FALSE,
  ...
)
```

Arguments

data	Matrix or data frame. Should consist only of variables to be used in the analysis. Can be raw data or correlation matrix
n	Numeric (length = 1). Sample size for when a correlation matrix is input into data. Defaults to NULL. n is not necessary and is provided for better functionality in EGAnet
corr	Character (length = 1). Method to compute correlations. Defaults to "auto". Available options: <ul style="list-style-type: none"> "auto" — Automatically computes appropriate correlations for the data using Pearson's for continuous, polychoric for ordinal, tetrachoric for binary, and polyserial/biserial for ordinal/binary with continuous. To change the number of categories that are considered ordinal, use <code>ordinal.categories</code> (see polychoric.matrix for more details) "cor_auto" — Uses <code>cor_auto</code> to compute correlations. Arguments can be passed along to the function

- "cosine" — Uses [cosine](#) to compute cosine similarity
- "pearson" — Pearson's correlation is computed for all variables regardless of categories
- "spearman" — Spearman's rank-order correlation is computed for all variables regardless of categories

For other similarity measures, compute them first and input them into data with the sample size (n)

na.data	Character (length = 1). How should missing data be handled? Defaults to "pairwise". Available options: <ul style="list-style-type: none"> • "pairwise" — Computes correlation for all available cases between two variables • "listwise" — Computes correlation for all complete cases in the dataset
partial	Boolean (length = 1). Whether partial correlations should be output. Defaults to FALSE. The TMFG method is based on the zero-order correlations; the Local-Global Inversion Method (LoGo; see Barfuss et al., 2016 for more details) uses the decomposability of the TMFG network to obtain the inverse covariance structure of the network (which is then converted to partial correlations). Set to TRUE to obtain the partial correlations from the LoGo method
returnAllResults	Boolean (length = 1). Whether all results should be returned. Defaults to FALSE (network only). Set to TRUE to access separators and cliques
verbose	Boolean (length = 1). Whether messages and (insignificant) warnings should be output. Defaults to FALSE (silent calls). Set to TRUE to see all messages and warnings for every function call
...	Additional arguments to be passed on to auto.correlate

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 network or list containing:

network	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**Local-Global Inversion Method**

Barfuss, W., Massara, G. P., Di Matteo, T., & Aste, T. (2016). Parsimonious modeling with information filtering networks. *Physical Review E*, *94*, 062306.

Psychometric network introduction to TMFG

Christensen, A. P., Kenett, Y. N., Aste, T., Silvia, P. J., & Kwapil, T. R. (2018). Network structure of the Wisconsin Schizotypy Scales-Short Forms: Examining psychometric network filtering approaches. *Behavior Research Methods*, *50*, 2531-2550.

Triangulated Maximally Filtered Graph

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

```
# TMFG filtered network
TMFG(wmt2[,7:24])

# Partial correlations using the LoGo method
TMFG(wmt2[,7:24], partial = TRUE)
```

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

Description

Computes the total correlation of a dataset

Usage

```
totalCor(data, base = 2.718282)
```

Arguments

data	Matrix or data frame. Should consist only of variables to be used in the analysis
base	Numeric (length = 1). Base to use for entropy. Defaults to exp(1) or 2.718282

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
Normalized	Total correlation divided by the sum of the individual entropies minus the maximum of the individual entropies

Author(s)

Hudson F. Golino <hfg9s at virginia.edu>

References**Formalization of total correlation**

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

Applied 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 ([totalCor](#)) for a dataset

Usage

```
totalCorMat(data, base = 2.718282, normalized = FALSE)
```

Arguments

data	Matrix or data frame. Should consist only of variables to be used in the analysis
base	Numeric (length = 1). Base to use for entropy. Defaults to $\exp(1)$ or 2.718282
normalized	Boolean (length = 1). Should the normalized total correlation be computed? Defaults to FALSE

Value

Returns a symmetric matrix with pairwise total correlations

Author(s)

Hudson F. Golino <hfg9s at virginia.edu>

References**Formalization of total correlation**

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

Applied 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 matrix
totalCorMat(wmt2[,7:24])
```

UVA

Unique Variable Analysis

Description

Identifies locally dependent (redundant) variables in a multivariate dataset using the [EBICglasso.qgraph](#) network estimation method and weighted topological overlap (see Christensen, Garrido, & Golino, 2023 for more details)

Usage

```
UVA(  
  data = NULL,  
  network = NULL,  
  n = NULL,  
  key = NULL,  
  uva.method = c("MBR", "EJP"),  
  cut.off = 0.25,  
  reduce = TRUE,  
  reduce.method = c("latent", "mean", "remove", "sum"),  
  auto = TRUE,  
  verbose = FALSE,  
  ...  
)
```

Arguments

data	Matrix or data frame. Should consist only of variables to be used in the analysis. Can be raw data or a correlation matrix. Defaults to NULL
network	Symmetric matrix or data frame. A symmetric network. Defaults to NULL If both data and network are provided, then UVA will use the network with the data (rather than estimating a network from the data)
n	Numeric (length = 1). Sample size if data provided is a correlation matrix. Defaults to NULL
key	Character vector (length = ncol(data)). Item key for labeling variables in the results
uva.method	Character (length = 1). Whether the method described in Christensen, Garrido, and Golino (2023) publication in <i>Multivariate Behavioral Research</i> ("MBR") or Christensen, Golino, and Silvia (2020) publication in <i>European Journal of Personality</i> ("EJP") should be used. Defaults to "MBR" Based on simulation and accumulating empirical evidence, the methods described in Christensen, Golino, and Silvia (2020) such as adaptive alpha are outdated . Evidence supports using a single cut-off value (regardless of continuous, polytomous, or dichotomous data; Christensen, Garrido, & Golino, 2023)
cut.off	Numeric (length = 1). Cut-off used to determine when pairwise <i>wto</i> values are considered locally dependent (or redundant). Must be values between 0 and 1. Defaults to 0.25 This cut-off value is recommended and based on extensive simulation (Christensen, Garrido, & Golino, 2023). Printing the result will provide a gradient of pairwise redundancies in increments of 0.20, 0.25, and 0.30. Use <code>print</code> or <code>summary</code> on the output rather than adjusting this cut-off value
reduce	Logical (length = 1). Whether redundancies should be reduced in data. Defaults to TRUE
reduce.method	Character (length = 1). Method to reduce redundancies. Available options: <ul style="list-style-type: none"> • "latent" — Computes latent variables using <code>cfa</code> when there are three or more redundant variables. If variables are not all coded in the same direction, then they will be recoded as necessary. A warning will be produced for all variables that are flipped • "mean" — Computes mean of redundant variables. If variables are not all coded in the same direction, then they will be recoded as necessary. A warning will be produced for all variables that are flipped • "remove" — Removes all but one variable from a set of redundant variables • "sum" — Computes sum of redundant variables. If variables are not all coded in the same direction, then they will be recoded as necessary. A warning will be produced for all variables that are flipped
auto	Logical (length = 1). Whether reduce should occur automatically. For <code>reduce.method = "remove"</code> , the automated decision process is as follows: <ul style="list-style-type: none"> • Two variables — The variable with the lowest maximum <i>wto</i> to all other variables (other than the one it is redundant with) is retained and the other is removed

- Three or more variables — The variable with the highest mean `wto` to all other variables that are redundant with one another is retained and all others are removed
- verbose Boolean (length = 1). Whether messages and (insignificant) warnings should be output. Defaults to FALSE (silent calls). Set to TRUE to see all messages and warnings for every function call
- ... Additional arguments that should be passed on to old versions of UVA or to [EGA](#) and [cfa](#)

References

Most recent simulation and implementation

Christensen, A. P., Garrido, L. E., & Golino, H. (2023). Unique variable analysis: A network psychometrics method to detect local dependence. *Multivariate Behavioral Research*.

Conceptual foundation and outdated methods

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.

Weighted topological overlap

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(3), 354-373.

Examples

```
# Perform UVA
uva.wmt <- UVA(wmt2[,7:24])

# Show summary
summary(uva.wmt)
```

vn.entropy

Entropy Fit Index using Von Neumann's entropy (Quantum Information Theory) for correlation matrices

Description

Computes the fit of a dimensionality structure using Von Neumann'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	Matrix or data frame. Contains variables to be used in the analysis
structure	Numeric or character vector (length = ncol(data)). 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 Neumann's entropy

Total.Correlation
The total correlation of the dataset

Average.Entropy
The average entropy of the dataset

Author(s)

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

References**Initial formalization and simulation**

Golino, H., Moulder, R. G., Shi, D., Christensen, A. P., Garrido, L. E., Nieto, M. D., Nesselrode, 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*.

Examples

```
# Get EGA result
ega.wmt <- EGA(
  data = wmt2[,7:24], model = "glasso",
  plot.EGA = FALSE # no plot for CRAN checks
)

# Compute Von Neumann entropy
vn.entropy(ega.wmt$correlation, ega.wmt$wc)
```

wmt2

WMT-2 Data

Description

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

Usage

```
data(wmt2)
```

Format

A 1185x24 response matrix

Examples

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

wto

Weighted Topological Overlap

Description

Computes weighted topological overlap following the Novick et al. (2009) definition

Usage

```
wto(network, signed = TRUE, diagonal.zero = TRUE)
```

Arguments

network	Symmetric matrix or data frame. A symmetric network
signed	Boolean (length = 1). Whether the signed version should be used. Defaults to TRUE. Use FALSE for absolute values
diagonal.zero	Boolean (length = 1). Whether diagonal of overlap matrix should be set to zero. Defaults to TRUE. Use FALSE to allow overlap of a node with itself

Value

A symmetric matrix of weighted topological overlap values between each pair of variables

References

Original formalization

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.

Examples

```
# Obtain network
network <- network. estimation(wmt2[,7:24], model = "glasso")

# Compute wTO
wto(network)
```

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