

# Package: factoextra (via r-universe)

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**Type** Package

**Title** Extract and Visualize the Results of Multivariate Data Analyses

**Version** 1.0.7.999

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**Description** Provides some easy-to-use functions to extract and visualize the output of multivariate data analyses, including 'PCA' (Principal Component Analysis), 'CA' (Correspondence Analysis), 'MCA' (Multiple Correspondence Analysis), 'FAMD' (Factor Analysis of Mixed Data), 'MFA' (Multiple Factor Analysis) and 'HMFA' (Hierarchical Multiple Factor Analysis) functions from different R packages. It contains also functions for simplifying some clustering analysis steps and provides 'ggplot2' - based elegant data visualization.

**License** GPL-2

**LazyData** true

**Encoding** UTF-8

**Depends** R (>= 3.1.2), ggplot2 (>= 2.2.0)

**Imports** abind, cluster, dendextend, FactoMineR, ggpubr(>= 0.1.5), grid, stats, reshape2, ggrepel, tidyr

**Suggests** ade4, ca, igraph, MASS, knitr, mclust

**URL** <http://www.sthda.com/english/rpkgs/factoextra>

**BugReports** <https://github.com/kassambara/factoextra/issues>

**Collate** 'cluster\_utilities.R' 'decathlon2.R' 'print.factoextra.R'  
'get\_hmfa.R' 'fviz\_hmfa.R' 'get\_mfa.R' 'fviz\_mfa.R'  
'deprecated.R' 'fviz\_add.R' 'eigenvalue.R' 'utilities.R'  
'dist.R' 'fviz\_dend.R' 'hcut.R' 'get\_pca.R' 'fviz\_cluster.R'  
'eclust.R' 'facto\_summarize.R' 'fviz.R' 'fviz\_ca.R'  
'fviz\_contrib.R' 'fviz\_cos2.R' 'fviz\_ellipses.R' 'fviz\_famd.R'  
'get\_mca.R' 'fviz\_mca.R' 'fviz\_mclust.R' 'fviz\_nbclust.R'  
'fviz\_pca.R' 'fviz\_silhouette.R' 'get\_ca.R'  
'get\_clust\_tendency.R' 'get\_famd.R' 'hkmeans.R' 'housetasks.R'  
'multishapes.R' 'poison.R' 'zzz.R'

**RoxygenNote** 7.1.0

**Repository** <https://kassambara.r-universe.dev>

**RemoteUrl** <https://github.com/kassambara/factoextra>

**RemoteRef** HEAD

**RemoteSha** 1689fc74b938510b773e4a595999e82bdf37104e

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decathlon2	<i>Athletes' performance in decathlon</i>
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**Description**

Athletes' performance during two sporting meetings

**Usage**

```
data("decathlon2")
```

**Format**

A data frame with 27 observations on the following 13 variables.

X100m a numeric vector

Long.jump a numeric vector

Shot.put a numeric vector

High.jump a numeric vector

X400m a numeric vector

X110m.hurdle a numeric vector

Discus a numeric vector

Pole.vault a numeric vector

Javeline a numeric vector

X1500m a numeric vector

Rank a numeric vector corresponding to the rank

Points a numeric vector specifying the point obtained

Competition a factor with levels Decastar OlympicG

**Source**

This data is a subset of decathlon data in FactoMineR package.

**Examples**

```
data(decathlon2)
decathlon.active <- decathlon2[1:23, 1:10]
res.pca <- prcomp(decathlon.active, scale = TRUE)
fviz_pca_biplot(res.pca)
```

**Description**

Deprecated functions. Will be removed in the next version.

- `get_mfa_var_quanti()`. Deprecated. Use `get_mfa_var(res.mfa, "quanti.var")` instead.
- `get_mfa_var_quali()`. Deprecated. Use `get_mfa_var(res.mfa, "quali.var")` instead.
- `get_mfa_group()`. Deprecated. Use `get_mfa_var(res.mfa, "group")` instead.
- `fviz_mfa_ind_starplot()`: Star graph of individuals (draws partial points). Deprecated. Use `fviz_mfa_ind(res.mfa, partial = "All")` instead.
- `fviz_mfa_quanti_var()`: Graph of quantitative variables. Deprecated. Use `fviz_mfa(X, "quanti.var")` instead.
- `fviz_mfa_quali_var()`: Graph of qualitative variables. Deprecated. Use `fviz_mfa(X, "quali.var")` instead.
- `get_hmfa_var_quanti()`. Deprecated. Use `get_hmfa_var(res.hmfa, "quanti.var")` instead.
- `get_hmfa_var_quali()`. Deprecated. Use `get_hmfa_var(res.hmfa, "quali.var")` instead.
- `get_hmfa_group()`. Deprecated. Use `get_hmfa_var(res.hmfa, "group")` instead.
- `fviz_hmfa_ind_starplot()`: Graph of partial individuals. Deprecated. Use `fviz_hmfa_ind(X, partial = "all")` instead.
- `fviz_hmfa_quanti_var()`: Graph of quantitative variables. Deprecated. Use `fviz_hmfa_var(X, "quanti.var")` instead.
- `fviz_hmfa_quali_var()`: Graph of qualitative variables. Deprecated. Use `fviz_hmfa_var(X, "quali.var")` instead.
- `fviz_hmfa_group()`: Graph of the groups representation. Deprecated. Use `fviz_hmfa_var(X, "group")` instead.

**Usage**

```
get_mfa_quanti_var(res.mfa)
```

```
get_mfa_quali_var(res.mfa)
```

```
get_mfa_group(res.mfa)
```

```
fviz_mfa_ind_starplot(X, ...)
```

```
fviz_mfa_group(X, ...)
```

```
fviz_mfa_quanti_var(X, ...)
```

```
fviz_mfa_quali_var(X, ...)
```

```
get_hmfa_quanti_var(res.hmfa)
get_hmfa_quali_var(res.hmfa)
get_hmfa_group(res.hmfa)
fviz_hmfa_quanti_var(X, ...)
fviz_hmfa_quali_var(X, ...)
fviz_hmfa_ind_starplot(X, ...)
fviz_hmfa_group(X, ...)
```

### Arguments

res.mfa	an object of class MFA [FactoMineR].
X	an object of class MFA or HMFA [FactoMineR].
...	Other arguments.
res.hmfa	an object of class HMFA [FactoMineR].

### Author(s)

Alboukadel Kassambara <alboukadel.kassambara@gmail.com>

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dist

*Enhanced Distance Matrix Computation and Visualization*

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### Description

Clustering methods classify data samples into groups of similar objects. This process requires some methods for measuring the distance or the (dis)similarity between the observations. Read more: [STHDA website - clarifying distance measures..](#)

- `get_dist()`: Computes a distance matrix between the rows of a data matrix. Compared to the standard `dist()` function, it supports correlation-based distance measures including "pearson", "kendall" and "spearman" methods.
- `fviz_dist()`: Visualizes a distance matrix

### Usage

```
get_dist(x, method = "euclidean", stand = FALSE, ...)
fviz_dist(
  dist.obj,
```

```

order = TRUE,
show_labels = TRUE,
lab_size = NULL,
gradient = list(low = "red", mid = "white", high = "blue")
)

```

### Arguments

x	a numeric matrix or a data frame.
method	the distance measure to be used. This must be one of "euclidean", "maximum", "manhattan", "canberra", "binary", "minkowski", "pearson", "spearman" or "kendall".
stand	logical value; default is FALSE. If TRUE, then the data will be standardized using the function scale(). Measurements are standardized for each variable (column), by subtracting the variable's mean value and dividing by the variable's standard deviation.
...	other arguments to be passed to the function dist() when using get_dist().
dist.obj	an object of class "dist" as generated by the function dist() or get_dist().
order	logical value. if TRUE the ordered dissimilarity image (ODI) is shown.
show_labels	logical value. If TRUE, the labels are displayed.
lab_size	the size of labels.
gradient	a list containing three elements specifying the colors for low, mid and high values in the ordered dissimilarity image. The element "mid" can take the value of NULL.

### Value

- get\_dist(): returns an object of class "dist".
- fviz\_dist(): returns a ggplot2

### Author(s)

Alboukadel Kassambara <alboukadel.kassambara@gmail.com>

### See Also

[dist](#)

### Examples

```

data(USArrests)
res.dist <- get_dist(USArrests, stand = TRUE, method = "pearson")

fviz_dist(res.dist,
  gradient = list(low = "#00AFBB", mid = "white", high = "#FC4E07"))

```

## Description

Provides solution for enhancing the workflow of clustering analyses and ggplot2-based elegant data visualization. Read more: [Visual enhancement of clustering analysis](#).

## Usage

```
eclust(  
  x,  
  FUNcluster = c("kmeans", "pam", "clara", "fanny", "hclust", "agnes", "diana"),  
  k = NULL,  
  k.max = 10,  
  stand = FALSE,  
  graph = TRUE,  
  hc_metric = "euclidean",  
  hc_method = "ward.D2",  
  gap_maxSE = list(method = "firstSEmax", SE.factor = 1),  
  nboot = 100,  
  verbose = interactive(),  
  seed = 123,  
  ...  
)
```

## Arguments

x	numeric vector, data matrix or data frame
FUNcluster	a clustering function including "kmeans", "pam", "clara", "fanny", "hclust", "agnes" and "diana". Abbreviation is allowed.
k	the number of clusters to be generated. If NULL, the gap statistic is used to estimate the appropriate number of clusters. In the case of kmeans, k can be either the number of clusters, or a set of initial (distinct) cluster centers.
k.max	the maximum number of clusters to consider, must be at least two.
stand	logical value; default is FALSE. If TRUE, then the data will be standardized using the function scale(). Measurements are standardized for each variable (column), by subtracting the variable's mean value and dividing by the variable's standard deviation.
graph	logical value. If TRUE, cluster plot is displayed.
hc_metric	character string specifying the metric to be used for calculating dissimilarities between observations. Allowed values are those accepted by the function dist() [including "euclidean", "manhattan", "maximum", "canberra", "binary", "minkowski"] and correlation based distance measures ["pearson", "spearman" or "kendall"]. Used only when FUNcluster is a hierarchical clustering function such as one of "hclust", "agnes" or "diana".

hc_method	the agglomeration method to be used (?hclust): "ward.D", "ward.D2", "single", "complete", "average", ...
gap_maxSE	a list containing the parameters (method and SE.factor) for determining the location of the maximum of the gap statistic (Read the documentation ?cluster::maxSE).
nboot	integer, number of Monte Carlo ("bootstrap") samples. Used only for determining the number of clusters using gap statistic.
verbose	logical value. If TRUE, the result of progress is printed.
seed	integer used for seeding the random number generator.
...	other arguments to be passed to FUNcluster.

### Value

Returns an object of class "eclust" containing the result of the standard function used (e.g., kmeans, pam, hclust, agnes, diana, etc.).

It includes also:

- cluster: the cluster assignement of observations after cutting the tree
- nbclust: the number of clusters
- silinfo: the silhouette information of observations, including \$widths (silhouette width values of each observation), \$clus.avg.widths (average silhouette width of each cluster) and \$avg.width (average width of all clusters)
- size: the size of clusters
- data: a matrix containing the original or the standardized data (if stand = TRUE)

The "eclust" class has method for `fviz_silhouette()`, `fviz_dend()`, `fviz_cluster()`.

### Author(s)

Alboukadel Kassambara <alboukadel.kassambara@gmail.com>

### See Also

[fviz\\_silhouette](#), [fviz\\_dend](#), [fviz\\_cluster](#)

### Examples

```
# Load and scale data
data("USArrests")
df <- scale(USArrests)

# Enhanced k-means clustering
# nboot >= 500 is recommended
res.km <- eclust(df, "kmeans", nboot = 2)
# Silhouette plot
fviz_silhouette(res.km)
# Optimal number of clusters using gap statistics
res.km$nbclust
```



```
# Print result
res.km

## Not run:
# Enhanced hierarchical clustering
res.hc <- eclust(df, "hclust", nboot = 2) # compute hclust
fviz_dend(res.hc) # dendrogram
fviz_silhouette(res.hc) # silhouette plot

## End(Not run)
```

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eigenvalue

*Extract and visualize the eigenvalues/variances of dimensions*

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## Description

Eigenvalues correspond to the amount of the variation explained by each principal component (PC).

- `get_eig()`: Extract the eigenvalues/variances of the principal dimensions
- `fviz_eig()`: Plot the eigenvalues/variances against the number of dimensions
- `get_eigenvalue()`: an alias of `get_eig()`
- `fviz_screplot()`: an alias of `fviz_eig()`

These functions support the results of Principal Component Analysis (PCA), Correspondence Analysis (CA), Multiple Correspondence Analysis (MCA), Factor Analysis of Mixed Data (FAMD), Multiple Factor Analysis (MFA) and Hierarchical Multiple Factor Analysis (HMFA) functions.

## Usage

```
get_eig(X)
```

```
get_eigenvalue(X)
```

```
fviz_eig(
  X,
  choice = c("variance", "eigenvalue"),
  geom = c("bar", "line"),
  barfill = "steelblue",
  barcolor = "steelblue",
  linecolor = "black",
  ncp = 10,
  addlabels = FALSE,
  hjust = 0,
  main = NULL,
  xlab = NULL,
  ylab = NULL,
```

```

  ggtheme = theme_minimal(),
  ...
)

fviz_screplot(...)

```

### Arguments

<code>x</code>	an object of class PCA, CA, MCA, FAMD, MFA and HMFA [FactoMineR]; prcomp and princomp [stats]; dudi, pca, coa and acm [ade4]; ca and mjca [ca package].
<code>choice</code>	a text specifying the data to be plotted. Allowed values are "variance" or "eigenvalue".
<code>geom</code>	a text specifying the geometry to be used for the graph. Allowed values are "bar" for barplot, "line" for lineplot or c("bar", "line") to use both types.
<code>barfill</code>	fill color for bar plot.
<code>barcolor</code>	outline color for bar plot.
<code>linecolor</code>	color for line plot (when geom contains "line").
<code>ncp</code>	a numeric value specifying the number of dimensions to be shown.
<code>addlabels</code>	logical value. If TRUE, labels are added at the top of bars or points showing the information retained by each dimension.
<code>hjust</code>	horizontal adjustment of the labels.
<code>main, xlab, ylab</code>	plot main and axis titles.
<code>ggtheme</code>	function, ggplot2 theme name. Default value is theme_pubr(). Allowed values include ggplot2 official themes: theme_gray(), theme_bw(), theme_minimal(), theme_classic(), theme_void(), ....
<code>...</code>	optional arguments to be passed to the function <a href="#">ggpar</a> .

### Value

- `get_eig()` (or `get_eigenvalue()`): returns a data.frame containing 3 columns: the eigenvalues, the percentage of variance and the cumulative percentage of variance retained by each dimension.
- `fviz_eig()` (or `fviz_screplot()`): returns a ggplot2

### Author(s)

Alboukadel Kassambara <[alboukadel.kassambara@gmail.com](mailto:alboukadel.kassambara@gmail.com)>

### References

<http://www.sthda.com/english/>

### See Also

[fviz\\_pca](#), [fviz\\_ca](#), [fviz\\_mca](#), [fviz\\_mfa](#), [fviz\\_hmfa](#)

**Examples**

```

# Principal Component Analysis
# ++++++
data(iris)
res.pca <- prcomp(iris[, -5], scale = TRUE)

# Extract eigenvalues/variances
get_eig(res.pca)

# Default plot
fviz_eig(res.pca, addlabels = TRUE, ylim = c(0, 85))

# Scree plot - Eigenvalues
fviz_eig(res.pca, choice = "eigenvalue", addlabels=TRUE)

# Use only bar or line plot: geom = "bar" or geom = "line"
fviz_eig(res.pca, geom="line")

## Not run:
# Correspondence Analysis
# ++++++
library(FactoMineR)
data(housetasks)
res.ca <- CA(housetasks, graph = FALSE)
get_eig(res.ca)
fviz_eig(res.ca, linecolor = "#FC4E07",
         barcolor = "#00AFBB", barfill = "#00AFBB")

# Multiple Correspondence Analysis
# ++++++
library(FactoMineR)
data(poison)
res.mca <- MCA(poison, quanti.sup = 1:2,
              quali.sup = 3:4, graph=FALSE)
get_eig(res.mca)
fviz_eig(res.mca, linecolor = "#FC4E07",
         barcolor = "#2E9FDF", barfill = "#2E9FDF")

## End(Not run)

```

---

facto\_summarize

*Subset and summarize the output of factor analyses*


---

**Description**

Subset and summarize the results of Principal Component Analysis (PCA), Correspondence Analysis (CA), Multiple Correspondence Analysis (MCA), Factor Analysis of Mixed Data (FAMD), Multiple Factor Analysis (MFA) and Hierarchical Multiple Factor Analysis (HMFA) functions from several packages.

**Usage**

```
facto_summarize(
  X,
  element,
  node.level = 1,
  group.names,
  result = c("coord", "cos2", "contrib"),
  axes = 1:2,
  select = NULL
)
```

**Arguments**

<code>X</code>	an object of class PCA, CA, MCA, FAMD, MFA and HMFA [FactoMineR]; prcomp and princomp [stats]; dudi, pca, coa and acm [ade4]; ca [ca package]; expoOutput [ExPosition].
<code>element</code>	the element to subset from the output. Possible values are "row" or "col" for CA; "var" or "ind" for PCA and MCA; "mca.cor" for MCA; 'quanti.var', 'quali.var', 'group' or 'ind' for FAMD, MFA and HMFA.
<code>node.level</code>	a single number indicating the HMFA node level.
<code>group.names</code>	a vector containing the name of the groups (by default, NULL and the group are named group.1, group.2 and so on).
<code>result</code>	the result to be extracted for the element. Possible values are the combination of c("cos2", "contrib", "coord")
<code>axes</code>	a numeric vector specifying the axes of interest. Default values are 1:2 for axes 1 and 2.
<code>select</code>	a selection of variables. Allowed values are NULL or a list containing the arguments name, cos2 or contrib. Default is list(name = NULL, cos2 = NULL, contrib = NULL): <ul style="list-style-type: none"> <li>• name: is a character vector containing variable names to be selected</li> <li>• cos2: if cos2 is in [0, 1], ex: 0.6, then variables with a cos2 &gt; 0.6 are selected. if cos2 &gt; 1, ex: 5, then the top 5 variables with the highest cos2 are selected</li> <li>• contrib: if contrib &gt; 1, ex: 5, then the top 5 variables with the highest cos2 are selected.</li> </ul>

**Details**

If  $\text{length}(\text{axes}) > 1$ , then the columns contrib and cos2 correspond to the total contributions and total cos2 of the axes. In this case, the column coord is calculated as  $x^2 + y^2 + \dots$ ; x, y, ... are the coordinates of the points on the specified axes.

**Value**

A data frame containing the (total) coord, cos2 and the contribution for the axes.

**Author(s)**

Alboukadel Kassambara <alboukadel.kassambara@gmail.com>

**References**

<http://www.sthda.com/english/>

**Examples**

```
# Principal component analysis
# ++++++
data(decathlon2)
decathlon2.active <- decathlon2[1:23, 1:10]
res.pca <- prcomp(decathlon2.active, scale = TRUE)

# Summarize variables on axes 1:2
facto_summarize(res.pca, "var", axes = 1:2)[,-1]
# Select the top 5 contributing variables
facto_summarize(res.pca, "var", axes = 1:2,
  select = list(contrib = 5))[,-1]
# Select variables with cos2 >= 0.6
facto_summarize(res.pca, "var", axes = 1:2,
  select = list(cos2 = 0.6))[,-1]
# Select by names
facto_summarize(res.pca, "var", axes = 1:2,
  select = list(name = c("X100m", "Discus", "Javeline")))[,-1]

# Summarize individuals on axes 1:2
facto_summarize(res.pca, "ind", axes = 1:2)[,-1]

# Correspondence Analysis
# ++++++
# Install and load FactoMineR to compute CA
# install.packages("FactoMineR")
library("FactoMineR")
data("housetasks")
res.ca <- CA(housetasks, graph = FALSE)
# Summarize row variables on axes 1:2
facto_summarize(res.ca, "row", axes = 1:2)[,-1]
# Summarize column variables on axes 1:2
facto_summarize(res.ca, "col", axes = 1:2)[,-1]

# Multiple Correspondence Analysis
# ++++++
library(FactoMineR)
data(poison)
res.mca <- MCA(poison, quanti.sup = 1:2,
  quali.sup = 3:4, graph=FALSE)
# Summarize variables on axes 1:2
res <- facto_summarize(res.mca, "var", axes = 1:2)
head(res)
# Summarize individuals on axes 1:2
```

```

res <- facto_summarize(res.mca, "ind", axes = 1:2)
head(res)

# Multiple factor Analysis
# ++++++
library(FactoMineR)
data(poison)
res.mfa <- MFA(poison, group=c(2,2,5,6), type=c("s","n","n","n"),
              name.group=c("desc","desc2","symptom","eat"),
              num.group.sup=1:2, graph=FALSE)
# Summarize categorcial variables on axes 1:2
res <- facto_summarize(res.mfa, "quali.var", axes = 1:2)
head(res)
# Summarize individuals on axes 1:2
res <- facto_summarize(res.mfa, "ind", axes = 1:2)
head(res)

```

**Description**

Generic function to create a scatter plot of multivariate analyse outputs, including PCA, CA, MCA and MFA.

**Usage**

```

fviz(
  X,
  element,
  axes = c(1, 2),
  geom = "auto",
  label = "all",
  invisible = "none",
  labelsize = 4,
  pointsize = 1.5,
  pointshape = 19,
  arrowsize = 0.5,
  habillage = "none",
  addEllipses = FALSE,
  ellipse.level = 0.95,
  ellipse.type = "norm",
  ellipse.alpha = 0.1,
  mean.point = TRUE,
  color = "black",
  fill = "white",
  alpha = 1,
  gradient.cols = NULL,

```

```

col.row.sup = "darkblue",
col.col.sup = "darkred",
select = list(name = NULL, cos2 = NULL, contrib = NULL),
title = NULL,
axes.linetype = "dashed",
repel = FALSE,
col.circle = "grey70",
circlesize = 0.5,
ggtheme = theme_minimal(),
ggp = NULL,
font.family = "",
...
)

```

### Arguments

<code>X</code>	an object of class PCA, CA, MCA, FAMD, MFA and HMFA [FactoMineR]; prcomp and princomp [stats]; dudi, pca, coa and acm [ade4]; ca [ca package]; expoOutput [ExPosition].
<code>element</code>	the element to subset from the output. Possible values are "row" or "col" for CA; "var" or "ind" for PCA and MCA; "mca.cor" for MCA; 'quanti.var', 'quali.var', 'group' or 'ind' for FAMD, MFA and HMFA.
<code>axes</code>	a numeric vector specifying the axes of interest. Default values are 1:2 for axes 1 and 2.
<code>geom</code>	a text specifying the geometry to be used for the graph. Default value is "auto". Allowed values are the combination of c("point", "arrow", "text"). Use "point" (to show only points); "text" to show only labels; c("point", "text") or c("arrow", "text") to show both types.
<code>label</code>	a text specifying the elements to be labelled. Default value is "all". Allowed values are "none" or the combination of c("ind", "ind.sup", "quali", "var", "quanti.sup", "group.sup"). "ind" can be used to label only active individuals. "ind.sup" is for supplementary individuals. "quali" is for supplementary qualitative variables. "var" is for active variables. "quanti.sup" is for quantitative supplementary variables.
<code>invisible</code>	a text specifying the elements to be hidden on the plot. Default value is "none". Allowed values are the combination of c("ind", "ind.sup", "quali", "var", "quanti.sup", "group.sup").
<code>labelsize</code>	font size for the labels
<code>pointsize</code>	the size of points
<code>pointshape</code>	the shape of points
<code>arrowsize</code>	the size of arrows. Controls the thickness of arrows.
<code>habillage</code>	an optional factor variable for coloring the observations by groups. Default value is "none". If X is a PCA object from FactoMineR package, habillage can also specify the supplementary qualitative variable (by its index or name) to be used for coloring individuals by groups (see ?PCA in FactoMineR).

<code>addEllipses</code>	logical value. If TRUE, draws ellipses around the individuals when <code>habillage != "none"</code> .
<code>ellipse.level</code>	the size of the concentration ellipse in normal probability.
<code>ellipse.type</code>	Character specifying frame type. Possible values are "convex", "confidence" or types supported by <code>stat_ellipse()</code> including one of <code>c("t", "norm", "euclid")</code> for plotting concentration ellipses. <ul style="list-style-type: none"> <li>• "convex": plot convex hull of a set o points.</li> <li>• "confidence": plot confidence ellipses around group mean points as <code>coord.ellipse()</code> [in FactoMineR].</li> <li>• "t": assumes a multivariate t-distribution.</li> <li>• "norm": assumes a multivariate normal distribution.</li> <li>• "euclid": draws a circle with the radius equal to level, representing the euclidean distance from the center. This ellipse probably won't appear circular unless <code>coord_fixed()</code> is applied.</li> </ul>
<code>ellipse.alpha</code>	Alpha for ellipse specifying the transparency level of fill color. Use <code>alpha = 0</code> for no fill color.
<code>mean.point</code>	logical value. If TRUE (default), group mean points are added to the plot.
<code>color</code>	color to be used for the specified geometries (point, text). Can be a continuous variable or a factor variable. Possible values include also : "cos2", "contrib", "coord", "x" or "y". In this case, the colors for individuals/variables are automatically controlled by their qualities of representation ("cos2"), contributions ("contrib"), coordinates ( $x^2+y^2$ , "coord"), x values ("x") or y values ("y"). To use automatic coloring (by cos2, contrib, ...), make sure that <code>habillage = "none"</code> .
<code>fill</code>	same as the argument <code>color</code> , but for point fill color. Useful when <code>pointshape = 21</code> , for example.
<code>alpha</code>	controls the transparency of individual and variable colors, respectively. The value can variate from 0 (total transparency) to 1 (no transparency). Default value is 1. Possible values include also : "cos2", "contrib", "coord", "x" or "y". In this case, the transparency for the individual/variable colors are automatically controlled by their qualities ("cos2"), contributions ("contrib"), coordinates ( $x^2+y^2$ , "coord"), x values("x") or y values("y"). To use this, make sure that <code>habillage = "none"</code> .
<code>gradient.cols</code>	vector of colors to use for n-colour gradient. Allowed values include brewer and ggsci color palettes.
<code>col.col.sup, col.row.sup</code>	colors for the supplementary column and row points, respectively.
<code>select</code>	a selection of individuals/variables to be drawn. Allowed values are NULL or a list containing the arguments name, cos2 or contrib: <ul style="list-style-type: none"> <li>• name: is a character vector containing individuals/variables to be drawn</li> <li>• cos2: if cos2 is in [0, 1], ex: 0.6, then individuals/variables with a cos2 &gt; 0.6 are drawn. if cos2 &gt; 1, ex: 5, then the top 5 individuals/variables with the highest cos2 are drawn.</li> <li>• contrib: if contrib &gt; 1, ex: 5, then the top 5 individuals/variables with the highest contrib are drawn</li> </ul>



title	the title of the graph
axes.linetype	linetype of x and y axes.
repel	a boolean, whether to use ggrepel to avoid overplotting text labels or not.
col.circle	a color for the correlation circle. Used only when X is a PCA output.
circlesize	the size of the variable correlation circle.
ggtheme	function, ggplot2 theme name. Default value is theme_pubr(). Allowed values include ggplot2 official themes: theme_gray(), theme_bw(), theme_minimal(), theme_classic(), theme_void(), ....
ggp	a ggplot. If not NULL, points are added to an existing plot.
font.family	character vector specifying font family.
...	Arguments to be passed to the functions ggpubr::ggscatter() & ggpubr::ggpar().

**Value**

a ggplot

**Author(s)**

Alboukadel Kassambara <alboukadel.kassambara@gmail.com>

**Examples**

```
# Principal component analysis
# ++++++
data(decathlon2)
decathlon2.active <- decathlon2[1:23, 1:10]
res.pca <- prcomp(decathlon2.active, scale = TRUE)
fviz(res.pca, "ind") # Individuals plot
fviz(res.pca, "var") # Variables plot

# Correspondence Analysis
# ++++++
# Install and load FactoMineR to compute CA
# install.packages("FactoMineR")
library("FactoMineR")
data("housetasks")
res.ca <- CA(housetasks, graph = FALSE)
fviz(res.ca, "row") # Rows plot
fviz(res.ca, "col") # Columns plot

# Multiple Correspondence Analysis
# ++++++
library(FactoMineR)
data(poison)
res.mca <- MCA(poison, quanti.sup = 1:2,
               quali.sup = 3:4, graph=FALSE)

fviz(res.mca, "ind") # Individuals plot
fviz(res.mca, "var") # Variables plot
```

fviz\_add

*Add supplementary data to a plot***Description**

Add supplementary data to a plot

**Usage**

```
fviz_add(
  ggp,
  df,
  axes = c(1, 2),
  geom = c("point", "arrow"),
  color = "blue",
  addlabel = TRUE,
  labelsize = 4,
  pointsize = 2,
  shape = 19,
  linetype = "dashed",
  repel = FALSE,
  font.family = "",
  ...
)
```

**Arguments**

<code>ggp</code>	a ggplot2 plot.
<code>df</code>	a data frame containing the x and y coordinates
<code>axes</code>	a numeric vector of length 2 specifying the components to be plotted.
<code>geom</code>	a character specifying the geometry to be used for the graph Allowed values are "point" or "arrow" or "text"
<code>color</code>	the color to be used
<code>addlabel</code>	a logical value. If TRUE, labels are added
<code>labelsize</code>	the size of labels. Default value is 4
<code>pointsize</code>	the size of points
<code>shape</code>	point shape when geom ="point"
<code>linetype</code>	the linetype to be used when geom ="arrow"
<code>repel</code>	a boolean, whether to use ggrepel to avoid overplotting text labels or not.
<code>font.family</code>	character vector specifying font family.
<code>...</code>	Additional arguments, not used

**Value**

a ggplot2 plot

**Author(s)**

Alboukadel Kassambara <alboukadel.kassambara@gmail.com>

**References**

<http://www.sthda.com>

**Examples**

```
# Principal component analysis
data(decathlon2)
decathlon2.active <- decathlon2[1:23, 1:10]
res.pca <- prcomp(decathlon2.active, scale = TRUE)

# Visualize variables
p <- fviz_pca_var(res.pca)
print(p)

# Add supplementary variables
coord <- data.frame(PC1 = c(-0.7, 0.9), PC2 = c(0.25, -0.07))
rownames(coord) <- c("Rank", "Points")
print(coord)
fviz_add(p, coord, color = "blue", geom="arrow")
```

---

fviz\_ca

*Visualize Correspondence Analysis*

---

**Description**

Correspondence analysis (CA) is an extension of Principal Component Analysis (PCA) suited to analyze frequencies formed by two categorical variables. `fviz_ca()` provides ggplot2-based elegant visualization of CA outputs from the R functions: `CA` [in `FactoMineR`], `ca` [in `ca`], `coa` [in `ade4`], `correspondence` [in `MASS`] and `expOutput/epCA` [in `ExPosition`]. Read more: [Correspondence Analysis](#)

- `fviz_ca_row()`: Graph of row variables
- `fviz_ca_col()`: Graph of column variables
- `fviz_ca_biplot()`: Biplot of row and column variables
- `fviz_ca()`: An alias of `fviz_ca_biplot()`

**Usage**

```
fviz_ca_row(  
  X,  
  axes = c(1, 2),  
  geom = c("point", "text"),  
  geom.row = geom,  
  shape.row = 19,  
  col.row = "blue",  
  alpha.row = 1,  
  col.row.sup = "darkblue",  
  select.row = list(name = NULL, cos2 = NULL, contrib = NULL),  
  map = "symmetric",  
  repel = FALSE,  
  ...  
)  
  
fviz_ca_col(  
  X,  
  axes = c(1, 2),  
  shape.col = 17,  
  geom = c("point", "text"),  
  geom.col = geom,  
  col.col = "red",  
  col.col.sup = "darkred",  
  alpha.col = 1,  
  select.col = list(name = NULL, cos2 = NULL, contrib = NULL),  
  map = "symmetric",  
  repel = FALSE,  
  ...  
)  
  
fviz_ca_biplot(  
  X,  
  axes = c(1, 2),  
  geom = c("point", "text"),  
  geom.row = geom,  
  geom.col = geom,  
  label = "all",  
  invisible = "none",  
  arrows = c(FALSE, FALSE),  
  repel = FALSE,  
  title = "CA - Biplot",  
  ...  
)  
  
fviz_ca(X, ...)
```

**Arguments**

<code>X</code>	an object of class CA [FactoMineR], ca [ca], coa [ade4]; correspondence [MASS] and expOutput/epCA [ExPosition].
<code>axes</code>	a numeric vector of length 2 specifying the dimensions to be plotted.
<code>geom</code>	a character specifying the geometry to be used for the graph. Allowed values are the combination of c("point", "arrow", "text"). Use "point" (to show only points); "text" to show only labels; c("point", "text") or c("arrow", "text") to show both types.
<code>geom.row, geom.col</code>	as geom but for row and column elements, respectively. Default is geom.row = c("point", "text"), geom.col = c("point", "text").
<code>shape.row, shape.col</code>	the point shapes to be used for row/column variables. Default values are 19 for rows and 17 for columns.
<code>map</code>	character string specifying the map type. Allowed options include: "symmetric", "rowprincipal", "colprincipal", "sympbiplot", "rowgab", "colgab", "rowgreen" and "colgreen". See details
<code>repel</code>	a boolean, whether to use ggrepel to avoid overplotting text labels or not.
<code>...</code>	Additional arguments. <ul style="list-style-type: none"> <li>• in <code>fviz_ca_row()</code> and <code>fviz_ca_col()</code>: Additional arguments are passed to the functions <code>fviz()</code> and <code>ggpubr::ggpar()</code>.</li> <li>• in <code>fviz_ca_biplot()</code> and <code>fviz_ca()</code>: Additional arguments are passed to <code>fviz_ca_row()</code> and <code>fviz_ca_col()</code>.</li> </ul>
<code>col.col, col.row</code>	color for column/row points. The default values are "red" and "blue", respectively. Can be a continuous variable or a factor variable. Allowed values include also : "cos2", "contrib", "coord", "x" or "y". In this case, the colors for row/column variables are automatically controlled by their qualities ("cos2"), contributions ("contrib"), coordinates ( $x^2 + y^2$ , "coord"), x values("x") or y values("y")
<code>col.col.sup, col.row.sup</code>	colors for the supplementary column and row points, respectively.
<code>alpha.col, alpha.row</code>	controls the transparency of colors. The value can variate from 0 (total transparency) to 1 (no transparency). Default value is 1. Allowed values include also : "cos2", "contrib", "coord", "x" or "y" as for the arguments <code>col.col</code> and <code>col.row</code> .
<code>select.col, select.row</code>	a selection of columns/rows to be drawn. Allowed values are NULL or a list containing the arguments name, <code>cos2</code> or <code>contrib</code> : <ul style="list-style-type: none"> <li>• name is a character vector containing column/row names to be drawn</li> <li>• <code>cos2</code> if <code>cos2</code> is in [0, 1], ex: 0.6, then columns/rows with a <code>cos2 &gt; 0.6</code> are drawn. if <code>cos2 &gt; 1</code>, ex: 5, then the top 5 columns/rows with the highest <code>cos2</code> are drawn.</li> <li>• <code>contrib</code> if <code>contrib &gt; 1</code>, ex: 5, then the top 5 columns/rows with the highest <code>contrib</code> are drawn</li> </ul>

label	a character vector specifying the elements to be labelled. Default value is "all". Allowed values are "none" or the combination of c("row", "row.sup", "col", "col.sup"). Use "col" to label only active column variables; "col.sup" to label only supplementary columns; etc
invisible	a character value specifying the elements to be hidden on the plot. Default value is "none". Allowed values are the combination of c("row", "row.sup", "col", "col.sup").
arrows	Vector of two logicals specifying if the plot should contain points (FALSE, default) or arrows (TRUE). First value sets the rows and the second value sets the columns.
title	the title of the graph

### Details

The default plot of (M)CA is a "symmetric" plot in which both rows and columns are in principal coordinates. In this situation, it's not possible to interpret the distance between row points and column points. To overcome this problem, the simplest way is to make an asymmetric plot. This means that, the column profiles must be presented in row space or vice-versa. The allowed options for the argument map are:

- "rowprincipal" or "colprincipal": asymmetric plots with either rows in principal coordinates and columns in standard coordinates, or vice versa. These plots preserve row metric or column metric respectively.
- "symbiplot": Both rows and columns are scaled to have variances equal to the singular values (square roots of eigenvalues), which gives a symmetric biplot but does not preserve row or column metrics.
- "rowgab" or "colgab": Asymmetric maps, proposed by Gabriel & Odoroff (1990), with rows (respectively, columns) in principal coordinates and columns (respectively, rows) in standard coordinates multiplied by the mass of the corresponding point.
- "rowgreen" or "colgreen": The so-called contribution biplots showing visually the most contributing points (Greenacre 2006b). These are similar to "rowgab" and "colgab" except that the points in standard coordinates are multiplied by the square root of the corresponding masses, giving reconstructions of the standardized residuals.

### Value

a ggplot

### Author(s)

Alboukadel Kassambara <alboukadel.kassambara@gmail.com>

### References

<http://www.sthda.com>

### See Also

[get\\_ca](#), [fviz\\_pca](#), [fviz\\_mca](#)

**Examples**

```

# Correspondence Analysis
# ++++++
# Install and load FactoMineR to compute CA
# install.packages("FactoMineR")

library("FactoMineR")
data(housetasks)
head(housetasks)
res.ca <- CA(housetasks, graph=FALSE)

# Biplot of rows and columns
# ++++++
# Symetric Biplot of rows and columns
fviz_ca_biplot(res.ca)

# Asymetric biplot, use arrows for columns
fviz_ca_biplot(res.ca, map = "rowprincipal",
  arrow = c(FALSE, TRUE))

# Keep only the labels for row points
fviz_ca_biplot(res.ca, label = "row")

# Keep only labels for column points
fviz_ca_biplot(res.ca, label = "col")

# Select the top 7 contributing rows
# And the top 3 columns
fviz_ca_biplot(res.ca,
  select.row = list(contrib = 7),
  select.col = list(contrib = 3))

# Graph of row variables
# ++++++

# Control automatically the color of row points
# using the "cos2" or the contributions "contrib"
# cos2 = the quality of the rows on the factor map
# Change gradient color
# Use repel = TRUE to avoid overplotting (slow if many points)
fviz_ca_row(res.ca, col.row = "cos2",
  gradient.cols = c("#00AFBB", "#E7B800", "#FC4E07"),
  repel = TRUE)

# You can also control the transparency
# of the color by the "cos2" or "contrib"
fviz_ca_row(res.ca, alpha.row="contrib")

# Select and visualize some rows with select.row argument.
# - Rows with cos2 >= 0.5: select.row = list(cos2 = 0.5)
# - Top 7 rows according to the cos2: select.row = list(cos2 = 7)

```

```

# - Top 7 contributing rows: select.row = list(contrib = 7)
# - Select rows by names: select.row = list(name = c("Breakfast", "Repairs", "Holidays"))

# Example: Select the top 7 contributing rows
fviz_ca_row(res.ca, select.row = list(contrib = 7))

# Graph of column points
# ++++++

# Control colors using their contributions
fviz_ca_col(res.ca, col.col = "contrib",
  gradient.cols = c("#00AFBB", "#E7B800", "#FC4E07"))

# Select columns with select.col argument
# You can select by contrib, cos2 and name
# as previously described for ind
# Select the top 3 contributing columns
fviz_ca_col(res.ca, select.col = list(contrib = 3))

```

---

fviz\_cluster

*Visualize Clustering Results*


---

## Description

Provides ggplot2-based elegant visualization of partitioning methods including kmeans [stats package]; pam, clara and fanny [cluster package]; dbSCAN [fpc package]; Mclust [mclust package]; HCPC [FactoMineR]; hkmeans [factoextra]. Observations are represented by points in the plot, using principal components if  $\text{ncol}(\text{data}) > 2$ . An ellipse is drawn around each cluster.

## Usage

```

fviz_cluster(
  object,
  data = NULL,
  choose.vars = NULL,
  stand = TRUE,
  axes = c(1, 2),
  geom = c("point", "text"),
  repel = FALSE,
  show.clust.cent = TRUE,
  ellipse = TRUE,
  ellipse.type = "convex",
  ellipse.level = 0.95,
  ellipse.alpha = 0.2,

```



```

    shape = NULL,
    pointsize = 1.5,
    labelsize = 12,
    main = "Cluster plot",
    xlab = NULL,
    ylab = NULL,
    outlier.color = "black",
    outlier.shape = 19,
    outlier.pointsize = pointsize,
    outlier.labelsize = labelsize,
    ggtheme = theme_grey(),
    ...
  )

```

### Arguments

object	an object of class "partition" created by the functions pam(), clara() or fanny() in cluster package; "kmeans" [in stats package]; "dbscan" [in fpc package]; "Mclust" [in mclust]; "hkmeans", "eclust" [in factoextra]. Possible value are also any list object with data and cluster components (e.g.: object = list(data = mydata, cluster = myclust)).
data	the data that has been used for clustering. Required only when object is a class of kmeans or dbscan.
choose.vars	a character vector containing variables to be considered for plotting.
stand	logical value; if TRUE, data is standardized before principal component analysis
axes	a numeric vector of length 2 specifying the dimensions to be plotted.
geom	a text specifying the geometry to be used for the graph. Allowed values are the combination of c("point", "text"). Use "point" (to show only points); "text" to show only labels; c("point", "text") to show both types.
repel	a boolean, whether to use ggrepel to avoid overplotting text labels or not.
show.clust.cent	logical; if TRUE, shows cluster centers
ellipse	logical value; if TRUE, draws outline around points of each cluster
ellipse.type	Character specifying frame type. Possible values are 'convex', 'confidence' or types supported by <a href="#">stat_ellipse</a> including one of c("t", "norm", "euclid").
ellipse.level	the size of the concentration ellipse in normal probability. Passed for ggplot2::stat_ellipse's level. Ignored in 'convex'. Default value is 0.95.
ellipse.alpha	Alpha for frame specifying the transparency level of fill color. Use alpha = 0 for no fill color.
shape	the shape of points.
pointsize	the size of points
labelsize	font size for the labels
main	plot main title.

`xlab, ylab` character vector specifying x and y axis labels, respectively. Use `xlab = FALSE` and `ylab = FALSE` to hide `xlab` and `ylab`, respectively.

`outlier.pointsize, outlier.color, outlier.shape, outlier.labelsize` arguments for customizing outliers, which can be detected only in DBSCAN clustering.

`ggtheme` function, ggplot2 theme name. Default value is `theme_pubr()`. Allowed values include ggplot2 official themes: `theme_gray()`, `theme_bw()`, `theme_minimal()`, `theme_classic()`, `theme_void()`, ....

... other arguments to be passed to the functions `ggscatter` and `ggpar`.

**Value**

return a ggplot.

**Author(s)**

Alboukadel Kassambara <alboukadel.kassambara@gmail.com>

**See Also**

[fviz\\_silhouette](#), [hcut](#), [hkmeans](#), [eclust](#), [fviz\\_dend](#)

**Examples**

```
set.seed(123)

# Data preparation
# ++++++
data("iris")
head(iris)
# Remove species column (5) and scale the data
iris.scaled <- scale(iris[, -5])

# K-means clustering
# ++++++
km.res <- kmeans(iris.scaled, 3, nstart = 10)

# Visualize kmeans clustering
# use repel = TRUE to avoid overplotting
fviz_cluster(km.res, iris[, -5], ellipse.type = "norm")

# Change the color palette and theme
fviz_cluster(km.res, iris[, -5],
  palette = "Set2", ggtheme = theme_minimal())

## Not run:
# Show points only
fviz_cluster(km.res, iris[, -5], geom = "point")
# Show text only
fviz_cluster(km.res, iris[, -5], geom = "text")
```

```

# PAM clustering
# ++++++
require(cluster)
pam.res <- pam(iris.scaled, 3)
# Visualize pam clustering
fviz_cluster(pam.res, geom = "point", ellipse.type = "norm")

# Hierarchical clustering
# ++++++
# Use hcut() which compute hclust and cut the tree
hc.cut <- hcut(iris.scaled, k = 3, hc_method = "complete")
# Visualize dendrogram
fviz_dend(hc.cut, show_labels = FALSE, rect = TRUE)
# Visualize cluster
fviz_cluster(hc.cut, ellipse.type = "convex")

## End(Not run)

```

---

fviz\_contrib

*Visualize the contributions of row/column elements*


---

## Description

This function can be used to visualize the contribution of rows/columns from the results of Principal Component Analysis (PCA), Correspondence Analysis (CA), Multiple Correspondence Analysis (MCA), Factor Analysis of Mixed Data (FAMD), and Multiple Factor Analysis (MFA) functions.

## Usage

```

fviz_contrib(
  X,
  choice = c("row", "col", "var", "ind", "quanti.var", "quali.var", "group",
    "partial.axes"),
  axes = 1,
  fill = "steelblue",
  color = "steelblue",
  sort.val = c("desc", "asc", "none"),
  top = Inf,
  xtckslab.rt = 45,
  ggtheme = theme_minimal(),
  ...
)

fviz_pca_contrib(

```

```

X,
choice = c("var", "ind"),
axes = 1,
fill = "steelblue",
color = "steelblue",
sortcontrib = c("desc", "asc", "none"),
top = Inf,
...
)

```

### Arguments

<code>X</code>	an object of class PCA, CA, MCA, FAMD, MFA and HMFA [FactoMineR]; <code>precomp</code> and <code>princomp</code> [stats]; <code>dudi</code> , <code>pca</code> , <code>coa</code> and <code>acm</code> [ade4]; <code>ca</code> [ca package].
<code>choice</code>	allowed values are "row" and "col" for CA; "var" and "ind" for PCA or MCA; "var", "ind", "quanti.var", "quali.var" and "group" for FAMD, MFA and HMFA.
<code>axes</code>	a numeric vector specifying the dimension(s) of interest.
<code>fill</code>	a fill color for the bar plot.
<code>color</code>	an outline color for the bar plot.
<code>sort.val</code>	a string specifying whether the value should be sorted. Allowed values are "none" (no sorting), "asc" (for ascending) or "desc" (for descending).
<code>top</code>	a numeric value specifying the number of top elements to be shown.
<code>xtickslab.rt</code>	Same as <code>x.text.angle</code> and <code>y.text.angle</code> , respectively. Will be deprecated in the near future.
<code>ggtheme</code>	function, ggplot2 theme name. Default value is <code>theme_pubr()</code> . Allowed values include ggplot2 official themes: <code>theme_gray()</code> , <code>theme_bw()</code> , <code>theme_minimal()</code> , <code>theme_classic()</code> , <code>theme_void()</code> , ....
<code>...</code>	other arguments to be passed to the function <code>ggpar</code> .
<code>sortcontrib</code>	see the argument <code>sort.val</code>

### Details

The function `fviz_contrib()` creates a barplot of row/column contributions. A reference dashed line is also shown on the barplot. This reference line corresponds to the expected value if the contribution were uniform.

For a given dimension, any row/column with a contribution above the reference line could be considered as important in contributing to the dimension.

### Value

a ggplot2 plot

### Functions

- `fviz_pca_contrib`: deprecated function. Use `fviz_contrib()`

**Author(s)**

Alboukadel Kassambara <alboukadel.kassambara@gmail.com>

**References**

<http://www.sthda.com/english/>

**Examples**

```
# Principal component analysis
# ++++++
data(decathlon2)
decathlon2.active <- decathlon2[1:23, 1:10]
res.pca <- prcomp(decathlon2.active, scale = TRUE)

# variable contributions on axis 1
fviz_contrib(res.pca, choice="var", axes = 1, top = 10 )

# Change theme and color
fviz_contrib(res.pca, choice="var", axes = 1,
             fill = "lightgray", color = "black") +
  theme_minimal() +
  theme(axis.text.x = element_text(angle=45))

# Variable contributions on axis 2
fviz_contrib(res.pca, choice="var", axes = 2)
# Variable contributions on axes 1 + 2
fviz_contrib(res.pca, choice="var", axes = 1:2)

# Contributions of individuals on axis 1
fviz_contrib(res.pca, choice="ind", axes = 1)

## Not run:
# Correspondence Analysis
# ++++++
# Install and load FactoMineR to compute CA
# install.packages("FactoMineR")
library("FactoMineR")
data("housetasks")
res.ca <- CA(housetasks, graph = FALSE)

# Visualize row contributions on axes 1
fviz_contrib(res.ca, choice = "row", axes = 1)
# Visualize column contributions on axes 1
fviz_contrib(res.ca, choice = "col", axes = 1)

# Multiple Correspondence Analysis
# ++++++
library(FactoMineR)
data(poison)
res.mca <- MCA(poison, quanti.sup = 1:2,
              quali.sup = 3:4, graph=FALSE)
```

```

# Visualize individual contributions on axes 1
fviz_contrib(res.mca, choice = "ind", axes = 1)
# Visualize variable categorie contributions on axes 1
fviz_contrib(res.mca, choice = "var", axes = 1)

# Multiple Factor Analysis
# ++++++
library(FactoMineR)
data(poison)
res.mfa <- MFA(poison, group=c(2,2,5,6), type=c("s","n","n","n"),
              name.group=c("desc","desc2","symptom","eat"),
              num.group.sup=1:2, graph=FALSE)

# Visualize individual contributions on axes 1
fviz_contrib(res.mfa, choice = "ind", axes = 1, top = 20)
# Visualize catecorical variable categorie contributions on axes 1
fviz_contrib(res.mfa, choice = "quali.var", axes = 1)

## End(Not run)

```

---

fviz\_cos2

---

*Visualize the quality of representation of rows/columns*


---

## Description

This function can be used to visualize the quality of representation (cos2) of rows/columns from the results of Principal Component Analysis (PCA), Correspondence Analysis (CA), Multiple Correspondence Analysis (MCA), Factor Analysis of Mixed Data (FAMD), Multiple Factor Analysis (MFA) and Hierarchical Multiple Factor Analysis (HMFA) functions.

## Usage

```

fviz_cos2(
  X,
  choice = c("row", "col", "var", "ind", "quanti.var", "quali.var", "group"),
  axes = 1,
  fill = "steelblue",
  color = "steelblue",
  sort.val = c("desc", "asc", "none"),
  top = Inf,
  xtckslab.rt = 45,
  ggtheme = theme_minimal(),
  ...
)

```

**Arguments**

<code>X</code>	an object of class PCA, CA, MCA, FAMD, MFA and HMFA [FactoMineR]; prcomp and princomp [stats]; dudi, pca, coa and acm [ade4]; ca [ca package].
<code>choice</code>	allowed values are "row" and "col" for CA; "var" and "ind" for PCA or MCA; "var", "ind", "quanti.var", "quali.var" and "group" for FAMD, MFA and HMFA.
<code>axes</code>	a numeric vector specifying the dimension(s) of interest.
<code>fill</code>	a fill color for the bar plot.
<code>color</code>	an outline color for the bar plot.
<code>sort.val</code>	a string specifying whether the value should be sorted. Allowed values are "none" (no sorting), "asc" (for ascending) or "desc" (for descending).
<code>top</code>	a numeric value specifying the number of top elements to be shown.
<code>xtickslab.rt</code>	Same as <code>x.text.angle</code> and <code>y.text.angle</code> , respectively. Will be deprecated in the near future.
<code>ggtheme</code>	function, ggplot2 theme name. Default value is <code>theme_pubr()</code> . Allowed values include ggplot2 official themes: <code>theme_gray()</code> , <code>theme_bw()</code> , <code>theme_minimal()</code> , <code>theme_classic()</code> , <code>theme_void()</code> , ....
<code>...</code>	not used

**Value**

a ggplot

**Author(s)**

Alboukadel Kassambara <alboukadel.kassambara@gmail.com>

**References**

<http://www.sthda.com/english/>

**Examples**

```
# Principal component analysis
# ++++++
data(decathlon2)
decathlon2.active <- decathlon2[1:23, 1:10]
res.pca <- prcomp(decathlon2.active, scale = TRUE)

# variable cos2 on axis 1
fviz_cos2(res.pca, choice="var", axes = 1, top = 10 )

# Change color
fviz_cos2(res.pca, choice="var", axes = 1,
          fill = "lightgray", color = "black")

# Variable cos2 on axes 1 + 2
fviz_cos2(res.pca, choice="var", axes = 1:2)
```

```

# cos2 of individuals on axis 1
fviz_cos2(res.pca, choice="ind", axes = 1)

## Not run:
# Correspondence Analysis
# ++++++
library("FactoMineR")
data("housetasks")
res.ca <- CA(housetasks, graph = FALSE)

# Visualize row cos2 on axes 1
fviz_cos2(res.ca, choice = "row", axes = 1)
# Visualize column cos2 on axes 1
fviz_cos2(res.ca, choice = "col", axes = 1)

# Multiple Correspondence Analysis
# ++++++
library(FactoMineR)
data(poison)
res.mca <- MCA(poison, quanti.sup = 1:2,
               quali.sup = 3:4, graph=FALSE)

# Visualize individual cos2 on axes 1
fviz_cos2(res.mca, choice = "ind", axes = 1, top = 20)
# Visualize variable categorie cos2 on axes 1
fviz_cos2(res.mca, choice = "var", axes = 1)

# Multiple Factor Analysis
# ++++++
library(FactoMineR)
data(poison)
res.mfa <- MFA(poison, group=c(2,2,5,6), type=c("s","n","n","n"),
              name.group=c("desc","desc2","symptom","eat"),
              num.group.sup=1:2, graph=FALSE)
# Visualize individual cos2 on axes 1
# Select the top 20
fviz_cos2(res.mfa, choice = "ind", axes = 1, top = 20)
# Visualize catecorical variable categorie cos2 on axes 1
fviz_cos2(res.mfa, choice = "quali.var", axes = 1)

## End(Not run)

```



## Description

Draws easily beautiful dendrograms using either R base plot or ggplot2. Provides also an option for drawing a circular dendrogram and phylogenetic trees.

## Usage

```
fviz_dend(
  x,
  k = NULL,
  h = NULL,
  k_colors = NULL,
  palette = NULL,
  show_labels = TRUE,
  color_labels_by_k = TRUE,
  label_cols = NULL,
  labels_track_height = NULL,
  repel = FALSE,
  lwd = 0.7,
  type = c("rectangle", "circular", "phylogenetic"),
  phylo_layout = "layout.auto",
  rect = FALSE,
  rect_border = "gray",
  rect_lty = 2,
  rect_fill = FALSE,
  lower_rect,
  horiz = FALSE,
  cex = 0.8,
  main = "Cluster Dendrogram",
  xlab = "",
  ylab = "Height",
  sub = NULL,
  ggtheme = theme_classic(),
  ...
)
```

## Arguments

x	an object of class dendrogram, hclust, agnes, diana, hcut, hkmeans or HCPC (FactoMineR).
k	the number of groups for cutting the tree.
h	a numeric value. Cut the dendrogram by cutting at height h. (k overrides h)
k_colors, palette	a vector containing colors to be used for the groups. It should contains k number of colors. Allowed values include also "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; and scientific journal palettes from ggsci R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty".

<code>show_labels</code>	a logical value. If TRUE, leaf labels are shown. Default value is TRUE.
<code>color_labels_by_k</code>	logical value. If TRUE, labels are colored automatically by group when <code>k != NULL</code> .
<code>label_cols</code>	a vector containing the colors for labels.
<code>labels_track_height</code>	a positive numeric value for adjusting the room for the labels. Used only when <code>type = "rectangle"</code> .
<code>repel</code>	logical value. Use <code>repel = TRUE</code> to avoid label overplotting when <code>type = "phylogenetic"</code> .
<code>lwd</code>	a numeric value specifying branches and rectangle line width.
<code>type</code>	type of plot. Allowed values are one of "rectangle", "triangle", "circular", "phylogenetic".
<code>phylo_layout</code>	the layout to be used for phylogenetic trees. Default value is "layout.auto". Allowed values include: <code>layout.auto</code> , <code>layout_with_drl</code> , <code>layout_as_tree</code> , <code>layout.gem</code> , <code>layout.mds</code> and <code>layout_with_lgl</code> .
<code>rect</code>	logical value specifying whether to add a rectangle around groups. Used only when <code>k != NULL</code> .
<code>rect_border, rect_lty</code>	border color and line type for rectangles.
<code>rect_fill</code>	a logical value. If TRUE, fill the rectangle.
<code>lower_rect</code>	a value of how low should the lower part of the rectangle around clusters. Ignored when <code>rect = FALSE</code> .
<code>horiz</code>	a logical value. If TRUE, an horizontal dendrogram is drawn.
<code>cex</code>	size of labels
<code>main, xlab, ylab</code>	main and axis titles
<code>sub</code>	Plot subtitle. If NULL, the method used hierarchical clustering is shown. To remove the subtitle use <code>sub = ""</code> .
<code>ggtheme</code>	function, ggplot2 theme name. Default value is <code>theme_classic()</code> . Allowed values include ggplot2 official themes: <code>theme_gray()</code> , <code>theme_bw()</code> , <code>theme_minimal()</code> , <code>theme_classic()</code> , <code>theme_void()</code> , ....
<code>...</code>	other arguments to be passed to the function <code>plot.dendrogram()</code>

### Value

an object of class `fviz_dend` which is a ggplot with the attributes "dendrogram" accessible using `attr(x, "dendrogram")`, where `x` is the result of `fviz_dend()`.

### Examples

```
# Load and scale the data
data(USArrests)
df <- scale(USArrests)

# Hierarchical clustering
```

```

res.hc <- hclust(dist(df))

# Default plot
fviz_dend(res.hc)

# Cut the tree
fviz_dend(res.hc, cex = 0.5, k = 4, color_labels_by_k = TRUE)

# Don't color labels, add rectangles
fviz_dend(res.hc, cex = 0.5, k = 4,
  color_labels_by_k = FALSE, rect = TRUE)

# Change the color of tree using black color for all groups
# Change rectangle border colors
fviz_dend(res.hc, rect = TRUE, k_colors = "black",
  rect_border = 2:5, rect_lty = 1)

# Customized color for groups
fviz_dend(res.hc, k = 4,
  k_colors = c("#1B9E77", "#D95F02", "#7570B3", "#E7298A"))

# Color labels using k-means clusters
km.clust <- kmeans(df, 4)$cluster
fviz_dend(res.hc, k = 4,
  k_colors = c("blue", "green3", "red", "black"),
  label_cols = km.clust[res.hc$order], cex = 0.6)

```

---

fviz\_ellipses

*Draw confidence ellipses around the categories*


---

## Description

Draw confidence ellipses around the categories

## Usage

```

fviz_ellipses(
  X,
  habillage,
  axes = c(1, 2),
  addEllipses = TRUE,
  ellipse.type = "confidence",
  palette = NULL,
  pointsize = 1,
  geom = c("point", "text"),
  ggtheme = theme_bw(),
  ...
)

```

**Arguments**

<code>X</code>	an object of class MCA, PCA or MFA.
<code>habillage</code>	a numeric vector of indexes of variables or a character vector of names of variables. Can be also a data frame containing grouping variables.
<code>axes</code>	a numeric vector specifying the axes of interest. Default values are 1:2 for axes 1 and 2.
<code>addEllipses</code>	logical value. If TRUE, draws ellipses around the individuals when <code>habillage != "none"</code> .
<code>ellipse.type</code>	Character specifying frame type. Possible values are "convex", "confidence" or types supported by <code>stat_ellipse()</code> including one of <code>c("t", "norm", "euclid")</code> for plotting concentration ellipses. <ul style="list-style-type: none"> <li>• "convex": plot convex hull of a set o points.</li> <li>• "confidence": plot confidence ellipses around group mean points as <code>coord.ellipse()</code> [in FactoMineR].</li> <li>• "t": assumes a multivariate t-distribution.</li> <li>• "norm": assumes a multivariate normal distribution.</li> <li>• "euclid": draws a circle with the radius equal to level, representing the euclidean distance from the center. This ellipse probably won't appear circular unless <code>coord_fixed()</code> is applied.</li> </ul>
<code>palette</code>	the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. <code>c("blue", "red")</code> ; and scientific journal palettes from ggsci R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmarty". Can be also a numeric vector of length(groups); in this case a basic color palette is created using the function <code>palette</code> .
<code>pointsize</code>	the size of points
<code>geom</code>	a text specifying the geometry to be used for the graph. Allowed values are the combination of <code>c("point", "text")</code> . Use "point" (to show only points); "text" to show only labels; <code>c("point", "text")</code> to show both types.
<code>ggtheme</code>	function, ggplot2 theme name. Default value is <code>theme_pubr()</code> . Allowed values include ggplot2 official themes: <code>theme_gray()</code> , <code>theme_bw()</code> , <code>theme_minimal()</code> , <code>theme_classic()</code> , <code>theme_void()</code> , ....
<code>...</code>	Arguments to be passed to the functions <code>ggpubr::ggscatter()</code> & <code>ggpubr::ggpar()</code> .

**Value**

a ggplot

**Author(s)**

Alboukadel Kassambara <alboukadel.kassambara@gmail.com>

## Examples

```
# Multiple Correspondence Analysis
# ++++++
library(FactoMineR)
data(poison)
res.mca <- MCA(poison, quanti.sup = 1:2,
               quali.sup = 3:4, graph=FALSE)

fviz_ellipses(res.mca, 1:4, geom = "point",
              palette = "jco")
```

---

fviz\_famd

*Visualize Factor Analysis of Mixed Data*


---

## Description

Factor analysis of mixed data (FAMD) is, a particular case of MFA, used to analyze a data set containing both quantitative and qualitative variables. `fviz_famd()` provides ggplot2-based elegant visualization of FAMD outputs from the R function: FAMD [FactoMineR].

- `fviz_famd_ind()`: Graph of individuals
- `fviz_famd_var()`: Graph of variables
- `fviz_famd()`: An alias of `fviz_famd_ind(res.famd)`

## Usage

```
fviz_famd_ind(
  X,
  axes = c(1, 2),
  geom = c("point", "text"),
  repel = FALSE,
  habillage = "none",
  palette = NULL,
  addEllipses = FALSE,
  col.ind = "blue",
  col.ind.sup = "darkblue",
  alpha.ind = 1,
  shape.ind = 19,
  col.quali.var = "black",
  select.ind = list(name = NULL, cos2 = NULL, contrib = NULL),
  gradient.cols = NULL,
  ...
```

```

)

fviz_famd_var(
  X,
  choice = c("var", "quanti.var", "quali.var"),
  axes = c(1, 2),
  geom = c("point", "text"),
  repel = FALSE,
  col.var = "red",
  alpha.var = 1,
  shape.var = 17,
  col.var.sup = "darkgreen",
  select.var = list(name = NULL, cos2 = NULL, contrib = NULL),
  ...
)

fviz_famd(X, ...)

```

### Arguments

<code>X</code>	an object of class FAMD [FactoMineR].
<code>axes</code>	a numeric vector of length 2 specifying the dimensions to be plotted.
<code>geom</code>	a text specifying the geometry to be used for the graph. Allowed values are the combination of <code>c("point", "arrow", "text")</code> . Use "point" (to show only points); "text" to show only labels; <code>c("point", "text")</code> or <code>c("arrow", "text")</code> to show arrows and texts. Using <code>c("arrow", "text")</code> is sensible only for the graph of variables.
<code>repel</code>	a boolean, whether to use <code>ggrepel</code> to avoid overplotting text labels or not.
<code>habillage</code>	an optional factor variable for coloring the observations by groups. Default value is "none". If X is an MFA object from FactoMineR package, <code>habillage</code> can also specify the index of the factor variable in the data.
<code>palette</code>	the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. <code>c("blue", "red")</code> ; and scientific journal palettes from ggsci R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty". Can be also a numeric vector of length(groups); in this case a basic color palette is created using the function <a href="#">palette</a> .
<code>addEllipses</code>	logical value. If TRUE, draws ellipses around the individuals when <code>habillage != "none"</code> .
<code>col.ind, col.var</code>	color for individuals and variables, respectively. Can be a continuous variable or a factor variable. Possible values include also : "cos2", "contrib", "coord", "x" or "y". In this case, the colors for individuals/variables are automatically controlled by their qualities ("cos2"), contributions ("contrib"), coordinates ( $x^2 + y^2$ , "coord"), x values("x") or y values("y"). To use automatic coloring (by <code>cos2</code> , <code>contrib</code> , ...), make sure that <code>habillage = "none"</code> .
<code>col.ind.sup</code>	color for supplementary individuals

<code>alpha.ind</code> , <code>alpha.var</code>	controls the transparency of individuals and variables, respectively. The value can vary from 0 (total transparency) to 1 (no transparency). Default value is 1. Possible values include also : "cos2", "contrib", "coord", "x" or "y". In this case, the transparency for individual/variable colors are automatically controlled by their qualities ("cos2"), contributions ("contrib"), coordinates ( $x^2 + y^2$ , "coord"), x values("x") or y values("y"). To use this, make sure that <code>habillage = "none"</code> .
<code>shape.ind</code> , <code>shape.var</code>	point shapes of individuals, variables, groups and axes
<code>col.quali.var</code>	color for qualitative variables in <code>fviz_mfa_ind()</code> . Default is "black".
<code>select.ind</code> , <code>select.var</code>	a selection of individuals and variables to be drawn. Allowed values are NULL or a list containing the arguments name, <code>cos2</code> or <code>contrib</code> : <ul style="list-style-type: none"> <li>• <code>name</code> is a character vector containing individuals/variables to be drawn</li> <li>• <code>cos2</code> if <code>cos2</code> is in <math>[0, 1]</math>, ex: 0.6, then individuals/variables with a <code>cos2 &gt; 0.6</code> are drawn. if <code>cos2 &gt; 1</code>, ex: 5, then the top 5 individuals/variables with the highest <code>cos2</code> are drawn.</li> <li>• <code>contrib</code> if <code>contrib &gt; 1</code>, ex: 5, then the top 5 individuals/variables with the highest <code>cos2</code> are drawn</li> </ul>
<code>gradient.cols</code>	vector of colors to use for n-colour gradient. Allowed values include brewer and ggsci color palettes.
...	Arguments to be passed to the function <code>fviz()</code>
<code>choice</code>	The graph to plot in <code>fviz_mfa_var()</code> . Allowed values include one of <code>c("var", "quanti.var", "quali.var")</code> .
<code>col.var.sup</code>	color for supplementary variables.

**Value**

a ggplot

**Author(s)**

Alboukadel Kassambara <alboukadel.kassambara@gmail.com>

**Examples**

```
# Compute FAMD
library("FactoMineR")
data(wine)
res.famd <- FAMD(wine[,c(1,2, 16, 22, 29, 28, 30,31)], graph = FALSE)

# Eigenvalues/variances of dimensions
fviz_screplot(res.famd)
# Graph of variables
fviz_famd_var(res.famd)
# Quantitative variables
fviz_famd_var(res.famd, "quanti.var", repel = TRUE, col.var = "black")
```

```
# Qualitative variables
fviz_famd_var(res.famd, "quali.var", col.var = "black")
# Graph of individuals colored by cos2
fviz_famd_ind(res.famd, col.ind = "cos2",
  gradient.cols = c("#00AFBB", "#E7B800", "#FC4E07"),
  repel = TRUE)
```

---

fviz\_hmfa

*Visualize Hierarchical Multiple Factor Analysis*

---

## Description

Hierarchical Multiple Factor Analysis (HMFA) is, an extension of MFA, used in a situation where the data are organized into a hierarchical structure. `fviz_hmfa()` provides `ggplot2`-based elegant visualization of HMFA outputs from the R function: `HMFA` [FactoMineR].

- `fviz_hmfa_ind()`: Graph of individuals
- `fviz_hmfa_var()`: Graph of variables
- `fviz_hmfa_quali_biplot()`: Biplot of individuals and qualitative variables
- `fviz_hmfa()`: An alias of `fviz_hmfa_ind()`

## Usage

```
fviz_hmfa_ind(
  X,
  axes = c(1, 2),
  geom = c("point", "text"),
  repel = FALSE,
  habillage = "none",
  addEllipses = FALSE,
  shape.ind = 19,
  col.ind = "blue",
  col.ind.sup = "darkblue",
  alpha.ind = 1,
  select.ind = list(name = NULL, cos2 = NULL, contrib = NULL),
  partial = NULL,
  col.partial = "group",
  group.names = NULL,
  node.level = 1,
  ...
)

fviz_hmfa_var(
```



```

X,
choice = c("quanti.var", "quali.var", "group"),
axes = c(1, 2),
geom = c("point", "text"),
repel = FALSE,
col.var = "red",
alpha.var = 1,
shape.var = 17,
col.var.sup = "darkgreen",
select.var = list(name = NULL, cos2 = NULL, contrib = NULL),
...
)

fviz_hmfa_quali_biplot(
  X,
  axes = c(1, 2),
  geom = c("point", "text"),
  repel = FALSE,
  habillage = "none",
  title = "Biplot of individuals and qualitative variables - HMFA",
  ...
)

fviz_hmfa(X, ...)

```

### Arguments

<code>X</code>	an object of class HMFA [FactoMineR].
<code>axes</code>	a numeric vector of length 2 specifying the dimensions to be plotted.
<code>geom</code>	a text specifying the geometry to be used for the graph. Allowed values are the combination of <code>c("point", "arrow", "text")</code> . Use "point" (to show only points); "text" to show only labels; <code>c("point", "text")</code> or <code>c("arrow", "text")</code> to show arrows and texts. Using <code>c("arrow", "text")</code> is sensible only for the graph of variables.
<code>repel</code>	a boolean, whether to use <code>ggrepel</code> to avoid overplotting text labels or not.
<code>habillage</code>	an optional factor variable for coloring the observations by groups. Default value is "none". If X is an HMFA object from FactoMineR package, <code>habillage</code> can also specify the index of the factor variable in the data.
<code>addEllipses</code>	logical value. If TRUE, draws ellipses around the individuals when <code>habillage != "none"</code> .
<code>shape.ind, shape.var</code>	point shapes of individuals and variables, respectively.
<code>col.ind, col.var</code>	color for individuals, partial individuals and variables, respectively. Can be a continuous variable or a factor variable. Possible values include also : "cos2", "contrib", "coord", "x" or "y". In this case, the colors for individuals/variables

are automatically controlled by their qualities ("cos2"), contributions ("contrib"), coordinates ( $x^2 + y^2$ , "coord"), x values("x") or y values("y"). To use automatic coloring (by cos2, contrib, ....), make sure that habillage="none".

col.ind.sup	color for supplementary individuals
alpha.ind, alpha.var	controls the transparency of individual, partial individual and variable, respectively. The value can vary from 0 (total transparency) to 1 (no transparency). Default value is 1. Possible values include also : "cos2", "contrib", "coord", "x" or "y". In this case, the transparency for individual/variable colors are automatically controlled by their qualities ("cos2"), contributions ("contrib"), coordinates ( $x^2 + y^2$ , "coord"), x values("x") or y values("y"). To use this, make sure that habillage="none".
select.ind, select.var	a selection of individuals and variables to be drawn. Allowed values are NULL or a list containing the arguments name, cos2 or contrib: <ul style="list-style-type: none"> <li>• name is a character vector containing individuals/variables to be drawn</li> <li>• cos2 if cos2 is in [0, 1], ex: 0.6, then individuals/variables with a cos2 &gt; 0.6 are drawn. if cos2 &gt; 1, ex: 5, then the top 5 individuals/variables with the highest cos2 are drawn.</li> <li>• contrib if contrib &gt; 1, ex: 5, then the top 5 individuals/variables with the highest cos2 are drawn</li> </ul>
partial	list of the individuals for which the partial points should be drawn. (by default, partial = NULL and no partial points are drawn). Use partial = "All" to visualize partial points for all individuals.
col.partial	color for partial individuals. By default, points are colored according to the groups.
group.names	a vector containing the name of the groups (by default, NULL and the group are named group.1, group.2 and so on).
node.level	a single number indicating the HMFA node level to plot.
...	Arguments to be passed to the function fviz() and ggpubr::ggpar()
choice	the graph to plot. Allowed values include one of c("quanti.var", "quali.var", "group") for plotting quantitative variables, qualitative variables and group of variables, respectively.
col.var.sup	color for supplementary variables.
title	the title of the graph

### Value

a ggplot

### Author(s)

Fabian Mundt <f.mundt@inventionate.de>

Alboukadel Kassambara <alboukadel.kassambara@gmail.com>

## References

<http://www.sthda.com/english/>

## Examples

```
# Hierarchical Multiple Factor Analysis
# ++++++
# Install and load FactoMineR to compute MFA
# install.packages("FactoMineR")
library("FactoMineR")
data(wine)
hierar <- list(c(2,5,3,10,9,2), c(4,2))
res.hmfa <- HMFA(wine, H = hierar, type=c("n",rep("s",5)), graph = FALSE)

# Graph of individuals
# ++++++
# Color of individuals: col.ind = "#2E9FDF"
# Use repel = TRUE to avoid overplotting (slow if many points)
fviz_hmfa_ind(res.hmfa, repel = TRUE, col.ind = "#2E9FDF")

# Color individuals by groups, add concentration ellipses
# Remove labels: label = "none".
# Change color palette to "jco". See ?ggpubr::ggpar
grp <- as.factor(wine[,1])
p <- fviz_hmfa_ind(res.hmfa, label="none", habillage=grp,
  addEllipses=TRUE, palette = "jco")
print(p)

# Graph of variables
# ++++++
# Quantitative variables
fviz_hmfa_var(res.hmfa, "quanti.var")
# Graph of categorical variable categories
fviz_hmfa_var(res.hmfa, "quali.var")
# Groups of variables (correlation square)
fviz_hmfa_var(res.hmfa, "group")

# Biplot of categorical variable categories and individuals
# ++++++
fviz_hmfa_quali_biplot(res.hmfa)

# Graph of partial individuals (starplot)
# ++++++
fviz_hmfa_ind(res.hmfa, partial = "all", palette = "Dark2")
```

## Description

Multiple Correspondence Analysis (MCA) is an extension of simple CA to analyse a data table containing more than two categorical variables. `fviz_mca()` provides ggplot2-based elegant visualization of MCA outputs from the R functions: `MCA` [in `FactoMineR`], `acm` [in `ade4`], and `expOutput/epMCA` [in `ExPosition`]. Read more: [Multiple Correspondence Analysis Essentials](#).

- `fviz_mca_ind()`: Graph of individuals
- `fviz_mca_var()`: Graph of variables
- `fviz_mca_biplot()`: Biplot of individuals and variables
- `fviz_mca()`: An alias of `fviz_mca_biplot()`

## Usage

```
fviz_mca_ind(  
  X,  
  axes = c(1, 2),  
  geom = c("point", "text"),  
  geom.ind = geom,  
  repel = FALSE,  
  habillage = "none",  
  palette = NULL,  
  addEllipses = FALSE,  
  col.ind = "blue",  
  col.ind.sup = "darkblue",  
  alpha.ind = 1,  
  shape.ind = 19,  
  map = "symmetric",  
  select.ind = list(name = NULL, cos2 = NULL, contrib = NULL),  
  ...  
)
```

```
fviz_mca_var(  
  X,  
  choice = c("var.cat", "mca.cor", "var", "quanti.sup"),  
  axes = c(1, 2),  
  geom = c("point", "text"),  
  geom.var = geom,  
  repel = FALSE,  
  col.var = "red",  
  alpha.var = 1,  
  shape.var = 17,  
  col.quanti.sup = "blue",
```

```

    col.quali.sup = "darkgreen",
    map = "symmetric",
    select.var = list(name = NULL, cos2 = NULL, contrib = NULL),
    ...
  )

fviz_mca_biplot(
  X,
  axes = c(1, 2),
  geom = c("point", "text"),
  geom.ind = geom,
  geom.var = geom,
  repel = FALSE,
  label = "all",
  invisible = "none",
  habillage = "none",
  addEllipses = FALSE,
  palette = NULL,
  arrows = c(FALSE, FALSE),
  map = "symmetric",
  title = "MCA - Biplot",
  ...
)

fviz_mca(X, ...)

```

### Arguments

<code>X</code>	an object of class MCA [FactoMineR], acm [ade4] and expOutput/epMCA [Ex-Position].
<code>axes</code>	a numeric vector of length 2 specifying the dimensions to be plotted.
<code>geom</code>	a text specifying the geometry to be used for the graph. Allowed values are the combination of <code>c("point", "arrow", "text")</code> . Use "point" (to show only points); "text" to show only labels; <code>c("point", "text")</code> or <code>c("arrow", "text")</code> to show arrows and texts. Using <code>c("arrow", "text")</code> is sensible only for the graph of variables.
<code>geom.ind, geom.var</code>	as <code>geom</code> but for individuals and variables, respectively. Default is <code>geom.ind = c("point", "text"), geom.var = c("point", "text")</code> .
<code>repel</code>	a boolean, whether to use <code>ggrepel</code> to avoid overplotting text labels or not.
<code>habillage</code>	an optional factor variable for coloring the observations by groups. Default value is "none". If <code>X</code> is an MCA object from FactoMineR package, <code>habillage</code> can also specify the index of the factor variable in the data.
<code>palette</code>	the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. <code>c("blue", "red")</code> ; and scientific journal palettes from ggsci R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago",

	"simpsons" and "rickandmorty". Can be also a numeric vector of length(groups); in this case a basic color palette is created using the function <a href="#">palette</a> .
addEllipses	logical value. If TRUE, draws ellipses around the individuals when habillage != "none".
col.ind, col.var	color for individuals and variables, respectively. Can be a continuous variable or a factor variable. Possible values include also : "cos2", "contrib", "coord", "x" or "y". In this case, the colors for individuals/variables are automatically controlled by their qualities ("cos2"), contributions ("contrib"), coordinates ( $x^2 + y^2$ , "coord"), x values("x") or y values("y"). To use automatic coloring (by cos2, contrib, ...), make sure that habillage ="none".
col.ind.sup	color for supplementary individuals
alpha.ind, alpha.var	controls the transparency of individual and variable colors, respectively. The value can variate from 0 (total transparency) to 1 (no transparency). Default value is 1. Possible values include also : "cos2", "contrib", "coord", "x" or "y". In this case, the transparency for individual/variable colors are automatically controlled by their qualities ("cos2"), contributions ("contrib"), coordinates ( $x^2 + y^2$ , "coord"), x values("x") or y values("y"). To use this, make sure that habillage ="none".
shape.ind, shape.var	point shapes of individuals and variables.
map	character string specifying the map type. Allowed options include: "symmetric", "rowprincipal", "colprincipal", "symbiplot", "rowgab", "colgab", "rowgreen" and "colgreen". See details
select.ind, select.var	a selection of individuals/variables to be drawn. Allowed values are NULL or a list containing the arguments name, cos2 or contrib: <ul style="list-style-type: none"> <li>• name is a character vector containing individuals/variables to be drawn</li> <li>• cos2 if cos2 is in [0, 1], ex: 0.6, then individuals/variables with a cos2 &gt; 0.6 are drawn. if cos2 &gt; 1, ex: 5, then the top 5 individuals/variables with the highest cos2 are drawn.</li> <li>• contrib if contrib &gt; 1, ex: 5, then the top 5 individuals/variables with the highest contrib are drawn</li> </ul>
...	Additional arguments. <ul style="list-style-type: none"> <li>• in fviz_mca_ind(), fviz_mca_var() and fviz_mca_cor(): Additional arguments are passed to the functions fviz() and ggpubr::ggpar().</li> <li>• in fviz_mca_biplot() and fviz_mca(): Additional arguments are passed to fviz_mca_ind() and fviz_mca_var().</li> </ul>
choice	the graph to plot. Allowed values include: i) "var" and "mca.cor" for plotting the correlation between variables and principal dimensions; ii) "var.cat" for variable categories and iii) "quanti.sup" for the supplementary quantitative variables.
col.quanti.sup, col.quali.sup	a color for the quantitative/qualitative supplementary variables.

label	a text specifying the elements to be labelled. Default value is "all". Allowed values are "none" or the combination of c("ind", "ind.sup", "var", "quali.sup", "quanti.sup"). "ind" can be used to label only active individuals. "ind.sup" is for supplementary individuals. "var" is for active variable categories. "quali.sup" is for supplementary qualitative variable categories. "quanti.sup" is for quantitative supplementary variables.
invisible	a text specifying the elements to be hidden on the plot. Default value is "none". Allowed values are the combination of c("ind", "ind.sup", "var", "quali.sup", "quanti.sup").
arrows	Vector of two logicals specifying if the plot should contain points (FALSE, default) or arrows (TRUE). First value sets the rows and the second value sets the columns.
title	the title of the graph

### Details

The default plot of MCA is a "symmetric" plot in which both rows and columns are in principal coordinates. In this situation, it's not possible to interpret the distance between row points and column points. To overcome this problem, the simplest way is to make an asymmetric plot. The argument "map" can be used to change the plot type. For more explanation, read the details section of `fviz_ca` documentation.

### Value

a ggplot

### Author(s)

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### See Also

[get\\_mca](#), [fviz\\_pca](#), [fviz\\_ca](#), [fviz\\_mfa](#), [fviz\\_hmfa](#)

### Examples

```
# Multiple Correspondence Analysis
# ++++++
# Install and load FactoMineR to compute MCA
# install.packages("FactoMineR")
library("FactoMineR")
data(poison)
poison.active <- poison[1:55, 5:15]
head(poison.active)
res.mca <- MCA(poison.active, graph=FALSE)

# Graph of individuals
# ++++++

# Default Plot
```

```

# Color of individuals: col.ind = "steelblue"
fviz_mca_ind(res.mca, col.ind = "steelblue")

# 1. Control automatically the color of individuals
# using the "cos2" or the contributions "contrib"
# cos2 = the quality of the individuals on the factor map
# 2. To keep only point or text use geom = "point" or geom = "text".
# 3. Change themes: http://www.sthda.com/english/wiki/ggplot2-themes

fviz_mca_ind(res.mca, col.ind = "cos2", repel = TRUE)

## Not run:
# You can also control the transparency
# of the color by the cos2
fviz_mca_ind(res.mca, alpha.ind="cos2")

## End(Not run)

# Color individuals by groups, add concentration ellipses
# Remove labels: label = "none".
grp <- as.factor(poison.active[, "Vomiting"])
p <- fviz_mca_ind(res.mca, label="none", habillage=grp,
  addEllipses=TRUE, ellipse.level=0.95)
print(p)

# Change group colors using RColorBrewer color palettes
# Read more: http://www.sthda.com/english/wiki/ggplot2-colors
p + scale_color_brewer(palette="Dark2") +
  scale_fill_brewer(palette="Dark2")

# Change group colors manually
# Read more: http://www.sthda.com/english/wiki/ggplot2-colors
p + scale_color_manual(values=c("#999999", "#E69F00"))+
  scale_fill_manual(values=c("#999999", "#E69F00"))

# Select and visualize some individuals (ind) with select.ind argument.
# - ind with cos2 >= 0.4: select.ind = list(cos2 = 0.4)
# - Top 20 ind according to the cos2: select.ind = list(cos2 = 20)
# - Top 20 contributing individuals: select.ind = list(contrib = 20)
# - Select ind by names: select.ind = list(name = c("44", "38", "53", "39") )

# Example: Select the top 40 according to the cos2
fviz_mca_ind(res.mca, select.ind = list(cos2 = 20))

# Graph of variable categories
# ++++++
# Default plot: use repel = TRUE to avoid overplotting
fviz_mca_var(res.mca, col.var = "#FC4E07")

# Control variable colors using their contributions

```



```

# use repel = TRUE to avoid overplotting
fviz_mca_var(res.mca, col.var = "contrib",
  gradient.cols = c("#00AFBB", "#E7B800", "#FC4E07"))

# Biplot
# ++++++
grp <- as.factor(poison.active[, "Vomiting"])
fviz_mca_biplot(res.mca, repel = TRUE, col.var = "#E7B800",
  habillage = grp, addEllipses = TRUE, ellipse.level = 0.95)

## Not run:
# Keep only the labels for variable categories:
fviz_mca_biplot(res.mca, label = "var")

# Keep only labels for individuals
fviz_mca_biplot(res.mca, label = "ind")

# Hide variable categories
fviz_mca_biplot(res.mca, invisible = "var")

# Hide individuals
fviz_mca_biplot(res.mca, invisible = "ind")

# Control automatically the color of individuals using the cos2
fviz_mca_biplot(res.mca, label = "var", col.ind="cos2")

# Change the color by groups, add ellipses
fviz_mca_biplot(res.mca, label="var", col.var = "blue",
  habillage=grp, addEllipses=TRUE, ellipse.level=0.95)

# Select the top 30 contributing individuals
# And the top 10 variables
fviz_mca_biplot(res.mca,
  select.ind = list(contrib = 30),
  select.var = list(contrib = 10))

## End(Not run)

```

---

fviz\_mclust

*Plot Model-Based Clustering Results using ggplot2*


---

## Description

Plots the classification, the uncertainty and the BIC values returned by the Mclust() function.

## Usage

```
fviz_mclust(
```

```

    object,
    what = c("classification", "uncertainty", "BIC"),
    ellipse.type = "norm",
    ellipse.level = 0.4,
    ggtheme = theme_classic(),
    ...
)

fviz_mclust_bic(
  object,
  model.names = NULL,
  shape = 19,
  color = "model",
  palette = NULL,
  legend = NULL,
  main = "Model selection",
  xlab = "Number of components",
  ylab = "BIC",
  ...
)

```

### Arguments

<code>object</code>	an object of class <code>Mclust</code>
<code>what</code>	choose from one of the following three options: "classification" (default), "uncertainty" and "BIC".
<code>ellipse.type</code>	Character specifying frame type. Possible values are 'convex', 'confidence' or types supported by <a href="#">stat_ellipse</a> including one of c("t", "norm", "euclid").
<code>ellipse.level</code>	the size of the concentration ellipse in normal probability. Passed for <code>ggplot2::stat_ellipse</code> 's level. Ignored in 'convex'. Default value is 0.95.
<code>ggtheme</code>	function, <code>ggplot2</code> theme name. Default value is <code>theme_pubr()</code> . Allowed values include <code>ggplot2</code> official themes: <code>theme_gray()</code> , <code>theme_bw()</code> , <code>theme_minimal()</code> , <code>theme_classic()</code> , <code>theme_void()</code> , ....
<code>...</code>	other arguments to be passed to the functions <a href="#">fviz_cluster</a> and <a href="#">ggpar</a> .
<code>model.names</code>	one or more model names corresponding to models fit in <code>object</code> . The default is to plot the BIC for all of the models fit.
<code>shape</code>	point shape. To change point shape by model names use <code>shape = "model"</code> .
<code>color</code>	point and line color.
<code>palette</code>	the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. c("blue", "red"); and scientific journal palettes from <code>ggsci</code> R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty". Can be also a numeric vector of length(groups); in this case a basic color palette is created using the function <a href="#">palette</a> .

legend	character specifying legend position. Allowed values are one of c("top", "bottom", "left", "right", "none"). To remove the legend use legend = "none". Legend position can be also specified using a numeric vector c(x, y); see details section.
main	plot main title.
xlab	character vector specifying x axis labels. Use xlab = FALSE to hide xlab.
ylab	character vector specifying y axis labels. Use ylab = FALSE to hide ylab.

### Functions

- fviz\_mclust: Plots classification and uncertainty.
- fviz\_mclust\_bic: Plots the BIC values.

### Examples

```
if(require("mclust")){
  # Compute model-based-clustering
  require("mclust")
  data("diabetes")
  mc <- Mclust(diabetes[, -1])

  # Visualize BIC values
  fviz_mclust_bic(mc)

  # Visualize classification
  fviz_mclust(mc, "classification", geom = "point")
}
```

---

 fviz\_mfa

*Visualize Multiple Factor Analysis*


---

### Description

Multiple factor analysis (MFA) is used to analyze a data set in which individuals are described by several sets of variables (quantitative and/or qualitative) structured into groups. `fviz_mfa()` provides ggplot2-based elegant visualization of MFA outputs from the R function: `MFA` [FactoMineR].

- `fviz_mfa_ind()`: Graph of individuals
- `fviz_mfa_var()`: Graph of variables
- `fviz_mfa_axes()`: Graph of partial axes
- `fviz_mfa()`: An alias of `fviz_mfa_ind(res.mfa, partial = "all")`
- `fviz_mfa_quali_biplot()`: Biplot of individuals and qualitative variables

**Usage**

```
fviz_mfa_ind(  
  X,  
  axes = c(1, 2),  
  geom = c("point", "text"),  
  repel = FALSE,  
  habillage = "none",  
  palette = NULL,  
  addEllipses = FALSE,  
  col.ind = "blue",  
  col.ind.sup = "darkblue",  
  alpha.ind = 1,  
  shape.ind = 19,  
  col.quali.var.sup = "black",  
  select.ind = list(name = NULL, cos2 = NULL, contrib = NULL),  
  partial = NULL,  
  col.partial = "group",  
  ...  
)  
  
fviz_mfa_quali_biplot(  
  X,  
  axes = c(1, 2),  
  geom = c("point", "text"),  
  repel = repel,  
  title = "Biplot of individuals and qualitative variables - MFA",  
  ...  
)  
  
fviz_mfa_var(  
  X,  
  choice = c("quanti.var", "group", "quali.var"),  
  axes = c(1, 2),  
  geom = c("point", "text"),  
  repel = FALSE,  
  habillage = "none",  
  col.var = "red",  
  alpha.var = 1,  
  shape.var = 17,  
  col.var.sup = "darkgreen",  
  palette = NULL,  
  select.var = list(name = NULL, cos2 = NULL, contrib = NULL),  
  ...  
)  
  
fviz_mfa_axes(  
  X,  
  axes = c(1, 2),
```

```

    geom = c("arrow", "text"),
    col.axes = NULL,
    alpha.axes = 1,
    col.circle = "grey70",
    select.axes = list(name = NULL, contrib = NULL),
    repel = FALSE,
    ...
)

fviz_mfa(X, partial = "all", ...)

```

## Arguments

<code>X</code>	an object of class MFA [FactoMineR].
<code>axes</code>	a numeric vector of length 2 specifying the dimensions to be plotted.
<code>geom</code>	a text specifying the geometry to be used for the graph. Allowed values are the combination of <code>c("point", "arrow", "text")</code> . Use "point" (to show only points); "text" to show only labels; <code>c("point", "text")</code> or <code>c("arrow", "text")</code> to show arrows and texts. Using <code>c("arrow", "text")</code> is sensible only for the graph of variables.
<code>repel</code>	a boolean, whether to use <code>ggrepel</code> to avoid overplotting text labels or not.
<code>habillage</code>	an optional factor variable for coloring the observations by groups. Default value is "none". If X is an MFA object from FactoMineR package, <code>habillage</code> can also specify the index of the factor variable in the data.
<code>palette</code>	the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. <code>c("blue", "red")</code> ; and scientific journal palettes from ggsci R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty". Can be also a numeric vector of length(groups); in this case a basic color palette is created using the function <a href="#">palette</a> .
<code>addEllipses</code>	logical value. If TRUE, draws ellipses around the individuals when <code>habillage != "none"</code> .
<code>col.ind, col.var, col.axes</code>	color for individuals, variables and <code>col.axes</code> respectively. Can be a continuous variable or a factor variable. Possible values include also : "cos2", "contrib", "coord", "x" or "y". In this case, the colors for individuals/variables are automatically controlled by their qualities ("cos2"), contributions ("contrib"), coordinates ( $x^2 + y^2$ , "coord"), x values("x") or y values("y"). To use automatic coloring (by <code>cos2</code> , <code>contrib</code> , ...), make sure that <code>habillage = "none"</code> .
<code>col.ind.sup</code>	color for supplementary individuals
<code>alpha.ind, alpha.var, alpha.axes</code>	controls the transparency of individual, variable, group and axes colors, respectively. The value can vary from 0 (total transparency) to 1 (no transparency). Default value is 1. Possible values include also : "cos2", "contrib", "coord", "x" or "y". In this case, the transparency for individual/variable colors are automatically controlled by their qualities ("cos2"), contributions ("contrib"), coordinates

	( $x^2 + y^2$ , "coord"), x values("x") or y values("y"). To use this, make sure that <code>habillage = "none"</code> .
<code>shape.ind</code> , <code>shape.var</code>	point shapes of individuals, variables, groups and axes
<code>col.quali.var.sup</code>	color for supplementary qualitative variables. Default is "black".
<code>select.ind</code> , <code>select.var</code> , <code>select.axes</code>	a selection of individuals/partial individuals/ variables/groups/axes to be drawn. Allowed values are NULL or a list containing the arguments name, <code>cos2</code> or <code>contrib</code> : <ul style="list-style-type: none"> <li>• <code>name</code> is a character vector containing individuals/variables to be drawn</li> <li>• <code>cos2</code> if <code>cos2</code> is in <math>[0, 1]</math>, ex: 0.6, then individuals/variables with a <code>cos2</code> &gt; 0.6 are drawn. if <code>cos2</code> &gt; 1, ex: 5, then the top 5 individuals/variables with the highest <code>cos2</code> are drawn.</li> <li>• <code>contrib</code> if <code>contrib</code> &gt; 1, ex: 5, then the top 5 individuals/variables with the highest <code>cos2</code> are drawn</li> </ul>
<code>partial</code>	list of the individuals for which the partial points should be drawn. (by default, <code>partial = NULL</code> and no partial points are drawn). Use <code>partial = "All"</code> to visualize partial points for all individuals.
<code>col.partial</code>	color for partial individuals. By default, points are colored according to the groups.
...	Arguments to be passed to the function <code>fviz()</code>
<code>title</code>	the title of the graph
<code>choice</code>	the graph to plot. Allowed values include one of <code>c("quanti.var", "quali.var", "group")</code> for plotting quantitative variables, qualitative variables and group of variables, respectively.
<code>col.var.sup</code>	color for supplementary variables.
<code>col.circle</code>	a color for the correlation circle. Used only when X is a PCA output.

**Value**

a ggplot2 plot

**Author(s)**

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**References**

<http://www.sthda.com/english/>

**Examples**

```

# Compute Multiple Factor Analysis
library("FactoMineR")
data(wine)
res.mfa <- MFA(wine, group=c(2,5,3,10,9,2), type=c("n",rep("s",5)),
              ncp=5, name.group=c("orig","olf","vis","olfag","gust","ens"),
              num.group.sup=c(1,6), graph=FALSE)

# Eigenvalues/variances of dimensions
fviz_screplot(res.mfa)
# Group of variables
fviz_mfa_var(res.mfa, "group")
# Quantitative variables
fviz_mfa_var(res.mfa, "quanti.var", palette = "jco",
             col.var.sup = "violet", repel = TRUE)
# Graph of individuals colored by cos2
fviz_mfa_ind(res.mfa, col.ind = "cos2",
             gradient.cols = c("#00AFBB", "#E7B800", "#FC4E07"),
             repel = TRUE)
# Partial individuals
fviz_mfa_ind(res.mfa, partial = "all")
# Partial axes
fviz_mfa_axes(res.mfa)

# Graph of categorical variable categories
# ++++++
data(poison)
res.mfa <- MFA(poison, group=c(2,2,5,6), type=c("s","n","n","n"),
              name.group=c("desc","desc2","symptom","eat"),
              num.group.sup=1:2, graph=FALSE)

# Plot of qualitative variables
fviz_mfa_var(res.mfa, "quali.var")

# Biplot of categorical variable categories and individuals
# ++++++
# Use repel = TRUE to avoid overplotting
grp <- as.factor(poison[, "Vomiting"])
fviz_mfa_quali_biplot(res.mfa, repel = FALSE, col.var = "#E7B800",
                    habillage = grp, addEllipses = TRUE, ellipse.level = 0.95)

```

## Description

Partitioning methods, such as k-means clustering require the users to specify the number of clusters to be generated.

- `fviz_nbclust()`: Determines and visualize the optimal number of clusters using different methods: **within cluster sums of squares**, **average silhouette** and **gap statistics**.
- `fviz_gap_stat()`: Visualize the gap statistic generated by the function `clusGap()` [in cluster package]. The optimal number of clusters is specified using the "firstmax" method (`?cluster::clusGap`).

Read more: [Determining the optimal number of clusters](#)

## Usage

```
fviz_nbclust(
  x,
  FUNcluster = NULL,
  method = c("silhouette", "wss", "gap_stat"),
  diss = NULL,
  k.max = 10,
  nboot = 100,
  verbose = interactive(),
  barfill = "steelblue",
  barcolor = "steelblue",
  linecolor = "steelblue",
  print.summary = TRUE,
  ...
)

fviz_gap_stat(
  gap_stat,
  linecolor = "steelblue",
  maxSE = list(method = "firstSEmax", SE.factor = 1)
)
```

## Arguments

- |                         |  |
|-------------------------|--|
| <code>x</code>          | numeric matrix or data frame. In the function <code>fviz_nbclust()</code> , <code>x</code> can be the results of the function <code>NbClust()</code> .   |
| <code>FUNcluster</code> | a partitioning function which accepts as first argument a (data) matrix like <code>x</code> , second argument, say <code>k</code> , $k \geq 2$ , the number of clusters desired, and returns a list with a component named <code>cluster</code> which contains the grouping of observations. Allowed values include: <code>kmeans</code> , <code>cluster::pam</code> , <code>cluster::clara</code> , <code>cluster::fanny</code> , <code>hcut</code> , etc. This argument is not required when <code>x</code> is an output of the function <code>NbClust::NbClust()</code> . |
| <code>method</code>     | the method to be used for estimating the optimal number of clusters. Possible values are "silhouette" (for average silhouette width), "wss" (for total within sum of square) and "gap_stat" (for gap statistics).  |



diss	dist object as produced by dist(), i.e.: <code>diss = dist(x, method = "euclidean")</code> . Used to compute the average silhouette width of clusters, the within sum of square and hierarchical clustering. If NULL, <code>dist(x)</code> is computed with the default method = "euclidean"
k.max	the maximum number of clusters to consider, must be at least two.
nboot	integer, number of Monte Carlo ("bootstrap") samples. Used only for determining the number of clusters using gap statistic.
verbose	logical value. If TRUE, the result of progress is printed.
barfill, barcolor	fill color and outline color for bars
linecolor	color for lines
print.summary	logical value. If true, the optimal number of clusters are printed in <code>fviz_nbclust()</code> .
...	optionally further arguments for <code>FUNcluster()</code>
gap_stat	an object of class "clusGap" returned by the function <code>clusGap()</code> [in cluster package]
maxSE	a list containing the parameters (method and SE.factor) for determining the location of the maximum of the gap statistic (Read the documentation <code>?cluster::maxSE</code> ). Allowed values for <code>maxSE\$method</code> include: <ul style="list-style-type: none"> <li>• "globalmax": simply corresponds to the global maximum, i.e., <code>is which.max(gap)</code></li> <li>• "firstmax": gives the location of the first local maximum</li> <li>• "Tibs2001SEmax": uses the criterion, Tibshirani et al (2001) proposed: "the smallest k such that <math>\text{gap}(k) \geq \text{gap}(k+1) - s_{k+1}</math>". It's also possible to use "the smallest k such that <math>\text{gap}(k) \geq \text{gap}(k+1) - \text{SE.factor} * s_{k+1}</math>" where SE.factor is a numeric value which can be 1 (default), 2, 3, etc.</li> <li>• "firstSEmax": location of the first <code>f()</code> value which is not larger than the first local maximum minus <code>SE.factor * SE.f[]</code>, i.e, within an "f S.E." range of that maximum.</li> <li>• see <code>?cluster::maxSE</code> for more options</li> </ul>

**Value**

- `fviz_nbclust`, `fviz_gap_stat`: return a `ggplot2`

**Author(s)**

Alboukadel Kassambara <[alboukadel.kassambara@gmail.com](mailto:alboukadel.kassambara@gmail.com)>

**See Also**

[fviz\\_cluster](#), [eclust](#)

**Examples**

```
set.seed(123)

# Data preparation
```

```

# ++++++
data("iris")
head(iris)
# Remove species column (5) and scale the data
iris.scaled <- scale(iris[, -5])

# Optimal number of clusters in the data
# ++++++
# Examples are provided only for kmeans, but
# you can also use cluster::pam (for pam) or
# hcut (for hierarchical clustering)

### Elbow method (look at the knee)
# Elbow method for kmeans
fviz_nbclust(iris.scaled, kmeans, method = "wss") +
geom_vline(xintercept = 3, linetype = 2)

# Average silhouette for kmeans
fviz_nbclust(iris.scaled, kmeans, method = "silhouette")

### Gap statistic
library(cluster)
set.seed(123)
# Compute gap statistic for kmeans
# we used B = 10 for demo. Recommended value is ~500
gap_stat <- clusGap(iris.scaled, FUN = kmeans, nstart = 25,
  K.max = 10, B = 10)
print(gap_stat, method = "firstmax")
fviz_gap_stat(gap_stat)

# Gap statistic for hierarchical clustering
gap_stat <- clusGap(iris.scaled, FUN = hcut, K.max = 10, B = 10)
fviz_gap_stat(gap_stat)

```

---

fviz\_pca

*Visualize Principal Component Analysis*


---

## Description

Principal component analysis (PCA) reduces the dimensionality of multivariate data, to two or three that can be visualized graphically with minimal loss of information. `fviz_pca()` provides ggplot2-based elegant visualization of PCA outputs from: i) `prcomp` and `princomp` [in built-in R stats], ii) PCA [in `FactoMineR`], iii) `dudi.pca` [in `ade4`] and `epPCA` [ExPosition]. Read more: [Principal Component Analysis](#)

- `fviz_pca_ind()`: Graph of individuals
- `fviz_pca_var()`: Graph of variables

- `fviz_pca_biplot()`: Biplot of individuals and variables
- `fviz_pca()`: An alias of `fviz_pca_biplot()`

Note that, `fviz_pca_xxx()` functions are wrapper around the core function `fviz()`, which is also a wrapper around the function `ggscatter()` [in `ggpubr`]. Therefore, further arguments, to be passed to the function `fviz()` and `ggscatter()`, can be specified in `fviz_pca_ind()` and `fviz_pca_var()`.

## Usage

```
fviz_pca(X, ...)
```

```
fviz_pca_ind(
  X,
  axes = c(1, 2),
  geom = c("point", "text"),
  geom.ind = geom,
  repel = FALSE,
  habillage = "none",
  palette = NULL,
  addEllipses = FALSE,
  col.ind = "black",
  fill.ind = "white",
  col.ind.sup = "blue",
  alpha.ind = 1,
  select.ind = list(name = NULL, cos2 = NULL, contrib = NULL),
  ...
)
```

```
fviz_pca_var(
  X,
  axes = c(1, 2),
  geom = c("arrow", "text"),
  geom.var = geom,
  repel = FALSE,
  col.var = "black",
  fill.var = "white",
  alpha.var = 1,
  col.quanti.sup = "blue",
  col.circle = "grey70",
  select.var = list(name = NULL, cos2 = NULL, contrib = NULL),
  ...
)
```

```
fviz_pca_biplot(
  X,
  axes = c(1, 2),
  geom = c("point", "text"),
  geom.ind = geom,
```

```

geom.var = c("arrow", "text"),
col.ind = "black",
fill.ind = "white",
col.var = "steelblue",
fill.var = "white",
gradient.cols = NULL,
label = "all",
invisible = "none",
repel = FALSE,
habillage = "none",
palette = NULL,
addEllipses = FALSE,
title = "PCA - Biplot",
...
)

```

## Arguments

<code>X</code>	an object of class PCA [FactoMineR]; pcomp and princomp [stats]; dudi and pca [ade4]; expOutput/epPCA [ExPosition].
<code>...</code>	Additional arguments. <ul style="list-style-type: none"> <li>• in <code>fviz_pca_ind()</code> and <code>fviz_pca_var()</code>: Additional arguments are passed to the functions <code>fviz()</code> and <code>ggpubr::ggpar()</code>.</li> <li>• in <code>fviz_pca_biplot()</code> and <code>fviz_pca()</code>: Additional arguments are passed to <code>fviz_pca_ind()</code> and <code>fviz_pca_var()</code>.</li> </ul>
<code>axes</code>	a numeric vector of length 2 specifying the dimensions to be plotted.
<code>geom</code>	a text specifying the geometry to be used for the graph. Allowed values are the combination of <code>c("point", "arrow", "text")</code> . Use "point" (to show only points); "text" to show only labels; <code>c("point", "text")</code> or <code>c("arrow", "text")</code> to show arrows and texts. Using <code>c("arrow", "text")</code> is sensible only for the graph of variables.
<code>geom.ind, geom.var</code>	as <code>geom</code> but for individuals and variables, respectively. Default is <code>geom.ind = c("point", "text)</code> , <code>geom.var = c("arrow", "text")</code> .
<code>repel</code>	a boolean, whether to use <code>ggrepel</code> to avoid overplotting text labels or not.
<code>habillage</code>	an optional factor variable for coloring the observations by groups. Default value is "none". If <code>X</code> is a PCA object from FactoMineR package, <code>habillage</code> can also specify the supplementary qualitative variable (by its index or name) to be used for coloring individuals by groups (see ?PCA in FactoMineR).
<code>palette</code>	the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. <code>c("blue", "red")</code> ; and scientific journal palettes from ggsci R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty". Can be also a numeric vector of length(groups); in this case a basic color palette is created using the function <a href="#">palette</a> .
<code>addEllipses</code>	logical value. If TRUE, draws ellipses around the individuals when <code>habillage != "none"</code> .

<code>col.ind, col.var</code>	color for individuals and variables, respectively. Can be a continuous variable or a factor variable. Possible values include also : "cos2", "contrib", "coord", "x" or "y". In this case, the colors for individuals/variables are automatically controlled by their qualities of representation ("cos2"), contributions ("contrib"), coordinates ( $x^2+y^2$ , "coord"), x values ("x") or y values ("y"). To use automatic coloring (by cos2, contrib, ....), make sure that <code>habillage = "none"</code> .
<code>fill.ind, fill.var</code>	same as <code>col.ind</code> and <code>col.var</code> but for the fill color.
<code>col.ind.sup</code>	color for supplementary individuals
<code>alpha.ind, alpha.var</code>	controls the transparency of individual and variable colors, respectively. The value can variate from 0 (total transparency) to 1 (no transparency). Default value is 1. Possible values include also : "cos2", "contrib", "coord", "x" or "y". In this case, the transparency for the individual/variable colors are automatically controlled by their qualities ("cos2"), contributions ("contrib"), coordinates ( $x^2+y^2$ , "coord"), x values("x") or y values("y"). To use this, make sure that <code>habillage = "none"</code> .
<code>select.ind, select.var</code>	a selection of individuals/variables to be drawn. Allowed values are NULL or a list containing the arguments name, cos2 or contrib: <ul style="list-style-type: none"> <li>• name: is a character vector containing individuals/variables to be drawn</li> <li>• cos2: if cos2 is in [0, 1], ex: 0.6, then individuals/variables with a cos2 &gt; 0.6 are drawn. if cos2 &gt; 1, ex: 5, then the top 5 individuals/variables with the highest cos2 are drawn.</li> <li>• contrib: if contrib &gt; 1, ex: 5, then the top 5 individuals/variables with the highest contrib are drawn</li> </ul>
<code>col.quanti.sup</code>	a color for the quantitative supplementary variables.
<code>col.circle</code>	a color for the correlation circle. Used only when X is a PCA output.
<code>gradient.cols</code>	vector of colors to use for n-colour gradient. Allowed values include brewer and ggsci color palettes.
<code>label</code>	a text specifying the elements to be labelled. Default value is "all". Allowed values are "none" or the combination of c("ind", "ind.sup", "quali", "var", "quanti.sup"). "ind" can be used to label only active individuals. "ind.sup" is for supplementary individuals. "quali" is for supplementary qualitative variables. "var" is for active variables. "quanti.sup" is for quantitative supplementary variables.
<code>invisible</code>	a text specifying the elements to be hidden on the plot. Default value is "none". Allowed values are the combination of c("ind", "ind.sup", "quali", "var", "quanti.sup").
<code>title</code>	the title of the graph

**Value**

a ggplot

**Author(s)**

Alboukadel Kassambara <alboukadel.kassambara@gmail.com>

**See Also**

[fviz\\_ca](#), [fviz\\_mca](#)

**Examples**

```
# Principal component analysis
# ++++++
data(iris)
res.pca <- prcomp(iris[, -5], scale = TRUE)

# Graph of individuals
# ++++++

# Default plot
# Use repel = TRUE to avoid overplotting (slow if many points)
fviz_pca_ind(res.pca, col.ind = "#00AFBB",
             repel = TRUE)

# 1. Control automatically the color of individuals
#   # using the "cos2" or the contributions "contrib"
#   # cos2 = the quality of the individuals on the factor map
# 2. To keep only point or text use geom = "point" or geom = "text".
# 3. Change themes using ggtheme: http://www.sthda.com/english/wiki/ggplot2-themes

fviz_pca_ind(res.pca, col.ind="cos2", geom = "point",
             gradient.cols = c("white", "#2E9FDF", "#FC4E07" ))

# Color individuals by groups, add concentration ellipses
# Change group colors using RColorBrewer color palettes
# Read more: http://www.sthda.com/english/wiki/ggplot2-colors
# Remove labels: label = "none".
fviz_pca_ind(res.pca, label="none", habillage=iris$Species,
             addEllipses=TRUE, ellipse.level=0.95, palette = "Dark2")

# Change group colors manually
# Read more: http://www.sthda.com/english/wiki/ggplot2-colors
fviz_pca_ind(res.pca, label="none", habillage=iris$Species,
             addEllipses=TRUE, ellipse.level=0.95,
             palette = c("#999999", "#E69F00", "#56B4E9"))

# Select and visualize some individuals (ind) with select.ind argument.
# - ind with cos2 >= 0.96: select.ind = list(cos2 = 0.96)
# - Top 20 ind according to the cos2: select.ind = list(cos2 = 20)
# - Top 20 contributing individuals: select.ind = list(contrib = 20)
# - Select ind by names: select.ind = list(name = c("23", "42", "119" ) )

# Example: Select the top 40 according to the cos2
fviz_pca_ind(res.pca, select.ind = list(cos2 = 40))
```

```

# Graph of variables
# ++++++

# Default plot
fviz_pca_var(res.pca, col.var = "steelblue")

# Control variable colors using their contributions
fviz_pca_var(res.pca, col.var = "contrib",
  gradient.cols = c("white", "blue", "red"),
  ggtheme = theme_minimal())

# Biplot of individuals and variables
# ++++++
# Keep only the labels for variables
# Change the color by groups, add ellipses
fviz_pca_biplot(res.pca, label = "var", habillage=iris$Species,
  addEllipses=TRUE, ellipse.level=0.95,
  ggtheme = theme_minimal())

```

---

fviz\_silhouette

*Visualize Silhouette Information from Clustering*


---

### Description

Silhouette (Si) analysis is a cluster validation approach that measures how well an observation is clustered and it estimates the average distance between clusters. `fviz_silhouette()` provides ggplot2-based elegant visualization of silhouette information from i) the result of `silhouette()`, `pam()`, `clara()` and `fanny()` [in cluster package]; ii) `eclust()` and `hcut()` [in factoextra].

Read more: [Clustering Validation Statistics](#).

### Usage

```
fviz_silhouette(sil.obj, label = FALSE, print.summary = TRUE, ...)
```

### Arguments

<code>sil.obj</code>	an object of class silhouette: pam, clara, fanny [in cluster package]; eclust and hcut [in factoextra].
<code>label</code>	logical value. If true, x axis tick labels are shown
<code>print.summary</code>	logical value. If true a summary of cluster silhouettes are printed in <code>fviz_silhouette()</code> .
<code>...</code>	other arguments to be passed to the function <code>ggpubr::ggpar()</code> .

**Details**

- Observations with a large silhouette Si (almost 1) are very well clustered.
- A small Si (around 0) means that the observation lies between two clusters.
- Observations with a negative Si are probably placed in the wrong cluster.

**Value**

return a ggplot

**Author(s)**

Alboukadel Kassambara <alboukadel.kassambara@gmail.com>

**See Also**

[fviz\\_cluster](#), [hcut](#), [hkmeans](#), [eclust](#), [fviz\\_dend](#)

**Examples**

```
set.seed(123)

# Data preparation
# ++++++
data("iris")
head(iris)
# Remove species column (5) and scale the data
iris.scaled <- scale(iris[, -5])

# K-means clustering
# ++++++
km.res <- kmeans(iris.scaled, 3, nstart = 2)

# Visualize kmeans clustering
fviz_cluster(km.res, iris[, -5], ellipse.type = "norm")+
theme_minimal()

# Visualize silhouette information
require("cluster")
sil <- silhouette(km.res$cluster, dist(iris.scaled))
fviz_silhouette(sil)

# Identify observation with negative silhouette
neg_sil_index <- which(sil[, "sil_width"] < 0)
sil[neg_sil_index, , drop = FALSE]
## Not run:
# PAM clustering
# ++++++
require(cluster)
pam.res <- pam(iris.scaled, 3)
# Visualize pam clustering
fviz_cluster(pam.res, ellipse.type = "norm")+
```



```

theme_minimal()
# Visualize silhouette information
fviz_silhouette(pam.res)

# Hierarchical clustering
# ++++++
# Use hcut() which compute hclust and cut the tree
hc.cut <- hcut(iris.scaled, k = 3, hc_method = "complete")
# Visualize dendrogram
fviz_dend(hc.cut, show_labels = FALSE, rect = TRUE)
# Visualize silhouette information
fviz_silhouette(hc.cut)

## End(Not run)

```

---

get\_ca

*Extract the results for rows/columns - CA*


---

### Description

Extract all the results (coordinates, squared cosine, contributions and inertia) for the active row/column variables from Correspondence Analysis (CA) outputs.

- `get_ca()`: Extract the results for rows and columns
- `get_ca_row()`: Extract the results for rows only
- `get_ca_col()`: Extract the results for columns only

### Usage

```
get_ca(res.ca, element = c("row", "col"))
```

```
get_ca_col(res.ca)
```

```
get_ca_row(res.ca)
```

### Arguments

<code>res.ca</code>	an object of class CA [FactoMineR], ca [ca], coa [ade4]; correspondence [MASS].
<code>element</code>	the element to subset from the output. Possible values are "row" or "col".

### Value

a list of matrices containing the results for the active rows/columns including :

<code>coord</code>	coordinates for the rows/columns
<code>cos2</code>	cos2 for the rows/columns
<code>contrib</code>	contributions of the rows/columns
<code>inertia</code>	inertia of the rows/columns

**Author(s)**

Alboukadel Kassambara <alboukadel.kassambara@gmail.com>

**References**

<http://www.sthda.com>

**Examples**

```
# Install and load FactoMineR to compute CA
# install.packages("FactoMineR")
library("FactoMineR")
data("housetasks")
res.ca <- CA(housetasks, graph = FALSE)

# Result for column variables
col <- get_ca_col(res.ca)
col # print
head(col$coord) # column coordinates
head(col$cos2) # column cos2
head(col$contrib) # column contributions

# Result for row variables
row <- get_ca_row(res.ca)
row # print
head(row$coord) # row coordinates
head(row$cos2) # row cos2
head(row$contrib) # row contributions

# You can also use the function get_ca()
get_ca(res.ca, "row") # Results for rows
get_ca(res.ca, "col") # Results for columns
```

---

get\_clust\_tendency      *Assessing Clustering Tendency*

---

**Description**

Before applying cluster methods, the first step is to assess whether the data is clusterable, a process defined as the **assessing of clustering tendency**. `get_clust_tendency()` assesses clustering tendency using Hopkins' statistic and a visual approach. An ordered dissimilarity image (ODI) is shown. Objects belonging to the same cluster are displayed in consecutive order using hierarchical clustering. For more details and interpretation, see [STHDA website: Assessing clustering tendency](#).

**Usage**

```
get_clust_tendency(  
  data,  
  n,  
  graph = TRUE,  
  gradient = list(low = "red", mid = "white", high = "blue"),  
  seed = 123  
)
```

**Arguments**

data	a numeric data frame or matrix. Columns are variables and rows are samples. Computation are done on rows (samples) by default. If you want to calculate Hopkins statistic on variables, transpose the data before.
n	the number of points selected from sample space which is also the number of points selected from the given sample(data).
graph	logical value; if TRUE the ordered dissimilarity image (ODI) is shown.
gradient	a list containing three elements specifying the colors for low, mid and high values in the ordered dissimilarity image. The element "mid" can take the value of NULL.
seed	an integer specifying the seed for random number generator. Specify seed for reproducible results.

**Details**

**Hopkins statistic:** If the value of Hopkins statistic is close to 1 (far above 0.5), then we can conclude that the dataset is significantly clusterable.

**VAT (Visual Assessment of cluster Tendency):** The VAT detects the clustering tendency in a visual form by counting the number of square shaped dark (or colored) blocks along the diagonal in a VAT image.

**Value**

A list containing the elements:

- hopkins\_stat for Hopkins statistic value
- plot for ordered dissimilarity image. This is generated using the function [fviz\\_dist](#)(dist.obj).

**Author(s)**

Alboukadel Kassambara <[alboukadel.kassambara@gmail.com](mailto:alboukadel.kassambara@gmail.com)>

**See Also**

[fviz\\_dist](#)

**Examples**

```

data(iris)

# Clustering tendency
gradient_col = list(low = "steelblue", high = "white")
get_clust_tendency(iris[,-5], n = 50, gradient = gradient_col)

# Random uniformly distributed dataset
# (without any inherent clusters)
set.seed(123)
random_df <- apply(iris[, -5], 2,
                  function(x){runif(length(x), min(x), max(x))}
                  )
get_clust_tendency(random_df, n = 50, gradient = gradient_col)

```

---

`get_famd`*Extract the results for individuals and variables - FAMD*

---

**Description**

Extract all the results (coordinates, squared cosine and contributions) for the active individuals and variables from Factor Analysis of Mixed Date (FAMD) outputs.

- `get_famd()`: Extract the results for variables and individuals
- `get_famd_ind()`: Extract the results for individuals only
- `get_famd_var()`: Extract the results for quantitative and qualitative variables only

**Usage**

```

get_famd(res.famd, element = c("ind", "var", "quanti.var", "quali.var"))

get_famd_ind(res.famd)

get_famd_var(res.famd, element = c("var", "quanti.var", "quali.var"))

```

**Arguments**

`res.famd` an object of class FAMD [FactoMineR].

`element` the element to subset from the output. Possible values are "ind", "quanti.var" or "quali.var".

**Value**

a list of matrices containing the results for the active individuals and variables, including :

coord	coordinates of individuals/variables.
cos2	cos2 values representing the quality of representation on the factor map.
contrib	contributions of individuals / variables to the principal components.

**Author(s)**

Alboukadel Kassambara <alboukadel.kassambara@gmail.com>

**Examples**

```
# Compute FAMD
library("FactoMineR")
data(wine)
res.famd <- FAMD(wine[,c(1,2, 16, 22, 29, 28, 30,31)], graph = FALSE)

# Extract the results for qualitative variable categories
quali.var <- get_famd_var(res.famd, "quali.var")
print(quali.var)
head(quali.var$coord) # coordinates of qualitative variables

# Extract the results for quantitative variables
quanti.var <- get_famd_var(res.famd, "quanti.var")
print(quanti.var)
head(quanti.var$coord) # coordinates

# Extract the results for individuals
ind <- get_famd_ind(res.famd)
print(ind)
head(ind$coord) # coordinates of individuals
```

---

get_hmfa	<i>Extract the results for individuals/variables/group/partial axes - HMFA</i>
----------	--

---

**Description**

Extract all the results (coordinates, squared cosine and contributions) for the active individuals/quantitative variables/qualitative variable categories/groups/partial axes from Hierarchical Multiple Factor Analysis (HMFA) outputs.

- get\_hmfa(): Extract the results for variables and individuals

- `get_hmfa_ind()`: Extract the results for individuals only
- `get_hmfa_var()`: Extract the results for variables (quantitatives, qualitatives and groups)
- `get_hmfa_partial()`: Extract the results for partial.node.

### Usage

```
get_hmfa(
  res.hmfa,
  element = c("ind", "quanti.var", "quali.var", "group", "partial.node")
)
```

```
get_hmfa_ind(res.hmfa)
```

```
get_hmfa_var(res.hmfa, element = c("quanti.var", "quali.var", "group"))
```

```
get_hmfa_partial(res.hmfa)
```

### Arguments

<code>res.hmfa</code>	an object of class HMFA [FactoMineR].
<code>element</code>	the element to subset from the output. Possible values are "ind", "quanti.var", "quali.var", "group" or "partial.node".

### Value

a list of matrices containing the results for the active individuals, variables, groups and partial nodes, including :

<code>coord</code>	coordinates
<code>cos2</code>	cos2
<code>contrib</code>	contributions

### Author(s)

Alboukadel Kassambara <alboukadel.kassambara@gmail.com>  
 Fabian Mundt <f.mundt@inventionate.de>

### Examples

```
# Multiple Factor Analysis
# ++++++
# Install and load FactoMineR to compute MFA
# install.packages("FactoMineR")
library("FactoMineR")
data(wine)
hierar <- list(c(2,5,3,10,9,2), c(4,2))
res.hmfa <- HMFA(wine, H = hierar, type=c("n",rep("s",5)), graph = FALSE)

# Extract the results for qualitative variable categories
```

```

var <- get_hmfa_var(res.hmfa, "quali.var")
print(var)
head(var$coord) # coordinates of qualitative variables
head(var$cos2) # cos2 of qualitative variables
head(var$contrib) # contributions of qualitative variables

# Extract the results for individuals
ind <- get_hmfa_ind(res.hmfa)
print(ind)
head(ind$coord) # coordinates of individuals
head(ind$cos2) # cos2 of individuals
head(ind$contrib) # contributions of individuals

# You can also use the function get_hmfa()
get_hmfa(res.hmfa, "ind") # Results for individuals
get_hmfa(res.hmfa, "quali.var") # Results for qualitative variable categories

```

---

get\_mca

---

*Extract the results for individuals/variables - MCA*


---

## Description

Extract all the results (coordinates, squared cosine and contributions) for the active individuals/variable categories from Multiple Correspondence Analysis (MCA) outputs.

- `get_mca()`: Extract the results for variables and individuals
- `get_mca_ind()`: Extract the results for individuals only
- `get_mca_var()`: Extract the results for variables only

## Usage

```
get_mca(res.mca, element = c("var", "ind", "mca.cor", "quanti.sup"))
```

```
get_mca_var(res.mca, element = c("var", "mca.cor", "quanti.sup"))
```

```
get_mca_ind(res.mca)
```

## Arguments

<code>res.mca</code>	an object of class MCA [FactoMineR], acm [ade4], expoOutput/epMCA [Ex-Position].
<code>element</code>	the element to subset from the output. Possible values are "var" for variables, "ind" for individuals, "mca.cor" for correlation between variables and principal dimensions, "quanti.sup" for quantitative supplementary variables.

**Value**

a list of matrices containing the results for the active individuals/variable categories including :

coord	coordinates for the individuals/variable categories
cos2	cos2 for the individuals/variable categories
contrib	contributions of the individuals/variable categories
inertia	inertia of the individuals/variable categories

**Author(s)**

Alboukadel Kassambara <alboukadel.kassambara@gmail.com>

**References**

<http://www.sthda.com/english/>

**Examples**

```
# Multiple Correspondence Analysis
# ++++++
# Install and load FactoMineR to compute MCA
# install.packages("FactoMineR")
library("FactoMineR")
data(poison)
poison.active <- poison[1:55, 5:15]
head(poison.active[, 1:6])
res.mca <- MCA(poison.active, graph=FALSE)

# Extract the results for variable categories
var <- get_mca_var(res.mca)
print(var)
head(var$coord) # coordinates of variables
head(var$cos2) # cos2 of variables
head(var$contrib) # contributions of variables

# Extract the results for individuals
ind <- get_mca_ind(res.mca)
print(ind)
head(ind$coord) # coordinates of individuals
head(ind$cos2) # cos2 of individuals
head(ind$contrib) # contributions of individuals

# You can also use the function get_mca()
get_mca(res.mca, "ind") # Results for individuals
get_mca(res.mca, "var") # Results for variable categories
```



---

get\_mfa

*Extract the results for individuals/variables/group/partial axes - MFA*


---

### Description

Extract all the results (coordinates, squared cosine and contributions) for the active individuals/quantitative variables/qualitative variable categories/groups/partial axes from Multiple Factor Analysis (MFA) outputs.

- `get_mfa()`: Extract the results for variables and individuals
- `get_mfa_ind()`: Extract the results for individuals only
- `get_mfa_var()`: Extract the results for variables (quantitatives, qualitatives and groups)
- `get_mfa_partial_axes()`: Extract the results for partial axes only

### Usage

```
get_mfa(
  res.mfa,
  element = c("ind", "quanti.var", "quali.var", "group", "partial.axes")
)
```

```
get_mfa_ind(res.mfa)
```

```
get_mfa_var(res.mfa, element = c("quanti.var", "quali.var", "group"))
```

```
get_mfa_partial_axes(res.mfa)
```

### Arguments

<code>res.mfa</code>	an object of class MFA [FactoMineR].
<code>element</code>	the element to subset from the output. Possible values are "ind", "quanti.var", "quali.var", "group" or "partial.axes".

### Value

a list of matrices containing the results for the active individuals/quantitative variable categories/qualitative variable categories/groups/partial axes including :

<code>coord</code>	coordinates for the individuals/quantitative variable categories/qualitative variable categories/groups/partial axes
<code>cos2</code>	cos2 for the individuals/quantitative variable categories/qualitative variable categories/groups/partial axes
<code>contrib</code>	contributions of the individuals/quantitative variable categories/qualitative variable categories/groups/partial axes
<code>inertia</code>	inertia of the individuals/quantitative variable categories/qualitative variable categories/groups/partial axes

**Author(s)**

Alboukadel Kassambara <alboukadel.kassambara@gmail.com>

Fabian Mundt <f.mundt@inventionate.de>

**Examples**

```
# Multiple Factor Analysis
# ++++++
# Install and load FactoMineR to compute MFA
# install.packages("FactoMineR")
library("FactoMineR")
data(poison)
res.mfa <- MFA(poison, group=c(2,2,5,6), type=c("s","n","n","n"),
name.group=c("desc","desc2","symptom","eat"), num.group.sup=1:2,
graph = FALSE)

# Extract the results for qualitative variable categories
var <- get_mfa_var(res.mfa, "quali.var")
print(var)
head(var$coord) # coordinates of qualitative variables
head(var$cos2) # cos2 of qualitative variables
head(var$contrib) # contributions of qualitative variables

# Extract the results for individuals
ind <- get_mfa_ind(res.mfa)
print(ind)
head(ind$coord) # coordinates of individuals
head(ind$cos2) # cos2 of individuals
head(ind$contrib) # contributions of individuals

# You can also use the function get_mfa()
get_mfa(res.mfa, "ind") # Results for individuals
get_mfa(res.mfa, "quali.var") # Results for qualitative variable categories
```

---

get\_pca

*Extract the results for individuals/variables - PCA*

---

**Description**

Extract all the results (coordinates, squared cosine, contributions) for the active individuals/variables from Principal Component Analysis (PCA) outputs.

- `get_pca()`: Extract the results for variables and individuals
- `get_pca_ind()`: Extract the results for individuals only
- `get_pca_var()`: Extract the results for variables only

**Usage**

```
get_pca(res.pca, element = c("var", "ind"))

get_pca_ind(res.pca, ...)

get_pca_var(res.pca)
```

**Arguments**

res.pca	an object of class PCA [FactoMineR]; prcomp and princomp [stats]; pca, dudi [ade4]; epPCA [ExPosition].
element	the element to subset from the output. Allowed values are "var" (for active variables) or "ind" (for active individuals).
...	not used

**Value**

a list of matrices containing all the results for the active individuals/variables including:

coord	coordinates for the individuals/variables
cos2	cos2 for the individuals/variables
contrib	contributions of the individuals/variables

**Author(s)**

Alboukadel Kassambara <alboukadel.kassambara@gmail.com>

**References**

<http://www.sthda.com/english/>

**Examples**

```
# Principal Component Analysis
# ++++++
data(iris)
res.pca <- prcomp(iris[, -5], scale = TRUE)
# Extract the results for individuals
ind <- get_pca_ind(res.pca)
print(ind)
head(ind$coord) # coordinates of individuals
head(ind$cos2) # cos2 of individuals
head(ind$contrib) # contributions of individuals

# Extract the results for variables
var <- get_pca_var(res.pca)
print(var)
head(var$coord) # coordinates of variables
head(var$cos2) # cos2 of variables
head(var$contrib) # contributions of variables
```

```
# You can also use the function get_pca()
get_pca(res.pca, "ind") # Results for individuals
get_pca(res.pca, "var") # Results for variable categories
```

---

hcut

*Computes Hierarchical Clustering and Cut the Tree*


---

### Description

Computes hierarchical clustering (hclust, agnes, diana) and cut the tree into k clusters. It also accepts correlation based distance measure methods such as "pearson", "spearman" and "kendall".

### Usage

```
hcut(
  x,
  k = 2,
  isdiss = inherits(x, "dist"),
  hc_func = c("hclust", "agnes", "diana"),
  hc_method = "ward.D2",
  hc_metric = "euclidean",
  stand = FALSE,
  graph = FALSE,
  ...
)
```

### Arguments

x	a numeric matrix, numeric data frame or a dissimilarity matrix.
k	the number of clusters to be generated.
isdiss	logical value specifying whether x is a dissimilarity matrix.
hc_func	the hierarchical clustering function to be used. Default value is "hclust". Possible values is one of "hclust", "agnes", "diana". Abbreviation is allowed.
hc_method	the agglomeration method to be used (?hclust) for hclust() and agnes(): "ward.D", "ward.D2", "single", "complete", "average", ...
hc_metric	character string specifying the metric to be used for calculating dissimilarities between observations. Allowed values are those accepted by the function dist() [including "euclidean", "manhattan", "maximum", "canberra", "binary", "minkowski"] and correlation based distance measures ["pearson", "spearman" or "kendall"].
stand	logical value; default is FALSE. If TRUE, then the data will be standardized using the function scale(). Measurements are standardized for each variable (column), by subtracting the variable's mean value and dividing by the variable's standard deviation.

graph	logical value. If TRUE, the dendrogram is displayed.
...	not used.

### Value

an object of class "hcut" containing the result of the standard function used (read the documentation of `hclust`, `agnes`, `diana`).

It includes also:

- `cluster`: the cluster assignement of observations after cutting the tree
- `nbclust`: the number of clusters
- `silinfo`: the silhouette information of observations (if  $k > 1$ )
- `size`: the size of clusters
- `data`: a matrix containing the original or the standardized data (if `stand = TRUE`)

### See Also

[fviz\\_dend](#), [hkmeans](#), [eclust](#)

### Examples

```
data(USArrests)

# Compute hierarchical clustering and cut into 4 clusters
res <- hcut(USArrests, k = 4, stand = TRUE)

# Cluster assignements of observations
res$cluster
# Size of clusters
res$size

# Visualize the dendrogram
fviz_dend(res, rect = TRUE)

# Visualize the silhouette
fviz_silhouette(res)

# Visualize clusters as scatter plots
fviz_cluster(res)
```

---

hkmeans

*Hierarchical k-means clustering*


---

## Description

The final k-means clustering solution is very sensitive to the initial random selection of cluster centers. This function provides a solution using an hybrid approach by combining the hierarchical clustering and the k-means methods. The procedure is explained in "Details" section. Read more: [Hybrid hierarchical k-means clustering for optimizing clustering outputs](#).

- `hkmeans()`: compute hierarchical k-means clustering
- `print.hkmeans()`: prints the result of `hkmeans`
- `hkmeans_tree()`: plots the initial dendrogram

## Usage

```
hkmeans(
  x,
  k,
  hc.metric = "euclidean",
  hc.method = "ward.D2",
  iter.max = 10,
  km.algorithm = "Hartigan-Wong"
)

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

hkmeans_tree(hkmeans, rect.col = NULL, ...)
```

## Arguments

<code>x</code>	a numeric matrix, data frame or vector
<code>k</code>	the number of clusters to be generated
<code>hc.metric</code>	the distance measure to be used. Possible values are "euclidean", "maximum", "manhattan", "canberra", "binary" or "minkowski" (see <code>?dist</code> ).
<code>hc.method</code>	the agglomeration method to be used. Possible values include "ward.D", "ward.D2", "single", "complete", "average", "mcquitty", "median" or "centroid" (see <code>?hclust</code> ).
<code>iter.max</code>	the maximum number of iterations allowed for k-means.
<code>km.algorithm</code>	the algorithm to be used for kmeans (see <code>?kmeans</code> ).
<code>...</code>	others arguments to be passed to the function <code>plot.hclust()</code> ; (see <code>?plot.hclust</code> )
<code>hkmeans</code>	an object of class <code>hkmeans</code> (returned by the function <code>hkmeans()</code> )
<code>rect.col</code>	Vector with border colors for the rectangles around clusters in dendrogram

## Details

The procedure is as follow:

1. Compute hierarchical clustering
2. Cut the tree in k-clusters
3. compute the center (i.e the mean) of each cluster
4. Do k-means by using the set of cluster centers (defined in step 3) as the initial cluster centers. Optimize the clustering.

This means that the final optimized partitioning obtained at step 4 might be different from the initial partitioning obtained at step 2. Consider mainly the result displayed by `fviz_cluster()`.

## Value

hkmeans returns an object of class "hkmeans" containing the following components:

- The elements returned by the standard function `kmeans()` (see `?kmeans`)
- `data`: the data used for the analysis
- `hclust`: an object of class "hclust" generated by the function `hclust()`

## Examples

```
# Load data
data(USArrests)
# Scale the data
df <- scale(USArrests)

# Compute hierarchical k-means clustering
res.hk <-hkmeans(df, 4)

# Elements returned by hkmeans()
names(res.hk)

# Print the results
res.hk

# Visualize the tree
hkmeans_tree(res.hk, cex = 0.6)
# or use this
fviz_dend(res.hk, cex = 0.6)

# Visualize the hkmeans final clusters
fviz_cluster(res.hk, frame.type = "norm", frame.level = 0.68)
```

---

housetasks	<i>House tasks contingency table</i>
------------	--------------------------------------

---

**Description**

A data frame containing the frequency of execution of 13 house tasks in the couple. This table is also available in ade4 package.

**Usage**

```
data("housetasks")
```

**Format**

A data frame with 13 observations (house tasks) on the following 4 columns.

Wife a numeric vector

Alternating a numeric vector

Husband a numeric vector

Jointly a numeric vector

**Source**

This data is from FactoMineR package.

**Examples**

```
library(FactoMineR)
data(housetasks)
res.ca <- CA(housetasks, graph=FALSE)
fviz_ca_biplot(res.ca, repel = TRUE)+
  theme_minimal()
```

---

multishapes	<i>A dataset containing clusters of multiple shapes</i>
-------------	---

---

**Description**

Data containing clusters of any shapes. Useful for comparing density-based clustering (DBSCAN) and standard partitioning methods such as k-means clustering.

**Usage**

```
data("multishapes")
```



**Format**

A data frame with 1100 observations on the following 3 variables.

x a numeric vector containing the x coordinates of observations

y a numeric vector containing the y coordinates of observations

shape a numeric vector corresponding to the cluster number of each observations.

**Details**

The dataset contains 5 clusters and some outliers/noises.

**Examples**

```
data(multishapes)
plot(multishapes[,1], multishapes[, 2],
      col = multishapes[, 3], pch = 19, cex = 0.8)
```

---

poison

*Poison*

---

**Description**

This data is a result from a survey carried out on children of primary school who suffered from food poisoning. They were asked about their symptoms and about what they ate.

**Usage**

```
data("poison")
```

**Format**

A data frame with 55 rows and 15 columns.

**Source**

This data is from FactoMineR package.

**Examples**

```
library(FactoMineR)
data(poison)
res.mca <- MCA(poison, quanti.sup = 1:2, quali.sup = c(3,4),
               graph = FALSE)
fviz_mca_biplot(res.mca, repel = TRUE)+
  theme_minimal()
```

---

print.factoextra      *Print method for an object of class factoextra*

---

**Description**

Print method for an object of class factoextra

**Usage**

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

**Arguments**

x                    an object of class factoextra  
...                  further arguments to be passed to print method

**Author(s)**

Alboukadel Kassambara <alboukadel.kassambara@gmail.com>

**Examples**

```
data(iris)  
res.pca <- prcomp(iris[, -5], scale = TRUE)  
ind <- get_pca_ind(res.pca, data = iris[, -5])  
print(ind)
```

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