

# Package ‘EFAfactors’

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**Type** Package

**Title** Determining the Number of Factors in Exploratory Factor Analysis

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**Description** Provides a collection of standard factor retention methods in Exploratory Factor Analysis (EFA), making it easier to determine the number of factors. Traditional methods such as the scree plot by Cattell (1966) <[doi:10.1207/s15327906mbr0102\\_10](https://doi.org/10.1207/s15327906mbr0102_10)>, Kaiser-Guttman Criterion (KGC) by Guttman (1954) <[doi:10.1007/BF02289162](https://doi.org/10.1007/BF02289162)> and Kaiser (1960) <[doi:10.1177/001316446002000116](https://doi.org/10.1177/001316446002000116)>, and Parallel Analysis (PA) by Horn (1965) <[doi:10.1007/BF02289447](https://doi.org/10.1007/BF02289447)> based on eigenvalues from PCA or EFA are readily available. This package also implements several newer methods, such as the Empirical Kaiser Criterion (EKC) by Braeken and van Assema (2017) <[doi:10.1037/met0000074](https://doi.org/10.1037/met0000074)>, Comparison Data (CD) by Ruscio and Roche (2012) <[doi:10.1037/a0025697](https://doi.org/10.1037/a0025697)>, and Hull method by Lorenzo-Seva et al. (2011) <[doi:10.1080/00273171.2011.564527](https://doi.org/10.1080/00273171.2011.564527)>, as well as some AI-based methods like Comparison Data Forest (CDF) by Goretzko and Ruscio (2024) <[doi:10.3758/s13428-023-02122-4](https://doi.org/10.3758/s13428-023-02122-4)> and Factor Forest (FF) by Goretzko and Buhner (2020) <[doi:10.1037/met0000262](https://doi.org/10.1037/met0000262)>. Additionally, it includes a deep neural network (DNN) trained on large-scale datasets that can efficiently and reliably determine the number of factors.

**License** GPL-3

**Depends** R (>= 4.1.0)

**Imports** BBmisc, checkmate, ddpccr, ineq, MASS, Matrix, mlr, proxy, psych, ranger, reticulate, Rcpp, RcppArmadillo, SimCorMultRes, xgboost

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**Collate** 'CD.R' 'CDF.R' 'check\_python\_libraries.R' 'data.bfi.R'  
 'data.datasets.R' 'data.scaler.R' 'DNN\_predictor.R'  
 'EFAhclust.R' 'EFAindex.R' 'EFAkmeans.R' 'EFAvote.R' 'EKC.R'  
 'EFAscree.R' 'EFAsim.data.R' 'extractor.feature.DNN.R'  
 'extractor.feature.FF.R' 'factor.analysis.R' 'FF.R' 'GenData.R'  
 'get.runs.R' 'Hull.R' 'KGC.R' 'load.R' 'model.xgb.R'  
 'normalizor.R' 'PA.R' 'ParamHelpers.R' 'plot.R' 'print.R'  
 'RcppExports.R' 'af.softmax.R' 'utils.R' 'zzz.R'

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## Contents

af.softmax . . . . .	3
CD . . . . .	4
CDF . . . . .	6
check_python_libraries . . . . .	9
data.bfi . . . . .	10
data.datasets . . . . .	12
data.scaler . . . . .	12
DNN_predictor . . . . .	13
EFAhclust . . . . .	16
EFAindex . . . . .	19
EFAkmeans . . . . .	20
EFAscree . . . . .	22
EFAsim.data . . . . .	24
EFAvote . . . . .	26
EKC . . . . .	27
extractor.feature.DNN . . . . .	30
extractor.feature.FF . . . . .	32
factor.analysis . . . . .	34
FF . . . . .	36
GenData . . . . .	40
Hull . . . . .	42
KGC . . . . .	44
load_DNN . . . . .	46
load_scaler . . . . .	47
load_xgb . . . . .	48
model.xgb . . . . .	49
normalizor . . . . .	49
PA . . . . .	50
plot.CD . . . . .	53
plot.CDF . . . . .	54
plot.DNN_predictor . . . . .	55

<i>af.softmax</i>	3
plot.EFAhclust . . . . .	56
plot.EFAkmeans . . . . .	57
plot.EFAscreet . . . . .	58
plot.EFAvote . . . . .	59
plot.EKC . . . . .	60
plot.FF . . . . .	61
plot.Hull . . . . .	62
plot.KGC . . . . .	63
plot.PA . . . . .	64
predictLearner.classif.xgboost.earlystop . . . . .	65
print.CD . . . . .	66
print.CDF . . . . .	66
print.DNN_predictor . . . . .	67
print.EFAdata . . . . .	68
print.EFAhclust . . . . .	68
print.EFAkmeans . . . . .	69
print.EFAscreet . . . . .	69
print.EFAvote . . . . .	70
print.EKC . . . . .	70
print.FF . . . . .	71
print.Hull . . . . .	71
print.KGC . . . . .	72
print.PA . . . . .	73
<b>Index</b>	<b>74</b>

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<i>af.softmax</i>	<i>An Activation Function: Softmax</i>
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**Description**

This function computes the softmax of a numeric vector. The softmax function transforms a vector of real values into a probability distribution, where each element is between 0 and 1 and the sum of all elements is 1. @seealso [DNN\\_predictor](#)

**Usage**

`af.softmax(x)`

**Arguments**

`x`                      A numeric vector for which the softmax transformation is to be computed.

### Details

The softmax function is calculated as:

$$\text{softmax}(x_i) = \frac{\exp(x_i)}{\sum_j \exp(x_j)}$$

In the case of overflow (i.e., when  $\exp(x_i)$  is too large), this function handles Inf values by assigning 1 to the corresponding positions and 0 to the others before Softmax. @seealso [DNN\\_predictor](#)

### Value

A numeric vector representing the softmax-transformed values of x.

### Examples

```
x <- c(1, 2, 3)
af.softmax(x)
```

---

CD	<i>the Comparison Data (CD) Approach</i>
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### Description

This function runs the comparison data (CD) approach of Ruscio & Roche (2012).

### Usage

```
CD(
  response,
  nfact.max = 10,
  N.pop = 10000,
  N.Samples = 500,
  Alpha = 0.3,
  cor.type = "pearson",
  use = "pairwise.complete.obs",
  vis = TRUE,
  plot = TRUE
)
```

### Arguments

response	A required $N \times I$ matrix or data.frame consisting of the responses of N individuals to $\times I$ items.
nfact.max	The maximum number of factors discussed by CD approach. (default = 10)
N.pop	Size of finite populations of simulating.. (default = 10,000)
N.Samples	Number of samples drawn from each population. (default = 500)

Alpha	Alpha level when testing statistical significance (Wilcoxon Rank Sum and Signed Rank Tests) of improvement with additional factor. (default = .30)
cor.type	A character string indicating which correlation coefficient (or covariance) is to be computed. One of "pearson" (default), "kendall", or "spearman". @seealso <a href="#">cor</a> .
use	an optional character string giving a method for computing covariances in the presence of missing values. This must be one of the strings "everything", "all.obs", "complete.obs", "na.or.complete", or "pairwise.complete.obs" (default). @seealso <a href="#">cor</a> .
vis	A Boolean variable that will print the factor retention results when set to TRUE, and will not print when set to FALSE. (default = TRUE)
plot	A Boolean variable that will print the CD plot when set to TRUE, and will not print it when set to FALSE. @seealso <a href="#">plot.CD</a> . (Default = TRUE)

### Details

Ruscio and Roche (2012) proposed a method for determining the number of factors through comparison data (CD). This method identifies the appropriate number of factors by finding the solution that best reproduces the pattern of eigenvalues. CD employs an iterative procedure when generating comparison data with a known factor structure, taking into account previous factors. Initially, CD compares whether the simulated comparison data with one latent factor ( $j=1$ ) reproduces the empirical eigenvalue pattern significantly worse than the two-factor solution ( $j+1$ ). If so, CD increases the value of  $j$  until further improvements are no longer significant or a preset maximum number of factors is reached. Specifically, CD involves five steps:

1. Generate random data with either  $j$  or  $j+1$  latent factors and calculate the eigenvalues of the respective correlation matrices.
2. Compute the root mean square error (RMSE) of the difference between the empirical and simulated eigenvalues using the formula

$$RMSE = \sqrt{\sum_{i=1}^p (\lambda_{emp,i} - \lambda_{sim,i})^2}$$

, where:

- $\lambda_{emp,i}$ : The  $i$ -th empirical eigenvalue.
- $\lambda_{sim,i}$ : The  $i$ -th simulated eigenvalue.
- $p$ : The number of items or eigenvalues.

. This step produces two RMSEs, corresponding to the different numbers of latent factors.

3. Repeat steps 1 and 2, 500 times ( default in the Package ).
4. Use a one-sided Wilcoxon test ( $\alpha = 0.30$ ) to assess whether the RMSE is significantly reduced under the two-factor condition.
5. If the difference in RMSE is not significant, CD suggests selecting  $j$  factors. Otherwise,  $j$  is increased by 1, and steps 1 to 4 are repeated.

The code is implemented based on the resources available at:

- <https://ruscio.pages.tcnj.edu/quantitative-methods-program-code/>
- [https://osf.io/gqma2/?view\\_only=d03efba1fd0f4c849a87db82e6705668](https://osf.io/gqma2/?view_only=d03efba1fd0f4c849a87db82e6705668)
- <https://osf.io/mvrau/>

Since the CD approach requires extensive data simulation and computation, C++ code is used to speed up the process.

**Value**

An object of class CD is a list containing the following components:

nfact	The number of factors to be retained.
RMSE.Eigs	A matrix containing the root mean square error (RMSE) of the eigenvalues produced by each simulation for every discussed number of factors.
Sig	A boolean variable indicating whether the significance level of the Wilcoxon Rank Sum and Signed Rank Tests has reached Alpha.

**Author(s)**

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**References**

Auerswald, M., & Moshagen, M. (2019). How to determine the number of factors to retain in exploratory factor analysis: A comparison of extraction methods under realistic conditions. *Psychological methods*, 24(4), 468-491. <https://doi.org/https://doi.org/10.1037/met0000200>.

Goretzko, D., & Buhner, M. (2020). One model to rule them all? Using machine learning algorithms to determine the number of factors in exploratory factor analysis. *Psychol Methods*, 25(6), 776-786. <https://doi.org/10.1037/met0000262>.

Ruscio, J., & Roche, B. (2012). Determining the number of factors to retain in an exploratory factor analysis using comparison data of known factorial structure. *Psychological Assessment*, 24, 282–292. <http://dx.doi.org/10.1037/a0025697>.

**See Also**

[GenData](#)

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CDF	<i>the Comparison Data Forest (CDF) Approach</i>
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**Description**

The Comparison Data Forest (CDF; Goretzko & Ruscio, 2019) approach is a combination of Random Forest with the comparison data (CD) approach.

**Usage**

```
CDF(
  response,
  num.trees = 500,
  mtry = "sqrt",
  nfact.max = 10,
  N.pop = 10000,
  N.Samples = 500,
  cor.type = "pearson",
  use = "pairwise.complete.obs",
  vis = TRUE,
  plot = TRUE
)
```

**Arguments**

response	A required $N \times I$ matrix or data.frame consisting of the responses of $N$ individuals to $\times I$ items.
num.trees	the number of trees in the Random Forest. (default = 500) See details.
mtry	the maximum depth for each tree, can be a number or a character ("sqrt"). When <code>mtry = "sqrt"</code> , it means that the maximum depth of each tree will be determined by the square root of the number of available features (converted to an integer by <a href="#">round</a> ).default = "sqrt". See details.
nfact.max	The maximum number of factors discussed by CD approach. (default = 10)
N.pop	Size of finite populations of simulating.. (default = 10,000)
N.Samples	Number of samples drawn from each population. (default = 500)
cor.type	A character string indicating which correlation coefficient (or covariance) is to be computed. One of "pearson" (default), "kendall", or "spearman". @seealso <a href="#">cor</a> .
use	an optional character string giving a method for computing covariances in the presence of missing values. This must be one of the strings "everything", "all.obs", "complete.obs", "na.or.complete", or "pairwise.complete.obs" (default). @seealso <a href="#">cor</a> .
vis	A Boolean variable that will print the factor retention results when set to TRUE, and will not print when set to FALSE. (default = TRUE)
plot	A Boolean variable that will print the CDF plot when set to TRUE, and will not print it when set to FALSE. @seealso <a href="#">plot.CDF</a> . (Default = TRUE)

**Details**

The Comparison Data Forest (CDF; Goretzko & Ruscio, 2019) Approach is a combination of random forest with the comparison data (CD) approach. Its basic steps involve using the method of Ruscio & Roche (2012) to simulate data with different factor counts, then extracting features from this data to train a random forest model. Once the model is trained, it can be used to predict the number of factors in empirical data. The algorithm consists of the following steps:

1. **\*\*Simulation Data:\*\***

- (1) For each value of  $n_{fact}$  in the range from 1 to  $n_{fact_{max}}$ , generate a population data using the [GenData](#) function.
- (2) Each population is based on  $n_{fact}$  factors and consists of  $N_{pop}$  observations.
- (3) For each generated population, repeat the following for  $N_{rep}$  times, For the  $j$ -th in  $N_{rep}$ : a. Draw a sample  $N_{sam}$  from the population that matches the size of the empirical data; b. Compute a feature set  $\mathbf{fea}_{n_{fact},j}$  from each  $N_{sam}$ .
- (4) Combine all the generated feature sets  $\mathbf{fea}_{n_{fact},j}$  into a data frame as  $\mathbf{data}_{train,n_{fact}}$ .
- (5) Combine all  $\mathbf{data}_{train,n_{fact}}$  into a final data frame as the training dataset  $\mathbf{data}_{train}$ .

## 2. \*\*Training RF:\*\*

Train a Random Forest model  $RF$  using the combined  $\mathbf{data}_{train}$ .

## 3. \*\*Prediction the Empirical Data:\*\*

- (1) Calculate the feature set  $\mathbf{fea}_{emp}$  for the empirical data.
- (2) Use the trained Random Forest model  $RF$  to predict the number of factors  $n_{fact_{emp}}$  for the empirical data:

$$n_{fact_{emp}} = RF(\mathbf{fea}_{emp})$$

According to Goretzko & Ruscio (2024) and Breiman (2001), the number of trees in the Random Forest `num. trees` is recommended to be 500. The Random Forest in CDF performs a classification task, so the recommended maximum depth for each tree `mtry` is  $\sqrt{q}$  (where  $q$  is the number of features), which results in  $m_{try} = \sqrt{181} = 13$ .

Since the CDF approach requires extensive data simulation and computation, which is much more time consuming than the [CD](#) Approach, C++ code is used to speed up the process.

## Value

An object of class CDF is a list containing the following components:

<code>nfact</code>	The number of factors to be retained.
<code>RF</code>	the trained Random Forest model
<code>probability</code>	A matrix containing the probabilities for factor numbers ranging from 1 to <code>nfact.max</code> ( <code>1xnfact.max</code> ), where the number in the $f$ -th column represents the probability that the number of factors for the response is $f$ .
<code>features</code>	A matrix ( <code>1x181</code> ) containing all the features for determining the number of factors. @seealso <a href="#">extractor.feature.FF</a>

## Author(s)

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## References

- Breiman, L. (2001). Random Forests. *Machine Learning*, 45(1), 5-32. <https://doi.org/10.1023/A:1010933404324>
- Goretzko, D., & Ruscio, J. (2024). The comparison data forest: A new comparison data approach to determine the number of factors in exploratory factor analysis. *Behavior Research Methods*, 56(3), 1838-1851. <https://doi.org/10.3758/s13428-023-02122-4>
- Ruscio, J., & Roche, B. (2012). Determining the number of factors to retain in an exploratory factor analysis using comparison data of known factorial structure. *Psychological Assessment*, 24, 282-292. <http://dx.doi.org/10.1037/a0025697>.

## See Also

[GenData](#)

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check\_python\_libraries

*Check and Install Python Libraries (numpy and onnxruntime)*

---

## Description

This function checks whether the Python libraries ‘numpy’ and ‘onnxruntime’ are installed. If not, it will prompt the user to decide whether to install them. If the user chooses ‘y’, the required library will be installed using the ‘reticulate’ package. If the user chooses ‘n’, the installation will be skipped. @seealso [DNN\\_predictor](#)

## Usage

```
check_python_libraries()
```

## Value

A list indicating whether ‘numpy’ and ‘onnxruntime’ are installed. The list contains the following logical elements:

numpy\_installed

TRUE if ‘numpy’ is installed, FALSE otherwise.

onnxruntime\_installed

TRUE if ‘onnxruntime’ is installed, FALSE otherwise.

data.bfi

*25 Personality Items Representing 5 Factors***Description**

This dataset includes 25 self-report personality items sourced from the International Personality Item Pool (ipip.ori.org) as part of the Synthetic Aperture Personality Assessment (SAPA) web-based personality assessment project. The dataset contains responses from 2,800 examinees. Additionally, three demographic variables (sex, education, and age) are included.

**Format**

A data frame with 2,800 observations on 28 variables. The variables include:

- A1 - Am indifferent to the feelings of others. (q\_146)
- A2 - Inquire about others' well-being. (q\_1162)
- A3 - Know how to comfort others. (g\_1206)
- A4 - Love children. (g\_1364)
- A5 - Make people feel at ease. (q\_1419)
- C1 - Am exacting in my work. (q\_124)
- C2 - Continue until everything is perfect. (q\_530)
- C3 - Do things according to a plan. (q\_619)
- C4 - Do things in a half-way manner. (g\_626)
- C5 - Waste my time. (g\_1949)
- E1 - Don't talk a lot. (q\_712)
- E2 - Find it difficult to approach others. (q\_901)
- E3 - Know how to captivate people. (q\_1205)
- E4 - Make friends easily. (q\_1410)
- E5 - Take charge. (g\_1768)
- N1 - Get angry easily. (q\_952)
- N2 - Get irritated easily. (q\_974)
- N3 - Have frequent mood swings. (q\_1099)
- N4 - Often feel blue. (g\_1479)
- N5 - Panic easily. (q\_1505)
- O1 - Am full of ideas. (q\_128)
- O2 - Avoid difficult reading material. (g\_316)
- O3 - Carry the conversation to a higher level. (q\_492)
- O4 - Spend time reflecting on things. (g\_1738)
- O5 - Will not probe deeply into a subject. (q\_1964)
- gender - Gender: Males = 1, Females = 2
- education - Education level: 1 = High School, 2 = Finished High School, 3 = Some College, 4 = College Graduate, 5 = Graduate Degree
- age - Age in years

## Details

The 25 items are organized by five factors: Agreeableness, Conscientiousness, Extraversion, Neuroticism, and Openness. The scoring key is created using `make.keys`, and scores are calculated using `score.items`. These factors are useful for IRT-based latent factor analysis of the polychoric correlation matrix. Endorsement plots and item information functions reveal variations in item quality. Responses were collected on a 6-point scale: 1 = Very Inaccurate, 2 = Moderately Inaccurate, 3 = Slightly Inaccurate, 4 = Slightly Accurate, 5 = Moderately Accurate, 6 = Very Accurate, as part of the Synthetic Aperture Personality Assessment (SAPA) project (<https://www.sapa-project.org/>). For examples of data collection techniques, visit <https://www.sapa-project.org/> or the International Cognitive Ability Resource at <https://icar-project.org>. The items were sampled from the International Personality Item Pool of Lewis Goldberg using SAPA sampling techniques. This dataset is a sample from the larger SAPA data bank.

## Note

The data.bfi data set and items should not be confused with the BFI (Big Five Inventory) of Oliver John and colleagues (John, O. P, Donahue, E. M., & Kentle, R.L. (1991). The Big Five Inventory Versions 4a and 54. Berkeley, CA: University of California, Berkeley, Institute of Personality and Social Research.)

## Source

The items are from the ipip (Goldberg, 1999). The data are from the SAPA project (Revelle, Wilt and Rosenthal, 2010), collected Spring, 2010(<https://www.sapa-project.org/>).

## References

- Goldberg, L.R. (1999). A broad-bandwidth, public domain, personality inventory measuring the lower-level facets of several five-factor models. In Mervielde, I., Deary, I., De Fruyt, F., & Ostendorf, F. (Eds.), *Personality psychology in Europe* (Vol. 7, pp. 7-28). Tilburg University Press.
- Revelle, W., Wilt, J., & Rosenthal, A. (2010). Individual Differences in Cognition: New Methods for Examining the Personality-Cognition Link. In Gruszka, A., Matthews, G., & Szymura, B. (Eds.), *Handbook of Individual Differences in Cognition: Attention, Memory and Executive Control* (pp. 117-144). Springer.
- Revelle, W., Condon, D., Wilt, J., French, J.A., Brown, A., & Elleman, L.G. (2016). Web and phone-based data collection using planned missing designs. In Fielding, N.G., Lee, R.M., & Blank, G. (Eds.), *SAGE Handbook of Online Research Methods* (2nd ed., pp. 100-116). Sage Publications.

## Examples

```
data(data.bfi)
head(data.bfi)
```

---

data.datasets	<i>Subset Dataset for Training the Pre-Trained Deep Neural Network (DNN)</i>
---------------	--

---

### Description

This dataset is a subset of the full datasets, consisting of 1,000 samples from the original 10,000,000-sample datasets.

### Format

A 1,000×55 matrix, where the first 54 columns represent feature values and the last column represents the labels, which correspond to the number of factors associated with the features.

### Note

Methods for generating and extracting features from the dataset can be found in [DNN\\_predictor](#).

### See Also

[DNN\\_predictor](#), [load\\_scaler](#), [data.scaler](#), [normalizer](#)

### Examples

```
data(data.datasets)
head(data.datasets)
```

---

data.scaler	<i>the Scaler for the Pre-Trained Deep Neural Network (DNN)</i>
-------------	---

---

### Description

This dataset contains the means and standard deviations of the 10,000,000 datasets for training Pre-Trained Deep Neural Network (DNN), which can be used to determine the number of factors.

### Format

A list containing two vectors, each of length 54:

**means** A numeric vector representing the means of the 54 features extracted from the 10,000,000 datasets.

**sds** A numeric vector representing the standard deviations of the 54 features extracted from the 10,000,000 datasets.

**See Also**

[DNN\\_predictor](#), [load\\_scaler](#), [data.datasets](#), [normalizer](#)

**Examples**

```
data(data.scaler)
print(data.scaler)

data.scaler <- load_scaler()
print(data.scaler)
```

---

DNN_predictor	<i>A Pre-Trained Deep Neural Network (DNN) for Determining the Number of Factors</i>
---------------	--

---

**Description**

This function will invoke a pre-trained deep neural network that can reliably perform the task of determining the number of factors. The maximum number of factors that the network can discuss is 10. The DNN model is implemented in Python and trained on PyTorch (<https://pytorch.org/>) with CUDA 11.8 for acceleration. After training, the DNN was saved as a DNN.onnx file. The *DNN\_predictor* function performs inference by loading the DNN.onnx file in both Python and R environments. Therefore, please note that Python (suggested  $\geq 3.10$ ) and the libraries numpy and onnxruntime are required. @seealso [check\\_python\\_libraries](#)

To run this function, Python is required, along with the installation of numpy and onnxruntime. See more in Details and Note.

**Usage**

```
DNN_predictor(
  response,
  cor.type = "pearson",
  use = "pairwise.complete.obs",
  vis = TRUE,
  plot = TRUE
)
```

**Arguments**

response	A required $N \times I$ matrix or data.frame consisting of the responses of $N$ individuals to $I$ items.
cor.type	A character string indicating which correlation coefficient (or covariance) is to be computed. One of "pearson" (default), "kendall", or "spearman". @seealso <a href="#">cor</a> .

use	An optional character string giving a method for computing covariances in the presence of missing values. This must be one of the strings "everything", "all.obs", "complete.obs", "na.or.complete", or "pairwise.complete.obs" (default). @seealso <a href="#">cor</a> .
vis	A Boolean variable that will print the factor retention results when set to TRUE, and will not print when set to FALSE. (default = TRUE)
plot	A Boolean variable that will print the DNN_predictor plot when set to TRUE, and will not print it when set to FALSE. @seealso <a href="#">plot.DNN_predictor</a> . (Default = TRUE)

## Details

Due to the improved performance of deep learning models with larger datasets (Chen et al., 2017), a total of 10,000,000 datasets ([data.datasets](#)) were simulated to extract features for training deep learning neural networks. Each dataset was generated following the methods described by Auerwald & Moshagen (2019) and Goretzko & Buhner (2020), with the following specifications:

- Factor number:  $F \sim U[1,10]$
- Sample size:  $N \sim U[100,1000]$
- Number of variables per factor:  $vpf \sim [3,20]$
- Factor correlation:  $fc \sim U[0.0,0.4]$
- Primary loadings:  $pl \sim U[0.35,0.80]$
- Cross-loadings:  $cl \sim U[-0.2,0.2]$

A population correlation matrix was created for each data set based on the following decomposition:

$$\Sigma = \Lambda \Phi \Lambda^T + \Delta$$

where  $\Lambda$  is the loading matrix,  $\Phi$  is the factor correlation matrix, and  $\Delta$  is a diagonal matrix, with  $\Delta = 1 - \text{diag}(\Lambda \Phi \Lambda^T)$ . The purpose of  $\Delta$  is to ensure that the diagonal elements of  $\Sigma$  are 1.

The response data for each subject was simulated using the following formula:

$$X_i = L_i + \epsilon_i, \quad 1 \leq i \leq I$$

where  $L_i$  follows a normal distribution  $N(0, \sigma)$ , representing the contribution of latent factors, and  $\epsilon_i$  is the residual term following a standard normal distribution.  $L_i$  and  $\epsilon_i$  are uncorrelated, and  $\epsilon_i$  and  $\epsilon_j$  are also uncorrelated.

For each simulated dataset, a total of 6 types of features (which can be classified into 2 types; @seealso [extractor.feature.DNN](#)) are extracted and compiled into a feature vector, consisting of 54 features:  $8 + 8 + 8 + 10 + 10 + 10$ . These features are as follows:

### 1. Clustering-Based Features

- (1) Hierarchical clustering is performed with correlation coefficients as dissimilarity. The top 9 tree node heights are calculated, and all heights are divided by the maximum height. The heights from the 2nd to 9th nodes are used as features. @seealso [EFAhclust](#)
- (2) Hierarchical clustering with Euclidean distance as dissimilarity is performed. The top 9 tree node heights are calculated, and all heights are divided by the maximum height. The heights from the 2nd to 9th nodes are used as features. @seealso [EFAhclust](#)

- (3) K-means clustering is applied with the number of clusters ranging from 1 to 9. The within-cluster sum of squares (WSS) for clusters 2 to 9 are divided by the WSS for a single cluster. @seealso [EFAkmeans](#)

These three features are based on clustering algorithms. The purpose of division is to normalize the data. These clustering metrics often contain information unrelated to the number of factors, such as the number of items and the number of respondents, which can be avoided by normalization. The reason for using the 2nd to 9th data is that only the top F-1 data are needed to determine the number of factors F. The first data point is fixed at 1 after the division operations, so it is excluded. This approach helps in model simplification.

## 2. Traditional Exploratory Factor Analysis Features (Eigenvalues)

- (4) The top 10 largest eigenvalues.
- (5) The ratio of the top 10 largest eigenvalues to the corresponding reference eigenvalues from Empirical Kaiser Criterion (EKC; Braeken & van Assen, 2017). @seealso [EKC](#)
- (6) The cumulative variance proportion of the top 10 largest eigenvalues.

Only the top 10 elements are used to simplify the model.

The DNN model is implemented in Python and trained on PyTorch (<https://download.pytorch.org/whl/cu118>) with CUDA 11.8 for acceleration. After training, the DNN was saved as a DNN.onnx file. The *DNN\_predictor* function performs inference by loading the DNN.onnx file in both Python and R environments.

## Value

An object of class *DNN\_predictor* is a list containing the following components:

<code>nfact</code>	The number of factors to be retained.
<code>features</code>	A matrix (1×54) containing all the features for determining the number of factors by the DNN.
<code>probability</code>	A matrix containing the probabilities for factor numbers ranging from 1 to 10 (1×10), where the number in the f-th column represents the probability that the number of factors for the response is f.

## Note

Note that Python and the libraries `numpy` and `onnxruntime` are required.

First, please ensure that Python is installed on your computer and that Python is included in the system's PATH environment variable. If not, please download and install it from the official website (<https://www.python.org/>).

If you encounter an error when running this function stating that the `numpy` and `onnxruntime` modules are missing:

```
Error in py_module_import(module, convert = convert) :
```

```
ModuleNotFoundError: No module named 'numpy'
```

or

```
Error in py_module_import(module, convert = convert) :
```

ModuleNotFoundError: No module named 'onnxruntime'

this means that the numpy or onnxruntime library is missing from your Python environment. The [check\\_python\\_libraries](#) function can help you install these two dependency libraries. @seealso [check\\_python\\_libraries](#)

Of course, you can also choose not to use the [check\\_python\\_libraries](#) function. You can directly install the numpy or onnxruntime library using the appropriate commands. If you are using Windows or macOS, please run the command `pip install numpy` or `pip install onnxruntime` in Command Prompt or Windows PowerShell (Windows), or Terminal (macOS). If you are using Linux, please ensure that pip is installed and use the command `pip install numpy` or `pip install onnxruntime` to install the missing libraries.

### Author(s)

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### References

- Auerswald, M., & Moshagen, M. (2019). How to determine the number of factors to retain in exploratory factor analysis: A comparison of extraction methods under realistic conditions. *Psychological methods*, 24(4), 468-491. <https://doi.org/10.1037/met0000200>.
- Braeken, J., & van Assen, M. A. L. M. (2017). An empirical Kaiser criterion. *Psychological methods*, 22(3), 450-466. <https://doi.org/10.1037/met0000074>.
- Goretzko, D., & Buhner, M. (2020). One model to rule them all? Using machine learning algorithms to determine the number of factors in exploratory factor analysis. *Psychol Methods*, 25(6), 776-786. <https://doi.org/10.1037/met0000262>.

---

EFAhclust

*Hierarchical Clustering for EFA*

---

### Description

A function performs clustering on items by calling [hclust](#). Hierarchical cluster analysis on a set of dissimilarities and methods for analyzing it. The items will be continuously clustered in pairs until all items are grouped into a single cluster, at which point the process will stop.

### Usage

```
EFAhclust(
  response,
  dissimilarity.type = "R",
  method = "ward.D",
  cor.type = "pearson",
  use = "pairwise.complete.obs",
  nfact.max = 10,
  plot = TRUE
)
```

## Arguments

<code>response</code>	A required $N \times I$ matrix or <code>data.frame</code> consisting of the responses of $N$ individuals to $I$ items.
<code>dissimilarity.type</code>	A character indicating which kind of dissimilarity is to be computed. One of "R" or "E" (default) for the correlation coefficient or Euclidean distance.
<code>method</code>	the agglomeration method to be used. This should be (an unambiguous abbreviation of) one of "ward.D", "ward.D2", "single", "complete", "average" (= UPGMA), "mcquitty" (= WPGMA), "median" (= WPGMC) or "centroid" (= UPGMC). (default = "ward.D") @seealso <a href="#">hclust</a>
<code>cor.type</code>	A character string indicating which correlation coefficient (or covariance) is to be computed. One of "pearson" (default), "kendall", or "spearman". @seealso <a href="#">cor</a> .
<code>use</code>	An optional character string giving a method for computing covariances in the presence of missing values. This must be one of the strings "everything", "all.obs", "complete.obs", "na.or.complete", or "pairwise.complete.obs" (default). @seealso <a href="#">cor</a> .
<code>nfact.max</code>	The maximum number of factors discussed by the Second-Order Difference (SOD) approach. (default = 10)
<code>plot</code>	A Boolean variable that will print the EFAhclust plot when set to TRUE, and will not print it when set to FALSE. @seealso <a href="#">plot.EFAhclust</a> . (Default = TRUE)

## Details

Hierarchical cluster analysis always merges the two nodes with the smallest dissimilarity, forming a new node in the process. This continues until all nodes are merged into one large node, at which point the algorithm terminates. This method undoubtedly creates a hierarchical structure by the end of the process, which encompasses the relationships between all items: items with high correlation have short connecting lines between them, while items with low correlation have longer lines. This hierarchical structure is well-suited to be represented as a binary tree. In this representation, the dissimilarity between two nodes can be indicated by the height of the tree nodes; the greater the difference between nodes, the higher the height of the tree nodes connecting them (the longer the line). Researchers can decide whether two nodes belong to the same cluster based on the height differences between nodes, which, in exploratory factor analysis, represents whether these two nodes belong to the same latent factor.

The Second-Order Difference (SOD) approach is a commonly used method for finding the "elbow" (the point of greatest slope change). According to the principles of exploratory factor analysis, items belonging to different latent factors have lower correlations, while items under the same factor are more highly correlated. In hierarchical clustering, this is reflected in the height of the nodes in the dendrogram, with differences in node heights representing the relationships between items. By sorting all node heights in descending order and applying the SOD method to locate the elbow, the number of factors can be determined. @seealso [EFAkmeans](#)

## Value

An object of class `EFAhclust` is a list containing the following components:

<code>hc</code>	An object of class <code>hclust</code> that describes the tree produced by the clustering process. @seealso <a href="#">hclust</a>
<code>cor.response</code>	A matrix of dimension $I \times I$ containing all the correlation coefficients of items.
<code>clusters</code>	A list containing all the clusters.
<code>heights</code>	A vector containing all the heights of the cluster tree. The heights are arranged in descending order.
<code>nfact.SOD</code>	The number of factors to be retained by the Second-Order Difference (SOD) approach.

## References

Batagelj, V. (1988). Generalized Ward and Related Clustering Problems. In H. H. Bock, Classification and Related Methods of Data Analysis the First Conference of the International Federation of Classification Societies (IFCS), Amsterdam.

Murtagh, F., & Legendre, P. (2014). Ward's Hierarchical Agglomerative Clustering Method: Which Algorithms Implement Ward's Criterion? *Journal of Classification*, 31(3), 274-295. <https://doi.org/10.1007/s00357-014-9161-z>.

## Examples

```
library(EFAfactors)
set.seed(123)

##Take the data.bfi dataset as an example.
data(data.bfi)

response <- as.matrix(data.bfi[, 1:25]) ## loading data
response <- na.omit(response) ## Remove samples with NA/missing values

## Transform the scores of reverse-scored items to normal scoring
response[, c(1, 9, 10, 11, 12, 22, 25)] <- 6 - response[, c(1, 9, 10, 11, 12, 22, 25)] + 1

## Run EFAhclust function with default parameters.

EFAhclust.obj <- EFAhclust(response)

plot(EFAhclust.obj)

## Get the heights.
heights <- EFAhclust.obj$heights
print(heights)

## Get the nfact retained by SOD
nfact.SOD <- EFAhclust.obj$nfact.SOD
print(nfact.SOD)
```

EFAindex

*Various Indices in EFA***Description**

A function performs clustering on items by calling [VSS](#) and [fa](#). Apply the Very Simple Structure (VSS), Comparative Fit Index (CFI), MAP, and other criteria to determine the appropriate number of factors.

**Usage**

```
EFAindex(
  response,
  nfact.max = 10,
  cor.type = "cor",
  use = "pairwise.complete.obs"
)
```

**Arguments**

<code>response</code>	A required $N \times I$ matrix or data.frame consisting of the responses of $N$ individuals to $I$ items.
<code>nfact.max</code>	The maximum number of factors discussed by CD approach. (default = 10)
<code>cor.type</code>	How to find the correlations: "cor" is Pearson, "cov" is covariance, "tet" is tetrachoric, "poly" is polychoric, "mixed" uses mixed cor for a mixture of tetrachorics, polychorics, Pearsons, biserials, and polyserials, Yuleb is Yulebonett, Yuleq and YuleY are the obvious Yule coefficients as appropriate.
<code>use</code>	an optional character string giving a method for computing covariances in the presence of missing values. This must be one of the strings "everything", "all.obs", "complete.obs", "na.or.complete", or "pairwise.complete.obs" (default). @seealso <a href="#">cor</a> .

**Value**

A matrix with the following components:

**CFI** the Comparative Fit Index

**RMSEA** Root Mean Square Error of Approximation (RMSEA) for each number of factors.

**SRMR** Standardized Root Mean Square Residual.

**MAP** Velicer's MAP values (lower values are better).

**BIC** Bayesian Information Criterion (BIC) for each number of factors.

**SABIC** Sample-size Adjusted Bayesian Information Criterion (SABIC) for each number of factors.

**chisq** Chi-square statistic from the factor analysis output.

**df** Degrees of freedom.

**prob** Probability that the residual matrix is greater than 0.  
**eChiSq** Empirically found chi-square statistic.  
**eCRMS** Empirically found mean residual corrected for degrees of freedom.  
**eBIC** Empirically found BIC based on the empirically found chi-square statistic.  
**vss** VSS fit with complexity 1.  
**sqresid** Squared residual correlations.  
**fit** Factor fit of the complete model.

### Examples

```
library(EFAfactors)
set.seed(123)

##Take the data.bfi dataset as an example.
data(data.bfi)

response <- as.matrix(data.bfi[, 1:25]) ## loading data
response <- na.omit(response) ## Remove samples with NA/missing values

## Transform the scores of reverse-scored items to normal scoring
response[, c(1, 9, 10, 11, 12, 22, 25)] <- 6 - response[, c(1, 9, 10, 11, 12, 22, 25)] + 1

## Run EFAindex function with default parameters.

EFAindex.matrix <- EFAindex(response)

print(EFAindex.matrix)
```

---

EFAkmeans

*K-means for EFA*


---

### Description

A function performs K-means algorithm on items by calling [kmeans](#).

### Usage

```
EFAkmeans(response, nfact.max = 10, plot = TRUE)
```

**Arguments**

response	A required $N \times I$ matrix or data.frame consisting of the responses of $N$ individuals to $I$ items.
nfact.max	The maximum number of factors discussed by EFAkmeans approach. (default = 10)
plot	A Boolean variable that will print the EFAkmeans plot when set to TRUE, and will not print it when set to FALSE. @seealso <a href="#">plot.EFAkmeans</a> . (Default = TRUE)

**Details**

K-means is a well-established and widely used classical clustering algorithm. It is an unsupervised machine learning algorithm that requires the number of clusters  $K$  to be specified in advance. After K-means terminates, the total within-cluster sum of squares (WSS) can be calculated to represent the goodness of fit of the clustering:

$$WSS = \sum_{C_k \in C} \sum_{i \in C_k} \|i - \mu_k\|^2$$

where  $C$  is the set of all clusters.  $C_k$  is the  $k$ -th cluster.  $i$  represents each item in the cluster  $C_k$ .  $\mu_k$  is the centroid of cluster  $C_k$ .

Similar to the scree plot where eigenvalues decrease as the number of factors increases, WSS also decreases as  $K$  increases. A "significant reduction" in WSS at a particular  $K$  may suggest that  $K$  is the most appropriate number of clusters, which in exploratory factor analysis implies that the number of factors is  $K$ . The "significant reduction" can be identified using the Second-Order Difference (SOD) approach. @seealso [EFAkmeans](#)

**Value**

An object of class EFAkmeans is a list containing the following components:

wss	A vector containing all within-cluster sum of squares (WSS).
nfact.SOD	The number of factors to be retained by the Second-Order Difference (SOD) approach.

**Examples**

```
library(EFAfactors)
set.seed(123)

##Take the data.bfi dataset as an example.
data(data.bfi)

response <- as.matrix(data.bfi[, 1:25]) ## loading data
response <- na.omit(response) ## Remove samples with NA/missing values

## Transform the scores of reverse-scored items to normal scoring
response[, c(1, 9, 10, 11, 12, 22, 25)] <- 6 - response[, c(1, 9, 10, 11, 12, 22, 25)] + 1
```

```
## Run EFAkmeans function with default parameters.

EFAkmeans.obj <- EFAkmeans(response)

plot(EFAkmeans.obj)

## Get the heights.
wss <- EFAkmeans.obj$wss
print(wss)

## Get the nfact retained by SOD
nfact.SOD <- EFAkmeans.obj$nfact.SOD
print(nfact.SOD)
```

---

EFAscreet

*Scree Plot*


---

## Description

This function generates a scree plot to display the eigenvalues of the correlation matrix computed from the given response data. The scree plot helps in determining the number of factors to retain in exploratory factor analysis by examining the point at which the eigenvalues start to level off, indicating less variance explained by additional factors.

## Usage

```
EFAscreet(
  response,
  fa = "pc",
  nfact = 1,
  cor.type = "pearson",
  use = "pairwise.complete.obs"
)
```

## Arguments

response	A required $N \times I$ matrix or data.frame consisting of the responses of $N$ individuals to $I$ items.
fa	A string that determines the method used to obtain eigenvalues. If 'pc', it represents Principal Component Analysis (PCA); if 'fa', it represents Principal Axis Factoring (a widely used Factor Analysis method; @seealso <a href="#">factor.analysis</a> ; Auerswald & Moshagen, 2019). (Default = 'pc')

<code>nfact</code>	A numeric value that specifies the number of factors to extract, only effective when <code>fa = 'fa'</code> . (Default = 1)
<code>cor.type</code>	A character string indicating which correlation coefficient (or covariance) is to be computed. One of "pearson" (default), "kendall", or "spearman". @seealso <a href="#">cor</a> .
<code>use</code>	An optional character string giving a method for computing covariances in the presence of missing values. This must be one of the strings "everything", "all.obs", "complete.obs", "na.or.complete", or "pairwise.complete.obs" (default). @seealso <a href="#">cor</a> .

### Value

An object of class `EFAscreet` is a list containing the following components:

`eigen.value`      A vector containing the empirical eigenvalues

### See Also

[plot.EFAscreet](#)

### Examples

```
library(EFAfactors)
set.seed(123)

##Take the data.bfi dataset as an example.
data(data.bfi)

response <- as.matrix(data.bfi[, 1:25]) ## loading data
response <- na.omit(response) ## Remove samples with NA/missing values

## Transform the scores of reverse-scored items to normal scoring
response[, c(1, 9, 10, 11, 12, 22, 25)] <- 6 - response[, c(1, 9, 10, 11, 12, 22, 25)] + 1

## Run EFAscreet function with default parameters.

EFAscreet.obj <- EFAscreet(response)

plot(EFAscreet.obj)
```

---

EFAsim.data

---

*Simulate Data that Conforms to the theory of Exploratory Factor Analysis.*


---

## Description

This function is used to simulate data that conforms to the theory of exploratory factor analysis, with a high degree of customization for the variables involved.

## Usage

```
EFAsim.data(
  nfact,
  vpf,
  N = 500,
  distri = "normal",
  fc = "R",
  pl = "R",
  cl = "R",
  low.vpf = 5,
  up.vpf = 15,
  a = NULL,
  b = NULL,
  vis = TRUE
)
```

## Arguments

nfact	A numeric value specifying the number of factors to simulate.
vpf	A numeric or character value specifying the number of items under each factor. If a numeric value is provided, the numeric must be larger than 2, and the number of items under each factor will be fixed to this value. If a character value is provided, it must be one of 'S', 'M', 'L', or 'R'. These represent random selection of items under each factor from $U(5, 10)$ , $U(5, 15)$ , $U(5, 20)$ , or $U(low.vpf, up.vpf)$ , respectively.
N	A numeric value specifying the number of examinees to simulate.
distri	A character, either 'normal' or 'beta', indicating whether the simulated data will follow a standard multivariate normal distribution or a multivariate beta distribution.
fc	A numeric or character value specifying the degree of correlation between factors. If a numeric value is provided, it must be within the range of 0 to 0.75, and the correlation between all factors will be fixed at this value. If a character value is provided, it must be 'R', and the correlations between factors will be randomly selected from $U(0.0, 0.5)$ .

pl	A numeric or character value specifying the size of the primary factor loadings. If a numeric value is provided, it must be within the range of 0 to 1, and all primary factor loadings in the loading matrix will be fixed at this value. If a character value is provided, it must be one of 'L', 'M', 'H', or 'R', representing $pl \sim U(0.35, 0.50)$ , $pl \sim U(0.50, 0.65)$ , $pl \sim U(0.65, 0.80)$ , or $pl \sim U(0.35, 0.80)$ , respectively, consistent with the settings in Goretzko & Buhner (2020).
cl	A numeric or character value specifying the size of cross-loadings. If a numeric value is provided, it must be within the range of 0 to 0.5, and all cross-loadings in the loading matrix will be fixed at this value. If a character value is provided, it must be one of 'L', 'H', 'None', or 'R', representing $cl \sim U(-0.1, 0.1)$ , $cl \sim U(-0.2, -0.1) \cup U(0.1, 0.2)$ , $cl = 0$ , or $cl \sim U(-0.2, 0.2)$ , respectively, consistent with the settings in Auerswald & Moshagen (2019).
low.vpf	A numeric value specifying the minimum number of items per factor, must be larger than 2, effective only when vpf is 'R'. (default = 5)
up.vpf	A numeric value specifying the maximum number of items per factor, effective only when vpf is 'R'. (default = 15)
a	A numeric or NULL specifying the 'a' parameter of the beta distribution, effective only when <code>distri = 'beta'</code> . If a numeric value is provided, it will be used as the 'a' parameter of the beta distribution. If NULL, a random integer between 1 and 10 will be used. (default = NULL)
b	A numeric or NULL specifying the 'b' parameter of the beta distribution, effective only when <code>distri = 'beta'</code> . If a numeric value is provided, it will be used as the 'b' parameter of the beta distribution. If NULL, a random integer between 1 and 10 will be used. (default = NULL)
vis	A logical value indicating whether to print process information. (default = TRUE)

## Details

A population correlation matrix was created for each data set based on the following decomposition:

$$\Sigma = \Lambda \Phi \Lambda^T + \Delta$$

where  $\Lambda$  is the loading matrix,  $\Phi$  is the factor correlation matrix, and  $\Delta$  is a diagonal matrix, with  $\Delta = 1 - \text{diag}(\Lambda \Phi \Lambda^T)$ . The purpose of  $\Delta$  is to ensure that the diagonal elements of  $\Sigma$  are 1.

The response data for each subject was simulated using the following formula:

$$X_i = L_i + \epsilon_i, \quad 1 \leq i \leq I$$

where  $L_i$  follows a standard normal distribution (`distri = 'normal'`) or a beta distribution (`distri = 'beta'`), representing the contribution of latent factors. And  $\epsilon_i$  is the residual term following a standard normal distribution (`distri = 'normal'`) or a beta distribution (`distri = 'beta'`).  $L_i$  and  $\epsilon_i$  are uncorrelated, and  $\epsilon_i$  and  $\epsilon_j$  are also uncorrelated.

## Value

An object of class `EFadata` is a list containing the following components:

loadings	A simulated loading matrix.
items	A list containing all factors and the item indices under each factor.
cor.factors	A simulated factor correlation matrix.
cor.items	A simulated item correlation matrix.
response	A simulated response data matrix.

## References

Goretzko, D., & Buhner, M. (2020). One model to rule them all? Using machine learning algorithms to determine the number of factors in exploratory factor analysis. *Psychological Methods*, 25(6), 776-786. <https://doi.org/10.1037/met0000262>.

Auerswald, M., & Moshagen, M. (2019). How to determine the number of factors to retain in exploratory factor analysis: A comparison of extraction methods under realistic conditions. *Psychological methods*, 24(4), 468-491. <https://doi.org/https://doi.org/10.1037/met0000200>

## Examples

```
library(EFAfactors)

## Run EFAsim.data function with default parameters.
data.obj <- EFAsim.data(nfact = 3, vpf = 5, N=500, distri="normal", fc="R", pl="R", cl="R",
                      low.vpf = 5, up.vpf = 15, a = NULL, b = NULL, vis = TRUE)

head(data.obj$loadings)
```

---

EFAvote

---

*Voting Method for Number of Factors in EFA*


---

## Description

This function implements a voting method to determine the most appropriate number of factors in exploratory factor analysis (EFA). The function accepts a vector of votes, where each value represents the number of factors suggested by different EFA approaches. If there is a clear winner (a single number of factors with the most votes), that number is returned. In case of a tie, the function returns the first value among the tied results and outputs a message. The result is returned as an object of class `vote`, which can be printed and plotted.

## Usage

```
EFAvote(votes, vis = TRUE, plot = TRUE)
```

**Arguments**

<code>votes</code>	A vector of integers, where each element corresponds to the number of factors suggested by an EFA method.
<code>vis</code>	Logical, whether to print the results of the voting. Defaults to TRUE.
<code>plot</code>	Logical, whether to display a pie chart of the voting results. Defaults to TRUE.

**Value**

An object of class `EFAvote`, which is a list containing:

<code>nfact</code>	The number of factors with the most votes. If there is a tie, the first one in the order is returned.
<code>votes</code>	The original vector of votes.

**See Also**

[plot.EFAvote](#)

**Examples**

```
library(EFAfactors)

nfacts <- c(5, 5, 5, 6, 6, 4)
names(nfacts) <- c("Hull", "CD", "PA", "EKC", "FF", "DNN")

EFAvote.obj <- EFAvote(votes = nfacts)

# Visualize the voting results
plot(EFAvote.obj)
```

---

EKC

---

*Empirical Kaiser Criterion*


---

**Description**

This function will apply the Empirical Kaiser Criterion (Braeken & van Assen, 2017) method to determine the number of factors. The method assumes that the distribution of eigenvalues asymptotically follows a Marcenko-Pastur distribution (Marcenko & Pastur, 1967). It calculates the reference eigenvalues based on this distribution and determines whether to retain a factor by comparing the size of the empirical eigenvalues to the reference eigenvalues.

**Usage**

```
EKC(
  response,
  cor.type = "pearson",
  use = "pairwise.complete.obs",
  vis = TRUE,
  plot = TRUE
)
```

**Arguments**

<code>response</code>	A required $N \times I$ matrix or <code>data.frame</code> consisting of the responses of $N$ individuals to $I$ items.
<code>cor.type</code>	A character string indicating which correlation coefficient (or covariance) is to be computed. One of "pearson" (default), "kendall", or "spearman". @seealso <a href="#">cor</a> .
<code>use</code>	An optional character string giving a method for computing covariances in the presence of missing values. This must be one of the strings "everything", "all.obs", "complete.obs", "na.or.complete", or "pairwise.complete.obs" (default). @seealso <a href="#">cor</a> .
<code>vis</code>	A Boolean variable that will print the factor retention results when set to TRUE, and will not print when set to FALSE. (default = TRUE)
<code>plot</code>	A Boolean variable that will print the EKC plot when set to TRUE, and will not print it when set to FALSE. @seealso <a href="#">plot.EKC</a> . (Default = TRUE)

**Details**

The Empirical Kaiser Criterion (EKC; Auerswald & Moshagen, 2019; Braeken & van Assen, 2017) refines Kaiser-Guttman Criterion by accounting for random sample variations in eigenvalues. At the population level, the EKC is equivalent to the original Kaiser-Guttman Criterion, extracting all factors whose eigenvalues from the correlation matrix are greater than one. However, at the sample level, it adjusts for the distribution of eigenvalues in normally distributed data. Under the null model, the eigenvalue distribution follows the Marčenko-Pastur distribution (Marčenko & Pastur, 1967) asymptotically. The upper bound of this distribution serves as the reference eigenvalue for the first eigenvalue  $\lambda$ , so

$$\lambda_{1,ref} = \left(1 + \sqrt{\frac{I}{N}}\right)^2$$

, which is determined by  $N$  individuals and  $I$  items. For subsequent eigenvalues, adjustments are made based on the variance explained by previous factors. The  $j$ -th reference eigenvalue is:

$$\lambda_{j,ref} = \max \left[ \frac{I - \sum_{i=0}^{j-1} \lambda_i}{I - j + 1} \left(1 + \sqrt{\frac{I}{N}}\right)^2, 1 \right]$$

The  $j$ -th reference eigenvalue is reduced according to the magnitude of earlier eigenvalues since higher previous values mean less unexplained variance remains. As in the original Kaiser-Guttman Criterion, the reference eigenvalue cannot drop below one.

$$F = \sum_{i=1}^I I(\lambda_i > \lambda_{i,ref})$$

Here,  $\lfloor F \rfloor$  represents the number of factors determined by the EKC, and  $I(\cdot)$  is the indicator function, which equals 1 when the condition is true, and 0 otherwise.

### Value

An object of class EKC is a list containing the following components:

nfact	The number of factors to be retained.
eigen.value	A vector containing the empirical eigenvalues
eigen.ref	A vector containing the reference eigenvalues

### Author(s)

Haijiang Qin <Haijiang133@outlook.com>

### References

- Auerswald, M., & Moshagen, M. (2019). How to determine the number of factors to retain in exploratory factor analysis: A comparison of extraction methods under realistic conditions. *Psychological methods*, 24(4), 468-491. <https://doi.org/10.1037/met0000200>.
- Braeken, J., & van Assen, M. A. L. M. (2017). An empirical Kaiser criterion. *Psychological methods*, 22(3), 450-466. <https://doi.org/10.1037/met0000074>.
- Marcenko, V. A., & Pastur, L. A. (1967). Distribution of eigenvalues for some sets of random matrices. *Mathematics of the USSR-Sbornik*, 1, 457-483. <http://dx.doi.org/10.1070/SM1967v001n04ABEH001994>

### Examples

```
library(EFAfactors)
set.seed(123)

##Take the data.bfi dataset as an example.
data(data.bfi)

response <- as.matrix(data.bfi[, 1:25]) ## loading data
response <- na.omit(response) ## Remove samples with NA/missing values

## Transform the scores of reverse-scored items to normal scoring
response[, c(1, 9, 10, 11, 12, 22, 25)] <- 6 - response[, c(1, 9, 10, 11, 12, 22, 25)] + 1

## Run EKC function with default parameters.
```

```
EKC.obj <- EKC(response)

print(EKC.obj)

plot(EKC.obj)

## Get the eigen.value, eigen.ref and nfact results.
eigen.value <- EKC.obj$eigen.value
eigen.ref <- EKC.obj$eigen.ref
nfact <- EKC.obj$nfact

print(eigen.value)
print(eigen.ref)
print(nfact)
```

---

extractor.feature.DNN *Extracting features for the Pre-Trained Deep Neural Network (DNN)*

---

## Description

This function is used to extract the features required by the Pre-Trained Deep Neural Network (DNN). @seealso [DNN\\_predictor](#)

## Usage

```
extractor.feature.DNN(
  response,
  cor.type = "pearson",
  use = "pairwise.complete.obs"
)
```

## Arguments

response	A required $N \times I$ matrix or data.frame consisting of the responses of $N$ individuals to $I$ items.
cor.type	A character string indicating which correlation coefficient (or covariance) is to be computed. One of "pearson" (default), "kendall", or "spearman". @seealso <a href="#">cor</a> .
use	An optional character string giving a method for computing covariances in the presence of missing values. This must be one of the strings "everything", "all.obs", "complete.obs", "na.or.complete", or "pairwise.complete.obs" (default). @seealso <a href="#">cor</a> .

## Details

A total of two types of features (6 kinds, making up 54 features in total) will be extracted, and they are as follows: 1. Clustering-Based Features

- (1) Hierarchical clustering is performed with correlation coefficients as dissimilarity. The top 9 tree node heights are calculated, and all heights are divided by the maximum height. The heights from the 2nd to 9th nodes are used as features. @seealso [EFAhclust](#)
- (2) Hierarchical clustering with Euclidean distance as dissimilarity is performed. The top 9 tree node heights are calculated, and all heights are divided by the maximum height. The heights from the 2nd to 9th nodes are used as features. @seealso [EFAhclust](#)
- (3) K-means clustering is applied with the number of clusters ranging from 1 to 9. The within-cluster sum of squares (WSS) for clusters 2 to 9 are divided by the WSS for a single cluster. @seealso [EFAkmeans](#)

These three features are based on clustering algorithms. The purpose of division is to normalize the data. These clustering metrics often contain information unrelated to the number of factors, such as the number of items and the number of respondents, which can be avoided by normalization. The reason for using the 2nd to 9th data is that only the top F-1 data are needed to determine the number of factors F. The first data point is fixed at 1 after the division operations, so it is excluded. This approach helps in model simplification.

## 2. Traditional Exploratory Factor Analysis Features (Eigenvalues)

- (4) The top 10 largest eigenvalues.
- (5) The ratio of the top 10 largest eigenvalues to the corresponding reference eigenvalues from Empirical Kaiser Criterion (EKC; Braeken & van Assen, 2017). @seealso [EKC](#)
- (6) The cumulative variance proportion of the top 10 largest eigenvalues.

Only the top 10 elements are used to simplify the model.

## Value

A matrix (1×54) containing all the features for determining the number of factors by the DNN.

## Author(s)

Haijiang Qin <Haijiang133@outlook.com>

## See Also

[DNN\\_predictor](#)

## Examples

```
library(EFAfactors)
set.seed(123)

##Take the data.bfi dataset as an example.
data(data.bfi)
```

```

response <- as.matrix(data.bfi[, 1:25]) ## loading data
response <- na.omit(response) ## Remove samples with NA/missing values

## Transform the scores of reverse-scored items to normal scoring
response[, c(1, 9, 10, 11, 12, 22, 25)] <- 6 - response[, c(1, 9, 10, 11, 12, 22, 25)] + 1

## Run extractor.feature.DNN function with default parameters.

features <- extractor.feature.DNN(response)

print(features)

```

---

extractor.feature.FF    *Extracting features According to Goretzko & Buhner (2020)*

---

## Description

This function will extract 181 features from the data according to the method by Goretzko & Buhner (2020).

## Usage

```

extractor.feature.FF(
  response,
  cor.type = "pearson",
  use = "pairwise.complete.obs"
)

```

## Arguments

response	A required $N \times I$ matrix or data.frame consisting of the responses of $N$ individuals to $I$ items.
cor.type	A character string indicating which correlation coefficient (or covariance) is to be computed. One of "pearson" (default), "kendall", or "spearman". @seealso <a href="#">cor</a> .
use	An optional character string giving a method for computing covariances in the presence of missing values. This must be one of the strings "everything", "all.obs", "complete.obs", "na.or.complete", or "pairwise.complete.obs" (default). @seealso <a href="#">cor</a> .

## Details

The code for the `extractor.feature.FF` function is implemented based on the publicly available code by Goretzko & Buhner (2020) (<https://osf.io/mvrau/>). The extracted features are completely consistent with the 181 features described in the original text by Goretzko & Buhner (2020). These features include:

- 1. - Number of examinees
- 2. - Number of items
- 3. - Number of eigenvalues greater than 1
- 4. - Proportion of variance explained by the 1st eigenvalue
- 5. - Proportion of variance explained by the 2nd eigenvalue
- 6. - Proportion of variance explained by the 3rd eigenvalue
- 7. - Number of eigenvalues greater than 0.7
- 8. - Standard deviation of the eigenvalues
- 9. - Number of eigenvalues accounting for 50
- 10. - Number of eigenvalues accounting for 75
- 11. - L1-norm of the correlation matrix
- 12. - Frobenius-norm of the correlation matrix
- 13. - Maximum-norm of the correlation matrix
- 14. - Average of the off-diagonal correlations
- 15. - Spectral-norm of the correlation matrix
- 16. - Number of correlations smaller or equal to 0.1
- 17. - Average of the initial communality estimates
- 18. - Determinant of the correlation matrix
- 19. - Measure of sampling adequacy (MSA after Kaiser, 1970)
- 20. - Gini coefficient (Gini, 1921) of the correlation matrix
- 21. - Kolm measure of inequality (Kolm, 1999) of the correlation matrix
- 22-101. - Eigenvalues from Principal Component Analysis (PCA), padded with -1000 if insufficient
- 102-181. - Eigenvalues from Factor Analysis (FA), fixed at 1 factor, padded with -1000 if insufficient

## Value

A matrix (1×181) containing all the 181 features (Goretzko & Buhner, 2020).

## References

Goretzko, D., & Buhner, M. (2020). One model to rule them all? Using machine learning algorithms to determine the number of factors in exploratory factor analysis. *Psychol Methods*, 25(6), 776-786. <https://doi.org/10.1037/met0000262>.

factor.analysis

*Factor Analysis by Principal Axis Factoring***Description**

This function performs factor analysis using the Principal Axis Factoring (PAF) method. The process involves extracting factors from an initial correlation matrix and iteratively refining the factor estimates until convergence is achieved.

**Usage**

```
factor.analysis(
  data,
  nfact = 1,
  iter.max = 1000,
  criterion = 0.001,
  cor.type = "pearson",
  use = "pairwise.complete.obs"
)
```

**Arguments**

data	A data.frame or matrix of response. If the matrix is square, it is assumed to be a correlation matrix. Otherwise, correlations (with pairwise deletion) will be computed.
nfact	The number of factors to extract. (default = 1)
iter.max	The maximum number of iterations for the factor extraction process. Default is 1000.
criterion	The convergence criterion for the iterative process. The extraction process will stop when the change in communalities is less than this value. Default is 0.001
cor.type	A character string indicating which correlation coefficient (or covariance) is to be computed. One of "pearson" (default), "kendall", or "spearman". @seealso <a href="#">cor</a> .
use	An optional character string giving a method for computing covariances in the presence of missing values. This must be one of the strings "everything", "all.obs", "complete.obs", "na.or.complete", or "pairwise.complete.obs" (default). @seealso <a href="#">cor</a> .

**Details**

The Principal Axis Factoring (PAF) method involves the following steps:

Step 1. **\*\*Basic Principle\*\***: The core principle of factor analysis using Principal Axis Factoring (PAF) is expressed as:

$$\mathbf{R} = \mathbf{\Lambda}\mathbf{\Lambda}^T + \mathbf{\Phi}$$

$$R_{ii} = H_i^2 + \Phi_{ii}$$

where  $\Lambda$  is the matrix of factor loadings, and  $\Phi$  is the diagonal matrix of unique variances. Here,  $H_i^2$  represents the portion of the i-th item's variance explained by the factor model.  $\mathbf{H}^2$  reflects the amount of total variance in the variable accounted for by the factors in the model, indicating the explanatory power of the factor model for that variable.

Step 2. **\*\*Factor Extraction by Iteration\*\***:

- Initial Communalities: Compute the initial communalities as the squared multiple correlations:

$$H_{i(t)}^2 = R_{ii(t)}$$

where  $H_{i(t)}^2$  is the communality of i-th item in the  $t$ -th iteration, and  $R_{ii(t)}$  is the i-th diagonal element of the correlation matrix in the  $t$ -th iteration.

- Extract Factors and Update Communalities:

$$\Lambda_{ij} = \sqrt{\lambda_j} \times v_{ij}$$

$$H_{i(t+1)}^2 = \sum_j \Lambda_{ij}^2$$

$$R_{ii(t+1)} = H_{i(t+1)}^2$$

where  $\Lambda_{ij}$  represents the j-th factor loading for the i-th item,  $\lambda_j$  is the j-th eigenvalue,  $H_{i(t+1)}^2$  is the communality of i-th item in the  $t + 1$ -th iteration, and  $v_{ij}$  is the j-th value of the i-th item in the eigen vector matrix  $\mathbf{v}$ .

Step 3. **\*\*Iterative Refinement\*\***:

- Calculate the Change between  $\mathbf{H}_t^2$  and  $\mathbf{H}_{t+1}^2$ :

$$\Delta H_i^2 = |H_{i(t+1)}^2 - H_{i(t)}^2|$$

where  $\Delta H_i^2$  represents the change in communalities between iterations  $t$  and  $t + 1$ .

- Convergence Criterion: Continue iterating until the change in communalities is less than the specified criterion *criterion*:

$$\sum_i \Delta H_i^2 < \text{criterion}$$

The iterative process is implemented using C++ code to ensure computational speed.

## Value

A list containing:

loadings	The extracted factor loadings.
eigen.value	The eigenvalues of the correlation matrix.
H2	A vector that contains the explanatory power of the factor model for all items.

## Author(s)

Haijiang Qin <Haijiang133@outlook.com>

### Examples

```
library(EFAfactors)
set.seed(123)

##Take the data.bfi dataset as an example.
data(data.bfi)

response <- as.matrix(data.bfi[, 1:25]) ## loading data
response <- na.omit(response) ## Remove samples with NA/missing values

## Transform the scores of reverse-scored items to normal scoring
response[, c(1, 9, 10, 11, 12, 22, 25)] <- 6 - response[, c(1, 9, 10, 11, 12, 22, 25)] + 1

## Run factor.analysis function to extract 5 factors

PAF.obj <- factor.analysis(response, nfact = 5)

## Get the loadings, eigen.value and H2 results.
loadings <- PAF.obj$loadings
eigen.value <- PAF.obj$eigen.value
H2 <- PAF.obj$H2

print(loadings)
print(eigen.value)
print(H2)
```

---

FF

---

*Factor Forest (FF) Powered by An Tuned XGBoost Model for Determining the Number of Factors*


---

### Description

This function will invoke a tuned XGBoost model (Goretzko & Buhner, 2020; Goretzko, 2022; Goretzko & Ruscio, 2024) that can reliably perform the task of determining the number of factors. The maximum number of factors that the network can discuss is 8.

### Usage

```
FF(
  response,
  cor.type = "pearson",
  use = "pairwise.complete.obs",
```

```

    vis = TRUE,
    plot = TRUE
  )

```

### Arguments

response	A required $N \times I$ matrix or data.frame consisting of the responses of $N$ individuals to $I$ items.
cor.type	A character string indicating which correlation coefficient (or covariance) is to be computed. One of "pearson" (default), "kendall", or "spearman". @seealso <a href="#">cor</a> .
use	An optional character string giving a method for computing covariances in the presence of missing values. This must be one of the strings "everything", "all.obs", "complete.obs", "na.or.complete", or "pairwise.complete.obs" (default). @seealso <a href="#">cor</a> .
vis	A Boolean variable that will print the factor retention results when set to TRUE, and will not print when set to FALSE. (default = TRUE)
plot	A Boolean variable that will print the FF plot when set to TRUE, and will not print it when set to FALSE. @seealso <a href="#">plot.FF</a> . (Default = TRUE)

### Details

A total of 500,000 datasets were simulated to extract features for training the tuned XGBoost model (Goretzko & Buhner, 2020; Goretzko, 2022). Each dataset was generated according to the following specifications:

- Factor number:  $F \sim U[1,8]$
- Sample size:  $N \sim U[200,1000]$
- Number of variables per factor:  $v_{pf} \sim U[3,10]$
- Factor correlation:  $fc \sim U[0.0,0.4]$
- Primary loadings:  $pl \sim U[0.35,0.80]$
- Cross-loadings:  $cl \sim U[0.0,0.2]$

A population correlation matrix was created for each data set based on the following decomposition:

$$\Sigma = \Lambda \Phi \Lambda^T + \Delta$$

where  $\Lambda$  is the loading matrix,  $\Phi$  is the factor correlation matrix, and  $\Delta$  is a diagonal matrix, with  $\Delta = 1 - \text{diag}(\Lambda \Phi \Lambda^T)$ . The purpose of  $\Delta$  is to ensure that the diagonal elements of  $\Sigma$  are 1.

The response data for each subject were simulated using the following formula:

$$X_i = L_i + \epsilon_i, \quad 1 \leq i \leq I$$

where  $L_i$  follows a normal distribution  $N(0, \sigma)$ , representing the contribution of latent factors, and  $\epsilon_i$  is the residual term following a standard normal distribution.  $L_i$  and  $\epsilon_i$  are uncorrelated, and  $\epsilon_i$  and  $\epsilon_j$  are also uncorrelated.

For each simulated dataset, a total of 184 features are extracted and compiled into a feature vector. These features include:

- 1. - Number of examinees
- 2. - Number of items
- 3. - Number of eigenvalues greater than 1
- 4. - Proportion of variance explained by the 1st eigenvalue
- 5. - Proportion of variance explained by the 2nd eigenvalue
- 6. - Proportion of variance explained by the 3rd eigenvalue
- 7. - Number of eigenvalues greater than 0.7
- 8. - Standard deviation of the eigenvalues
- 9. - Number of eigenvalues accounting for 50
- 10. - Number of eigenvalues accounting for 75
- 11. - L1-norm of the correlation matrix
- 12. - Frobenius-norm of the correlation matrix
- 13. - Maximum-norm of the correlation matrix
- 14. - Average of the off-diagonal correlations
- 15. - Spectral-norm of the correlation matrix
- 16. - Number of correlations smaller or equal to 0.1
- 17. - Average of the initial communality estimates
- 18. - Determinant of the correlation matrix
- 19. - Measure of sampling adequacy (MSA after Kaiser, 1970)
- 20. - Gini coefficient (Gini, 1921) of the correlation matrix
- 21. - Kolm measure of inequality (Kolm, 1999) of the correlation matrix
- 21. - Number of factors retained by the PA method @seealso [PA](#)
- 23. - Number of factors retained by the EKC method @seealso [EKC](#)
- 24. - Number of factors retained by the CD method @seealso [CD](#)
- 25-104. - Eigenvalues from Principal Component Analysis (PCA), padded with -1000 if insufficient
- 105-184. - Eigenvalues from Factor Analysis (FA), fixed at 1 factor, padded with -1000 if insufficient

The code for the FF function is implemented based on the publicly available code by Goretzko & Buhner (2020) (<https://osf.io/mvrau/>). The Tuned XGBoost Model is also obtained from this site. However, to meet the requirements for a streamlined R package, we can only save the core components of the Tuned XGBoost Model. Although these non-core parts do not affect performance, they include a lot of information about the model itself, such as the number of features, subsets of samples, and data from the training process, among others. For the complete Tuned XGBoost Model, please download it from <https://osf.io/mvrau/>.

## Value

An object of class FF is a list containing the following components:

nfact	The number of factors to be retained.
probability	A matrix containing the probabilities for factor numbers ranging from 1 to 8 (1x8), where the number in the f-th column represents the probability that the number of factors for the response is f.
features	A matrix (1x184) containing all the features for determining the number of factors by the tuned XGBoost Model.

## References

Goretzko, D., & Buhner, M. (2020). One model to rule them all? Using machine learning algorithms to determine the number of factors in exploratory factor analysis. *Psychol Methods*, 25(6), 776-786. <https://doi.org/10.1037/met0000262>.

Goretzko, D. (2022). Factor Retention in Exploratory Factor Analysis With Missing Data. *Educ Psychol Meas*, 82(3), 444-464. <https://doi.org/10.1177/00131644211022031>.

## Examples

```
library(EFAfactors)
set.seed(123)

##Take the data.bfi dataset as an example.
data(data.bfi)

response <- as.matrix(data.bfi[, 1:25]) ## loading data
response <- na.omit(response) ## Remove samples with NA/missing values

## Transform the scores of reverse-scored items to normal scoring
response[, c(1, 9, 10, 11, 12, 22, 25)] <- 6 - response[, c(1, 9, 10, 11, 12, 22, 25)] + 1

## Run FF function with default parameters.
## Not run:
FF.obj <- FF(response)

print(FF.obj)

plot(FF.obj)

## Get the probability and nfact results.
probability <- FF.obj$probability
nfact <- FF.obj$nfact

print(probability)
print(nfact)

## End(Not run)
```

## Description

This function simulates data with  $n_{fact}$  factors based on empirical data. It represents the simulation data part of the [CD](#) function and the [CDF](#) function. This function improves upon [GenDataPopulation](#) by utilizing C++ code to achieve faster data simulation.

## Usage

```
GenData(
  response,
  nfact = 1,
  N.pop = 10000,
  Max.Trials = 5,
  lr = 1,
  cor.type = "pearson",
  use = "pairwise.complete.obs",
  isSort = FALSE
)
```

## Arguments

response	A required $N \times I$ matrix or data.frame consisting of the responses of $N$ individuals to $I$ items.
nfact	The number of factors to extract in factor analysis. (default = 1)
N.pop	Size of finite populations for simulating. (default = 10,000)
Max.Trials	The maximum number of consecutive trials without obtaining a lower RMSR. (default = 5)
lr	The learning rate for updating the correlation matrix during iteration. (default = 1)
cor.type	A character string indicating which correlation coefficient (or covariance) is to be computed. One of "pearson" (default), "kendall", or "spearman". @seealso <a href="#">cor</a> .
use	An optional character string specifying a method for computing covariances in the presence of missing values. This must be one of the strings "everything", "all.obs", "complete.obs", "na.or.complete", or "pairwise.complete.obs" (default). @seealso <a href="#">cor</a> .
isSort	Logical, determines whether the simulated data needs to be sorted in descending order. (default = FALSE)

### Details

The core idea of GenData is to start with the empirical data's correlation matrix and iteratively approach data with  $n_{fact}$  factors. Any value in the simulated data must come from the empirical data. The specific steps of GenData are as follows:

- (1) Use the empirical data ( $\mathbf{Y}_{emp}$ ) correlation matrix as the target,  $\mathbf{R}_{targ}$ .
- (2) Simulate scores for  $N.pop$  examinees on  $n_{fact}$  factors using a multivariate standard normal distribution:

$$\mathbf{S}_{(N.pop \times n_{fact})} \sim \mathcal{N}(0, 1)$$

Simulate noise for  $N.pop$  examinees on  $I$  items:

$$\mathbf{U}_{(N.pop \times I)} \sim \mathcal{N}(0, 1)$$

- (3) Initialize  $\mathbf{R}_{temp} = \mathbf{R}_{targ}$ , and set the minimum Root Mean Square Residual  $RMSR_{min} = \text{Inf}$ . Start the iteration process.
- (4) Extract  $n_{fact}$  factors from  $\mathbf{R}_{temp}$ , and obtain the factor loadings matrix  $\mathbf{L}_{share}$ . Ensure that the first element of  $\mathbf{L}_{share}$  is positive to standardize the direction.
- (5) Calculate the unique factor matrix  $\mathbf{L}_{uniq, (I \times 1)}$ :

$$L_{uniq,i} = \sqrt{1 - \sum_{j=1}^{n_{fact}} L_{share,i,j}^2}$$

- (6) Calculate the simulated data  $\mathbf{Y}_{sim}$ :

$$Y_{sim,i,j} = \mathbf{S}_i \mathbf{L}_{share,j}^T + U_{i,j} L_{uniq,i}$$

- (7) Compute the correlation matrix of the simulated data,  $\mathbf{R}_{simu}$ .
- (8) Calculate the residual correlation matrix  $\mathbf{R}_{resi}$  between the target matrix  $\mathbf{R}_{targ}$  and the simulated data's correlation matrix  $\mathbf{R}_{simu}$ :

$$\mathbf{R}_{resi} = \mathbf{R}_{targ} - \mathbf{R}_{simu}$$

- (9) Calculate the current RMSR:

$$RMSR_{cur} = \sqrt{\frac{\sum_{i < j} \mathbf{R}_{resi,i,j}^2}{0.5 \times (I^2 - I)}}$$

- (10) If  $RMSR_{cur} < RMSR_{min}$ , update  $\mathbf{R}_{temp} = \mathbf{R}_{temp} + lr \times \mathbf{R}_{resi}$ ,  $RMSR_{min} = RMSR_{cur}$ , set  $\mathbf{R}_{min, resi} = \mathbf{R}_{resi}$ , and reset the count of consecutive trials without improvement  $cou = 0$ . If  $RMSR_{cur} \geq RMSR_{min}$ , update  $\mathbf{R}_{temp} = \mathbf{R}_{temp} + 0.5 \times cou \times lr \times \mathbf{R}_{min, resi}$  and increment  $cou = cou + 1$ .
- (11) Repeat steps (4) through (10) until  $cou \geq Max.Trials$ .

Of course C++ code is used to speed up.

### Value

A  $N.pop \times I$  matrix containing the simulated data.

## References

Ruscio, J., & Roche, B. (2012). Determining the number of factors to retain in an exploratory factor analysis using comparison data of known factorial structure. *Psychological Assessment*, 24, 282–292. <http://dx.doi.org/10.1037/a0025697>.

---

Hull

---

*the Hull Approach*


---

## Description

The Hull method is a heuristic approach used to determine the optimal number of common factors in factor analysis. It evaluates models with increasing numbers of factors and uses goodness-of-fit indices relative to the model degrees of freedom to select the best-fitting model. The method is known for its effectiveness and reliability compared to other methods like the scree plot.

## Usage

```
Hull(
  response,
  fa = "pc",
  nfact = 1,
  cor.type = "pearson",
  use = "pairwise.complete.obs",
  vis = TRUE,
  plot = TRUE
)
```

## Arguments

response	A required $N \times I$ matrix or data.frame consisting of the responses of $N$ individuals to $\times I$ items.
fa	A string that determines the method used to obtain eigenvalues in PA. If 'pc', it represents Principal Component Analysis (PCA); if 'fa', it represents Principal Axis Factoring (a widely used Factor Analysis method; @seealso <a href="#">factor.analysis</a> ; Auerswald & Moshagen, 2019). (Default = 'pc')
nfact	A numeric value that specifies the number of factors to extract, only effective when <code>fa = 'fa'</code> . (Default = 1)
cor.type	A character string indicating which correlation coefficient (or covariance) is to be computed. One of "pearson" (default), "kendall", or "spearman". @seealso <a href="#">cor</a> .
use	an optional character string giving a method for computing covariances in the presence of missing values. This must be one of the strings "everything", "all.obs", "complete.obs", "na.or.complete", or "pairwise.complete.obs" (default). @seealso <a href="#">cor</a> .

vis	A Boolean variable that will print the factor retention results when set to TRUE, and will not print when set to FALSE. (default = TRUE)
plot	A Boolean variable that will print the CD plot when set to TRUE, and will not print it when set to FALSE. @seealso <a href="#">plot.Hull</a> . (Default = TRUE)

## Details

The Hull method (Lorenzo-Seva & Timmerman, 2011) is a heuristic approach used to determine the number of common factors in factor analysis. This method is similar to non-graphical variants of Cattell's scree plot but relies on goodness-of-fit indices relative to the model degrees of freedom. The Hull method finds the optimal number of factors by following these steps:

1. Calculate the goodness-of-fit index (CFI) and model degrees of freedom (df; Lorenzo-Seva & Timmerman, 2011;  $df = IF - 0.5F * (F - 1)$ ,  $I$  is the number of items, and  $F$  is the number of factors) for models with an increasing number of factors, up to a prespecified maximum, which is equal to the
2. nfact of [PA](#) method. the GOF will always be Comparative Fit Index (CFI), for it performs best under various conditions than other GOF (Auerswald & Moshagen, 2019; Lorenzo-Seva & Timmerman, 2011), such as RMSEA and SRMR. @seealso [EFAindex](#)
3. Identify and exclude solutions that are less complex (with fewer factors) but have a higher fit index.
4. Further exclude solutions if their fit indices fall below the line connecting adjacent viable solutions.
5. Determine the number of factors where the ratio of the difference in goodness-of-fit indices to the difference in degrees of freedom is maximized.

## Value

A list with the following components:

nfact	The optimal number of factors according to the Hull method.
CFI	A numeric vector of CFI values for each number of factors considered.
df	A numeric vector of model degrees of freedom for each number of factors considered.
Hull.CFI	A numeric vector of CFI values with points below the convex Hull curve removed.
Hull.df	A numeric vector of model degrees of freedom with points below the convex Hull curve removed.

## Author(s)

Haijiang Qin <Haijiang133@outlook.com>

## References

Auerswald, M., & Moshagen, M. (2019). How to determine the number of factors to retain in exploratory factor analysis: A comparison of extraction methods under realistic conditions. *Psychological methods*, 24(4), 468-491. <https://doi.org/https://doi.org/10.1037/met0000200>.

Lorenzo-Seva, U., Timmerman, M. E., & Kiers, H. A. L. (2011). The Hull Method for Selecting the Number of Common Factors. *Multivariate Behavioral Research*, 46(2), 340-364. <https://doi.org/10.1080/00273171.2011.564>

## Examples

```
library(EFAfactors)
set.seed(123)

##Take the data.bfi dataset as an example.
data(data.bfi)

response <- as.matrix(data.bfi[, 1:25]) ## loading data
response <- na.omit(response) ## Remove samples with NA/missing values

## Transform the scores of reverse-scored items to normal scoring
response[, c(1, 9, 10, 11, 12, 22, 25)] <- 6 - response[, c(1, 9, 10, 11, 12, 22, 25)] + 1

## Run EKC function with default parameters.

Hull.obj <- Hull(response)

print(Hull.obj)

plot(Hull.obj)

## Get the CFI, df and nfact results.
CFI <- Hull.obj$CFI
df <- Hull.obj$df
nfact <- Hull.obj$nfact

print(CFI)
print(df)
print(nfact)
```

## Description

This function implements the Kaiser-Guttman criterion (Guttman, 1954; Kaiser, 1960) for determining the number of factors to retain in factor analysis. It is based on the eigenvalues of the correlation matrix of the responses. According to the criterion, factors are retained if their corresponding eigenvalues are greater than 1.

## Usage

```
KGC(
  response,
  fa = "pc",
  nfact = 1,
  cor.type = "pearson",
  use = "pairwise.complete.obs",
  vis = TRUE,
  plot = TRUE
)
```

## Arguments

<code>response</code>	A required $N \times I$ matrix or <code>data.frame</code> consisting of the responses of $N$ individuals to $I$ items.
<code>fa</code>	A string that determines the method used to obtain eigenvalues. If 'pc', it represents Principal Component Analysis (PCA); if 'fa', it represents Principal Axis Factoring (a widely used Factor Analysis method; @seealso <a href="#">factor.analysis</a> ; Auerswald & Moshagen, 2019). (Default = 'pc')
<code>nfact</code>	A numeric value that specifies the number of factors to extract, only effective when <code>fa = 'fa'</code> . (Default = 1)
<code>cor.type</code>	A character string indicating which correlation coefficient (or covariance) is to be computed. One of "pearson" (default), "kendall", or "spearman". @seealso <a href="#">cor</a> .
<code>use</code>	An optional character string giving a method for computing covariances in the presence of missing values. This must be one of the strings "everything", "all.obs", "complete.obs", "na.or.complete", or "pairwise.complete.obs" (default). @seealso <a href="#">cor</a> .
<code>vis</code>	A Boolean variable that will print the factor retention results when set to TRUE, and will not print when set to FALSE. (default = TRUE)
<code>plot</code>	A Boolean variable that will print the KGC plot when set to TRUE, and will not print it when set to FALSE. @seealso <a href="#">plot.KGC</a> . (Default = TRUE)

## Value

An object of class `KGC` is a list containing the following components:

<code>nfact</code>	The number of factors to be retained.
<code>eigen.value</code>	A vector containing the empirical eigenvalues

## References

Guttman, L. (1954). Some necessary conditions for common-factor analysis. *Psychometrika*, 19, 149–161. <http://dx.doi.org/10.1007/BF02289162>.

Kaiser, H. F. (1960). The application of electronic computers to factor analysis. *Educational and Psychological Measurement*, 20, 141–151. <http://dx.doi.org/10.1177/001316446002000116>.

## Examples

```
library(EFAfactors)
set.seed(123)

##Take the data.bfi dataset as an example.
data(data.bfi)

response <- as.matrix(data.bfi[, 1:25]) ## loading data
response <- na.omit(response) ## Remove samples with NA/missing values

## Transform the scores of reverse-scored items to normal scoring
response[, c(1, 9, 10, 11, 12, 22, 25)] <- 6 - response[, c(1, 9, 10, 11, 12, 22, 25)] + 1

## Run KGC function with default parameters.

KGC.obj <- KGC(response)

print(KGC.obj)

plot(KGC.obj)

## Get the eigen.value, eigen.ref and nfact results.
eigen.value <- KGC.obj$eigen.value
nfact <- KGC.obj$nfact

print(eigen.value)
print(nfact)
```

---

load\_DNN

---

*Load the Trained Deep Neural Network (DNN)*


---

## Description

Loads the Pre-Trained Deep Neural Network (DNN) from the `DNN.onnx`. The function uses the `reticulate` package to import the `onnxruntime` Python library and create an inference session for the model.

**Usage**

```
load_DNN()
```

**Value**

An ONNX runtime inference session object for the DNN model.

**Note**

Note that Python and the libraries numpy and onnxruntime are required.

First, please ensure that Python is installed on your computer and that Python is included in the system's PATH environment variable. If not, please download and install it from the official website (<https://www.python.org/>).

If you encounter an error when running this function stating that the numpy and onnxruntime modules are missing:

```
Error in py_module_import(module, convert = convert) :
```

```
ModuleNotFoundError: No module named 'numpy'
```

or

```
Error in py_module_import(module, convert = convert) :
```

```
ModuleNotFoundError: No module named 'onnxruntime'
```

this means that the numpy or onnxruntime library is missing from your Python environment. If you are using Windows or macOS, please run the command `pip install numpy` or `pip install onnxruntime` in Command Prompt or Windows PowerShell (Windows), or Terminal (macOS). If you are using Linux, please ensure that pip is installed and use the command `pip install numpy` or `pip install onnxruntime` to install the missing libraries.

**See Also**

[DNN\\_predictor](#)

---

load\_scaler

---

*Load the Scaler for the Pre-Trained Deep Neural Network (DNN)*


---

**Description**

Loads the scaler object within the EFAfactors package. This object is a list containing a mean vector and a standard deviation vector, which were computed from the 10,000,000 datasets [data.datasets](#) used for training the Pre-Trained Deep Neural Network (DNN). It serves as a tool for normalizing features in [DNN\\_predictor](#).

**Usage**

```
load_scaler()
```

**Value**

scaler objective.

**See Also**

[DNN\\_predictor](#), [normalizer](#), [data.datasets](#), [data.scaler](#)

**Examples**

```
library(EFAfactors)

scaler <- load_scaler()
print(scaler)
```

---

load\_xgb

*Load the Tuned XGBoost Model*

---

**Description**

Loads the tuned XGBoost model object within the EFAfactors package into the global environment and retrieves it for use. Only the core model is retained to reduce the size.

**Usage**

```
load_xgb()
```

**Value**

The tuned XGBoost model object

**Examples**

```
library(EFAfactors)

xgb_model <- load_xgb()
print(xgb_model)
```

---

model.xgb	<i>the Tuned XGBoost Model for Determining the Number of Facotrs</i>
-----------	--

---

**Description**

the Tuned XGBoost Model for Determining the Number of Facotrs

**Format**

An object of class TuneModel is the Tuned XGBoost Model for Determining the Number of Facotrs

**See Also**

[FF](#), [load\\_xgb](#)

**Examples**

```
data(model.xgb)
print(model.xgb)

model.xgb <- load_xgb()
print(model.xgb)
```

---

normalizor	<i>Feature Normalization</i>
------------	------------------------------

---

**Description**

This function normalizes a matrix of features using precomputed means and standard deviations. The function automatically runs [load\\_scaler](#) to read the standard deviations and means of the features, which are organized into a list object named scaler. These means and standard deviations are computed from the 10,000,000 datasets [data.datasets](#) for training the Pre-Trained Deep Neural Network (DNN).

**Usage**

```
normalizor(features)
```

**Arguments**

features	A numeric matrix where each row represents an observation and each column represents a feature.
----------	---

### Details

The function applies z-score normalization to each element in the features matrix. It uses the `scaler` object, which is expected to contain precomputed means and standard deviations for each feature. The normalized value for each element is computed as:

$$z = \frac{x - \mu}{\sigma}$$

where  $x$  is the original value,  $\mu$  is the mean, and  $\sigma$  is the standard deviation.

### Value

A matrix of the same dimensions as features, where each feature has been normalized.

### See Also

[DNN\\_predictor](#), [load\\_scaler](#), [data.datasets](#), [data.scaler](#)

---

PA

*Parallel Analysis*

---

### Description

This function performs Parallel Analysis (PA), which is a method used to determine the number of factors to retain in exploratory factor analysis. It compares the empirical eigenvalues with those obtained from simulated random data to identify the point where the observed eigenvalues are larger than those expected by chance. The number of empirical eigenvalues that are greater than the corresponding reference eigenvalues is the number of factors recommended to be retained by the PA method.

### Usage

```
PA(  
  response,  
  fa = "pc",  
  n.iter = 100,  
  type = "quant",  
  nfact = 1,  
  quant = 0.95,  
  cor.type = "pearson",  
  use = "pairwise.complete.obs",  
  vis = TRUE,  
  plot = TRUE  
)
```

### Arguments

response	A required $N \times I$ matrix or data.frame consisting of the responses of $N$ individuals to $I$ items.
fa	A string that determines the method used to obtain eigenvalues in PA. If 'pc', it represents Principal Component Analysis (PCA); if 'fa', it represents Principal Axis Factoring (a widely used Factor Analysis method; @seealso <a href="#">factor.analysis</a> ; Auerswald & Moshagen, 2019). (Default = 'pc')
n.iter	A numeric value that determines the number of simulations for the random data. (Default = 100)
type	A string that determines the method used to calculate the reference eigenvalues from the simulated data. If 'mean', the reference eigenvalue (eigen.ref) is the mean of the simulated eigenvalues (eigen.sim); if 'quant', the reference eigenvalue is the quant percentile of eigen.sim. (Default = 'quant')
nfact	A numeric value that specifies the number of factors to extract, only effective when fa = 'fa'. (Default = 1)
quant	A numeric value between 0 and 1, representing the quantile to be used for the reference eigenvalues calculation when type = 'quant'. (Default = 0.95)
cor.type	A character string indicating the correlation coefficient (or covariance) to be computed. One of "pearson" (default), "kendall", or "spearman". @seealso <a href="#">cor</a> .
use	An optional character string specifying the method for computing covariances when there are missing values. This must be one of "everything", "all.obs", "complete.obs", "na.or.complete", or "pairwise.complete.obs" (default). @seealso <a href="#">cor</a> .
vis	A Boolean that determines whether to print the factor retention results. Set to TRUE to print, or FALSE to suppress output. (Default = TRUE)
plot	A Boolean that determines whether to display the PA plot. Set to TRUE to show the plot, or FALSE to suppress it. @seealso <a href="#">plot.PA</a> . (Default = TRUE)

### Details

This function performs Parallel Analysis (PA; Horn, 1965; Auerswald & Moshagen, 2019) to determine the number of factors to retain. PA is a widely used method and is considered the "gold standard" for factor retention due to its high accuracy and stability, although it may underperform compared to methods like CD or EKC under certain conditions. The core idea of PA is to simulate random data multiple times, for example, 100 times, and compute the eigenvalues from each simulation. These simulated eigenvalues are then processed using either the mean or a quantile method to obtain the reference eigenvalues, such as the  $i$ -th reference eigenvalue  $\lambda_{i,ref}$ . The relationship between the  $i$ -th empirical eigenvalue  $\lambda_i$  and  $\lambda_{i,ref}$  indicates whether the  $i$ -th factor should be retained. If  $\lambda_i > \lambda_{i,ref}$ , it suggests that the explanatory power of the  $i$ -th factor from the original data is stronger than that of the  $i$ -th factor from the random data, and therefore the factor should be retained. Conversely, if  $\lambda_i \leq \lambda_{i,ref}$ , it indicates that the explanatory power of the  $i$ -th factor from the original data is weaker or equal to that of the random data, making it indistinguishable from noise, and thus the factor should not be retained. So,

$$F = \sum_{i=1}^I I(\lambda_i > \lambda_{i,ref})$$

Here,  $\lfloor F \rfloor$  represents the number of factors determined by the EKC, and  $I(\cdot)$  is the indicator function, which equals 1 when the condition is true, and 0 otherwise.

Auerswald & Moshagen (2019) found that the most accurate results for PA were obtained when using PCA to extract eigenvalues and using the 95th percentile of the simulated eigenvalues to calculate the reference eigenvalues. Therefore, the recommended settings for this function are `fa = 'pc'`, `type = 'quant'`, and `quant = 0.95`.

### Value

An object of class PA, which is a list containing the following components:

<code>nfact</code>	The number of factors to retain.
<code>fa</code>	Indicates the method used to obtain eigenvalues in PA. 'pc' represents Principal Component Analysis, and 'fa' represents Principal Axis Factoring.
<code>type</code>	Indicates the method used to calculate <code>eigen.ref</code> . If 'mean', <code>eigen.ref</code> is the mean of <code>eigen.sim</code> ; if 'quant', <code>eigen.ref</code> is the quant percentile of <code>eigen.sim</code> .
<code>eigen.value</code>	A vector containing the empirical eigenvalues.
<code>eigen.ref</code>	A vector containing the reference eigenvalues, which depend on <code>type</code> .
<code>eigen.sim</code>	A matrix containing the simulated eigenvalues for all iterations.

### Author(s)

Haijiang Qin <Haijiang133@outlook.com>

### References

- Auerswald, M., & Moshagen, M. (2019). How to determine the number of factors to retain in exploratory factor analysis: A comparison of extraction methods under realistic conditions. *Psychological methods*, 24(4), 468-491. <https://doi.org/10.1037/met0000200>.
- Horn, J. L. (1965). A rationale and test for the number of factors in factor analysis. *Psychometrika*, 30, 179-185. <http://dx.doi.org/10.1007/BF02289447>.

### Examples

```
library(EFAfactors)
set.seed(123)

##Take the data.bfi dataset as an example.
data(data.bfi)

response <- as.matrix(data.bfi[, 1:25]) ## loading data
response <- na.omit(response) ## Remove samples with NA/missing values

## Transform the scores of reverse-scored items to normal scoring
response[, c(1, 9, 10, 11, 12, 22, 25)] <- 6 - response[, c(1, 9, 10, 11, 12, 22, 25)] + 1

## Run PA function with default parameters.
```

```
PA.obj <- PA(response)

print(PA.obj)

plot(PA.obj)

## Get the eigen.value, eigen.ref and nfact results.
eigen.value <- PA.obj$eigen.value
eigen.ref <- PA.obj$eigen.ref
nfact <- PA.obj$nfact

print(eigen.value)
print(eigen.ref)
print(nfact)
```

---

plot.CD

*Plot Comparison Data for Factor Analysis*

---

### Description

This function generates a Comparison Data plot to visualize the Root Mean Square Error (RMSE) of eigenvalues for various numbers of factors. This plot helps in evaluating the fit of different factor models and identifying the optimal number of factors based on RMSE values.

### Usage

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

### Arguments

x	An object of class CD, representing the results to be plotted.
...	Additional arguments to be passed to the plotting function.

### Value

None. This function is used for side effects (plotting).

### See Also

[CD](#)

### Examples

```
library(EFAfactors)
set.seed(123)

##Take the data.bfi dataset as an example.
data(data.bfi)

response <- as.matrix(data.bfi[, 1:25]) ## loading data
response <- na.omit(response) ## Remove samples with NA/missing values

## Transform the scores of reverse-scored items to normal scoring
response[, c(1, 9, 10, 11, 12, 22, 25)] <- 6 - response[, c(1, 9, 10, 11, 12, 22, 25)] + 1

CD.obj <- CD(response)

## CD plot
plot(CD.obj)
```

---

plot.CDF

---

*Plot Comparison Data Forest (CDF) Classification Probability Distribution*


---

### Description

This function generates a bar plot of the classification probabilities predicted by the Comparison Data Forest for determining the number of factors. The plot displays the probability distribution across different numbers of factors, with each bar representing the probability for a specific number of factors.

### Usage

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

### Arguments

**x** An object of class CDF, representing the results to be plotted.

**...** Additional arguments to be passed to the plotting function.

### Value

None. This function is used for side effects (plotting).

**See Also**[CDF](#)**Examples**

```

library(EFAfactors)
set.seed(123)

## Take the data.bfi dataset as an example.
data(data.bfi)

response <- as.matrix(data.bfi[, 1:25]) ## Load data
response <- na.omit(response) ## Remove samples with NA/missing values

## Transform the scores of reverse-scored items to normal scoring
response[, c(1, 9, 10, 11, 12, 22, 25)] <- 6 - response[, c(1, 9, 10, 11, 12, 22, 25)] + 1

CDF.obj <- CDF(response)

## Plot the CDF probabilities
plot(CDF.obj)

```

---

plot.DNN_predictor	<i>Plot DNN Predictor Classification Probability Distribution</i>
--------------------	---

---

**Description**

This function generates a bar plot of the classification probabilities predicted by the pre-trained deep neural network for determining the number of factors. The plot displays the probability distribution across different numbers of factors, with each bar representing the probability for a specific number of factors. The maximum number of factors that the network can evaluate is 10. The function also annotates each bar with its probability value.

**Usage**

```

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

```

**Arguments**

x	An object of class DNN_predictor, representing the results to be plotted.
...	Additional arguments to be passed to the plotting function.

**Value**

None. This function is used for side effects (plotting).

**See Also**

[DNN\\_predictor](#)

---

plot.EFAhclust

*Plot Hierarchical Cluster Analysis Dendrogram*

---

**Description**

This function generates a dendrogram from hierarchical cluster analysis results. The hierarchical clustering method merges the two nodes with the smallest dissimilarity at each step, forming a new node until all nodes are combined into a single hierarchical structure. The resulting dendrogram represents the hierarchical relationships between items, where items with high correlation are connected by shorter lines, and items with low correlation are connected by longer lines. The height of the tree nodes indicates the dissimilarity between nodes: a greater height reflects a larger difference. Researchers can use this representation to determine if two nodes belong to the same cluster, which in exploratory factor analysis, helps identify whether items belong to the same latent factor.

**Usage**

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

**Arguments**

x	An object of class EFAhclust, representing the results to be plotted.
...	Additional arguments to be passed to the plotting function.

**Value**

None. This function is used for side effects (plotting).

**See Also**

[EFAhclust](#)

**Examples**

```
library(EFAfactors)
set.seed(123)

## Take the data.bfi dataset as an example.
data(data.bfi)

response <- as.matrix(data.bfi[, 1:25]) ## Load data
```

```

response <- na.omit(response) ## Remove samples with NA/missing values

## Transform the scores of reverse-scored items to normal scoring
response[, c(1, 9, 10, 11, 12, 22, 25)] <- 6 - response[, c(1, 9, 10, 11, 12, 22, 25)] + 1

EFAhclust.obj <- EFAhclust(response)

## Plot the hierarchical clustering dendrogram
plot(EFAhclust.obj)

```

---

plot.EFAkmeans

*Plot EFA K-means Clustering Results*


---

## Description

This function creates a plot to visualize the Within-cluster Sum of Squares (WSS) for different numbers of clusters (K) in the context of exploratory factor analysis. The plot helps identify the most appropriate number of factors by showing how WSS decreases as the number of factors (or clusters) increases.

## Usage

```

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

```

## Arguments

x	An object of class EFAkmeans, representing the results to be plotted.
...	Additional arguments to be passed to the plotting function.

## Value

None. This function is used for side effects (plotting).

## See Also

[EFAkmeans](#)

## Examples

```

library(EFAfactors)
set.seed(123)

## Take the data.bfi dataset as an example.
data(data.bfi)

```

```

response <- as.matrix(data.bfi[, 1:25]) ## Load data
response <- na.omit(response) ## Remove samples with NA/missing values

## Transform the scores of reverse-scored items to normal scoring
response[, c(1, 9, 10, 11, 12, 22, 25)] <- 6 - response[, c(1, 9, 10, 11, 12, 22, 25)] + 1

EFAkmeans.obj <- EFAkmeans(response)

## Plot the EFA K-means clustering results
plot(EFAkmeans.obj)

```

---

plot.EFAscreet

*Plots the Scree Plot*


---

## Description

Plots the Scree Plot from an object of class EFAscreet. The scree plot visualizes the eigenvalues of the correlation matrix in descending order and helps in identifying the optimal number of factors by showing where the eigenvalues start to plateau.

## Usage

```

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

```

## Arguments

x	An object of class EFAscreet, which contains the eigenvalues from the factor analysis.
...	Additional arguments to be passed to the plot function (not used).

## Details

The scree plot is a graphical tool used in exploratory factor analysis. It shows the eigenvalues corresponding to the factors. The number of factors is typically determined by finding the point where the plot levels off ("elbow" point).

## Value

A scree plot displaying the eigenvalues against the number of factors.

**Examples**

```

library(EFAfactors)
set.seed(123)

##Take the data.bfi dataset as an example.
data(data.bfi)

response <- as.matrix(data.bfi[, 1:25]) ## loading data
response <- na.omit(response) ## Remove samples with NA/missing values

## Transform the scores of reverse-scored items to normal scoring
response[, c(1, 9, 10, 11, 12, 22, 25)] <- 6 - response[, c(1, 9, 10, 11, 12, 22, 25)] + 1

## Run EFAscreet function with default parameters.

EFAscreet.obj <- EFAscreet(response)

plot(EFAscreet.obj)

```

plot.EFAvote

*Plot Voting Results for Number of Factors***Description**

This function creates a pie chart to visualize the results of a voting method used to determine the number of factors in exploratory factor analysis (EFA). The voting method combines the results from multiple EFA techniques, and the pie chart displays the proportions of votes each number of factors received. Each slice of the pie represents the percentage of votes for a specific number of factors, providing a clear visual representation of the most commonly suggested number of factors.

**Usage**

```

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

```

**Arguments**

x	An object of class EFAvote, representing the results to be plotted.
...	Additional arguments to be passed to the plotting function.

**Value**

None. This function is used for side effects (plotting).

**See Also**[EFAvote](#)**Examples**

```
library(EFAfactors)

nfacts <- c(5, 5, 5, 6, 6, 4)
names(nfacts) <- c("Hull", "CD", "PA", "EKC", "XGB", "DNN")

EFAvote.obj <- EFAvote(votes = nfacts)
plot(EFAvote.obj)
```

plot.EKC

*Plot Empirical Kaiser Criterion (EKC) Plot***Description**

This function generates an Empirical Kaiser Criterion (EKC) plot to visualize the eigenvalues of the actual data. The EKC method helps in determining the number of factors to retain by identifying the point where the eigenvalues exceed the reference eigenvalue. The plot provides a graphical representation to assist in factor selection.

**Usage**

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

**Arguments**

**x**                      An object of class EKC, representing the results to be plotted.

**...**                   Additional arguments to be passed to the plotting function.

**Value**

None. This function is used for side effects (plotting).

**See Also**[EKC](#)

**Examples**

```

library(EFAfactors)
set.seed(123)

##Take the data.bfi dataset as an example.
data(data.bfi)

response <- as.matrix(data.bfi[, 1:25]) ## loading data
response <- na.omit(response) ## Remove samples with NA/missing values

## Transform the scores of reverse-scored items to normal scoring
response[, c(1, 9, 10, 11, 12, 22, 25)] <- 6 - response[, c(1, 9, 10, 11, 12, 22, 25)] + 1

EKC.obj <- EKC(response)

## EKC plot
plot(EKC.obj)

```

---

plot.FF

---

*Plot Factor Forest (FF) Classification Probability Distribution*


---

**Description**

This function generates a bar plot of the classification probabilities predicted by the Factor Forest for determining the number of factors. The plot displays the probability distribution across different numbers of factors, with each bar representing the probability for a specific number of factors. Unlike the deep neural network (DNN) model, the Factor Forest can evaluate up to a maximum of 8 factors. The function also annotates each bar with its probability value.

**Usage**

```

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

```

**Arguments**

**x** An object of class FF, representing the results to be plotted.

**...** Additional arguments to be passed to the plotting function.

**Value**

None. This function is used for side effects (plotting).

**See Also**[FF](#)**Examples**

```
library(EFAfactors)
set.seed(123)

## Take the data.bfi dataset as an example.
data(data.bfi)

response <- as.matrix(data.bfi[, 1:25]) ## Load data
response <- na.omit(response) ## Remove samples with NA/missing values

## Transform the scores of reverse-scored items to normal scoring
response[, c(1, 9, 10, 11, 12, 22, 25)] <- 6 - response[, c(1, 9, 10, 11, 12, 22, 25)] + 1

FF.obj <- FF(response)

## Plot the FF probabilities
plot(FF.obj)
```

---

plot.Hull

---

*Plot Hull Plot for Factor Analysis***Description**

This function creates a Hull plot to visualize the relationship between the Comparative Fit Index (CFI) and the degrees of freedom (df) for a range of models with different numbers of factors. The Hull plot helps in assessing model fit and identifying optimal models.

**Usage**

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

**Arguments**

x	An object of class Hull, representing the results to be plotted.
...	Additional arguments to be passed to the plotting function.

**Value**

None. This function is used for side effects (plotting).

**See Also**[Hull](#)**Examples**

```

library(EFAfactors)
set.seed(123)

##Take the data.bfi dataset as an example.
data(data.bfi)

response <- as.matrix(data.bfi[, 1:25]) ## loading data
response <- na.omit(response) ## Remove samples with NA/missing values

## Transform the scores of reverse-scored items to normal scoring
response[, c(1, 9, 10, 11, 12, 22, 25)] <- 6 - response[, c(1, 9, 10, 11, 12, 22, 25)] + 1

Hull.obj <- CD(response)

## Hull plot
plot(Hull.obj)

```

plot.KGC

*Plot Kaiser-Guttman Criterion (KGC) Plot***Description**

This function generates a Kaiser-Guttman Criterion (KGC) plot to visualize the eigenvalues of the actual data. The Kaiser-Guttman Criterion, also known as the Kaiser criterion, suggests retaining factors with eigenvalues greater than 1. The plot shows the eigenvalues and includes a reference line at 1 to indicate the threshold for factor retention.

**Usage**

```

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

```

**Arguments**

x	An object of class KGC, representing the results to be plotted.
...	Additional arguments to be passed to the plotting function.

**Value**

None. This function is used for side effects (plotting).

**See Also**[KGC](#)**Examples**

```

library(EFAfactors)
set.seed(123)

## Take the data.bfi dataset as an example.
data(data.bfi)

response <- as.matrix(data.bfi[, 1:25]) ## Load data
response <- na.omit(response) ## Remove samples with NA/missing values

## Transform the scores of reverse-scored items to normal scoring
response[, c(1, 9, 10, 11, 12, 22, 25)] <- 6 - response[, c(1, 9, 10, 11, 12, 22, 25)] + 1

KGC.obj <- KGC(response)

## Plot the Kaiser-Guttman Criterion
plot(KGC.obj)

```

plot.PA

*Plot Parallel Analysis Scree Plot***Description**

This function creates a Parallel Analysis (PA) scree plot to compare the eigenvalues of the actual data with the eigenvalues from simulated data. The plot helps in determining the number of factors by visualizing where the eigenvalues of the actual data intersect with those from simulated data. It provides a graphical representation of the results from a parallel analysis to aid in factor selection.

**Usage**

```

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

```

**Arguments**

x	An object of class PA, representing the results to be plotted.
...	Additional arguments to be passed to the plotting function.

**Value**

None. This function is used for side effects (plotting).

**See Also**[PA](#)**Examples**

```
library(EFAfactors)
set.seed(123)

##Take the data.bfi dataset as an example.
data(data.bfi)

response <- as.matrix(data.bfi[, 1:25]) ## loading data
response <- na.omit(response) ## Remove samples with NA/missing values

## Transform the scores of reverse-scored items to normal scoring
response[, c(1, 9, 10, 11, 12, 22, 25)] <- 6 - response[, c(1, 9, 10, 11, 12, 22, 25)] + 1

PA.obj <- PA(response)

## PA plot
plot(PA.obj)
```

---

```
predictLearner.classif.xgboost.earlystop
```

*Prediction Function for the Tuned XGBoost Model with Early Stopping*

---

**Description**

This function performs predictions using a trained XGBoost model with early stopping. The function itself does not have any specific purpose; its existence is solely to ensure the proper operation of [FF](#).

**Usage**

```
## S3 method for class 'classif.xgboost.earlystop'
predictLearner(.learner, .model, .newdata, ...)
```

**Arguments**

<code>.learner</code>	An object representing the learner.
<code>.model</code>	The trained XGBoost model used to make predictions.

`.newdata`      A data frame or matrix containing new observations for which predictions are to be made.

`...`            Additional parameters passed to the predict function in XGBoost.

**Value**

A vector of predicted class labels or a matrix of predicted probabilities.

---

<code>print.CD</code>	<i>Print Comparison Data Method Results</i>
-----------------------	---

---

**Description**

This function prints the number of factors suggested by the Comparison Data (CD) method.

**Usage**

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

**Arguments**

`x`                An object of class CD, representing the results to be printed.

`...`            Additional arguments to be passed to the plotting function.

**Value**

None. This function is used for side effects (printing).

**See Also**

[CD](#)

---

<code>print.CDF</code>	<i>Print Comparison Data Forest (CDF) Results</i>
------------------------	---

---

**Description**

This function prints the number of factors suggested by the Comparison Data Forest.

**Usage**

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

**Arguments**

- x                      An object of class CDF, representing the results to be printed.
- ...                    Additional arguments to be passed to the plotting function.

**Value**

None. This function is used for side effects (printing).

**See Also**

[CDF](#)

---

print.DNN_predictor	<i>Print DNN Predictor Method Results</i>
---------------------	---

---

**Description**

This function prints the number of factors suggested by the deep neural network (DNN) predictor.

**Usage**

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

**Arguments**

- x                      An object of class DNN\_predictor, representing the results to be printed.
- ...                    Additional arguments to be passed to the plotting function.

**Value**

None. This function is used for side effects (printing).

**See Also**

[DNN\\_predictor](#)

---

print.EFadata	<i>Print the EFAsim.data</i>
---------------	------------------------------

---

**Description**

This function prints the items in factors.

**Usage**

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

**Arguments**

x	An object of class EFA.data, representing the results to be printed.
...	Additional arguments to be passed to the plotting function.

**Value**

None. This function is used for side effects (printing).

**See Also**

[EFAsim.data](#)

---

print.EFAhclust	<i>Print EFAhclust Method Results</i>
-----------------	---------------------------------------

---

**Description**

This function prints the number of factors suggested by the EFAhclust method using the Second-Order Difference (SOD) approach.

**Usage**

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

**Arguments**

x	An object of class EFAhclust, representing the results to be printed.
...	Additional arguments to be passed to the plotting function.

**Value**

None. This function is used for side effects (printing).

**See Also**[EFAhclust](#)


---

print.EFAkmeans	<i>Print EFAkmeans Method Results</i>
-----------------	---------------------------------------

---

**Description**

This function prints the number of factors suggested by the EFAkmeans method using the Second-Order Difference (SOD) approach.

**Usage**

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

**Arguments**

x	An object of class EFAkmeans, representing the results to be printed.
...	Additional arguments to be passed to the plotting function.

**Value**

None. This function is used for side effects (printing).

**See Also**[EFAkmeans](#)


---

print.EFAscree	<i>Print the Scree Plot</i>
----------------	-----------------------------

---

**Description**

Prints a brief summary of an object of class EFAscree. This function will display the scree plot of the eigenvalues when called, providing a visual representation of the factor analysis results.

**Usage**

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

**Arguments**

x	An object of class EFAscree, which contains the eigenvalues from the factor analysis.
...	Additional arguments to be passed to the plotting function.

**Value**

None. This function is used for side effects (printing).

---

<code>print.EFAvote</code>	<i>Print Voting Method Results</i>
----------------------------	------------------------------------

---

**Description**

This function prints the number of factors suggested by the voting.

**Usage**

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

**Arguments**

- `x` An object of class EFAvote, representing the results to be printed.
- `...` Additional arguments to be passed to the plotting function.

**Value**

None. This function is used for side effects (printing).

**See Also**

[EFAvote](#)

---

<code>print.EKC</code>	<i>Print Empirical Kaiser Criterion Results</i>
------------------------	---

---

**Description**

This function prints the number of factors suggested by the Empirical Kaiser Criterion (EKC).

**Usage**

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

**Arguments**

- `x` An object of class EKC, representing the results to be printed.
- `...` Additional arguments to be passed to the plotting function.

**Value**

None. This function is used for side effects (printing).

**See Also**

[EKC](#)

---

print.FF	<i>Print Factor Forest (FF) Results</i>
----------	---

---

**Description**

This function prints the number of factors suggested by the Factor Forest.

**Usage**

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

**Arguments**

x	An object of class FF, representing the results to be printed.
...	Additional arguments to be passed to the plotting function.

**Value**

None. This function is used for side effects (printing).

**See Also**

[FF](#)

---

print.Hull	<i>Print Hull Method Results</i>
------------	----------------------------------

---

**Description**

This function prints the number of factors suggested by the Hull method.

**Usage**

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

**Arguments**

`x`                      An object of class `Hull`, representing the results to be printed.

`...`                    Additional arguments to be passed to the `print` function.

**Value**

None. This function is used for side effects (printing).

**See Also**

[Hull](#)

---

`print.KGC`

*Print Kaiser-Guttman Criterion Results*

---

**Description**

This function prints the number of factors suggested by the Kaiser-Guttman Criterion (KGC).

**Usage**

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

**Arguments**

`x`                      An object of class `KGC`, representing the results to be printed.

`...`                    Additional arguments to be passed to the plotting function.

**Value**

None. This function is used for side effects (printing).

**See Also**

[KGC](#)

---

print.PA	<i>Print Parallel Analysis Method Results</i>
----------	---

---

**Description**

This function prints the number of factors suggested by the Parallel Analysis (PA) method.

**Usage**

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

**Arguments**

x	An object of class PA, representing the results to be printed.
...	Additional arguments to be passed to the plotting function.

**Value**

None. This function is used for side effects (printing).

**See Also**

[PA](#)

# Index

af.softmax, 3

CD, 4, 8, 38, 40, 53, 66  
CDF, 6, 40, 55, 67  
check\_python\_libraries, 9, 13, 16  
cor, 5, 7, 13, 14, 17, 19, 23, 28, 30, 32, 34, 37, 40, 42, 45, 51

data.bfi, 10  
data.datasets, 12, 13, 14, 47–50  
data.scaler, 12, 12, 48, 50  
DNN\_predictor, 3, 4, 9, 12, 13, 13, 30, 31, 47, 48, 50, 56, 67

EFAhclust, 14, 16, 31, 56, 69  
EFAindex, 19, 43  
EFAkmeans, 15, 17, 20, 21, 31, 57, 69  
EFAscreet, 22  
EFAsim.data, 24, 68  
EFAvote, 26, 60, 70  
EKC, 15, 27, 31, 38, 60, 71  
extractor.feature.DNN, 14, 30  
extractor.feature.FF, 8, 32

fa, 19  
factor.analysis, 22, 34, 42, 45, 51  
FF, 36, 49, 62, 65, 71

GenData, 6, 8, 9, 40  
GenDataPopulation, 40

hclust, 16–18  
Hull, 42, 63, 72

KGC, 44, 64, 72  
kmeans, 20

load\_DNN, 46  
load\_scaler, 12, 13, 47, 49, 50  
load\_xgb, 48, 49

model.xgb, 49

normalizor, 12, 13, 48, 49

PA, 38, 43, 50, 65, 73  
plot.CD, 5, 53  
plot.CDF, 7, 54  
plot.DNN\_predictor, 14, 55  
plot.EFAhclust, 17, 56  
plot.EFAkmeans, 21, 57  
plot.EFAscreet, 23, 58  
plot.EFAvote, 27, 59  
plot.EKC, 28, 60  
plot.FF, 37, 61  
plot.Hull, 43, 62  
plot.KGC, 45, 63  
plot.PA, 51, 64  
predictLearner.classif.xgboost.earlystop, 65  
print.CD, 66  
print.CDF, 66  
print.DNN\_predictor, 67  
print.EFAdata, 68  
print.EFAhclust, 68  
print.EFAkmeans, 69  
print.EFAscreet, 69  
print.EFAvote, 70  
print.EKC, 70  
print.FF, 71  
print.Hull, 71  
print.KGC, 72  
print.PA, 73

round, 7

VSS, 19